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Flexible fluorinated graphite foils with high contents of the $(C_2F)_n$ phase for slow neutron reflectors†

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In order to prepare self-standing and flexible slow neutron reflectors made of graphite fluoride (GF) with high contents of $(C_2F)_n$ structural phase, graphite foils of different thicknesses were used as starting materials for gas $(F_2)/\text{solid}$ fluorination. The maximal interlayer distance of GF was obtained with this phase thanks to the stacking sequence FCCF/FCCF; this is mandatory for the efficient reflection of slow neutrons. 71 and 77% of the $(C_2F)_n$ phase were achieved for graphite foils with thicknesses of 1.0 and 0.1 mm, respectively. The interlayer distances were 8.6 Å as expected. The fluorination conditions (static mode, a long duration of 24 h, annealing in pure F_2 gas for 24 h, and temperatures in the 390–460 °C range) were adapted to large pieces of graphite foils $(7 \times 7 \text{ cm}^2)$ in order to both avoid exfoliation and achieve a homogeneous dispersion of fluorine atoms. This process was also efficient for thinner (0.01 mm thick) graphitized graphene oxide foil. 56% of the $(C_2F)_n$ phase and an interlayer of 8.6 Å were achieved for this foil when fluorination was performed at 430 °C. Whatever the nature and the thickness of the foil, their flexibilities are maintained.

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Introduction

Neutrons are of great importance in fundamental and applied research, and neutron reflectors are an indispensable tool in this field. In order to design neutron reflectors that are efficient in the complete range of neutron wavelengths, new methods and materials have to be developed. Thanks to a specialized diffractometer adapted to work with slower neutrons at the PF1B instrument 11,12 at ILL, Grenoble, France, powdered $(C_2F)_n$ -rich graphite fluoride, with an interplanar distance d of 9 Å, was demonstrated to be an effective reflector for neutrons with wavelengths in the range 6–18 Å, completely covering the gap in the reflectivity of modern neutron reflectors. As a matter of fact, the efficiency of standard neutron

gas) starting from flakes of natural graphite. 12

reflectors, like graphite, decreases sharply at neutron energies

below the so-called Bragg cut-off associated with the coherent

scattering of neutrons on crystal planes. The value of the energy depends on the reflector material; the neutron wavelength λn corresponding to this energy value is equal to 2 interplanar distances d. For graphite, ¹³ it is $\lambda_n^{\text{Bragg}(Gr)} = 2 \cdot d_{Gr} = 6.708 \text{ Å}$, (where $d_{\rm Gr}$ = 3.354 Å). To overcome this limitation for the slowest neutrons, we have developed a reflector method based on fluoridetonation nanodiamond (F-DND) powders. 14-19 Nevertheless, prior to ref. 12, there had been still a gap in reflectivity in the neutron wavelength range $[\lambda_n^{\text{F-DND}} - \lambda_n^{\text{Bragg}}]$, only partially covered with less efficient materials.20 The high interplanar distance d (9 Å) in the $(C_2F)_n$ phase is related to the stacking sequence FCCF/FCCF where half of the carbon atoms is bound to fluorine, whereas each carbon atom is fluorinated in the $(CF)_n$ phase, resulting in an FCF/FCF stacking sequence with an interplanar distance equal to 6 Å. In addition to this advantage, graphite fluoride of the $(C_2F)_n$ structural phase consists of atoms with very small capture cross sections (σ_{abs}^{C} = 3.5 mb and $\sigma_{\rm abs}^{\rm F}$ = 9.6 mb at the thermal neutron velocity, respectively) and large coherent scattering lengths ($b^{C} = 6.65$ fm and $b^{\rm F}$ = 5.65 fm, respectively). Graphite is relatively free from impurities and can be further purified to a significant extent.²¹ According to Kita's work, 22 GF with a (C2F)n phase content of 96% was prepared using the reaction of molecular fluorine (F2

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Up to now, only powders or small flakes (200-400 µm) of graphite have been considered as starting materials for fluorination.²³ The next step was the compaction of the resulting GF in order to increase the density of the reflector; the higher its density, the higher its reflectivity, for the realistic geometries of reflectors. The high electronegativity of fluorine atoms

appears as a drawback because it decreases the surface energy of fluorocarbon layers and hinders compaction. In the present work, we have chosen an alternative strategy with the fluorination of graphite foil in order to prepare GF foil with a high content of (C₂F)_n. Nuclear grade was selected for the foils to avoid impurities that would increase neutron losses and be activated in high radiation fluxes. The main experimental difficulty was encountered in avoiding the exfoliation of the graphite foil during fluorination in order to maintain the density as high as possible. Because large foils are necessary for neutron reflection, the reaction conditions must be adapted to achieve a homogeneous distribution of the fluorine atoms in the bulk of the foils. Unlike powder reflectors, intercalated foils exhibit anisotropic neutron scattering. This process requires a special design to obtain an isotropic reflector. However, it opens up an additional opportunity for more efficient directional neutron scattering.

Experimental

Paper

Fluorination of graphite foil

High purity premium grade or nuclear grade flexible graphite foils (Papyex) with thicknesses of 0.1 and 1.0 mm were kindly provided by Mersen. Following chemical preparation with acids and oxidizing agents, natural graphite was expanded using a violent heat treatment procedure. Self-bonded by a rolling operation, the expanded graphite particles form a continuous strip, without the addition of a binder. A multi-step process was used to achieve ultra-low levels of sulfur (typical value 500 ppm), chloride (20 ppm, a leachable chlorine content of 10 ppm) and other impurity contents. The carbon rate and ash content were 99.9% and 0.1%, respectively.

Our previous works on powdered $(C_2F)_n$ GF evidenced that long durations were necessary in the temperature range of 390-450 °C.12 An up-scalable static mode was used to perform fluorination for 48 h in a 5 L horizontal tubular reactor made with nickel alloy and passivated with NiF2. In this mode, fluorine was added in a closed reactor starting from a primary vacuum. Foils of 70 × 70 mm² size and 1.0 or 0.1 mm thickness were used (see Fig. 1). Before the reaction, moisture and oxygen were removed by applying a primary vacuum (10-3 mbar) at 200 °C. In order to avoid the exfoliation during addition of F2 gas, a two-step fluorination was carried out as follows: after the preliminary step (primary vacuum at 200 °C, addition of F2 at 270 °C, step at 360 °C for 2 h), the reaction temperature was first maintained at 390 °C for 24 h to form the $(C_2F)_n$ phase and then increased to TF (390, 410, 430 and 460 °C) to enhance the structural homogeneity by annealing. The annealing temperature was maintained for 24 h. The

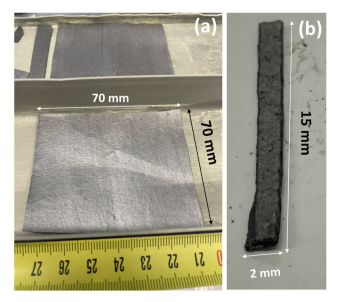


Fig. 1 Representative picture of the fluorinated foils (a) and cut pieces for XRD analysis (b).

sample is denoted by the temperature of annealing TA and foil's thickness (0.1 or 1.0 mm), i.e. FTF-Foil0.1 and FTF Foil1.0. Addition of F₂ gas was performed when the pressure drop was at least 0.2 bar. After the fluorination and during the cooling of the reactor, the reactive atmosphere was flushed with nitrogen gas. For comparison, GF was synthesized with the same fluorination process and TA = 440 °C using natural graphite (Sigma Aldrich) with small flakes of 200-400 µm as a starting material. With the same aim of comparison, a $(CF)_n$ GF was prepared at 600 °C for 3 h using graphite powder (Timrex KS4 graphite with 4 µm granulometry). Its composition is CF₁.

Graphitized graphene oxide (GO) foil was also used as a starting material in order to increase the structural order and then the density. A GO film was first synthesized by exfoliating graphite oxide using the modified Hummers' method. 23-25 The resulting GO was dispersed in water to achieve a concentration of 0.5 mg mL⁻¹. GO films were prepared by vacuumdriven filtration of 10 mL of the aqueous GO dispersion through mixed cellulose ester (MCE) filters (Whatman™, 10401712) with a pore size of 0.2 µm and a diameter of 47 mm. Subsequently, the GO films were dried in ambient air and carefully separated from the filters by using isopropyl alcohol. To remove any residual impurities from the MCE filters, the films were washed with acetone. After graphitization at 2750 °C, its interlayer distance was then 3.36 nm and its coherence length along the c-axis was 390 Å corresponding to 116 stacked sheets. The graphene film is carbon-pure, uniform, and flexible, with good mechanical performance, and high electrical and thermal conductivities (~1200 W m⁻¹ K⁻¹). The resulting sample is denoted as gGO. Its thickness was around 0.1 mm. The same fluorination conditions as for the foils of 0.1 and 1.0 mm thickness were applied for the FTFgGO series ($T_{\rm F}$ = 390, 410 and 430 °C).

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Physicochemical characterization

NMR experiments were carried out with a Bruker Avance spectrometer, with working frequencies for ¹³C and ¹⁹F of 73.4 and 282.2 MHz, respectively. A Magic Angle Spinning (MAS) probe (Bruker) operating with 2.5 mm rotors was used. For MAS spectra, a simple sequence was performed with single $\pi/2$ pulse lengths of 4.0 and 3.5 μs for ¹⁹F and ¹³C, respectively. For MAS measurements, the samples must be ground. 13C chemical shifts were externally referenced to tetramethylsilane (TMS), while ¹⁹F chemical shifts were referenced with respect to CFCl₃. X-ray diffraction (XRD) patterns were recorded using both a PANalytical X'Pert X-ray reflection diffractometer with Cu (K α) radiation ($\lambda_{K\alpha_1}$ = 1.5406 Å) and a Bruker D8 Advance transmission diffractometer using a molybdenum anticathode $(\lambda_{K\alpha} = 0.7093 \text{ Å})$ equipped with a 2D LynxEye detector and a 2.5° divergence Soller slit. Regarding the XRD patterns recorded with Mo radiation, diffractograms were accumulated for 1 h in the 2-45° (2 θ) angular range. This measurement was repeated 5 times for both (00l) and (hk0) diffractograms, in order to increase the signal-to-noise ratio. Fluorinated foil samples were placed in Lindemann glass capillaries (1.5 mm diameter). Due to the extremely oriented nature of the samples, 00l and hk0 reflections can be recorded separately depending on the orientation of samples versus the incident beam. Fourier-transform infrared (FTIR) spectra were recorded on a Nicolet 6700 FT-IR (Thermo scientific) spectrometer in

transmission mode at a resolution of 4 cm⁻¹ and 256 scans were taken for each spectrum. Transmission electron microscopy (TEM) specimens were prepared by disintegrating the films in ethanol and drop casting the suspensions onto lacey carbon-coated Cu grids (SPI Supplies) and dried in air. TEM analysis was performed using a JEOL JEM-F200 Multi-Purpose FEG-S/TEM operating at an accelerating voltage of 200 kV. Image J and Pathfinder were used for processing the TEM and elemental map images.

Results and discussion

In accordance with the expected value for the $(C_2F)_n$ phase (F/C=0.5), an F/C molar ratio close to 0.6 was obtained by weight uptake for the GF starting whatever the temperature of fluorination and initial thickness. The composition will be extracted from ¹³C NMR data. XRD was performed as a systematic characterization study in order to study the annealing effect over the $(C_2F)_n$ structure (Fig. 2). The four annealed F-Foil0.1 samples present the same XRD pattern with four diffraction peaks clearly visible (Fig. 2a and b) and indexed as (001), (002), (003) and (004) planes. A magnification of the 15–70° region is presented in Fig. 2b (highlighted in yellow in Fig. 2a). A shoulder is visible around 41° on the (004) peaks and corresponds to the (110) plane. It is noted that (002), (003)

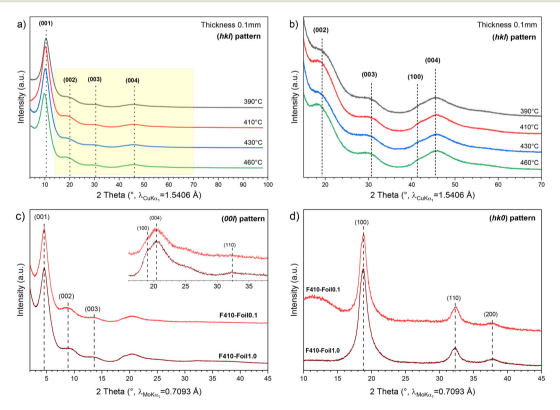


Fig. 2 (a) XRD patterns of fluorinated graphite foils (Cu Kα radiation with $\lambda_{K\alpha_1} = 1.5406$ Å) and (b) with magnification in the 15–70° 2 θ range (shown as a yellow area on (a)). (c) and (d) are respectively the (00l) and (hk0) patterns of foils of 0.1 (F410-Foil0.1, red) and 1.0 mm (F410-Foil1.0, dark red) thickness fluorinated at 390 °C and then annealed under F₂ gas at 410 °C. The latter were acquired with a Mo Kα source ($\lambda_{K\alpha_1} = 0.7093$ Å).

Table 1 $d_{(hkl)}$ (in Å), FWHM (°) and L_c for the (001) peak (in Å) of the four annealed FT_A-Foil0.1 samples ($T_A = 390$, 410, 430 and 460 °C). The XRD patterns were acquired using a Cu K α_1 source ($\lambda = 1.5406$ Å)

| | F390-Foil0.1 | | | F410-Foil0.1 | | | F430-Foil0.1 | | | F460-Foil0.1 | | |
|-------|-----------------|----------|----------------------------------------------|------------------------------------|----------|---------------------------------------------------------|-----------------|----------|---------------------------------------------------------|-----------------|----------|------------------|
| | $d_{(00l)}$ (Å) | FWHM (°) | $L_{\mathrm{c}(001)}(\mathring{\mathrm{A}})$ | $d_{(00l)}(\mathring{\mathrm{A}})$ | FWHM (°) | $L_{\mathrm{c}(001)}\left(\mathring{\mathrm{A}}\right)$ | $d_{(00l)}$ (Å) | FWHM (°) | $L_{\mathrm{c}(001)}\left(\mathring{\mathrm{A}}\right)$ | $d_{(00l)}$ (Å) | FWHM (°) | $L_{c(001)}$ (Å) |
| (001) | 8.41 | 3.41 | 24.4 | 8.63 | 3.62 | 22.0 | 8.63 | 3.54 | 22.5 | 8.90 | 3.48 | 22.9 |
| (002) | 4.35 | 4.08 | _ | 4.48 | 4.27 | _ | 4.53 | 3.73 | _ | 4.59 | 4.46 | _ |
| (003) | 2.91 | 3.93 | _ | 2.95 | 4.06 | _ | 2.95 | 4.18 | _ | 2.94 | 3.79 | _ |
| (004) | 1.97 | | _ | 1.98 | | _ | 1.98 | | _ | 1.98 | _ | _ |

and (004) peaks are particularly broad, which complicates the determination of the d-space. After background subtraction, the d-space and the crystallite size along the c-axis $L_{\rm c}$ (from the full width at half maximum FWHM) were calculated precisely and the obtained values are presented in Table 1. It is seen that the d-space increases with the annealing temperature, otherwise the FWHM and $L_{\rm c}$ do not seem to be particularly affected by F_2 annealing. F390-Foil0.1 has a slightly lower d-space (8.41 Å) than that found in the literature (8.72–9.05 Å), 12,26 and compared to a powder with 96% of the $(C_2F)_n$ phase (9.2 Å), while F460-Foil0.1 has a d-space (8.98 Å) closer to what is expected. Even though the d-space of F390-Foil0.1 is slightly lower than that expected, no other diffraction peak appears, meaning that no $(CF)_n$ pattern is necessary to fully index this diffractogram.

The foil thickness effect on the synthesis of the $(C_2F)_n$ phase was also investigated. The (00l) and (hk0) XRD patterns of F410-Foil1.0 and F410-Foil0.1 were acquired (Mo anticathode) and are respectively shown in Fig. 2c and d. In the (00l) XRD pattern, the same diffractograms as those obtained previously are observed and the d-space, FWHM and L_c are presented in Table 2. Generally, F410-Foil0.1 shows a slightly higher d-space than that observed for the 1 mm thick foil, but both samples remain quite similar. The (00l) pattern confirms the high interlayer distance where the value is 8.9 Å, which is in accordance with a high content of the $(C_2F)_n$ phase. Nonetheless, two (hk0) planes are observed in the (00l) XRD pattern: (100) at 2.18 Å $(\sim19^\circ)$ and (110) at 1.28 Å $(\sim32^\circ)$. This can be attributed to the crumpled nature of Papyex on edges with some layers being folded.

Regarding the (hk0) pattern (Fig. 2d), three diffraction peaks are visible: (100), (110) and (200) planes at 2.17, 1.28

Table 2 Comparison of $d_{(hkl)}$ (in Å), FWHM (°) and L_c (in Å) between F410-Foil0.1 and F410-Foil1.0. The XRD patterns were acquired using a Mo K α_1 source ($\lambda = 0.7093$ Å)

| | F410-F | oil0.1 | | F410-Foil1.0 | | | | |
|-------|---------------------------|-------------|------------------|---------------------------|-------------|-------------------------|--|--|
| | d _(00l) (Å) | FWHM (°) | $L_{c(001)}$ (Å) | d _(00l) (Å) | FWHM (°) | L _{c(001)} (Å) | | |
| (001) | 8.90 | 1.41 | 26.0 | 8.80 | 1.27 | 28.8 | | |
| (002) | 4.50 | 2.07 | _ | 4.33 | 1.92 | _ | | |
| (003) | 2.95 | 1.74 | _ | 2.97 | 1.86 | _ | | |
| (004) | 1.99 | | _ | 1.99 | | _ | | |

and 1.09 Å, respectively. As expected, no significant difference between samples can be noted regarding their initial thickness, as peak widths and positions of the hk0 reflections remain equivalent. The a parameter and the distance between two carbon atoms $(d_{\rm C-C})$ can be extracted from the (hk0) planes. Thus, considering $({\rm C_2F})_n$ as a hexagonal structure, 32 a = 2.53 Å and $d_{\rm C-C}$ = 1.46 Å compared to the a and $d_{\rm C-C}$ of graphite equals 2.46 and 1.42 Å, respectively. As expected, the fluorination of graphite extends the distance between carbon atoms, increasing the in-plane lattice parameter.

Thus, whatever the foil thickness and more surprisingly whatever the annealing temperature between 390 and 460 °C, the XRD patterns corresponding to fluorinated graphite can be indexed as $(C_2F)_n$ phase and no additional diffraction peak is needed to fully index the observed features. In particular, no diffraction peak corresponding to pristine graphite or (CF)n phase is noticed. Also, studying separately the (hk0) pattern allowed us to determine the lattice parameter a and the d_{C-C} , which are higher than those of graphite.

In order to confirm the conclusion from XRD data, solid state NMR was performed for the investigation of the structural type in the short range. The $^{13}\mathrm{C}$ NMR spectra of the foils fluorinated at 390 °C and annealed at 410 °C are composed of two main lines at 84 and 42 ppm which are assigned to the carbon atoms involved in the C–F bonds (denoted C–F in Fig. 3a and d for experimental and fitting data) and non-fluorinated carbons with sp³ hybridization (C–C–F). $^{27,29-32}\mathrm{A}$ very few amount of non-fluorinated sp² C is also evidenced by the line with a weak intensity at 135 ppm. In order to evidence CF2 groups with an expected line at 110 ppm, $^{19}\mathrm{F} \rightarrow ^{13}\mathrm{C}$ crosspolarization MAS spectra were recorded. Their content is close to zero.

According to the integrated surface of the lines $S_{\text{C-F}}$ and $S_{\text{C-C-F}}$ (see Fig. 3c for the representative example of F410-Foil0.1), the ratio $100 \times (\text{C}_2\text{F})_n/(\text{CF})_n = 100 \times S_{\text{C-C-F}}/S_{\text{C-F}}$ gives the percentage of the $(\text{C}_2\text{F})_n$ structural type. 77 ± 2% of C-F bonds are included in the $(\text{C}_2\text{F})_n$ structural type (23% in $(\text{CF})_n$) for F410-Foil0.1. The F/C ratio can be extracted from the fit parameters thanks to the formula F/C = $S_{\text{C-F}}/(S_{\text{C-F}} + S_{\text{C-C-F}} + S_{\text{Csp2}})$ and a value of 0.55 ± 0.01 is obtained for F410-Foil0.1 in good accordance with the weight uptake method (close to 0.6). When the initial thickness is 1.0 mm, very close results are obtained. In order to evidence the effect of the fluorination temperature, Foil0.1 was also treated at 390, 430 and 460 °C, *i.e.* in the " $(\text{C}_2\text{F})_n$ -rich" range defined by Kita.²² The data

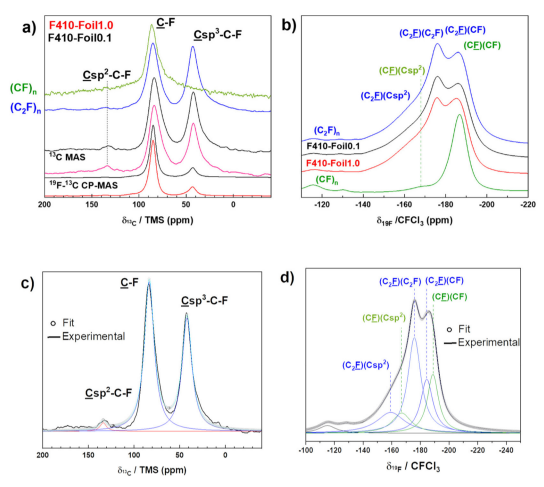


Fig. 3 MAS NMR spectra of F410-Foil0.1 (black line) and F410-Foil1.0 (red line) on 13 C (a) and 19 F nuclei (b). The spinning rates are 10 and 30 kHz, respectively. Sample F440-NG from ref. 21 is used as a $(C_2F)_n$ reference for both 13 C and 19 F spectra (in blue). The spectra of a $(C_2F)_n$ reference were added (in green). Representative examples of fits for 13 C (c) and 19 F (d) spectra are shown (F410-Foil0.1).

extracted from 13 C NMR (Fig. SI1a and b in ESI†) evidence very few differences, 9 (C_2F) $_n$ is around 76% for F390-Foil0.1 and F430-Foil0.1, as for F410-Foil0.1 (see Table SI1 in the ESI†). The slightly lower value of 69 \pm 2% for F460-Foil0.1 is explained by the formation of a high amount of the (CF) $_n$ phase but the main phase is still (C_2F) $_n$. A line at 132 ppm related to non-fluorinated carbons with neighboring C–F is observed both in MAS and $^{19}F^{-13}C$ CP-MAS only at the lowest fluorination temperature of 390 $^{\circ}C$. Nevertheless, its intensity is very low. F410-Foil1.0 exhibits a F/C of 0.58 and a content of (C_2F) $_n$ phase equal to 71%. The applied fluorination and annealing treatment conditions were also efficient for thicker graphite foils.

Fig. 3b and d show respectively the $^{19}\mathrm{F}$ experimental spectra of both F410-Foils where several lines are evidenced. The spectrum of F410-Foil0.1 was simulated as a representative example (Fig. 3d). The presence of CF₂ (-120 ppm) and CF₃ (-80 ppm) is excluded regarding $^{19}\mathrm{F} \rightarrow ^{13}\mathrm{C}$ cross-polarization data and the absence of the corresponding lines. $^{28,29-31}\mathrm{By}$ analogy with the chemical shift at -178 ppm of fluorinated nanodiamonds, *i.e.* Csp³-C- $\underline{\mathrm{F}}$ (their $^{13}\mathrm{C}$ chemical shift is also

42 ppm),¹⁵ the line at -178 ppm is related to the C-F bonds in $(C_2F)_n$ with only C_2F ($(C_2F)_n$) as the neighbor (denoted as $(C_2\underline{F})$ (C_2F)). The presence of the $(CF)_n$ phase results in the second main line at -188 ppm which is the superimposition of two components: C-F bonds in $(C_2F)_n$ with neighboring CF ($C_2\underline{F}$) (CF) and C-F in $(CF)_n$ with neighboring C-F ($C\underline{F}$)(CF). Finally, the shoulder at -164 ppm is assigned to C-F bonds in $(C_2F)_n$ and $(CF)_n$ types with their neighboring non-fluorinated sp² carbon (denoted $(C_2\underline{F})(Csp^2)$ and $(C\underline{F})(Csp^2)$). This assignment follows the same trend as the slightly lower d-space of the samples annealed at 390, 410 and 430 °C. Because the content of $(C_2F)_n$ is high regardless of the fluorination temperature, the ¹⁹F MAS spectra (14 kHz) are very similar to those of F390-Foil0.1, F430-Foil0.1 and F460-Foil0.1 (Fig. SI1c†). The assignments of the lines are similar to those of F410-Foil0.1.

Natural graphite fluorinated at 440 °C with the same reaction conditions was identified as the quasi-perfect $(C_2F)_n$ phase with 96% of this phase.²¹ F440-NG was used as a reference and labelled $(C_2F)_n$ in Fig. 3. The ¹³C and ¹⁹F spectra of both foils fluorinated at 410 °C exhibit strong similarities with our reference. The characteristic lines of $(C_2F)_n$ at 42 and

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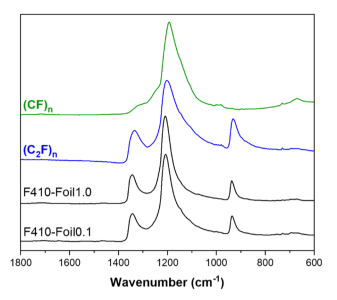


Fig. 4 IR spectra of fluorinated graphite foils (0.1 and 1.0 mm annealed at 410 °C) and reference compounds of both $(C_2F)_n$ (F440-NG sample) and (CF)_n structural types.

-178 ppm for 13 C and 19 F nuclei, respectively, are of lower intensity for both F410-Foil0.1 and F410-Foil1.0 compared to the $(C_2F)_n$ reference, which was expected since the $(C_2F)_n$ content is lower for these samples (96% for the F440-NG reference, 77% for F410-Foil0.1, and 71% for F410-Foil1.0). When compared to a typical $(CF)_n$ type, these two lines are absent.

It is also interesting to compare the infrared spectra of our reference with those of F410-Foil0.1 and F410-Foil1.0. Graphite fluorides with mainly $(CF)_n$ and $(C_2F)_n$ phases are also considered. All infrared spectra are shown in Fig. 4. The vibration of the covalent C–F bond is observed at around 1200 cm⁻¹ for both phases^{28,30} The spectra with high content of $(C_2F)_n$ are characterized by strong vibrational bands at 1350 and 940 cm⁻¹ which are absent in $(CF)_n$ type graphite fluoride. The latter band cannot be assigned to the antisymmetric elongation of >CF₂ groups, as in the literature, ³² because of its high intensity. Indeed, the NMR study was not able to evidence those groups.

Towards the densification of $(C_2F)_n$ graphite fluoride

To increase the density of fluorinated graphite, we prepared gGO foils with a high degree of graphitization, which were fluorinated under the same conditions as those used previously to promote the formation of the $(C_2F)_n$ phase. The

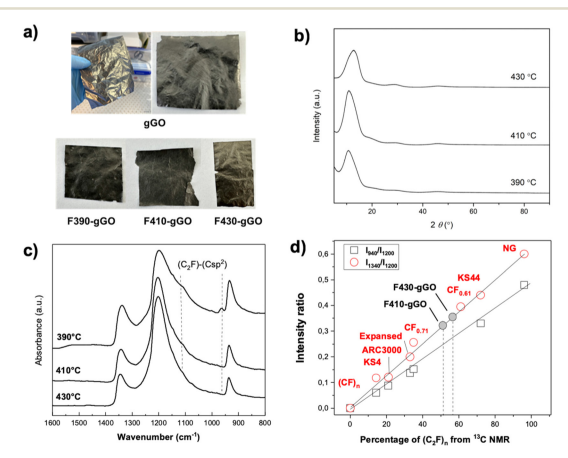


Fig. 5 Photographs of gGO foils before and after fluorination (a). XRD patterns of fluorinated gGO foils (b). IR spectra of the fluorinated gGO films (c). The intensity ratio of the lines typical of the $(C_2F)_n$ phase as a function of the percentage of the $(C_2F)_n$ phase from ¹³C NMR is plotted in (d). The CF_x compositions in (a) are extracted by the weight uptake method.

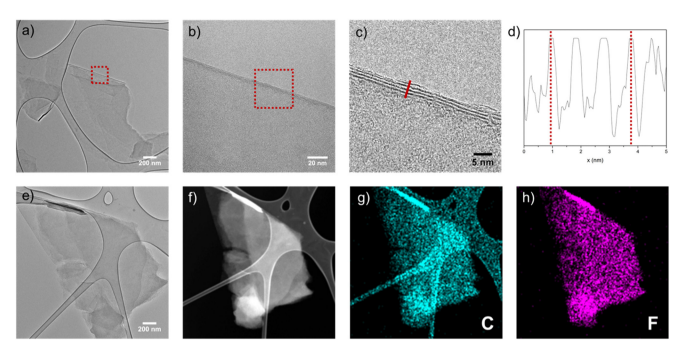


Fig. 6 (a) TEM image of a typical exfoliated platelet from F410-gGO. (b) Magnified TEM image from the indicated area in (a), showing the edge of the platelet. (c) Further magnified TEM image from the indicated area in (b), with (d) showing a plot of the profile along the line in (c). (e) TEM image of another exfoliated platelet from F410-gGO, with (f) showing the corresponding STEM-HAADF image and (g and h) showing the elemental maps of C and F.

shape/size of the foils remained unchanged after fluorination at all tested temperatures (Fig. 5a). However, we noted that the sub-metallic appearance was lost after fluorination, resulting in a greyish color for the FTF-gGO series. Despite this change, the mechanical properties were maintained, and the foils did not become more fragile after treatment. XRD analysis revealed interlayer distances of 8.2 Å and 8.1 Å for F390-gGO and F410gGO, respectively (Fig. 5b). These values are slightly lower than the expected interlayer distance for the $(C_2F)_n$ phase (9 Å). The conversion of $(C_2F)_n$ into the $(CF)_n$ phase is favored in the case of F430-gGO as revealed by the interlayer distance of 7.0 Å, which is closer to the value expected for $(CF)_n$ (6 Å). IR spectra (Fig. 5c) further confirmed the high content of the $(C_2F)_n$ phase, with prominent bands at 1350 cm⁻¹ and 940 cm⁻¹, in addition to the main band at 1200 cm⁻¹. F390-gGO showed singularities with an inclined baseline, an additional line at 964 cm⁻¹, and a shoulder at 1110 cm⁻¹, indicating the presence of non-fluorinated carbon with sp² hybridization. The line at 964 cm⁻¹ is typically present when two conditions are met: the presence of a $(C_2F)_n$ phase and a significant amount of sp² C¹². While IR spectroscopy data confirmed the high content of the $(C_2F)_n$ phase, it was unable to estimate the corresponding content accurately. Solid-state NMR analysis of FTF-gGO required milling, which could be destructive. To avoid this approach, an alternative method was employed. By utilizing the linear relationship between the intensity ratio I_{940}/I_{1200}^{21} as a function of the percentage of the $(C_2F)_n$ phase, the latter value could be graphically deduced. F410-gGO and F430-gGO exhibited percentages of the $(C_2F)_n$ phase equal to 52% and 46%, respectively. As expected from XRD data, this

content is lower for F430-gGO because of the higher conversion into the (CF)_n phase. However, this method was not suitable for F390-gGO due to both the non-zero baseline and the shoulder at 1100 cm⁻¹. The F/C ratio, *i.e.* F/C = x in CF_x, can be calculated as F/C = $1/(1 + \%(C_2F)_n)$; the values are 0.66 and 0.68 for F410-gGO and F430-gGO, respectively.

Fig. 6a shows a TEM image of a typical exfoliated platelet from F410-gGO with the highest $(C_2F)_n$ content in the series, i.e. 52%. A magnified TEM image (Fig. 6b), from the indicated area in Fig. 6a, shows the edge of the platelet, with a further zoomed-in image in Fig. 6c showing the presence of 5-6 layers. Fig. 6d shows a plot of the profile along the line in Fig. 6c. The distance between the two dotted lines is ~27 Å, indicating an average interlayer distance of ~9 Å within the lattices, consistent with the expected interlayer distance for the $(C_2F)_n$ phase. Another platelet was observed (Fig. 6e), where a high-angle annular dark-field (HAADF) image in the scanning transmission electron microscopy (STEM) mode (Fig. 6f) was acquired and the corresponding elemental maps of carbon and fluorine were obtained (Fig. 6g and h). These maps indicate the uniform presence of carbon and fluorine within the exfoliated $(C_2F)_n$.

Conclusions

Using minute control of the fluorination conditions, the partial exfoliation of graphite foil was avoided irrespective of its initial thickness, 0.1 or 1.0 mm. Long duration fluorination (24 h) at 390 °C followed by annealing (24 h) at 410 °C under

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pure F₂ gas for the same time period allowed homogeneous fluorination to be achieved. The higher content of the $(C_2F)_n$ phase, i.e. 56%, was achieved at 430 °C for thinner (0.01 mm) graphitized GO foil. Keeping the temperature in the " $(C_2F)_n$ rich domain" defined by Kita et al., 22 high contents of this phase were obtained for the two types of foils, i.e. 77 and 71% for graphite foils of 0.1 and 1.0 mm thicknesses respectively. The percentages of the $(C_2F)_n$ phase $(\%(C_2F)_n)$ were obtained from 13C NMR data, whereas the values for FFT-gGO series were estimated using the linear relationship between the IR intensity ratio I_{1340}/I_{1200} and %(C₂F)_n.²² All the samples prepared exhibit an IR spectrum typical of the $(C_2F)_n$ phase, i.e. with 3 bands at 1350, 1200 and 940 cm⁻¹. The features resulted in an interlayer distance d close to 9 Å for the 0.1 and 1.0 mm thick foils (8.6 for the FTF-gGO series). Furthermore, the content of non-fluorinated sp² carbons was very low regarding 13C NMR data. All these characteristics made the fluorinated graphite foils with mainly the $(C_2F)_n$ structural phase very promising as reflectors for slow neutrons. As evidenced by ¹⁹F NMR spectra, a substantial (CF)_n phase was observed in all samples, indicating that the $(C_2F)_n$ regions were surrounded by $(CF)_n$ ones which then appear in the XRD patterns of the samples (interlayer distance of 6 Å). Nonetheless, the average d value was close to 9 Å, slightly lower, as targeted. Graphite fluorides in the form of flexible foils of the $(C_2F)_n$ structural phase are efficient reflectors for neutrons with wavelengths higher than $\lambda_n^{\text{Bragg}(Gr)12}$. Moreover, the flexibility of the foils was maintained after fluorination which allowed the building of curved walls with the prepared fluorinated graphite foils for the desired slow neutron reflectors.

Author contributions

The investigation was mainly performed by Killian Henry, with the help of Marc Dubois, Brigitte Vigolo, Claire Herold and Sébastien Cahen. Marie Colin, Gabin Chambery, Valery Nesvizhevsky, Sylvie Le Floch, Vittoria Pischedda and Sam Chen have contributed to fluorination, characterization, neutron science, high pressure measurements, high pressure measurements and the work on GO membranes, respectively. Writing - original draft - was performed by Marc Dubois and Killian Henry by adding also parts from Valery Nesvizhevsky and Sam Chen. Writing - review and editing - was performed by Marc Dubois and Sébastien Cahen. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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