There is disclosed an aqueous dispersion comprising:

a) inorganic particles;

b) at least one fatty acid or a salt thereof,

c) a polymeric binder, and

d) water, wherein said aqueous dispersion is free of organic solvent. There is also disclosed use of an aqueous dispersion comprising:

a) inorganic particles;

b) at least one fatty acid or a salt thereof;

c) a polymeric binder, and
d) water, as a coating on a substrate surface, wherein said surface after coating displays an equilibrium contact angle higher than 120° degrees, preferably more than 135°, most preferably more than 150° for a drop of water on the surface. There is also disclosed a method for coating a substrate comprising contacting said substrate with the aqueous dispersion. The coating allows application in one step, it is non-toxic, safe for food packaging, and environmentally friendly as well as inexpensive.
An aqueous dispersion, a coated subject and use of an aqueous dispersion

Technical field of the invention

The present invention relates to the field highly hydrophobic and superhydrophobic coatings.

Background of the invention

Hydrophobic, highly hydrophobic and superhydrophobic coatings are used for many kinds of surfaces. In the art many approaches have been used to manufacture highly hydrophobic coatings.

US 3,940,385 discloses glossy coating compositions where a pigment for instance with a particle size 0.2-0.5 µm is coated with an organic compound to render the pigment hydrophobic. The pigment is emulsified and renders the coating hydrophobic. TiO₂ is mentioned as a pigment. Fatty acids are mentioned as compounds to be used to coat the pigment.

DE 1033561 shows the use of both hydrophilic and hydrophobic substances to coat a hydrophobic surface. Examples of substances which can be included in the compositions are salts of fatty acids and hydrophobic and/or hydrophilic pigments. Also a binder is disclosed.

US 2002/0002932 describes a paint composition which may comprise components such as TiO₂, silica, a fatty acid, a surfactant, and an acrylic latex binder.

WO 2005/042655 presents a aqueous ink and coating composition which consists essentially of: 20-60 wt% of an acrylic resin dispersion, 5-30 wt% of a pigment, 0.5-10
wt% of an alcohol, 20-75 wt% of an aqueous solvent, 0.5-5 wt% of a hydrogenated vegetable derived wax. Examples of pigment include calcium carbonate, and examples of wax include stearic acid.

GB 1452674 mentions a composition comprising calcium carbonate. There is mentioned paper coated with a composition including filler. In one embodiment there is mentioned a coating paste for paper comprising calcium carbonate, calcium stearate, an acrylic emulsion and water.

US 2,576,914 discusses coated paper and a coating composition for paper. In one embodiment it comprises a pigment, a fatty acid salt or a salt thereof, a copolymer of styrene, and water.

JP 2005-036173 discloses an ink and in particular one embodiment comprising precipitated calcium carbonate, a polymer, and a surfactant. It is also mentioned that a fatty acid can be used to treat particles.

US 2006/0141223 A1 relates to textile sheet-like constructions having enhanced watertight properties and to a process for producing them. This fibre modification process requires solvent.

WO 2001/062863 A1 demonstrates an aqueous lacquer dispersion suitable for hydrophobic coatings. The material is mainly carboxylated polystyrene pigments, wherein part of the carboxylic groups are esterified with fluorinated aliphatic alcohols.
US 2006/0257643 A1 illustrates a method of producing hydrophobic composites and aggregates. The process requires several process steps and is not suited to preparing continuous coatings.

FR 2 852 966 concerns an aqueous composition for treating surfaces and making them superhydrophobic, comprising a thermoplastic polymer in an aqueous emulsion and mineral particles having a size from 5 to 500 µm. The mineral can be for instance calcium carbonate, quartz, mica, talc, titanium dioxide, barium sulphate, calcium sulphate etc. The polymer can be for instance polystyrene, polymethacrylate, polyvinyl butyral, and polyurethane.

US 6,712,932 shows a paper or a paper-like material with a structure that comprises particles of for instance metal oxides and carbonates, which are fixed to the paper by means of a wet-laying method using a binder together with a water-repelling agent.

US 6,660,363 concerns self-cleaning surface comprising elevations made of hydrophobic polymers or permanently hydrophobized materials.

US 2005/0136217 A1 relates to a self-cleaning object with a layer of hydrophobic material having protrusions and recesses, which layer is applied with a solution, dispersion or emulsion containing hydrophobic material and a liquid where the liquid is evaporated. The mixture may also comprise other solid particles.

US 2006/0257643 mentions hydrophobic composites, particularly hydrophobic particulates and free-flowing aggregates and methods utilizing the same.
Highly hydrophobic wet-laid coatings may be divided into two main types of coating treatments, either solvent-borne or water-borne. Solvent-borne treatments are subject to controls and/or regulatory limitations in some countries. On the other hand, water-borne coating treatments are usually more difficult to implement in order to obtain hydrophobic coatings and thus more limited than solvent-borne coating treatments, due to the dual requirement for a stable coating dispersion in the aqueous phase and for hydrophobicity in the dry state of the final coating layer. In particular, the application of the superhydrophobic coating often involves multiple steps by creating surface structure and low surface energy coating in different steps. Moreover, standard oil-in-water emulsion-based strategies for encapsulating and delivering hydrophobic species tend to leave an emulsifier (e.g. surfactant) on the coated surface upon drying, which in turn tends to enhance wetting.

Other disadvantages of prior art methods include that they may involve toxic components, that coatings may be made of expensive materials and cannot easily be applied to for instance paper using existing processes and equipment.

Other problems in the prior art related to highly hydrophobic coatings include the use of silane treated and/or fluorinated components, which are expensive and may have negative impact on environment in both producing the components and during the life cycle of the coatings containing these components. Another problem in the prior art is that hydrophobic coatings require multiple steps for the application, which often leads to use of more material and a more complicated process for the application. Thus there is a need for an alternative
coating composition, which would be effective in rendering surfaces hydrophobic without having the above-mentioned drawbacks.

5 **Summary of the invention**

The present invention relates to an aqueous dispersion suitable for use in the manufacture of a hydrophobic coating, said aqueous dispersion comprises inorganic particles, at least one fatty acid or a salt thereof, a polymeric binder, and water, wherein said aqueous dispersion is free of organic solvent.

Moreover the present invention concerns a subject at least partly coated by contacting said substrate with an aqueous dispersion according to the present invention, wherein said subject has an equilibrium contact angle higher than 120° degrees, preferably more than 135°, more preferably more than 150° for a drop of water on the surface.

20 The present invention moreover relates to use of an aqueous dispersion comprising a) inorganic particles, b) at least one fatty acid or a salt thereof, c) a polymeric binder, and d) water, as a coating on a substrate surface, wherein said surface after coating displays an equilibrium contact angle higher than 120° degrees, preferably more than 135°, most preferably more than 150° for a drop of water on the surface.

The present invention further relates to a method for coating a substrate comprising contacting said substrate with an aqueous dispersion according to the invention.
Further embodiments of the present invention are defined in the appended dependent claims, which are specifically incorporated by reference herein.

It is one object of the present invention to obviate at least some of the disadvantages in the prior art and to provide an aqueous dispersion and a subject coated with the aqueous dispersion as well a use of an aqueous dispersion.

Advantages of the present invention include that the coating allows application in one step; it is non-toxic, safe for food packaging, and relatively environmentally friendly and inexpensive. A further advantage is that existing industrial coating processes can be used for applying the coating. Another advantage is that it is possible to obtain highly hydrophobic or superhydrophobic coatings without use of silanes or fluorinated components.

Definitions

Before the invention is disclosed and described in detail, it is to be understood that this invention is not limited to particular ingredients, configurations, method steps, substrates, and materials disclosed herein as such ingredients, configurations, method steps, substrates, and materials may vary somewhat. It is also to be understood that the terminology employed herein is used for the purpose of describing particular embodiments only and is not intended to be limiting since the scope of the present invention is limited only by the appended claims and equivalents thereof.

It must be noted that, as used in this specification and the appended claims, the singular forms "a", "an" and
"the" include plural referents unless the context clearly dictates otherwise.

The term "about" as used in connection with a value throughout the description and the claims means that the true value can be up to 10% higher or down to 10% lower than the indicated value.

If nothing else is defined, any terms and scientific terminology used herein are intended to have the meanings commonly understood by those of skill in the art to which this invention pertains.

The following terms are used throughout the description and the claims.

"Acicular" is used herein to denote a needle-like shape.

"Antioxidant" as used herein denotes a substance capable of preventing, slowing down, and/or suppressing oxidation.

"Apparent density" as used herein denotes dry mass per unit volume of a material including voids inherent in the material.

"Aqueous dispersion" as used herein encompasses a mixture comprising water.

"Aragonite" as used herein denotes a carbonate mineral, one of the two common, naturally occurring polymorphs of calcium carbonate. The other polymorph is the mineral calcite. Aragonite's crystal lattice differs from that of calcite, resulting in a different crystal shape, an orthorombic system with acicular crystals. Repeated
twinning results in pseudo-hexagonal forms. Aragonite may be columnar or fibrous, occasionally in branching stalactitic forms.

"Biocide" as used herein denotes a substance capable of preventing, slowing down, or suppressing growth of living organisms.

"Coalescence agent" as used herein denotes an agent that causes or promotes coalescence.

"Crosslink" as used herein denotes any bond linking one polymer chain to another.

"D_{50}" as used herein denotes the 50\textsuperscript{th} percentile of the mass-weighted size distribution of particles. Accordingly 50% of the inorganic particles have a size greater than D_{50} and 50% of the inorganic particles have a size of less than D_{50}. The particle size is determined for the primary particles if the particles are not aggregated in larger agglomerates, but if the particles are aggregated in larger agglomerates the size of the agglomerates is measured.

"Defoaming agents" as used herein denotes a substance capable of preventing, slowing down, or suppressing foaming.

"Fungicide" as used herein denotes a substance capable of preventing, slowing down, or suppressing growth of fungi.

"Highly hydrophobic" is used herein to denote a surface with an equilibrium contact angle between 120 degrees and 150 degrees for a drop of water on the surface.
"Hydrophobic" as used herein denotes the property to repel water. A hydrophobic surface is a surface with a contact angle of more than 90 degrees but less than 120 degrees.

"Inorganic particle" as used herein encompasses an inorganic particle of any shape.

"Optical brighteners" as used herein include dyes that absorb light in the ultraviolet and violet region of the electromagnetic spectrum and re-emit light in the blue region.

"Polymeric binder" as used herein denotes a binder that is a polymer.

"Rheology modifiers" as used herein denotes a substance with the capability to modify rheological properties of a fluid.

"Scalenohedral" as used herein denotes a pyramidal form under the rhombohedral system, enclosed by twelve faces, each a scalene triangle.

"Substance" as used herein denotes a pure or a non-pure chemical compound or a mixture of chemical compounds, thus for instance a mineral is encompassed within the term.

"Superhydrophobic" as used herein denotes a surface with an equilibrium contact angle higher than 150 degrees for a drop of water on the surface.
Detailed description of the invention

According to the present invention there is provided an aqueous dispersion comprising a) inorganic particles, b) at least one fatty acid or a salt thereof, c) a polymeric binder, and d) water, wherein said aqueous dispersion is free of organic solvent.

In this context free of organic solvent means that the aqueous dispersion is essentially free of organic solvent. An organic solvent in this context is a carbon containing chemical. Essentially free of organic solvent means that very small amounts of organic solvent can be present, although such small amounts are not added, they may nevertheless be present because of impurities in the ingredients. Examples of amounts of accidentally occurring organic solvents include but are not limited to 0.001 wt%, and 0.01 wt%. In one embodiment the amount of organic solvents in the aqueous dispersion is less than 0.01 wt%. In one embodiment there are no organic solvents at all. It is an advantage that the dispersion is free of organic solvent since it is environmentally friendly and safe for food applications.

According to the present invention there is provided an aqueous dispersion comprising a) inorganic particles comprising aragonite, wherein said inorganic particles have a D₅₀ of less than 20 µm, b) at least one fatty acid or a salt thereof, c) a polymeric binder, and d) water.

In one embodiment the amount of inorganic particles is from about 20 to about 55 wt% of the aqueous dispersion. In another embodiment the amount of inorganic particles is from about 30 to about 45 wt%. In a further embodiment the
amount of inorganic particles is from about 30 to about 40 wt%.

In one embodiment the amount of at least one fatty acid or a salt thereof is from about 0.1 to about 2 wt% of the aqueous dispersion. In another embodiment the amount of at least one fatty acid or a salt thereof is from about 0.5 to about 1 wt%. In a further embodiment the amount of at least one fatty acid or a salt thereof, is from about 0.7 to about 0.8 wt%.

In one embodiment the amount of a polymeric binder is from about 5 to about 20 wt% of the aqueous dispersion. In another embodiment the amount of a polymeric binder is from about 8 to about 15 wt%. In a further embodiment the amount of the amount of a polymeric binder is from about 10.5 to about 12 wt%.

The remaining parts up to 100 wt% are water and optional additives.

In one embodiment the inorganic particles according to the present invention have an apparent density from about 0.30 g/ml to about 4 g/ml, and a BET specific surface area from about 1 to 20 m²/g.

In one embodiment of the present invention the inorganic particles have the following properties:

- Apparent density from about 0.30 g/ml to about 2.7 g/ml, preferably from about 0.30 g/ml to about 0.80 g/ml and most preferably about 0.30 g/ml to about 0.65 g/ml.
- BET specific surface area from about 1 to 20 m²/g, preferably more than about 3 m²/g and more preferably more than 5 m²/g.
• \( D_{50} \) less than about 20 µm, preferably from about 1 to about 10 µm, most preferably from about 2 to 5 µm.

In an alternative embodiment, of the present invention the inorganic particles the following properties:

• Apparent density from about 1 g/ml to about 4 g/ml, preferably from about 2 to about 3.5 g/ml and most preferably about 2.5 g/ml to about 2.9 g/ml.
• BET specific surface area from about 1 to 20 m\(^2\)/g, preferably more than about 6 m\(^2\)/g and more preferably more than 9 m\(^2\)/g.
• \( D_{50} \) less than about 10 µm, preferably from about 0.1 to about 5 µm, most preferably from about 0.2 to 2 µm.

The inorganic particles used in the present invention may comprise a mixture of different inorganic particles with different properties.

In one embodiment the inorganic particles used in the present invention comprise at least one substance selected from the group consisting of anatase, calcinated clay, kaolin, and talc, the surface of which has been made basic.

In one embodiment the inorganic particles used in the present invention comprise at least one substance selected from the group consisting of aluminium hydroxide, barium sulphate, calcite, calcium sulphate, dolomite, magnesium hydroxide, magnesium carbonate, magnesite, titanium dioxide (rutile), and vaterite.

In one embodiment the inorganic particles used in the present invention comprise at least one substance selected...
from the group consisting of zincite, corundum, hematite, magnetite, ilmenite, and cassiterite.

In one embodiment the inorganic particles used in the present invention comprise at least one substance selected from the group consisting of dispore, boehmite, goethite, lepidocrocite, rhodocrosite, siderite, baryte, strontianite, apatite, feldspar and fluorite.

In one embodiment the inorganic particles used in the present invention comprise SiO₂.

The inorganic particles preferably comprise calcium carbonate particles, more preferably precipitated calcium carbonate and most preferably aragonite.

In one embodiment the inorganic particles are surface treated with a fatty acid or a salt thereof. Advantages of using one or more fatty acids include that fatty acids are inexpensive compared to silanes and fluorinated polymers and are readily available and used in many industries. Fatty acids interact in a suitable way with inorganic particles comprising calcium carbonate and many fatty acids are approved for contact with food. The inorganic particles are coated by contacting them with an aqueous solution or dispersion comprising a fatty acid or a salt thereof. In one embodiment the aqueous solution or dispersion that comprises the fatty acid or a salt thereof further comprises the binder.

In an alternative embodiment the coating is performed in a separate aqueous solution or dispersion comprising a fatty acid or a salt thereof.
In one embodiment the coating of the inorganic particles is performed in a separate aqueous solution or dispersion.

In one embodiment the inorganic particles are coated with several different fatty acids of salts thereof, optionally in several steps.

In one embodiment the fatty acid or salt thereof forms a layer on the entire inorganic particle surface; alternatively the fatty acid or salt thereof forms a layer on a part of the surface.

In one embodiment the coating comprises at least one surfactant.

The amount of fatty acid or salt thereof should be high enough so that the inorganic particles become dispersible in water.

In one embodiment the amount of fatty acid corresponds to a double layer of molecules on the surface of the inorganic particles.

Thus the inorganic particles become dispersible in water or alternatively the ability to be dispersed in water is improved.

Inorganic particles with relatively poor packing are preferred, which yields a suitable roughness of the coating. Inorganic particles with narrow particle size distribution are used in one embodiment. In another embodiment inorganic particles are combined with a tendency to aggregate to larger secondaries.
In one embodiment the inorganic particles are acicular or scalenohedral. The shape of the particles is however not limited to these two shapes. Also other thorny, spiky and needle like shapes are used in other embodiments according to the present invention. Other possible shapes include but are not limited to chestnut husk shapes.

In one embodiment particles with a small size, low density and high specific surface are used.

In one embodiment inorganic particles without any appreciable residues of dispersants are used. It was found in some cases that dispersant added during the manufacture of particles comprising aragonite can interfere with the adsorption of fatty acid or a salt thereof on the particle surface. Examples of such undesired dispersants include but are not limited to sodium polyacrylate polymers and copolymers thereof. For certain embodiments it is therefore important to use inorganic particles with no or only very low amounts of added dispersants.

In one embodiment acicular aragonite is used for the inorganic particles, especially acicular aragonite with a D₅₀ from about 0.1 to about 20 µm, preferably from about 0.2 to about 10 µm. It has turned out that aragonite gives coatings with high contact angle for a drop of water on the surface.

In an alternative embodiment, PCC (precipitated calcium carbonate) and/or GCC (ground calcium carbonate) is used.

A saturated or unsaturated fatty acid or salt thereof is used. A fatty acid or salt thereof with linear or branched hydrocarbon chain is used. Preferably the fatty acid or
salt thereof has 8 to 22, more preferably 10 to 18 carbon
atoms. In one embodiment the fatty acids are selected from
the group consisting of oleic acid, stearic acid and
palmitic acid. In one embodiment the salts of the above
mentioned fatty acids are used. The counter ions of a
fatty acid salt can be any suitable ion. Examples include
but are not limited to sodium ions and ammonium ions,
which are common and inexpensive salts.

In one embodiment a fatty acid and a salt thereof are used
together.

Examples of binders include but are not limited to
carboxylated latex, styrene-butadiene latex and styrene
acrylate. Such carboxylated latex is a latex or emulsion
polymer stabilised predominantly by carboxylation. In one
embodiment the glass transition temperature of the binder
is in the range from about -40 to about 50°C. In another
embodiment the glass transition temperature of the binder
is in the range from about 0 to about 50°C. Examples of a
polymeric binder include but are not limited to
commercially available binders supplied by Dow Chemical
Company® under the trade name DL 940® or experimental
latex Dow/HPQ73® or by Rhodia® under the trade name
Ultradia® 7100, 7300 or 7400.

Synthetic latex, as is well known, is an aqueous
dispersion of polymer particles prepared by emulsion
polymerization of one or more monomers.

In one embodiment the monomer composition employed in the
preparation of the latex comprises from about 10 to 95
pphm of a first monomer (A), from about 40 to 90 pphm of a
second monomer (B), and from 0 to about 5 pphm of a
functional monomer (C). As used herein, the term "pphm" means parts per hundred monomer, a term well known to those skilled in the art. Accordingly, the total parts monomer employed is 100 parts monomer, on a weight basis.

The first monomer (A) is a low Tg monomer comprising an alkyl acrylate or butadiene. The low Tg monomer is used in amounts of from about 10 pphpm to about 95 pphpm, preferably 15 pphpm to 40 pphpm. Examples of low Tg monomers include but are not limited to monomers having a Tg of less than 10°C that are Ci-Cio alkyl esters of acrylic acid, C2-C10 alkyl esters of alpha, beta-ethylenically unsaturated C4-C6 monocarboxylic acids, C4-C10 dialkyl esters of alpha, beta-ethylenically unsaturated C4-Cs dicarboxylic acids, and vinyl esters of carboxylic acids, including, without limitation, vinyl isobutyrate, vinyl-2-ethyl-hexanoate, vinyl propionate, vinyl isoctanoate and vinyl versatate and butadiene. The low Tg monomer can be selected from the group consisting of C1-C10 alkyl esters of (meth)acrylic acid, i.e. alkyl (meth)acrylates, and C4-Cs dialkyl esters of maleic, itaconic and fumaric acids. Preferably, at least one C2-C8 alkyl ester of acrylic acid is utilized. Particularly preferred low Tg monomers include but are not limited to ethyl acrylate, butyl acrylate, 2-ethyl hexyl acrylate, decyl acrylate, dibutyl maleate, dioctyl maleate, and butadiene with butadiene being most preferred. Mixtures of first monomers can be employed.

The second monomer (B) is a high Tg monomer having a Tg greater than 10°C such as, for example, vinyl esters of carboxylic acids, the acid having from two to about 13 carbon atoms and styrene. Representative high Tg comonomers include but are not limited to methyl methacrylate, dimethyl maleate, t-butyl methacrylate, t-
butyl isobornyl acrylate, phenyl methacrylate, acrylonitrile and vinyl esters of carboxylic acids having Tg of greater than 100°C and styrene. Examples of such vinyl esters include but are not limited to vinyl pivalate, vinyl neodecanoate, vinyl neononanoate, and mixtures of branched vinyl esters such as the commercially available VeoVa 11 and EXXAR Neo-12. The second monomer is in one embodiment employed in an amount of from about 40 pphm to about 90 pphm, preferably 60 pphm to 85 pphm. Mixtures of high Tg comonomers can be employed.

It may also be desired to incorporate in the binder polymer minor amounts of one or more functional comonomers (C). Suitable copolymerizable comonomers (C) include but are not limited to, for example: acrylic acid; methacrylic acid; itaconic acid; fumaric acid; the half esters of maleic acid, such as monoethyl, monobutyl or monooctyl maleate; acrylamide; tertiary octylacrylamide; N-methylol (meth)acrylamide; N-vinylpyrrolidinone; diallyl adipate; triallyl cyanurate; butanediol diacrylate; allyl methacrylate; etc.; as well as C2-C3 hydroxyalkyl esters such as hydroxyethyl acrylate, hydroxy propyl acrylate and corresponding methacrylates. The comonomer (C) generally is used at levels of less than 5 pphm, preferably less than 2.5 pphm, depending upon the nature of the specific comonomer. Mixtures of comonomer (C) can be employed.

In addition, certain copolymerizable monomers that assist in the stability of the binder, e.g., vinyl sulfonic acid, sodium vinyl sulfonate, sodium styrene sulfonate, sodium allyl ether sulfate, sodium 2-acrylamide-2-methyl-propane sulfonate (AMPS), 2-sulfoethyl methacrylate, and 2-sulfopropyl methacrylate, can be employed as emulsion stabilizers. These optional monomers, if employed, are
added in very low amounts of from 0.1 pphm to about 2 pphm.

Methods for preparing synthetic latexes are well known in the art and any of these procedures can be used.

Suitable free radical polymerization initiators are the initiators known to promote emulsion polymerization and include but are not limited to water-soluble oxidizing agents, such as, organic peroxides (e.g., t-butyl hydroperoxide, cumene hydroperoxide, etc.), inorganic oxidizing agents (e.g., hydrogen peroxide, potassium persulfate, sodium persulfate, ammonium persulfate, etc.) and those initiators that are activated in the water phase by a water-soluble reducing agent. Such initiators are employed in an amount sufficient to cause polymerization. As a general rule, a sufficient amount is from about 0.1 to about 5 pphm. Alternatively, redox initiators may be employed, especially when polymerization is carried out at lower temperatures. For example, reducing agents may be used in addition to the persulfate and peroxide initiators mentioned above. Typical reducing agents include but are not limited to: alkali metal salts of hydrosulfites, sulfoxylates, thiosulfates, sulfites, bisulfites, reducing sugars such as glucose, sorbose, ascorbic acid, erythorubic acid, and the like. In general, the reducing agents are used at levels from about 0.01 pphm to about 5 pphm.

The emulsifying agents are those generally used in emulsion polymerization. The emulsifiers can be anionic, cationic, surface-active compounds or mixtures thereof.

Suitable nonionic emulsifiers include but are not limited to polyoxyethylene condensates. Exemplary polyoxyethylene
condensates that can be used include but are not limited to polyoxyethylene aliphatic ethers, such as polyoxyethylene lauryl ether and polyoxyethylene oleyl ether; polyoxyethylene alkaryl ethers, such as polyoxyethylene nonylphenol ether and polyoxyethylene octylphenol ether; polyoxyethylene esters of higher fatty acids, such as polyoxyethylene laurate and polyoxyethylene oleate, as well as condensates of ethylene oxide with resin acids and tall oil acids; polyoxyethylene amide and amine condensates such as N-polyoxyethylene lauramide, and N-lauryl-N-polyoxyethylene amine and the like; and polyoxyethylene thio-ethers such as polyoxyethylene n-dodecyl thio-ether.

Nonionic emulsifying agents that can be used also include but are not limited to a series of surface active agents available from BASF® under the Pluronic® and Tetronic® trade names. In addition, a series of ethylene oxide adducts of acetylenic glycols, sold commercially by Air Products® under the Surfynol® trade name, are suitable as nonionic emulsifiers.

Representative anionic emulsifiers include but are not limited to the alkyl aryl sulfonates, alkali metal alkyl sulfates, the sulfonated alkyl esters, and fatty acid soaps. Specific examples include but are not limited to sodium dodecylbenzene sulfonate, sodium butynaphthalene sulfonate, sodium lauryl sulfate, disodium dodecyl diphenyl ether disulfonate, N-octadecyl sulfo succinate and dioctyl sodiumsulfosuccinate. The emulsifiers are employed in amounts effective to achieve adequate emulsification of the polymer in the aqueous phase and to provide desired particle size and particle size distribution.
Other ingredients known in the art to be useful for various specific purposes in emulsion polymerization, such as, acids, salts, chain transfer agents, chelating agents, buffering agents, neutralizing agents, defoamers and plasticizers also may be employed in the preparation of the polymer. For example, if the polymerizable constituents include a monoethylenically unsaturated carboxylic acid monomer, polymerization under acidic conditions (pH 2 to 7, preferably 2 to 5) is preferred. In such instances the aqueous medium can include those known weak acids and their salts that are commonly used to provide a buffered system at the desired pH range.

Various protective colloids may also be used in place of or in addition to the emulsifiers described above. Suitable colloids include but are not limited to casein, hydroxyethyl starch, carboxymethyl cellulose, carboxymethyl cellulose, hydroxyethylcellulose, gum arabic, alginate, poly (vinyl alcohol), polyacrylates, polymethacrylates, styrene-maleic anhydride copolymers, polyvinylpyrrolidones, polyacrylamides, polyethers, and the like, as known in the art of emulsion polymerization technology. In general, when used, these colloids are used at levels of 0.05 to 10% by weight based on the total weight of the reactor contents.

The manner of combining the polymerization ingredients can be by various known monomer feed methods, such as, continuous monomer addition, incremental monomer addition, or addition in a single charge of the entire amounts of monomers. The entire amount of the aqueous medium with polymerization additives can be present in the polymerization vessel before introduction of the monomers, or alternatively, the aqueous medium, or a portion of it,
can be added continuously or incrementally during the course of the polymerization.

Final particle size of the latex can vary from 30nm to 1500nm.

The amount of binder must be high enough so that the coating exhibits the desired adhesion, mechanical strength and hydrophobicity, but on the other hand the amount of binder must not be too high so that the hydrophobicity of the coating is reduced by the binder submerging the inorganic particles. A person skilled in the art can in the light of this description adjust the amount of binder within the ranges described in the description.

The degree of carboxylation for carboxylated latex should be adapted relative to the amount of fatty acid in the coating composition. It is undesired that the total number of carboxyl groups of the binder is higher than the total number of carboxyl groups of the fatty acid. Therefore, in general, low-carboxylated latex should perform best for low amounts of fatty acid.

The process for making the above-defined coating composition can be carried out in several ways according to the present invention. In one embodiment the process comprises the step of mixing an aqueous solution of a polymeric binder and a mixture of fatty acid and inorganic particles. In one embodiment the process comprises the step of coating the inorganic particles with a fatty acid. Said coating occurs in the mixture of fatty acid and inorganic particles.
In one embodiment the aqueous dispersion is prepared by mixing an aqueous dispersion of a polymeric binder with a mixture of inorganic particles and at least one fatty acid or a salt thereof.

In one embodiment the mixture of inorganic particles and at least one fatty acid or a salt thereof is prepared by
a) mixing said at least one fatty acid or a salt thereof with water,
b) mixing said inorganic particles with water, and then
c) mixing the mixtures from step a) and step b).

In an alternative embodiment said mixture of inorganic particles and at least one fatty acid or a salt thereof is prepared by
a) mixing at least one fatty acid or a salt thereof with water, and then
b) mixing the mixture from step a) with said inorganic particles.

In one embodiment the polymeric binder is mixed with said inorganic particles at least 15 minutes after the mixing of said at least one fatty acid or a salt thereof with said inorganic particles.

In one embodiment the aqueous dispersion according to the present invention further comprises surfactants. If a surfactant is used in the aqueous dispersion it can be added before, at the same time as, or after the fatty acid or salt thereof. An optional surfactant can also be added before, at the same time as, or after the polymeric binder. The surfactant is chosen so that it does not adversely affect the coating. Cationic surfactants are less preferred. Examples of surfactants include but are
not limited to phosphoric acid alkyl ester and diphosphonate surfactants, silicone based surfactants, fluorosurfactants, and salts thereof.

In one embodiment the aqueous dispersion further comprises additives. Examples of such additives include but are not limited to at least one additive selected from antioxidants, biocides, coalescence agents, coloured inorganic particles, crosslinkers, defoaming agents, dyes, coalescence agents, fungicides, lubricants, optical brighteners, rheology modifiers, or any combination thereof. Preferably such additives are compatible with the other components of the aqueous dispersion.

The present invention provides a method for coating a substrate with a highly hydrophobic or superhydrophobic coating comprising: a) preparing an aqueous dispersion, b) contacting said substrate with said aqueous dispersion. Thereby the surface of a substrate is rendered hydrophobic, highly hydrophobic or superhydrophobic.

Thus in one embodiment there is disclosed a method for coating a substrate comprising contacting the substrate with an aqueous dispersion, wherein the aqueous dispersion comprises a) inorganic particles comprising aragonite, wherein said inorganic particles have a D₅₀ of less than 20 µm, b) at least one fatty acid or a salt thereof, c) a polymeric binder, and d) water.

The aqueous dispersion as described above is contacted with the substrate to be coated. After contacting the substrate with the aqueous dispersion the substrate is in one embodiment dried. The thickness of the dried coating on the substrate is in one embodiment from about 3 to
about 40 µm. In another embodiment the thickness of the dried coating on the substrate is from about 8 to about 25 µm.

The amount of polymeric binder depends on several variables including the surface area of the inorganic particles. The larger the surface area, the more polymeric binder is required for strength properties.

After the substrate has been contacted with the aqueous dispersion according to the present invention it is in one embodiment heated. Heating is particularly useful if a short drying time and high process speed is needed. The properties of the polymeric binder are in one embodiment improved by curing by heat. If paper or paper-like material is the substrate, heating is in one embodiment used to decrease the water content of the coated paper. Heating is also used in one embodiment to decrease the water content of any other substrate.

In an alternative embodiment the object is dried without additional heating. In one embodiment a combination of drying and heating is used.

In one embodiment the method of coating a substrate comprises at least one method step selected from the group consisting of spray coating, dip coating, roll application, free jet application, blade metering, rod metering, metered film press coating, air knife coating, curtain coating, flexography printing, roll coating, and powder coating.

The coating according to the present invention may be applied to a large variety of substrates.
According to the present invention there is provided objects coated with the method according to the present invention.

Coating for medical devices is possible.

In one embodiment the coating according to the present invention is highly hydrophobic, i.e. it displays an equilibrium contact angle between 120 degrees and 150 degrees. In another embodiment the contact angle is higher than 135 degrees. Using the present invention it is even possible to manufacture superhydrophobic coatings, which display an equilibrium contact angle greater than 150 degrees.

Thus there is provided a subject at least partly coated by contacting said substrate with an aqueous dispersion according to the present invention, wherein said subject has an equilibrium contact angle higher than 120° degrees, preferably more than 135°, more preferably more than 150° for a drop of water on the surface.

In one embodiment the aqueous dispersion used for the coating comprises a) inorganic particles comprising aragonite, wherein said inorganic particles have a D₅₀ of less than 20 µm, b) at least one fatty acid or a salt thereof, c) a polymeric binder, and d) water.

In one embodiment the subject is partly coated. In an alternative embodiment the subject is entirely coated.

Advantages of the present invention include that the coating allows application in one step, it is non-toxic,
approved for food contact, inexpensive and it can be produced in an environmentally friendly manner. A further advantage is that existing industrial coating processes can be used for applying the coating. Another advantage is that a hydrophobic surface is created without any need for stamping or etching.

It is to be understood that this invention is not limited to the particular embodiments shown here. The following examples are provided for illustrative purposes and are not intended to limit the scope of the invention since the scope of the present invention is limited only by the appended claims and equivalents thereof.

**Examples**
The following methods apply to all examples mentioned below.

**Dry stain size measurement**

In the stain test 5 drops of an exact amount (9 µl, i.e. drop diameter 2.58 mm) of a blue dye aqueous solution are auto-pipetted (from a fixed height of 1.9 mm from drop bottom to coat surface) on the coated surface. The blue dye is added to aid visual inspection of stain size after complete evaporation of the water. The surface tension of the colored water is the same as the non-colored deionized water. The samples are stored at 23°C and 50% relative humidity, and the final size of the dry stain after complete evaporation is measured with a sliding gauge, both in machine direction MD and cross direction CD. The values given below correspond to the mean of the set of 5 drops measured in these two directions. They are expressed in a dimensionless form by dividing the stain diameter by the drop diameter prior to contact (i.e. 2.58 mm). This
measure relates to the total ability of the substrate to resist both surface spreading and sub-surface penetration and spreading (within the top coating layer and layers below) over long times. A hydrophobic surface leads to a smaller stain diameter than the initial droplet diameter. This method can be used to rank the samples' performance regarding hydrophobicity.

*Initial contact angle measurement*

Short-time contact angles of drops of deionized water (i.e. without the blue dye) on the coated sheets are measured manually with a Rame-Hart goniometer, using the same autopipette, drop volume and procedure (i.e. 5 drops at different places) as in the staining experiments described above. The time from contact to measurement of advancing angle is approximately 10 s. This is a standard measure of short-term hydrophobicity, reflecting the ability of the substrate to reject water drops on first contact. The use of a manual goniometer is convenient because the drop can roll or hop on initial contact with a highly hydrophobic or superhydrophobic substrate.

Measurement of contact angles are described in further detail by Strom et al. in J. Colloid Interface Sci., Vol. 123, No. 2, pages 324-338, 1988, which is explicitly incorporated herein by reference in its entirety.

*Rolling angle measurement*

The drop rolling tests are performed using a tilt table. The same blue dye solution as mentioned above is autopipetted in a similar manner as in the stain test on the coated samples pre-inclined at 5 fixed angles (2.5°, 5°, 10°, 15° and 20° from horizontal). The lowest angle for which free rolling occurs, i.e. the drop rolls the entire distance of the sample size (around 10 cm), is the
value assigned to the substrate. Failure to roll freely at 20° is regarded as a no-score, despite the fact that free rolling may occur at higher angles not tested (e.g. approaching vertical). It is expected that drop rolling is closely dependent on advancing initial contact angle (see above).

Example 1
Precipitated calcium carbonate (PCC) (Sturcal® F, Specialty Minerals Inc.®) (Particle size D50 ca 2.5 μm, apparent density 0.32-0.43 g/ml, and BET surface area ca 6 m²/g) (Aragonite content is minimum 50%) particles were mixed together with water and a sodium oleate solution in a glass beaker with a magnetic stirrer. The total content of calcium carbonate in water was 30 wt% and the content of sodium oleate was 1 wt% per pigment weight (dry on dry). This suspension was mixed until it was essentially homogenous. Further, 30 wt% per pigment weight (dry on dry) of commercially available styrene-butadiene (SB) latex (DL 940, The Dow Chemical Company®) was added to the above-mentioned aqueous suspension containing sodium oleate and calcium carbonate. This suspension was again mixed with a magnetic stirrer to obtain an essentially homogenous mixture. The aqueous dispersion was coated on paper (Performa Natura®, 255 g/m², Stora Enso®). The coating was performed using a bench coater from RK Print-Coat Instruments Ltd.® Several sheets of paper were coated and then dried in an oven at 70 °C for 2 minutes.

Water contact angle on coated paper was characterized by Rame Hart goniometer. The analysis performed for coatings are explained thoroughly above. For the above-mentioned coating the contact angle was 140° and the normalized stain size was 1.0. The rms roughness of the coated
substrate measured by Zygo white light interferometric profilometer (NewView 5010, Zygo Corporation®) was below 1.2 µm for following length scales: 0-5 µm, 5-10 µm, 10-20 µm, 20-40 µm, 40-80 µm and 80-170 µm.

Example 2
The formulation of Example 1 was modified by having 50 wt% per pigment weight of the same SB-latex binder (DL 940, The Dow Chemical Company®) instead of 30 wt% per pigment weight as used in Example 1. The coating and the analysis were performed as in Example 1. For this coated substrate the contact angle was 120° and the normalized stain size was 1.4. The rms roughness measured by Zygo® white light interferometric profilometer was below 1.2 µm for following length scales: 0-5 µm, 5-10 µm, 10-20 µm, 20-40 µm, 40-80 µm and 80-170 µm.

Example 3
The formulation of Example 1 was modified by having 17 wt% per pigment of the same SB-latex binder (DL 940, The Dow Chemical Company®) instead of 30 wt% per pigment as used in Example 1. The coating and the analysis were performed as in Example 1. For this coated substrate the contact angle was 145° and the normalized stain size was 2.5.

Example 4
The formulation of Example 1 was modified by having 2 wt% per pigment weight of sodium oleate instead of 1 wt% per pigment weight as used in Example 1. The coating and the analysis were performed as in Example 1. For this coated substrate the contact angle was 142° and the normalized stain size was 0.8. Water droplets (9 µl) showed rolling at 15° inclination.
Example 5

The formulation of Example 1 was modified by having 3 wt% per pigment weight of sodium oleate instead of 1 wt% per pigment weight as used in Example 1. The coating and the analysis were performed as in Example 1. For this formulation the contact angle was 154° and the normalized stain size was 0.8. Water droplets (9 µl) showed rolling at 10° inclination.

Example 6

The formulation of Example 1 was modified by having 2 wt% per pigment weight of sodium oleate instead of 1 wt% per pigment weight as used in Example 1. Further, the formulation in Example 1 was modified by having 40 wt% per pigment weight of the same SB-latex binder (DL 940, The Dow Chemical Company®) instead of 30 wt% per pigment. The coating and the analysis were performed as in Example 1. For this coated substrate the contact angle was 146° and the normalized stain size was 0.9. Water droplets (9 µl) showed rolling at 20° inclination.

Example 7

The formulation of Example 1 was modified by changing the pigment to another type of PCC calcium carbonate, Sturcal® H (Specialty Minerals Inc.®). (Particle size D₅₀ 4.0 µm, apparent density 0.48–0.61 g/ml, and BET surface area ca 5 m²/g) (Aragonite content is minimum 50%) Further, the formulation in Example 1 was modified by adding 2 wt% per pigment of sodium oleate instead of 1 wt% per pigment as used in Example 1. The coating and the analysis were performed as in Example 1. For this coated object the contact angle was 153° and the normalized stain size was 0.6. Water droplets (9 µl) showed rolling at 5° inclination.
Example 8
The formulation of Example 7 was modified by changing the SB-latex binder to an experimental grade SB-latex (SB/HPQ73, The Dow Chemical Company®). The SB-latex content was again 30 wt% per pigment weight. The coating was performed on commercially available paper (Cupforma Classic®, 230 g/m², Stora Enso®). Otherwise the coating and analysis were performed as in Example 1. For this coated substrate the contact angle was 143° and the normalized stain size was 0.6. Water droplets (9 µl) showed rolling at 10° inclination.

Example 9
The formulation of Example 8 was modified by increasing the total solids content of the coating from 34 wt% to 51 wt%. Otherwise the coating and analysis were performed as in Example 8. For this coated substrate the contact angle was 160° and the normalized stain size was 0.5. Water droplets (9 µl) showed rolling at 2.5° inclination.

Example 10
The formulation of Example 8 was modified by adding commercially available ammonium zirconium crosslinker (Allicross® AZC-R, Allinova®). The crosslinker content was 4 wt% dry based on dry binder weight. Otherwise the coating and analysis were performed as in Example 8. For this coated substrate the contact angle was 141° and the normalized stain size was 0.6. Water droplets (9 µl) showed rolling at 15° inclination.

Example 11
The formulation of Example 9 was modified by changing the binder to the commercially available latex Rhodopas
Ultradia® 7100 (Rhodia®). The binder content was 30 wt% per pigment weight (dry on dry). The coating and analysis were performed as in Example 8. For this coated substrate the contact angle was 155° and the normalized stain size was 0.6. Water droplets (9 µl) showed rolling at 2.5° inclination.

Example 12
The formulation of Example 11 was modified by changing the binder to the commercially available latex Rhodopas Ultradia® 7300 (Rhodia). The binder content was again 30 wt% per pigment weight. The coating and analysis were performed as in Example 8. For this coated substrate the contact angle was 154° and the normalized stain size was 0.6. Water droplets (9 µl) showed rolling at 10° inclination.

Example 13
The formulation of Example 9 was modified by using diphosphonate alkyl surfactant instead of the sodium oleate. The content of diphosphonate alkyl surfactant was 1 wt% per pigment weight (dry on dry). The coating and analysis were performed as in Example 8. For this coated substrate the contact angle was 139° and the normalized stain size was 0.9.

Example 14
The formulation of Example 13 was modified by changing the diphosphonate alkyl surfactant content to 2 wt% per pigment weight. The coating and analysis were performed as in Example 8. For this coated substrate the contact angle was 136° and the normalized stain size was 0.9.
Example 15

The formulation of Example 8 was modified by changing the Sturcal® H pigment to the PCC Opacarb® A40 (Specialty Minerals Inc.®) (Particle size D_{50} 0.4 \mu m, density 0.48-0.61 g/ml, and BET surface area 12 m^2/g). (Opacarb® A40 comprises aragonite) The total solids content of the formulation was 30 wt%. The coating and analysis were performed as in Example 8. For this formulation the contact angle was 148° and the normalized stain size was 0.6. Water droplets (9 \mu l) showed rolling at 10° inclination.

Example 16

Aluminium sulfate, Al₂(SO₄)₃·18H₂O was mixed together with water. This solution was mixed with kaolin (Kaolin C, ECC), (Particle size D_{50} <2 \mu m, density 2.6 g/ml, and BET surface area 10 m^2/g) stirring until it was essential homogenous. Then a 10% solution of sodium carbonate was added, and the mixture stirred until again homogenous. To this slurry a 5% solution of sodium oleate was added. The total content of kaolin in the aqueous dispersion was 29 wt%, the content of sodium oleate was 2.4 wt% per pigment weight (dry on dry), the content of sodium carbonate was 4.3 wt% on pigment (dry on dry), and the content of Al₂(SO₄)₃·18H₂O was 9.4 wt% per pigment weight (dry on dry).

The aqueous dispersion was coated on paperboard (Cupforma Classic®, 230 g/m², Stora Enso®). The coating was performed using a bench coater from RK Print-Coat Instruments Ltd.® Several sheets of paper were coated and then dried in an oven at 90 °C for 2 minutes. For the above-mentioned coating the contact angle with water was 132°.
Example 17
To the aqueous dispersion from Example 16, a styrene-
butadiene latex binder (HPQ 73, Dow® Europe) was added,
and the dispersion was stirred until homogeneous. The
content of latex was 30 wt% per pigment weight (dry on
dry). The coating and the analysis were performed as in
Example 16. For this coated substrate the contact angle
was 113°.

Example 18
The formulation in Example 17 was modified by increasing
the level of the latex HPQ 73 to 30 wt% per pigment weight
(dry on dry). The coating and the analysis were performed
as in Example 16. For this coated substrate the contact
angle was 90°.

Example 19
Precipitated calcium carbonate (PCC) (Sturcal® H,
Specialty Minerals Inc.®) (Particle size D_{50} 4.0 \mu m,
apparent density 0.48-0.61 g/ml, and BET surface area ca 5
m²/g) (Aragonite content is minimum 50%) particles were
mixed together with water and a sodium oleate solution in
a glass beaker with a magnetic stirrer. The total content
of calcium carbonate in water was 30 wt% and the content
of sodium oleate was 2 wt% per pigment weight (dry on
dry). This suspension was mixed until it was essentially
homogenous and then added to the aqueous dispersion in
Example 16 and again mixed until homogeneous. The
resulting ratio of PCC pigment to clay pigment was 70% PCC
and 30% clay. To this aqueous dispersion, a styrene-
butadiene latex binder (HPQ 73, Dow® Europe) was added,
and the dispersion was stirred until homogeneous. The
content of latex was 15 wt% per pigment weight (dry on
dry). The coating and the analysis were performed as in
Example 16. For this coated substrate the contact angle was 123°.

Example 20
The formulation in Example 19 was modified by changing the ratio of PCC pigment to clay pigment to 50% PCC and 50% clay (dry on dry). The coating and the analysis were performed as in Example 16. For this coated substrate the contact angle was 115°.

Example 21
The formulation of Example 1 was modified by having 2 wt% per pigment of sodium oleate instead of 1 wt% per pigment weight as used in Example 1. Further, the formulation in Example 1 was modified by having 30 wt% per pigment weight of another SB-latex binder (SHY-7, Dow® Europe) instead of DL 940. The aqueous dispersion was coated on paperboard (Cupforma Classic®, 230 g/m², Stora Enso®). The coating was performed using a bench coater from RK Print-Coat Instruments Ltd.® Several sheets of paper were coated and then dried in an oven at 90 °C for 2 minutes. The coating and the analysis were performed as in Example 1. For this coated substrate the contact angle was 148° and the normalized stain size was 0.7. Water droplets (9 µl) showed rolling at 2.5° inclination.

Example 22 (comparative, not according to the present invention)
A comparative formulation was prepared by utilising commercially available dispersant (Dispex N40, Ciba Specialty Chemicals®). As particles, PCC Opacarb® A40 (Specialty Minerals Inc.®) (Particle size D₅₀ 0.4 µm, density 2.8 g/ml, and BET surface area 12 m²/g) were used. The content of Dispex® N40 was 0.05 wt% per pigment weight
and no fatty acid or other surfactant were added. The coating and analysis were performed as in Example 8. For this formulation the contact angle was 71° and the normalized stain size was 2.1. Water droplets (9 µl) did not show rolling at any inclination.
Claims

1. An aqueous dispersion comprising:
   a) inorganic particles
   b) at least one fatty acid or a salt thereof,
   c) a polymeric binder, and
   d) water,

   wherein said aqueous dispersion is free of organic solvent.

2. An aqueous dispersion according to claim 1,

   comprising:
   a) inorganic particles comprising aragonite, wherein
       said inorganic particles have a D_{50} of less than 20
       μm,
   b) at least one fatty acid or a salt thereof,
   c) a polymeric binder, and
   d) water.

3. The aqueous dispersion according to any one of claims

   1-2, wherein said inorganic particles have an apparent density
   from about 0.30 g/ml to about 4 g/ml, and wherein
   said inorganic particles have a BET specific surface area
   from about 1 m²/g to about 20 m²/g.

4. The aqueous dispersion according to any one of claims

   1-3, wherein said fatty acid or salt thereof has 8 to 22
   carbon atoms.

5. The aqueous dispersion according to any one of claims

   1-4, wherein said polymeric binder is selected from the
   group consisting of a carboxylated latex, a styrene-
   butadiene latex and a styrene acrylate.
6. The aqueous dispersion according to any one of claims 1-5 further comprising at least one additive selected from the group consisting of an antioxidant, a biocide, a coalescence agent, a coloured inorganic particle, a crosslinker, a defoaming agent, a dye, a fungicide, a lubricant, an optical brightener, a rheology modifier, or any combination thereof.

7. The aqueous dispersion according to any one of claims 1-6 further comprising a zirconium crosslinker.

8. A subject at least partly coated by contacting said substrate with an aqueous dispersion according to any one of claims 1-7, wherein said subject has an equilibrium contact angle higher than 120° degrees, preferably more than 135°, more preferably more than 150° for a drop of water on the surface.

9. Use of an aqueous dispersion comprising:
   a) inorganic particles
   b) at least one fatty acid or a salt thereof,
   c) a polymeric binder, and
   d) water,
as a coating on a substrate surface, wherein said surface after coating displays an equilibrium contact angle higher than 120° degrees, preferably more than 135°, most preferably more than 150° for a drop of water on the surface.

10. Use according to claim 9, wherein said inorganic particles are acicular or scalenohedral.

11. Use according to any one of claims 9-10, wherein said inorganic particles comprise at least one entity selected
from ground calcium carbonate, and precipitated calcium carbonate.

12. Use according to any one of claims 9-11, wherein said inorganic particles comprise aragonite.

13. Use according to any one of claims 9-12, wherein said substrate is contacted with said aqueous dispersion and thereafter heated.


15. The method according to claim 14, wherein said substrate further is heated after contacting said substrate with said aqueous dispersion.

16. The method according to any one of claims 14-15, wherein said method comprises at least one method step selected from the group consisting of spray coating, dip coating, roll application, free jet application, blade metering, rod metering, metered film press coating, air knife coating, curtain coating, flexography printing, roll coating, and powder coating.
# INTERNATIONAL SEARCH REPORT

**Date**
10 July 2008

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European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016
Matthijsen, J-J

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## A. CLASSIFICATION OF SUBJECT MATTER

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

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols):

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

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

EPO-Internal, WPI Data

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

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Further documents are listed in the continuation of Box C.

See patent family annex.

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"A" document defining the general state of the art which is not considered to be of particular relevance.

"E" earlier document but published on or after the International filing date.

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"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.

"R" document member of the same patent family.
### DOCUMENTS CONSIDERED TO BE RELEVANT

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