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A process for producing an ink jet recording material

Herstellungsverfahren zu einem Tintenstrahlaufzeichnungsmaterial

Méthode de fabrication d’un matériau d’enregistrement par jet d’encre

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Background of the invention

1. Field of the Invention

[0001] The present invention relates to processes for producing an ink jet recording material. More particularly, the present invention relates to processes for producing an ink jet recording material having excellent gloss and ink absorption and capable of recording ink images with excellent color density.

2. Description of the Related Art

[0002] The ink jet recording system is an ink image recording system wherein ink droplets are jetted through an ink-jetting nozzle toward a recording material at a high speed, and ink jetted droplets are absorbed in and fixed on the recording material, to form ink images. This ink jet recording system is advantageous in that full colored images can be easily formed and the printing noise is low.

[0003] The ink usable for the ink jet recording system contains a large amount of a solvent in which a coloring material is dissolved or dispersed and thus, to obtain a high color density of the recorded ink images, a large amount of the ink must be absorbed in the recording material. For the complete absorption of the ink droplets which have reached the recording material, a certain length of time is necessary. This feature of the ink jet recording system causes such a disadvantage that when the ink droplets are continuously jetted imagewise to form ink dots on the recording material surface, sometimes the ink droplets reach a target dot before an ink dot adjacent to the target dot has been completely absorbed in the recording material, and the target ink dot is connected to the adjacent ink dot so that the resultant ink image becomes unclear.

[0004] Accordingly if the recording material for the ink jet recording system is to have such an advantage that the ink dots formed thereon have high color density and brightness and a high clarity, the ink droplets must be rapidly absorbed therein and even if the ink dots are overlapped on each other, substantially no blotting of the ink may occur.

[0005] Currently, due to the rapid spread of ink jet printers, various ink jet prints with a high gloss are demanded for publications and packing paper sheets. Particularly, in colored prints, film type or coated sheet type ink jet recording sheets which have an appropriate dot form (true circle), a high dot sharpness, and high ink-receiving properties such as high ink-absorbing and fixing rates and a high ink absorption are in great demand.

[0006] To meet to the above-mentioned demands, a large number of types of ink jet recording sheets in which a coating layer containing an ink-absorbing pigment, for example, silica and alumina, and a binder is formed on a surface of a support sheet, are available. In these recording sheets, the pigments have the very small particle size of several µm, and thus the surface of the resultant recording sheet is rough and a high gloss in the resultant prints is difficult to obtain. Also, since the coating layer is quite opaque, the applied ink is easily embedded in the coating layer and thus the color density of the resultant ink images is low. Namely, the droplets of ink easily spread between the fine pigment particles over a wide area and the color density in the image decreases with an increase in the distance from the center of the image. Also, the ink-spread area becomes unnecessarily large. Therefore, the color density of the ink image becomes low as a whole and the sharpness and clearness of the image decrease. Thus, undesirable unevenness and blotting of the image occur.

[0007] To solve the above-mentioned problems, Japanese Unexamined Patent Publication No. 62-111,782 proposed an ink jet recording material in which upper and under porous coating layers, different in porosity from each other, are formed on a support sheet, to control the ink-spread and to prevent unevenness and blotting of ink images. However, since the upper and under coating layers contain pigments having a relatively large particle size, in the µm order, the resultant recording sheet was unsatisfactory in gloss and the color density of ink image.

[0008] Also, Japanese Unexamined Patent Publication No. 62-244,689 discloses an ink jet recording material in which a support is coated with an under coating layer containing a while pigment having a refraction of 1.50 or more and then with an upper coating layer containing a silicon pigment having a refraction of 1.43 to 1.48. This ink jet recording sheet can record ink images having a bright color and a high clarity. However, in this type of recording material, the pigments have a large particle size of 1 µm or more, and thus the recording material exhibits an unsatisfactory gloss and the resultant ink images have an unsatisfactory color density.

[0009] Further, Japanese Unexamined Patent Publication No. 63-104,878 discloses an ink jet recording material in which a support is coated with a two-layered recording layer. In this recording layer, the pigments contained in an upper layer and a under layer are different in particle size, and one of the upper and under coating layers contains spherical silica particles, to enhance the ink-coloring property and the dot properties, for example, the true circle form of dots. However, the ink jet recording material is still unsatisfactory in gloss and color density of ink images.

[0010] For the purpose of enhancing the gloss of ink images and the ink-absorbing property of the ink jet recording
material, currently, for example, Japanese Unexamined Patent Publication No. 7-101,142 and No. 7-117,335 disclose a two layered recording layer in which an upper coating layer serves as a high gloss-exhibiting layer. In these publications, since the high gloss-exhibiting layer contains primary particles of a pigment, the porosity of the high gloss-exhibiting layer is very low. Therefore, the high gloss-exhibiting layer has very small or substantially no space for receiving and fixing the ink, and thus can serve only as an ink-passing layer. Almost all of the applied ink can be fixed only in the coating layer. The undercoat layer contains pigment particles having a particle size of a µm order. Thus, the resultant undercoating layer has a poor transparency, and thus cannot record ink images having a high color density. Therefore, this type of ink jet recording material substantially cannot record photographic image-like ink images having a high gloss and color density.

Further, EP-A-803374 discloses an inkjet recording material in which a support is coated with an ink receiving layer containing secondary particles having an average secondary particle size of 10 to 300 nm and consisting of primary particles having an average particle size of 3 to 40 nm and agglomerated with each other. This inkjet recording material can record photographic image-like ink images having a high gloss and a high color density. However, in the current inkjet printers, since the printing speed and ink-jetting rate are greatly increased, the blotting and unevenness of the printed ink images must be further prevented.

EP 0 759 365 A discloses a process for producing an inkjet recording material comprising a support and an ink receiving layer, with an intermediate layer therebetween, in which process the ink receiving layer is formed on a casting surface; separately the intermediate layer is formed on the surface of the support; the surface of the intermediate layer not adjacent to the support is then laminated to the ink receiving layer formed on the casting surface; and the resulting laminate is peeled from the casting surface.

SUMMARY OF THE INVENTION

An object of the present invention is to provide an inkjet recording material having a high gloss, a high ink-absorbing property and a high resistance to blotting of ink and capable of recording clear ink images having excellent color density and sharpness, and processes for producing the same.

The above-mentioned object can be attained by a process for producing an inkjet recording material comprising a support and a multi-layered ink receiving layer comprising at least one underrecording layer formed on the support and at least one upperrecording layer formed on the underrecording layer, the under and upperrecording layers comprising secondary particles of a pigment having an average secondary particle size of 10 to 400 nm, and each consisting essentially of a plurality of primary particles agglomerated with each other and having an average primary particle size of 3 to 40 nm, and the average primary particle size of the primary particles of the pigment in the underrecording layer being larger than the average primary particle size of the primary particles of the pigment in the upperrecording layer, in which process at least one first coating layer corresponding to the at least one upperrecording layer is formed on a casting surface; separately at least one second coating layer corresponding to the at least one underrecording layer is formed on a surface of a support; the second coating layer-coated support is laminated on the first coating layer formed on the casting surface in such a manner that the surface of the second coating layer is brought into contact with and bonded to the surface of the first coating layer; and the resultant laminate is separated from the casting surface.

Another embodiment of the process of the present invention for producing an inkjet recording material comprises:

- forming at least one dried coating layer for said upperrecording layer on a casting surface;
- separately forming at least one coating layer for said underrecording layer on a surface of a support;
- laminating, while the surface of the coating layer formed on the support is kept in a wetted condition, the coating layer-coated support on the dried coating layer formed on the casting surface in such a manner that the wetted coating liquid layer surface on the support is brought into contact with and bonded to the dried coating layer on the casting surface;
- and separating the resultant laminate from the casting surface.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows an explanatory cross-sectional profile of another embodiment of the inkjet recording material produced according to the present invention, and Fig. 2 shows an explanatory cross-sectional profile of another embodiment of the inkjet recording material produced according to the present invention.
DESCRIPTION OF THE PREFERRED EMBODIMENTS

[0017] The inventors of the present invention have extensively studied an ink jet recording materials to solve the above-mentioned problems of the conventional ink jet recording material and have found that when a multi-layered ink receiving layer which comprises at least one underrecording layer and at least one upperrecording layer each containing secondary particles of a pigment, each consisting essentially of a plurality of primary particles agglomerated with each other, is formed on a support, the average secondary particles size of the secondary particles is controlled within the range of from 10 to 400 nm, the average primary particle size of the primary particles is controlled within the range of from 3 to 40 nm, and the average primary particle size of the primary particles in the underrecording layer is controlled to be larger than the average primary particle size of the primary particles in the upperrecording layer. In this case, an ink jet recording material having a high ink-absorbing capacity and a significantly high resistance to blotting of the ink, is obtained.

[0018] When the particle size of the primary particles, from which secondary particles of a pigment having an average secondary particle size of 10 to 400 nm are formed, is small, the resultant recording layer exhibits a high transparency and the pores formed between the particles are relatively small. Also, the specific surface area of the small particles is large, and thus the ink dye is easily fixed on the particle surfaces and a printed material having a high gloss and a high color density of printed ink images can be easily obtained. However, the small size of the primary particle causes the film-forming property of the primary particles to be low and a binder must be employed in a large amount to obtain a recording layer having a satisfactory basis weight. When the content of the binder in the recording layer is too high, the resultant recording layer exhibits a significantly decreased ink-absorbing rate and capacity. Current ink jet printers exhibit a high ink-jetting rate and capacity. Therefore, the decreased ink-absorbing rate and capacity may cause the ink droplets applied to the recording material to flow or blot to the outside of the ink dots.

[0019] To solve the above-mentioned problems, in the ink jet recording material produced according to the present invention, the ink receiving layer is formed in a two or more layered form and the average particle size of the primary particles in the underrecording layer is controlled to be larger than that in the upperrecording layer.

[0020] When the size of the primary particles, from which the secondary particles in the upper recording layer are formed, is small, the resultant ink receiving layer exhibits a high gloss and is capable of recording ink images having a high color density. When the size of the primary particles, from which the primary particles in the underrecording layer are formed, is large, the underrecording layer exhibits an enhanced film-forming property and can be formed in a large basis weight. Also, it becomes possible to control the ink absorbing rate of the underrecording layer to be higher than that of the upperrecording layer. In this case, an ink jet recording material having a high ink-absorbing capacity and a significantly high resistance to blotting of the ink, is obtained.

[0021] In the ink jet recording material, when a cationic compound is contained in the ink receiving layer, an anionic dye of the ink can be easily fixed in the ink receiving layer, and the resultant prints exhibit a significantly enhanced water resistance and durability in storage.

[0022] Further, when at least the upperrecording layer is formed on a casting surface and then transferred to a support or an underrecording layer formed on the support, the resultant ink jet recording material has a photographic paper-like high gloss.

[0023] The ink jet recording material produced according to the present invention comprises a support and a multi-layered ink-receiving layer comprising at least one underrecording layer formed on the support and at least one upperrecording layer formed on the underrecording layer, wherein, the under and upperrecording layers comprise secondary particles of a pigment having an average secondary particle size of 10 to 400 nm and each consisting essentially of a plurality of primary particles agglomerated with each other and having an average primary particle size of 3 to 40 nm, and the average primary particle size of the primary particles contained in the underrecording layer is larger than the average primary particle size of the primary particles contained in the upperrecording layer.

[0024] In the ink jet recording material produced according to the present invention, the average primary particle size of the primary particles contained in the underrecording layer may be 1.2 to 10 times the average primary particle size of the primary particles contained in the upperrecording layer.

[0025] In the ink jet recording material produced according to the present invention, the upperrecording layer optionally further comprises a cationic compound.

[0026] In the ink jet recording material produced according to the present invention, the ink-receiving layer optionally further comprises an additional recording layer comprising, as a principal component, a polymeric material. The addi-
In the ink jet recording material produced according to the present invention, the upperrecording layer may be one provided in such a manner that a coating layer corresponding to the upperrecording layer is formed on a casting surface, a underrecording layer formed on a support is superposed on and bonded to the coating layer located on the casting surface, and the resultant laminate is separated from the casting surface.

In the process for producing an ink jet recording material comprising a support and a multi-layered ink receiving layer comprising at least one underrecording layer formed on the support and at least one upperrecording layer formed on the underrecording layer, the under and upper-recording layers comprising secondary particles of a pigment having an average secondary particle size of 10 to 400 nm and each consisting essentially of a plurality of primary particles agglomerated with each other and having an average primary particle size of 3 to 40 nm, and the average primary particle size of the primary particles contained in the under recording layer is larger than the average primary particle size of the primary particles contained in the upperrecording layer, at least one first coating layer corresponding to the at least one upperrecording layer is formed on a casting surface; separately at least one second coating layer corresponding to the at least one underrecording layer is formed on a surface of a support;

the second coating layer-coated support is laminated on the first coating layer formed on the casting surface in such a manner that the surface of the second coating layer is brought into contact with and bonded to the surface of the first coating layer; and the resultant laminate is separated from the casting surface.

Another process for producing an ink jet recording material comprises:

forming at least one dried coating layer containing a binder and secondary particles of a pigment having an average secondary particle size of 10 to 400 nm and each consisting essentially of a plurality of primary particles agglomerated with each other and having an average primary particle size of 3 to 40 nm, on a casting surface; separately forming at least one coating layer containing a binder and secondary particles of a pigment having an average secondary particle size of 10 to 400 nm and each consisting essentially of a plurality of primary particles agglomerated with each other and having an average primary particle size of 3 to 40 nm and larger than the average primary particle size of the primary particles contained in the dried coating layer located on the casting surface, on a surface of a support;

laminating, while the surface of the coating layer formed on the support is kept in a wetted condition, the coating layer-coated support on the dried coating layer formed on the casting surface in such a manner that the wetted coating layer surface on the support is brought into contact with and bonded to the dried coating layer on the casting surface;

and separating the resultant laminate from the casting surface.

In the ink jet recording material produced according to the present invention, the pigment preferably comprises at least one member selected from the group consisting of amorphous silica and alumina silicate.

In the ink jet recording material produced according to the present invention, the pigment preferably comprises amorphous silica.

In the ink jet recording material produced according to the present invention, at least one recording layer other than a uppermost recording layer of the ink-receiving layer may have cracks formed to an extent such that when an ink jet dot printing is applied to the crack-formed recording layer, and printed ink dots having a diameter of 50 μm are observed through an optical microscope at a magnification of 150, a crack proportion, which is represented by a ratio in % of the number of printed dots in which cracks are formed to the total number of the printed dots, is 30% or more.

In the ink jet recording material produced according to the present invention, the uppermost recording layer may have a crack proportion smaller than that of the other recording layers.

In the ink jet recording material produced according to the present invention, the uppermost recording layer may have a pore volume of 0.2 to 3.0 ml/g.

In the ink jet recording material produced according to the present invention, at least the uppermost recording layer other than a uppermost recording layer of the ink-receiving layer may have cracks formed to an extent such that when an ink jet dot printing is applied to the crack-formed recording layer, and printed ink dots having a diameter of 50 μm are observed through an optical microscope at a magnification of 150, a crack proportion, which is represented by a ratio in % of the number of printed dots in which cracks are formed to the total number of the printed dots, is 30% or more.

The above-mentioned type of the ink jet recording material produced according to the present invention has a high gloss, exhibits an excellent ink-absorbing property and can record ink images with a high color density.

In the ink jet recording material produced according to the present invention, there is no limitation to the type, form and size of the support. The support may be transparent or opaque. For example, the support can be formed from at least one member selected from cellulose films, plastic film, for example polyethylene, polypropylene, soft polyvinyl chloride, hard polyvinyl chloride and polyester films, paper sheets, for example, woodfree paper sheets, neutral paper sheets, photographic paper support sheets, art paper sheets, coated paper sheets, cast-coated paper sheets, metallic foil-laminated paper sheets, kraft paper sheets, polyethylene-laminated paper sheets, impregnated paper sheets, metallized paper sheets and water-soluble paper sheets, metal foils and synthetic paper sheets.
The binder comprises at least one member selected from, for example, water-soluble polymeric material, for example, alcohols, cation-modified polyvinyl alcohols, casein, soybean protein, synthetic proteins, starch, and cellulose derivatives.

In the present invention, to increase the ink-absorbing property, unless the gloss and color density of the ink images are decreased.

When the average particle size of the secondary particles in the underrecording layer is larger than the average particle size of the upperrecording layer. In the present invention, the average particle size of the secondary particles for the underrecording layer are formed is larger than the average particle size of the upperrecording layer. Preferably, the average primary particle size in the underrecording layer is 1.2 to 10 times, more preferably 1.5 to 5 times, that in the upperrecording layer.

When the average primary particle size in the underrecording layer is too close to the average primary particle size in the upperrecording layer, the color density of the ink images and the ink-absorbing property of the ink-receiving layer are difficult to balance. When the average particle size of the primary particles in the underrecording layer is too large, the resultant underrecording layer exhibits a low mechanical strength and is not preferred.

In the present invention, the average particle size of the secondary particles in the underrecording layer is larger than the average particle size of the secondary particles in the upperrecording layer. In the present invention, the average particle size of the secondary particles in the upperrecording layer is in the range of from 10 to 400 nm, preferably 10 to 300 nm, more preferably 15 to 200 nm, still more preferably 20 to 150 nm. When the particle size of the secondary particles is large, the resultant secondary particles formed from the primary particles agglomerated with each other are large, the resultant recording layer exhibits a reduced transparency and thus there is a risk that the ink images having a high color density cannot be formed.

The pigment comprises at least one member selected from, for example, amorphous silica, cation-modified silica such as alumina-modified silica, alumina silicate, kaolin, clay, calcined clay, zinc oxide, tin oxides, magnesium sulfate, aluminum oxide, aluminum hydroxide, alumina, pseudo-boehmite, calcium carbonate, satin white, aluminum silicate, smectite, zeolite, magnesium silicate, magnesium carbonate, magnesia, diatomaceous earth, styrene resin plastic pigments, urea resin plastic pigments and benzoguanamine resin plastic pigment, which are commonly known and widely used in practice. However, to obtain the ink jet recording material produced according to the present invention having high gloss and ink-absorbing property and capable of recording ink images having a high color density, the pigment secondary particles having an average secondary particle size of 10 to 400 nm and constituted from a plurality of primary particles having an average primary particle size of 3 to 40 nm must be used. To obtain the secondary particle of a pigment having an average secondary particle size of 10 to 400 nm, trade pigment particle having a particle size in µm order are subjected to a high mechanical force, namely, a cracking down method or a breaking down method is applied to the pigment particles. The cracking down method is utilized to finely pulverize a lump-shaped material. The mechanical means include ultrasonic homogenizers, pressure-type homogenizers, nanomizers, high speed revolution mills, roller mills, container-driven medium mills, medium-stirring mills, jet mills, mortars, and sand grinders. The fine particles of the pigment usable for the present invention may be colloidal particles and in the state of a slurry.

The fine particles of the pigment usable for the present invention may be colloidal particles and in the state of a slurry. To attain the effect of the present invention as much as possible, the pigment is preferably selected from silica, alumina silicate, alumina (including pseudo-boehmite) and calcium carbonate pigments, more preferably amorphous silica and alumina silicate. Particularly, the amorphous silica is preferred. Unless specifically mentioned, the average particle size used in the present invention is a particle diameter (Martin size) determined by using an electron microscope (both SEM and TEM) (Asakura Shoten, "Fine Particle Handbook" page 52).

Each recording layer comprises a binder and a pigment. The under- and upperrecording layers produced according to the present invention usually contain a binder.

Conventional pigments which are commonly known and widely used in the general coated paper sheet field, and have a particle size of an µm order, can be employed together with the specific pigment particles of the present invention, to increase the ink-absorbing property, unless the gloss and color density of the ink images are decreased.

The under- and upperrecording layers produced according to the present invention usually contain a binder. The binder comprises at least one member selected from, for example, water-soluble polymeric material, for example, polyvinyl alcohol which will be referred to as PVA hereinafter, modified polyvinyl alcohols such as silyl-modified polyvinyl alcohols, cation-modified polyvinyl alcohols, casein, soybean protein, synthetic proteins, starch, and cellulose derivatives such as carboxymethyl cellulose and methylcellulose; and water-dispersible or emulsionizable polymeric mate-
In the ink-receiving layer, the cationic resin is preferably contained in an amount of 1 to 30 parts by weight, more preferably 5 to 20 parts by weight, per 100 parts by weight of the pigment. The cationic resin may be mixed into the fine pigment particle dispersion produced, for example, by the breaking down method. When the fine pigment particles are anionic, the mixing of the cationic resin with the anionic fine pigment particles results in an agglomeration of the fine pigment particles. Therefore, the mixing of the cationic resin with the anionic pigment particles is preferably carried out in such a manner that fine pigment particles having a particle size in a 1 µm order is mixed with a cationic resin, and the resultant mixture is dispersed and further mechanically pulverized and dispersed. Otherwise, secondary particles of a pigment prepared by a disperse-pulverizing procedure and having a secondary particle size of 10 to 400 nm are mixed with a cationic resin, to provide a coagulated mixture having an increased viscosity, and the coagulated mixture is mechanically pulverized and dispersed. Also, the ink-receiving layer optionally contains a conventional additive comprising at least one member selected from dispersants, thickening agents, defoaming agents, coloring materials, antistatics and preservatives.

There are no limitations to the amounts of the upper-recording layer and the underrecording layer. Preferably, the amount of the upper-recording layer is controlled to 1 to 50 g/m², more preferably 2 to 20 g/m². Also, the amount of the under-recording layer is preferably controlled to 2 to 80 g/m², more preferably 5 to 70 g/m². When the amount of the upper- or underrecording layer is small, the layer may be difficult to form uniformly. Also, when the layer amount is large, the resultant layer may have large cracks.

To increase the ink-absorbing capacity, for example, the upper-recording layer may be formed in two or more layers. Also, the ink receiving layer may have an additional recording layer comprising, as a principal component, an ink-absorbing resin, and may be located between the support and the underrecording layer.

The ink-absorbable resin may comprise at least one member selected from, for example, polyalkyleneoxides, PVA, modified polyvinyl alcohols, cellulosic derivatives, casein, gelatin, and polyvinyl pyrrolidone. Among these resins, cross-linked thermoplastic polyalkyleneoxide resins are preferably utilized. Compared with other resins, the thermoplastic polyalkyleneoxide resins are advantageous in high processability, high ink absorbing capacity, and high sharpness of resultant ink images. Particularly, the cross-linked polyalkyleneoxide resin is advantageously mixed with a polyamide resin. In this mixture, a high ink-absorbing performance derived from the cross-linked polyalkyleneoxide resin and high heat resistance and water resistance derived from the polyamide resin are not lost and a synergistic effect can be attained.

In an embodiment of the process of the ink jet recording material of the present invention, the upper-recording layer, is formed on a casting surface, and is transferred onto the underrecording layer formed on a support, and the resultant laminate is separated from the casting surface. In this process, the resultant upper-recording layer surface has a high gloss.

The casting surface can be selected from high smoothness surfaces of polymeric films, for example, regen-
erated cellulosic films, polyethylene films, polypropylene films, soft polyvinyl chloride films, hard polyvinyl chloride films, and polyester films surface-smoothed papers, for example, polyethylene laminated paper sheets, glassine paper sheets, impregnated paper sheets, and metallized paper sheets, metal foils, and thermoplastic sheets, for example, synthetic paper sheets, inorganic glass plates and surface-smoothed metallic and plastic drums and plates. In consideration of ease in production and separation of the resultant recording layer from the casting surface, a polymer film, for example, a polyethylene, polypropylene or polyester film, or a metallic drum having a high surface smoothness is used as a casting surface.

[0058] For the purpose of imparting a high gloss to the upper recording layer, the casting surface preferably has a high smoothness. Namely, the surface roughness Ra of the casting surface is preferably 0.5 µm or less, more preferably 0.05 µm or less, determined in accordance with JIS B 0601. Also, the surface of the upper recording layer may be semi-gloss or mat. The gloss or mat surface of the upper recording layer can be obtained by controlling the surface roughness Ra of the casting surface.

[0059] The casting surface may be a non-treated one. However, to control the bonding strength between the casting surface and the upper recording layer to a level lower than the bonding strength between the support and the ink receiving layer, the casting surface may be coated with a releasing agent, for example, a silicone or a fluorine-containing resin.

[0060] The bonding method between the upper recording layer formed on the casting surface and the under recording layer formed on the support, and the support is not limited to a specific method as long as they are firmly bonded to each other. The bonding method can be selected from direct bonding method in which they are directly press-bonded to each other under pressure, and indirect bonding method in which the surface of the support or of the under recording layer formed on the support is coated with an intermediate layer (including an adhesive layer, a adhesive recording layer or a tacky or adhesive ink-absorbable resin layer), and then laminated to the coating layer formed on the casting surface through the intermediate layer. Otherwise, the intermediate layer is coated on the coating layer formed on the casting surface and then, the support or the under recording layer-coated support is laminated to the coating layer on the casting surface through the intermediate layer. After the lamination is completed, the resultant laminate is separated from the casting surface.

[0061] The recording layers and intermediate layer can be formed by using conventional coating means, for example, a die coater, blade coater, air knife coater, roll coater, bar coater, gravure coater, rodblade coater, lip coater, or curtain coater.

[0062] The ink usable for the ink jet recording material produced according to the present invention comprises, as indispensable components, a coloring material for forming colored images and a liquid medium for dissolving or dispersing the coloring material, and as optional components, a dispersant; surfactant, viscosity modifier, specific resistance-modifier, pH-adjuster, mildewcide, and a solution or a dispersion-stabilizer for a recording agent.

[0063] The recording agent for the ink comprises at least one member selected from direct dyes, acid dyes, basic dyes, reactive dyes, food coloring matters, disperse dyes, oil dyes and pigments. Conventional recording agents can be used for the ink without any restriction. The content of the coloring material in the ink is established in response to the type of the liquid medium and the properties required to the coloring material. In the ink usable for the ink jet recording material of the present invention, the content of the coloring material is about 0.1 to 20% by weight which is quite usual for conventional inks.

[0064] In a preferred embodiment of the present invention, an ink jet recording material having a high surface smoothness and gloss, and a high ink-absorbing capacity and capable of recording ink images having a high color density and grade and a process for producing the same are provided.

[0065] Generally, a dispersion containing particles having a particle size of 300 nm or more causes a resultant recording layers formed therefrom to exhibit an unsatisfactory transparency and a poor surface smoothness and thus is not suitable for producing an ink jet recording material having a high gloss and capable of recording ink images with a high color density. Regarding the upper recording layer formed on a casting surface, there are no specifically severe limitations to the viscosity and particle concentration of the coating liquid for the upper recording layer. The upper recording layer having a high surface smoothness and gloss can be obtained by forming it from a coating liquid containing a binder and secondary particles of a pigment having an average secondary particle size of 10 to 400 nm, preferably 10 to 300 nm, and each consisting essentially a plurality of primary particles having an average particle size of 3 to 40 nm. When the casting surface has a high smoothness, a high smoothness of the upper recording layer surface can be obtained. Also, when the secondary pigment particles having an average secondary particle size of 10 to 400 nm are used, the resultant upper recording layer exhibit a high transparency and can record ink images having a high color density.

[0066] Also, when the upper recording layer is bonded to the under recording layer by a wet laminating method, no bubble or pore is formed in the surface portion of the upper recording layer. Further, when the upper recording layer is bonded, after drying, to the under recording layer, there is no risk of mixing the coating liquid for the upper recording layer with the coating liquid for the under recording layer. By increasing the total amount of the ink receiving layer, the
ink-absorbing capacity of the resultant ink jet recording material can be increased. The above-mentioned capability
was confirmed.

[0067] The present invention includes the following embodiments but is not limited to the embodiments.

[0068] In the ink jet recording material produced according to the present invention, preferably, the pigment particles
comprise at least one member selected from amorphous silica and alumina silicale.

[0069] In the embodiment of the ink jet recording material produced according to the present invention, the support
is preferably formed from at least one member selected from moisture-permeable (air-permeable) sheets, for example,
air-permeable polymer films, synthetic paper sheets and paper sheets. The moisture-permeable paper sheets are
preferred. For example, woodfree paper sheets, neutral paper sheets, acidic paper sheets, support sheets for photo-
graphic sheets, art paper sheets, coated paper sheets, cast-coated paper sheets, kraft paper sheets, and impregnated
paper sheets are preferably used for the support. Particularly, in view of the surface smoothness and gloss, woodfree
paper sheets, support sheets for photographic sheets, art paper sheets, coated paper sheets and cast-coated paper
sheets which have a high surface smoothness are more preferably employed for the support.

[0070] The pigment to be contained in each recording layer comprises, for example, amorphous silia (including cat-
ion-modified silica, for example, alumina-modified silica), kaolin, clay, calcined clay, zinc oxide, tin oxides, magnesium
sulfates, aluminum oxide, aluminum hydroxide, alumina, pseudo-boehmite, calcium carbonate, satin white, aluminum
silicate, smectites, zeolites, magnesium silicate, magnesium carbonate, magnesium oxide, diatomaceous earth, sty-
rene polymer plastic pigments, urea resin plastic pigments and benzoguanamine resin plastic pigments which are
widely used in the conventional coated paper sheet field.

[0071] In the upperrecording layer, fine pigment particles having, an average secondary particle size of 10 to 400
nm, preferably 10 to 300 nm and an average primary particle size of 3 to 40 nm. When the average secondary particle
size is in the range of from 10 to 400 nm, the resultant upperrecording layer exhibits a high surface smoothness and
can record ink images having a high sharpness and clarity. More preferably, the average primary particle size is 5 to
30 nm and the average secondary particle size is 20 to 200 nm.

[0072] The fine pigment particles having an average secondary particle size of 10 to 400 nm can be produced by
pulverizing and dispersing a pigment material by mechanical means, for example, a breaking down method in which
a lump-formed pigment material is finely divided. The mechanical means include an ultrasonic homogenizer, high
speed revolution mill, sand grinder and pressure-applying homogenizer.

[0073] The lower limits of the average secondary particle size is 10 nm, preferably 20 nm. Also, when the average
primary particle size is in the range of from 3 to 40 nm, the resultant upperrecording layer exhibits an excellent ink-
absorbing property.

[0074] Particularly, the pigment is preferably selected from amorphous silica (including cation-modified silica, for
example, alumina-modified silica) and alumina silicale.

[0075] In the formation of each recording layer, a binder is used to adhere the fine pigment particles to each other
and to the support.

[0076] The binder preferably comprises at least one member selected from, for example, water-soluble resins, for
example, polyvinyl alchol (PVA)), casein, soybean protein, synthetic proteins, starch, cellulose derivatives, for example,
carboxymethyl cellulose and methylcellulose, conjugated diene polymer latices, for example, styrene-butadiene co-
polymer and methyl methacrylate-butadiene copolymer latices, acrylic polymer latices and vinyl polymer latices, for
example, styrene-vinyl acetate copolymer latices, which are widely employed for the coated paper sheet production
and can be used alone in a mixture of two or more thereof.

[0077] To enhance the bonding strength between the underrecording layer and the upperrecording layer formed on
a casting surface, the surface smoothness, the ink-absorbing property, the color density of printed ink images, and
the water resistance, the binder is preferably selected from water-soluble resins, for example, PVA.

[0078] There is no specific limitation to the solid weight ratio of the pigment to the binder in each recording layer.
Preferably, the solid weight ratio (pigment/binder) is in the range of from 100/5 to 100/200, more preferably 100/10 to
100/100. When the proportion of the binder is too high, the volumes of the pores formed between the pigment particles
may be small and the ink-absorbing rate may be small. When the proportion of the binder is too low, the resultant
recording layer may exhibit a low resistance to cracking.

[0079] Each recording layer optionally comprises a cationic resin for the purpose of enhancing the ink-fixing property
of the layer. The cationic resin can be selected from those mentioned above. The content of the cationic resin is not
limited to a specific level. Usually, the cationic resin is present in an amount of 1 to 30 parts by weight, preferably 5 to
20 parts by weight, per 100 parts by weight of the pigment, in each recording layer. The recording layers optionally
contain at least one additive selected from, for example, disperses, viscosity-modifiers, anti-foaming agents, coloring
materials, antistatics, and preservatives.

[0080] There is no specific limitation to the amounts of the recording layers. The amounts of the recording layers are
preferably 1 to 100 g/m², more preferably 2 to 50 g/m². When the amount of the upperrecording layer is less than 1 g/
m², the resultant gloss and smoothness may be unsatisfactory. When the amount of the upperrecording layer is more
than 100 g/m², the resultant layer may exhibit an unsatisfactory resistance to cracking. When the amount of the under-
recording layer is less than 1 g/m², the resultant underrecording layer may be difficult to serve as a bonding layer 
between the support and the upperrecording layer and thus cannot firmly bond the support to the upperrecording layer 
by the wet lamination method. Also, when the amount of the underrecording layer is more than 100 g/m², the bonding 
procedure between the underrecording layer and the upperrecording layer under pressure may be difficult to carry out 
smoothly.

[0081] In this embodiment of the process of the present invention, a wet lamination method is utilized. The wet 
lamination method is a conventional method for laminating, for example, a paper sheet to a paper sheet or to an 
aluminum foil or to a regenerated cellulose film. In this embodiment, an aqueous solution of a water-soluble binder, 
which may be a coating liquid for forming a recording layer, is coated on a support surface (or a surface of a recording 
layer formed on the support) by a conventional coating procedure, and, while the coated binder is in a wetted condition, 
the binder-coated support is laminated to another recording layer, and then the laminate is passed through a pair of 
rolls under pressure and then dried to remove water from the binder solution. By these procedures, a support or a 
recording layer-coated support is bonded to another recording layer.

[0082] The underrecording layer in the wetted condition preferably contains water in an amount of 350 parts by weight 
or more, more preferably 500 parts by weight or more, per 100 parts by weight of the total solid of the underrecording 
layer.

[0083] In the process of the present invention, the lamination by the wet lamination method is carried out, for example, 
by bringing the underrecording layer formed on a support into contact with the upperrecording layer formed on a casting 
surface, and the resultant laminate is pressed with a pressing roll under nipping pressure to bond the underrecording 
layer to the upper-recording layer. The nipping linear pressure is preferably controlled to 1 to 200 kg/cm, more preferably 
3 to 70 kg/cm. More preferably, the nipping linear pressure is controlled to 5 to 30 kg/cm in consideration of the rela-
tionship between the coating amount and the surface gloss and smoothness.

[0084] In this embodiment, an excellent gloss and smoothness can be obtained by forming an upperrecording layer 
as defined in present claim 1 comprising at least one layer containing a binder and pigment particles having an average 
primary particle size of 3 to 40 nm and an average secondary particle size of 10 to 400 nm on a casting surface; 
transferring and bonding the resultant upperrecording layer to a recording layer formed on the support coated with an 
underrecording layer as defined in present claim 1, while the underrecording layer is in a wetted condition, by a wet 
lamination method; then drying the resultant laminate; and separating the dried laminate from the casting surface. In 
this case, in consideration of the drying property of the laminate, at least one of the casting surface and the support is 
preferably formed from a moisture-permeable material.

[0085] The wetted coating layer can be formed by a method other than the above-mentioned method in which the 
wetted coating layer is formed on the support. In the other method, the wetted coating layer is formed on a recording 
layer (upperrecording layer) formed and dried on a casting surface, a support is superposed and bonded to the wetted 
casting layer, and the resultant laminate is separated from the casting surface.

[0086] The casting surface is preferably formed from plastic polymeric films, for example, regenerated cellulose, 
polyurethane, polyethylene, polypropylene, soft polyvinyl chloride, hard polyvinyl chloride, and polyester films; paper 
sheets, for example, polyethylene-laminated paper sheets, glassine paper sheets, impregnated paper sheets and met-
alized paper sheets; metallic foils and synthetic paper sheets; and glass articles, drums and plates having a high 
surface smoothness and made from an inorganic glass, a metal or a plastic resin. To obtain a further enhanced gloss 
and smoothness, preferably, the casting surface is formed from plastic polymeric films, for example, regenerated cel-
lulose, polyethylene, polypropylene, soft polyvinyl chloride, hard polyvinyl chloride, and polyester films; and drums and 
plates made from an inorganic glass, metal or plastic resin and having a high surface smoothness. Particularly, in 
consideration of the production process and the releasing property between the casting surface and the recording 
layer, a polymeric film (for example, polyethylene, polypropylene, or polyester film) or a metallic drum having a high 
surface smoothness periphery surface is used for the casting surface.

[0087] To provide the high gloss and smoothness to the upperrecording layer surface, the casting surface preferably 
has a high smoothness. The casting surface preferably has a surface roughness Ra of 0.5 µm or less, more preferably 
0.05 µm or less, determined in accordance with JIS B 0601. However, the casting surface may be a semi-gloss surface 
or mat-like surface formed by controlling the surface roughness.

[0088] The casting surface may be not surface-treated. Alternatively, to control the bonding strength between the 
casting surface and the upperrecording layer to lower than the bonding strength between the upperrecording layer and 
the underrecording layer or between the underrecording layer and the support which may have a coating layer, the 
casting surface may be coated with a compound having a releasing property, for example, a silicon or fluorine-containing 
resin.

[0089] The recording layer may be formed by a conventional coater, for example, blade, air knife, roll, bar, gravule, 
rodblade, lip, die or curtain coater.

[0090] In the ink jet recording material produced according to the present invention, since the under and upper-
recording layers contain very fine pigment secondary particles having an average secondary particle size of 10 to 400 nm, preferably 10 to 300 nm, and consisting essentially of primary particles having an average primary particle size of 4 to 40 nm and agglomerated with each other, the ink-receiving layer exhibits a high gloss and can record ink images with a high color density.

[0091] When a cationic resin is contained in the upper-recording layer, anionic dyes can be easily fixed in the upper-recording layer and the resultant ink images exhibit an enhanced water resistance and durability in storage.

[0092] To fix the dye in the ink in the uppermost recording layer, the uppermost recording layer preferably has a pore volume of 0.2 to 3.0 ml/g, more preferably 0.5 to 2.5 ml/g. The pore volume was determined by forming the recording layer on a plastic film to prevent an influent of the support, removing the film from the recording layer by using, for example, a cutter, and measuring the pore volume of the recording layer by a mercury penetration method using Poresizer 9320 made by SHIMAZU SEISAKUSHO.

[0093] The ink usable for the ink jet recording material produced according to the present invention contains a solvent which comprises at least one member selected from water and water-soluble organic solvents, for example, alkyl alcohols having 1 to 4 carbon atoms, for example, methyl alcohol, ethyl alcohol, n-propyl alcohol, isopropyl alcohol, n-butyl alcohol, and isobutyl alcohol; ketones and ketone alcohols, for example, acetone and diacetone-alcohol; poly-alkyleneglycols, for example, polyethylene glycol and polypropylene glycol; alkylene glycols having 1 to 6 alkylene groups, for example, ethylene glycol, propylene glycol, butylene glycol, triethylene glycol, tetraethylene glycol, hexylene glycol, and diethylene glycol; amides, for example, dimethylformamide; ethers, for example, tetrahydrofuran; glycerol; and lower alkyl ethers of polyhydric alcohols, for example, ethyleneglycolmonoethyl ether, diethyleneglycolmethyl(ethyl)ether, and triethyleneglycolmonomethyl ether.

[0094] The present invention will be further illustrated by the following examples which are merely representative and do not restrict the scope of the present invention in any way.

[0095] In the examples and comparative examples, the “part” and “%” are --part by solid weight-- and --% by solid weight“, unless specifically indicated otherwise.

[0096] In the group of Reference Examples II-1 to 3 and Comparative Examples II-1 and 2, the following pigments were employed. Reference Examples II-1 to 3 are not according to the present invention.

[Silica sol II-A]

[0097] A synthetic amorphous silica (made by NIPPON SILICA INDUSTRIAL CO., LTD.) having an average primary particle size of 11 nm and an average secondary particle size of 3 µm was pulverized and dispersed by a sand grinder and then by a pressure-type homogenizer (made by SMT COMPANY, trademark: SUPER HIGH PRESSURE TYPE HOMOGENIZER GM-1) and the above-mentioned pulverizing and dispersing procedures were repeatedly carried out until the average secondary particle size reached 70 nm. A 5% aqueous dispersion of the silica pigment was obtained. No change occurred in the average primary particle size of the silica pigment.

[Silica sol II-AK]

[0098] When 100 parts of silica sol II-A was mixed with 15 parts of a trade cationic resin (trademark: SUMILASE RESIN SR-1001, made by SUMITOMO CHEMICAL CO., LTD.), the silica pigment particles agglomerated. The agglomerated pigment mixture was pulverized and dispersed by a sand grinder and then with a pressure type homogenizer (trademark: SUPER HIGH PRESSURE HOMOGENIZER GM-1, made by SMT COMPANY), and the dispersing procedures were repeatedly carried on until the average secondary particle size reached 150 nm. A 6% dispersion of the silica pigment-cation resin mixture was obtained.

[Alumina silicate fine particles II]

[0099] Isopropyl alcohol in an amount of 100g was placed in a glass reactor (separable flask) having a capacity of 2 liters and equipped with a stirrer having a diameter of 3 cm and comprising three stirring rings and a thermometer, and heated to a liquid temperature of 60°C by using an oil bath heater. While the stirrer was rotated at a rotation speed of 100 rpm to agitate the liquid in the reactor, 5g of aluminum isopropoxide (made by WAKO PURE CHEMICAL INDUSTRIES, INC.) were added, and then 1g of an acid catalyst consisting of acetic acid (made by WAKO PURE CHEMICAL INDUSTRIES, LTD.) was further added to the isopropyl alcohol. The reaction mixture was refluxed, while maintaining the refluxing temperature constant, for 24 hours.

[0100] Separately, in the same type of glass reactor as mentioned above, was charged 100g of ion-exchanged water and the charge was heated to a temperature of 60°C, and 1.8g ethyl orthosilicate (made by WAKO PURE CHEMICAL INDUSTRIES, LTD.) was added and then 1g of an acid catalyst consisting of nitric acid (made by WAKO PURE CHEMICAL INDUSTRIES, LTD.) was added to the ion-exchanged water. The mixture was refluxed for 24 hours while main-
taining the refluxing temperature constant.

[0101] The ethyl orthosilicate-nitric acid-ion-exchanged water solution was mixed with the aluminum isopropoxide-acetic acid-isopropyl alcohol solution, and the mixture was stirred and heated at a temperature of 60°C for 6 hours to prepare fine particles of alumina silicate. Then, the reaction mixture was concentrated by evaporation at a temperature of 60°C, to provide agglomerated particles of alumina silicate. In the resultant particles, the composition ratio of alumina to silica was 3:2 and the average primary particle size was 10 nm. The agglomerated particles were mixed with water and subjected to repeated pulverizing and dispersing procedures with a sand grinder and then with a ultrasonic homogenizer until the average secondary particle size reached 60 nm, to provide an 8% aqueous dispersion.

Reference Example II-1

[0102] A 7% aqueous coating liquid was prepared by mixing 100 parts of silica sol II-AK with 40 parts of a polyvinyl alcohol (trademark: PVA-135H, made by KURARAY CO., LTD.) having a degree of polymerization of 3500 and a degree of saponification of 99% or more. This coating liquid will be referred to as "coating liquid α". The coating liquid α was coated on a casting surface consisting of a surface of a PET film (trademark: LUMILAR T, made by TORAY INDUSTRIES, INC.) having a thickness of 75 µm and a surface roughness Ra of 0.02 µm, and dried to form a coating layer 2 having a weight of 30 g/m² and corresponding to an upper recording layer.

[0103] Separately, a 7% aqueous coating liquid was prepared by mixing 100 parts of synthetic amorphous silica particles (trademark: NIPSIL G-300, made by NIPPON SILICA INDUSTRIAL CO., LTD.) having an average secondary particle size of 2.3 µm and an average primary particle size of 22 nm with 40 parts of a polyvinyl alcohol (trademark: R-1130, made by KURARAY CO., LTD.) having a degree of polymerization of 1800. This aqueous coating liquid will be referred to as "coating liquid β".

[0104] The coating liquid β was coated on a surface of a trade paper sheet (trademark: OK PRINCE, made by OJI PAPER CO., LTD., pulp composition: L wood pulp 100%, freeness: 430 ml) having a basis weight of 127.9 g/m² to form a coating layer 1 corresponding an underrecording layer and having a dry solid weight of 10 g/m².

[0105] The coating layer 1 was, while the coating layer 1 was kept in a wetted condition, laminated and bonded to the coating layer 2, and dried. The resultant dry laminate was separated from the casting PET film surface.

[0106] An ink jet recording material obtainable by the present invention was obtained.

Reference Example II-2

[0107] An ink jet recording material of the present invention was produced by the same procedures as in Example II-1, except that silica sol II-AK was replaced by silica sol II-A.

Reference Example II-3

[0108] An ink jet recording material of the present invention was produced by the same procedures as in Example II-1, except that silica sol II-AK was replaced by alumina silicate fine particles II.

Comparative Example II-1

[0109] A comparative ink jet recording material was produced by coating the same coating liquid α as in Example II-1 on a trade paper sheet (trademark: OK PRINCE, made by OJI PAPER CO., LTD., pulp composition: L wood pulp 100%, freeness: 430 ml) having a basis weight of 127.9 g/m² and drying, to form an ink-receiving layer having a weight of 30 g/m².

Comparative Example II-2

[0110] A comparative ink jet recording material was prepared by the same procedures as in Comparative Example II-1, except that the coating liquid α was replaced by the same coating liquid β as in Example II-1.

[Evaluation]

[0111] The gloss of ink images, smoothness of ink receiving layer, ink-absorbing property, water resistance of ink images and color density of ink images of each of the ink jet recording materials of Examples II-1 to II-3 and Comparative Examples II-1 and II-2 were tested and evaluated by the methods shown below.

[0112] The image gloss, ink absorption, ink water resistance and image color density were tested and evaluated on the solid images printed by a trade ink jet printer (trademark: PM-700C, made by EPSON CORP., LTD.)
The gloss of image was determined by observing the solid image at an inclined angle of 20 degrees by the naked eye, into the following four classes.

<table>
<thead>
<tr>
<th>class</th>
<th>Gloss</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>High gloss similar to the gloss of photographic colored images</td>
</tr>
<tr>
<td>3</td>
<td>Satisfactory gloss which is slightly lower than the gloss of photographic colored images</td>
</tr>
<tr>
<td>2</td>
<td>Surface is roughened and smoothness is unsatisfactory</td>
</tr>
<tr>
<td>1</td>
<td>Surface is significantly roughened and appearance is very bad</td>
</tr>
</tbody>
</table>

The smoothness of an ink receiving layer was evaluated by naked eye observation of the surface of the coating layer, into the following three classes.

<table>
<thead>
<tr>
<th>class</th>
<th>Surface condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>Surface roughness is small. Smoothness is high.</td>
</tr>
<tr>
<td>2</td>
<td>Surface is roughened. Smoothness is low.</td>
</tr>
<tr>
<td>1</td>
<td>Surface is severely roughened. Appearance is bad</td>
</tr>
</tbody>
</table>

A sample of an ink jet recording material having dimensions of 10 cm x 10 cm was attached to a center portion of a A4 size PPC paper sheet, a black ink was extruded in an amount of 30 g/m² onto the sample to form a solid print. The ink absorbing condition on the sample was observed. Also, a woodfree paper sheet was superposed on the ink-applied sample to determine an ink drying time after which time was passed, no ink was transferred from the sample to the woodfree paper sheet.

The ink absorbing property was evaluated into the following four classes.

<table>
<thead>
<tr>
<th>class</th>
<th>In-drying time</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Less than one minute</td>
</tr>
<tr>
<td>3</td>
<td>One minute or more but less than 2 minutes</td>
</tr>
<tr>
<td>2</td>
<td>Two minutes or more but less than 5 minutes</td>
</tr>
<tr>
<td>1</td>
<td>Five minutes or more</td>
</tr>
</tbody>
</table>

An ink jet recording material was printed with an ink and left to stand for 24 hours. Thereafter, a water drop was placed on the ink image and removed by wiping one minute after the water dropping.

The condition of the water-wetted portion was evaluated by naked eye into the following four classes.

<table>
<thead>
<tr>
<th>class</th>
<th>Water resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>No blotting of ink is found</td>
</tr>
<tr>
<td>3</td>
<td>Substantially no ink blotting is found</td>
</tr>
<tr>
<td>2</td>
<td>Ink is slightly blotted. Usable in practice</td>
</tr>
<tr>
<td>1</td>
<td>Ink is blotted Practical use is difficult</td>
</tr>
</tbody>
</table>

A black-colored ink solid print on an ink jet recording material was subjected to a measurement of color density by a MACBETH REFLECTION COLOR DENSITY TESTER (RD-920). The color density of the image was shown by an average of five measurement data.
The test results are shown in Table 2.

<table>
<thead>
<tr>
<th>Example No.</th>
<th>Reference Example</th>
<th>Comparative Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>II-1</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>II-2</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>II-3</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>

Table 2 shows that the above ink jet recording materials had excellent image gloss, ink-absorbing property, color density of image, satisfactory smoothness of ink receiving layer, and water resistance of image.

In the group of Examples III-3 and III-4 and Comparative Examples III-1 and III-2, the cracks formed in the ink receiving layer were checked in the following manner.

For each ink jet recording material, ten samples were collected and each was printed with 100 ink dots per mm$^2$. The dots were evenly distributed, each of the dots had a diameter of 50 µm, and the dots were produced by a trade ink jet printer (model: PM-700C made by EPSON CORP., LTD.) having a dot density of 720 dots/2.54 cm$^2$. The printed ink dots were observed by an optical microscope at a magnification of 150. When cracks were found in the dots in the number corresponding to 60% or more of the total number of the dots printed in the 10 samples, the ink jet recording material was recognized to have sufficient cracks formed in at least one recording layer.

When, for example, cracks were formed in the dots in the number corresponding to 50% of the total number of the dots printed in the 10 samples, the degree of cracks of the ink jet recording material is 50%.

In Examples III-3 and III-4 and Comparative Examples III-1 and III-2, the following pigments were employed.

**Silica sol III-A**

- Synthetic amorphous silica particles (trademark: NIPSIL HD-2, made by NIPPON SILICA INDUSTRIAL CO., LTD.) having an average secondary particle size of 3 µm and an average primary particle size of 11 nm were pulverized and dispersed by a sand grinder and then by a pressure type homogenizer, and pulverizing and dispersing procedures with a sand grinder and then with a pressure type homogenizer were repeatedly applied to the silica particles until the average secondary particle size reached 60 nm. A 7% silica particle dispersion was prepared.

**Silica sol III-B**

- Synthetic amorphous silica particles (trademark: NIPSIL LP, made by NIPPON SILICA INDUSTRIAL CO., LTD.) having an average secondary particle size of 3 µm and an average primary particle size of 16 nm were pulverized and dispersed by a sand grinder and then by a pressure type homogenizer, and pulverizing and dispersing procedures with a sand grinder and then with a pressure type homogenizer were repeatedly applied to the silica particles until the average secondary particle size reached 100 nm. A 9% silica particle dispersion was prepared.

**Silica sol III-C**

- Synthetic amorphous silica particles (trademark: NIPSIL E-1011, made by NIPPON SILICA INDUSTRIAL CO., LTD.) having an average secondary particle size of 1.5 µm and an average primary particle size of 24 nm were pulverized and dispersed by a sand grinder and then by a pressure type homogenizer, and pulverizing and dispersing procedures with a sand grinder and then with a pressure type homogenizer were repeatedly applied to the silica particles until the average secondary particle size reached 150 nm. A 12% silica particle dispersion was prepared.
Comparative Example III-1

[0128] As a support, a laminated paper sheet prepared by laminating polyethylene layer, at a thickness of 25 µm by an extrusion-lamination method onto a trade coated paper sheet (trademark: OK COAT, made by OJI PAPER CO., LTD.) having a basis weight of 127.9 g/m², was employed. This laminated paper sheet will be referred to as a "lamine sheet" hereinafter.

[0129] A 10% aqueous coating liquid comprising a mixture of 100 parts of silica sol III-C with 35 parts of a polyvinyl alcohol (trademark: PVA-135H, made by KURARAY CO., LTD.) having a degree of polymerization of 3500 and a degree of saponification of 99% or more, was coated on the polyethylene-coated surface of the laminate sheet and dried at a temperature of 120°C, to form an underrecording layer having a dry weight of 20 g/m². The resultant underrecording layer had a degree of cracks of 80%.

[0130] The underrecording layer was wetted with water in an amount of 20 g/m², and then coated with a 7% aqueous coating liquid prepared from a mixture of 100 parts of silica sol III-A with 30 parts of a PVA (PVA-135H), and dried at a temperature of 50°C to form an upperrecording layer having a degree of cracks of 30% and a pore volume of 0.9 ml/g.

Comparative Example III-2

[0131] Silica sol III-A in an amount of 100 parts was mixed with 15 parts of cationic resin consisting of diallyl-dimethyl ammonium chloride-acrylamide copolymer (trademark: PAS-J-81, made by NITTO BOSEKI CO., LTD.). The resultant mixture exhibited an increased viscosity and silica particles were agglomerated. The mixture was pulverized and dispersed by a pressure type homogenizer until the average agglomerated particle size reached 100 nm. Then, the resultant cationic resin-containing silica sol was further mixed with 20 parts of a polyvinyl alcohol (PVA-135H), to provide an 8% aqueous coating liquid.

[0132] The same support as in Comparative Example III-1 was coated with the same silica sol III-C-containing underrecording layer as in Comparative Example III-1. After wetting the underrecording layer with water in an amount of 20 g/m², the above-mentioned coating liquid was coated on the water-wetted underrecording layer, and dried at a temperature of 50°C, to form an upperrecording layer having a dry weight of 8 g/m², a degree of cracks of 30%, and a pore volume of 0.83 ml/g.

Example III-3

[0133] The same diallyl-dimethyl ammonium chloride-acrylamide copolymer-containing silica sol III-A as in Comparative Example III-2 and in amount of 115 parts was mixed with 20 parts of a polyvinyl alcohol (PVA-135H), to provide an aqueous coating liquid having a solid content of 8%.

[0134] The coating liquid was coated on a surface of a PET film (trademark: LUMILAR T, made by TORAY INDUSTRIES INC.) having a thickness of 50 µm and a surface roughness Ra of 0.02 µm, used as a casting surface, and dried at a temperature of 40°C, to form a coating layer corresponding to an upperrecording layer having a weight of 8 g/m² and a degree of cracks of 10% or less.

[0135] Separately, a 10% aqueous coating liquid containing a mixture of 100 parts of silica sol III-C with 30 parts of a polyvinyl alcohol (PVA-135H) was coated on a surface of a trade woodfree paper sheet having a basis weight of 157 g/m², and dried at a temperature of 120°C, to form an underrecording layer having a weight of 20 g/m², and a degree of cracks of 90%.

[0136] After the cationic resin-containing coating layer formed on the PET film was wetted with 5 g/m² of water, the underrecording layer was laminated and bonded to the wetted coating layer at a calendering temperature of 75°C under a calendering linear pressure of 20 kg/cm. After drying, the resultant laminate was separated from the PET film. An ink jet recording material obtainable by the present invention was obtained. In the recording material, the upperrecording layer had a pore volume of 0.76 ml/g.

Example III-4

[0137] The same diallyl-dimethyl ammonium chloride-acrylamide copolymer-containing silica sol III-A as in Comparative Example III-2 and in amount of 115 parts was mixed with 20 parts of a polyvinyl alcohol (PVA-135H), to provide an aqueous coating liquid having a solid content of 8%.

[0138] The coating liquid was coated on a surface of a PET film (trademark: LUMILAR T, made by TORAY INDUSTRIES INC.) having a thickness of 50 µm, used as a casting surface, and dried at a temperature of 40°C to form a coating layer corresponding to an upperrecording layer having a weight of 8 g/m² and a degree of cracks of 10% or less.

[0139] The coating layer was coated with a 10% aqueous coating liquid containing a mixture of 100 parts of silica sol III-B with 35 parts of a polyvinyl alcohol (PVA-135H) and dried at a temperature of 80°C to form an intermediate
recording layer having a dry weight of 10 g/m² and a degree of cracks of 20%.

[0140] Separately, a 10% aqueous coating liquid containing a mixture of 100 parts of silica sol III-C with 25 parts of a polyvinyl alcohol (PVA-135H), was coated on a surface of a trade woodfree paper sheet having a basis weight of 157 g/m², to form a coating liquid layer in a dry amount of 10 g/m².

[0141] The coating liquid layer was superposed on and bonded to the intermediate recording layer, and dried at a temperature of 130°C, to form an underrecording layer. The resultant laminate was separated from the PET film. An ink jet recording material was obtained by the present invention.

[0142] In the obtained recording material, the silica sol III-C-containing underrecording layer had a degree of cracks of 95% which was measured by the method as mentioned above, after the upperrecording layer and the intermediate recording layer were removed by shaving and printed with the ink. The upperrecording layer had a pore volume of 0.76 ml/g.

[Evaluation]

[0143] Each of the ink jet recording materials of Examples III-3 and III-4 and Comparative Examples III-1 and III-2 was subjected to evaluations of ink-absorbing rate, ink-absorbing capacity, gloss, and color density of ink image by the methods as shown below. In the evaluations of the gloss and color density of image, solid ink prints were formed by a trade ink jet printer (model: PM-700C, made by EPSON CORP., LTD.), and the gloss and color density of the solid prints were determined.

[Ink-absorbing rate]

[0144] An yellow-colored ink, a magenta-colored ink and a cyan-colored ink were printed and overlapped on each other to form a black-colored solid print on an ink jet recording material. Every five seconds after the printing, a woodfree paper sheet was superposed on the solid print, to determine an ink-drying time after which time no ink was transferred from the solid print to the paper sheet. The ink-absorbing rate was evaluated in four classes as shown below.

<table>
<thead>
<tr>
<th>class</th>
<th>Ink drying time</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Less than 5 seconds</td>
</tr>
<tr>
<td>3</td>
<td>Five seconds or more but less than 10 seconds</td>
</tr>
<tr>
<td>2</td>
<td>Ten seconds or more but less than 30 seconds</td>
</tr>
<tr>
<td>1</td>
<td>Thirty seconds or more</td>
</tr>
</tbody>
</table>

[Ink-absorbing capacity]

[0145] A sample of an ink jet recording material in dimensions of 10 cm × 10 cm was attached to a center portion of an A4 size PPC paper sheet, and a black-colored ink was jetted toward the sample at an ink-extrusion amount of 40 g/m² to form a black-colored solid print. Every 10 seconds after the printing, a woodfree paper sheet was superposed on the solid print on the sample, and an ink-drying time after which time no ink was transferred from the solid print from the woodfree paper sheet, was determined.

[0146] The ink-absorbing capacity was evaluated in the four classes as shown below.

<table>
<thead>
<tr>
<th>class</th>
<th>Ink-drying time</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Less than 10 seconds</td>
</tr>
<tr>
<td>3</td>
<td>Ten seconds or more but less than 30 seconds</td>
</tr>
<tr>
<td>2</td>
<td>Thirty seconds or more but less than one minute</td>
</tr>
<tr>
<td>1</td>
<td>One minute or more</td>
</tr>
</tbody>
</table>

[Gloss (luster) of ink image]

[0147] Black-colored images, yellow-colored images, cyan-colored images and magenta-colored images formed on an ink jet recording material were observed by the naked eye at an inclined angle and gloss of the images was evaluated in the following four classes.
A black-colored solid print formed on an ink jet recording material was subjected to a color density measurement by MACBETH REFLECTION COLOR DENSITY TESTER (model RD-920). The color density was shown by an average of five measurement data.

An ink-printed sample was stored in a constant temperature, constant humidity vessel at a temperature of 40°C at a humidity of 90% for 48 hours. Then, the blotting of ink on the sample was checked and evaluated in the following four classes.

On a sample of an ink jet recording material, a black-colored solid print was formed by overlapping-yellow, cyan and magenta-colored inks, and a corrugation of the solid printed sample was checked and evaluated in the following three classes.

Table 3 shows that the ink jet recording materials produced in accordance with the present invention exhibited excellent ink-absorbing property and satisfactory gloss and color density of an ink image.
Claims

1. A process for producing an ink jet recording material comprising a support and a multi-layered ink receiving layer comprising at least one underrecording layer formed on the support and at least one upperrecording layer formed on the underrecording layer, the under and upperrecording layers comprising secondary particles of a pigment having an average secondary particle size of 10 to 400 nm, and each consisting essentially of a plurality of primary particles agglomerated with each other and having an average primary particle size of 3 to 40 nm, and the average primary particle size of the primary particles of the pigment in the underrecording layer being, larger than the average primary particle size of the primary particles of the pigment in the upperrecording layer.

in which process at least one first coating layer corresponding to the at least one upperrecording layer is formed on a casting surface; separately at least one second coating layer corresponding to the at least one underrecording layer is formed on a surface of the support; the second coating layer-coated support is laminated on the first coating layer formed on the casting surface in such a manner that the surface of the second coating layer is brought into contact with and bonded to the surface of the first coating layer; and the resultant laminate is separated from the casting surface.

2. The process according to claim 1 comprising:

forming at least one dried coating layer for said upperrecording layer on a casting surface;

separately forming at least one coating layer for said underrecording layer on a surface of a support;

laminating, while the surface of the coating layer formed on the support is kept in a wetted condition, the coating layer-coated support on the dried coating layer formed on the casting surface in such a manner that the wetted coating liquid layer surface on the support is brought into contact with and bonded to the dried coating layer on the casting surface;

and separating the resultant laminate from the casting surface.

Patentansprüche

1. Verfahren zur Herstellung eines Tintenstrahlaufzeichnungsmaterials, das einen Träger und eine mehrlagige Tintenaufnahmeschicht umfaßt, die mindestens eine untere Aufzeichnungsschicht, die auf dem Träger ausgebildet ist, und mindestens eine obere Aufzeichnungsschicht, die auf der unteren Aufzeichnungsschicht ausgebildet ist umfaßt, wobei die untere und obere Aufzeichnungsschicht Sekundärteilchen eines Pigments umfassen, die eine mittlere Sekundärteilchengröße von 10 bis 400 nm aufweisen und jeweils aus im wesentlichen einer Vielzahl an Primärteilchen bestehen, die miteinander agglomeriert sind und eine mittlere Primärteilchengröße von 3 bis 40 nm aufweisen, und wobei die mittlere Primärteilchengröße der Primärteilchen des Pigments in der unteren Aufzeichnungsschicht größer ist als die mittlere Primärteilchengröße der Primärteilchen des Pigments in der oberen Aufzeichnungsschicht,

wobei in dem Verfahren mindestens eine erste Beschichtungsschicht, die der mindestens einen oberen Aufzeichnungsschicht entspricht, auf einer Formgebungsoberfläche ausgebildet wird; getrennt mindestens eine zweite Beschichtungsschicht, die der mindestens einen unteren Aufzeichnungsschicht entspricht, auf einer Oberfläche des Trägers ausgebildet wird; der mit der zweiten Beschichtungsschicht beschichtete Träger auf die erste Beschichtungsschicht, die auf der Formgebungsoberfläche ausgebildet ist, auf eine solche Weise laminiert wird, daß die Oberfläche der zweiten Beschichtungsschicht in Kontakt mit der Oberfläche der ersten Beschichtungsschicht gebracht und daran gebunden wird; und das resultierende Laminat von der Formgebungsoberfläche getrennt wird.

2. Verfahren nach Anspruch 1, umfassend:

Ausbilden von mindestens einer getrockneten Beschichtungsschicht für die obere Aufzeichnungsschicht auf einer Formgebungsoberfläche;

getrenntes Ausbilden von mindestens einer Beschichtungsschicht für die untere Aufzeichnungsschicht auf einer Oberfläche eines Trägers;

Laminieren des mit einer Beschichtungsschicht beschichteten Trägers, während die Oberfläche der auf dem
Träger ausgebildeten Beschichtungsschicht in einem nassen Zustand gehalten wird, auf die getrocknete, auf der Formgebungsoberfläche ausgebildete Beschichtungsschicht, auf eine solche Weise, daß die Oberfläche der nassen Beschichtungsfließigkeitsschicht auf dem Träger mit der getrockneten Beschichtungsschicht auf der Formgebungsoberfläche in Kontakt gebracht und daran gebunden wird;

und Trennen des resultierenden Laminats von der Formgebungsoberfläche.

Revendications

1. Procédé de fabrication d'une matière d'enregistrement à jet d'encre comportant un support et une couche de réception d'encre multicouche comportant au moins une couche en dessous de l'enregistrement formée sur le support et au moins une couche au-dessus de l'enregistrement formée sur la couche en dessous de l'enregistrement, les couches en dessous et au-dessus de l'enregistrement comportant des particules secondaires d'un pigment ayant une taille moyenne de particule secondaire de 10 à 400 nm, et chacune se composant essentiellement d'une multiplicité de particules principales agglomérées les unes aux autres et ayant une taille moyenne de particule principale de 3 à 40 nm, et la taille moyenne de particule principale des particules principales du pigment dans la couche en dessous de l'enregistrement étant plus grande que la taille moyenne de particule principale des particules principales du pigment dans la couche au-dessus de l'enregistrement,
dans lequel procédé au moins une première couche de revêtement correspondant à la au moins une couche au-dessus de l'enregistrement est formée sur une surface de couche brillant; au moins une deuxième couche de revêtement correspondant à la au moins une couche en dessous de l'enregistrement est formée séparément sur une surface du support; le support revêtu de la deuxième couche de revêtement est stratifié sur la première couche de revêtement formée sur la surface de couche brillant d'une manière telle que la surface de la deuxième couche de revêtement est amenée en contact avec et collée sur la surface de la première couche de revêtement; et le stratifié résultant est séparé de la surface de couche brillant.

2. Procédé selon la revendication 1, comportant le fait de :

former au moins une couche de revêtement séchée pour ladite couche au-dessus de l'enregistrement sur une surface de couche brillant;
former séparément au moins une couche de revêtement pour ladite couche en dessous de l'enregistrement sur une surface d'un support;
stratifier, alors que la surface de la couche de revêtement formée sur le support est maintenue dans une condition humide, le support revêtu de la couche de revêtement sur la couche de revêtement séchée formée sur la surface de couche brillant d'une manière telle que la surface de couche de liquide de revêtement humide sur le support est amenée en contact avec et collée sur la couche de revêtement séchée de la surface de couche brillant;
et séparer le stratifié résultant de la surface de couche.