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**Ultrathin Diamond-like Carbon Films Deposited by Filtered Carbon Vacuum Arcs**

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Ultrathin Diamond-like Carbon Films Deposited By Filtered Carbon Vacuum Arcs

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ABSTRACT

Ultrathin (< 5 nm) hard carbon films are of great interest to the magnetic storage industry as the areal density approaches 100 Gbit/in². These films are used as overcoats to protect the magnetic layers on disk media and the active elements of the read-write slider. Tetrahedral amorphous carbon films can be produced by filtered cathodic arc deposition, but the films will only be accepted by the storage industry only if the “macroparticle” issue has been solved. Better plasma filters have been developed over recent years. Emphasis is put on the promising twist filter system - a compact, open structure that operates with pulsed arcs and high magnetic field. Based on corrosion tests it is shown that the macroparticle reduction by the twist filter is satisfactory for this demanding application, while plasma throughput is very high. Ultrathin hard carbon films have been synthesized using S-filter and twist filter systems. Film properties such as hardness, elastic modulus, wear, and corrosion resistance have been tested.

Index Terms: diamond-like carbon, cathodic vacuum arc, macroparticle filtering, magnetic storage, carbon overcoats
I. INTRODUCTION

Ultrathin (< 5 nm) hard carbon films are used as protective overcoats on hard disks and read-write heads. The tribological properties of the head-disk interface are not only of mechanical but also of chemical nature: the overcoat should protect the magnetic layers against wear and corrosion [1, 2]. The areal density of information stored in disk drive products increased at an amazing rate of over 65% for many years, and continues to accelerate to even greater rates [3]. To accomplish this, the “magnetic spacing” between the magnetic layers of the disk and read/write sensor of the head must continue to decrease. The magnetic spacing includes the overcoats, lubrication, and the fly height. Thinner overcoats allow the read-write head to be closer to the magnetic layer of the disk, and hence, the size of individual bits on the disk to be smaller. As we work toward an areal density of 100 Gbit/in², the magnetic spacing approaches the sub-10-nm regime (pseudo-contact recording) and overcoat thickness must shrink to 1-2 nm. Overcoats currently used are sputtered or ion-beam deposited diamond-like carbon films, typically doped with hydrogen or nitrogen. They cease to provide acceptable levels of corrosion protection when the film thickness is less than 5 nm.

Overcoats need to be tough (hard and elastic), chemically inert, pinhole-free, and compatible with the lubricant. The challenge of ultrathin film synthesis is to make the films as thin as possible and still continuous, as opposed to an assembly of islands.

Nucleation and growth of ultrathin films is a field of intense research. For carbon, it is well known that films that are rich in sp³ (diamond) bonds should be grown in a kinetic mode at low temperature (less than 200°C, preferably room temperature), with film-forming carbon atoms or ions having an energy of about 100 eV. This energy is optimized for subplantation film growth of diamond-like carbon [4-10]. If the energy is too low, say less than 20 eV, ions do not have the kinetic energy to penetrate the surface; if the energy is too high, thermal spikes and a general enhancement of the temperature will allow the carbon atoms of the film to move to configurations of smaller energy. In both cases, the films tend to be more graphitic with characteristic sp² bonds.

Diamond-like films are characterized by an outstanding combination of advantageous properties: they can be very hard, tough, super-smooth, chemically inert, well adherent to the substrate, and compatible with lubricants. They can be deposited fast, efficiently, at low cost, and on room temperature substrates. The various deposition methods result in a variety of diamond-like films. Widely used are hydrogenated diamond-like carbon (DLC or a-C:H) [11], nitrogen-doped amorphous carbon (a-C:N) [12] or amorphous carbon nitride (CNₙ) [13-15], hydrogenated carbon nitride (CH₃Nₓ), hydrogen-free amorphous carbon (a-C) [16], silicon-doped amorphous carbon (a-C:Si) or silicon carbide (SiC) [17], and metal-doped amorphous carbon (a-C:Me) [18]. Amorphous carbon films (a-C) that have more than 80% tetrahedral (sp³) bonding are referred to as tetrahedral amorphous carbon (ta-C) [7, 9, 19], or sometimes even as “amorphous diamond.” [20, 21]

There are large variations even within each class of materials, depending on the method and parameters of deposition. Diamond-like carbon films can be synthesized by argon sputter deposition using a graphite target [12, 22], by pulsed laser ablation of graphite [19], direct ion beam deposition [23], plasma beam deposition [24], mass-selected ion beam deposition [25, 26], electron-cyclotron-resonance plasma chemical vapor deposition (ECR plasma CVD) [27], radio-frequency-CVD [11], and by filtered cathodic arc plasma deposition [6, 16, 20, 21]. Research over the last years has identified filtered cathodic arc deposition as one of the promising techniques for hard carbon films thinner than 5 nm [28, 29]. The greatest challenge is the complete removal of “macroparticles” that are generated at the cathode spot.

This paper focuses on the deposition of ultrathin amorphous carbon films using filtered vacuum-arc carbon plasma. We discuss first the generation of plasma and macroparticles as well as methods of macroparticle removal by curved filters. Particular attention is given to the “twist filter”
system which is the latest advancement in this field. In the last section, we report about tests of the carbon films made by S-filtered and twist-filtered cathodic arc systems.

II. MACROPARTICLE-FREE CARBON PLASMA

A. Cathodic arc carbon plasma

Cathodic arc plasmas are characterized by a very high degree of ionization, with multiply charged ions present [30]. Carbon plasma is dominated by ions of charge state 1+. The carbon plasma flows with a high velocity of about $2 \times 10^4$ m/s, and even somewhat greater carbon ion velocities have been measured recently [31]. Both the high degree of ionization and plasma velocity result from explosive processes at cathode spots [32]. The “vacuum” arc is a non-stationary high-pressure phenomenon driven by a very high power density of at least $10^{13}$ W/m².

Solid cathode material is transformed to dense plasma that expands while moving away from the spot area. Not only plasma is produced but also debris particles that are liquid in case of metal cathodes and liquid or solid in case of a graphite cathode [33]. These particles, usually referred to as “macroparticles” emphasizing their massive nature compared to ions and electrons, have a wide range of size distribution. For carbon, large chunks of graphite (fractions of millimeters) are sometime ejected from the cathode – it is believed that shock-heated gas inclusions in the cathode contribute to these massive cathode losses. Due to their high temperature, carbon macroparticles are easily visible when leaving the cathode spot, and macroparticle velocities of 10-370 m/s have been determined with a CCD camera [34]. Smaller macroparticles are much more frequent than µm-size particles; the size distribution ranges well down to the nanoparticle range [35]. The presence of macro- and nanoparticles is detrimental to the quality of ultrathin films and, until recently, has prevented commercial application of cathodic arc plasma deposition in some fields such as the magnetic storage industry.

B. Macroparticle filters

Many attempts have been tried to reduce, and eventually eliminate macroparticles from the plasma and coating [36-40]. The term “macroparticles” is generically used for all particles, including nanoparticles. The most successful approach to macroparticle removal is based on the vast mass-to-charge difference of macroparticles and plasma particles. The plasma can be guided out of the line-of-sight from the cathode using curved magnetic fields, employing a combined magnetic and electric mechanism for electrons and ions, respectively. The motion of macroparticles is almost not influenced by the presence of fields, thus they move along straight trajectories, thereby being separated from the plasma.

In the literature, two basic constructions and filter philosophies have been described. They can be described as closed and open. The “classic” 90° filter duct [41] employs a duct, a closed tube, surrounded a by set of magnetic field coils. The interior wall of the duct is equipped with baffles designed to catch or reflect macroparticles. While metal macroparticles tend to stick to baffles, graphite particles usually “bounce” back. Graphite particles may suffer multiple reflections and can fracture into several “sub-macroparticles.” Consequently, there is a significant likelihood that carbon macroparticles, or fractions of them, arrive at the substrate via multiple reflections. Additionally, small nanoparticles may be transported through momentum exchanged with ions (“ion wind”). There are a numerous closed filters derived from the classic 90° filter, including the 45° filter [42], S-filter [43], segmented filter [44], filters of rectangular cross section [45, 46], and the out-of-plane double-bend (OPDB) filter [47, 48].

Open filters address the issue of particle reflection by removing the particle from the filter and plasma volume via openings [49-51]. Figure 1 shows an S-shaped open filter that consists of a bent
solenoid. It is obvious that most macroparticles can leave the plasma volume, and some are removed even if they hit a turn of the magnetic field coil.

We have produced thin and ultrathin carbon films using different types of filters: a closed S-filter [43], an open S-filter, a prototype twist filter (twisted S-filter) [52], and a second-generation twist filter. The closed and open S-filters in the present experiments had the following dimensions and parameters. Closed filter: two 90° bents (in-plane), wall material stainless steel (bellows) of 0.4 mm thickness, inner diameter 2.50 inches (6.35 cm), curvature radius 10 cm, magnetic field about 45 mT; open filter: two 90° bents of open coil (in-plane), round copper wire of 2.25 mm diameter, inner diameter 5.5 cm, curvature radius 6 cm, magnetic field about 50 mT. The relatively novel twist filter is described in more detail below.

C. Twist Filter System

In recent work [52], a new open filter design was presented that is essentially a “twisted” S-shaped open solenoid, or “twist filter” for short. The prototype of the twist filter system is shown in Fig. 2; the two open 90° sections of the filter are twisted by 90°. The design is based on several paradigms:

1) The filter must be short to minimize plasma losses [49].
2) The source (and thus the cathode) must be small to be compatible with the entrance size of the filter.
3) If the (consumable!) cathode is small, it needs to be replenished via a cathode feed mechanism.
4) If the source and filter are small, heating constraints can be alleviated by operating in pulsed mode.
5) Pulsed mode operation can be performed at relatively high arc currents, leading to reasonable average plasma production, possibly with enhanced plasma properties such as slightly enhanced average ion charge state and reduced macroparticle production compared to DC operation [53].
6) In the pulsed mode, the high arc current can be used to generate a strong magnetic filter field that is synchronized with plasma production.
7) Pulsed operation can use simple and very reliable arc initiation such as the “triggerless” mechanism [54].
8) The system needs a macroparticle “firewall”, i.e. a wall separating the source and filter area from the clean substrate area.
9) The system needs a designated region in which macroparticles are collected for removal on a regular maintenance schedule. This area is located under the plasma source and filter.
10) The plasma exiting the filtered is highly focused but a uniform deposition over a relatively large area (e.g. wafer of 95 mm diameter) is required. Therefore, a plasma expansion zone and a plasma homogenizer are needed.
11) All parts of the system (source, filter, power supply, firewall, homogenizer, etc) need to be well-matched.

The Twist Filter system is a consequent realization of the design paradigms. The prototype version shown in Figure 2 consists of two 90° coil bents that are twisted by 90°. The filter’s inner diameter is 30 mm, and its outer diameter is 48 mm. The coil “wire” had a flat cross section with a thickness of 1.0 mm. The material was oxygen-free copper; this choice was made with the idea to have very good electrical conductivity, minimizing ohmic losses, and very good thermal conductivity, ensuring effective heat removal from water-cooled holding points. The filter-matching arc source did not have its own magnetic field but the filter’s fringe field was found well-suited for source-filter coupling. The distance from the cathode surface to the filter entrance was 3 cm. The source had a 1/4 inch (6.25 mm) diameter graphite rod cathode that could be advanced as the cathode is eroded. The average cathode advancement was about 0.5 μm per arc pulse, were a pulse is
typically 1.5 kA with 1 ms duration. The likelihood of macroparticle transport is greatly reduced while the plasma transport is impressive compared to other filter designs: the filtered ion current may reach up to 4% of the arc current. The coil turns have a flat cross-section promoting macroparticle reflection towards the outside of the filter volume, i.e., the coil acts as a baffle. The Twist Filter has a strong magnetic field (0.25 mT/A) providing excellent plasma confinement. At high currents (> 1 kA), not only electrons but also carbon ions are magnetized.

A second generation twist filter was installed at an R&D deposition system in a class 100 cleanroom environment. The parameters were practically the same with the following exceptions: the twist of the two 90° sections was only 45° for reasons of space constrains (see Fig. 3), the anode was built as a pre-filter with baffle structure, the maximum total cathode advancement was increased to 15 cm, the arc pulse length was increased to 1.0 ms, and the substrates were mounted on a rotating substrate holder.

Arc initiation based on the “triggerless” principle [54] was used and found to be 100% reliable due to the high open-circuit voltage used (900 V). A total of about 10^6 arc pulses have been made with the system so far without failure.

The chamber base pressure was about 10^-5 Pa. When the pulsed arc was operating with about 1 pulse per second, the pressure in the chamber initially increased to 10^-3 Pa and settled at about 10^-4 Pa after several 100 arc pulses. A residual gas analyzer showed that most of the background gas is water vapor, as it is typical for high-vacuum systems.

The plasma at the exit of the twist filter is highly focussed. Under its own pressure, the streaming plasma expands on its way to the substrate. Without special means, a deposited film will have a Gaussian thickness distribution with the thickest area in the center. A magnetic multipole, also known as a magnetic bucket, can be used to flatten the plasma density distribution leading to uniform films [55, 56]. Magnetic multipoles can be made from strong permanent magnets or by using a wire structure as shown in Fig. 4. The electromagnetic homogenizer shown in Fig. 4 was installed in the center between the twist filter exit and the substrate location. Using the homogenizer and a substrate rotation of 20 rpm., film uniformity of ±2% could be achieved over an area of 95 mm diameter for films thicker than 10 nm. Films of 3 nm showed apparently greater non-uniformity which is most likely due to the limitations of ellipsometry. One has to realize that 3% of a nominal thickness of 3 nm is 0.09 nm, that is about half the average bond length between individual carbon atoms of amorphous carbon!

Film thickness and graphite cathode rod usage were very reproducible and could be predicted simply by counting arc pulses. The pulse number – thickness relationship was about linear although we found that the output per pulse increases slightly and reproducibly during a deposition run. We attribute this to increasing cathode temperature and erosion.

The carbon ion energy was measured specifically for the second generation twist filter. The method used is a time-of-flight measurement plus a correction due to the sheath in front of the substrate. The time delay between an intentional modulation of the arc current and Langmuir probe ion current gives the ion velocity in the plasma. The Langmuir probe was located at the substrate location 220 mm from the exit of the twist filter. The total path length between the cathode surface and the Langmuir probe was 425 mm. The measured time difference between arc current maximum and ion current maximum was 18 µs, resulting in a velocity of 2.36 x 10^4 m/s or an energy of 34.6 eV. In order to estimate the plasma potential, a second measurement was done with the Langmuir probe floating. The floating potential is close to the plasma potential and easy to measure. The floating potential for a vacuum arc carbon plasma is about 4 volts negative with respect to the plasma potential [57]. With some fluctuations, the floating potential was found to be about 2 V positive with respect to ground. In this system both the anode and the substrate were grounded. Therefore, the
additional energy singly charged carbon ions gain when they are accelerated from the plasma potential to the substrate potential is about 6 eV, and thus the average total kinetic energy is about 40 eV when ions hits the substrate surface.

III. SYNTHESIS AND PROPERTIES OF a-C FILMS SYNTHESIZED BY FILTERED CATHODIC PLASMA DEPOSITION

It is known that the properties of a-C films can be tuned by carefully choosing the ion energy during deposition. Properties such as sp³ content, hardness, Young’s modulus, density, and stress have a maximum when ions have a kinetic energy of about 100 eV [6, 58-60]. While in most experiments the energy could be manipulated by substrate biasing, the second generation twist filter system had a permanently grounded, rotating substrate holder. In the latter case, the above mentioned 40 eV ion energy applies.

A. Corrosion resistance

In a first round of experiments, closed and open S-filters were used with an arc current of 300 A, 5 ms pulse duration, and 1 pulse per second repetition rate. Bursts of pulsed substrate bias were used (-100 V, 2 µs on / 6 µs off). The first 10% of the deposition process were done at a pulsed bias of –2 kV to obtain good adhesion, however, this step may not be needed for ultrathin films. The substrates for the first experiments were low-resistivity Si <100> wafers of 2.5 cm diameter that were previously coated with a magnetic layer of 100 nm permalloy (80% Ni / 20% Fe) using industry-standard magnetron sputtering. On these substrates, three sets of a-C films were fabricated using a closed S-duct filter, an open S-filter, and the second generation of twist filter. Industry-standard a-C:H films were used as reference to compare film quality; the a-C:H films were obtained by magnetron sputter deposition using a graphite target with argon plasma containing 30 at.% H2. Film thickness was measured with an n&k reflectance spectrometer and a Dektak IID profilometer.

To test corrosion resistance, the films were evaluated after immersing them for 24 h in a solution of 0.5 mol NaCl, 0.5 mol (NH₄)H₂PO₄, 1 g Liquinox, and 1 liter of deionized H₂O. The specimens were rinsed in an ultrasonic bath of deionized water and dried with nitrogen. Inspection was done under an optical microscope at 100x magnification. The defect size and density was measured using a CCD camera, frame grabber, and Media Cybernetics’ Image-Pro Plus image analysis software.

Fig. 5 shows the corrosion pinhole count versus film thickness. No significant difference in corrosion performance was noted for films thicker than 4 nm, indicating that the a-C films produced were continuous down to 4 nm thickness. The pinhole count rises an order of magnitude at thickness less than 4 nm for closed S-filter films and also significantly for films produced with the open S-filter. However, films of 3 nm thickness produced by the twist filter system still met the requirements that are normally applied to the industry-standard 7 nm a-C:H film. This shows that the twist filter system has indeed an improvement in filter capability.

In another round of experiments, focusing on twist-filter deposited carbon films, actual sliders were used as substrates to be coated with ultrathin a-C films. The sliders used in the study at UC Berkeley and IBM San Jose were commercially available pico-sliders (1 mm x 1.25 mm) made of 70% Al₂O₃ – 30% TiC material with a magnetoresistive (MR) element. In studies at Read-Rite Corporation, the sliders had dual synthetic heads, a form of the GMR (giant magnetoresistive) class of recording heads. The sliders were clamped to an aluminum substrate holder at ground potential.

For these experiments, 3 nm and 4 nm a-C coatings were deposited on sets of sliders with read/write heads. For corrosion tests, a variation of the “Battelle test” was applied: 48 h exposure at 30°C, 65% relative humidity, in the presence of 10 ppb hydrogen sulfide, 10 ppb chlorine gas, and
100 ppb nitrous oxide. As shown in Fig. 6, the sliders coated with the Twist Filter survived the corrosion test, as shown in images obtained using a Nomarsky optical microscope. Confirming the first-round dip-corrosion tests, the Battelle tests indicate that the 3 nm a-C films were continuous. The tests show that the twist filter system can indeed deliver films of 3 nm that provide corrosion protection.

**B. Hardness and elastic modulus**

Hardness and elastic modulus were measured using a force/displacement transducer coupled to an AFM. A corner of a cubic diamond tip with a nominal radius of 50 nm was used to indent the sample at loads of 1 µN to 10 mN. The measurement took indenter deformation [61] into account. The results are shown in Fig. 7. Hardness and elastic modulus decrease with decreasing film thickness, in agreement with independent measurements [6]. If the data may be linearly fitted showing the well-known proportionality $H \sim E^{\alpha}$ [62].

Nanoindentation of ultrathin films is difficult because the effect of the substrate increases with decreasing film thickness, and the indenter tip radius is much larger than the indentation depth. Other methods such as dispersion measurements of ultrasonic waves are being perfected for ultrathin films [63]. Young’s modulus measurements performed this way often give greater values than nanoindentation.

Although the hardness of the film in the sub-10-nm regime appears to be relatively low, one must be careful in equating hardness with the film’s wear resistance. Fracture toughness may be more relevant than hardness, especially in the case of sliding wear [1]. Scratch tests of ultrathin, ion-beam-deposited carbon nitride films indicate film failure by brittle fracture followed by abrasive wear [14]. Nanoindentation and nanoscratching is being used to evaluate fracture toughness of ultrathin films [11, 64-66].

**C. Drag tests in ultra-high vacuum tribochamber**

An UHV tribochamber [67] was used to detect the gaseous products generated at the head-disk interface during drag tests. A quadrupole mass spectrometer monitored 15 different species, and friction and temperature data were recorded by strain gauge transducers and a thermocouple, respectively. Background intensities were recorded before the drag tests. The drag speed was 0.2 m/s and the load 15 mN.

Initial drag tests with 5 nm cathodic-arc carbon overcoats (closed S-filter) on disks revealed promising tribo-chemical properties against uncoated Al$_2$O$_3$-TiC sliders [2]. The cathodic-arc carbon overcoat reduced the catalytic decomposition of perfluoropolyether ZDOL 2000 lubricant commonly used in disk drive applications [68]. According to the catalytic decomposition mechanism [2], hydrogen generated from the overcoat is a key component of the reaction that occurs during sliding. Since cathodic-arc carbon films have relatively low hydrogen content, the catalytic decomposition should be hindered. This was confirmed in subsequent drag tests that involved 5 nm a-C:H and cathodic-arc a-C coated disks in contact with uncoated Al$_2$O$_3$-TiC sliders [69].

**D. Measurements of the magnetically dead layer**

To investigate the dead layer thickness, a layer of 20 nm NiFe (18% Fe) was coated on Si substrate. The saturated magnetic flux density $B_s$ was measured before and after carbon deposition using a B-H looper. From the loss of $B_s$ one can calculate the approximate thickness of the dead layer. This technique was applied to a-C films synthesized with the twist filter system, and the magnetically dead layer was found to be about 0.5 nm which is close to the resolution limit of the method.
E. Hydrogen content

The hydrogen content of twist-filtered a-C film was determined by Nuclear Resonance Analysis (NRA). A silicon specimen deposited with a twist filter a-C film of about 10 nm thickness yielded the surprisingly high H-concentration of 9-11%. From other experiments we have evidence that a pulsed vacuum arc process with a strong magnetic field in a high-vacuum environment does indeed produce significant amounts of hydrogen. Schneider and co-workers [70, 71] have shown that residual water vapor can be efficiently ionized when a magnetic field is present. The hydrogen content in alumina films deposited by filtered cathodic arcs was found to be about 8% [72]. Interestingly, cathodic arc ta-C films made several years ago at Berkeley Lab with a 90° filter at much lower magnetic field strength showed only 1-3% hydrogen content. These (unpublished) measurements were made with Elastic Recoil Spectroscopy (ERS). The difference in the results could be associated with not only possibly different vacuum base pressures but also by the degree of ionization of the water vapor due to the higher magnetic field of the twist filter.

IV. CONCLUSIONS AND OUTLOOK

The development of improved macroparticle filters enables cathodic arc plasma deposition to produce thin and ultrathin films of good quality. Open filters address the reflection issue that has plagued traditional closed filter ducts. The most complete removal of carbon macroparticles was achieved by an open twisted S-filter dubbed “twist filter”. It has been shown that ultrathin a-C films down to 3 nm could meet the requirements of the magnetic data storage industry. For storage densities approaching 100 Gbit/in², current research focuses on the synthesis and analysis of even thinner films as well as on the improvement of uniformity and deposition rate.

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REFERENCES


Figure Captions
Figure 1. S-shaped open filter guiding carbon plasma from a cathodic arc plasma source (right) to the substrate region (left). Carbon macroparticles are hot and can therefore easily be identified by their bright traces. Open-shutter photograph, arc current 1000 A, pulse length 1 ms.
Figure 2. Photograph of the prototype twist filter system (two 90° sections, twisted by 90°, ID 30 mm, OD 48 mm) with its holding structure; the cathodic arc source with advancing cathode, and the arc power supply. For scale: the arc supply can be mounted in a standard 19” (48.3 cm) rack.
Figure 3. Mounting of the second generation twist filter (two 90° sections twisted by 45°).
Figure 4. A simple electromagnetic multipole or “bucket” used to flatten the plasma density distribution in the alpha-version of the Twist Filter System. As with the Twist Filter, the arc current itself was used to generate the magnetic field. The pen is for size comparison.
Figure 5. Corrosion pinhole count in an area of 0.8 x 1 mm² versus thickness of a-C films synthesized with three filter systems. The horizontal line indicates the average pinhole count for a 7-nm RF-sputtered a-C:H reference film that is considered acceptable.
Figure 6. Battelle corrosion test of 3-nm a-C coated read/write heads. Optical micrographs before (left column) and after (right column) the test. For comparison, the top four rows show heads that were coated using a DC carbon vacuum arc with a closed filter; the bottom four rows show heads coated with the twist filter system.
Figure 7. Hardness and elastic modulus of a-C films as a function of film thickness as determined by nanoindentation.
Figure 1
Figure 3

- anode/prefilter
- grounded holding plate
- twist filter
Figure 4
Figure 5

- closed S-filter
- open S-filter
- twist filter

permitted limit for 7 nm
a-C:H reference film
Figure 6

closed filter

open filter

50 μm