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Size-controlled growth of nanoparticles in a highly ionized pulsed plasma

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Copper nanoparticles (NPs) were synthesized using a high power pulsed hollow cathode technique and the produced NPs were studied as a function of pulse parameters, i.e., frequency, peak current, and pulse width. It was found that the particle size can be altered in a range from 10 to 40 nm by changing any one of the pulse parameters. The mechanisms of NP synthesis with respect to a pulsed discharge and a high degree of ionization of the sputtered material are discussed. © 2013 American Institute of Physics. [http://dx.doi.org/10.1063/1.4788739]

Nanoparticles (NPs) are of interest in many applications due to their physical and chemical properties that greatly differ from bulk properties. Examples include catalytic effects because of their high surface-to-volume ratio,1,2 and interaction with light due to plasmonic activity.3,4 For both examples, the response is a function of NP size and the geometry of the NP. Therefore, a technique for controlling these parameters to accurately synthesize NPs—in sufficient number and without agglomeration—is desired.

One method for synthesizing NPs is by using plasma discharges where the material for NP growth is supplied to the plasma via sputtering the source material from a target.5–7 A benefit of this method is that NPs of any material, which can be sputtered, can in principle be synthesized. When the NPs have grown to a certain size, they attain a negative charge.8,9 The electrostatic repulsion between such charged NPs hinders further agglomeration inside the plasma. Using dc discharges, it has been shown that the size of the NPs can be varied by changing process parameters such as gas flow, gas pressure, and discharge power.10–12 As a way of obtaining additional control parameters, pulsing has been used in the related field of magnetron sputtering for thin film growth, where it has been shown to give a wider accessible range of plasma parameters.13 Most important in the present application is that the momentary density of sputtered material can be increased in a pulse while keeping the gas temperature low, and that a high degree of ionization of the sputtered material can be achieved.14 Nanoparticle synthesis using pulsed plasma discharges is a growing research field.15–17 For instance, Stranák et al.15 demonstrated particle size control—8 to 10 nm in diameter—in a pulsed magnetron over a frequency range from 0.1 to 25 kHz. In this paper, we report a NP synthesis method that makes use of a high power pulsed hollow cathode (HC) discharge, a technique that can exploit a high degree of ionization of the sputtered species. We demonstrate that it is possible to tailor the size of NPs with spherical shape and diameters in the range of 10 to 40 nm by varying the pulse frequency, the pulse amplitude, and the pulse length. As a special case, we demonstrate that size control is achievable with constant average power.

The NPs are formed by sputtered vapor in the gas phase, and the growth can be divided into three main stages:9 nucleation, coagulation, and accretion by attachment of atoms and ions. The nucleation starts when two metal particles (atoms or ions) form a dimer by a three-body collision with a third particle. These dimers then grow by attachment of individual metal atoms and ions to small clusters. In the second stage, the density of such clusters has become large enough for coagulation between them. The small clusters have a charge that is determined by a stochastic process and they can be neutral, positively, or negatively charged. However, when the clusters attain a certain size, of the order of a few to ten nanometers,9,18 the charge becomes determined by the balance of ion and electron currents to the cluster, i.e., the floating condition, and the clusters approach the floating potential19,20

\[ U_{\text{fi}} = -K_1 \frac{k_B T_e}{e}, \]  

where \( k_B \) is the Boltzmann constant, \( e \) the elementary charge, and \( T_e \) the electron temperature. The factor \( K_1 \) depends on the ion mass and temperature ratio of electrons and ions, but can be assumed to be a constant factor of 2.41 for \( T_e/T_i \approx 100 \). Nanoparticles that attained floating potential repel each other, further growth by coagulation is suppressed and stage three begins with growth only by attachment of metal atoms and ions. For the atoms, this process is independent of the charge and their attachment cross section is given by the geometrical cross section of the NP,

\[ \sigma_{\text{att, atom}} = \pi R_{n}^2, \]  

where \( R_n \) is its radius. The attachment cross section for ions with opposite charge to the NP is given by,19,21

\[ \sigma_{\text{att, ion}} = \pi R_{n}^2 \left( 1 + \frac{|eU_{\text{att}}|}{E_{\text{kin}}} \right), \]  

where \( E_{\text{kin}} \) is the kinetic energy of ions that is of the order of \( k_BT_i \) for low drift velocities. Combining these expressions with the floating potential gives the effective collection cross section for singly charged, thermally distributed ions as,

\[ \sigma_{\text{att, ion}} = \pi R_{n}^2 \left( 1 + K_1 \frac{T_e}{T_i} \right). \]
For typical discharge parameters, e.g., $T_e = 2$ eV and $T_i = 0.03$ eV, this is more than two orders of magnitude larger than the atom attachment cross section, Eq. (2).

By varying the plasma parameters (sputtered density and degree of ionization, gas temperature, and electron temperature), there are two possibilities to influence the NP growth. In the first two stages of nucleation and coagulation, the most important parameters are the density of sputtered material and the gas temperature, the latter because a high gas temperature suppresses nucleation. In the third stage of growth, the most important parameters are the degree of ionization of the sputtered species and the electron temperature. In an environment with a given number density of sputtered material, variations in these two parameters—from the difference in the attachment rates due to different cross-sections [Eqs. (2) and (3)]—can influence the rate of NP growth by several orders of magnitude.

For achieving a plasma in which the above identified key discharge parameters can be flexibly varied, a setup was developed using a hollow cathode in pulsed operation. Due to the hollow cathode effect, a high degree of ionization can be achieved. The pulsed technique applied to the hollow cathode is similar to high power impulse magnetron sputtering, which has shown to increase the degree of ionization substantially as compared to dc magnetron sputtering. The pulsed technique applied to the hollow cathode was developed using a hollow cathode in pulsed operation. Due to the hollow cathode effect, a high degree of ionization can be achieved. The pulsed technique applied to the hollow cathode is similar to high power impulse magnetron sputtering, which has shown to increase the degree of ionization substantially as compared to dc magnetron sputtering. Further, the small orifice of the hollow cathode leads to an ejection of the sputtered source vapor into a smaller region and maintains a higher material density.

A sketch of the experimental setup is shown in Fig. 1(a). The chamber is cylindrical with a length of 430 mm and a diameter of 290 mm. The target is a hollow cathode made of copper which is mounted in the center of the lid. The cathode (length: 54 mm, inner diameter: 5 mm) is water-cooled. A grounded anode ring (AR, diameter: 30 mm) is placed 45 mm below the cathode. The motivation for placing the anode ring in this location is to force the discharge current through the region in which the NPs grow [Fig. 1(b)], thus providing a means to influence the plasma parameters during the growth process. A stainless steel mesh cage encloses the anode ring. The substrate table is made of stainless steel and can be loaded with six substrates. The distance between the substrate table and the cathode is 160 mm. The substrates have a size of 10 × 10 mm and are made of silicon coated with a 200 nm thick titanium layer; they were positively biased at 10 V during the experiments. The base pressure of the chamber was 5.3 × 10⁻⁴ Pa, and argon (99.9999%) at a pressure of 107 Pa was used as the working gas. The pulsed parameters were varied for the frequency in a range between 250 and 1300 Hz and the pulse width between 10 and 77 μs; a peak current of up to 20 A was used.

The NPs were analyzed using a scanning electron microscope (SEM, LEO 1550 Gemini) by taking a number of plan-view micrographs with different magnifications at several positions on the substrate. From each micrograph, the nanoparticle sizes were determined and the measured size distributions were combined to improve the statistics. The mean size and the standard deviation were determined by fitting a lognormal distribution. An example of a micrograph is given in Fig. 1(c) ($f = 698$ Hz, $I_H = 10.2$ A ($U_H = 576$ V), and $t_{PW} = 30$ μs) together with the corresponding size distribution determined from the micrograph, Fig. 1(d). The NPs are the white features that have a spherical shape. Surface features from the titanium coating of the substrate are visible as plates sticking out of the surface.

The dynamics of the atom and ion motion in this type of discharge has recently been studied by means of simulations by Hasan et al. The general behavior can be summarized as follows: Initially, a puff containing sputtered material is ejected with each pulse, see Fig. 1(b). The metal atoms form one cloud, which drifts with the gas flow and expands by classical diffusion. At the time of a pulse, the atom cloud from the previous pulse is typically between the hollow cathode and the anode ring, and has a diameter of a few cm. The ejected atom cloud will then partially overlap with the remnants of the atom clouds from the previous pulses, i.e., the fractions that have not been consumed in the NP growth process. The ions form a separate cloud [not shown in Fig. 1(b)] that is subject to ambipolar diffusion, a much faster process than the classical diffusion of the metal atoms. When a pulse is applied, the center of the ion cloud from the previous pulse is, therefore, typically below the anode ring. Both for neutral and ionized sputtered material, there is thus a direct interaction of overlapping sputtered material from subsequent pulses. The clouds containing neutrals merge with each other, and the ions ejected in subsequent pulses flow through these overlapping neutral clouds providing new material. In addition, there is an indirect interaction when the discharge current of subsequent pulses penetrates the region between the hollow cathode and the anode ring. This can affect both the degree of ionization of the neutral material and, through the electron temperature [see Eq. (3)], the growth rate of NPs in the attachment phase.

Consider a typical NP which is initiated by the formation of a dimer in the ejected puff of sputtered material, at a

![FIG. 1. (a) Sketch of the setup. The gas is fed through the hollow cathode (HC). The hollow cathode is biased and a grounded anode ring (AR) is placed below the hollow cathode and enclosed by a mesh (M). The NPs are collected on a positively biased substrate (S). (b) Sketch of the sputtered metal atoms ejected during consecutive pulses. (c) Example of a SEM plan-view micrograph of spherical NPs. The structure of the titanium film can be seen as plates sticking out of the surface. (d) The corresponding size distribution calculated from the micrograph.](image-url)
time when it is close to the cathode opening. Assuming that
in its subsequent trajectory, it moves with the gas as modeled
by Hasan et al.26 and will—during the first ms—be pushed
10 to 20 mm from the cathode by the fast gas flow associated
with the puff. Approximately at this distance, it will then be
picked up by the slower steady feed gas flow pattern and
take an additional 10 to 15 ms to reach the anode ring. For
the frequency \( f = 700 \text{ Hz} \), the NP will be passed by typically
ten ejected ion clouds, and also becomes exposed to ten dis-
charge current pulses with enhanced ionization and elevated
electron temperature. The densities of the sputtered metal
atom and ion clouds will, due to the NP growth itself, be
reduced by an unknown fraction below those modeled in
Ref. 26. We are, therefore, limited to qualitative analysis and
identify four mechanisms that would favor the growth of
NPs: (1) Pulse strength: strong pulses will give a higher den-
sity of single-pulse sputtered material close to the cathode.
This enhances the two first phases of nucleation and coagula-
tion. This also holds true for (2) neutral overlap: mixing
with the atom clouds from the previous and subsequent
pulse(s). (3) Ion overlap: we define as NPs being swept over
by the ion clouds from subsequent pulses. This makes the
faster ion attachment to NPs possible in the third phase of
growth, and this, in turn, is enhanced by an increase in elec-
tron temperature due to (4) discharge overlap: penetration of
the discharge current from subsequent pulses. Below we
present four experiments demonstrating that the trends of NP
sizes with process parameters \((f, I_{H}, \text{ and } t_{PW})\) can all be qual-
itatively understood in terms of these four mechanisms.

In the first two experiments, the pulse amplitude and the
pulse length were varied, while the pulse frequency was kept
constant at \( f = 700 \text{ Hz} \). As described, most of the drift time
of a NP down to the anode ring is spent in the feed gas flow
which is almost independent of the pulse strength. Therefore,
a NP is exposed to approximately the same number of pulses
in the overlap mechanisms.

Figure 2(a) shows the effect of varying the peak current.
Two size populations are found. At the highest current, still a
tiny fraction of smaller NPs is produced; this quantity is too
small to obtain reliable statistics for determining a size distri-
bution. For the dominating population of NP size data, there
is an approximately linear relation between the size and the
peak current. The sputter yield can be assumed to be constant
which allows also to assume a linear increase of the amount
of sputtered material with the peak current. The variation in
NP size with peak current is, therefore, proposed to be
mainly a pulse strength effect due to the amount of material
available for growth. However, with stronger pulses there
will also be a stronger discharge overlap, and possibly also a
higher degree of ionization. Both effects would enhance NP
growth in the third phase.

Figure 2(b) shows the effect of varying the pulse width
\( t_{PW} \). The peak current was kept constant by adjusting the
discharge voltage in the range \( 740 \text{ V} \quad (t_{PW} = 77 \mu s) \) to \( 472 \text{ V} \quad (t_{PW} = 10 \mu s) \). The combined effect of longer pulse width,
costant peak current, and lower applied voltage is that the
amount of sputtered material increases, but slower than line-
early with the pulse width. This is consistent with the trend in
size in Fig. 2(b) which, therefore, is proposed to be primarily
a pulse strength effect. The fraction of small NPs disappears,
and instead the width of the size distribution becomes
broader as compared to Fig. 2(a). Such broadening of the
size distribution could be explained by having a situation
where different stages of the NP growth may take place at
the same time. This could be the case when assuming that
the ejected cloud is more spread out for longer pulses and,
therefore, the material density might be less concentrated.

In both aforementioned experiments, the pulse strength
effect is proposed to be the primary mechanism to describe
the trends in NP size. We stress that this does not mean that
we propose the three types of overlap to be inconsequential,
only that we do not see them as directly linked to the size
trends. The effect of overlaps is more directly studied by
changing the frequency, shown in Fig. 2(c). The model calcu-
lations26 show that the production, the degree of ioniza-
tion, and the extraction of sputtered material in each
individual pulse are only minimally influenced by the fre-
cuency, apart from a minor interaction between pulses for
the highest frequencies in the studied range. Still, it was pos-
sible to vary the NP size in the range 10 to 40 nm. This is a
clear demonstration that there must be some kind of interac-
tion between the pulses—i.e., some type of overlap(s)—
because the amount of sputtered material in each pulse is
constant. The “neutral overlap” of metal atom clouds
increases with the frequency, which would enhance the for-
formation of dimers and small clusters. “Ion overlap” also
increases since a larger number of metal ion clouds pass by
the NP growth region delivering new material. Further, a NP
in the third stage of growth will be subject to a number of
“discharge overlaps” in which the ion attachment efficiency

\[
\begin{align*}
\text{average power (W)} & \quad 0 \quad 20 \quad 40 \quad 60 \\
\text{peak current (A)} & \quad 20 \quad 40 \quad 60 \\
\text{pulse width (ms)} & \quad 0.5 \quad 1.0 \quad 1.5 \\
\text{frequency (kHz)} & \quad 0 \quad 20 \quad 40 \quad 60 \\
\text{frequency (kHz)} & \quad 0 \quad 20 \quad 40 \quad 60
\end{align*}
\]

FIG. 2. Mean NP diameters obtained with the pulsed hollow cathode for varying process parameters of (a) peak current \((f = 700 \text{ Hz}, t_{PW} = 30 \mu s)\), (b) pulse
width \((f = 700 \text{ Hz}, I_{H} = 10 \text{ A})\), (c) frequency at constant energy per pulse \((I_{H} = 10 \text{ A}, t_{PW} = 30 \mu s)\), and (d) frequency at constant average power
\((P_{av} = 30 \text{ W}, t_{PW} = 30 \mu s)\). The peak current was adjusted to keep the average power constant with frequency. The error bar indicates the standard deviation
of the size distribution and for points with no error bar the statistic was too poor to estimate it.
could increase dramatically for negatively charged NPs. With several independent mechanisms, the combination is likely to give a size growth faster than a linear $f$-trend, at least up to the point where most of the sputtered material is consumed for NP formation. The fast rise and saturation in NP size, seen around 700 Hz in Fig. 2(c), might be caused by such effects.

Figure 2(d) shows an example where the NPs are varied at constant average power. In this case, the pulse length is kept constant at 30 μs, while the frequency and peak current were varied to give a constant average power $P_{av} = 30$ W. This involves two opposing trends and two counteracting mechanisms. With increasing frequency, we have the increasing size trend in Fig. 2(c). Higher frequency, however, is now compensated by a lower peak current which, according to Fig. 2(a), would give smaller NPs. Figure 2(d) shows that these two effects do not cancel. A variation in NP size is possible over the full range from 10 to 40 nm, and with several attractive features: a smooth size variation rather than the step-like function in Fig. 2(c), a smaller spread in size than in Fig. 2(b), and without the two-size distribution in Fig. 2(a).

It has been found that the NP size can be adjusted in a range between 10 and 40 nm by using a high power pulsed hollow cathode and altering the process conditions. It was found that the NP size can be controlled by adjusting one of the pulse parameters, i.e., frequency, pulse width, or peak current. In these cases, the NP size always increases with average discharge power. In addition, it was shown that the NPs size could be controlled also when keeping the average discharge power constant, by changing the frequency and adjusting the peak current. The growing NPs are transported together with the gas flow and, hence, in a cloud of metal atoms which drift with the background gas, and expands by diffusion. Subsequent neutral clouds of this type overlap, and are also periodically flushed by ion clouds as the ion ambipolar diffusion takes place on a faster time scale than neutral convection and diffusion. Four basic mechanisms affecting the growth dynamics due to the pulsing of the discharge are proposed, which provide control parameters for the growth: pulse strength, neutral overlap, ion overlap, and discharge overlap. The pulse strength determines the amount of material sputtered in each pulse, while the frequency is the most important pulse parameter to influence the degree of overlaps between the pulses. A high degree of ionization is a key feature in the process since it is expected to dramatically increase the growth rate for NP growth as soon as the NPs become negatively charged.

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