

# (54) METHOD FOR PRODUCING A METAL PARTICLE

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- Prior Publication Data

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# Related U.S. Application Data

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# (21) Appl. No.: 14/644,579 FOREIGN PATENT DOCUMENTS



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# ( 57 ) ABSTRACT

A method for producing a metal particle which includes the steps of: mixing a metal salt and a polycarboxylic acid in a liquid phase; adding a reducing agent to the resultant mixture to deposit metal particles; and drying the deposited metal particles.

# 5 Claims, 8 Drawing Sheets



 $\begin{array}{c} {\rm JP} \\ {\rm JP} \\ {\rm JP} \end{array}$ 

 $\overline{JP}$  $\frac{J}{JP}$  $JP$  $\rm JP$  $\rm{JP}$ 

 $J\bar{P}$  $\rm{JP}$  $\rm{JP}$  $JP$  $\rm{JP}$ 

JP JP UP



CPC .................. **B22F 9/18** (2013.01); **C22C 1/02**  $(2013.01)$ ; C22C  $5/02$  (2013.01); C22C  $5/04$  $(2013.01)$ ; C22C 5/06 (2013.01); C22C 9/00 (2013.01); C22C 19/03 (2013.01); H01B 1/02 (2013.01); H01B 1/22 (2013.01); Y10T 428/12  $(2015.01)$  $(52)$ 

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Fig. 6







Fig. 8











Fig . 11







Fig. 13



PRIOR ART

Fig . 14



PRIOR ART

Fig . 15



Magnification: 20,000					
Magnification: 2,000					
Magnification: 5,000					
Magnification: 10,000					
Measured value	$\begin{bmatrix} \text{Particle size} \\ \text{distribution} (\mu n) \end{bmatrix}$ $\cos/9$ dej BET(m/g) 0.47 $\overline{0.67}$ 0.99 K Value 5.88 145 S, $\frac{10}{050}$ <b>D90</b>	Particle size distribution (um) $\cos\theta$ del 4.29 $\overline{\mathsf{BET}(\mathsf{m}'\hspace{-0.2pt}/_{\hspace{-0.1em}\mathsf{g}})}$ $\frac{1}{2}$ $\frac{33}{2}$ <b>K</b> Value 3.19 $\frac{5.62}{ }$ 2.82 $\overline{510}$ D90 <b>D50</b>	$\frac{\text{Particle size}}{\text{distribution}(\mu\text{m})}$ $\frac{1}{2}$ (g/cc) 10.17 $\overline{\text{BET}(\text{m/g})}$ 5.84 7.97 <b>K</b> Value 2.09 4.48 $\overline{3.43}$ $\frac{100}{100}$		
	Example 2	Example 3	Example 4		

Fig. 16

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9,186,727), which is a United States national phase appli-<br>cation of International application PCT/JP2011/075508 filed  $^{10}$  The metal particle described in patent document 3 is also<br>Nov. 11, 2011. The entire contents of Nov. 11, 2011. The entire contents of each of application Ser. formed from dendritic crystals which have grown from a<br>No. 13/884,013 and PCT/JP20111075508 are incorporated nucleating agent as a center, and hence inevitably No. 13/884,013 and PCT/JP20111075508 are incorporated nucleating agent as a center, and hence inevitably requires a<br>by reference herein.

The present invention relates to a metal particle which is<br>a non-nucleated, spherical porous material having continu-<br>a non-nucleating agent; however, the powder has<br>an a non-nucleating continuation of the particle of the ous open pores, and a method for producing the same. More requires no nucleating agent, nowever, the powder has<br>noticularly the present invention is concerned with a motel 20 dendritic portions formed from needle-like crys particularly, the present invention is concerned with a metal 20 dendritic portions formed from needle-like crystals which<br>needle formed from dendritic crystals which have grown particle formed from dendritic crystals which have grown have thinly grown, and therefore the thin and needle-like<br>uniformly outward from the center without requiring a dendritic portions are likely to tangle, causing aggr uniformly outward from the center without requiring a<br>nucleating agent so that the metal particle has a fine uneven of the silver powder. Further, the silver powder has the nucleating agent so that the metal particle has a fine uneven of the silver powder. Further, the silver powder has the structure in the spherical surface, and a method for produc-<br>dendritic portions formed from needle-like structure in the spherical surface, and a method for producing the same. <sup>1</sup> <sup>25</sup> have thinly grown, and therefore has a relatively sparse

der obtained by allowing dendritic crystals of silver or  $30$  particle is unlikely to suffer bonding or aggregation of the copper to grow on an electrode plate by an electrolytic metal particles and exhibits excellent dis copper to grow on an electrode plate by an electrolytic metal particles and exhibits excellent dispersibility, but also method (patent document 1). There have also been known a in that the metal particle has an appropriate method (patent document 1). There have also been known a metal particle obtained by allowing dendritic crystals of silver or copper to grow from a nucleating agent as a center to the specific surface area, and a method for producing the<br>by an electroless method so that the metal particle has <sup>35</sup> same. An object of the present inventio by an electroless method so that the metal particle has <sup>35</sup> same. An object of the present invention is to provide a metal radially extending protrusions and depressions between the particle which is advantageous in that radially extending protrusions and depressions between the protrusions (patent document 2), and a metal particle having conductive composition, such as a conductive paste, the a plurality of protrusions which protrude like a chestnut bur<br>composition can be cured at a relatively lo (patent document 3). Further, a dendritic silver powder (for example, at 120 to 200 $^{\circ}$  C.), and the obtained cured obtained by an electroless wet process has been known  $^{40}$  product can exhibit satisfactory conductio obtained by an electroless wet process has been known  $40$  (patent document 4).

## PRIOR ART REFERENCES

Patent document 1: Japanese Unexamined Patent Publication No. 2007-204795

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However, the above-mentioned fine silver powder  $\omega$  particle which is a non-nucleated described in patent document 1 is obtained by a method in having continuous open pores. which silver particles deposited on an electrode plate by an The present invention is directed to the metal particle electrolytic method are removed from the electrode plate by which has a volume cumulative particle diame dendritic silver powder. Therefore, the dendrite growth is 65 relatively non-uniform, making it difficult to obtain a true spherical fine silver powder. In addition, the resultant silver

**METHOD FOR PRODUCING A METAL** powder has a small tap density, and therefore it is difficult to<br>**PARTICLE** form a uniform sintered film from the silver powder.

Pharticle described in patent document 2 is<br>CROSS-REFERENCE TO RELATED formed from dendritic crystals which have grown from a EFERENCE TO RELATED formed from dendritic crystals which have grown from a<br>APPLICATIONS  $\frac{5}{100}$  nucleating agent as a center, and hence inevitably requires a nucleating agent as a center, and hence inevitably requires a nucleating agent, and the obtained metal particle has a relatively sparse structure such that the content of voids due This application is a Divisional application of application relatively sparse structure such that the content of voids due<br>Ser. No. 13/884,013 filed May 8, 2013 (U.S. Pat. No. to depressions is preferably more than 40% by

> nucleating agent, and the obtained metal particle has a FIELD OF THE INVENTION  $15$  number of protrusions in a chestnut bur form, and therefore the protrusions in a chestnut bur form are likely to tangle,

> > structure such that the tap density is as small as  $0.4$  to  $0.7$

BACKGROUND ART  $g/cm<sup>3</sup>$ . An object of the present invention is to provide a metal<br>there has been known a fine silver pow-<br>particle which is advantageous not only in that the metal Conventionally, there has been known a fine silver pow-<br>
r obtained by allowing dendritic crystals of silver or  $30$  particle is unlikely to suffer bonding or aggregation of the a large specific surface area and the density is large relative to the specific surface area, and a method for producing the composition can be cured at a relatively low temperature ing it possible to easily control the specific gravity or resistance, and a method for producing the same.

### Means to Solve the Problems

Patent Documents<br>
The present invention for solving the above problems is a<br>
Japanese Unexamined Patent Publica-<br>
The present invention for solving the above problems is a dvanta-<br>
The present invention for solving the abo tion No. 2007-204795<br>Patent document 2: Japanese Unexamined Patent Publica-<br>Patent document 2: Japanese Unexamined Patent Publica-<br>bonding or aggregation of the metal particles and exhibits bonding or aggregation of the metal particles and exhibits tion No. 2004-149903<br>tent document 3: Japanese Unexamined Patent Publica-<br>an appropriate tap density and a large specific surface area Patent document 3: Japanese Unexamined Patent Publica - an appropriate tap density and a large specific surface area and the density is large relative to the specific surface area. Patent document 4: Japanese Unexamined Patent Publica-<br>tion No. 2005-146387<br>such as a conductive paste, the composition can be cured at 55 a relatively low temperature (for example, at  $120$  to  $200^{\circ}$  C.), and the obtained cured product can exhibit satisfactory SUMMARY OF THE INVENTION C.), and the obtained cured product can exhibit satisfactory<br>conduction properties, making it possible to easily control<br>Problems to be Solved by the Invention the specific gravity or resistance.

Accordingly, the present invention is directed to a metal particle which is a non-nucleated, spherical porous material

to 15  $\mu$ m as measured by a particle size distribution measurement method using image analysis, a tap density of 1 to 6 g/cm<sup>3</sup>, or a specific surface area of 0.25 to 8 m<sup>2</sup>/g as measured by a BET method.

wherein the value K determined from a specific surface area particle obtained by the above - SS and a specific surface area BS and represented by the method particle. general formula (2) below satisfies the relationship:<br> $3 \le K \le 72$  wherein the specific surface area SS is represented <sup>5</sup> Effect of the Invention  $3 \le K \le 72$ , wherein the specific surface area SS is represented  $5$ by the formula (1) below wherein particle diameter d is a volume cumulative particle diameter  $D_{50}$  as measured by a The present invention is a metal particle which is a narticle size distribution measurement method using image non-nucleated, substantially true spherical porou particle size distribution measurement method using image non-nucleated, substantially true spherical porous material<br>analysis and 0 is a theoretical density of the metal particle having continuous open pores, and comprise analysis and  $\rho$  is a theoretical density of the metal particle, having continuous open pores, and comprises a metal par-<br>and the specific surface area BS is a specific surface area as  $10$  ticle formed from dendritic cr

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SS=6/\rho d \tag{1}
$$

$$
(SS/BS) \times 100 = K \tag{2}
$$

of a scanning electron microscope, magnified 20,000 times,<br>ties can be obtained, making it possible to easily control the<br>the morphology of the appearance of the metal particle has<br>specific pravity or resistance, and a met the morphology of the appearance of the metal particle has specific gravity or resistance, and a method for producing an aegagropila form. The present invention is directed to the the same. an aegagropila form. The present invention is directed to the the same.<br>
metal particle, wherein, in the image of the metal particle Further, in the present invention, a metal particle which is<br>
taken by means of a scannin

which is selected from the group consisting of silver, copper, surface.<br>gold, nickel, and palladium.<br>Further, the present invention is directed to a conductive 40 BRIEF DESCRIPTION OF THE DRAWINGS

composition comprising the metal particle which is a non nucleated, spherical porous material having continuous open FIG. 1 A SEM photomicrograph of the cross-section of pores, and a resin, a conductor comprising a cured product the metal (silver) particle of the present inventi obtained by curing the conductive composition, and an 20,000 times.<br>electronic part having the conductor.<br>The present invention is directed to a method for produc-<br>the metal (silver) particle of the present invention, magn

ing a metal particle, which comprises the steps of: mixing a<br>metal salt and a polycarboxylic acid in a liquid phase; adding FIG. 3 A SEM photomicrograph of the metal (silver)<br>a reducing agent to the resultant mixture to de

producing a metal particle, wherein the temperature for the FIG. 5 A SEM photomicrograph of the metal (silver) mixing step and the depositing step is 10 to 30° C., and the particle of the present invention magnified 40,00

The present invention is directed to the method for 55 particle of the present invention magnified 5,000 times.<br>producing a metal particle, wherein the metal constituting FIG. 7 A SEM photomicrograph of the metal (silver)<br> copper, gold, nickel, and palladium, or wherein the metal FIG. 8 A SEM photomicrograph of the cross-section of salt is selected from the group consisting of a nitrate, a the metal (silver) particle of the present invention salt is selected from the group consisting of a nitrate, a the metal (silver) particle of the present invention, magnified sulfate, a carbonate, and a chloride.  $60\ 20,000$  times, showing the region SA of void portions

producing a metal particle, wherein the polycarboxylic acid FIG. 9 A diagrammatic view showing the growth of a is at least one polycarboxylic acid selected from the group<br>consisting of citric acid, malic acid, maleic acid, and<br>malonic acid. The present invention is directed to the 65 FIG. 10 A SEM photomicrograph enlarged view of t agent is ascorbic acid or an isomer thereof.

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The present invention is directed to the metal particle,<br>therein the value K determined from a specific surface area particle obtained by the above-mentioned the method for

make the state and by a specific structure and the structure of the center without requiring a<br>measured by a BET method:<br> $SS=6/\rho d$  (1) and intervention, the metal particle has dendritic portions formed from crystals which 15 structure in the spherical surface , and therefore is unlikely to suffer bonding or aggregation of the metal particles and exhibits excellent dispersibility, and has an appropriate tap The present invention is directed to the metal particle,<br>wherein the region SA of void portions obtained by subject-<br>ing the image of the cross-section of the metal particle taken<br>by means of a scanning electron microscop

a non-nucleated, spherical porous material having continufied 10,000 times, the morphology of the cross-section of the ous open pores can be obtained by mixing a metal salt and metal particle has a non-nucleated coral form. metal particle has a non-nucleated coral form. a polycarboxylic acid in a liquid phase to effect a reaction,<br>The present invention is directed to the metal particle,<br>wherein the cross-sectional structure of the metal parti taken by means of a scanning electron microscope, magni- 35 dendritic crystals which have grown uniformly outward fied 20,000 times, has a structure shown in FIG. 1. d 20,000 times, has a structure shown in FIG. 1. from the center without requiring a nucleating agent so that The present invention is directed to the metal particle the metal particle has a fine uneven structure in the sp

particles; and drying the deposited metal particles.<br>The present invention is directed to the method for particle of the present invention magnified 20,000 times.

Ifate, a carbonate, and a chloride.<br>The present invention is directed to the method for obtained by image processing.

metal (silver) particle of the present invention magnified 5,000 times.

FIG. 11 A SEM photomicrograph enlarged view of the persed and hence exhibits excellent dispersibility. The rea-<br>etal (silver) particle of the present invention magnified son why the metal particle is unlikely to suffer bon metal (silver) particle of the present invention magnified

particle in Comparative Example 1, magnified 5,000 times.

eters and SEM photomicrographs of these particles magni- 15 As mentioned above, the metal particle of the present<br>fied 10,000 times, 5,000 times, 2,000 times, and 20,000 invention is unlikely to suffer bonding or aggregati

particle of the present invention taken by means of a 25 scanning electron microscope (SEM), magnified 20,000 particle which is substantially true spherical. By virtue of times. The cross-sectional structure of the metal particle of having the fine uneven structure, the metal particle is fused the present invention has a structure shown in FIG. 1. at a low temperature (for example, at 80 t

invention is a non-nucleated, spherical porous material hav- 30 such as a conductive paste, using the metal particle of the ing continuous open pores, and comprises a metal particle present invention is heated at a relativ formed from dendritic crystals which have grown uniformly (for example, at 120 to 200° C.), the metal particle is fused, outward from the center without requiring a nucleating exhibiting excellent conduction properties. On have dendritic portions formed from needle-like crystals 35 which have thinly grown but has dendritic portions formed have grown in a relatively sparse state. Therefore, the from crystals which have radially grown so that the metal pointed needle-like dendritic portions are likely particle has a fine uneven structure in the spherical surface. and to be strongly fused together, causing aggregation of the In the present specification, the term "non-nucleated" means particles, so that the dispersibilit that a nucleating agent separately added for causing nucle- 40

a scanning electron microscope, magnified 10,000 times. As The metal particle of the present invention preferably has shown in FIG. 2, the morphology of the cross-section of the 45 a volume cumulative particle diameter  $D$ 

present invention taken by means of a scanning electron In the present invention, the particle size distribution microscope (SEM), magnified 10,000 times, 20,000 times, 50 measurement method using image analysis is a metho microscope (SEM), magnified  $10,000$  times,  $20,000$  times,  $50$  and  $40,000$  times, respectively. As shown in FIG.  $4$ , the morphology of the appearance of the metal particle of the present invention has an aegagropila form.

As shown in FIGS 3, 4, and 5, the metal particle is size distribution with respect to the resultant image is substantially true spherical and has dendritic portions 55 measured using a particle size distribution image anal substantially true spherical and has dendritic portions 55 measured using a particle size distribution image analysis<br>formed from crystals which have radially grown substan-<br>system (for example, trade name: Mac-VIEW ver 1. tially uniformly, and therefore has fine unevenuess in the manufactured by Mountech Co., Ltd.), and the volume spherical surface. The unevenuess in the spherical surface of cumulative particle diameter  $D_{50}$  indicates a spherical surface. The unevenness in the spherical surface of cumulative particle diameter  $D_{50}$  indicates a particle diam-<br>the metal particle of the present invention has a fine structure eter at volume accumulation 50 the metal particle of the present invention has a fine structure eter at volume accumulation 50% as measured by a particle comprising protrusions and spaces (depressions) between 60 size distribution measurement method usi

present invention taken by means of a scanning electron to 12  $\mu$ m, more preferably 0.99 to 11  $\mu$ m, as measured by a microscope (SEM), magnified 5,000 times and 2,000 times, particle size distribution measurement method respectively. As shown in FIGS. 6 and 7, the metal particle 65 analysis, and preferably has a volume cumulative particle<br>of the present invention is unlikely to suffer bonding or diameter  $D_{10}$  of 0.45 to 7.8 µm, more p

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5,000 times.<br>FIG. 12 A diagrammatic view showing the growth of a particle of the present invention has dendritic portions metal (silver) particle produced by the method in Compara- 5 formed from crystals which have densely and uniformly<br>grown, and thus has a fine uneven form such that the uneven<br>FIG. 13 A SEM photomicrograph of the metal (sil FIG. 13 A SEM photomicrograph of the metal (silver) structures in the spherical surfaces do not mesh with each ricle in Comparative Example 1, magnified 5,000 times. other, so that the metal particle is unlikely to suffer FIG. 14 A SEM photomicrograph of the metal (silver) or aggregation of the metal particles. Further, the crystals in <br>
rticle in Comparative Example 2, magnified 5,000 times. 10 the metal particle have radially grown outwar particle in Comparative Example 2, magnified 5,000 times. 10 the metal particle have radially grown outward from the<br>FIG. 15 A SEM photomicrograph of a flake-form silver enter, and the metal particles are prevented from be FIG. 15 A SEM photomicrograph of a flake-form silver center, and the metal particles are prevented from being particle magnified 5,000 times.<br>FIG. 16 Analysis values of the metal (silver) particles the crystal growth, so t FIG. 16 Analysis values of the metal (silver) particles the crystal growth, so that the bonding force between the having different volume cumulative average particle diam- metal particles becomes weak.

times. The metal particles is the metal particles of the metal particle has excellent dispersibility in a medium, such as a resin, and further the MODE FOR CARRYING OUT THE INVENTION dendritic portions suffer no breakage upon dispersing the<br>20 metal particle, and it is expected that when the metal particle Hereinbelow, a mode for carrying out the present inven-<br>tion will be described in detail with reference to the accom-<br>conductive composition, such as a conductive paste, it is tion will be described in detail with reference to the accom-<br>possible to easily control the specific gravity or resistance. possible to easily control the specific gravity or resistance. FIG. 1 shows an image of the cross-section of the metal Further, the metal particle of the present invention has a fine ricle of the present invention taken by means of a 25 uneven portion formed in the spherical surface o the present invention has a structure shown in FIG. 1. at a low temperature (for example, at 80 to 100°C.). For this As shown in FIG. 1, the metal particle of the present reason, it is expected that when a conductive compo present invention is heated at a relatively low temperature hand, a conventional dendritic metal particle has dendritic portions formed from pointed needle-like crystals which particles, so that the dispersibility of the particle in, e.g., a resin becomes poor. Further, it is expected that the pointed ation is not present.<br>FIG. 2 is a SEM photomicrograph of the cross-section of a resin, making it difficult to control the specific gravity or FIG. 2 is a SEM photomicrograph of the cross-section of a resin, making it difficult to control the specific gravity or the metal particle of the present invention taken by means of resistance.

metal particle of the present invention has a non-nucleated more preferably 0.3 to 10  $\mu$ m, further preferably 0.5 to 9  $\mu$ m, coral form.<br>
FIGS. 3, 4, and 5 are images of the metal particle of the method using image ana

which an image of the metal particle taken by means of a scanning electron microscope (SEM) at a predetermined esent invention has an aegagropila form.<br>
As shown in FIGS. 3, 4, and 5, the metal particle is size distribution with respect to the resultant image is

the protrusions. Further, the metal particle of the present invention pref-<br>FIGS. 6 and 7 are images of the metal particle of the erably has a volume cumulative particle diameter  $D_{90}$  of 0.5 present invention taken by  $\mu$ m, as measured by a particle size distribution measurement

ticle diameters  $D_{90}$ ,  $D_{10}$  indicate particle diameters at vol-<br>ume accumulations 90% and 10%, respectively, as measured excellent dispersibility upon being dispersed in a resin.

particle size distribution measurement method using image scanning electron microscope, magnified 20,000 times, to analysis is preferably 1.2 to 1.98 more preferably 1.22 to  $\frac{1}{2}$  to  $\frac{1}{2}$  to  $\frac{1}{2}$  to  $\frac{1}{2}$ 

image analysis, is preferably 1.05 to 1.5, more preferably the metal particle taken by means of a scaning electron<br>incroscope, magnified 20,000 times, to analysis using an<br>1.06 to 1.45. Thus, the meal particle diameter an a conductive composition, such as a conductive paste, the<br>metal particle of the present invention exhibits satisfactory<br>conduction properties even at a small content, as compared<br>to the metal particle having no void there

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method using image analysis. The volume cumulative par-<br>ticle diameters of the value K represented by formula (2) above is within<br>ticle diameters  $D_{\text{oo}}$ ,  $D_{\text{oo}}$  indicate particle diameters at vol-<br>the above range, t

by a particle size distribution measurement method using<br>in the metal particle of the present invention, the region<br>image analysis.<br>The ratio of D<sub>80</sub> to D<sub>50</sub> (D<sub>00</sub>/D<sub>50</sub>), as measured by a<br>cross-section of the metal pa The ratio of  $D_{90}$  to  $D_{50}$  ( $D_{90}/D_{50}$ ), as measured by a cross-section of the metal particle taken by means of a scanning electron microscope, magnified 20,000 times, to analysis, is preferably 1.2 to 1.98, more preferably 1.22 to image processing preferably satisfies the relationship:<br> $1.65$  Eurther the ratio of D to D (D D) as monetical  $20 \leq SA \leq 40$ . The region SA of void portions ind 1.65. Further, the ratio of  $D_{50}$  to  $D_{10}$  ( $D_{50}/D_{10}$ ), as measured 20 s 20 ASAS40. The region SA of void portions indicates a value by a particle size distribution measurement method using  $\frac{10}{2}$  determined by subjecting the image of a securitor image analysis, is preferably the cross-section of the metal particle taken by means of a security elec

sme diamed and polycarboxylic acid in a liquid<br>sme diamental sme and the metal particle of the present invention.<br>Invention.<br>The metal particle of the present invention preferably has  $\frac{45}{3}$  formly mixed with each oth

collected by filtration and then dried. With respect to the  $SS = 6/2d$  (1) 65 drying temperature, there is no particular limitation, but the drying temperature is preferably 0 to  $80^{\circ}$  C., more preferably  $(SSSB) \times 100 = K$  (2) 10 to 60° C. The drying time varies depending on the drying

from the group consisting of silver, copper, gold, nickel, and the reducing agent ( $100\%$  by mass) (in terms of a solids palladium. By using the above metal, a metal particle having 5 content). the characteristic features of the present invention can be Further, in the method for producing a metal particle of obtained. The metal salt is preferably selected from the the present invention, if necessary, an additive obtained. The metal salt is preferably selected from the the present invention, if necessary, an additive may be group consisting of a nitrate, a sulfate, a carbonate, and a added. chloride, more preferably a nitrate. Specifically, the metal Examples of additives include cationic dispersants, such salt is preferably selected from the group consisting of silver 10 as higher alkylmonoamine salts, alkyldiamine salts, and nitrate, copper nitrate, gold nitrate, nickel nitrate, palladium quaternary ammonium salts; anionic nitrate, copper nitrate, gold nitrate, nickel nitrate, palladium nitrate, silver sulfate, copper sulfate, gold sulfate, nickel nitrate, silver sulfate, copper sulfate, gold sulfate, nickel carboxylic acid salts, sulfate salts, and phosphate salts; and<br>sulfate, palladium sulfate, silver carbonate, copper carbon-fatty acids, such as lauric acid, ste ate, nickel carbonate, silver chloride, copper chloride, gold<br>chut the additive is not particularly limited to these.<br>chloride, nickel chloride, and palladium chloride. The metal 15 FIG. 9 is a diagrammatic view showing th

particular limitation, and examples include aliphatic poly- 20 As shown in FIG. 9, in the metal particle produced by the carboxylic acids, such as dicarboxylic acids and oxypoly- method of the present invention, there is n carboxylic acids. Examples of dicarboxylic acids include rately add a nucleating agent, and a reducing agent is added malonic acid, succinic acid, maleic acid, and fumaric acid, to a mixture containing a metal salt and a p and examples of polycarboxylic acids include oxydicarbox-<br>victor deposit metal particles in the solution, and then<br>vic acids, such as tartaric acid and malic acid, and oxytri- 25 dendritic crystals grow uniformly outward f carboxylic acids, such as citric acid. Of these, preferred is at ited metal as a center. The crystals radially grow outward least one polycarboxylic acid selected from the group con-<br>from the center so that the resultant m least one polycarboxylic acid selected from the group con-<br>sisting of citric acid, malic acid, maleic acid, and malonic uneven structure in the spherical surface. As shown in FIGS. acid, and more preferred is citric acid, malic acid, or maleic 10 and 11, the ends of the dendritic portions of the metal acid. The polycarboxylic acids may be used individually or 30 particles, each of which is a non-nucl acid. The polycarboxylic acids may be used individually or 30

boxylic acid are mixed with each other is a solvent capable the boundaries between the adjacent metal particles. There-<br>of dissolving therein both the metal salt and the polycar-<br>fore, the metal particle of the present inv boxylic acid, preferably pure water or ion-exchanged water. 35

isomer thereof. Examples of isomers of ascorbic acid include L-ascorbic acid and isoascorbic acid. With respect to

It is preferred that the metal salt, the polycarboxylic acid, control the specific gravity or resistance. Moreover, the and the reducing agent are individually dissolved in pure metal particle obtained by the method of the and the reducing agent are individually dissolved in pure metal particle obtained by the method of the present invenwater or ion-exchanged water and used in the form of an ion has a fine uneven structure formed from dendri water or ion-exchanged water and used in the form of an tion has a fine uneven structure formed from dendritic aqueous solution. The aqueous metal salt solution preferably portions in the spherical surface of the metal par aqueous solution. The aqueous metal salt solution preferably portions in the spherical surface of the metal particle which has a concentration of 3 to 20 mol %/L. The aqueous 45 is substantially true spherical, and therefo has a concentration of 3 to 20 mol %/L. The aqueous 45 is substantially true spherical, and therefore is expected to be polycarboxylic acid solution preferably has a concentration fused at a relatively low temperature and

reducing agent solution fall in the above-mentioned respec-<br>tive ranges, a metal particle which is a non-nucleated,<br>specifical productive composition, and an electronic part<br>spherical provis material having continuous open there can be obtained a metal particle formed from dendritic 55 crystals which have grown uniformly outward from the

vary depending on their respective concentrations. For  $60$  example, it is preferred that, relative to 100 parts by mass of example, it is preferred that, relative to 100 parts by mass of phenol F, phenolic novolak, or alicyclic epoxy resins; oxet-<br>the metal salt, 10 to 100 parts by mass of the polycarboxylic ane resins; resol or novolak phenol acid is incorporated. Further, for example, it is preferred modified organic resins, such as silicone epoxy and silicone that, relative to 100 parts by mass of the metal salt, 60 to 600 polyester. These resins may be used that, relative to 100 parts by mass of the metal salt, 60 to 600 polyester. The parts by mass of the reducing agent is incorporated. Further, 65 combination. it is preferred that the amount of the metal salt incorporated . In the conductive composition, the metal particle: resin<br>is 10 to 60% by mass, the amount of the polycarboxylic acid weight ratio is preferably 90:10 to 70:3

temperature and is not particularly limited, but is preferably incorporated is 10 to 40% by mass, and the amount of the 1 to 20 hours, more preferably 3 to 18 hours. 1 to 20 hours, more preferably 3 to 18 hours.<br>
1 to 18 hours reducing agent incorporated is 30 to 80% by mass, based on The metal constituting the metal salt is a metal selected the total mass of the metal salt, the polyca The metal constituting the metal salt is a metal selected the total mass of the metal salt, the polycarboxylic acid, and<br>from the group consisting of silver, copper, gold, nickel, and the reducing agent (100% by mass) (in

metal particle produced by the method of the present invennitrate, nickel nitrate, or palladium nitrate, further preferably<br>silver nitrate, or gold nitrate.<br>With respect to the polycarboxylic acid, there is no<br>fied 5,000 times.

dendritic crystals grow uniformly outward from the deposin combination.<br>The liquid phase in which the metal salt and the polycar-<br>Inter the metal particles easily separate from each other at<br> $\frac{1}{2}$ The liquid phase in which the metal salt and the polycar-<br>boxylic acid are mixed with each other is a solvent capable<br>the boundaries between the adjacent metal particles. Therefore, the metal particle of the present invention is unlikely to suffer strong bonding or aggregation of the metal particles The reducing agent is preferably ascorbic acid or an and exhibits excellent dispersibility. Further, the ends of the oner thereof. Examples of isomers of ascorbic acid dendritic portions suffer no breakage when dispersed i include L-ascorbic acid and isoascorbic acid. With respect to medium, such as a resin, and it is expected that when the the reducing agent, ascorbic acid and the isomers thereof metal particle is dispersed in a medium, suc the reducing agent, ascorbic acid and the isomers thereof metal particle is dispersed in a medium, such as a resin, to may be used individually or in combination. 40 produce, e.g., a conductive paste, it is possible to eas ay be used individually or in combination. 40 produce, e.g., a conductive paste, it is possible to easily It is preferred that the metal salt, the polycarboxylic acid, control the specific gravity or resistance. Moreover, fused at a relatively low temperature and exhibit excellent

of 0.7 to 40 mol %/L. The aqueous reducing agent solution conduction properties.<br>
preferably has a concentration of 3 to 10 mol %/L. Further, the present invention is a conductive composition<br>
When the concentrations of th

resin. Examples of thermoplastic resins include an acrylic resin, ethyl cellulose, a polyester, a polysulfone, a phenoxy center. resin, ethyl cellulose, a polyester, a polysulfone, a phenoxy<br>The amounts of the metal salt, polycarboxylic acid, and resin, and a polyimide. Preferred examples of thermosetting<br>reducing agent incorporated (in term ane resins; resol or novolak phenolic resins; and silicone-

weight ratio is preferably 90:10 to 70:30. When the metal

particle: resin weight ratio is within the above range, a metal of a thin film having a uniform thickness of about 25  $\mu$ m and film, which is obtained by applying the conductive compo-<br>having excellent conduction properti sition to a substrate to form a film, and heating the formed<br>film, can maintain a desired specific resistance.<br>he effectively formed into a conductor, such as an electronic

Further, in the present invention, by virtue of the method 5 circuit or an electrode, particularly a patterned conductor on in which a metal salt and a polycarboxylic acid are mixed in the surface of a substrate. Further, in which a metal salt and a polycarboxylic acid are mixed in the surface of a substrate. Further, the conductive compo-<br>a liquid phase to effect a reaction, and then a reducing agent sition of the present invention can be a liquid phase to effect a reaction, and then a reducing agent<br>is added to the resultant mixture, the obtained metal particle<br>has dendritic portions formed from crystals which have<br>electrode, a semiconductor sealing agent, has dendritic portions formed from crystals which have electrode, a semiconductor sealing agent, or a conductive grown radially outward from the center without requiring a 10 adhesive, such as a die attach adhesive. grown radially outward from the center without requiring a 10 adhesive, such as a die attach adhesive.<br>
nucleating agent so that the metal particle has a fine uneven<br>
The conductor comprising a cured product obtained by<br>
s particle is unlikely to suffer bonding or aggregation of the is useful as an electronic part for a chip capacitor, an end metal particles, and the metal particle is easily fused at a face under electrode for chip resistanc metal particles, and the metal particle is easily fused at a<br>race under electrode for chip resistance, a variable resistor,<br>relatively low temperature (for example, at 120 to 200° C.), 15 or a film substrate circuit.<br>and,

lent specific resistance can be maintained.<br>The conductive composition of the present invention can aromatic hydrocarbons, such as toluene and xylene; ketones,<br>such as imiting the scope of the present<br>such as methyl ethyl ketone, methyl isobutyl ketone, and<br>cyclohexanone; ethylene glycol monomethyl ether, ethylene<br>glycol diethylene glycol monomethyl ether, diethylene glycol 25 monoethyl ether, diethylene glycol monobutyl ether, and 10 kg of an aqueous silver nitrate solution (concentration: esters thereof, such as acetic esters; and terpineol. It is 10 mol %/L), 4 kg of an aqueous citric acid so esters thereof, such as acetic esters; and terpineol. It is 10 mol %/L), 4 kg of an aqueous citric acid solution preferred that, relative to 100 parts by mass of the total of the (concentration: 10 mol %/L), and 20 kg of metal particle and the resin, 2 to 10 parts by mass of the solvent is incorporated. **30** (L) stainless steel tank and agitated using an agitator (manu-

further comprise at least one member selected from the name: JET TYPE AJITER) at room temperature  $(25\pm10^{\circ} \text{ C})$  group consisting of an inorganic pigment, an organic pig-<br>for 30 minutes to prepare a mixture of silver group consisting of an inorganic pigment, an organic pig-<br>ment, a silane coupling agent, a leveling agent, a thixotropic acid.

be produced by charging the metal particle which is a kg of pure water at 25° C. were individually weighed, and non-nucleated, spherical porous material having continuous then placed in a 450 L stainless steel reaction tan non-nucleated, spherical porous material having continuous then placed in a 450 L stainless steel reaction tank and<br>open pores, a resin, and other components into a mixing agitated using an agitator (manufactured by Shimaz machine, such as a planetary stirring machine, a dissolver, a 40 bead mill, a Raikai mixer, a three-roll mill, a rotary mixer, at room temperature  $(25\pm10^{\circ} \text{ C})$  for 30 minutes to prepare or a twin-screw mixer, and mixing them with one another. an aqueous ascorbic acid solution. Thus, the conductive composition having an apparent vis-<br>
Then, using an agitator having four stainless steel blades<br>
cosity suitable for screen printing, dipping, or another having a diameter of 600 mm (500 rpm), the mixt cosity suitable for screen printing, dipping, or another having a diameter of 600 mm (500 rpm), the mixture of desired film forming method can be prepared. 45 silver nitrate and citric acid was poured at once into the

used as a conductive paste, and applied to a support of, e.g., of silver nitrate and citric acid and the aqueous ascorbic acid polyethylene terephthalate (PET) or indium tin oxide (ITO) solution with each other. by, e.g., a printing or coating method to form a film, and the The aqueous ascorbic acid solution was added to the formed film is cured at, for example, 150° C., obtaining a 50 mixture of silver nitrate and citric acid and formed film is cured at, for example, 150° C., obtaining a 50 mixture of silver nitrate and citric acid and then, after several conductor comprising the resultant cured product. The con-<br>seconds, a reduction reaction was s conductor comprising the resultant cured product. The conductor comprising the cured product preferably has a specific resistance of  $35 \times 10^{-4}$   $\Omega$  cm or less. The temperature for was terminated, the agitation was continued for 30 minutes, heating the conductive composition varies depending on the and then the agitation was stopp type of the resin constituting the conductive composition 55 silver nitrate, citric acid, and ascorpinal is not particularly limited. When the resin is a thermo-reduction reaction had a pH of 2. plastic resin, the conductive composition is preferably The resultant reaction mixture was allowed to stand, and heated to 60 to 350° C., more preferably 80 to 300° C., and, then the supernatant was removed, and the settl heated to 60 to 350° C., more preferably 80 to 300° C., and, then the supernatant was removed, and the settling silver when the resin is a thermosetting resin, the conductive particles were collected by filtration using a when the resin is a thermosetting resin, the conductive particles were collected by filtration using a Nutsche, and the composition is preferably heated to 60 to 350 $^{\circ}$  C., more 60 collected silver particles were sprea

As described above, the conductive composition of the silver particles having a specific surface area of 3.2 m<sup>2</sup>/g, as present invention contains the metal particle which is a measured by a BET method, shown in SEM photo non-nucleated, spherical porous material having continuous graphs of FIGS. 1 to 8, 10, and 11 were obtained. The SA open pores, and therefore the metal particle is fused at a 65 value was 30, which was determined by subjec open pores, and therefore the metal particle is fused at a 65 relatively low temperature (for example, at  $120$  to  $200^{\circ}$  C.), so that a conductor comprising a cured product in the form means of a SEM, magnified 20,000 times, to image pro-

m, can maintain a desired specific resistance.<br>Further, in the present invention, by virtue of the method  $\frac{1}{2}$  circuit or an electrode, particularly a patterned conductor on

The conductive composition of the present invention can Hereinbelow, the present invention will be described in further comprise a solvent , and examples of solvents include 20 more detail with reference to the following Examples , which

(concentration: 10 mol %/L), and 20 kg of pure water at  $25^{\circ}$ <br>C. were individually weighed, and then placed in a 50 liter The conductive composition of the present invention can factured by Shimazaki Mixing Engineering Co., Ltd.; trade

agent, and an anti-foaming agent. <br>Then, 17 kg of an aqueous ascorbic acid solution (aqueous The conductive composition of the present invention can<br>L-ascorbic acid solution; concentration: 5 mol %/L) and 300 The conductive composition of the present invention can L-ascorbic acid solution; concentration: 5 mol %/L) and 300 be produced by charging the metal particle which is a kg of pure water at  $25^{\circ}$  C, were individually w agitated using an agitator (manufactured by Shimazaki Mixing Engineering Co., Ltd.; trade name: JET TYPE AJITER)

sired film forming method can be prepared. 45 silver nitrate and citric acid was poured at once into the The conductive composition of the present invention is prepared aqueous ascorbic acid solution to mix the mixture The conductive composition of the present invention is prepared aqueous ascorbic acid solution to mix the mixture used as a conductive paste, and applied to a support of, e.g., of silver nitrate and citric acid and the aqu

foaming phenomenon accompanying the reduction reaction was terminated, the agitation was continued for 30 minutes,

composition is preferably heated to 60 to 350° C., more 60 collected silver particles were spread over a stainless steel preferably 80 to 300° C.<br>vat, and dried in a dryer at 60° C. for 15 hours. After drying, measured by a BET method, shown in SEM photomicrographs of FIGS. 1 to  $\delta$ , 10, and 11 were obtained. The SA image of the cross-section of the silver particle taken by

WinROOF, manufactured by Mitani Corporation). As shown nitrate solution and the aqueous and the and the acid solution and the acid solution and the and the acid solution and the acid solution with  $\text{FIG. 8}$ , in the imag the cross-section of the silver particle taken by means of a<br>scanning electron microscope, magnified 20,000 times, to <sup>5</sup> after several seconds, a reduction reaction was started, and,<br>image processing, the colored area is portions and the white area is the portion other than the voids.

As shown in FIGS. 1 to 8, 10, and 11, the silver particle containing silver nitrate and ascorbin Example 1 is a non-nucleated, spherical porous material  $10$  reduction reaction had a pH of 2. in Example 1 is a non-nucleated, spherical porous material <sup>10</sup> reduction reaction had a pH of 2.<br>having continuous open pores, and has dendritic portions formed from crystals which have grown uniformly outward<br>from the c

6 L of an aqueous silver nitrate solution (concentration:  $_{20}$  0.15 mol/L) and 200 ml of aqueous ammonia (concentra-0.15 mol/L) and 200 ml of aqueous ammonia (concentra-<br>tion: 25 wt %) were mixed with each other to effect a nointed needle-like crystals which have grown outward from tion: 25 wt %) were mixed with each other to effect a pointed needle-like crystals which have grown outward from reaction, obtaining an aqueous solution of a silver ammine the center in a relatively sparse state, and there reaction, obtaining an aqueous solution of a silver ammine the center in a relatively sparse state, and therefore the complex. To the obtained aqueous solution was added 20 g pointed needle-like dendritic portions are like of hydrated hydrazine (concentration: 80 wt %) as a reduc- 25 causing aggregation of the particles. Further, it is expected<br>ing agent to effect a reduction, depositing silver particles, that the pointed needle-like portion and drying to obtain a spherical silver powder. The mixture Comparative Example 2 is used in a conductive paste, a containing the silver ammine complex and hydrazine uniform metal film cannot be formed from the paste at a containing the silver ammine complex and hydrazine uniform metal film cannot be formed from the paste at a obtained after the reduction reaction had a pH of 2. 30 relatively low temperature and satisfactory conduction prop obtained after the reduction reaction had a pH of 2.

growth of a metal particle produced by the conventional specific gravity or resistance.<br>
method in Comparative Example 1. FIG. 13 is a SEM . With respect to the silver particles in Example 1 and<br>
photomicrograph of the sil photomicrograph of the silver particle in Comparative Comparative Examples 1 and 2, the following measure<br>Example 1 magnified 5,000 times.<br>35 ments were performed. The results are shown in Table 1.

As shown in FIG. 12, in the metal particle produced by the Specific surface area as measured by a BET method no dendrite grows, but crystals grow Tap density determined by a method using a tap density conventional method, no dendrite grows, but crystals grow Tap density determined by a method using a tap density<br>thick so that layers of them are stacked on one another. The measurement apparatus (manufactured by Kuramochi thick so that layers of them are stacked on one another. Therefore, as shown in FIG. 13, the silver particle in Scientific Instruments), in which 10 g of a sample is Therefore, as shown in FIG. 13, the silver particle in Scientific Instruments), in which 10 g of a sample is Comparative Example 1 has dispersion in the particle diam- 40 percisely weighed and placed in a 10 mL settling tu Comparative Example 1 has dispersion in the particle diam-40 precisely weighed and placed in a 10 mL settling tube eter, and it is likely that the silver particles are strongly fused and subjected to 400-time tapping to ca together at their surfaces, causing aggregation of the particles are strongly to 400  $\mu$  to 400  $\mu$ ticles. In the silver particle in Comparative Example 1, no Volume cumulative particle diameters  $D_{10}$ ,  $D_{50}$ ,  $D_{90}$  as dendrite grows and there is almost no void in the metal measured by a particle size distributio dendrite grows and there is almost no void in the metal measured by a particle size distribution measurement particle, and therefore an SA value could not be measured. 45 method using image analysis (particle size distribu

Comparative Example 2 1.00, manufactured by Mountech Co., Ltd.)<br>Particle size distributions  $D_{90}/D_{50}$ ,  $D_{50}/D_{10}$ <br>10 kg of an aqueous silver nitrate solution (concentration: SA Value measured by subjecting the image 10 mol %/L) and 20 kg of pure water at 25° C. were 50 section of the silver particle taken by means of a SEM weighed, and then placed in a 50 L stainless steel tank and magnified 20,000 times to image processing using an weighed, and then placed in a 50 L stainless steel tank and magnified 20,000 times to image processing using an agitated using an agitator (manufactured by Shimazaki Mix-<br>image analysis software (trade name: WinROOF, ing Engineering Co., Ltd.; trade name: JET TYPE AJITER manufactured by Mitani Corporation at room temperature ( $25\pm10^{\circ}$  C.) for 30 minutes. K Value determined from a specific surf

kg of pure water at 25° C. were individually weighed, and area SS is represented by the formula (1) below then placed in a 450 L stainless steel reaction tank and wherein particle diameter d is a volume cumulative then placed in a 450 L stainless steel reaction tank and agitated using an agitator (manufactured by Shimazaki Mixing Engineering Co., Ltd.; trade name: JET TYPE AJITER) at room temperature ( $25\pm10^{\circ}$  C.) for 30 minutes to prepare at room temperature ( $25 \pm 10^{\circ}$  C.) for 30 minutes to prepare and p is a theoretical density of the metal particle, and an aqueous ascorbic acid solution.

having a diameter of 600 mm (manufactured by Shimazaki<br>Mixing Four steel by a diameter of 600 mm ( manufactured by Shimazaki Mixing Engineering Co., Ltd.; trade name: JET TYPE 65 AJITER) at 500 rpm, the aqueous solution of silver nitrate dissolved in pure water was poured at once into the prepared  $(SSBS)x100=K$  (2)

cessing using an image analysis software (trade name: aqueous ascorbic acid solution to mix the aqueous silver WinROOF, manufactured by Mitani Corporation). As shown intrate solution and the aqueous ascorbic acid solution

minutes, and then the agitation was stopped. The mixture containing silver nitrate and ascorbic acid obtained after the

Shown in FIG. 14.<br>FIG. 14 is a SEM photomicrograph of the silver particle<br>in Comparative Example 2 magnified 5,000 times. As seen<br>cous silver nitrate solution (concentration: <sub>20</sub> in FIG. 14, the silver particle produced w upon mixing into a resin, and that when the silver particle in Comparative Example 2 is used in a conductive paste, a FIG. 12 is a diagrammatic view showing the expected erties cannot be obtained, making it difficult to control the owen to with of a metal particle produced by the conventional specific gravity or resistance.

- 
- particle, and therefore an SA value could not be measured. 45 method using image analysis (particle size distribution image analysis system, trade name: Mac-VIEW ver

- 10 SA Value measured by subjecting the image of the cross-section of the silver particle taken by means of a SEM
- K Value determined from a specific surface area SS and a specific surface area BS and represented by the Then, 17 kg of an aqueous ascorbic acid solution (aqueous 55 a specific surface area BS and represented by the L-ascorbic acid solution; concentration: 5 mol %/L) and 300 general formula (2) below, wherein the specific sur particle diameter  $D_{50}$  as measured by a particle size distribution measurement method using image analysis aqueous ascorbic acid solution.<br>
Then, using an agitator having four stainless steel blades measured by a BET method.

 $(1)$ 



As can be seen from Table 1, the silver particle in [Specific Resistance]<br>
Simple 1 has a larger specific surface area than those of the  $15$  Using a 250-mesh stainless steel screen, the conductive Example 1 has a larger specific surface area than those of the 15 metal particles in Comparative Examples 1 and 2. In addimetal particles in Comparative Examples 1 and 2. In addi-<br>
tion, the silver particle in Example 1 has dendritic portions<br>
Comparative Examples 1 and 3 were individually subjected formed from crystals which have densely and uniformly to 71 mm×1 mm zigzag pattern printing on a 20 mm square grown, and therefore has a smaller tap density than that of alumina substrate, and cured under heating condition the silver particle in Comparative Example 1, in which no  $_{20}$  150° C. for 30 minutes. After curing, a resistance was dendrite grows, and has a larger tap density than that of the measured by an LCR meter four-terminal dendrite grows, and has a larger tap density than that of the silver particle in Comparative Example 2, which is formed from needle-like crystals which have thinly grown to cause  $50 \pm 15\%$ . A specific resistance was determined from the larger voids. Further, the silver particle in Example 1 has a specific resistance and the thickness of larger voids. Further, the silver particle in Example 1 has a specific surface area about three times that of the silver particle in Comparative Example 2, but has a K value almost

Comparative Examples 1 and  $3$  were individually subjected alumina substrate, and cured under heating conditions at  $150^{\circ}$  C. for 30 minutes. After curing, a resistance was temperature of  $20±3°$  C. and at a relative humidity of (thickness of the cured film:  $30 \mu m$ ). The results are shown in Table 2.

TABLE 2

		Example 1	(Silver citrate powder)	Comparative Example 1 (Spherical powder)		Comparative Example 3 (Flake powder)	
	Silver/resin ratio (wt $\%$ )	Cured at $150^\circ$ C.	Cured at $200^\circ$ C.	Cured at $150^{\circ}$ C.	Cured at $200^\circ$ C.	Cured at $150^\circ$ C.	Cured at $200^{\circ}$ C.
Specific resistance $(x 10^{-4})$ $\Omega$ · cm)	90/10 80/20 70/30 60/40 50/50	6.81 9.16 24.51	4.15 6.6 20.45	1.15 5.05 Not conductive Not conductive Not conductive Not conductive Not conductive Not conductive Not conductive Not conductive Not conductive Not conductive Not conductive Not conductive Not conductive	0.79 3.09 159.8	0.35 5.18 Not conductive	0.39 3.55 54.1

from particle diameter d and theoretical density  $\rho$  to the when the silver particle: phenoxy resin (silver particle: phenoxy resin) ratio is 70:30, that is, the silver particle weight specific surface area as measured by a BET method. This<br>value confirms that the silver particle in Example 1 has a 45 more excellent specific resistance than those of the conduc-<br>large specific surface area, as compared to in Comparative Example 2 and the density is large relative Examples 1 and 3, and a conductor comprising a cured<br>to the specific surface area and that the silver particle in product obtained by curing the conductive composi to the specific surface area, and that the silver particle in product obtained by curing the conductive composition in Example 1 had a specific resistance of  $24.51 \times 10^{-4} \Omega$  cm or Example 1 has dendritic portions formed from crystals  $\frac{EXan}{50}$  less.

Next, using the silver particles in Example 1 and Com-<br>prepared by the method shown below. With respect to the<br>parative Example 1 and the flake-form silver particle (Com-<br>obtained silver particles in Examples 2, 3, and 4, parative Example 1 and the flake-form silver particle (Com-<br>parative in Examples 2, 3, and 4, a specific<br>parative Example 3) and a phenoxy resin, conductive com- 55 surface area, a tap density, a K value, and volume cumula parative Example 3) and a phenoxy resin, conductive com- 55 surface area, a tap density, a K value, and volume cumula-<br>positions were individually prepared so that the silver tive particle diameters  $D_{\text{tot}} D_{\text{tot}} D_{\text{tot}}$ positions were individually prepared so that the silver tive particle diameters  $D_{10}$ ,  $D_{50}$ ,  $D_{90}$  were measured by the particle phenoxy resin weight ratio (silver particle/phenoxy same methods as those in Example particle : phenoxy resin weight ratio (silver particle/phenoxy same methods as those in Example 1. The specific surface resin) became 90/10, 80/20, 70/30, 60/40, or 50/50, and a area, tap density, K value, and volume cumul specific resistance of each composition was measured by the diameters  $D_{10}$ ,  $D_{50}$ ,  $D_{90}$  of the silver particles in Examples method shown below. The flake-form silver particle used as 60 2, 3, and 4 and SEM photomi Comparative Example 3 has an average particle diameter of magnified  $10,000$  times,  $5,000$  times,  $2,000$  times, and  $10 \mu$ m. The average particle diameter of the flake-form silver  $20,000$  times are shown in FIG. 16. 10 particle indicates an average diameter with respect to the flat<br>
10 surface of the particle. In Table 2, the indication "Not **Example 2** surface of the particle. In Table 2, the indication "Not conductive" means that no electric conduction is made. FIG. 65<br>15 shows a SEM photomicrograph of the flake-form silver 15 shows a SEM photomicrograph of the flake-form silver  $\mu$  A silver particle having a volume cumulative particle particle magnified 5,000 times.

equivalent to that of Comparative Example 2, wherein K $40$  As can be seen from Table 2, with respect to the conduc-<br>value indicates a ratio of the specific surface area determined tive composition using the silver particl tive compositions using the silver particles in Comparative

which have densely and uniformly grown. Further, the silver Further, silver particles having different volume cumula-<br>particle in Example 1 has a sharp particle size distribution.<br>Next, using the silver particles in Examp

same manner as in Example 1 except that the pH of the average particle diameter and has an appropriate tap density<br>mixture containing silver nitrate, citric acid, and ascorbic and a large specific surface area, and further acid obtained after the reduction reaction was adjusted to large relative to the specific surface area, and thus the metal<br>more than 3. SA value of the silver particle in Example 2 as particle can be advantageously used in measured in the same manner as in Example 1 was 20. 5 a conductive paste, a sintering auxiliary, a semiconductor

diameter  $D_{50}$  of 3.32 µm was obtained in substantially the 10 same manner as in Example 1 except that the pH of the same manner as in Example 1 except that the pH of the acid in a liquid phase; adding a reducing agent to the mixture containing silver nitrate, citric acid, and ascorbic resultant mixture to deposit metal particles; and dr acid obtained after the reduction reaction was adjusted to deposited metal particles to obtain a silver particle which is<br>more than 2 to 3 or less. The SA value of the silver particle a non-nucleated, spherical porous mate more than 2 to 3 or less. The SA value of the silver particle in Example 3 which was measured in the same manner as in 15 ous open pores and has :

A silver particle having a volume cumulative particle 20 a specific surface area of 2 to 6 m<sup>2</sup>/g measured by a BET ameter D<sub>50</sub> of 7.97  $\mu$ m was obtained in substantially the method; and diameter  $D_{50}$  of 7.97 µm was obtained in substantially the method; and same manner as in Example 1 except that the pH of the a value K of 3≤K≤15, mixture containing silver nitrate, citric acid, and ascorbic wherein the value K is determined from a specific surface acid obtained after the reduction reaction was adjusted to 2 area SS and a specific surface area BS, an acid obtained after the reduction reaction was adjusted to 2 area SS and a specific surface area BS, and is repre-<br>or less. SA value of the silver particle in Example 4 which  $25$  sented by a formula (2), was measured in the same manner as in Example 1, was wherein the specific surface area SS is represented by a 39.5.

diameters  $D_{50}$ , are individually a non-nucleated, spherical 30 porous material having continuous open pores and have silver particle, and in the formula (2), the specific<br>dendritic portions formed from crystals which have radially surface area BS is a specific surface area as measured a fine uneven structure in the spherical surface. As seen from FIG.  $16$ , in the silver particles in Examples 2 to 4, the ends  $35$ FIG. 16, in the silver particles in Examples 2 to 4, the ends 35  $SS=6/pd$  of the dendritic portions do not tangle, and further the silver particles easily separate from each other at the boundaries  $(SSSB S) \times 100 = K$ between the adjacent silver particles. Therefore, the silver particles in Examples 2 to 4 are unlikely to suffer bonding

particle which is a non-interested, spherical porous material and maximum of the method according to claim 1, wherein the reducing having continuous open pores, and has dendritic portions agent is ascorbic acid or an isome formed from dendritic crystals which have radially grown<br>uniformly outward from the center so that the metal particle 5. The method according to claim 1, wherein the poly-<br>has a fine uneven structure in the spherical surfa has a fine uneven structure in the spherical surface. The  $_{50}$  carboxylic acid is at least one polycarboxylic acid selected motel portiol of the present invention is educated and from the group consisting of citric acid metal particle of the present invention is advantageous not<br>acid and malonic acid; and wherein the reducing agent is<br>acid and malonic acid; and wherein the reducing agent is only in that the metal particle is unlikely to suffer bonding<br>or aggregation of the metal particles and exhibits excellent ascorbic acid or an isomer thereof. dispersibility, but also in that the particle has a uniform

more than 3. SA value of the silver particle in Example 2 as particle can be advantageously used in applications, such as sealing agent, a conductive adhesive, a catalyst, and a medical product. Example 3 medical product.<br>The invention claimed is:<br>
ing a volume cumulative particle 1. A method for producing a metal particle, which com-

A silver particle having a volume cumulative particle 1. A method for producing a metal particle, which com-<br>ameter  $D_{50}$  of 3.32  $\mu$ m was obtained in substantially the 10 prises the steps of: mixing silver nitrate and resultant mixture to deposit metal particles; and drying the deposited metal particles to obtain a silver particle which is

> a volume cumulative particle diameter  $D_{50}$  of 0.5 to 9 um measured by a particle size distribution measurement<br>Example 4 method using an image analysis;

a tap density of 1.8 to 4.5  $g/cm^3$ ;

- 
- 
- 
- 3.5. formula (1), wherein in the formula (1), d is a volume<br>As seen from FIG. 16, the silver particles in Examples 2 cumulative particle diameter  $D_{\epsilon_0}$  measured by a par-As seen from FIG. 16, the silver particles in Examples 2 cumulative particle diameter  $D_{50}$  measured by a particle 4, though they have different volume cumulative particle size distribution measurement method using an ticle size distribution measurement method using an image analysis and p is a theoretical density of the by a BET method, wherein formula  $(1)$  and formula  $(2)$  are as follows:

 $(1)$  $(2)$ .

particles in Examples 2 to 4 are unlikely to suffer bonding<br>or aggregation of the silver particles and exhibit excellent  $\frac{40}{40}$  step and the depositing are each carried out at a temperature<br>or aggregation of the silv

INDUSTRIAL APPLICABILITY 3. The method according to claim 1, wherein the poly-<br>carboxylic acid is at least one polycarboxylic acid selected The metal particle of the present invention is a metal  $45 \text{ from the group consisting of citric acid, malic acid, magnetic particle which is a non-nucleated, spherical porous material.}$ 

\* \* \* \* \*