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Synthesis and properties of CS$_x$F$_y$ thin films deposited by reactive magnetron sputtering in an Ar/SF$_6$ discharge

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Abstract

A theoretical and experimental study on the growth and properties of a ternary carbon-based material, CS$_x$F$_y$, synthesized from SF$_6$ and C as primary precursors is reported. The synthetic growth concept was applied to model the possible species resulting from the fragmentation of SF$_6$ molecules and the recombination of S-F fragments with atomic C. The possible species were further evaluated for their contribution to the film growth. Corresponding solid CS$_x$F$_y$ thin films were deposited by reactive direct current magnetron sputtering from a C target in mixed Ar/SF$_6$ discharge with different SF$_6$ partial pressures ($P_{SF6}$). Properties of the films were determined by X-ray photoelectron spectroscopy, X-ray reflectivity, and nanoindentation. A reduced mass density in the CS$_x$F$_y$ films is predicted due to incorporation of precursor species with a more pronounced steric effect, which also agrees with the low density values observed for the films. Increased $P_{SF6}$ leads to decreasing deposition rate and increasing density, as explained by enhanced fluorination and etching on the deposited surface by a larger concentration of F/F$_2$ species during the growth, as supported by an increment of the F relative content in the films. Mechanical properties indicating superelasticity were obtained from the film with lowest F content, implying a fullerene-like structure in CS$_x$F$_y$ compounds.
1. Introduction

Carbon-based thin films with concentration of $p$-elements below 30 at.% have become increasingly attractive due to their unique mechanical and tribological properties [1] [2] [3] [4] [5] [6] [7] as well as their good biocompatibility [8] [9]. A well-known example is the discovery of C-N thin films with fullerene-like (FL) nanostructure deposited by reactive direct current magnetron sputtering (rDCMS) [10]. The CN$_x$ thin films exhibit high hardness, high resilience and low wear rate [11]. These properties are explained by buckling and cross-linking of the curved graphitic C-N planes in the FL structure, mixing $sp^2$ and $sp^3$ bonded carbon atoms [10] [12]. Following the discovery of CN$_x$ thin films, the class of FL and nanostructured C-based thin solid films grew as a result of exploration of different binary systems, such as C-P, C-F, and C-S which also provide a wide range of possibilities for tuning the mechanical, electrical, optical, and chemical properties.

Studying the growth and the properties of ternary carbon-based materials, such as CS$_x$F$_y$, is also motivated by combining the effects of both hetero-elements, e.g., the increased electron mobility and the enhanced chemical inertness of S- and F-doped carbon, respectively. The differences in the atomic radius and in the electronic structure between the heteroatoms also affect the microstructure and, thus, the mechanical properties of the material. The ternary systems, therefore, provide an extra parameter to tune the properties of the carbon-based thin films compared to the binary systems.

The experimental attempts for synthesizing the CP$_x$, CF$_x$, and CS$_x$ carbon-based thin films has been preceded by mapping out the relationship between their chemical composition and their nanostructure via computer simulations applying the synthetic growth concept (SGC). The SGC is an approach for simulation of thin film formation during vapor phase deposition based on density functional theory (DFT) [12] [13] [14] in a recursive application of candidate precursors. It has been shown that the FL structure can be obtained in all of the above mentioned systems,
but within different ranges of heteroatom concentration (up to: 30 at.% N [12], 15 at.% P [13], 15-18 at.% S [15], or 8-12 at. % F [16] [17]). This variation between the compounds is mainly explained by differences in the number of valence electrons, the electronegativity, and the steric repulsion when introducing heteroatoms belonging to different chemical elements into C-basal planes as in graphite or graphene. For example, P and S belong to the third period of the periodic table thus possessing larger covalent radii in comparison to C and N (which belong to the second period), leading to larger energy costs when the heteroatoms substitute a carbon atom in the hexagonal network [18] [19]. Fluorine, on the other hand, does not bond to more than one C atom and therefore tends to introduce severe disruptions into the network [16]. Moreover, its three other non-bonding valence electron pairs cause strong repulsion on other atoms, making the network to distort in out-of-plane directions. Thus, higher concentration of F atoms in the system introduces higher density of large-scale defects (e.g., larger rings in the network) and consequently results in different nanostructures compared to graphene-like or FL kinds of nanostructure, such as polymer-like CF$_x$[17].

While the theoretical understanding of the structure evolution with respect to the chemical composition was achieved for some compounds, a wide range of structural and mechanical relations of carbon-based thin films remain to be investigated. The diversity of possible process techniques/conditions also complicate the prediction and explanation of the nanostructure evolution. This implies that cases of carbon-based nanostructured films within certain range of chemical compositions should be considered in the context of the deposition process parameters. For example, the ratio between $sp^2$ and $sp^3$ bonded carbon atoms in thin films has been shown to be a function of the energetic ion bombardment and the growth temperature [20] [21] [22] [23] [24].

It is also noteworthy that the source of the heteroatom species can influence the formation of the structure in deposited thin films. As corroborated by SGC simulations, Furlan et al.
observed intercalation of phosphorous between graphite-like sheets in CP \(_x\) \((x = 0.1 \text{ and } 0.24)\)
thin films deposited by DCMS [25] [26], as \(P_n\) clusters \((n\) is normally 2 or 4) can be incorporated
through evaporation from the surface of the compound target instead of pure sputtering
deposition [27]. Furthermore, Schmidt et al. reported that the mechanical properties can be
different for CF \(_x\) \((0.15 \leq x \leq 0.26)\) thin films deposited by reactive high power impulse
magnetron sputtering (HiPIMS) in Ar/CF \(_4\) and Ar/C\(_4\)F\(_8\) discharges, respectively [28]. An
extensive SGC-based theoretical investigation was carried out for the same compounds
indicating that different dissociation behaviors of the reactive gas species lead to different
bonding conditions for C-atoms, thus defining different structure and properties.

Previously, the ternary FL structure was predicted by SGC simulations as feasible to be
obtained in the C-S-F system for \(([S] + [F]) < 10\) at.\%. Meanwhile, indications of graphitic
structure were observed in the CS\(_x\)F\(_y\) thin films with the mentioned range of heteroatom
concentrations deposited by rDCMS with a mixed Ar/SF\(_6\) discharge [29]. Nevertheless, it
remains to be determined how SF\(_6\), as the primary precursor, affects the growth of CS\(_x\)F\(_y\) thin
films, and resulting physical and mechanical properties. Also, even though the SF\(_6\) gas is a
common dry etchant for Si-based (e.g., Si [30] and SiC [31]) or metallic (e.g., W [32]) materials
in plasma processes, this gas is less reported as a single source for both F and S species in
reactive deposition. Thus, studies dedicated to the precursor’s dissociation and reaction
behaviors, as well as the relationship between the obtained properties in the CS\(_x\)F\(_y\) films and
the process parameters, are needed in order to understand the formation of the C-S-F networks
in the reactive deposition process.

Here, theoretical calculations were carried out for studying the dissociation of SF\(_6\) gas into S-
F fragments and the following formation of different C-S-F precursor species in a C target
magnetron sputtering process. Also, CS\(_x\)F\(_y\) thin films were deposited at three different SF\(_6\)
partial pressures, \(P_{SF6} = 1.0\) mPa, 1.5 mPa and 2.0 mPa. Their chemical composition was
quantified by X-ray photoelectron spectroscopy (XPS). The deposition rate and density of the samples were acquired from X-ray reflectivity (XRR) patterns while their mechanical properties were studied by nanoindentation. The effects of SF$_6$, in terms of its partial pressure during depositions, on the microstructure of the CS$_x$F$_y$ samples were qualitatively elaborated based on the theoretical precursor studies and the measured properties.

2. Computational Details

The SGC [12] [13] [33] was used to model the precursor species most relevant to film formation during vapor phase deposition. The C$_p$S$_q$F$_r$ ($p, q \leq 2, r \leq 6$) species were studied by geometry optimization and calculation of the corresponding cohesive energy. The cohesive energy per atom ($E_{coh/\text{at.}}$) is defined as the energy needed to break the system into isolated atoms normalized by the total number of atoms, and is calculated according to:

$$E_{coh} = \frac{E_{total} - \sum N_j E_j}{\sum N_j}$$

, where $E_{total}$ is the total energy of the relaxed species, $E_j$ and $N_j$ correspond to the energy of the isolated atoms and the total number of atoms for different species $j = \{C, S, \text{and} F\}$.

The framework adopted for the calculations is density functional theory (DFT) within the generalized gradient approximation (GGA) as implemented in Gaussian 09 [34]. The results reported here were obtained using the Perdew-Wang exchange and correlation functional (PW91) [35] with the 6-31G** basis set. These choices have been shown to provide an accurate description for similar systems [36] [37].
3. Experimental Details

3.1 Reactive sputtering of thin films

CS$_x$F$_y$ thin films were synthesized onto Si(001) substrate by rDCMS in a high vacuum system with a base pressure below 0.1 mPa, as the system previously described in Ref. [25] and [26]. All depositions were carried out with a 2-inch C target (99.9% purity, pyrolytic graphite with density $\sim$1.8 g/cm$^3$) attached to a type II unbalanced magnetron operated at a constant power of 90 W, a deposition time of 120 min, and floating potential to the substrates. The magnetron was placed at a distance of 12 cm right below a substrate holder. The Si substrates were cleaned sequentially in acetone, ethanol, and isopropanol ultrasonic baths for 10 min. each, then blown dry with N$_2$ gas. Prior to depositions, the base pressure of the system was throttled up to 0.5 mPa and the reactive gaseous precursor SF$_6$ (99.993 % purity, Solvay Fluor GmbH) was supplied from a pressurized gas bottle through a leak valve up to the desired $P_{SF6}$, followed by additional Ar gas filled up to a working pressure of 533 mPa (4 mTorr). The CS$_x$F$_y$ samples were deposited at 350 °C and at three different SF$_6$ partial pressures: one sample at $P_{SF6} = 1.0$ mPa and 1.5 mPa, and two samples at $P_{SF6} = 2.0$ mPa.

3.2 Sample characterization

3.2.1 X-ray photoelectron spectroscopy (XPS)

The XPS were performed with the AXIS Ultra$^{\text{DLD}}$ system from Kratos Analytical Ltd. using monochromatic Al K$_\alpha$ radiation. Relative chemical composition of C, S, F, and O in the thin films was obtained by elemental quantitative analysis based on the peak areas acquired from XPS high resolution scans of the C1s, S2p, F1s, and O1s regions together with the elemental sensitivity factors, provided by Kratos Analytical Ltd. In order to reduce surface contamination, the samples were directly transferred from the deposition system and into the XPS system with an atmosphere exposure of only about 10 min. No sputter cleaning on the film surface was
performed before acquiring the XPS spectra, thus minimizes damage to the chemical bonding structure of the films.

3.2.2 X-ray reflectivity (XRR)

The XRR patterns of the films were acquired in an X-ray diffractometer (Empyrean, PANalytical B.V.) using CuKα radiation. The films were aligned for the height at 0° and for the tilt angle at 0.5° to improve the accuracy of the measurements. The acquired patterns were thereafter fitted by PANalytical X’Pert Reflectivity program with a single carbon layer on silicon substrate model in order to obtain the thickness and the density information of the films. This method has been used in Ref. [24] to determine the density of the CNx thin films. The film thickness is thereafter divided by the total deposition time (120 min.) giving the deposition rate of the samples.

3.2.3 Nanoindentation

Nanomechanical properties of the films were evaluated by quasi-static displacement-controlled nanoindentation experiments in a TI950 TriboindenterTM (Hysitron). Hardness ($H$) and reduced modulus ($Er$) measured with a Berkovich diamond probe with a blunt-ended tip curvature of radius of ~100 nm were calculated according to the method proposed by Oliver and Pharr [38]. The tip function area at low penetration depths was calibrated using a fused silica sample. In order to avoid substrate effects, penetration displacement was set to 10% of the total coating thickness. In all tests a total of 12 indents with a spacing of 10 μm were averaged to determine the mean value and standard deviations of $H$ and $Er$. 


4. Results and Discussion

4.1 Theoretical study on precursor species

Precursor species C_pS_qF_r (p, q ≤ 2; r ≤ 6) that are relevant to the here reported deposition process, (a graphite target and SF_6 reactive gas), were modeled in order to define their possible role during film formation of C_S_xF_y. These species are expected to originate from the fragmentation of the SF_6 molecule,

\[ \text{SF}_6 \rightarrow \text{SF}_r + (6 - r)\text{F}, \]  
\[ \text{(R1)} \]

and from recombination reactions in the plasma or at the target surface,

\[ P + P' \rightarrow PP', \]  
\[ \text{(R2)} \]

where both P and P’ are the precursor species C, S, F or SF_r fragments, and PP’ is a recombined species. For this particular study only those recombination reactions involving one collision were taken into account.

For all composition of precursor species considered here, alternative geometric isomers were taken into account in order to find the most stable precursor geometries. The following groups of possible precursor species were optimized, compared, and considered in the context of film formation:

(i) species resulting from the fragmentation of the SF_6 molecule: SF_5, SF_4, SF_3, SF_2, and SF (as in reaction R1);

(ii) dimers resulting from the recombination of two single atomic species C, S, and F, i.e., C_2, S_2, F_2, CS, CF, and SF (as in reaction R2);

(iii) medium-size species originating from the recombination of a single atomic species and a SF_r fragment: C–SF_r, S–SF_r, and F–SF_r (as in reaction R2);
(iv) larger species originating from the recombination of two equal or different SF$_r$ fragments: (SF$_r$)–(SF$_r'$) (as in reaction R2).

It is noteworthy that SF and F–SF$_r$ from groups (ii) and (iii), respectively, are resulting in the same species listed in group (i) plus SF$_6$ molecule, and hence they will be denoted below as SF$_r$. The resulting stable structures for the most relevant precursor species are shown in Fig. 1 and Fig. 2, and their cohesive energies are listed in Table 1 and Table 2.

**Figure 1.** The most stable geometries for SF$_r$ species resulting from the fragmentation of the SF$_6$ molecule.
Figure 2. The most stable geometries for possible precursor species resulting from recombination reactions in the plasma, that is likely to play a significant role in the film formation process. For the FSSF precursor species, the dihedral angle (FSSF) is 89°.

Table 1. $E_{\text{coh/at}}$ for the reactive gas molecule SF$_6$ and the possible precursor species resulting from its fragmentation, also shown in Fig 1.

<table>
<thead>
<tr>
<th>Species</th>
<th>SF$_6$</th>
<th>SF$_5$</th>
<th>SF$_4$</th>
<th>SF$_3$</th>
<th>SF$_2$</th>
<th>SF</th>
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<tbody>
<tr>
<td>$E_{\text{coh/at}}$ (eV/at)</td>
<td>-3.15</td>
<td>-2.94</td>
<td>-3.08</td>
<td>-2.80</td>
<td>-2.70</td>
<td>-2.00</td>
</tr>
</tbody>
</table>

Table 2. $E_{\text{coh/at}}$ for possible precursor species resulting from recombination reactions in the plasma, that is likely to play a significant role in the film formation process, also shown in Fig. 2.

<table>
<thead>
<tr>
<th>Species</th>
<th>CS</th>
<th>CF</th>
<th>C$_2$</th>
<th>F$_2$</th>
<th>S$_2$</th>
<th>S$_2$F</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_{\text{coh/at}}$ (eV/at)</td>
<td>-3.82</td>
<td>-3.21</td>
<td>-2.36</td>
<td>-1.29</td>
<td>-1.85</td>
<td>-2.66</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Species</th>
<th>S$_2$F$_2$</th>
<th>FSSF</th>
<th>CSF</th>
<th>CSF$_2$</th>
<th>CSF$_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_{\text{coh/at}}$ (eV/at)</td>
<td>-2.85</td>
<td>-2.83</td>
<td>-3.06</td>
<td>-2.86</td>
<td>-2.59</td>
</tr>
</tbody>
</table>
The role of these precursor species during film formation was evaluated according to their stabilities, relative size, and accessible dangling bonds. For example, those relatively large species (e.g., SF₅) were considered to have reduced contribution to the film formation, seen from their complete electron shells and relatively high number of bonded F atoms screen the valence electrons of C and S atoms that are responsible for interacting with the dangling bonds on the growing film surface. Consequently, these larger species become geometrically less accessible to bond to the surface. From these bonding conditions, we envisage the following situations in order of increasing importance for film formation:

1. (SF₇)–(SF₇') fragments (r, r' = 2 – 5) should have a reduced to no impact on the film formation process;

2. X–SF₅, X–SF₄, SF₅, and SF₄ (X = C or S) show a low number of dangling bonds in relation to their size, presenting lower likelihood of bonding onto the growing film surface;

3. X–SF₃, X–SF₂, SF₃, and SF₂ (X = C or S) show higher chances of bonding compared to (2) but represent medium size species that will sterically prevent other precursor species from bonding in their vicinity. The prevalence of these species hints to the formation of films with low density;

4. FS–SF, X–SF species (X = C or S), dimers and single atomic species are the most significant precursor species for growing well-structured and dense films due to their geometrical availability of dangling bonds and their smaller size.

Our evaluation shows that the precursor species with medium and smaller relative size, i.e., group (3) and (4), can contribute to most of the film growth, while the larger precursor species, i.e., group (1) and (2), have a much reduced probability to be deposited. Furthermore, properties of deposited CSₓFₘ, e.g., mass density and chemical composition, can be affected by relative
amount of the above classified groups of precursor species created during the growth due to the steric effect of differently sized species on the surface.

4.2 Synthesis and characterization of CSₓFᵧ thin films

Solid CSₓFᵧ thin films were prepared by rDMCS in an Ar/SF₆ discharge using three different P_{SF₆}. Fig. 3(a) summarizes the XPS quantified F and S atomic content, [F] and [S] respectively, of the CSₓFᵧ samples against the respective P_{SF₆} used during deposition. The similar chemical composition of the samples from repeated deposition at P_{SF₆} = 2.0 mPa suggests that the CSₓFᵧ thin film deposition by reactive sputtering with SF₆ gas is a reproducible process. The XPS spectra of C 1s, F 1s, and S 2p acquired from the samples are shown in Fig. S1 – S3 in the supplementary material. All samples have a close to constant O content in a range of 1.0 – 1.3 at.%. Notice that this [O], determined by XPS, is likely overestimated due to post-deposition oxidation on the film surface following a short exposure to air during sample transfer.

![Figure 3](image)

**Figure 3.** (a) The F and S atomic content ([F] and [S], respectively) and (b) the [F] / [S] ratio of the CSₓFᵧ samples plotted against P_{SF₆}. The dashed line in (b) indicates the [F] / [S] ratio of a SF₆ molecule. When P_{SF₆} = 2.0 mPa, solid and hollow symbols are used to distinguish the two samples.
The increment in concentration of both [F] and [S] from $P_{SF6} = 1.0 \text{ mPa}$ to 2.0 mPa suggests that the amount of the SF$_6$ partial pressure has a positive effect on incorporating both S and F atoms into the CS$_x$F$_y$ films. Individually, a steeper increment of [F] is observed between $P_{SF6} = 1.5 \text{ mPa}$ to 2.0 mPa compared to the [F] increment between $P_{SF6} = 1.0 \text{ mPa}$ to 1.5 mPa. In contrast, the amount of [S] in the films does not show a significant change after $P_{SF6} = 1.5 \text{ mPa}$.

The individual behaviors of F and S atoms in the reactive process are more visible by plotting the F-to-S atomic ratio of the samples against the SF$_6$ partial pressure, as shown in Fig. 3(b). In the figure, larger deviations from the chemical formula of the SF$_6$ precursor, where [F]/[S] = 6, are observed for the samples deposited at $P_{SF6} = 2.0 \text{ mPa}$ than those deposited at lower $P_{SF6}$. The increment in this composition ratio suggests a change in the behavior for the F- and S-containing precursor species to a regime permitting the former to remain in the films more abundantly, thus making higher and higher amount of F atoms to be incorporated during the deposition at higher $P_{SF6}$.

On one hand, such change of incorporation behavior can be explained by the differences in the sticking coefficient and in the dissociation behavior of the precursor species. According to the precursors’ evaluation in Section 4.1, the larger size species with less geometrically accessible dangling bonds, i.e., $X$–SF$_r$ and SF$_r$ ($r \geq 4$) in group (1) and (2), should be more easily impeded to stick to the surface, while the S-containing medium and smaller size species, i.e., $X$–SF$_r$ and SF$_r$ ($r \leq 3$), FS–SF, S$_2$, and S in group (3) and (4), are energetically less favorable to create [39]. It has been experimentally demonstrated that the most abundant species from SF$_6$ discharges are the neutral or ionized molecules of F, F$_2$ and SF$_r$ fragments ($r \geq 3$) [40] [41]. Hence, when depositing at higher $P_{SF6}$, the F$_2$, CF, and F species become relatively more abundant and, thus, more dominating in the growth than S-containing groups.
On the other hand, a more pronounced reactive etching process is expected on the deposited surface during growth at higher $P_{SF6}$, where the F/F$_2$ species, created by the fragmentation of SF$_6$ molecules in a plasma process, become more abundant in the deposition ambient [42] [43]. Exposing C-based materials to such environment leads to both fluorination and etching at the surface, while both reactions are enhanced as the partial pressure of SF$_6$ increases [44] [45]. As discussed in Section 4.1, the precursor species $X$–SF$_r$ and SF$_r$ ($r = 2$ or 3) from group (3) attached to the surface can locally reduce the density due to a more pronounced steric effect than group (4) species. Hence, the close-by regions are more permeable to the primary oxidants, F and F$_2$, and thus more susceptible to etching. As $P_{SF6}$ increases and the etching process is promoted, more group (3) species (than only F-containing species due to generally stronger bonded F atoms) are removed from the surface and consequently the [F]/[S] ratio raises.

The significant role played by the etching effect, is supported by a decrease of the deposition rate. In Fig. 4, the total thickness of the films determined by XRR is 225 nm, 190 nm and 164 – 165 nm for the samples deposited at $P_{SF6} = 1.0$ mPa, 1.5 mPa, and 2.0 mPa, respectively, which is corresponding to a decreasing deposition rate from 1.88 nm/min., 1.58 nm/min., down to 1.37 – 1.38 nm/min.

In addition to the deposition rate, the shift towards higher critical angle in Fig. 4 indicates an increment in the respective mass density, $D_x$, of the samples from 1.5 to 1.9 g/cm$^3$ along with increasing $P_{SF6}$ from 1.0 mPa to 2.0 mPa. The mass density of the CS$_x$F$_y$ samples is considered lower than the commonly reported values for graphite (2.3 g/cm$^3$) [46], for tetra-amorphous carbon (1.8 – 2.2 g/cm$^3$) [47], and for CF$_x$ thin films (~2.1 g/cm$^3$) with close chemical compositions ([F] < 26 at.%) deposited in mixed Ar/CF$_4$ and Ar/C$_4$F$_8$ discharges [28] [48].
Figure 4. The XRR patterns around the critical angle of the CS$_x$F$_y$ samples. The critical angles of the films and of the Si substrates are indicated with respective dashed lines.

The low mass density is similar to reported polymer-like C-based thin films, but only when the content of heteroatom species is above 25 to 30 at.% [28] [49]. Contrarily in this work, such low mass density (1.5 – 1.7 g/cm$^3$) of the samples deposited at low $P_{SF6}$ is explained by the incorporation of the medium size species $X$–SF$_r$ and SF$_r$ ($r = 2$ or 3), i.e., group (3) species in Section 4.1, which hinder the formation of dense structure in the films. These low density regions are more efficiently removed due to the promoted selective etching at higher $P_{SF6}$ (as explained in the previous paragraph) leading to more densified thin films (1.8 – 1.9 g/cm$^3$).

The nanoindentation load-displacement curves for the three samples of CS$_x$F$_y$ thin films (obtained at $P_{SF6} = 2.0$ mPa, 1.5 mPa, and 1.0 mPa, respectively) are presented in Figure 5(a). Changes on the mechanical properties caused by introducing S and F heteroatoms into the C network can be observed by comparing the shape of the curves with curves reported previously.
for pure C sample deposited at a similar set-up in the same deposition system. See Fig 3(b) of Ref. [25] where the pure C sample exhibits a recovery value of ~50 %, which is significantly lower than the recovery values of the CS$_x$F$_y$ samples in this work (> 75 %).

The hardness and the reduced elastic modulus of films determined by nanoindentation are summarized in Figure 5(b), where the values for the samples from repeated depositions at $P_{SF6}$ = 2.0 mPa are, again, in good agreement with each other. Both $H$ and $Er$ values follow a similar trend, which increases from $P_{SF6}$ = 1.0 mPa to 1.5 mPa and then decreases when $P_{SF6}$ further increases to 2.0 mPa, with the minimum values observed at the lowest SF$_6$ partial pressure. However, the behavior of the $H$ and $Er$ as a function of the $P_{SF6}$ is not monotonous as is the case with the chemical composition, the deposition rate, and the density.

**Figure 5.** (a) Nanoindentation load-displacement curves of CS$_x$F$_y$ thin films. (b) The hardness ($H$, squares) and the reduced elastic modulus ($Er$, circles) of CS$_x$F$_y$ thin films plotted against $P_{SF6}$. When $P_{SF6}$ = 2.0 mPa, solid and hollow symbols are used to distinguish between two samples, coupled to Figure 3.
The lower $H$ and $Er$ values of the sample deposited at $P_{\text{SF6}} = 1.0$ mPa, which possesses an overall heteroatom content of less than 5 at.%, are possibly due to its underdense structure. The enhancement of the mechanical properties between $P_{\text{SF6}} = 1.0 – 1.5$ mPa can therefore be attributed to the structural evolution of the film due to the removal of the medium size species $X$–SF$_r$ and SF$_r$ ($r = 2$ or $3$) from the film surface by the etching effect.

As shown by theoretical modelling, the incorporation of F and/or S atoms in the network introduces local distortions, consequently enhancing the cross-linking and buckling between neighboring hexagonal (originally graphene-like) networks [15] [17] [29]. In Ref. [29], an increasing curvature of a planar carbon network is observed, when the ([F] + [S]) content is increased from 4.8 at.% to 7.1 at.%. For compounds with ([F] + [S]) $\leq$ 10 at.%, the incorporated defects in the graphene-like network are predicted to have positive effects on their mechanical properties. This discussion corroborates the explanation for the improvement on the mechanical properties of the samples between $P_{\text{SF6}} = 1.0 – 1.5$ mPa.

When an even higher amount of heteroatoms is incorporated, corresponding to the transition from $P_{\text{SF6}} = 1.5$ mPa to 2.0 mPa, $H$ and $Er$ decrease even if higher mass density values were observed. Such results are explained by the fragmentation of the highly fluorinated networks due to the termination of, mainly, F atoms. Theoretical calculations have shown that a more disrupted and strongly distorted network is observed for values of ([F] + [S]) = 9.5 at.% or higher, especially when [F] > [S] (in contrast to the cases featuring lower heteroatom content (4.8 and 7.1 at.%) [29]. Consequently, it is reasonable to expect more defective structure when [F] > 18 at.% and [F] + [S] $\approx$ 20 at.%, and a degradation in the hardness and elasticity of the samples grown at highest SF$_6$ partial pressure. Exhibiting a very similar trend, in Ref. [28], the binary CF$_x$ thin films deposited with reactive gases of CF$_4$ or C$_4$F$_8$ soften when the F content increases above 15 at.%.
Despite that [F] for all samples considered in this work is significantly lower than its observed threshold for transition to polymer-like structure ([F] > 26 at.% reported in Ref. [28], both the values for $H$ (3 – 6 GPa) and $Er$ (15 – 65 GPa) belong to a similar range as in the case of the binary polymer-like $CF_x$ samples [28]. Again, this can be perceived as resulting from a less dense microstructure caused by the incorporation of the group (3) precursor species, which is also consistent with previous observation on the low mass density. In addition to the precursor’s effect, an inherent lack of energetic ion bombardment in DCMS can not only cause under-dense microstructure [50], but also leads to less $sp^3$-hybridized bonding between C atoms [22]. Hence, in general softer films are deposited by DCMS as compared to films deposited by, for example HiPIMS, as in Ref. [28].

The hardness-to-modulus ratio ($H/Er$), sometimes referred as the plasticity index of a material, is calculated to be 0.20, 0.09, and 0.15 for the samples deposited at $P_{SF6} = 1.0$ mPa, 1.5 mPa and 2.0 mPa, respectively. A material is considered to be more elastic and compliant when its ($H/Er$) ratio is higher. For example, ($H/Er$) ratio of an elastic material, such as diamond-like carbon, is commonly found to be about 0.1, while for a super-elastic material, such as FL-CN$_x$, values in the range of 1.5 – 1.9 have been reported [51]. Due to the dominating fluorine content to the sulfur content of the samples in this work, it is logical to compare their properties to those in another chemically similar binary system, i.e., C-F system. It turns out that the $CS_xF_y$ samples grown at higher $P_{SF6}$ partial pressures ($P_{SF6} \geq 1.5$ mPa) possess ($H/Er$) values comparable to those of $CF_x$ thin films with similar heteroatom content (10 – 30 at.%), e.g., of samples grown by HiPIMS (0.11 – 0.12) [28] and grown by chemical vapor deposition (~ 0.1) [52].

Additionally, a higher compliance, ($H/Er$) = 0.19, of the sample deposited at $P_{SF6} = 1.0$ mPa is obtained compared to the other samples in this work as well as the samples reported in Ref. [28] and [52]. Interestingly, this compliance value is comparable to that of the $CN_{0.13}$ thin film reported in Ref. [51], where the authors claimed superelasticity of that sample due to its FL-
structure. The similarity in their unique plasticity index may indicate the possibility for the existence of FL-structure (or at least close-order structure) in the \(\text{CS}_x\text{F}_y\) thin film deposited at \(P_{\text{SF}_6} = 1.0\) mPa. Such result is in agreement with the previous theoretical predictions on obtaining FL-structure in \(\text{CS}_x\text{F}_y\) samples with \([\text{F}] + [\text{S}] < 10\) at.\% and the experimental observation of graphite-like feature in the \(\text{CS}_x\text{F}_y\) thin film with \([\text{F}] + [\text{S}] \sim 3.8\) at.\% [29].

It should be noted that the mechanical properties of the sample deposited at \(P_{\text{SF}_6} = 1.0\) mPa are significantly inferior to those typically observed in FL thin films such as FL-CN\(_{0.13}\) from Ref. [51]. As discussed above, such inferior mechanical properties are due to the formation of under-dense microstructure at lower \(\text{SF}_6\) partial pressure. Our earlier transmission electron microscopy studies of the \(\text{CS}_x\text{F}_y\) sample with \([\text{F}] + [\text{S}] \sim 3.8\) at.\% showed a corresponding lack of long-range order [29]. In other words, the fullerene-likeness, i.e., the scale of FL-structural patterns, is reduced due to the incorporation of the relatively massive S-containing species in the \(\text{CS}_x\text{F}_y\) thin film.

5. Conclusions

Growth of the ternary carbon-based \(\text{CS}_x\text{F}_y\) thin films by rDCMS using \(\text{SF}_6\) as reactive gas was studied by characterizing the properties of the films and by evaluating the interaction of the predicted C-S-F reactant species with the structure formation of \(\text{CS}_x\text{F}_y\). The theoretical evaluation suggested that the film growth is mainly from the interaction of medium size (\(X-\text{SF}_r\) and \(\text{SF}_r; X = \text{C or S}; r = 2\) or \(3\)) and smaller size (\(\text{C-SF}, \text{S}_2\text{F}, \text{FS-SF}\), dimers, and atoms) species with more geometrically accessible dangling bonds to react with the surface. In addition, the medium sized species are likely to introduce under-dense structure to the surface due to the more pronounced steric effect, which is in a good agreement with experimentally observed low mass density values of the \(\text{CS}_x\text{F}_y\) thin films. The under-dense surface is expected to allow more effective fluorination or etching from \(\text{F}_2\), \(\text{CF}\), and \(\text{F}\) species, indicating by the increments in the
mass density and the decreased deposition rate when increasing $P_{SF6}$. Consequently, the under-dense structure and high amount of F-termination result in low values of the hardness and elastic modulus of the films deposited at low and high $P_{SF6}$, respectively. The mechanical compliance values indicate that most of the CS$_x$F$_y$ thin films are highly elastic, and especially that the sample with the lowest heteroatom content exhibits superelastic behavior, which is a characteristic of carbon thin films with the FL structure. The changes in micro- and the nanostructure, and consequently the properties, over a small $P_{SF6}$ change (1.0 mPa) indicate a strong dependence of the CS$_x$F$_y$ thin film formation on the SF$_6$ partial pressure and the dissociation characteristic of SF$_6$.

**Acknowledgments**

References


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Supplementary Material to

Synthesis and properties of CS$_x$F$_y$ thin films deposited by reactive magnetron sputtering in an Ar/SF$_6$ discharge

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XPS spectra of C 1s, S 2p, and F 1s regions from the CS$_x$F$_y$ thin film samples

Fig. S1 shows XPS spectra acquired around C 1s region of the CS$_x$F$_y$ thin films deposited with $P_{SF_6} = 1.0$ mPa, 1.5 mPa, and 2.0 mPa (from top to bottom of the figure). The most prominent features are marked with numbers (1) – (5) respectively from low to high binding energy (BE). The two samples grown at the same condition, i.e. with $P_{SF_6} = 2.0$ mPa, have very similar features in the C 1s spectra with each other, in consistence with other analysis techniques. The feature (1) at BE ~ 284.5 – 284.7 eV can be identified as the most characteristic C – C $sp^2$ component, which is shifting towards higher BE when increasing $P_{SF_6}$. This can be related to the increment of overall F content in the material and consequently the C – C $sp^2$ component is more affected by surrounding C – F bonds. The position of the feature (1) in the samples grown at $P_{SF_6} = 2.0$ mPa is close to the position reported by Schmidt et al. in the CF$_x$ samples with high F content (> 15.5 at.%) [1]. In addition, the low degree of asymmetry for (1) indicates a low contribution from feature (2), which can be assigned to C – C $sp^3$ component at BE ~ 285.9 eV, as compared to Ref. [1]. It is reasonable that in the growth condition lacking of energetic ion bombardment, e.g. in DCMS, the content of C – C $sp^3$ bond is low in the samples, as compared to growth condition utilizing high substrate bias or HiPIMS. The features (3) – (5) can be assigned to the components of C – F/C – CF$_2$ at ~ 287.2 eV, C – F$_2$/ C – CF$_3$ at ~ 290.0 eV, and C – F$_3$ at ~ 292.2 eV, respectively, and line up well with the same components reported in Ref. [1]. The increase in (3) – (5) and the decrease in (1) indicate that, upon increasing $P_{SF_6}$, more F atoms are bonded to C atoms in the network, inhibiting the formation of C – C $sp^2$ bond.

Due to similar electronegativity for C and S atoms, the C 1s feature is less affected by introducing S atoms into the C network. It has been reported by Chang that the C – C $sp^2$ component is located at BE ~ 284.3 – 284.5 eV in
C-S compound samples with S > 24 wt.% (~ 16 at.%) [2]. In our case, the S content is less than 1.5 at.% in all samples, and hence the effect of S atoms on C 1s components is possibly not distinguishable.

**Figure S1.** XPS spectra of the \( \text{CS}_x \text{F}_y \) samples acquired at C 1s regions. The main features are marked with numbers (1) – (5) and respective dashed lines.
Fig. S2 shows XPS spectra acquired around F 1s region of the CS$_x$F$_y$ samples deposited with $P_{SF6} = 1.0$ mPa, 1.5 mPa, and 2.0 mPa (from top to bottom of the figure), where the most prominent features are again marked with numbers (1) – (2) respectively from low to high BE. The main F 1s feature, marked as (1), is located at BE $\sim$ 687.0 – 687.2 eV, shifting towards higher BE and increasing in intensity along with increasing $P_{SF6}$. The position of (1) is similar to published values for covalent C – F bonds as in the organic compounds, while no sign of more ionic F bond with BE < 685 eV is observed [3]. Similar to the case in C 1s, the shift of (1) towards higher BE is related to the effect of higher F content in the material, and therefore a higher number of adjacent C – F bonds. The feature at higher BE side, marked as (2), is assigned to the components of perfluorinated groups or alternatively plasmon effects from (1) [3].
Figure S3. XPS spectra of the CS$_x$F$_y$ samples acquired at S 2p regions. The main features are marked with numbers (1) – (3) and respective dashed lines.

Fig. S3 shows XPS spectra acquired around S 2p region of the CS$_x$F$_y$ samples deposited with $P_{SF6} = 1.0$ mPa, 1.5 mPa, and 2.0 mPa (from top to bottom of the figure), where the most prominent features are again marked with numbers (1) – (3) respectively from low to high BE. Features (1) and (2) can be assigned to S 2p$_{3/2}$ at BE $\sim$ 163.8 eV and S 2p$_{1/2}$ at BE $\sim$ 165.0 eV, respectively for S bonded in C-S compounds [2]. No indication in the S 2p region shows other form of S in the samples, such as metal sulfides (160 – 163 eV), sulfates (168 – 170 eV), or elemental sulfur ($\sim$163.1 eV) [2] [4].
References


