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Structural characterization of Fe–C coatings prepared by reactive triode-magnetron sputtering

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Fe_{1-x}C_x coatings were synthesized by triode magnetron sputtering of an iron target in a methane/argon atmosphere with a large range of composition (x = 0.3 to 0.6 ± 0.06). Film surfaces were characterized by grazing incidence x-ray diffraction, scanning and transmission electron microscopies, and electron energy loss spectroscopy, to study effects of the variation of the methane gas flow rate on their structural properties. The coatings were constituted of the ε -Fe₃C carbide (x = 0.3 and 0.36), in which carbon atoms are in octahedral sites, and of nanocomposite structure constituted of disordered and crystalline carbide nanograins embedded in a carbon matrix made of an amorphous and poorly crystallized graphenelike material (x = 0.55 and 0.60). In situ annealing of the nanocomposite Fe_{0.45}C_{0.55} coating led to the formation of carbides θ -Fe₃C and Fe₇C₃ (with carbon atoms in prismatic sites) and C-rich cubic carbide possibly related to the τ_2 -Fe₂C₇ compound.

I. INTRODUCTION

Iron carbides have been intensively studied over the past several decades due to their mechanical, tribological, and magnetic properties.^{1–3} For several years, iron/carbon and iron carbides/carbon composite materials have also presented a strong interest for their magnetic properties like giant magnetoresistance for recording media applications.^{4–6} In these materials, ferromagnetic particles are embedded in carbon, which provides protection against outside degradation like oxidation or corrosion, and magnetic insulation between particles.^{4,6} Sputtering of a composite target in a neutral atmosphere, or of a metallic target in a reactive atmosphere, is a powerful process to produce metal/nonmetal films with a large range of composition.⁷ New iron carbon materials can be synthesized by this technique. As-sputtered states and film composition are strongly dependent on the elaboration parameters, like substrate temperature, working pressure, or composition of the composite target. Therefore, crystalline phases, like α -Fe(C), ϵ -Fe₃C, θ -Fe₃C, or χ -Fe₅C₂ compounds,^{3,8,9} amorphous³ or composite films can be prepared with the possibility to act on their structuration.

Kazama et al.¹⁰ first reported the formation of amorphous $\operatorname{Fe}_{1-x}C_x$ films (0.25 $\leq x \leq$ 0.50) deposited by conventional diode sputtering of Fe₂C/Fe₃C targets under variable Ar pressures at room temperature. Bauer-Grosse and Le Caer¹¹ obtained a set of $Fe_{1-x}C_x$ films by triode sputtering of composite targets keeping constant all the other deposition parameters: for $x \leq 0.19$, films consisted of a mixture of amorphous Fe-C zones and Ferich zones; for $0.19 \le x \le 0.32$, films were single-phased amorphous, and for $0.32 \le x$, they were a mixture of amorphous Fe-C zones and amorphous C-rich zones. It was indirectly shown that, with increasing of x, the Fe–C zones kept a carbon content of ~ 0.32 , but their size decreased (2000 nm for x = 0.32 to 50 nm for x = 0.45), indicating the possibility to produce Fe-C nanocomposites by nanostructuration of the two amorphous phases. Afterwards, nanocomposite $\text{Fe}_{1-x}C_x$ (0.26 $\leq x \leq 0.88$) thin films were effectively elaborated by ion-beam

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sputtering of an iron-graphite composite target^{6,12}: Fe_{0.74}C_{0.26} film, prepared at room temperature, was composed of Fe-rich connected nanograins surrounded by amorphous-like carbon, while films of composition Fe_{0.46}C_{0.54} and Fe_{0.26}C_{0.74} deposited at 573 and 773 K, respectively, consisted of spherical and well-separated α -Fe nanoparticles embedded in poorly crystallized graphite-like carbon layers. Lee et al.¹³ have also obtained nanocomposite films by sputtering of a composite target: the nanograins, embedded in an amorphous carbon matrix, were either Fe/ θ -Fe₃C or only θ -Fe₃C carbide according to the film composition (0.42 $\leq x \leq 0.85$).

Sputtering of a metallic target in a reactive atmosphere containing methane is therefore of a strong interest because the control of the gas ratio CH_4/Ar allows the control of the film structure and composition¹⁴ in a more flexible way than the use of composite target. It was shown in a previous work that the global carbon content (*x*) in films measured by energy dispersive spectroscopy (EDS) was proportional to the methane partial pressure and that films become x-ray amorphous with the increase of *x*.¹⁵

This work deals with the evolution of structural properties of the Fe–C film surfaces with the control of the gas ratio CH₄/Ar. Grazing incidence x-ray diffraction (GIXRD), scanning electron microscopy (SEM), transmission electron microscopy (TEM), and electron energy loss spectroscopy (EELS) were used for this purpose. A thermal in situ treatment study in the TEM was also performed on one of the richest carbon samples.

II. EXPERIMENTAL SECTION

A. Synthesis

Four Fe_{1-x}C_x films (denoted hereafter A, B, C, and D, see Table I) were deposited on copper substrates at 373 K by direct current (dc) sputtering of an iron target (50 mm in diameter, 1 mm thick, 99.5% purity) in a specific configuration detailed in Ref. 15 under argon/methane reactive mixtures. Before elaboration, the substrates were polished and heated in carbon tetrachloride until boiling, and then cleaned ultrasonically in an ethanol bath. The experimental sputtering device was a 30-L sputtering chamber equipped with a magnetron target substrate holder parallel to the substrate. A base vacuum of about 10^{-4} Pa was ensured by a secondary suction system composed of primary and secondary oil diffusion pumps.

The gas flow rates were controlled by digital MKS flowmeters (MKS Instruments, Inc., Andover, MA). They were kept constant at 30 standard cubic centimeters per minute (sccm) for Ar and regulated at 2.5, 5, 7.5, and 10 sccm for CH₄, for samples A, B, C, and D, respectively. The working pressure, measured using a MKS Baratron absolute gauge, was maintained around 0.2 Pa, with the help of a throttle valve. The Fe target, equipped with a balanced magnetron, was supplied with an Advanced Energy dc generator (Fort Collins, CO), which can deliver a maximal power of 1.5 kW. Target intensity was $I_{\rm T} = 0.2$ A. The coatings were prepared using a configuration called triode sputtering system, with a supplementary tungsten filament as second cathode and a ring placed between the substrate and the target as second anode. The intensity of the tungsten filament was $I_{\rm W} = 5$ A, the substrate-ring anode distance was $D_{S-A} = 20$ mm, while the anode-target distance was $D_{A-T} = 60$ mm. Time deposition was 7200 s and the rate deposition was about 1 μ m.h⁻¹. Therefore, the whole thickness of coatings was about $2 \,\mu m$.¹⁶

B. Analyzing methods

Identification of phases was realized by GIXRD using Co K_{α} ($\lambda = 0.17902$ nm) radiation with an incidence angle fixed at 0.5° (INEL CPS120, Artenay, France). SEM was performed with a Philips XL30 SFEG instrument (FEI Company, Eindhoven, The Netherlands), at 5.00 kV. Wedge samples for TEM were prepared by polishing the substrate by the classical tripod method¹⁷ until the sample was thin enough. A subsequent polishing was performed by ion milling (Gatan PIPS, Pleasanton, CA). By this method, only the top of the coating was analyzed by TEM and EELS (plane-view observation). Conventional TEM was performed on a LaB₆ Philips CM200 instrument with a voltage of 200 kV, and in situ thermal treatment was performed in a Philips CM12 TEM (120 kV) by using a Gatan heating holder. The sample was heated at a rate of 10 K/min, with stages of 15 min every 50 K, until the temperature 723 K was reached.

EELS was carried out with a PEELS 666 Gatan device, coupled to the TEM operating at 120 kV, with an entrance aperture of 2 mm, in diffraction mode with chamber length L = 100 mm, convergence semi-angle $\alpha_c = 6.7$ mrad, and acceptance semi-angle $\beta = 6.55$ mrad. Two energy dispersion values were chosen, namely

TABLE I. GIXRD, TEM, EELS, and EDS characterization results of Fe-C coatings.

Samples	D _{CH4} (sccm)	Identified phases by GIXRD	Identified phases by TEM	Chemical composition (EELS)	Chemical composition (EDS from Ref. 15)
А	2.5	ε-Fe ₃ C	ε-Fe ₃ C	$Fe_{0.70~\pm~0.07}C_{0.30~\pm~0.03}$	Fe _{0.72} C _{0.28}
В	5	ε-Fe ₃ C	ε-Fe ₃ C	$Fe_{0.64~\pm~0.06}C_{0.36~\pm~0.03}$	Fe _{0.63} C _{0.37}
С	7.5	Amorphous-like phase	Nanocomposite film	$Fe_{0.45~\pm~0.04}C_{0.55~\pm~0.05}$	Fe _{0.51} C _{0.49}
D	10	Amorphous-like phase	Nanocomposite film	$Fe_{0.40~\pm~0.04}C_{0.60~\pm~0.06}$	Fe _{0.37} C _{0.63}

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0.098 and 0.996 eV/channel, corresponding to the resolution values of 0.7 and 2 eV, respectively, with an underheated filament. All spectra were corrected for dark current and detector gain channel-to-channel variation. The low-loss spectra were deconvoluted of the zero-loss peak. To remove the plural inelastic scattering contribution, a Fourier ratio deconvolution procedure (Gatan EL/P software) was applied to each edge (C K and Fe L3L2), after background removal, according to a Ae^{-r} power-law. In addition, two reference samples were analyzed with the same procedure, namely an amorphous carbon thin film on a commercial copper grid and a pure iron sample (99.95%), also prepared by tripod polishing and ion milling.

III. RESULTS AND DISCUSSION

A. XRD measurements

Figure 1 shows XRD patterns of samples A, B, C, and D (the corresponding results are listed in Table I). The diffraction diagrams of samples A and B exhibit diffraction peaks corresponding to the metastable hexagonal ϵ -Fe₃C phase¹⁸ (a = 0.4767 nm, c = 0.4354 nm; SG: P6₃22; JCPDS No. 01-089-2005), which is a superstructure of the hexagonal ε -Fe₂C phase^{19,20} (a = 0.2754 nm, c = 0.4349 nm; SG: P6₃/mmc; JCPDS No. 00-36-1249). The lattice parameters are a = 0.4847 nm and c = 4.3884nm for sample A, and a = 0.4827 nm and c = 4.3704 nm for sample B, respectively, with the same c/a ratio, i.e., 0.905. Variation of lattice parameter values corresponds to an increasing occupation of the octahedral sites by carbon atoms in the hexagonal lattice of the iron atoms. Both patterns of samples C and D present an x-ray amorphous profile. Two corresponding wide peaks are observed at about $2\theta = 50.60^{\circ}$ (lattice spacing $d \approx 0.209$ nm) and 98.52° ($d \approx 0.118$ nm). These distances correspond to the {100} and {101} planes group of graphite



FIG. 1. Grazing incidence XRD patterns (wavelength $\lambda = K_{\alpha}Co)$ of samples A to D at incidence $\alpha = 0.5^{\circ}$ (* = copper substrate; $o = \epsilon$ -Fe₃C phase).

(or band 10 for turbostratic carbon) and to $\{110\}/\{112\}$ group (or band 11), respectively,²¹ and therefore indicate the possible presence of an amorphous carbon phase in the samples. However, the peaks are large enough to also include contributions of other amorphous phases or nanocrystalline Fe–C phases.

B. SEM

Figure 2 displays scanning electron micrographs of surfaces of the samples A to D. Surface of the sample A corresponds to facetted and elongated crystalline grains, with a broad size distribution between 50 and 200 nm. Sample B also presents crystalline grains with a larger size (roughly 100 to 500 nm). These grains are more isotropic and possess narrower size distribution than grains in sample A. It should be noted that a poisoning of the iron target has been observed in the presence of Ar/CH₄ gaseous mixture and that the poisoning level seemed to increase with the CH₄ flow rate.¹⁶ This phenomenon is well known in reactive sputtering. It leads to the modification of the discharge properties like the sputter yield, commonly lower in poisoning sputtering mode than in metallic mode. Thus, the surface properties are affected such as deposition rate, chemical composition, grain size, crystallinity, etc.^{7,22} The change in grain size and in grain morphology from sample A to B can therefore be explained by an increase of energy ion bombardment of the growing film leading to the higher mobility of adatoms at the surface. The sample C presents a glassy dense surface with wide curved domains (0.5 to $3 \mu m$), which corresponds to the amorphous-like state observed by GIXRD. The surface of the sample D is rougher and presents smaller domains than that of the sample C.

C. TEM observations

Bright-field and dark-field micrographs of the coatings A and B are shown on Figs. 3(a)-3(d). The sample A presents grains with an average size of 100 nm, while the sample B possesses the largest grain size, about 400 nm. This difference is also attributed to the poisoning of the target. Electron diffraction patterns (EDPs) of the ε-Fe₃C phase along [110] (sample A) and [110] (sample B) zone axes are displayed on Figs. 3(e) and 3(f) (Table I). On Fig. 3(e), the most intense reflections correspond to fundamental reflections of the ε -Fe₂C phase while weak spots correspond to the superstructure. Observation of several extinctions in this pattern demonstrates that the compound is nonstochiometric and presents carbon vacancies in its lattice. On Fig. 3(f), arrows indicate weak 001 reflections, with (1 = 2n + 1), that are not compatible with the P6₃22 space group. These reflections are understood as a result of double-diffraction phenomena, which are often encountered during studies of hexagonal lattices by transmission electron diffraction.



FIG. 2. Scanning electron micrographs of samples (a) A, (b) B, (c) C, and (d) D.

Bright-field micrographs of the samples C and D are shown on Figs. 4(a) and 4(b). These images reveal that the microstructure is constituted of dark nanograins (with an average size of 10 nm for sample C and 5 nm for sample D) embedded in a light matrix. The strong contrast difference between grains and matrix indicates that this last one is mainly constituted of carbon while the nanograins are richer in iron. Thickness of matrix between two grains is larger for sample D, corresponding to the higher content of carbon in this coating. In addition, the dark grains are connected in the sample C while they are mainly disconnected in the sample D, indicating that percolation threshold has not been reached.^{6,23}

Figures 4(e) and 4(f) present corresponding EDPs and their diffraction intensity profiles as a function of inverse of lattice spacing d. For both samples, the patterns show two diffuse and broad rings characteristic of an amorphous phase, at $d = 0.212 \pm 0.05$ nm and $d = 0.122 \pm$ 0.05 nm, in agreement with the distances measured by GIXRD (the additional sharp ring is an artefact and corresponds to the image of an upper diaphragm in the microscope). Figures 4(c) and 4(d) show dark-field images of both samples, obtained by selecting the shortest diffusive ring with an objective aperture of the TEM [circled on Figs. 4(e) and 4(f). The matrix and the nanograins are both illuminated by this method, the nanograins being brighter, indicating for them either a higher amount of iron and/or a partly crystalline state. Lattice fringe images (Fig. 5) were obtained on both samples. Within the resolution of our instrument, it can be seen that the matrix presents characteristic features of amorphous and poorly crystallized carbon layers with fringes corresponding to

bent graphenelike layers [i.e., (002) layers of graphite]. It is expected that diffraction of a graphitic amorphous phase results in a broad peak at $1/day = 2.7-2.8 \text{ nm}^{-1}$ corresponding to the interlayer spacing between graphene layers. For instance, the EDPs of Fe-C nanocomposite films in Refs. 6 and 12 exhibit an innermost halo ring corresponding to graphite basal (002) planes. In this study, such a feature is not seen in XRD patterns (at $\theta = 27^{\circ}$ - 30°) and in EDPs. The short stacking (2 or 3 layers) of highly faulted and strongly bent graphenelike layers, which partially compose the carbon matrix explains the absence of the (002) peak. Such distortion can enlarge the lattice spacing between subsequent graphitic sheets, introduce large stress, or avoid formation of short-order distances corresponding to interlayer bonds. It has been previously shown by Franklin²¹ that x-ray data of such graphite-like layers, which are well ordered in two dimensions but present disorder in the third direction, exhibit an extremely weak (002) intensity. In Fig. 5, most of the dark nanograins present short-order contrast while some of them present lattice fringes, with periodicities around 0.27 and 0.34 nm. These lattice distances could correspond to several carbides, but not to the α -Fe phase. At this point, we can conclude that the nanocomposite films are constituted of disordered and crystalline carbide nanograins embedded in a matrix made of amorphous and poorly crystallized graphenelike carbon.

D. Electron energy loss spectroscopy results

Local composition of the $Fe_{1-x}C_x$ coatings was measured by EELS and compared to previous global

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FIG. 3. Bright-field micrographs of samples (a) A and (b) B. Dark-field micrographs of samples (c) A and (d) B. EDPs of the ε -Fe₃C phase (a = 0.477 nm, c = 0.4365 nm) along the [110] zone axis [($11\overline{2}0$)* pattern] observed in (e) sample A and along the [$\overline{1}10$] zone axis [($\overline{1}100$)* pattern] observed in (f) sample B. Arrows indicate reflections arising from double-diffraction phenomena.

measurements by EDS (Table I). These measurements show that the amount of carbon in the coatings increases with the methane partial pressure, as previously reported in Ref. 15. Results from both techniques are similar. A small amount of oxygen was also detected by EELS in sample D at the detection limit of the technique (this amount could be roughly estimated to be below 3%). The presence of oxygen may be due to the tripod preparation, which involves a polishing process under water.

Figure 6 presents low-loss and core-loss spectra for samples A to D, compared to spectra of reference samples, i.e., pure iron and commercial amorphous carbon. In Fig. 6(a), the plasmon peak of the pure iron reference is located at 23.7 eV in agreement with the reported

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FIG. 4. (a, b) Bright-field micrographs of samples C and D; (c, d) dark-field micrographs of samples C and D obtained by selecting the first diffusive ring in the EDP; (e, f) corresponding EDPs (open circles correspond to the TEM objective aperture). Insets: electron diffraction intensity profile as a function of $1/day (nm^{-1})$.

value 24 eV for α -Fe.²⁴ A weak shoulder appears on the left of the peak at 16.8 eV. This feature is the surface plasmon, located at $1/\sqrt{2}$ times of the bulk plasmons energy and characteristic of a thin specimen.²⁵ The Fe M3,2 edge at 55 eV is clearly visible. The plasmons resonance of the amorphous carbon standard is located at 23.3 eV in agreement with literature results, in which

the collective excitation of valence electrons ($\pi + \sigma$ plasmons) appears between 21 to 25 eV.^{26,27} This corresponds to an amorphous carbon in which the atoms are mainly sp^2 bounded.²⁶ The broad resonance observed at 5 eV in this spectrum corresponds to the π plasmons related to the π - π * interband transition in graphitic materials.²⁶



FIG. 5. Lattice fringes electron micrographs of samples (a) C and (b) D.



FIG. 6. (a) Low-loss EELS spectra of amorphous carbon standard, pure iron standard, Fe–C samples A to D and annealed sample C; (b) core-loss (C K edge) EELS spectra of amorphous carbon standard, samples A to D and annealed sample C.

Concerning the samples A and B, both composed of the ϵ -Fe₃C phase, no significant difference with the pure iron spectrum is observed, except in the height of the Fe M3,2 edge. The relative intensity of this edge is more prominent for these samples than for the samples C and D, in agreement with the fact that the samples A and B are the richest in iron. Concerning the nanocomposite films, the plasmons resonance is located at 24.2 eV (sample C) and at 23.6 eV (sample D), respectively. In addition, the resonance is more intense for C than for D. The position

of this resonance does not correspond to that of the graphite (located at 27 eV according to literature),²⁶ but may correspond to amorphous carbon or to structures made of curved graphene sheets.^{28,29} No peaks relative to the π - π * bands transition is observed in spectra of sample C and D, although graphenelike layers were observed by TEM. It was demonstrated that due to the anisotropy of the graphite, the low-loss spectrum of this structure change as a function of the *c*-axis orientation regarding the electron beam.^{28,30} In particular, when the graphene layers are parallel to the beam, intensity of the interband resonance strongly decreases and the plasmons resonance is moved toward the lower energy values. The same behavior occurs for objects made of curved and defected graphene layers, as carbon-filled fullerenes, nanotubes, nanocones, etc., which present different spectra for the same unique object as a function of the zone, which is impacted by the beam.^{28,29} Nevertheless, when a large number of such randomly oriented objects are probed simultaneously, the mean spectrum presents the $\pi - \pi^*$ peak, as can be seen in Ref. 30 for nano-onions. Therefore, the absence of this peak in spectra of nanocomposite samples, together with the slight differences in position and height of the $(\pi + \sigma)$ plasmons peak between C and D, suggests a possible anisotropy of the C-rich domains observed in Fig. 5, with the graphenelike layers mainly parallel to the electron beam. Elongation of carbon domains of a nanocomposite Fe-C coating was previously evidenced by Babonneau et al.⁶ and considered the result of a columnar growth. Nevertheless, interpretation of low-loss spectra is limited because the samples are composite materials. In addition, no information concerning the iron-rich nanograins are available in these spectra.

Figure 6(b) shows the C K edge for the amorphous carbon standard and the samples A to D. The edge is characterized by a fine peak, noted (1) for all samples. This resonance at 284 eV results from transitions $1s-\pi^*$ and is characteristic of *sp* and *sp*2 bonds.^{31,32} Post-edge, energy-loss near-edge structures (ELNES) of the samples A and B are composed of two contributions (2) and (3), located at 8 and 15 eV from the edge, respectively. It is well known that the ELNES often exhibit a structure specific to the nearest neighbor environment of the atom undergoing excitation (also called coordination fingerprint).³³ In samples A and B, carbon atoms are located in octahedral sites of iron carbides. Post-edge contributions correspond to a fingerprint similar to that found for N K edges in isostructural hexagonal transition metal nitrides,³⁴ such as hexagonal Cr₂N, V₂N (both isostructural to ε -Fe₂₋₃C),^{35,36} and ε -Ti₂N (structurally related to ε -phase).³⁷ ELNES of these 3 nitrides are similar to the signal presented in Fig. 6(b). In particular, calculated spectra in Ref. 35 show the presence of the post-edge features at 7 and 14 eV behind the edge.

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The $1s-\pi^*$ peaks in spectra of the samples C and D are wider, while their magnitudes decrease. In addition, the ELNES structure is a single broad contribution (4), whose intensity, relatively to those of the edge is more and more important. For carbon materials, this wide peak at 295 eV results from transitions $1s-\sigma^*$.^{26,31,32,38} The shape of the ELNES structures in spectra of samples C and D is similar to that of the amorphous carbon standard, and different than that of the graphite.^{30,38} The height of the σ^* peak relative to that of the π^* edge demonstrates that the ratio sp2/sp3 bonds of the carbon material in the samples C and D is larger than in the standard. It is interesting to note that for carbon materials made of curved graphene layers, the sigma bonds are not purely of the sp2 type, but between sp2 and sp3 bonds,³⁸ while their spectrum could be similar to that of the amorphous phase.³⁹ Therefore, the ELNES structures of the samples C and D correspond either to amorphous carbon or to poorly crystallized graphene or to a mixture of both, as observed by TEM.

Concerning the Fe L3L2 edge, no significant difference has been observed between spectra of the iron standard and the Fe–C samples. Therefore, these results are not presented here. None of these spectra present characteristic features of iron oxidation (like large white line-intensity ratio L3/L2 or presence of broad peak at 37 eV from the edge).^{40–42}

E. In situ annealing treatment results

In situ annealing treatments in the TEM were undertaken to evaluate thermal stability of the nanocomposite Fe–C films and to identify the transformation products. Crystallization study of a disordered solid state is a way to discover new crystalline arrangements and this approach was already used in the Fe–C system.^{43–45} Furthermore, several new iron carbides were recently reported.^{46–48} Because of the presence of oxygen in the sample D, one of its crystallization products was the Fe₃O₄ oxide. Consequently, results of this experiment are not shown here. For the sample C, structural changes of the nanocomposite into large crystalline grains, with a size of about 0.5 µm, started around 523 K. The transformation was accomplished at 723 K.

This grain size is large enough to allow observation of the same grain in reciprocal space along several zone axes in microdiffraction conditions (i.e., with an almost parallel beam and a small probe size). Several grains have been analyzed and at least three crystalline structures have been identified in the same sample.

Figure 7 shows EDPs of these crystalline phases. The first one is the orthorhombic θ -Fe₃C (SG: Pnma; a = 0.5089 nm, b = 0.67433 nm, c = 0.45235 nm). Figure 7(a) presents the EDP along the [110] zone axis of θ -Fe₃C. The second observed structure is the orthorhombic Fe₇C₃ phase [Fig. 7(b): EDP along the [021] zone axis]. The









(b)

FIG. 7. (a) EDPs of the θ -Fe₃C phase along the [110] zone axis, (b) of the orthorhombic Fe₇C₃ phase along the [021] zone axis, (c) of a new fcc carbide (a = 0.784 nm, possibly related to the τ_2 -Fe₂C₇ phase) along the [110] zone axis.

lattice parameters of this phase are a = 0.453 nm, b = 0.689 nm, and c = 1.191 nm and its space group is Pnma. In these two carbides, the carbon atom coordination is prismatic. These results are in agreement with previous studies.^{43–45} Finally, EDPs of the third crystallization product were indexed using a face-centered-cubic (fcc) cell with a = 0.784 nm. EDP along [110] zone axis of this phase is shown on Fig. 7(c). From the lattice constant and Bravais lattice viewpoint, it is similar to the carbon-rich carbide called τ_2 -Fe₂C₇, one of the two new iron carbides discovered in 1995 by Komekyu et al.⁴⁶ by heating a mixture of iron and carbon powders in a stream of hydrocarbon gas at 1870 K. Moreover, the presence of a carbonrich compound with the two iron-rich θ -Fe₃C and Fe₇C₃ carbides allows preservation of the composition of the sample. Therefore, this new fcc phase is possibly related to the τ_2 -Fe₂C₇ carbide. The two lattice fringe measurements previously mentioned in Fig. 5 (≈ 0.24 and 0.274 nm) can be assigned to the {220} and {311} interplanar distances of this new fcc phase. This phase was possibly already present in nanograins before the thermal treatment. EELS was also performed on the annealed sample C. Only C K and Fe L3L2 edges are present in the spectrum registered with a dispersion value of $\sim 1 \text{ eV/channel}$ (not shown here), which means that no oxygen has been detected (at 532 eV) in the coating after annealing. Global chemical composition is Fe_{0.47 \pm 0.03C_{0.53 }\pm 0.03. All other spectra are presented in Fig. 6, to facilitate comparison with as-sputtered coatings. The plasmons resonance is located at 24.7 eV in the low-loss spectrum, which corresponds to a shift of 0.5 eV toward high energy values, compared to the sample C before annealing. This shift is expected since the annealed coating is constituted of carbides. The CK edge is presented in Fig. 6(b). This contribution is different from both crystalline E-Fe₃C and nanocomposite structures. The $1s-\pi^*$ resonance is broader and its intensity is smaller than the ELNES intensity, as for the nanocomposite structure. In the ELNES signal, several characteristic features are present. However, the present sample is a mixture of carbides, with possibly different carbon environments (prismatic coordination of carbon atoms in θ -Fe₃C and Fe₇C₃ phases, while iron atoms are octahedrally coordinated in τ_2 -Fe₂C₇, according to the structural model of this carbide, which was proposed from electron diffraction intensities⁴⁶ and if the new phase is the τ_2 -Fe₂C₇ compound), the corresponding spectrum is therefore more complex than previous ones. This ELNES structure should contain fingerprints of Fe₇C₃ and cementite-type carbides but it was difficult to compare with those found by previous studies for $(Cr,Fe)_7C_3^{34}$ and the cementite phase,⁴⁹ respectively. In all works, the π^* edge intensity is higher than the ELNES intensity. Concerning the Fe edge (spectrum not shown), no oxidation feature is observed.

has been investigated by GIXRD, SEM, TEM, and EELS. This study has demonstrated the ability to produce iron-carbon films by reactive sputtering of an iron target in partial methane atmosphere. Control of the methane flux allows the formation of more and less carbon-rich materials:

(1) iron carbide, in which the carbon atoms are in octahedral coordination. Electron low-loss and core-loss energy spectra have been investigated, and compared to that of amorphous carbon and pure iron standards, and to isostructural transition metal nitrides.

(2) nanocomposite microstructure consisting in disordered and crystalline carbide nanograins embedded in an amorphous and poorly crystallized sp2/sp3 carbon matrix. The graphenelike layers in the nanocomposite structures are strongly bent, have no tendency to parallel stacking, and are mainly parallel to the electron beam, indicating a columnar-like growth. The percolation threshold of the iron-rich grains is almost reached in the richest carbon coating (Fe_{0.40 \pm 0.03}C_{0.60 \pm 0.03}).

The crystallization products of the nanocomposite $Fe_{0.45~\pm~0.03}C_{0.55~\pm~0.03}$ film are the known $\theta\text{-}Fe_3C$ and Fe₇C₃ compounds, in which the carbon atoms are in prismatic coordination, and a new fcc carbide (a =0.784 nm), possibly related to the C-rich cubic τ_2 -Fe₂C₇ phase, in which the iron atoms are octahedrally coordinated.46

The results we obtained show the wide variety of materials that can be synthesized by reactive sputtering by varying only one elaboration parameter and suggest that it would be interesting to study the effects of other determining elaboration parameters, such as substrate temperature or gas reactivity, to produce either new carbides or to control composition and microstructure of nanocomposite films. Indeed, control of the size and of the insulation of individual particles of metals showing a low solubility for carbon (as Fe, Co, Ni, Cr, Au, Pt, Ag, etc.) and embedded in a carbon matrix is crucial for their mechanical, tribological, magnetic, and electronic properties.^{23,50–52} Reactive sputtering is nowadays an industrial process, the potential applications are high-density magnetic recording media, with magnetic Fe, Co, or Ni particles acting as granular materials, protective coatings against wear or hard cutting tools. Further investigations on properties of the new carbon-rich carbide are required to define a possible application of this material.

This study will be followed by the description of the effects of the poisoning of the target and the characterization of the coatings in cross-sectional view.

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IV. CONCLUSION

 $Fe_{1-x}C_x$ coatings have been prepared by reactive triode-magnetron sputtering. The surface of these coatings

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