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### Publication date

02-07-2024

### Published in

Communications Materials, 2024, 5 article number 115

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Konyashin, I., Muydinov, R., Cammarata, A., Bondarev, A., Rusu, M., Koliogiorgos, A., Polcar, T., Twitchen, D., Colard, P.-O., Szyszka, B. and Palmer, N. (2024) 'Face-centered cubic carbon as a fourth basic carbon allotrope with properties of intrinsic semiconductors and ultra-wide bandgap', available: <https://doi.org/10.34961/researchrepository-ul.26539858.v2>.

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<https://doi.org/10.1038/s43246-024-00547-8>

# Face-centered cubic carbon as a fourth basic carbon allotrope with properties of intrinsic semiconductors and ultra-wide bandgap



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Carbon is considered to exist in three basic forms: diamond, graphite/graphene/fullerenes, and carbyne, which differ in a type of atomic orbitals hybridization. Since several decades the existence of the fourth basic carbon allotropic form with the face-centered cubic (*fcc*) crystal lattice has been a matter of discussion despite clear evidence for its laboratory synthesis and presence in nature. Here, we obtain this carbon allotrope in form of epitaxial films on diamond in a quantity sufficient to perform their comprehensive studies. The carbon material has an *fcc* crystal structure, shows a negative electron affinity, and is characterized by a peculiar hybridization of the valence atomic orbitals. Its bandgap (~6 eV) is typical for insulators, whereas the noticeable electrical conductivity (~0.1 S m<sup>-1</sup>) increases with temperature, which is typical for semiconductors. Ab initio calculations explain this apparent contradiction by noncovalent sharing *p*-electrons present in the uncommon valence band structure comprising an intraband gap. This carbon allotrope can create a new pathway to ‘carbon electronics’ as the first intrinsic semiconductor with an ultra-wide bandgap.

For more than 60 years scientists have been aware of an unusual allotropic form of carbon with the face-centered cubic (*fcc*) structure but could not elucidate it and recognize it as an individual phase. Up to date a great number of studies indicating the existence of this carbon allotrope emerged<sup>1–22</sup>, nevertheless, according to ref. 1 its crystal structure “...remains a mystery”.

First crystallographic evidence for the presence of *fcc*-carbon in synthetic diamonds was reported in the classical work of ref. 2. The X-ray diffraction (XRD) from diamond grits obtained at high temperatures and ultra-high pressures showed, apart from conventional for diamond diffraction pattern, also strong (200)-, (222)-, and (420)-reflexes which are forbidden for the diamond crystal lattice. Besides that, there were side

satellite diffraction maxima found for all diamond reflexes. These additional maxima form the pattern of a face-centered cubic crystal structure.

Subsequently, an initially amorphous film, which was produced via carbon condensation assisted by the bombardment with Ar<sup>+</sup>-ions and annealed at a low temperature, contained nanoparticles of *fcc*-carbon<sup>3</sup>. According to electron diffraction results the lattice constant of the *fcc*-carbon particles was found to be 3.57 Å, which is close to that of diamond (3.567095 ± 0.000017 Å according to ref. 23). The authors of ref. 3 assumed that this carbon allotrope is a metallic form of carbon based on the crystal structure itself and closeness of the lattice parameter to that of typical *fcc*-metals. However, as it was shown in ref. 4, the metallic nature of *fcc*-carbon is out of the question due to the similarity of its crystal lattice parameter to that

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of typical fcc metals (e.g., Cu and Ni) at a significantly lower atomic radius of carbon compared to that of such fcc metals.

Later, this carbon allotrope was also synthesized at ultra-high pressures<sup>4–7</sup>. In 1991 Hirai and Kondo obtained nanocrystals of *fcc*-carbon by rapid cooling following the shock compression of graphite sheets<sup>5</sup>. The authors introduced for the first time the term “*n-diamond*” (new-diamond) to designate this carbon allotrope, when considering the closeness of its lattice parameter to that of diamond but ignoring the occurrence of the reflexes forbidden for the diamond crystal lattice. In 1994 the uncommon carbon allotrope was obtained by the dynamic shock compression of a diamond/amorphous carbon mixture<sup>6</sup>. In the same year *fcc*-carbon was reported to be produced from natural graphite at high temperatures and static ultra-high pressures<sup>7</sup>.

Chemical vapor deposition was also employed for obtaining this form of carbon<sup>8,9</sup>. Nanoparticles of *fcc*-carbon were produced in the early 2000s by etching a diamond single crystal in a hydrogen plasma and also by plasma assisted chemical vapor deposition<sup>10–12</sup>. In particular, experimental evidence based on the density of electron states and electron energy loss spectroscopy (EELS) was obtained in ref. 12 clearly indicating that *fcc*-carbon fundamentally differs from diamond with respect to the nature of its chemical bonds<sup>12</sup>.

Chlorination of SiC and other carbides also resulted in obtaining this carbon allotropic form in the early 2000s<sup>13,14</sup>. The authors of refs. 13,14 designated it as “diamond-structured carbon” and, later, the term “carbide-derived carbon” was introduced and is presently employed in the literature.

The first evidence for the existence of *fcc*-carbon in nature, particularly as nanoparticles in the asphaltene precipitated from a crude oil from Mexico, was reported in 2004<sup>15</sup>. Afterwards, in the late 2000s—early 2010s, a number of publications<sup>16–21</sup> provided evidence for the presence of *fcc*-carbon nanoparticles in nature (soils, rocks, Greenland ice, etc.). It was suggested that such nanoparticles are likely to form in Earth’s mantle and are transferred to Earth’s surface during volcanoes’ eruptions, so they have a terrestrial origin<sup>4</sup>.

According to the results of electron diffraction reported in the literature this carbon allotrope has a space group  $Fm\bar{3}m$  (group number 225) with a lattice parameter of about 3.54–3.57 Å. Nevertheless, because of the closeness of its lattice parameter to that of diamond, in many publications on its synthesis<sup>1,5–7,13,14</sup> and existence in nature<sup>15–21</sup>, the carbon allotrope was considered as a diamond polymorph and designated as “*n-diamond*”, so its crystal structure is still a matter of discussions<sup>22</sup>.

The issue of the existence of *fcc*-carbon was also a subject of several publications on ab initio calculations<sup>24–28</sup>. This structure was recognized as less stable than that of diamond, and the experimentally established crystal lattice parameter could not be confirmed by calculations from the first principles<sup>24–27</sup>. In one published work<sup>27</sup> the occurrence of *fcc*-carbon was explained by hydrogen doping of diamond; however, this assumption predicted the XRD pattern different from that obtained experimentally. In contrast to refs. 24–27, the basic theoretical results of ab initio calculations reported in ref. 28 justified the existence of the carbon allotrope with the face-centered cubic structure and predicted its lattice constant matching experimental data.

The major objective of the present work was to obtain *fcc*-carbon in a quantity sufficient to perform its comprehensive study by different methods and unambiguously establish its crystal and electronic structure. To explain the established peculiar electronic properties and electronic structure of this carbon allotrope ab initio calculations were carried out.

## Results

### Synthesis of *fcc*-carbon films

Thin films of *fcc*-carbon were obtained on single-crystalline diamond substrates by plasma-assisted chemical vapor deposition (PACVD) from CH<sub>4</sub>-H<sub>2</sub> gas mixtures at very special conditions given in Methods. An exclusive formation of *fcc*-carbon instead of diamond is explained by faster etching rate of the latter. This happens at insufficient for the diamond deposition presence of metastable C-H species combined with unusually low substrate

temperature. Still, the decomposition of such carbon species occurs with a deposition rate of ~0.14 nm/min. In a very narrow window of deposition conditions this decomposition results in the formation of the *fcc*-carbon phase, which presumably has lower susceptibility to etching by atomic hydrogen compared to that of diamond. The sample surface is shown in Supplementary Information, Fig. S1.

### Crystal structure

Figure 1a and Table 1 give the electron diffraction pattern from nm-sized fragments obtained from the single-crystalline film by the mechanical detachment (scratching) with a Cu grid. The pattern clearly indicates that the carbon film has a face-centered cubic crystal lattice with the space group  $Fm\bar{3}m$ , as it comprises all the reflections forbidden for the diamond crystal lattice. It is worth noticing that the material revealed a remarkable plasticity when it was mechanically detached from the film.

The XRD pattern also shows reflexes forbidden for the diamond crystal lattice (Fig. 1b, c and Table 1). The fact that some reflexes forbidden for the diamond crystal lattice are split (for example, the (200)-reflex shown in Fig. 1b) can be explained by the nonuniform distortion of the cubic crystal lattice. This phenomenon is discussed and explained in Supplementary Information, Fig. S2 and Fig. S3).

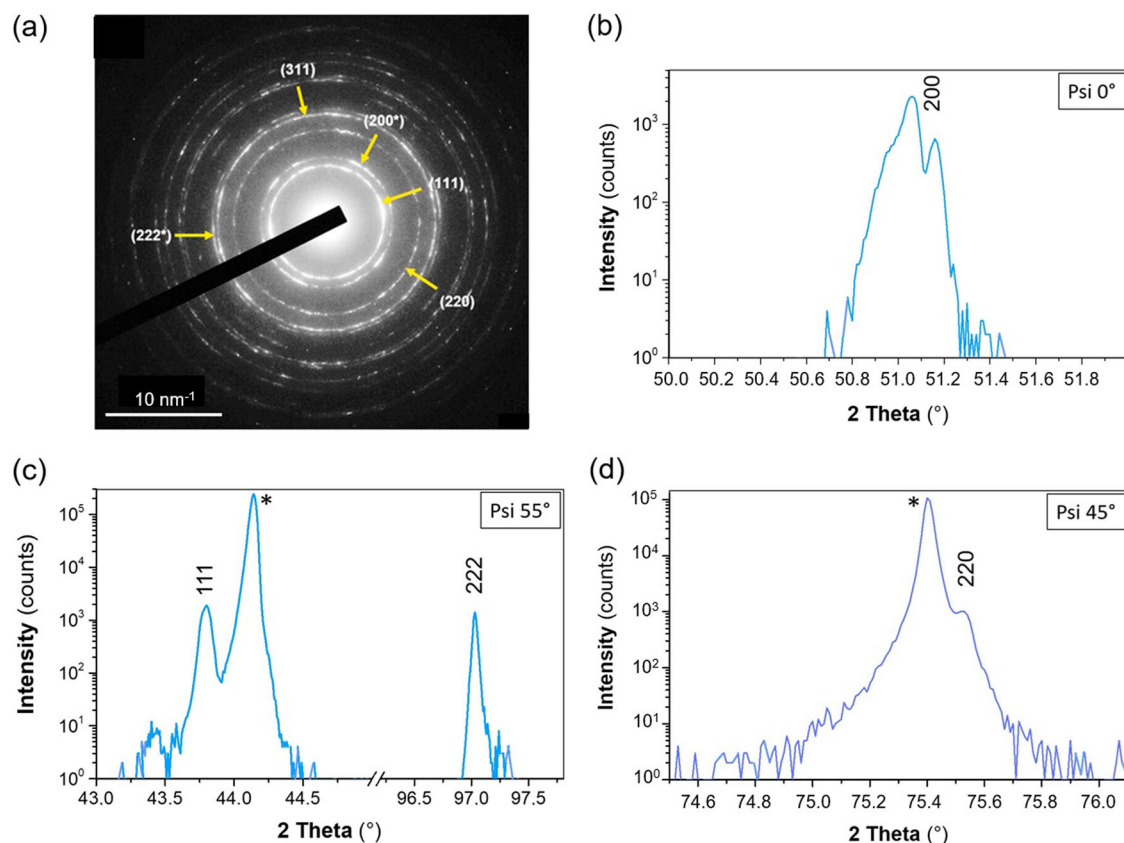
As one can see from all the acquired diffraction patterns (Fig. 1b–d), the *fcc*-carbon film grows on the (100)-diamond substrate epitaxially. No additional peaks were detected in the classical grazing incidence procedure (incident angle varied in the range 0.5–5°) or Bragg-Brentano XRD experiments. We also point out that no diffraction maxima in the 10°...30° 2θ-range, which could arise from graphite, graphene/carbon nanotubes or oxidized graphene, were detected. A comparison of the Phi-scans and rocking curves (see Supplementary Information, Fig. S4 and Fig. S5) for the diffraction maxima of the same system of planes in diamond and in *fcc*-carbon uniquely corroborates the epitaxial relation between the film and the substrate.

The high-resolution transmission electron microscopy (HRTEM) image (Fig. 2a, left) also confirms the face-centered cubic crystal structure of the film. It provides, as it is shown in Fig. 2a, the interplanar distance of roughly 2 Å, corresponding to the distance between (200) atomic planes in the face-centered cubic lattice schematically illustrated by the (110)-projection of an elementary cell (Fig. 2a, right). The fast Fourier transform (FFT) and inverse FFT images shown in Fig. 2b indicate a similar distance (~1.8 Å), which is even closer to the expected one. The determined from interatomic distances {111} and {200} planes build the angle of ~55°, that well corresponds to the face-centered crystal structure. A combination of the interatomic distances and angles determined from the HRTEM image unambiguously rules out the diamond structure.

The lattice spacing values calculated from our experimental diffraction patterns are summarized and compared with the literature data in Table 1. The free-standing *fcc*-carbon particles reveal the lattice spacings close to those reported in literature. On the contrary, the epitaxial film on diamond shows relatively large deviations of the lattice spacings, and they cannot be consistently explained by the existence of strains, which supports the abovementioned plasticity of the material.

### Spectroscopic characteristics

The Raman spectrum recorded from mechanically detached fragments of the film is shown in Fig. 3a. As one can see, the Raman spectrum does not comprise any signals. It is well known that materials with a face-centered cubic crystal lattice, the primitive unit cell of which contains only a single atom, have only acoustic phonons and no optical phonons<sup>29,30</sup>. It should be noted that the presence of nanodiamonds in the film would manifest a couple of broad vibrational modes and their presence can therefore be discarded. Accordingly, the Raman spectrum from the carbon film on the diamond substrate (Fig. 3b) comprises no material related signals; only the strong peak from the diamond substrate is visible due to the transparency of the *fcc*-carbon film. Some increase of FWHM after film deposition is consistent with the dramatic change of the substrate surface geometry and roughness. Diamond substrate reveals apart from the main, characteristic



**Fig. 1 | Diffraction patterns obtained from the carbon film.** **a** Electron diffraction pattern from nm-sized fragments obtained from the single-crystalline film by the mechanical detachment (scratching) with a Cu grid. **b–d** X-ray diffraction maxima obtained from the film at different tilting angles Psi. Note that the (200)- and (222)-

reflexes are forbidden for the diamond crystal lattice. The (111)- and (220)-reflexes exist for both phases; the peaks pointed by starlets and having significantly higher intensity correspond to the diamond substrate.

**Table 1 | Lattice spacings  $d$  and corresponding  $hkl$  indices attributed and those based on the diffraction patterns given in refs. 5,10 and in the present work to the fcc-carbon**

$hkl$	$d_{\text{exp}}$ (Å) from ref. 10	$d_{\text{calc}}$ (Å) adopted from ref. 10 at $a_0 = 3.563$ Å	$d$ (Å) from ref. 5	$d$ (Å) from the present work
111	2.067	2.057	2.06	$2.058^a/2.076 \pm 0.005^b/2.05017^c$
200 <sup>d</sup>	1.791	1.781	1.78	$1.786^a/1.787 \pm 0.0015^b/1.77550^c$
220	1.261	1.260	1.26	$1.266^a/1.257 \pm 0.0008^b/1.25546^c$
311	1.078	1.074	1.07	$1.074^a$
222 <sup>d</sup>	1.032	1.029	1.04	$1.030^a/1.0298 \pm 0.005^b/1.02508^c$
400	0.892	0.891	0.896	$0.893^a/0.891 \pm 0.0015^b/0.88775^c$
331	0.817	0.817	0.818	$0.812^a/-/1.07066^c$
420 <sup>d</sup>	0.796	0.797	0.796	$0.783^a$
422	0.727	0.727	0.726	$0.720^a$
331/511	0.686	0.686	0.683	$0.680^a$
440	0.630	0.630	-	$0.630^a$
531	0.601	0.602	-	$0.550^a$

<sup>a</sup>Based on the electron diffraction pattern shown in Fig. 1a.

<sup>b</sup>Based on the XRD results (Fig. 1b–d).

<sup>c</sup>Based on the ab initio calculations (simulated from the charge density of fcc-carbon).

<sup>d</sup>Reflexes forbidden for the diamond crystal lattice.

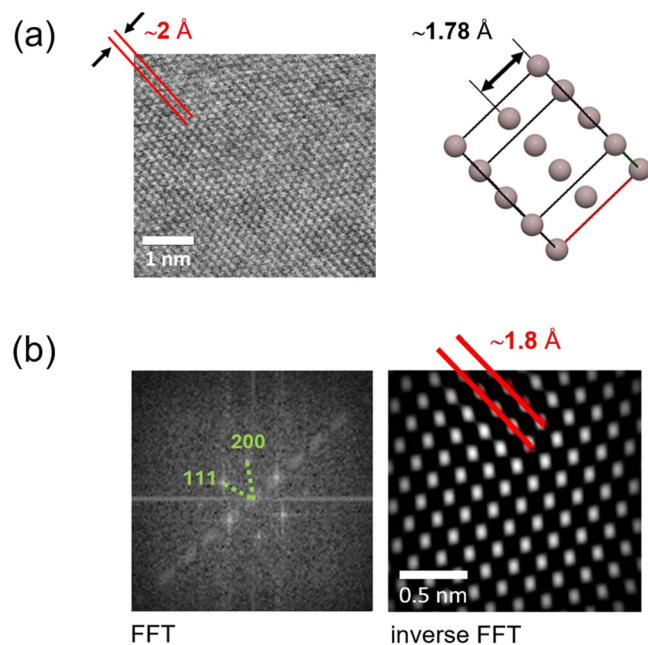
for diamond signal, some additional modes that have several orders of magnitude lower intensity and can be ascribed to the presence of structural defects at the diamond surface.

Vacuum ultraviolet reflection spectrum presented as measured for the film and as a Tauc-plot reconstructed based on the Kubelka-Munk theory is shown in Fig. 3b. Based on these results a value of bandgap of the carbon film

is found to be nearly 6.1 eV. As the transmittance data were not affordable the magnitude of absorption coefficient is not given here. We detected a nonzero background which is explained by a depleted occupation of the valence band (see *Ab initio calculations*).

The analysis of the XPS spectrum presented in Fig. 3d is nontrivial. The spectrum shows a significant degree of surface contamination since the





**Fig. 2 | Results of HRTEM studies of the fcc-carbon film.** **a** HRTEM image of the fcc-carbon crystal lattice in the (110) orientation (left) and schematic (110)-projection of the fcc-elementary cell (right); **b** fast Fourier transform (FFT) image with the indicated angle between the {111} and {200} planes (left) and inverse FFT image (right).

sample was exposed to air. No  $sp^2$  contribution was detected that allows excluding any influence of graphitized fraction on the film properties, particularly on the electrical conductivity. The major peak C1s is present at an untypically low value of 283.9 eV providing evidence that there are electrons in the system, which are weaker bound than if any kind of covalent bonds would exist. The contributions at 286.2 and 288.7 eV are attributed to the carbon chemical states respectively in the C-O-C and O-C=O species. The peak at  $\sim 284.5$  eV which we attributed to an adventitious carbon (C-C, C-H) has almost the same position as the  $sp^2$ -contribution, which was used for the calibration (see *Methods*). In the latter case, however, the  $(\pi-\pi)$ -satellite would show up and the good match of the binding energy (BE) seems to be incidental. Indeed, the BE measured for adventitious carbon was found to vary from 284.08 to 286.74 eV depending on the substrate<sup>31</sup>. The peak, which we ascribe as originating from fcc-carbon ( $\sim 283.9$  eV), was fit with the asymmetric line shape because of the material electrical conductivity (see, e.g., ref. 32). Such an asymmetry can arise from the multi-electron excitations and electron-hole pairs creation in the delocalized subvalence band, which is believed to comprise nearly unhybridized  $s$  and  $p$  electrons (see *Ab initio calculations*). Note that the symmetric peak at 284.8 eV, which is typical for diamond, and signals from  $\pi-\pi^*$  bonds in the range between  $\sim 290$  and 293 eV, which are typical for  $sp^2$ -hybridized carbon<sup>33</sup>, are absent in the spectrum.

The results of Fourier-transform infrared spectroscopy (FTIRS) also clearly indicate a difference in the nature of chemical bonds between diamond and fcc-carbon and provide evidence for the absence of carbon-hydrogen bonds in the electronic structure of fcc-carbon (see Supplementary Information, Fig. S6). Thus, the explanation of the crystal structure of fcc-carbon due to the presence of hydrogen atoms between adjacent carbon atoms suggested in ref. 27 is confirmed to be incorrect.

### Electrical conductivity and electronic structure

It was reported in ref. 22 that fcc-carbon is characterized by electrical conductivity. Based on its crystal structure it was suggested to have a metallic character of conductance. Our comprehensive measurements clearly demonstrated that the material is a semiconductor (Fig. 4). The conductivity

at room temperature is reasonably high ( $\sim 10^{-3} \text{ S cm}^{-1}$ ). As noted in *Methods* the Hall voltage could not be measured reliably. Besides the unfavorable for this method film shape, the concentration of free charge carriers in fcc-carbon is expected to be fairly low. Therefore, its reliable determination is the matter of a special sample preparation which is beyond the scope of this study.

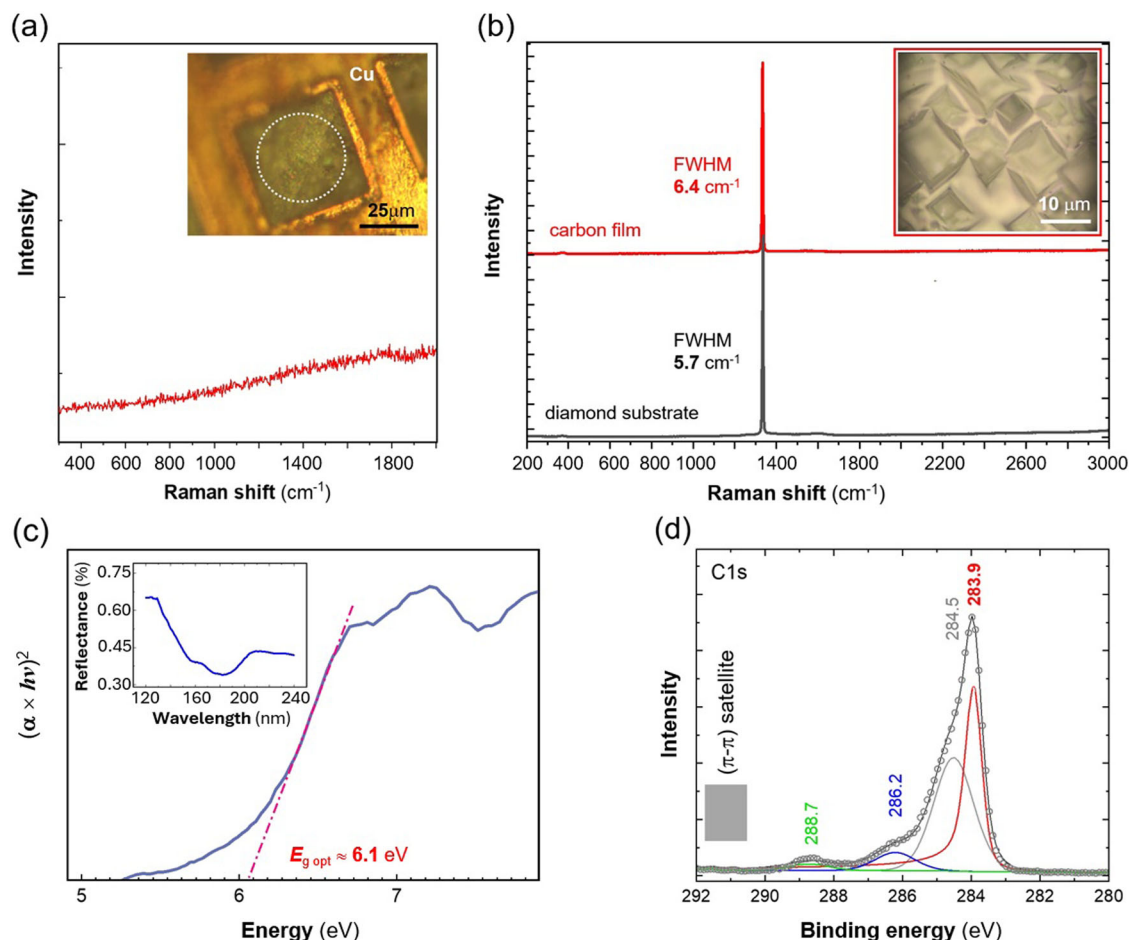
The extrapolated value of the activation energy ( $\sim 230$  meV) is unusually high if compared to classical semiconductors. It is known that both chemical state and size of crystallites (flakes) affect this energy. In polycrystalline and composite carbon materials, the activation energy is characterized by the hopping mechanism and electrons' tunneling<sup>34</sup>. The high value of  $E_a$  may be caused by a poor percolation of electron conducting domains<sup>35,36</sup>. Comparable magnitudes of  $E_a$  are demonstrated also by organic semiconductors, in which the ionization of molecules is a crucial stage of the conductivity thermal activation<sup>37</sup>; this is assumed to indicate the similarity of the nature of chemical bonds in fcc-carbon and inter-molecular bonds in organic semiconductors (see below).

The optical bandgap established for the carbon film is nearly 6.1 eV. The estimation based on the EELS published in our previous work<sup>12</sup> gives the bandgap value close to 7 eV. It is known that an electronic bandgap is usually larger than an optical gap due to the need to overcome the Coulomb repulsion energy when electrons are injected into the material. In our case the difference may also result from a crystal orientation which was (100) in case of the optical measurements and uncertain in electronic measurements. The established discrepancy between the values of bandgap obtained by different methods may also relate to the weird shape of the film (see *Supplementary Information* for details). In fact, the bandgap is strongly dependent on the interatomic distance and increases if atoms are remote from each other and the degree of electrons' delocalization decreases. In that sense, fcc-carbon shows consistently larger bandgap than that of diamond.

A precise determination of the work function ( $\phi$ ) is challenging for the diamond surface owing to several factors described in refs. 38–40. Among them possible O- or H-termination, presence of graphitic carbon and carbon-containing adsorbates. Moreover, the crystallinity (crystal size) and crystallographic orientation also have an impact on the ionization energy<sup>40</sup>. To avoid surface adsorbates originated from air and consequent changes of electronic properties, the Kelvin probe (KP) was performed and photoelectron yield spectroscopy (PYS) measurements according to ref. 39 in a dry nitrogen atmosphere were carried out. Prior to the KP measurements the surface of the samples were acid cleaned to eliminate dangling hydrogen atoms and weakly bound adsorbates from ambient air as described in *Methods*.

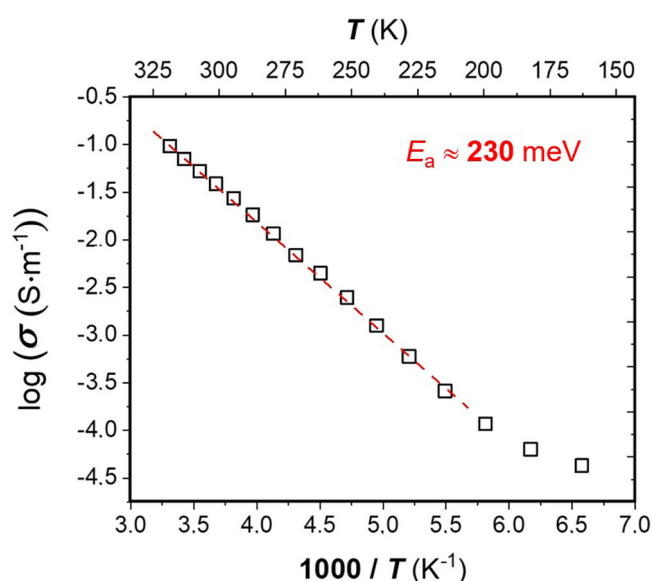
The results of the work function ( $\phi$ ) measurements by using the Kelvin probe method show quite a large difference between the measured curves for fcc-carbon and diamond (Fig. 5a). While a value of 4.74 eV is obtained for the diamond substrate, which agrees well with the published data (between 4.2 and 5.4 eV for the (100)-surface of diamond<sup>39</sup>), a significantly lower work function of 4.60 eV is determined for the fcc-carbon film. Note that no difference was observed between measurements in the dark and under illumination (Fig. 5a), namely, no surface photovoltage (SPV) related to the surface band bending is observed. This can be explained by the flat band conditions or Fermi level pinning at the film surface<sup>41</sup>.

The ionization energy, which determines the valence band maximum (VBM), was found from PYS spectra shown in Fig. 5b, which was obtained due to photoexcitation of electrons by photons in the energy range of 3.8–7.4 eV. Near the emission onset, the photoelectron yield  $PY(h\nu)$  can be approximated by  $PY(h\nu) \propto (h\nu - E_i)^3$ , where  $h\nu$  is the energy of the incident photon and  $E_i$  is the ionization energy of the surface<sup>42</sup>. The ionization energy of the surface is thus determined by a linear fit of the  $PY(h\nu)^{1/3}$  onset, as shown by the dashed lines in Fig. 5b; the results are summarized in Table 2. As it can be seen, compared to the VBM position of diamond, such a position for the carbon film is higher by 350 meV. Note that the work function of the carbon film is lower by 140 meV. Consequently, the calculated VBM position of the carbon film relatively to the Fermi level, according to  $E_F - E_{VBM} = \phi - E_i$ , is shifted downwards by about 500 meV, laying at



**Fig. 3 | Spectra measured for the *fcc*-carbon samples.** **a** Raman spectrum obtained for mechanically detached fragments of the film located in a cell of a Cu grid; the dashed circle marks the area of analysis. **b** A comparison of Raman spectra obtained for the as-deposited epitaxial film on diamond substrate and the bare diamond substrate. Insets show the corresponding optical images. **c** Vacuum ultraviolet

(VUV) reflection spectrum presented as measured for the film (inset) and as a Tauc-plot reconstructed on the basis of the Kubelka-Munk theory (see Methods). **d** X-ray photoelectron spectrum (XPS) obtained for the as-deposited carbon film after its fitting (the fitting procedure is described in Methods).



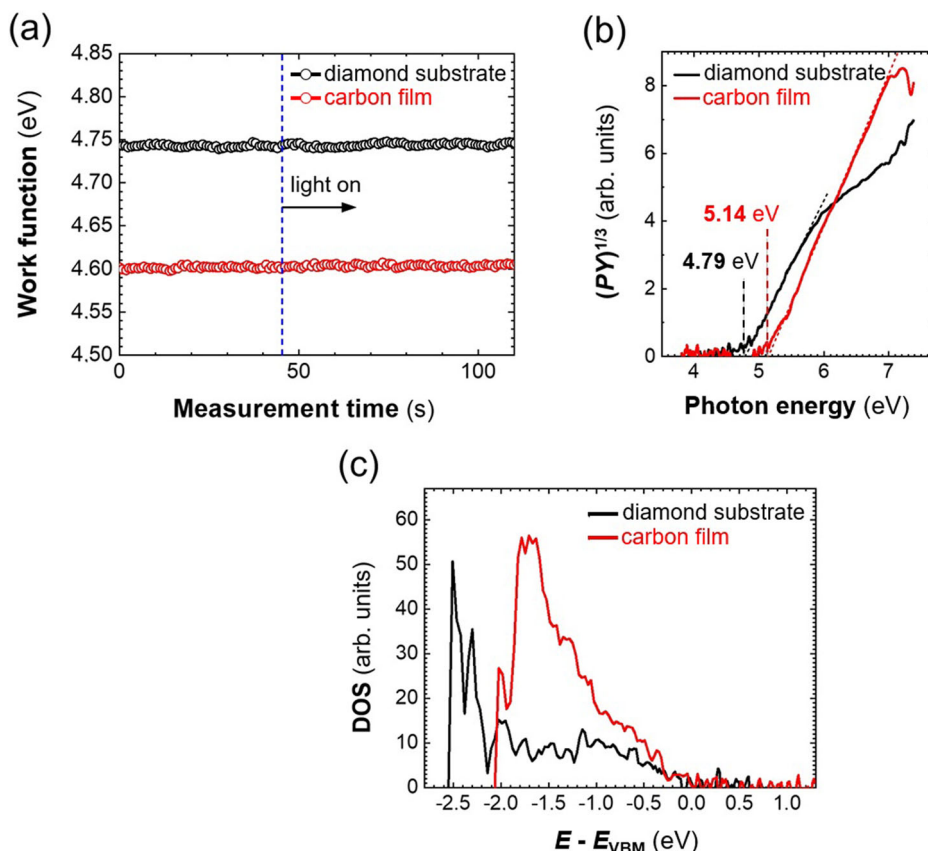
**Fig. 4 | Temperature dependence of electrical conductivity measured for the *fcc*-carbon film.** Activation energy was determined for the linear correlation range. The film thickness was set to 1 μm.

0.54 eV below the Fermi level. In contrast, the VBM of diamond lays only 0.05 eV below the Fermi level.

The electron affinity  $E_A$ , which defines the conduction band maximum, is determined as  $E_A = E_i - E_g$ , where  $E_g$  is the material bandgap. Considering the diamond bandgap of about 5.5 eV according to ref. 43 we obtained for diamond the value of  $E_A = -0.68$  eV. This negative value is in between the values of  $-0.82$  eV and  $-0.45$  eV measured on the same (100) crystal plane of diamond with H- and O-terminations respectively according to ref. 39. Other studies showed for H-terminated diamonds a strongly negative  $E_A$  value in vacuum of up to  $-1.3$  eV (see, e.g., ref. 44). On the other hand, O-terminated diamonds often showed a positive  $E_A$  value due to the strong electronegativity of the carbonyl groups involved<sup>45</sup>. A negative  $E_A$  value was also established for the *fcc*-carbon film by considering its bandgap of 6.1 eV determined from the optical measurements (see above). Thus, due to the negative electron affinity an easy emission of electrons from the *fcc*-carbon films is thought to be possible.

In addition to the ionization energy, the PYS spectra contain information on the electronic density of states (DOS) spreading from the Fermi level ( $E_F$ ) down to  $h\nu - \Phi$  below  $E_F$ . The DOS spectra were obtained from the  $PY(h\nu) - h\nu$  plots as the first derivative of the photoelectron yield  $PY(h\nu)$  with respect to the photon energy ( $h\nu$ ), thus as  $d(PY)/d(h\nu)$ . The experimentally obtained DOS are presented in Fig. 5c, which shows a higher density of states near the VBM and by about five times higher DOS at 1.5 eV below VBM for the *fcc*-carbon film. This can be an indication for higher holes effective mass in the carbon film compared to that of diamond. As it is

**Fig. 5 | Electronic properties of the *fcc*-carbon film compared to the diamond substrate.** **a** Work function measured by Kelvin probe (KP) in darkness and under illumination at  $\lambda = 172$  nm. **b** Photoemission spectra  $PY(h\nu)^{1/3}$  measured by the photoelectron yield spectroscopy (PYS), in which the thin dashed lines represent the linear fits for the determination of the ionization energy. **c** Experimentally established electronic density of states (DOS) spectra that were acquired from PYS measurements (the energy of the valence band maximum (VBM) is set to 0 eV).



shown in the next section, this is consistent with the calculated electronic band structure of the diamond and *fcc*-carbon structures.

### Ab initio calculations

The major objective of this part is to obtain a computational model of *fcc*-carbon describing its electronic structure and explaining its electrical conductivity at the large value of bandgap.

Our computational model was built within the framework of the Density Functional Theory. The goal of the model is to explain some properties of *fcc*-carbon that were experimentally observed. An extensive benchmark of different energy functionals was conducted in order to reproduce the electronic properties of the *fcc*-carbon (see Methods, Computational Models), and the revPBE DFT functional<sup>46</sup> was finally selected, which reproduces our preliminary experimental findings.

The electronic band dispersion of *fcc*-carbon resembles that of one of pure diamond (Fig. 6a–d).

The indirect band gap of diamond is found to be 4.0 eV, which is smaller than the experimental value of nearly 5.5 eV (see above). It is known that DFT functionals tend to underestimate the gap width<sup>47</sup>; however, here the focus is on the comparison of the valence band dispersion between the

two materials and, therefore, such difference does not affect the forthcoming discussion.

The valence band of *fcc*-carbon is separated from the conduction band by a wide band gap (about 7.1 eV), which is consistent with the value determined experimentally (see above). The empty valence states closest to the Fermi level are found to generate a very low density of delocalized electrons only in the region located between carbon atoms, that is, such electrons are free to move throughout the whole crystal lattice (Fig. 6e). For this reason, despite the large band gap value, *fcc*-carbon is expected to have properties of an intrinsic semiconductor that is confirmed by the conductivity measurements (Fig. 4).

Unlike diamond, the valence band in *fcc*-carbon is split into a lower and an upper part by an intraband gap of about 0.8 eV. Those sub-bands are occupied by *s*- and *p*-electrons respectively. It can therefore be expected that the photoelectron spectroscopy resolves two signals for *fcc*-carbon. The recorded XPS spectra (see Fig. 3d), show a broad asymmetric peak which can be explained as a result of the convolution of two peaks occurring at 283.9 and 284.5 eV, respectively.

According to the theoretical model, the specific constitution of the *s*- and *p*-electrons' hybridization leads to the formation of low-energy states in the range [−19.4, −11.2] eV with the predominant *s*-character, which is quantified as 93% contribution to the total density of states (see Fig. 7b). Moreover, this energy range is separated from the rest of the valence band by a small gap of 0.8 eV. It is known that a large symmetric distribution of the valence *s*-electrons induces a smaller binding energy of the core *s*-electrons<sup>48</sup>. For this reason, we detect the C1s peak with unusually low binding energy of 283.9 eV. The high energy (284.5 eV) peak represents then the impact of the shared (conducting) predominantly *p*-electrons from the upper valence sub-band. As mentioned above, this peak contains the signal from an adventitious carbon as well.

This leads to a peculiar type of  $|sp^3\rangle$  hybridization, which is derived from the analysis of the density of states (Fig. 7). The lowest band can be

**Table 2 | Surface work function  $\phi$  measured by Kelvin probe, ionization energy  $E_i$  determined by PYS, electron affinity  $E_A$ , and the VBM position in relation to the Fermi level**

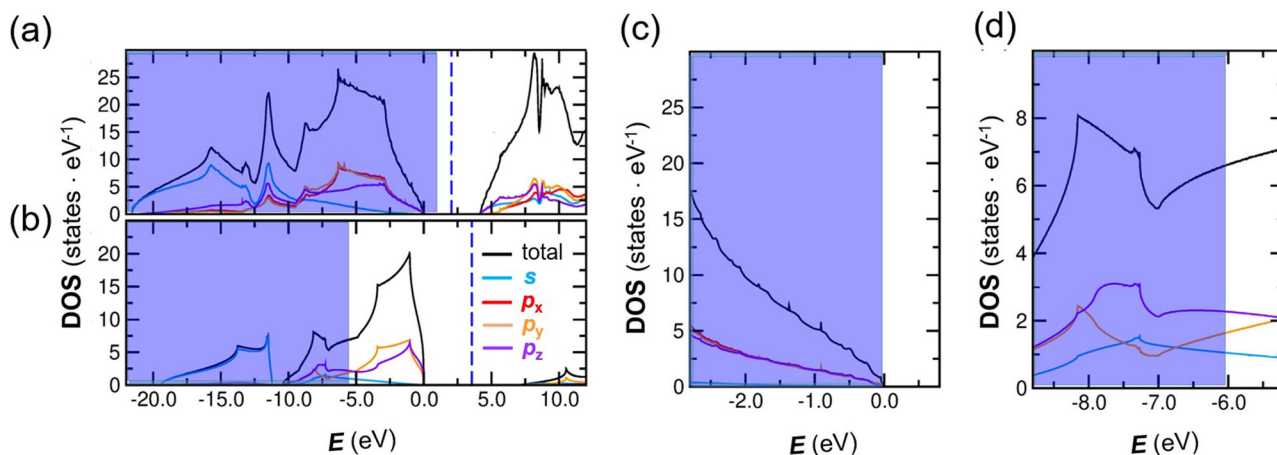
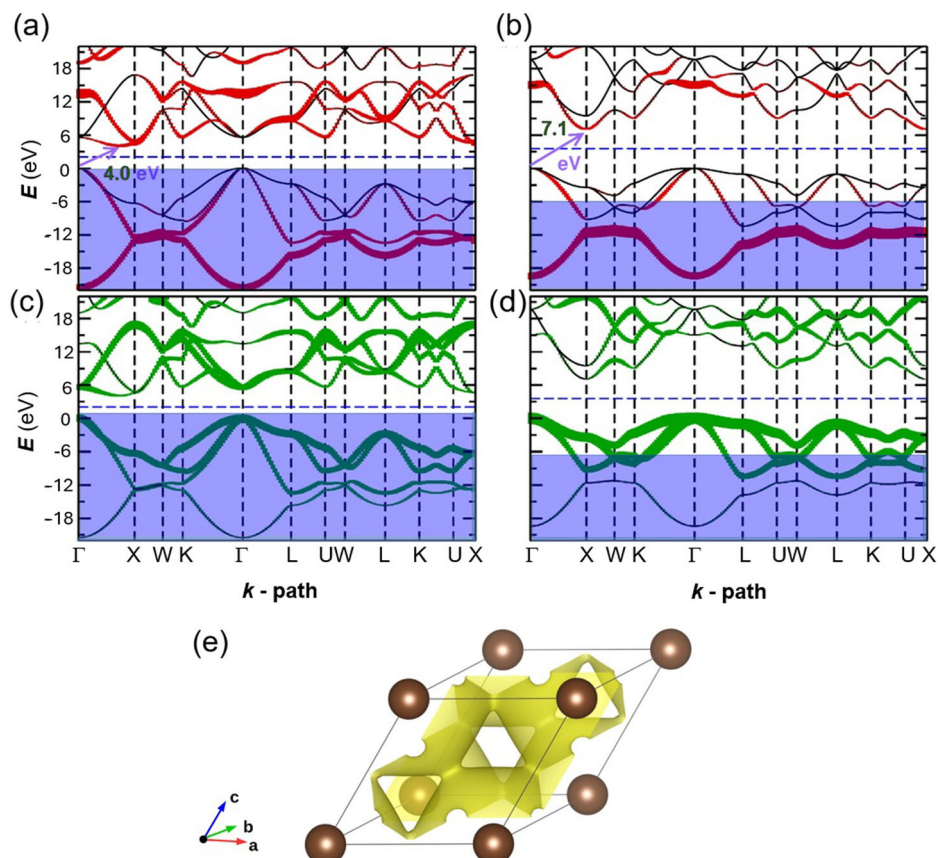
sample	$\phi$ ( $\pm 0.04$ eV)	$E_i$ ( $\pm 0.03$ eV)	$E_A$ ( $\pm 0.04$ eV) <sup>a)</sup>	$E_F - E_{VBM}$ ( $\pm 0.04$ eV) <sup>b)</sup>
diamond	4.74	4.79	-0.68	-0.05
carbon film	4.60	5.14	-0.96	-0.54

<sup>a)</sup>  $E_A = E_i - E_g$  with  $E_{g\text{-diamond}} = 5.47$  eV and  $E_{g\text{-carbon film}} = 6.1$  eV.

<sup>b)</sup>  $E_F - E_{VBM} = \phi - E_i$ .



**Fig. 6 | Calculated electronic band structure.** **a–c** Diamond and **b–d** *fcc*-carbon are compared. The red and green lines in the top and bottom panels show the *s*- and *p*-character of the bands along the *k*-path respectively, the thicker the colored line, the larger the character. The purple arrow indicates the predicted optical band gap with its value reported on the side. The blue shadowed area indicates the filling of the valence band. **e** Isosurface (0.018 e/Å<sup>3</sup>) of the *fcc*-carbon electron density generated by the upper part of the valence band; the shape of the isosurface shows the electron delocalization across the crystal lattice.



**Fig. 7 | Calculated total and orbital-resolved density of states (DOS).** DOS of **(a, c)** diamond and **(b, d)** *fcc*-carbon. In **(b)** and **(d)**, the curves corresponding to the *p<sub>x</sub>* and *p<sub>y</sub>* orbitals overlay. The VBM is set to 0 eV and the Fermi level as defined for intrinsic semiconductors is shown by a dashed line. The blue shadowed area indicates the

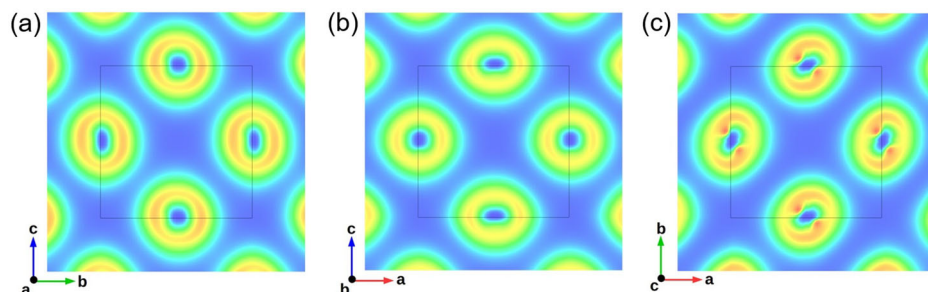
filling of the valence band as in Fig. 6. If the Bloch states of the valence band are projected on the hydrogen-like functions centered at the carbon site, it is observed that the lowest band below the intraband gap has a prevalent *s*-character (Fig. 6b), while the remaining three bands have a predominant *p*-character (Fig. 6d).

represented by the superposition  $|sp^3\rangle_{\text{lowest}} = 0.93|s\rangle + 0.01|p_x\rangle + 0.01|p_y\rangle + 0.05|p_z\rangle$ , while the remaining occupied states above the intraband gap can be described by the linear combination  $|sp^3\rangle_{\text{bond}} = 0.16|s\rangle + 0.24|p_x\rangle + 0.24|p_y\rangle + 0.36|p_z\rangle$ . The electronic configuration of *fcc*-carbon is thus characterized by a very low rate of hybridization of *s* and *p* electrons, in which each electron occupies one orbital in the configuration  $[2s^1 2p^3]$ . The overall hybridization for the valence band of *fcc*-carbon is  $|sp^3\rangle = 0.26|s\rangle + 0.25|p_x\rangle + 0.25|p_y\rangle + 0.24|p_z\rangle$ , while in the case of diamond is  $|sp^3\rangle = 0.31|s\rangle + 0.23|p_x\rangle + 0.23|p_y\rangle + 0.23|p_z\rangle$ ; equivalently, in *fcc*-carbon the character of the valence band is 26% of *s*-kind and 74% of *p*-kind, while

in the case of diamond is 31% and 69%, respectively. This is consistent with the fact that the C–C distance in *fcc*-carbon (2.51 Å) is larger than the interatomic distance in diamond (1.54 Å). In fact, it is known that the degree of hybridization, which is the percent contribution of the *s* and *p* orbitals to the hybridized ones, determines the C–C distance; in general, the larger the *s* character, the shorter and stronger the bond<sup>49–51</sup>. In other words, the unusual *s*–*p* hybridization is responsible for the larger C–C distances in *fcc*-carbon. Eventually, exactly the excess of *s* character in the hybridization ( $sp^3 \rightarrow sp^2 \rightarrow sp$ ) generates a nonisotropic spatial orientation of the electronic clouds.



**Fig. 8 | Electron localization function (ELF) sections.** **a** ELF corresponding to the (001)-; **b** – to the (010)-; and **c** – to the (100)-planes containing carbon atoms. The blue-green-red gradient represents ELF values in the range [0; 0.3]. The conventional unit cell is indicated by a thin black square, while the position of each carbon atom is at the center of each circular feature of the section.



One should recall here that the onset of a PYS signal is caused by the presence of electrons in occupied orbitals, and its magnitude is proportional to the density of states. In our systems, while the valence band of the diamond system is completely occupied, the valence band of the *fcc*-carbon is partially filled. It is thought that the measured 0 eV energy level (Fig. 5c) corresponds to the calculated energy levels of about 0 eV for diamond and –6 eV for *fcc*-carbon DOS (Fig. 7). Accordingly, it can be assumed that the DOS trends extracted from the PYS data (Fig. 5c) are fairly reproduced by the calculated DOS in the energy ranges [–2.5; 0.0] eV and [–8; –6] eV of the diamond and *fcc*-carbon systems, respectively (Fig. 7c, d). No perfect agreement is to expect here, since the computational model does not take into account the effect of the temperature on the electronic band structure and its occupancy.

To better understand how in *fcc*-carbon the 3D electronic cloud about each atom looks like, the Electron Localization Function (ELF)<sup>52</sup> was calculated. The ELF gives information on possible local accumulation of electrons. On the basis the Electron Localization Function sections shown in Fig. 8 it can be supposed that the distribution of electronic density around carbon atoms is inhomogeneous. Yet the maximum value of the ELF is found to be about 0.3, which indicates a large degree of delocalization of the overall electron density. Such a large rate of delocalization accounts for the electron sharing in the regions of overlapping the electronic clouds, where there is a non-negligible concentration of delocalized electrons (Fig. 6e). The same delocalization allows the accommodation of each carbon atom at the center of a regular polyhedron formed by 12 nearest neighboring carbon atoms, thus forming the Fm  $\bar{3}$  m face-centered cubic crystal lattice with coordination number of 12.

The X-ray diffraction pattern of the *fcc*-carbon was simulated by means of the calculated electronic density. The simulated pattern well reproduces the experimental electron and X-ray diffraction data obtained from the *fcc*-carbon film (see Table 1), which justifies our theoretical model.

## Discussion

### A formation of chemical bonds in *fcc*-carbon can be clarified in light of two problems

Firstly, the coordination number of carbon atoms in *fcc*-structure equal to 12, which cannot be obtained by the four valence electrons of carbon atoms. The peculiar hybridization of the *s* and *p* orbitals in the valence band established by the ab initio calculations generates a spherically symmetric electronic density distribution with a large degree of delocalization; this allows for a symmetric distribution of 12 carbon atoms near a reference one.

Secondly, the shortest interatomic distance in *fcc*-carbon was determined as ~2.5 Å that corresponds to a carbon atomic radius significantly larger than the covalent radius (0.77 Å), but smaller than the van der Waals radius (3.4 Å). Thus, the interaction among carbon atoms in *fcc*-carbon resembles the so-called “van der Waals gap” corresponding to unconventional chemical bonds, like  $\pi$ – $\pi$  stacking<sup>53</sup>.

A hypothesis suggested in ref. 4 explains the nature of chemical bonds in *fcc*-carbon based on their similarity to the well-known noncovalent intermolecular  $\pi$ – $\pi$  interactions, or  $\pi$ -stacking, in organic compounds and carbon-based materials. This kind of noncovalent interaction is typical for organic compounds, in particular, organic semiconductors which usually

contain aromatic rings or another system of conjugated double-bonds (see, e.g., refs. 37,54–57). Apparently, some degree of electron sharing between individual molecules must exist to ensure electrical conductivity in these materials. The parallel  $\pi$ -stacking interactions, also known as “ $\pi$ -stacking pancake bonding”, are likely to contain a component of electron sharing that can be viewed as noncovalent or “covalent-like”<sup>56</sup>. The intermolecular C–C distance in this case can be as short as 2.7 Å<sup>57</sup>, which is close to that in *fcc*-carbon.

In organic semiconductors the configuration of molecules and their mutual arrangement provide a 3D-system of conductive interrelated channels. It is also known that the triplet state of such molecules with nonhybridized *p*-states is often more stable than the singlet one<sup>58,59</sup>. This phenomenon stipulates a long-lived charge transfer event<sup>59</sup>, which is in turn responsible for the conductivity and other features of organic semiconductors<sup>37</sup>. The triplet states are also known to exist in carbon nanotubes and other carbon materials (see, e.g., ref. 58). It can be assumed that carbon atoms in *fcc*-carbon might be trapped in the triplet state that results in a valence band splitting and a charge separation; as a result, noncovalent electron sharing can take place among carbon atoms. Further theoretical investigations and ultrafast transient spectroscopic measurements are needed to confirm this hypothesis.

Intriguingly, other chemical elements of the IV group (Si and Ge) in the periodic table form allotropes with the *fcc*-structure. Indeed, as it was found in ref. 60, germanium crystallizes into a metastable phase with the face-centered cubic crystal lattice having a lattice parameter close to that of the common germanium allotrope with the diamond structure. Silicon was found to exist in the face-centered crystal structure at normal conditions so far only in the form of nanoparticles<sup>61</sup>. As it was established in this work the *fcc*-carbon films remain stable at ambient conditions for months and survives heating up to at least 100 °C in inert atmosphere. The reason of the formation and relative stability of *fcc*-carbon is still unclear and can be addressed to the following studies.

### It is worth giving the outlook for promising applications of *fcc*-carbon

Despite an existing variety of conductive carbon materials, the attractive area of “carbon electronics” remains unavailable. Several reasons can be named: dependency of properties on crystal lattice orientation (graphite), challenges with up-scaling and patterning (graphene), high sensitivity of electronic properties with respect to the substrate and impurities (graphene), instability/irreproducibility of doping and lack of shallow donors/acceptors (diamond). Diamond though has an exceptionally high thermal conductivity and wide bandgap making diamond an ideal material for its employment as wafers for microelectronic devices<sup>62</sup>. Technological progress in the past years made the mass production of diamond supports economically affordable.

Wide bandgap semiconductors are of great interest for high power/high frequency applications implying power electronics<sup>63</sup>. Such materials also open the area of high-temperature electronics which is required in automotive, aerospace, and energy production industries<sup>64</sup>. Transistors based on these materials should basically reveal larger breakdown field and electron saturation velocity<sup>65</sup>. Science and engineering were focused so far

on such materials as SiC and GaN with the band-gap energies ( $E_g$ ) somewhat above 3 eV. Conductors with  $E_g$  beyond these values, namely ultrawide-bandgap (UWBG) semiconductors, extend application fields to RF electronics, deep-UV optoelectronics, quantum devices and others<sup>66</sup>. Such materials (e.g.,  $\beta$ -Ga<sub>2</sub>O<sub>3</sub>, diamond and AlGaN) would allow increasing the Schottky barrier height and/or set up an energy barrier for traversing electrons<sup>65</sup>. There are still serious technological barriers to cope, which are the control of doping and of epitaxial growth itself. Despite a persistent need and efforts invested, *p*-type doping of diamond remains very challenging<sup>67</sup>. Speaking about conventional doping with boron, these acceptors are only weakly activated at room temperature due to the high ionization energy (0.36 eV). Therefore, higher boron concentration is needed that leads to undesirable effects and hinders the use of boron-doped diamond in microelectronics<sup>68</sup>.

In contrast, *fcc*-carbon possesses relatively high conductivity in its undoped state. Beyond that, it has an ultra-wide bandgap. It was proposed in ref. 69 to designate this carbon allotrope as “*metametallic carbon*”, as it possesses the crystal structure, that is typical for metals, but manifests itself as semiconductor with respect to the temperature dependence of electrical conductivity. As it was comprehensively studied and discussed in this work, *fcc*-carbon is presumably characterized by an uncommon type of conductivity which was not reported in the literature so far. Therefore, we propose to classify this material, together with the other ones possessing similar noncovalent sharing of the generally localized valent electrons, into a separate class of *quasivalent solids*. So *fcc*-carbon can be designated as *quasivalent carbon*.

The nature of chemical bonds in this carbon allotrope stipulates negative electron affinity and allows its high polarizability and very low thermal conductivity to be expected. These properties open a novel pathway to “carbon electronics” encompassing such high-end fields as high power/high-frequency electronics, transparent electronics, electron multipliers, field emitters, quantum memory and others<sup>70–74</sup>.

## Methods

### Deposition of carbon films

The diamond single-crystal substrates (plates) employed for the deposition of *fcc*-carbon films were produced with a microwave PACVD reactor operating at a frequency of 2.45 GHz. Such thin diamond plates were made from the initial diamond single crystal; they have the (100) orientation with <100> edges and dimensions of  $3.8 \times 3.8 \times 0.3$  mm with a polished surface subjected to acid cleaning with an RMS roughness <20 nm.

The *fcc*-carbon films were obtained of these substrates with the aid of PACVD by use of the following deposition parameters: temperature—around 600 °C, hydrogen content—98 vol%, methane content—2 vol%, total pressure—29.33 kPa, power—nearly 2.2 kW, deposition time—120 h. The plasma had the shape of ball of roughly 5 cm in diameter. Owing to simultaneous etching of diamond by plasma its surface was relatively rough (Fig. S1) and the film thickness was inhomogeneous. According to numerous SEM studies, the film thickness varies in the range of 100 nm–3 mm being on average ~1  $\mu$ m. The appearance of the films in an optical microscope indicates that the films are transparent.

After the deposition process, the coated samples were subjected to the treatment in diluted HNO<sub>3</sub> for 24 hours to remove hydrogen typically adsorbed on the diamond film surface in the form of dangling bonds leading to a noticeable surface conductivity<sup>75</sup>.

### Examination of the crystal structure

The carbon films on the diamond substrates were examined by different methods.

Obtaining electron diffraction patterns and high-resolution transmission electron microscopy (HRTEM) images were performed by use of a Tecnai TF20 X-Twin microscope (FEI). The FFT and inverse FFT images were obtained based on the HRTEM image by use of the ImageJ software<sup>76,77</sup>.

The atomic force microscopy (AFM) maps were acquired on the C3000 instrument (Nanosurf, Switzerland) using Tap190Al-G probes at

160 kHz frequency. Linear profiles were extracted in the free Gwyddion software. Optical images were obtained with the help of the LEXT OLS4100 confocal laser microscope (Olympus, Japan). The scanning electron microscopy (SEM) images were obtained on the GeminiSEM 500 instrument (Carl Zeiss GmbH, Germany) using 5 kV accelerating voltage, 20 mm aperture and 4 mm working distance.

The X-ray diffraction (XRD) measurements were performed on the D8 Discover system (Bruker, Germany) using Cu K $\alpha$  radiation. From the primary side Göbel mirror, 0.2 mm slit, 2-bounce 220-Ge monochromator and a 0.5 mm collimator were used. From the secondary side a combination of 1 mm, 0.6 mm and 0.2 mm slits was set. A 1D Lynxeye XE-T detector in a 0D mode and in a high-count rate for the alignment and in a high-resolution mode for actual measurement was used. Symmetric scans were conducted with 0.01°/2° steps. Rocking curves and Phi-scan were acquired with 0.005°/2° and 0.05°/0.5° steps respectively. The intensity of each diffraction maximum was optimized based on the following principal alignment steps: (1) setting detector to a maximum intensity of the beam; (2) parallelizing beamline to the sample surface in combination with Z-positioning; (3) fine-tilting around X/Y axes and linear X/Y translations for each series of diffraction maxima. Fitting of diffraction maxima was performed with the aid of a PeakFit program using pseudo-Voigt function and based on the area approximation. To establish the lattice constant, two standards were used: (1) stoichiometric, cubic single crystalline (100)-SrTiO<sub>3</sub> and (2) diamond substrate itself. In the former case the positions of reflexes within relevant 2 Theta-ranges for each tilting angle of interest ( $\Psi = 0^\circ, 45^\circ$  and  $55^\circ$ ) were determined. Comparing these data with the standard (PDF 01-070-8508) from the ICDD database, the instrumental error for each necessary Psi-2 Theta combination was determined. Using those errors, the lattice constant of diamond was calculated from the positions of the (111)-, (220)-, and (400)-maxima determined. A comparison of the obtained values with the standard (PDF 00-006-0675) provided the error of the lattice constant determination, which is given in Table 1.

### Computational models

Since the *fcc*-carbon is not widely accepted as an individual phase, we performed an extensive screening of as many Hamiltonian descriptions as possible, trying to obtain a model, which reproduces most of the experimental data at our disposal. To this aim, we fixed the lattice parameter to the experimental value of nearly 3.6 Å as detailed below, and do not perform any structural relaxation.

The model geometry of *fcc*-carbon consists of only one atom in the primitive unit cell (Supplementary Information, Fig. S7), which we locate at the origin of the cell. The space group is Fm  $\bar{3}$  m. The lattice parameters of the primitive unit cell are  $a = b = c = 2.511$  Å, realizing the lattice angles  $\alpha = \beta = \gamma = 60^\circ$ ; the corresponding conventional unit cell has lattice parameters  $a = b = c = 3.551$  Å and angles  $\alpha = \beta = \gamma = 90^\circ$ .

To model the interatomic interactions, we tested several DFT functionals within the LDA and GGA approximation<sup>46,78–87</sup> (Table 3), also including van der Waals corrections implicitly considered in the meta-GGA description<sup>88,89</sup>. Concerning the calculations with the meta-GGA energy functionals, we were able to achieve the SCF convergence at lattice parameters different than the experimental ones; since we aim to create a model with the atom geometry consistent with our experimental data, we did not consider the meta-GGA functionals in the comparison. We also considered the presence of electron screening effect within the GW description<sup>90</sup>, however, as expected, they were not showing any particular feature in the electronic structure which has not been already reproduced by some of the standard DFT functionals. The DOS and band structures generated with the considered DFT functionals showed similar features. After taking also into account the computational load, we finally selected the revPBE<sup>46</sup> DFT functional with no van der Waals corrections, which reproduces most of our preliminary experimental findings as discussed in the main text of the present manuscript. We perform our simulations by means of the ABINIT package with the PAW formalism<sup>90</sup>. We set the energy cut-off at 2000 eV and the PAW supplementary grid cut-off at 4600 eV, while we sample the

**Table 3 | Calculated bandgap (eV) at selected DFT energy functionals**

Energy Functional	Bandgap (eV)	Reference
CA	7.03	ref. 78
PZ	7.03	ref. 79
HL	7.02	ref. 80
PW	7.03	ref. 81
PBE	7.11	ref. 82
revPBE	7.09	ref. 46
RPBE	7.10	ref. 83
HTCH93	7.35	ref. 84
HTCH120	7.53	ref. 85
C09x	7.17	ref. 86
HTCH407	7.48	ref. 87

Brillouin zone with an 87x87x87 Monkhorst-Pack mesh. The SCF solution of the Schrödinger equation is considered converged when the difference between two consecutive cycles is less than  $3 \times 10^{-10}$  eV. Since the electronic energy levels display a band gap about the Fermi level right from the beginning of the SCF iterations, we did not experience any technical issue for the SCF convergence; for this reason, there has been no need to use any smearing method and the band occupancy is an integer number at any sampled point of the reciprocal space. We include the ABINIT input file “Supplementary Data 1” as supporting information in order to facilitate the reader in reproducing the results.

We calculated the electronic band dispersion of pure diamond for comparison. The space group of the diamond structure is Fd3m. The lattice parameters of the primitive unit cell are  $a = b = c = 2.518$  Å, realizing the lattice angles  $\alpha = \beta = \gamma = 60^\circ$ ; the corresponding conventional unit cell has lattice parameters  $a = b = c = 3.561$  Å and angles  $\alpha = \beta = \gamma = 90^\circ$ . We use the same computational settings as described above.

The atom-projected DOS provides us quantitative information on the  $s$  and  $p$  character of the valence band, that is the relative contribution of the  $s$  and  $p$  orbitals to the total dos. The character of each orbital is obtained by integrating the projected DOS from the bottom of the valence band up to the Fermi level, normalized by the integral of the total DOS. For example, the  $s$  character is obtained as:

$$\int_{-\infty}^{E_F} s(E) d(E) \quad (1)$$

$$\int_{-\infty}^{E_F} g(E) d(E) \quad (2)$$

where  $s(E)$  and  $g(E)$  are the  $s$ -projected and total density of states, respectively.

### Spectroscopic and electrical measurements

Raman spectroscopy was performed with the aid of a Horiba Xplora spectrometer equipped with a 532 nm laser (Horiba Scientific). The measurements were conducted using  $\sim 6$   $\mu$ W power, 1800 mm<sup>-1</sup> grating,  $\times 100$  lenses and 2 min acquisition time. Carbon particles of several microns in size were detached by a tiny diamond blade from the film's surface and placed into the cells of a Cu grid. The operation was performed under a light microscope.

X-ray photoelectron spectroscopy (XPS) analyses were performed using a Kratos XPS spectrometer (Shimadzu) equipped with a monochromated Al K $\alpha$  X-ray source ( $h\nu = 1486.6$  eV). During the measurements, the base pressure inside the XPS chamber was kept constant at around  $6.7 \times 10^{-7}$  Pa. High-resolution XPS spectra were recorded with a 0.1 eV step. The deconvolution of the spectra was conducted by use of the CasaXPS

software ([www.casaxps.com](http://www.casaxps.com)) using the Tougaard background. Charge correction was done based on the center position of the C  $sp^2$  spectral component at 284.5 eV<sup>91</sup>. This contribution was detected on the films' surface after the softest possible cleaning by argon ions. Fitting was performed using Shirley background, Gaussian-Lorentzian function for adventitious carbon peak at about 284.5 eV, and Doniach-Sunijic (DS) asymmetric tail function for the peak at nearly 283.9 eV<sup>92</sup>.

The UV spectrum from the carbon film was recorded using a vacuum ultraviolet spectrometer McPherson VUVAS 1000. The VUV reflectivity measurements were performed in the spectral band of 120–240 nm with 1 nm step; the PMT detector voltage was equal to 480 V. The spectrum was reconstructed according to the Tauc equation, and the curve obtained in such a way was extrapolated from the linear region by use of the Tauc method<sup>93–96</sup> to calculate the value of optical bandgap. Herewith the intersection with linear background was taken as described elsewhere<sup>97</sup>.

The FTIRS measurements were conducted on Vertex 70 (Bruker Optics, Germany) device using rock solid interferometer, mercury (Hg) IR-source, DLATGS IR-Detector, and the KBr Beam-splitter. Beam spot of 3 mm size was placed in the middle of the sample. There was no integration sphere installed. Sample was fixed on undoped (100)-silicon wafer by a carbon double-side tape. The attenuated total reflection measurement did not deliver enough intensity due to the extreme surface roughness. The attributed regions were taken from literature and database<sup>98–100</sup>.

KP and PYS measurements were performed. Prior to the measurements the samples were annealed at 100 °C for 1 h in a glovebox with water and oxygen contamination levels below 1 ppm. The transport of samples was in a sealed plastic bag whereas the installation onto the measurement table took place in a glovebox as well.

Electrical conductivity of carbon film and its temperature dependence were obtained by means of 4-point measurements in Van der Pauw geometry eliminating contact resistance. The film was bound by tiny indium wires from the corners of the sample which was in turn glued with the help of thermal paste onto sapphire holder to isolate sample electrically and provide good thermal conductivity. Electrical resistivity of the film was measured at different temperatures sweeping from 302 °K to 2 °K and back in 31 steps in each direction. The conductivity of the material was calculated implying film's average thickness of 1  $\mu$ m.

The Hall effect was measured in the same configuration. Due to low conductivity, the AC-mode with current of 0.01 mA and frequency of 57.1 Hz was used. The magnetic field was ranged from  $-1$  to  $+1$  T. The  $V_H$  was found to be inconsistent and negligibly low. We suggest the nonplanar geometry of the film (see Fig. S1, Fig. S2, Fig. S3) to be the main reason of a fault.

### Data availability

The experimental data are available and can be shared upon reasonable request. To facilitate the reader in the reproduction of the simulation results, we add the input file “Supplementary Data 1”.

Received: 14 December 2023; Accepted: 18 June 2024;

Published online: 02 July 2024

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## Acknowledgements

R.M. acknowledges the financial support in purchasing the X-ray diffractometer by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation), grant number AOBJ 649257. Prof. E. List-Kratochvil is appreciated for providing access to the various high-end analyses (SEM, AFM, Raman and fluorescence spectroscopy, laser microscopy). A.C. acknowledges the support of the Ministry of Education, Youth and Sports of the Czech Republic through the e-INFRA CZ (ID:90254) and access to the computational infrastructure of the OP VVV-funded project CZ.02.1.01/0.0/0.0/16\_019/0000765 “Research Center for Informatics”; A.C. also acknowledges that this work was co-funded by the European Union under the project “Robotics and advanced industrial production” (reg. no. CZ.02.01.01/00/22\_008/0004590). A.B. acknowledges CzechNanoLab Research Infrastructure supported by MEYS CR (LM2023051). Dr. D. Sidorenko and Dr. T. Sviridova are thanked for the contribution at the early stages of this work. Mr. M. Weise and Dr. S. Seeger are acknowledged for the FTIRS measurements.

## Author contributions

I.K. supervised the PACVD experiments and measurements, and wrote several parts of the manuscript; R.M. performed XRD, SEM, AFM, and Raman measurements and conducted treatments of their and some other results including work on texts; A.C. organized, supervised and performed the majority of the ab initio calculations, and wrote the corresponding part of the manuscript; A.B. conducted Raman, XPS, electron diffraction and HRTEM studies; M.R. performed the KP and PYS measurements and performed their analysis. A.K. performed some of the preliminary screening of the functionals; T.P. provided critical discussions on the theoretical results; D.T. organized and supervised the deposition trials and measurements of electric properties; N.P. performed the PACVD experiments and obtained samples of fcc-carbon needed for different examinations; B.S. and P.-O.C. were responsible for the acquisition of projects.

## Funding

Open Access funding enabled and organized by Projekt DEAL.

## Competing interests

The authors declare no competing interests.

## Additional information

**Supplementary information** The online version contains supplementary material available at <https://doi.org/10.1038/s43246-024-00547-8>.

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**Peer review information** *Communications Materials* thanks Xiaotian Zhang and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Primary Handling Editors: Zakaria Al Balushi and Aldo Isidori. A peer review file is available.

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