METHOD FOR FORMING NOBLE METAL NANOARTICLES ON A SUPPORT

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Cross-Reference to Related Application

[001] This application claims the benefit of priority of United States of America Provisional Patent Application No. 61/861,108, filed August 1, 2013, the contents of which being hereby incorporated by reference in its entirety for all purposes.

Technical Field

[002] The invention relates to a method for forming noble metal nanoparticles on a support. In particular, the method includes heating precursors of the noble metal nanoparticles in a spiral glass tube reactor to reduce the precursors to form the noble metal nanoparticles on the support.

Background

[003] Fuel cells can convert chemical energy resulting from the oxidation of fuels directly into electrical energy and are considered to be one of the most important clean energy conversion devices. Currently, low-temperature fuel cell research is quite active particularly in automobile-related fields, backup power, portable and mobile power source because low-temperature fuel cells have advantages in improved fuel efficiency, reduced emission, and more environmentally friendly compared to their internal combustion engines counterpart.

[004] Proton exchange membrane fuel cells (PEMFCs, including direct methanol/ethanol fuel cells, DMFCs) are characterized in that a wide operating temperature range from -20 °C to 180 °C (depending on the solid electrolyte properties), quick start-up and response, and high output power density which allows PEMFCs system to be readily smaller and lighter. PEMFCs are very suitable to be the power source for an automobile, unmanned aerial vehicle (UAV), auxiliary power units (APU), mobile market, portable, small stationary power applications, or a power supply for a small cogeneration system such as a combined heat and power (CHP) system.
[005] The core of PEMFCs is membrane-electrode assembly (MEA) which is composed of a solid electrolyte sandwiched in between two catalytic electrodes. The electrode used generally contains a catalyst layer, a macro-porous layer and a backing layer. The catalyst layer can be fixed directly on solid electrolyte, or supported on backing layer. Currently, PEMFCs employ noble metals especially scarce platinum or its alloy supported on carbon materials as electrode catalysts to promote the reactions of fuel electro-oxidation and oxygen reduction. In current hydrogen-fed PEMFCs, around 75% of precious metal is used as a cathode catalyst to accelerate the sluggish oxygen reduction reaction. Hence, it is imperative to reduce or eliminate the use of platinum in the cathode, which would lead to a more affordable fuel cell system as a whole and is made possible for large volume commercialization.

[006] In order for the fuel oxidation and oxygen reduction reactions in a fuel cell to occur at desired electrochemical kinetic rates and potentials, highly active and durable electro-catalysts are required. Due to the high catalytic nature of platinum and its chemical stability, platinum and platinum alloy materials, supported or unsupported, are preferred as electro-catalyst for the anode and cathodes in low-temperature fuel cells.

[007] Generally, to reduce the impact of costly platinum, conductive materials such as carbon and its derivatives are used to support platinum-based catalysts, which can also help to improve the stability and the dispersion of noble metals. There are two ways to reduce the electrode catalyst cost, which would lead to a reduction in the cost of a PEMFC stack as a whole. One way is to employ non-precious metal catalysts or non-metallic catalysts, which is more attractive and interesting. Another way is to decrease the Pt loading to increase the cost-effectiveness. Although, researchers are striving to improve the quality of the catalysts, the current use of non-precious metal catalysts is still limited by the limited activity in the acidic environment of solid polymer proton conducting electrolyte. Due to this reason, non-platinum catalysts that were
developed have little opportunity to replace platinum-based catalysts at least not in the foreseeable future.

[008] Alloying platinum with various less-expensive materials is one of the possible avenues to either reduce the amount of platinum required or increase the total activity of electrocatalysts, or both. Recently, platinum-based nanostructures such as platinum thin shells capped cheap metallic cores or metal-oxide cores were also suggested to reduce the use of platinum and increase the catalytic activity. Successive reduction procedures or in-situ displacement reactions are always used to achieve the core-shell nano-catalysts. Generally, it is believed that the interaction between the shell metal and core metals/oxides can enhance the activity and durability to some extent.

[009] The reduced noble metallic particle sizes, on the other hand, can produce more surface active sites and can therefore increase the available reaction sites in the electrode's catalyst layer or the so-called three-phase boundaries. Various methods have been reported to prepare supported Pt or Pt-based catalysts with high metallic loading.

[010] US Patent No. 7,713,902 reported a procedure in which lauric acid was used as a surfactant, and mesoporous alumina as template to prepare highly dispersed platinum. The resulting platinum particle size achieved with this method was ranging from 1.0 to 2.4 nm in diameter. In this patent, the procedure described needs to be carried out in butanol-water solution, and the catalyst activity was enhanced compared to the commercial E-TEK catalyst.

[011] In US Patent No. 5,759,944, metals were deposited by suspension of the support material in water, subsequently hydrolysis or precipitation of the corresponding noble metal salts and non-noble metal salts with aqueous reducing agent such as formaldehyde was carried out. In this procedure, a heat treatment was needed for the synthesized samples such as Pt, PtNi and PtNiAu in an inert or reducing atmosphere.
employed a home-made carbon black containing an H-content of greater than 4,000 ppm to synthesize the supported platinum catalyst. The resulting nanoparticle size was around 4.4 nm, which was carried out in aqueous solution at pH of 9. It was claimed that the H-content was helpful to the deposition of particle and activity improvement.

Support pre-treatment or functionalization can also play an important role in reducing the particle size of noble platinum and immobilizing metallic nanoparticles on support surface. US Patent Publication No. 2012/0149545 demonstrated that ammonia treatment of carbon carriers can reduce platinum particles to 1.28 nm on treated carbon powder from 2.26 nm on non-treated carbon powder with a 20 wt% platinum loading. The big increase in surface area results in larger electrochemical surface areas of platinum and the big improvement of activity.

Lin et al. presented in PCT Patent Publication No. 2008/048192 a method to functionalize carbon materials as carriers to support noble nanoparticles. The typical functional groups introduced in that invention were mainly the oxygen-containing radicals such as: -COOH (carboxyl), -CHO (aldehyde), -CO- (carbonyl), -O- (oxo) and -OH (hydroxyl). These surface oxygen-containing radicals can assist in immobilization of metallic precursors and therefore the dispersion of reduced metal particles. With this method, nanoparticle sizes of noble metal can be reduced with sparse surface particle density.

Therefore, there remains a need to provide for improved methods for preparing noble metal nanoparticles suitable for use as a catalyst for fuel cell applications, for example.

Summary

Present inventors have designed a feasible and facile procedure to synthesize ultrafine nanosized noble metal or noble metal-based catalysts with high metal loading and high surface particle density. In various embodiments, the noble metal may be platinum. The procedure is
scalable for mass production of supported or unsupported ultrafine nanosized platinum or platinum-based catalysts with high metal loading, or other metallic nanoparticles.

[017] In one aspect of the invention, there is provided a method for forming noble metal nanoparticles on a support. The method includes mixing a support ink with one or more noble metal precursor solution. The support ink includes a solution of a first polyhydric alcohol and the support. The one or more noble metal precursor solution includes a solution of a second polyhydric alcohol and the one or more noble metal precursor. The method further includes feeding the mixture of the support ink and the one or more noble metal precursor solution to a spiral glass tube reactor. The method further includes heating the spiral glass tube reactor containing the mixture of the support ink and the one or more noble metal precursor solution to reduce the noble metal precursor to form noble metal nanoparticles on the support.

[018] Another aspect of the invention relates to noble metal nanoparticles supported on a support. The average noble metal nanoparticle size is 2.0 nm or below, preferably about 1.8 nm. The noble metal loading based on the total mass of the noble metal nanoparticles and the support can be 40 wt% or more, preferably about 75 wt% or more.

[019] A further aspect of the invention relates to the use of the supported noble metal nanoparticles as a catalyst layer in an electrode for fuel cell applications.

**Brief Description of the Drawings**

[020] In the drawings, like reference characters generally refer to the same parts throughout the different views. The drawings are not necessarily drawn to scale, emphasis instead generally being placed upon illustrating the principles of various embodiments. In the following description, various embodiments of the invention are described with reference to the following drawings.

[021] Figure 1 shows a flow chart of present preparation procedure for ultra-fine nanosized catalysts.
Figure 2 shows a diagrammatic sketch of present designed reaction glass vessel.

Figure 3 shows TEM and particle size histogram of Pt supported on carbon powder samples (40 wt%): (a) Pt/C sample synthesized by common procedure; (b) Pt/C and the carbon powder is treated with citrate before used; (c) Pt/C synthesized by present method herein.

Figure 4 shows TEM and particle size histogram of Pt samples supported on multi-wall CNT and graphene (40 wt%).

Figure 5 shows TEM and particle size histogram of Pt/Graphene powder sample with Pt loading of 60 wt%.

Figure 6 shows electrochemical testing (rotating disc electrode, RDE) of Pt/C samples in oxygen-saturated 0.1 M HClO₄ supporting electrolyte at 25 °C.

Description

The following detailed description refers to the accompanying drawings that show, by way of illustration, specific details and embodiments in which the invention may be practised. These embodiments are described in sufficient detail to enable those skilled in the art to practise the invention. Other embodiments may be utilized and structural, chemical and other changes may be made without departing from the scope of the invention. The various embodiments are not necessarily mutually exclusive, as some embodiments can be combined with one or more other embodiments to form new embodiments.

A feasible and facile method to synthesize ultrafine nanosized noble metal or noble metal-based catalysts having a high metal loading and a high surface particle density is herein disclosed. The method involves a rapid and consecutive flow reduction which can save power, time and preparation cost. Such method further affords controllable and localized heating which can further save energy and improve effectiveness of the chemical reduction process. The total metal loading of the as-formed catalysts can be as high as 75 % of total catalyst mass. The average metal nanoparticle size is about 1.8 nm with narrow distribution and the specific mass
surface area of the noble metal, say platinum, can be as high as 151.0 m²/g. No expensive and cumbersome surfactants are needed for the reduction step, therefore making the synthesis method economically attractive.

[029] Accordingly, in one aspect of the invention, there is provided a method for forming noble metal nanoparticles on a support as illustrated in Figure 1.

[030] A "nanoparticle" refers to a particle having a characteristic length, such as diameter, in the range of up to 100 nm. Examples of a noble metal include platinum, ruthenium, palladium, gold, silver, rhenium, rhodium, iridium, osmium, or a combination thereof.

[031] In some embodiments, the noble metal nanoparticles are nanoparticles comprising or consisting essentially of platinum.

[032] In one embodiment, the noble metal nanoparticles are platinum nanoparticles.

[033] The noble metal nanoparticles may have a regular shape, or may be irregularly shaped. For example, the noble metal nanoparticle may be a sphere, a rod, a cube, or irregularly shaped. The size of the noble metal nanoparticles may be characterized by their mean diameter. The term "diameter" as used herein refers to the maximal length of a straight line segment passing through the center of a figure and terminating at the periphery. The term "mean diameter" refers to an average diameter of the nanoparticles, and may be calculated by dividing the sum of the diameter of each nanoparticle by the total number of nanoparticles. Although the term "diameter" is used normally to refer to the maximal length of a line segment passing through the centre and connecting two points on the periphery of a nanosphere, it is also used herein to refer to the maximal length of a line segment passing through the centre and connecting two points on the periphery of nanoparticles having other shapes, such as a nanocube or a nanotetrahedra, or an irregular shape.
As mentioned above, the present method affords the synthesis of noble metal nanoparticles having mean diameter of about 2 nm or less, such as about 1.9 nm, 1.8 nm, 1.7 nm, or even less, with narrow distribution.

In one embodiment, the mean diameter of the noble metal nanoparticles, such as platinum nanoparticles, is about 1.8 nm.

In various embodiments, the noble metal nanoparticles are essentially monodisperse.

By "support" when used in connection with the noble metal nanoparticles is meant a supporting structure or a supporting material for supporting the noble metal nanoparticles. Generally, any support capable of supporting and providing adequate dispersion for the noble metal nanoparticles may be used. The support is preferably stable in the local environment where the noble nanoparticles are to be used, for example as a catalyst layer in an electrode for low temperature fuel cell applications. The support preferably has a specific surface area and/or porosity sufficient to provide dispersion of the noble metal nanoparticles.

In various embodiments, the support may include one or more carbon material selected from the group consisting of carbon black, carbon nanotube, carbon fibre, graphene, graphite, and carbon mesosphere.

In preferred embodiments, the carbon support containing one or more carbon materials may include carbon materials treated by oxidizing or doping of other elements including nitrogen, sulphur, boron, halogens, and hydrogen. The treatment also includes the heat treatment in reducing atmosphere and/or inert atmosphere, solution, and surface functionalization by various chemicals before use.

In one embodiment, the support comprises carbon black.

In another embodiment, the support comprises carbon nanotube.

In yet another embodiment, the support comprises graphene.

In one embodiment, the support has surface area ranging from 20 to 3,000 m²/g.
The method includes mixing a support ink with one or more noble metal precursor solution. In Figure 1, it is illustrated as an example two separate (and different type) noble metal precursor solutions are mixed together. It is to be understood that in certain embodiments, only one noble metal precursor solution is used while in other embodiments, two, three, four, or even more (different type) noble metal solutions are mixed.

By "mixing" is meant contacting one component with another component. The order of mixing is generally not critical, unless stated otherwise. For example, a first noble metal precursor solution may be mixed with a second noble metal precursor solution before adding a third noble metal precursor. Alternatively, all three noble metal precursor solutions may be simultaneously added to and therefore mixed in a common container.

In one embodiment, the mixture of the support ink and the one or more noble metal precursor solution is conducted by intense agitation or ultrasonic agitation. The agitation is kept until all of mixture is used up.

The support ink includes a solution of a first polyhydric alcohol and the support.

In various embodiments, the solution of the first polyhydric alcohol comprises one or more polyhydric alcohol and water.

In certain preferred embodiments, the solution of the first polyhydric alcohol includes ethylene glycol and water. Preferably, the water content in the solution of the first polyhydric alcohol ranges from zero to 85 % volume.

The one or more noble metal precursor solution includes a solution of a second polyhydric alcohol and the one or more noble metal precursor.

In various embodiments, the solution of the second polyhydric alcohol and the solution of the first polyhydric alcohol are the same and as discussed above.
In various embodiments, the one or more noble metal precursor comprises an oxide, a halide, a nitrite, a sulphate, or a complex of platinum, ruthenium, palladium, gold, silver, rhenium, rhodium, iridium, osmium, or a combination thereof.

In one embodiment where the noble metal nanoparticles comprise platinum nanoparticles, the precursor solution comprises hexachloroplatinic acid ($\text{H}_2\text{PtCl}_6$).

In various embodiments, as illustrated in Figure 1, prior to mixing the support ink with the one or more noble metal precursor solution, the one or more noble metal precursor solution is first mixed with a polybasic carboxylic acid.

In various embodiments, the polybasic carboxylic acid comprises citric acid, tartaric acid, malic acid, oxalic acid, or their salts.

In preferred embodiments, the polybasic carboxylic acid comprises citric acid or a citrate of sodium or potassium. The molar ratio of citrate to total metals is generally ranging from 0.001 to 50.

It has been found that the addition of citrate acid or other polybasic carboxylic acids to the one or more noble metal precursor solution prior to mixing with the support ink aids in achieving formation of a uniformly dispersed noble metal nanoparticles on the support later.

The method further includes feeding the mixture of the support ink and the one or more noble metal precursor solution to a spiral glass tube reactor. The mixture may be fed to the spiral glass tube reactor by a pump, such as a peristaltic pump.

In various embodiments, the diameter of the spiral glass tube reactor is in a range of about 1 cm to 10 cm, preferably, about 2, 3, 4, or 5 cm. The spiral glass tube reactor is specially designed and aimed with a safety valve and thermocouple. The liquid in the spiral glass tube is heated by microwave where noble metal precursors are reduced and deposited on support particles. Figure 2 shows an example of a spiral glass tube reactor suitable for use in the present invention.
In certain embodiments, the method further comprises adding a base solution to the mixture of the support ink and the one or more noble metal precursor solution prior to feeding the mixture to the spiral glass tube reactor. This will increase the pH of the mixture to above 7, such as 7.5, 8, 8.5, 9, 9.5, 10, 10.5, or even higher.

The base solution may include a hydroxide of sodium or potassium, preferably sodium hydroxide.

The method further includes heating the spiral glass tube reactor containing the mixture of the support ink and the one or more noble metal precursor solution to reduce the noble metal precursor to form noble metal nanoparticles on the support.

By "heating" is meant that the temperature of the mixture of the support ink and the one or more noble metal precursor solution is deliberately raised such that the reduction process can take place.

By introducing the mixture into a spiral glass tube reactor, a continuous flow-reduction heating is achieved whereby only a small region is selectively heated. Thus, a continuous flow of small amount of mixed reactants is heated and chemically reduced simultaneously in a very small time frame. This leads to complete (i.e. energy savings) and uniform heat treatment of the noble metal nanoparticles, which is crucial in large volume production. It also enables the reduction of the noble metal nanoparticles and their uniform distribution of nanoparticles on the surface of supports. Advantageously, smaller particle size (1.8 nm) with narrow size distribution and a high metal loading (75 wt%) are achieved (see Examples section below).

In various embodiments, heating comprises exposing the spiral glass tube reactor to microwave heat or millimeter waves. The microwave heat is tune-able and therefore controllable.
Preferably, the heating comprises exposing the spiral glass tube reactor to microwave heat at 100 °C to 200 °C, such as 100 °C, 110 °C, 120 °C, 130 °C, 140 °C, 150 °C, 160 °C, 170 °C, 180 °C, 190 °C, or 200 °C.

In one embodiment, the heating comprises exposing the spiral glass tube reactor to microwave heat at 150 °C.

After reducing the noble metal precursors to the noble metal nanoparticles on the support, separation of the thus-formed supported noble metal nanoparticles may be carried out by known techniques, for example, filtering and drying as shown in Figure 1. In one illustration, the reduced mixture is led to a separation equipment such as vacuum filtration to separate the solid from liquid. After washing with copious deionized water, the filtered solid is freeze dried or dried in a vacuum oven overnight before it can be used directly or further treated.

The supported noble metal nanoparticles formed by the present method have demonstrated an average size of 2.0 nm or even less, such as 1.8 nm. Additionally, the noble metal loading based on the total mass of the noble metal nanoparticles and the support is 40 wt% and above, preferably 60 wt% and above, and in a case where the noble metal is platinum, a 75 wt% loading has been achieved.

The supported noble metal nanoparticles may be used as a catalyst layer in an electrode for low temperature fuel cell applications, for example.

By "comprising" it is meant including, but not limited to, whatever follows the word "comprising". Thus, use of the term "comprising" indicates that the listed elements are required or mandatory, but that other elements are optional and may or may not be present.

By "consisting of" is meant including, and limited to, whatever follows the phrase "consisting of. Thus, the phrase "consisting of indicates that the listed elements are required or mandatory, and that no other elements may be present.
[073] The inventions illustratively described herein may suitably be practiced in the absence of any element or elements, limitation or limitations, not specifically disclosed herein. Thus, for example, the terms "comprising", "including", "containing", etc. shall be read expansively and without limitation. Additionally, the terms and expressions employed herein have been used as terms of description and not of limitation, and there is no intention in the use of such terms and expressions of excluding any equivalents of the features shown and described or portions thereof, but it is recognized that various modifications are possible within the scope of the invention claimed. Thus, it should be understood that although the present invention has been specifically disclosed by preferred embodiments and optional features, modification and variation of the inventions embodied therein herein disclosed may be resorted to by those skilled in the art, and that such modifications and variations are considered to be within the scope of this invention.

[074] By "about" in relation to a given numerical value, such as for temperature and period of time, it is meant to include numerical values within 10% of the specified value.

[075] The invention has been described broadly and generically herein. Each of the narrower species and sub-generic groupings falling within the generic disclosure also form part of the invention. This includes the generic description of the invention with a proviso or negative limitation removing any subject matter from the genus, regardless of whether or not the excised material is specifically recited herein.

[076] Other embodiments are within the following claims and non-limiting examples. In addition, where features or aspects of the invention are described in terms of Markush groups, those skilled in the art will recognize that the invention is also thereby described in terms of any individual member or subgroup of members of the Markush group.

[077] In order that the invention may be readily understood and put into practical effect, particular embodiments will now be described by way of the following non-limiting examples.
Examples

[078] Materials used in the foregoing examples include:

- Hexachloroplatinic acid (H₂PtCl₆·6H₂O)
- Sodium hydroxide (NaOH)
- Ethylene glycol (HOCH₂CH₂OH)
- Hydrochloric acid (HCl)
- Citric acid monohydrate (C₆H₇O₇·H₂O)/salt citrate
- Low cost metal salt such as nickel (II) nitrate (Ni(NO₃)₂), cobalt (II) chloride (CoCl₂)
- Carbon powder (XC-72, Cabot)
- Graphene (home-made)

[079] Example 1

[080] 5.0 g of carbon powder (XC-72) was treated in 200 mL 5.0 M HNO₃ and 5.0 M HCl mixed aqueous solution by refluxing for 5 hours, separated, washed with copious DI water and then heated in oven at 120 °C overnight. The treatment can remove the impurity and enrich the support surface. The acid-treated carbon powder is labelled as XC-72R.

[081] Example 2: Citrate treated carbon powder

[082] 1.0 g of carbon black (XC-72R) and 1.0 g of citric acid monohydrate were mixed with 150 mL DI water with ultrasonic vibration to get highly dispersed carbon ink. After stirring for 2 hours, the mixed ink was dried to form a paste. The paste was placed in quartz boat and heated at 110 °C for 30 min in tube furnace. Then the solid was heated at 300 °C for another 30 min to get the citrate treated carbon powder which was labelled as XC-72RC.

[083] Example 3

[084] 0.6 g of carbon powder (XC-72R) was dispersed in 60 mL ethylene glycol to make carbon ink. 1.08 g of hexachloroplatinic acid was dissolved in 100 mL ethylene glycol. The carbon ink was added into metallic precursor solution and adequately stirred for one and a half
hour. NaOH was added into ethylene glycol to prepare 1.0 M solution and was used to change the above mixed ink pH level to 13. After adequate stirring, the mixed ink was heated to 150 °C and kept refluxing for 1 hour to get the metal precursor reduced adequately. The reduced black mixed slurry was centrifuged to separate the solid after pH level was adjusted in between 3 and 4 by HCl aqueous solution. The final catalyst Pt/C (40 %wt.) was produced after the solid was dried in vacuum oven at 60 °C overnight. Catalyst produced by this procedure is labelled as Pt/C-a.

Example 4

0.6 g of carbon powder which is treated with citrate in Example 2 (XC-72RC) was dispersed in 60 mL ethylene glycol to make carbon ink. 1.08 g of hexachloroplatinic acid was dissolved in 100 mL ethylene glycol. The carbon ink was added into metallic precursor solution and adequately stirred for one and a half hour. NaOH was added into ethylene glycol to prepare 1.0 M solution and was used to change the above mixed ink pH level to 13. After adequate stirring, the mixed ink was heated to 150 °C and kept refluxing for 1 hour to get the metal precursor reduced adequately. The reduced black mixed slurry was centrifuged to separate the solid after pH level was adjusted in between 3 and 4 by HCl aqueous solution. The final catalyst Pt/C (40 %wt.) was produced after the solid was dried in vacuum oven at 60 °C overnight. Catalyst produced by this procedure is labelled as Pt/C-b.

Example 5

0.6 g of carbon powder (XC-72R) was dispersed in 60 mL ethylene glycol to make carbon ink. 1.08 g of hexachloroplatinic acid was dissolved in 100 mL ethylene glycol and 1.6 g of citrate acid was dissolved in 50 mL ethylene glycol, respectively. Before the carbon ink was added, the citrate acid solution was mixed with metallic precursor solution and adequately stirred for one and a half hour. NaOH was added into ethylene glycol to prepare 1.0 M solution and was used to change the above mixed ink pH level to 13. After adequate stirring, the mixed
ink was heated to 150 °C and kept refluxing for 1 hour to get the noble metal precursor reduced adequately. The reduced black mixed slurry was centrifuged to separate the solid after pH level was adjusted in between 3 and 4 by HCl aqueous solution. The final catalyst Pt/C (40 %wt.) was produced after the solid was dried in vacuum oven at 60 °C overnight. Catalyst produced by this procedure is labelled as Pt/C-c.

[089] Same procedures are also applied to synthesize carbon nanotube (CNT) and graphene supported platinum catalysts labelled as Pt/CNT and Pt/graphene, respectively.

[090] Example 6: Catalyst sample characterization

[091] Samples synthesized with different procedures were characterized by Transmission Electron Microscope (TEM) and Figure 3 shows the typical TEM pictures and the particle size histogram. Samples "a" was synthesized by the previous ethylene glycol procedure when no citrate was involved. The general feature shows that this sample has quite uniform particle distribution on support surface without aggregation. The particle size range is also narrow and the average particle size is 2.3 nm with 40 %wt platinum loading.

[092] From the TEM result of sample Pt/C-b, it can be seen that further treatment of carbon powder with citrate can indeed decrease the platinum particle size greatly, from 2.3 nm to 1.8 nm. The surface functionalization of carbon powder proves useful in improving the dispersion of metallic particles. However, it is also found that the particle number density on the carbon surface is quite low compared to the sample Pt/C-a. This indicates that the immobilized citrate group on carbon surface may play a role of a double-edged sword. For comparison, the catalyst sample disclosed in PCT Publication No. WO 2008/048192 has similar results, namely, sparse platinum particles found on the support surface (Figure 1 of WO 2008/048192). The average particle size reported herein is much smaller than that disclosed in PCT Publication No. WO 2008/048192. Moreover, the latter has very low metallic content of only 16.3%wt.
Compared to sample Pt/C-b, the catalyst with the procedure disclosed herein, i.e. Pt/C-c has much more dense particles on the carbon surface. This indicates that most of platinum is reduced and deposited on the carrier surface. The specific mass surface area can reach 150 m²/g Pt. The current procedure in this disclosure therefore has the advantages of full depositing of metallic particles and obviously increasing metal surface area.

Figure 4 presents the TEM results of platinum particles supported on multi-walled CNT and graphene. It is amazing that the platinum particle sizes are almost the same even on different carriers with different surface area and structures. These results demonstrated the repeatability of the invention disclosed herein. The platinum loading on graphene was further increased from 40 %wt to 60 %wt and the TEM result is presented in Figure 5. From this figure, one can see that the average particle size is still around 1.8 nm without any obvious increase in particle size compared to the sample containing 40 %wt. of platinum. It is also found the particle sizes are quite uniform and the particle distribution on the support surface is also very homogeneous. Compared to the sample of 40 %wt Pt/graphene, the platinum particle numbers per unit area is obviously increased. This result has proved the feasibility of the procedure described above.

Example 7: Electrochemical testing of catalyst samples

5 mg of supported Pt catalyst prepared by this method was dispersed in 1.0 mL ethanol to get catalyst ink in ultrasonic bath for 20 min. 50 μL Nafion alcohol solution (5 %wt Nafion) was added into the ink, followed by ultrasonic oscillations for another 30 min. 20 μL catalyst ink was placed in the graphite disk electrode and dried in oven at 75 °C for 15 min to prepare working electrode. The electrochemical testing was conducted in a 4-neck electrochemical cell containing 0.1 M HClO₄ solution as supporting electrolyte, Ag/AgCl as reference electrode and platinum wire as counter electrode. The electrolyte was saturated by bubbling oxygen into solution and the potential was swept from 1.15 V to 0 V. The rotating disk electrode (RDE)
testing results at 1,600 rpm are shown in Figure 6, from which it can be seen that the catalysts prepared by this invented procedure are superior to commercial counterpart.

The current data at 0.85 V was selected to compare the catalyst activity towards oxygen reduction reaction (ORR) and included in Table 1.

Table 1. characterization summary of TEM and RDE testing results.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Average particle sizes (TEM)</th>
<th>Current @ 0.85V (RDE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt/C (40% wt, commercial)</td>
<td>2.30</td>
<td>1.72mA/mg Pt</td>
</tr>
<tr>
<td>Pt/C-c (40% wt)</td>
<td>1.81</td>
<td>1.85mA/mg Pt</td>
</tr>
<tr>
<td>Pt/Graphene (40% wt)</td>
<td>1.85</td>
<td>2.04mA/mg Pt</td>
</tr>
</tbody>
</table>
Claims

1. A method for forming noble metal nanoparticles on a support, comprising:
   - mixing a support ink with one or more noble metal precursor solution, wherein the support ink comprises a solution of a first polyhydric alcohol and the support, wherein the one or more noble metal precursor solution comprises a solution of a second polyhydric alcohol and the one or more noble metal precursor;
   - feeding the mixture of the support ink and the one or more noble metal precursor solution to a spiral glass tube reactor;
   - heating the spiral glass tube reactor containing the mixture of the support ink and the one or more noble metal precursor solution to reduce the noble metal precursor to form noble metal nanoparticles on the support.

2. Method according to claim 1, wherein heating comprises exposing the spiral glass tube reactor to microwave heat or millimeter waves.

3. Method according to claim 2, wherein heating comprises exposing the spiral glass tube reactor to microwave heat at 100 °C to 200 °C.

4. Method according to claim 3, wherein heating comprises exposing the spiral glass tube reactor to microwave heat at 150 °C.

5. Method according to any one of claims 1-4, wherein the one or more noble metal precursor solution further comprises a polybasic carboxylic acid.
6. Method according to claim 5, wherein the polybasic carboxylic acid comprises citric acid, tartaric acid, malic acid, oxalic acid, or their salts.

7. Method according to claim 6, wherein the polybasic carboxylic acid comprises citric acid or a citrate of sodium or potassium.

8. Method according to any one of claims 1-7, wherein the solution of the first polyhydric alcohol and the solution of the second polyhydric alcohol are the same.

9. Method according to claim 8, wherein the first polyhydric alcohol and the second polyhydric alcohol each comprises ethylene glycol.

10. Method according to any one of claims 1-9, further comprises adding a base solution to the mixture of the support ink and the one or more noble metal precursor solution prior to feeding the mixture to the spiral glass tube reactor.

11. Method according to claim 10, wherein the base solution comprises a hydroxide of sodium or potassium.

12. Method according to any one of claims 1-11, wherein the noble metal nanoparticles comprise nanoparticles of platinum, ruthenium, palladium, gold, silver, rhenium, rhodium, iridium, osmium, or a combination thereof.
13. Method according to claim 12, wherein the one or more noble metal precursor comprises an oxide, a halide, a nitrite, a sulphate, or a complex of platinum, ruthenium, palladium, gold, silver, rhenium, rhodium, iridium, osmium, or a combination thereof.

14. Method according to any one of claims 1-13, wherein the support comprises one or more carbon material selected from the group consisting of carbon black, carbon nanotube, carbon fibre, graphene, graphite, and carbon mesosphere.

15. Method according to any one of claims 1-14, further comprising filtering and drying the thus-formed supported noble metal nanoparticles.

16. Noble metal nanoparticles supported on a support, wherein the average noble metal nanoparticle size is 2.0 nm or below.

17. Supported noble metal nanoparticles of claim 16, wherein the noble metal loading based on the total mass of the noble metal nanoparticles and the support is 40 wt% or more.

18. Supported noble metal nanoparticles of claim 17, wherein the noble metal loading based on the total mass of the noble metal nanoparticles and the support is 60 wt% or more.

19. Supported noble metal nanoparticles of any one of claims 16-18, wherein the average noble metal nanoparticle size is 1.8 nm and the noble metal loading based on the total mass of the noble metal nanoparticles and the support is 75 wt%.
20. Use of the supported noble metal nanoparticles of any one of claims 16-19 as a catalyst layer in an electrode for fuel cell applications.
FIGURE 1

Ethylene glycol solution containing metal precursor 1

Sufficiently mixed

Mixture stirred ≥ 0.5 hr

Citrate salt

Support powders

Ethylene glycol

Ink

Sufficiently mixed

pH ≥ 10

Peristaltic pump

Spiral glass tube reactor

Vacuum filter

Freeze dry

Catalyst product

Controlled microwave
FIGURE 3 (cont.)

![Image of a microscope slide with a histogram showing particle diameter distribution.

- The histogram indicates that the average particle diameter is 1.81 nm.

- The x-axis represents the particle diameter in nanometers (nm), ranging from 1.0 to 3.0 nm.

- The y-axis represents the percentage of particles within each diameter range, ranging from 0 to 25%.

- The scale bar on the left indicates a size of 20 μm.
INTERNATIONAL SEARCH REPORT

International application No. PCT/SG2014/000338

A. CLASSIFICATION OF SUBJECT MATTER

BO1J 23/38 (2006.01) H01M 4/00 (2006.01) H01M 8/00 (2006.01) B82B 3/00 (2006.01)

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

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

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

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

WPI, EPDOC: IC/CC Marks: B82B 3/00, B82Y 40/00, B82Y 99/00, B01J 23/38, B01J 23/42, B01J 23/44, B01J 23/46, B01J 23/48, B01J 23/50, B01J 23/52, H01M 4/00, H01M 8/00; keywords: nano, micro, platinum, ruthenium, palladium, gold, silver, rhenium, rhodium, iridium, osmium, polyhydric alcohol, polyalcohol, ethylene glycol, metal precursor, halide, nitrate, sulphate, sulfate, oxide, complex, hexachloroplatinate (H2PtCl6). Other cited references: carboxylic acid, citrate, citric acid, tartaric acid, malic acid, oxalic acid, support, carbon nanotube, carbon black, carbon fibre, carbon fiber, graphene, graphite, carbon mesosphere, spiral reactor, heat, microwave, thermal, temperature and "Applicant/Inventor name search": Nanyang Technological University / ZHOU, Weijiang; LI, Miao; CHAN, Siew hwa" through AusPat and Esp@cenet; USPTO, Google Scholar, ScienceDirect & Google Patents using similar search terms.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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Date of the actual completion of the international search 1 October 2014

Date of mailing of the international search report 01 October 2014

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<td>US 6689505 B1 (ALBERS et al.) 10 February 2004 Abstract; claims 1, 4</td>
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