

Cite this: *Nanoscale Adv.*, 2024, 6, 5080

Covalent functionalization by using blue light activated radicals: on the reaction mechanisms of arylazo sulfone binding on graphene†

Alessandro Mameli,^{†ab} Alessandro Kovtun,^{†*b} Derek Jones,^b Vasiliki Benekou,^{ac} Vincenzo Palermo,^b Marco Bandini^{†a} and Manuela Melucci^b

Covalent functionalization of graphene presents a pivotal strategy to enhance its surface properties and overcome inherent chemical inertness. While diazonium salts have been extensively utilized for this purpose, their limitations necessitate exploration of alternative approaches. Arylazo sulfones, such as diazonium salt derivatives serving as chromophores, offer a promising solution, enabling photochemical reactions under visible light. Here, we propose a novel method for rapid covalent photofunctionalization of chemical vapor deposition (CVD) graphene on copper substrates using arylazo sulfones. The generation of aryl radicals – chlorobenzene in this case – was achieved through blue light LED irradiation of 4-chlorophenylazo methyl sulfone solution in acetonitrile. Efficient surface covalent modification of graphene was verified by observing (i) the photogeneration of radicals with a decrease in the $\pi-\pi^*$ band absorbance and an increase in the $n-\pi^*$ of arylazosulfone solution by UV-Vis spectroscopy; (ii) an increase in C sp^3 defects on graphene from the Raman D band, the Auger C KLL signal and graphene C 1s X-ray photoelectron spectroscopy (XPS); and (iii) the presence of the chlorobenzene XPS Cl 2p signal. The aryl radical generation was enhanced by the copper substrate during irradiation, with a possible double path reaction mechanism. This approach highlights the versatility of arylazo sulfones in covalently patterning graphene surfaces, thus unlocking opportunities by overcoming the current approach consisting of the deposition of resist materials with successive cycles of lithography and electrochemistry.

Received 30th April 2024
Accepted 31st July 2024

DOI: 10.1039/d4na00359d

rsc.li/nanoscale-advances

Introduction

Due to its exceptional properties, graphene has been extensively investigated over the past two decades. Graphene is a promising candidate for applications in electronics^{1,2} and energy storage,^{3,4} thanks to its electrical conductivity. Moreover, its biocompatibility and large surface area make it appealing for biomedical uses, especially in drug delivery and biosensing.⁵ Graphene is commonly produced on a large scale using chemical vapor deposition (CVD) on copper and nickel substrates, which is advantageous for catalytic applications. Although this method provides control over parameters such as thickness, quality, and

morphology, precise tuning of its properties through surface functionalization is needed to use it in a wider range of applications.

Surface functionalization of graphene and other 2D materials is now quite a well-established technique achieved through various methods, such as chemical activation,⁶ electrochemistry,^{7,8} thermal treatments,⁹ ultrasonication^{10,11} and more recently, visible-light photochemistry.¹² Surface functionalization appears to be an even harder challenge considering the control of surface properties over limited spaces at the micro-scale, *i.e.* for the fabrication of multiarray sensors; only a few examples are present in the literature which involve the use of lithography and diazonium salts.

Diazonium salts are amongst the most versatile and widely used chemical vectors for the covalent functionalization of graphene and various other materials, including carbon compounds, metals, semiconductors, and polymers.¹³ This approach is based on a charge transfer mechanism, leading to the generation of aryl radicals (Ar^\cdot) and their successive covalent bonding to the C sp^2 lattice, followed by a change in graphitic carbon atom hybridization from sp^2 to sp^3 .¹⁴ Among many, the most common approach relies on the use of an electrolytic cell, with graphene acting as a working electrode.¹⁵

^aDipartimento di Chimica "Giacomo Ciamician" Alma Mater Studiorum – Università di Bologna, Via P. Gobetti, 85, 40129 Bologna, Italy

^bIstituto per la Sintesi Organica e la Fotoreattività (ISOF), Consiglio Nazionale delle Ricerche (CNR), Via P. Gobetti, 101, 40129 Bologna, Italy. E-mail: alessandro.kovtun@isof.cnr.it

^cDipartimento di Scienze Fisiche, Informatiche e Matematiche (FIM), Università di Modena e Reggio Emilia (UNIMORE), Via G. Campi, 213/A, 41125 Modena, Italy

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4na00359d>

‡ Current affiliation: School of Science, RMIT University, Melbourne, Victoria 3001, Australia.



In this framework, light-mediated covalent functionalization of 2D materials has received much attention over the past 15 years. One interesting approach is using visible-light laser (514 nm) photolysis of silver trifluoroacetate, generating a radical able to covalently bond to graphene.¹⁶ Although this approach seems to be effective and potentially scalable, metallic silver residues could affect the performance of any final device or sensor. In a previous study, similar irradiation (514.5 nm laser) successfully induced the covalent bonding of aryl groups onto graphene in only a few minutes using benzoyl peroxide in toluene. However, detailed mechanistic investigations revealed the pivotal light absorption by graphene with subsequent charge transfer to benzoyl peroxide.¹⁷ In fact, the direct photolysis of benzoyl was excluded since the compound does not adsorb in the visible region. More recently, the use of multiple wavelengths (532 nm, 450 nm and 610 nm) to produce intermediates for covalent bonding to graphene was also found to be suitable for the “classic” diazonium salts,¹⁸ where light-induced charge transfer (CT) was exploited for patterning on the microscale. However, uncontrolled covalent bonding occurred in non-irradiated areas; control experiments in the absence of light or with lower energy red light (610 nm) showed covalent bonding even without exposure to 532 nm green light or 450 nm blue light. Moreover, graphene uniformly absorbs ~2.3% of light over the whole visible spectrum,¹⁹ and the photo-excited electrons with long wavelengths still have enough energy to overcome the reaction barrier.

In fact, the use of diazonium salts exhibits significant limitations for photo-induced covalent functionalization. As a matter of fact, even when functional groups containing oxygen or nitrogen are attached to the aromatic ring, these salts remain optically transparent to visible light. However, with sulfur-containing functional groups, in this case using 2,5-diethoxy-4-*n*-butylthiobenzenediazonium tetrafluoroborate, the diazonium salt shows a distinct absorption band around 400 nm.²⁰ While this molecular modification offers an optical absorption advantage, it can also pose difficulties when a sulfur group is not desired in conjunction with the aryl cation. To address this issue, a sulfur-substituted diazo moiety $-N_2SO_2CH_3$ has been proposed, as the N-S bond can be cleaved releasing nascent nitrogen. These compounds, known as arylazo sulfones, are chromophores which share a chemical structure similar to that of diazonium salts but exist in a covalent uncharged form. Originally studied in the early 1970s,²¹ their reaction mechanism was elucidated only more recently.^{22,23} Under the influence of heat or irradiation, arylazo sulfones can undergo decomposition or polymerization *via* either ionic or radical mechanisms, with the choice of mechanism favored in protic solvents (water) for the former and aprotic solvents (acetonitrile) for the latter.

These compounds have also found extensive use in metal-free coupling reactions both as radicals and electrophiles.^{24–26} More recently, they have been employed for photo-induced covalent bonding onto gold substrates,²⁷ graphene oxide, and reduced graphene oxide surfaces.¹² Similar photo-reactivity was also observed on aryl diazonium salts and arylazo sulfonates activated using blue light.²⁸ Arylazo sulfones are not unique for photografting on surfaces, and also iodonium salts present

analogous properties.²⁹ Additionally, successful covalent bonding onto graphene was recently achieved through the solvation of diazonium cations with dimethyl sulfoxide (DMSO) anions, which share a molecular structure similar to that of the arylazo sulfones presented here.³⁰

In fact, the present work introduces a quick method for the functionalization of CVD grown graphene on copper and graphite (Highly Oriented Pyrolytic Graphite, HOPG) surfaces with arylazo sulfones. The functionalization was investigated structurally and morphologically, with an emphasis on X-Ray Photoelectron Spectroscopy (XPS)³¹ and Raman spectroscopy.³² The influence of blue light and the presence of copper during the reactions were also investigated.

Experimental

The 4-chlorophenylazo methyl sulfone (IUPAC name 1-(4-chlorophenyl)-2-(methylsulfonyl)diazene) used in the present work was synthesized from diazonium salts using the procedure described by Protti *et al.*²² as described in our previous work.¹² Graphene monolayer films (CVD-G/Cu) on Cu were used as received (Graphenea, Spain). Highly Oriented Pyrolytic Graphite (HOPG) of ZYH grade (Advanced Ceramics, Cleveland, USA) was freshly cleaved before use.

The covalent functionalization of graphene and HOPG was achieved by irradiating the substrates in a 0.15 mM solution of 4-chlorophenylazo methyl sulfone in anhydrous acetonitrile with a commercial LED (462 nm) for 7 minutes under magnetic stirring. The temperature of the bath was kept below 30 °C and the irradiation power density of the photoreactor was 10–13 mW cm⁻² as measured using a photoradiometer.

The control samples were prepared using the same reaction conditions but avoiding any exposure to either the LED or ambient light. From a previous study of diazonium salts on gold³³ it might be expected that some reaction could occur spontaneously.

UV-Vis spectra of pre-irradiation and post-irradiation solutions of arylazo sulfones were recorded using a Cary 100 UV-Vis spectrophotometer (Agilent Technologies). The stability of the 4-chlorophenylazo methyl sulfone stock solution was checked daily and it showed stability for at least 15 days.

Raman spectroscopy

Micro-Raman spectra of substrates were recorded using a LabRAM HR Evolution spectrometer (HORIBA Jobin Yvon Ltd), equipped with a grating (600 grooves per mm), combined with an Olympus BXFM-ILHS microscope. The samples were exposed to a source of 532 nm wavelength (Nd-YAG laser, nominal power 100 mW), with a maximum measured radiation power on the sample of less than 8 mW and a spatial resolution of <2 μm and analyzed with a 100× objective lens, resulting in a 0.5 μm sample diameter exposed to <0.8 mW incident power on the sample. The measurements were performed over a 100–3000 cm⁻¹ spectral range with an overall acquisition time of 1.5 min distributed over five accumulations. The Raman spectrometer was calibrated using a silicon standard. For defect



distance and density estimation on functionalized samples we have followed the criteria described in the literature.³⁴

X-ray photoelectron spectroscopy

XPS spectra were acquired using Al K α radiation ($\hbar\omega = 1486.6$ eV, 125 W) with a Phoibos 100 hemispherical electron energy analyzer (Specs). Spectra were recorded in the constant analyzer energy (CAE) mode with analyzer pass energies of 10 eV for the high-resolution spectra. An overall resolution of 0.9 eV was determined on freshly Ar⁺-sputtered silver measured on the Ag 3d_{5/2} peak. Base pressure in the analysis chamber during measurements was 3×10^{-8} mbar. More details on C 1s peak fitting are given in the ESI.† Auger signals were resampled in 0.2 eV steps and smoothed using ~ 2 eV steps³⁵ before numerical differentiation.

Results and discussion

Photo-induced covalent bonding of chlorobenzene onto graphene was achieved using a commercial LED, with an emission peak at 462 nm, *via* the generation of radicals from 4-chlorophenylazo methyl sulfone (Scheme 1).

Radical generation was monitored using UV-Vis spectroscopy, where the presence of $\pi-\pi^*$ transitions at 305 nm and $n-\pi^*$ at 425 nm is associated with the pristine molecule (Fig. SI 2†). The latter state is populated selectively with visible light exposure (462 nm here) with consequent N-S bond homolysis which, after nitrogen loss from the diazenyl radical $\text{Ar}-\text{N}_2^{\cdot}$, generates an aryl (Ar^{\cdot})/methanesulfonyl ($\text{CH}_3\text{SO}_2^{\cdot}$) radical pair.²³ From the UV-Vis spectra it is possible to observe rapid degradation of the arylazo sulfone molecule over 5–10 minutes of LED irradiation, with a drastic decrease in the $\pi-\pi^*$ band and a flattening of the $n-\pi^*$ band. This implies that radical generation is relatively fast, with the reaction completing in only a few minutes. The kinetics observed by UV-Vis spectroscopy agree with previous results obtained on arylazo sulfonates irradiated at 456 nm,²⁸ where the generation of aryl radicals was confirmed by Electron Paramagnetic Resonance (EPR) and TEMPO radical traps.

The functionalization of CVD-G/Cu was monitored by Raman and XPS. The results from Raman spectroscopy are reported in Fig. 1. The Raman spectra of pristine CVD-G/Cu present the G (~ 1595 cm⁻¹) and 2D (~ 2713 cm⁻¹) bands,

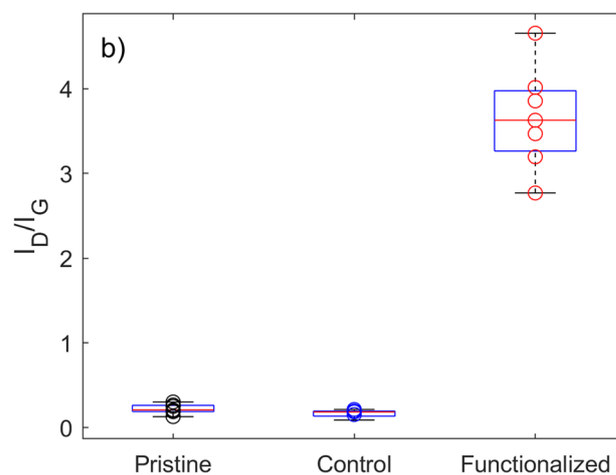
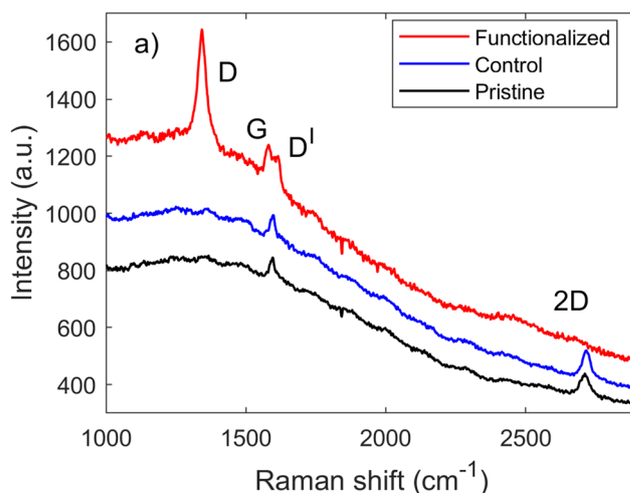
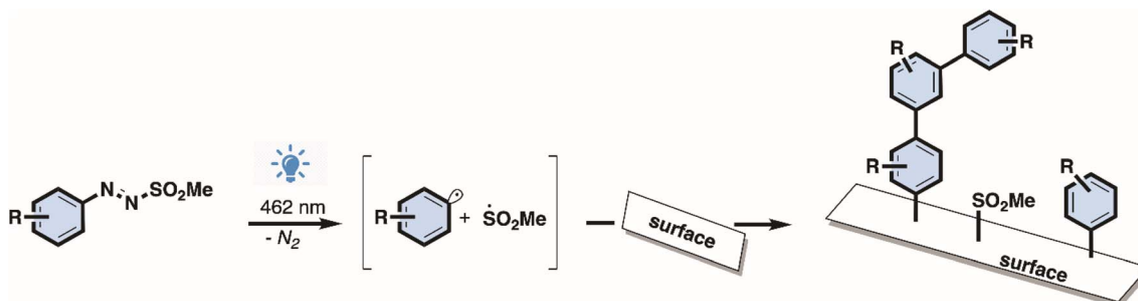


Fig. 1 (a) Representative Raman spectra of pristine, control and functionalized CVD-G/Cu. (b) Box plot of I_D/I_G of pristine, control and functionalized CVD-G/Cu. Functionalized samples were irradiated for 7 minutes in 0.15 mM solution of 4-chlorophenylazo methyl sulfone in acetonitrile.

which depend on the laser frequency³⁶ and correspond to the expected values for a 532 nm laser (2.33 eV). The typical D (~ 1350 cm⁻¹) peak associated with C sp³ defects on graphene gave an I_D/I_G ratio of only 0.2, where I_D and I_G are the intensities of D and G peaks, respectively. The presence of single layer



Scheme 1 Arylazo sulfone surface functionalization mechanism.



Table 1 Values of the main parameters obtained from Raman spectra of pristine, control and functionalized CVD-G/Cu

Parameter	Pristine	Control	Functionalized
D (cm ⁻¹)	1360 ± 4	1367 ± 7	1352 ± 7
G (cm ⁻¹)	1595 ± 8	1600 ± 4	1591 ± 5
2D (cm ⁻¹)	2713 ± 6	2712 ± 10	—
I _D /I _G	0.2 ± 0.1	0.19 ± 0.04	3.7 ± 0.6
FWHM 2D (cm ⁻¹)	38 ± 6	37 ± 5	—
A _{2D} /A _G	2.2 ± 0.4	1.7 ± 0.4	—

graphene was confirmed by observing the symmetry of the 2D peak, an FWHM of 38 cm⁻¹ and the ratio between the areas of 2D and G peaks (A_{2D}/A_G ratio) of ~2.³⁷

The covalent bonding of chlorobenzene onto CVD-G/Cu was confirmed by the appearance of an intense D peak, associated with the formation of C sp³ defects on graphene.¹⁴ The kinetics of the reaction was monitored by observing the evolution of the I_D/I_G peak ratio. As reported in Fig. S11,† the behavior of I_D/I_G as a function of time was in agreement with that expected for increasing defect density (or decreasing distance between defects).³⁴ All the spectroscopic (XPS and Auger) and microscopic (AFM) characterization was performed on the sample with an I_D/I_G peak ratio of 4, which was obtained after 7 minutes of reaction at a 0.15 mM concentration (see Table 1), with a high degree of functionalization being achieved within a relatively short time; longer reaction times lead to a further increase in functionalization, but after 15 minutes we observed a substantial plateau. It was possible to estimate that defect coverage after 7 minutes was of the order of magnitude of 1% and the corresponding distance between defects was close to 2 nm, similar to values found on chemical functionalization of different graphene-based materials.⁶ The control experiment was performed under the same reaction conditions, but without LED irradiation, presenting neither a D signal increase nor any other significant change in the Raman spectra. The initial I_D/I_G value for pristine graphene of ~0.2 was an overestimation, given the fluorescence signal present in the Raman signal. After functionalization, the D' peak appeared at ~1616 cm⁻¹ with the 2D peak almost disappearing, in agreement with what is usually observed for highly functionalized graphene. The Raman spectrum of the dry powder of 4-chlorophenylazo methyl sulfone is reported in the ESI† with a tentative assignment to each vibrational Raman mode with no signal being observed in the D region (Fig. 1).

The overall increase in C sp³ defects was also confirmed using XPS spectroscopy (Fig. 2), from the C 1s signals of graphene,³⁸ as well as the degree of functionalization (Cl 2p signal from bonded chlorobenzene) and the attenuated signals from the Cu substrate (XPS is surface sensitive due to the limited photoelectron escape depth). The presence of C sp³ relative shift was confirmed by C 1s analysis for covalent functionalization on graphene;³⁹ nevertheless, given the narrow chemical shift from the main component (C sp²), the quantification of C sp³ defects from C 1s fit is quite challenging and proper support of Auger signal analysis is necessary.⁴⁰ On fitting the C 1s signal, the

relative amount of C sp² over all carbon atoms decreased from 97% to 46%, while the C KLL Auger signal (Fig. 3) confirmed the overall trend: the first-derivative of the C KLL signal shows a distance (in eV) between the relative minimum and maximum which decreases from 19 eV in pristine CVD-G/Cu to 14 eV in functionalized graphene, which is quite close to the 13.2 eV value reported for C sp³.⁴¹ This parameter is called D (delta), (not to be confused with the D band in Raman), which increases linearly with the amount of C sp² present in the sample, as in the case of carbon nanotubes, amorphous diamonds and graphitic materials,⁴² although the presence of oxygen in C–O bonds may significantly affect the final C sp² estimation using the D parameter, with some deviation from linearity, as observed by Lesiak.⁴¹ Moreover, the different kinetic energies of C KLL and C 1s electrons lead to different escape depths, with the C KLL electrons with ~260 eV kinetic energy being more surface sensitive compared to the ~1200 eV kinetic energy of the C 1s photoelectrons.^{43,44} Thus, both C KLL and C 1s signals confirmed the overall increase in C sp³, with the C KLL signal being much more sensitive to the C sp³ present in the upper layer formed by the grafted molecular layer (comprising aryl and methanesulfonyl groups²⁷) compared to the C sp³ present on graphene itself, whereas the Raman signal derives only from C sp³ present on graphene.

XPS survey spectra (Fig. 2) also show strong Cl 2p_{3/2} signals peaking at 200.5 eV binding energy which arise from the chlorobenzene group (expected at 200.1 eV (ref. 45)) on functionalized samples, which are absent in control samples. The presence of chlorobenzene, combined with the results from the Raman spectroscopy (D band), confirms the effective covalent bonding of chlorobenzene onto graphene.

The XPS signals from the control samples seem to be affected by the reaction environment, contrary to what was observed in Raman data: C sp³ defects increase, as confirmed by both C KLL and C 1s, as well as by the O 1s signal increase (Table 2). The presence of the N 1s signal at ~400 eV with 0.4 at% (Fig. S14†) could be associated with adsorbed arylazo sulfone degraded on the graphene surface during or after the control experiment (no Cl 2p or S 2p XPS signals were present), while it can be excluded in the presence of residues from acetonitrile trapped between graphene and copper (absence of the N 1s signal after 7 minutes of immersion of CVD-G/Cu in acetonitrile, and the spectrum was not reported). On the functionalized graphene samples the presence of significant amounts of nitrogen (2.0 at%) and chlorine (7.0 at%), from N 1s (~400 eV) and Cl 2p signals, confirmed the presence of aryl (chlorobenzene) and nitrogen on the surface. The chemical state of nitrogen remains unclear: the broad signal (FWHM 3 eV) at *c.a.* 400 eV cannot be univocally associated with pyridinic (398.3 ± 0.3 eV), pyrrolic (400.1 ± 0.3 eV) or graphitic (401.5 ± 0.3 eV) nitrogen.⁴⁶ We cannot exclude the covalent grafting of diazenyl radical Ar–N₂ to graphene with the –N=N– group as reported by Nicchio for arylazo sulfonate sodium salts (Ar–N=N–SO₃Na²⁸), but this reaction path would be secondary compared to the main one: given the relative amount of chlorine and nitrogen, we can estimate that only one aryl every seven is bonded to graphene *via* nitrogen (Ar–N=N–graphene), and



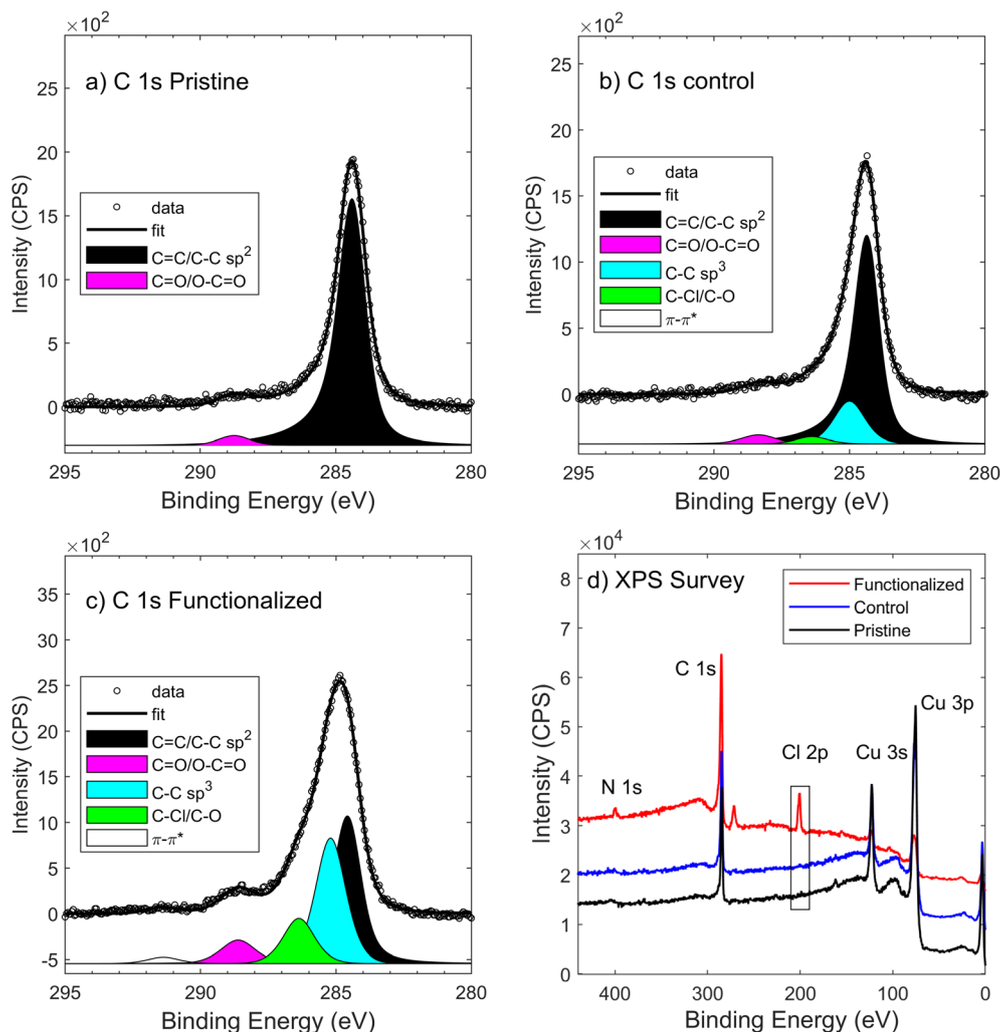


Fig. 2 (a) XPS C 1s of pristine CVD-G/Cu. (b) XPS C 1s of control CVD-G/Cu. (c) XPS C 1s of functionalized CVD-G/Cu. (d) XPS survey spectra of pristine, control and functionalized graphene. The reduced chi square values of C 1s fit were: 4.0 (pristine), 2.9 (control) and 3.8 (functionalized).

the other six are directly bonded to the graphene (Ar-graphene). The sulfur content (S $2p_{3/2}$ \sim 168 eV) on the surface is relatively low, ranging from 0.3 at% to 0.5 at%, indicating the presence of the methanesulfonyl group, due to the functionalization of graphene by the methanesulfonyl ($\text{CH}_3\text{SO}_2^\cdot$) radical. A related experiment, the covalent functionalization of gold substrates²⁷ using arylazo sulfones, showed a comparable level of oxidized sulfur content and the presence of N–Au bonds. These considerations about the presence of heteroatoms (N, Cl and S) in functionalized graphene are in agreement with the main reaction path reported in the literature.²³

XPS survey spectra also provide qualitative information on the increase in thickness of the carbon film (C 1s) on the copper substrate (Cu 2p): in pristine CVD-G/Cu the C/Cu ratio is close to 2, while after functionalization it increases to \sim 12, due to a Cu 2p signal decrease and a C 1s increase. The average thickness of the carbon film – graphene and functionalization – on copper was obtained by using the XPS C/Cu ratio as proposed by Cumpson,⁴⁷ obtaining a value in the range of 3.5 to 4 nm, at

least 3 times larger than the initial thickness (pristine and control samples) of 0.8–1.5 nm. The obtained thickness is of the same order of magnitude as the previously reported result of arylazo sulfone on gold.²⁷ These are average values calculated assuming a homogeneous film and considering only the major atoms present in the film (C) and substrate (Cu), thus excluding O, N and Cl. All these XPS results are compatible with a complete coverage of chlorobenzenes on graphene, where for each C–C sp^3 defect on graphene (about 1% from the Raman results) there is a large number of chlorobenzenes attached in series.

The topography of both the pristine and the functionalized CVD-G/Cu was obtained *via* atomic force microscopy (Fig. 4). Both samples under study showed the typical terrace structures of the metallic copper substrate. In addition, some amorphous and globular regions were present in minor amounts and they can be attributed to Cu_2O , typically present on Cu.⁴⁸ Here, we focus on the morphology of the functionalized CVD-G/Cu which appeared to have dot-like structures uniformly formed across



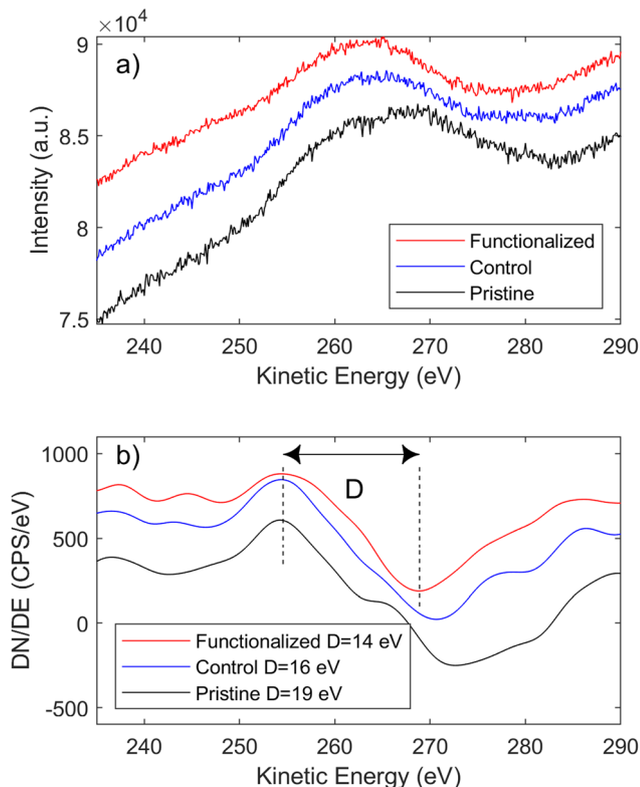
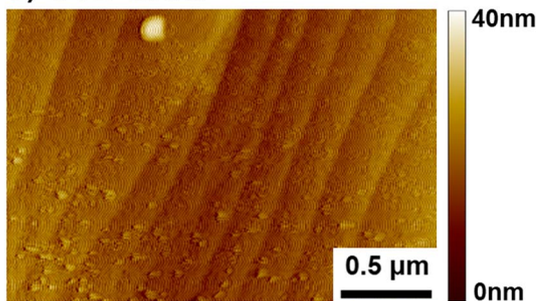


Fig. 3 (a) C KLL Auger signal. (b) First-derivative of C KLL of pristine, control and functionalized graphene on copper.

the surface of graphene, as shown in Fig. 4a. This granular network, as seen in the zoomed-in image in Fig. 4b, can be attributed to the functionalization of graphene as these dot-like structures are present in both metallic and oxidized regions, compatible with topologies observed for covalent functionalization of HOPG using N-heterocyclic carbenes;⁴⁹ A similar AFM topography was found on HOPG functionalized with 4-nitro phenyl diazonium,⁵⁰ where dendritic growth was observed. Unfortunately, from our data it is not possible to distinguish between dendritic and layer-by-layer growth.

a) CVD-G/Cu



b) Functionalized CVD-G/Cu

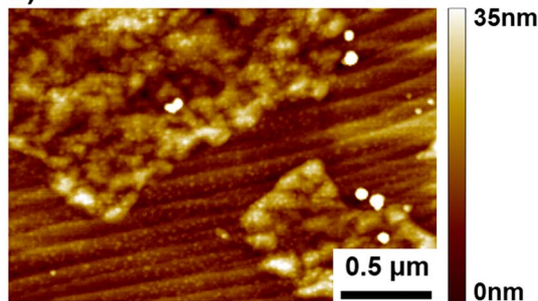


Fig. 4 AFM topography of (a) pristine CVD-G/Cu and (b) functionalized CVD-G/Cu. The functionalized CVD-G/Cu presents both metallic Cu (horizontal lines) and oxidized regions.

The AFM topography of pristine CVD-G/Cu can be seen in Fig. S15a.† As in the case of the functionalized sample under study, we observed the typical lines found when measuring Cu substrates with AFM as well as island-shaped amorphous structures attributed to Cu₂O formed underneath the layer of graphene. Because of the rough Cu surface, imaging with AFM is rather challenging; thus Fig. S15a† appears to show areas that cannot be flattened using the software and appear as black regions. As we zoom in over an area of interest, as in Fig. 4b, we observe the same lines of the substrate but no granular network is observed. This may be a further indication that the dot-like structures found in the functionalized sample under study

Table 2 XPS binding energies (B. E. in eV), the associated chemical states and relative atomic composition (at%) of pristine, control and functionalized CVD-G/Cu

XPS signal	Binding energy (eV)	Chemical state	Pristine (at%)	Control (at%)	Functionalized (at%)
C 1s	284.4	C=C/C-C sp ²	59.3	35.5	33.2
	285.1	C-C sp ³	—	7.3	24.5
	286.3	C-Cl/C-O	—	1.4	9.3
	288.7	C=O/O-C=O	2.0	1.4	5.0
O 1s	532.0	O-C/Cu-OH	3.0	8.5	10.2
	530.6	O-Cu/O=C	1.7	17.2	2.4
Cu 2p _{3/2}	932.8	Cu(0)	32.2	—	—
	932.3	Cu(I)	—	28.3	2.6
	935.0	Cu(II)	—	0	3.3
Cl 2p _{3/2}	200.5	Cl-C	—	—	7.0
N 1s	~400	N-C	—	0.4	2.0
S 2p _{3/2}	161.6	S-Cu	1.8	—	—
	168.6	S-O	—	—	0.5



may indeed be due to the functionalization itself. The chlorobenzene molecules bonded through the functionalization process lead to a non-homogeneous (dot-like) film with an average thickness of several nanometers.

Thermal desorption

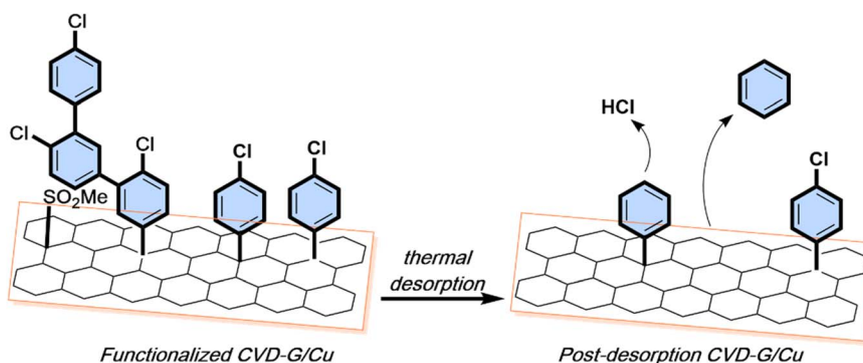
Thermal desorption experiments on post-reaction graphene constitute additional evidence of effective surface functionalization. A linear temperature ramp up to 500 °C allowed measurement of various fragments, including the aryl group ($C_6H_6^+/C_6H_5^+$) with m/z 78 and 77 which appeared above 400 °C and HCl with m/z 38 and 36 (confirmed by the $^{35}Cl/^{37}Cl$ isotopic ratio) above 200 °C. In contrast, no significant signal from the chlorobenzene moiety was detected (Fig. SI 21†). The presence of two complementary fragments from chlorobenzene at two different desorption temperatures could indicate the two-step decomposition of the chlorobenzene covalently bonded onto graphene, where first chlorine (in the form of HCl) was desorbed followed by desorption of the benzene ring. Similar results were reported for covalently bonded aryl groups on graphene by chemical activation of arylazo carboxylic *tert*-butyl esters⁶ and electrochemical functionalization *via* diazonium salts (the most common functionalization method) of the phenylsulfanyl group on CVD-G/Cu.⁵¹ The main m/z species in these papers were the aryl fragments of benzene (77 or 78), and only minor contributions were observed from the whole molecule used to bond onto graphene. After desorption, XPS spectra showed a significant decrease in the Cl 2p signal together with the transformation of C sp^3 into C sp^2 , confirmed through a reduction of the Raman D band. In Scheme 2 the structural properties of functionalized CVD-G/Cu are summarized before and after the thermal treatment.

The role of copper

The surface functionalization process employed in this study exhibited an improved efficiency characterized by rapid kinetics obtained at low concentrations, comparable with that of a previous study reported where visible light was used: 2–5 minutes when benzoyl peroxide is used with a 514.5 nm laser¹⁷ or only a few seconds with the laser-triggered photolysis of silver trifluoroacetate.¹⁶ Furthermore, the kinetics were an order of magnitude faster than those reported in prior investigations

involving graphene, *i.e.* 2 hours for ultrasonication assisted functionalization,¹⁰ 21 h for cycloaddition,⁵² 1 hour *via* a perfluorophenyl azide (PFPA)-mediated coupling⁵³ or 1/2 hours by using defect-activated sites using arylazocarboxylic *tert*-butyl esters.⁶ However, the kinetics also depend on the substrate; the same functionalization – the covalent immobilization of N-heterocyclic carbenes⁴⁹ – can be achieved in 10 minutes on HOPG or in 24 h in the bulk graphene nanoplatelets. The enhanced efficiency presented in this work is attributed to the role of copper, which influences the reaction dynamics. The interaction between copper and graphene improves the reactivity of the surface, changing its properties according to the surface crystal orientation of copper.⁵⁴

It is well known that a monolayer of graphene absorbs 2.3% of visible light.⁵⁵ This light absorption property stimulates a charge transfer process from the graphene to the arylazo molecule, with formation of a radical and consequent functionalization, in agreement with previous studies on the photochemical reactivity of graphene.¹⁷ The selection of the wavelength is of great importance in the electron transfer process for both the molecule and graphene, as shorter wavelengths can facilitate electron extraction from the surface. This wavelength-dependent behavior can be rationalized by considering the work function of graphene, which is 4.30 eV for pristine graphene and can be altered by defects, doping, or the substrate used.⁵⁶ The presence of copper beneath the graphene layer has been demonstrated to shift the work function to lower values (3.97–3.81 eV⁵⁴), thereby enhancing the reaction kinetics. Another significant aspect is the presence of oxidized copper regions in pristine CVD-G/Cu. These visible Cu_2O areas can be observed with an optical microscope (see Fig. SI 8 in the ESI†) and present a darker color than pristine metallic copper. For this reason, it is commonly referred to as a “dark” region (Cu_2O) in contrast with “bright” regions (metallic Cu). Oxygen can reach copper through the graphene grain boundary and the presence of Cu_2O has been confirmed by the presence of Raman signals at 149 and 218 cm^{-1} using a 514 nm laser⁴⁸ or 145 and 215 cm^{-1} with a 633 nm laser.⁵⁷ Our Raman spectra (Fig. SI 8†) confirmed the presence of Cu_2O showing 148 and 218 cm^{-1} signals only in “dark” regions with a 532 nm laser and 142 and 209 cm^{-1} signals with a 473 nm laser. The relative Raman intensity of these Cu_2O peaks is dramatically affected by the



Scheme 2 Functionalized CVD-G/Cu surface structure before and after thermal desorption in UHV at 500 °C.



laser wavelength (see Fig. SI 9†). The diffusion of oxygen between the graphene layer and copper forms a large Cu₂O region, where the graphene is partially detached from the substrate as previously observed by De Luca.⁵⁷ The downward shift of the Raman G and 2D bands to ~1570 and ~2650 (Table SI 4†), respectively, observed in these “dark” regions, is compatible with the “freestanding-like graphene”. The copper of pristine CVD-G/Cu is in a metallic “bright” state, but air and humidity exposure lead to the formation of these so-called “dark” (Cu₂O) regions.

The presence of both Cu states on a macroscopic scale (mm) influences the overall reactivity. Most of the pristine sample was metallic, as assessed by large area (3 × 7 mm²) XPS measurements (Cu 2p_{3/2} B. E. at 932.8 eV and Cu LMM at a K. E. of 918.5 eV) previously discussed. Fig. 1 shows the evolution of Raman spectra on the predominantly metallic “bright” regions, but Raman measurements were also made on the “dark” regions of the sample (see Fig. SI 10 and Table SI 4†), where the same evolution of the Raman signal was observed with an I_D/I_G peak ratio of ~1.0, indicative of successful covalent bonding. Contrary to “bright” regions, the Raman data for “dark” regions of the control samples also presented a significant increase in the D peak compared to pristine graphene (I_D/I_G ~0.2). When the reaction time was extended to 120 minutes, a minor peak (I_D/I_G ~1.0) became discernible even in the bright areas. It is crucial to note that accidental irradiation could occur during preparation of the experiments with exposure to ambient natural light. However, this side reaction is not relevant, because it takes place over an order of hours, while the main functionalization reaction is quite fast: after 30 s I_D/I_G values reach 0.6 over the bright regions (see Fig. SI 11†).

These findings indicate that metallic copper does not actively participate in catalysis without light activation. Conversely, cuprous copper(I) oxide demonstrates the capacity to engage in both reactions, albeit at a slower rate in the absence of light. Thus, we identify Cu(I) as the active species responsible for these reactions. Although aqueous Cu(I) tends to rapidly disproportionate into Cu(II) and Cu metal if not stabilized using complexing agents, acetonitrile solution of Cu(I) species did not undergo any modification of the chemical state due to specific interaction between the metal ion and acetonitrile⁵⁸ thus allowing rapid electron exchange with any copper(II) present. In fact, electron transfer in Cu(II/I) systems is aided by the reorganizational energies involved in changing from a tetrahedral d⁹ Cu(I) four-coordinate system to a six-coordinate d¹⁰ Cu(II) one.⁵⁹

Copper chemical states were further investigated using XPS after the reaction to observe the evolution of the Cu oxidation (see Fig. in SI 13†). XPS peaks for Cu 2p, the Auger Cu LMM and O 1s confirmed the presence of partial oxidation of copper from pristine (mainly metallic Cu and Cu(I)) to Cu(II) after the reaction. Despite this behavior suggesting a Single Electron Transfer (SET) mechanism, it remains unclear whether copper exclusively interacts with the graphene substrate or whether it also directly involves the arylazo sulphone enabling dinitrogen elimination as reported in the literature.⁶⁰ Since the surface exposed to the solution and light is composed of both CVD-G/Cu and pristine copper (localized at the edges of the sample),

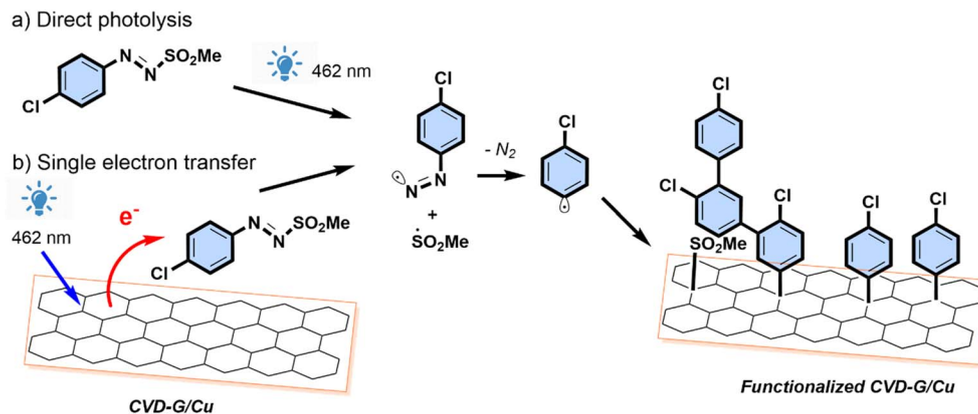
we are not able to discriminate between these two surfaces. Anyway, the observed oxidation of Cu prompts consideration of the potential release of copper atoms into the solution post-oxidation. Direct observation of copper release was achieved by subjecting a small (10 × 10 mm²) silicon substrate with 30 nm of a thermally evaporated copper thin film which was subjected to the same reaction conditions as the CVD-G/Cu samples. The result was the complete dissolution of the thin film in minutes, as shown photographically and from the Raman spectra (see Fig. SI 12†).

Summarizing, copper was a necessary, but not a sufficient element for the covalent bonding reaction. In order to prove this assumption, we performed the same functionalization reaction on a copper-free sample, using the surface of HOPG which is 100% C sp², and compared the results with those of HOPG immersed together with a copper wire. Both experiments were performed in acetonitrile 0.15 mM for several minutes of LED irradiation. The functionalization was extremely effective (I_D/I_G ~0.06) and homogeneous over the whole surface only in the presence of the copper wire, while no effective functionalization under copper-free conditions was observed (I_D/I_G << 0.01). The intensity of the observed I_D/I_G agreed with that in previous work on HOPG functionalization with diazonium salts^{61,62} and with that in our previous work on HOPG with arylazo sulfones, but at a high concentration (50 mM) and long LED exposure times (24 h).¹²

The proposed mechanism is illustrated in Scheme 3, wherein incident light simultaneously initiates the activation of the graphene surface and the cleavage of arylazo sulfones. This dual process results in the elimination of molecular nitrogen and the generation of aryl and CH₃SO₂ radicals which subsequently bond to the graphene surface, leading to surface functionalization. An alternative pathway for radical generation involves indirect activation attributed to charge transfer from copper. A similar mechanism is given by the single-electron transfer living radical polymerization (SET-LRP), where a radical anion initiator [Pn/P-X]⁻ interacts with Cu(0) and other copper electron donors to generate Pn[•] and X⁻ as Cu₂O is formed. Cu(0) undergoes conversion *via* single-electron transfer to Cu(I), which then undergoes disproportionation into Cu(0) and Cu(II), establishing a cyclic process that is deactivated by the generation of Cu(II).⁶³ The reaction is strongly affected by the solvent and it has a reduced rate in non-disproportionating solvents such as acetonitrile.⁶⁴

The control of covalent functionalization using visible light offers a novel method for the fabrication of patterned arrays and could overcome the current approach consisting of the deposition of resist materials with successive cycles of lithography and electrochemistry,⁶⁵ or be an alternative to using oxygen-sensitive photosensitizers such as iodonium salts.²⁹ Although direct patterning has been successfully employed for quantum dots,⁶⁶ produced through covalent reactions with a chosen ligand, no successful attempts have been reported for visible-light, specific photo-induced covalent bonding of generic organic molecules on surfaces such as silicon, gold or graphene. One of the versatile aspects of such wavelength-specific photo-induced reactions on surfaces lies in the possibility of





Scheme 3 Photoreaction mechanisms of arylazo sulfones on CVD graphene on Cu.

generating diverse reactive species even from the same starting molecule based on the wavelength of the light sources used.

Conclusions

We have presented a novel rapid photo-functionalization method for CVD graphene on copper using arylazo sulfones. The functionalization was proven by UV-Vis, Raman and XPS analyses showing the formation of covalent bonding on graphene. Furthermore, thermal desorption was carried out and the mass spectra of fragmentation during the heating process confirmed the covalent functionalization of the graphene surface. The efficiency of the method was primarily attributed to copper's role in enhancing the process, with the metal present as metallic copper (bright region) and cuprous oxide (dark region). Copper was able to activate the reaction for radical attack on the graphene surface, while insights into the molecular photo-dissociation provided help in understanding the overall mechanism.

The resemblance between this mechanistic pathway and the observed experimental findings presented in this work suggests the possibility for a photocatalytic enhancement of SET-LRP, using visible light and azosulfones. The approach could be reiterated by using different building blocks ultimately allowing fine tuning of graphene surfaces by molecular engineering with arylazo sulfones for micropatterning applications.

Data availability

The data supporting this article have been included as part of the ESI.† Atomic composition of samples obtained by X-ray photoelectron spectroscopy and all the main parameters of Raman spectra analysis are reported in the ESI.†

Author contributions

A. M. performed the functionalization and the desorption experiments and acquired Raman, UV-Vis and XPS spectra. A. M. and A. K. analyzed the Raman, XPS and Auger signals. V. B. acquired and analyzed the AFM topography. A. K. conceptualized and supervised the research. A. M., A. K., D. J., M.

B., M. M. and V. P. discussed the reaction mechanism and conditions. The manuscript was written with contributions from all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors are grateful to Graphenea (Spain), in particular to Dr Amaia Zurutuza, who kindly provided the CVD-G/Cu used in this work. A. K. and A. M. are grateful to Emanuele Treossi for helpful discussions. A. K. is grateful to Caterina Baldassini, Viola Ferretti and Leonardo Razzai, MSc Physics students at the University of Bologna for help with the control experiment on pristine CVD-G/Cu and scientific discussions on the related results. Raman measurements were performed at the interdepartmental instrument facility at the University of Modena and Reggio Emilia (Centro Interdipartimentale Grandi Strumenti, UNIMORE), with the financial support of Prof. Patrizia Fava of the Life Sciences Dept. (Dipartimento Scienze della Vita, UNIMORE) and with the technical support of Dr Fabio Bergamini. The authors are grateful to Cristian Bettini and Laura Favaretto for all their suggestions and active support. The authors are also grateful to Matteo Di Giosia and Matteo Calvaresi from the University of Bologna for the irradiation power density data. A. K. is particularly grateful to Dr Sebastiano Mantovani for the long and fruitful discussions on the chemistry of radicals and on the interpretation of NMR and EPR signals. This research was financially supported by the Italian Department for University, project PRIN 2017 NANO-CARBOCAT, grant number 2017W8KNZW. V. B. was financially supported by the H2020-MSCA-ITN ULTIMATE (grant number 813036).

Notes and references

- 1 J. W. Shin, J. H. Han, H. Cho, J. Moon, B. H. Kwon, S. Cho, T. Yoon, T. S. Kim, M. Suemitsu, J. I. Lee and N. S. Cho,



- Display process compatible accurate graphene patterning for OLED applications, *2D Mater.*, 2018, **5**, 014003.
- 2 T. Mahmoudi, Y. Wang and Y. B. Hahn, Graphene and its derivatives for solar cells application, *Nano Energy*, 2018, **47**, 51–65.
 - 3 H. Gwon, H. S. Kim, K. U. Lee, D. H. Seo, Y. C. Park, Y. S. Lee, B. T. Ahn and K. Kang, Flexible energy storage devices based on graphene paper, *Energy Environ. Sci.*, 2011, **4**, 1277–1283.
 - 4 H. Kim, K. Y. Park, J. Hong and K. Kang, All-graphene-battery: Bridging the gap between supercapacitors and lithium ion batteries, *Sci. Rep.*, 2014, **4**, 1–8.
 - 5 C. S. Park, H. Yoon and O. S. Kwon, Graphene-based nanoelectronic biosensors, *J. Ind. Eng. Chem.*, 2016, **38**, 13–22.
 - 6 C. E. Halbig, R. Lasch, J. Krüll, A. S. Pirzer, Z. Wang, J. N. Kirchhof, K. I. Bolotin, M. R. Heinrich and S. Eigler, Selective Functionalization of Graphene at Defect-Activated Sites by Arylazocarboxylic tert-Butyl Esters, *Angew. Chem., Int. Ed.*, 2019, **58**(11), 3599–3603.
 - 7 H. Chen, C. Li and L. Qu, Solution electrochemical approach to functionalized graphene: History, progress and challenges, *Carbon*, 2018, **140**, 41–56.
 - 8 Z. Xia, F. Leonardi, M. Gobbi, Y. Liu, V. Bellani, A. Liscio, A. Kovtun, R. Li, X. Feng, E. Orgiu, P. Samori, E. Treossi and V. Palermo, Electrochemical Functionalization of Graphene at the Nanoscale with Self-Assembling Diazonium Salts, *ACS Nano*, 2016, **10**, 7125–7134.
 - 9 J. E. Johnson, Process for the Surface Modification of a Polymeric Substrate, *US Pat.*, 6555175, 2003.
 - 10 B. Shen, W. Zhai, D. Lu, J. Wang and W. Zheng, Ultrasonication-assisted direct functionalization of graphene with macromolecules, *RSC Adv.*, 2012, **2**, 4713–4719.
 - 11 K. Muthosamy and S. Manickam, State of the art and recent advances in the ultrasound-assisted synthesis, exfoliation and functionalization of graphene derivatives, *Ultrason. Sonochem.*, 2017, **39**, 478–493.
 - 12 L. Lombardi, A. Kovtun, S. Mantovani, G. Bertuzzi, L. Favaretto, C. Bettini, V. Palermo, M. Melucci and M. Bandini, Visible-Light Assisted Covalent Surface Functionalization of Reduced Graphene Oxide Nanosheets with Arylazo Sulfones, *Chem.—Eur. J.*, 2022, **28**, e202200333.
 - 13 C. Combellas, D. E. Jiang, F. Kanoufi, J. Pinson and F. I. Podvorica, Steric effects in the reaction of aryl radicals on surfaces, *Langmuir*, 2009, **25**, 286–293.
 - 14 S. Niyogi, E. Bekyarova, M. E. Itkis, H. Zhang, K. Shepperd, J. Hicks, M. Sprinkle, C. Berger, C. N. Lau, W. A. Deheer, E. H. Conrad and R. C. Haddon, Spectroscopy of covalently functionalized graphene, *Nano Lett.*, 2010, **10**, 4061–4066.
 - 15 L. Gan, D. Zhang and X. Guo, Electrochemistry: An efficient way to chemically modify individual monolayers of graphene, *Small*, 2012, **8**, 1326–1330.
 - 16 T. Wei, S. Al-Fogra, F. Hauke and A. Hirsch, Direct Laser Writing on Graphene with Unprecedented Efficiency of Covalent Two-Dimensional Functionalization, *J. Am. Chem. Soc.*, 2020, **142**, 21926–21931.
 - 17 H. Liu, S. Ryu, Z. Chen, M. L. Steigerwald, C. Nuckolls and L. E. Brus, Photochemical reactivity of graphene, *J. Am. Chem. Soc.*, 2009, **131**, 17099–17101.
 - 18 Y. Li, W. Li, M. Wojcik, B. Wang, L. C. Lin, M. B. Raschke and K. Xu, Light-Assisted Diazonium Functionalization of Graphene and Spatial Heterogeneities in Reactivity, *J. Phys. Chem. Lett.*, 2019, **10**, 4788–4793.
 - 19 R. R. Nair, P. Blake, A. N. Grigorenko, K. S. Novoselov, T. J. Booth, T. Stauber, N. M. R. Peres and A. K. Geim, Fine structure constant defines visual transparency of graphene, *Science*, 2008, **320**, 1308.
 - 20 H. B. Ambroz, T. J. Kemp and G. K. Przybytniak, Optical spectroscopy of the aryl cation 3. Substituent effects on the production and electronic spectra of intermediates in the photodecomposition of ArN₂⁺; optical characterization of the reaction Ar⁺ + N₂ → ArN₂⁺, *J. Photochem. Photobiol., A*, 1992, **68**, 85–95.
 - 21 M. Kobayashi, S. Fujii and H. Minato, Photolysis of Phenylazo p-Tolyl Sulfones, *Bull. Chem. Soc. Jpn.*, 1972, **45**, 2039–2042.
 - 22 S. Crespi, S. Protti and M. Fagnoni, Wavelength Selective Generation of Aryl Radicals and Aryl Cations for Metal-Free Photoarylations, *J. Org. Chem.*, 2016, **81**, 9612–9619.
 - 23 D. Qiu, C. Lian, J. Mao, M. Fagnoni and S. Protti, Dydauxiliary Groups, an Emerging Approach in Organic Chemistry. The Case of Arylazo Sulfones, *J. Org. Chem.*, 2020, **85**, 12813–12822.
 - 24 S. Protti, D. Ravelli and M. Fagnoni, Wavelength dependence and wavelength selectivity in photochemical reactions, *Photochem. Photobiol. Sci.*, 2019, **18**, 2094–2101.
 - 25 Y. Zhao, S. Li, Y. Fan, C. Chen, X. Dong, R. Wang and Y. Y. Jiang, Mild diazenylation of Csp²-H and Csp³-H bonds via arylazo sulfones, *Org. Chem. Front.*, 2023, **10**, 5923–5932.
 - 26 A. Li, Y. Li, J. Liu, J. Chen, K. Lu, D. Qiu, M. Fagnoni, S. Protti and X. Zhao, Metal-Free Trifluoromethylthiolation of Arylazo Sulfones, *J. Org. Chem.*, 2021, **86**, 1292–1299.
 - 27 J. Médard, P. Decorse, C. Mangeney, J. Pinson, M. Fagnoni and S. Protti, Simultaneous Photografting of Two Organic Groups on a Gold Surface by using Arylazo Sulfones as Single Precursors, *Langmuir*, 2020, **36**, 2786–2793.
 - 28 L. Nicchio, J. Médard, P. Decorse, S. Gam-Derouich, A. Chevillot-Biraud, Y. Luo, C. Mangeney, A. Berisha, F. Averseng, M. Fagnoni, S. Protti and J. Pinson, Selective Nsp²- and Csp²- Photografting of Au-Surface by Aryldiazonium Salts and Arylazo Sulfonates, *Chem.—Eur. J.*, 2023, **29**, 1–11.
 - 29 J. Médard, C. Combellas, F. Kanoufi, J. Pinson, J. Chauvin and A. Deronzier, Patterning Surfaces through Photografting of Iodonium Salts, *J. Phys. Chem. C*, 2018, **122**, 19722–19730.
 - 30 Y. Xia, L. Sun, S. Eyley, B. Daelemans, W. Thielemans, J. Seibel and S. De Feyter, Grafting Ink for Direct Writing: Solvation Activated Covalent Functionalization of Graphene, *Advanced Science*, 2022, **9**, 1–11.
 - 31 T. Susi, Other spectroscopic methods for graphene characterization: X-ray and electron spectroscopies.



- Graphene: Properties, Preparation, Characterization and Applications*, 2nd edn, 2021, pp. 413–436.
- 32 A. J. Pollard, K. R. Paton, C. A. Clifford and E. Legge, Characterisation of the Structure of Graphene, in *Good Practice Guide*, National Physical Laboratory, UK, 2017, vol. 145.
- 33 A. Mesnage, X. Lefèvre, P. Jégou, G. Deniau and S. Palacin, Spontaneous grafting of diazonium salts: Chemical mechanism on metallic surfaces, *Langmuir*, 2012, **28**, 11767–11778.
- 34 L. G. Cançado, A. Jorio, E. H. M. Ferreira, F. Stavale, C. A. Achete, R. B. Capaz, M. V. O. Moutinho, A. Lombardo, T. S. Kulmala and A. C. Ferrari, Quantifying defects in graphene via Raman spectroscopy at different excitation energies, *Nano Lett.*, 2011, **11**, 3190–3196.
- 35 S. Kaciulis, A. Mezzi, P. Calvani and D. M. Trucchi, Electron spectroscopy of the main allotropes of carbon, *Surf. Interface Anal.*, 2014, **46**, 966–969.
- 36 M. S. Dresselhaus, A. Jorio, A. G. Souza Filho and R. Saito, Defect characterization in graphene and carbon nanotubes using Raman spectroscopy, *Philos. Trans. R. Soc., A*, 2010, **368**, 5355–5377.
- 37 P. Turner, K. R. Paton, E. J. Legge, A. De Luna Bugallo, A. K. S. Rocha-Robledo, A. A. Zahab, A. Centeno, A. Sacco, A. Pesquera, A. Zurutuza, A. M. Rossi, D. N. H. Tran, D. L. Silva, D. Losic, F. Farivar, H. Kerdoncuff, H. Kwon, J. Pirart, J. L. E. Campos, K. M. Subhedar, L. L. Tay, L. Ren, L. G. Cançado, M. Paillet, P. Finnie, P. L. Yap, R. Arenal, S. R. Dhakate, S. Wood, S. Jiménez-Sandoval, T. Batten, V. Nagyte, Y. Yao, A. R. Hight Walker, E. H. Martins Ferreira, C. Casiraghi and A. J. Pollard, International interlaboratory comparison of Raman spectroscopic analysis of CVD-grown graphene, *2D Mater.*, 2022, **9**, 035010.
- 38 A. Kovtun, D. Jones, S. Dell'Elce, E. Treossi, A. Liscio and V. Palermo, Accurate chemical analysis of oxygenated graphene-based materials using X-ray photoelectron spectroscopy, *Carbon*, 2019, **143**, 268–275.
- 39 S. Freddi, M. C. Rodriguez Gonzalez, A. Casotto, L. Sangaletti and S. De Feyter, Machine-Learning-Aided NO₂ Discrimination with an Array of Graphene Chemiresistors Covalently Functionalized by Diazonium Chemistry, *Chem.—Eur. J.*, 2023, **29**, e202302154.
- 40 A. J. Barlow, S. Popescu, K. Artyushkova, O. Scott, N. Sano, J. Hedley and P. J. Cumpson, Chemically specific identification of carbon in XPS imaging using Multivariate Auger Feature Imaging (MAFI), *Carbon*, 2016, **107**, 190–197.
- 41 B. Lesiak, L. Kövér, J. Tóth, J. Zemek, P. Jiricek, A. Kromka and N. Rangam, C sp²/sp³ hybridisations in carbon nanomaterials – XPS and (X)AES study, *Appl. Surf. Sci.*, 2018, **452**, 223–231.
- 42 A. Mezzi and S. Kaciulis, Surface investigation of carbon films: From diamond to graphite, *Surf. Interface Anal.*, 2010, **42**, 1082–1084.
- 43 B. Lesiak, G. Trykowski, J. Tóth, S. Biniak, L. Kövér, N. Rangam, L. Stobinski and A. Malolepszy, Chemical and structural properties of reduced graphene oxide—dependence on the reducing agent, *J. Mater. Sci.*, 2021, **56**, 3738–3754.
- 44 C. R. Brundle, Application of Electron Spectroscopy To Surface Studies, *J. Vac. Sci. Technol.*, 1974, **11**, 212–224.
- 45 D. T. Clark, D. Kilcast, D. B. Adams and W. K. R. Musgrave, An ESCA study of the molecular core binding energies of the chlorobenzenes, *J. Electron Spectrosc. Relat. Phenom.*, 1975, **6**, 117–134.
- 46 T. Susi, T. Pichler and P. Ayala, X-ray photoelectron spectroscopy of graphitic carbon nanomaterials doped with heteroatoms, *Beilstein J. Nanotechnol.*, 2015, **6**, 177–192.
- 47 P. J. Cumpson and P. C. Zalm, Thickogram: A method for easy film thickness measurement in XPS, *Surf. Interface Anal.*, 2000, **29**, 403–406.
- 48 P. V. Antonov, P. Restuccia, M. C. Righi and J. W. M. Frenken, Attractive curves: the role of deformations in adhesion and friction on graphene, *Nanoscale Adv.*, 2022, 4175–4184.
- 49 B. Daelemans, S. Bernaerts, S. Eyley, W. Thielemans, W. Dehaen and S. De Feyter, Covalent immobilization of N-heterocyclic carbenes on pristine carbon substrates: from nanoscale characterization to bulk catalysis, *Chem. Commun.*, 2024, **60**, 1432–1435.
- 50 T. M. T. Huynh, K. Tahara, S. De Feyter and T. H. Phan, On the role of functional groups in the formation of diazonium based covalent attachments: dendritic vs. layer-by-layer growth, *RSC Adv.*, 2023, **13**, 24576–24582.
- 51 P. Kovaříček, V. Vrkošlav, J. Plšek, Z. Bastl, M. Fridrichová, K. Drogowska and M. Kalbáč, Extended characterization methods for covalent functionalization of graphene on copper, *Carbon*, 2017, **118**, 200–207.
- 52 L. Daukiya, C. Mattioli, D. Aubel, S. Hajjar-Garreau, F. Vonau, E. Denys, G. Reiter, J. Fransson, E. Perrin, M. L. Bocquet, C. Bena, A. Gourdon and L. Simon, Covalent Functionalization by Cycloaddition Reactions of Pristine Defect-Free Graphene, *ACS Nano*, 2017, **11**, 627–634.
- 53 J. Park, X. Yang, D. Wickramasinghe, M. Sundhoro, N. Orbey, K. F. Chow and M. Yan, Functionalization of pristine graphene for the synthesis of covalent graphene-polyaniline nanocomposite, *RSC Adv.*, 2020, **10**, 26486–26493.
- 54 M. Kim, others, Direct Electrochemical Functionalization of Graphene Grown on Cu Including the Reaction Rate Dependence on the Cu Facet Type, *ACS Nano*, 2023, **17**, 18914–18923.
- 55 K. F. Mak, M. Y. Sfeir, Y. Wu, C. H. Lui, J. A. Misewich and T. F. Heinz, Measurement of the optical conductivity of graphene, *Phys. Rev. Lett.*, 2008, **101**, 2–5.
- 56 N. Dimov, A. Staykov, M. I. M. Kusdhany and S. M. Lyth, Tailoring the work function of graphene via defects, nitrogen-doping and hydrogenation: A first principles study, *Nanotechnology*, 2023, **34**, 415001.
- 57 O. De Luca, R. Grillo, M. Castriota, A. Policicchio, M. P. De Santo, G. Desiderio, A. Fasanella, R. G. Agostino, E. Cazzanelli, M. Giarola and G. Mariotto, Different spectroscopic behavior of coupled and freestanding



- monolayer graphene deposited by CVD on Cu foil, *Appl. Surf. Sci.*, 2018, **458**, 580–585.
- 58 S. E. Manahan, Electron exchange between copper(I) and copper(II) in acetonitrile, *Can. J. Chem.*, 1967, **45**, 2451–2452.
- 59 D. B. Rorabacher, Electron Transfer by Copper Centers, *Chem. Rev.*, 2004, **104**, 651–697.
- 60 A. Roglans, A. Pla-Quintana and M. Moreno-Mañas, Diazonium salts as substrates in palladium-catalyzed cross-coupling reactions, *Chem. Rev.*, 2006, **106**, 4622–4643.
- 61 M. C. R. González, A. Brown, S. Eyley, W. Thielemans, K. S. Mali and S. D. Feyter, Self-limiting covalent modification of carbon surfaces: Diazonium chemistry with a twist, *Nanoscale*, 2020, **12**, 18782–18789.
- 62 K. Tahara, T. Ishikawa, B. E. Hirsch, Y. Kubo, A. Brown, S. Eyley, L. Daukiya, W. Thielemans, Z. Li, P. Walke, S. Hirose, S. Hashimoto, S. De Feyter and Y. Tobe, Self-Assembled Monolayers as Templates for Linearly Nanopatterned Covalent Chemical Functionalization of Graphite and Graphene Surfaces, *ACS Nano*, 2018, **12**, 11520–11528.
- 63 G. Lligadas, S. Grama and V. Percec, Single-Electron Transfer Living Radical Polymerization Platform to Practice, Develop, and Invent, *Biomacromolecules*, 2017, **18**, 2981–3008.
- 64 G. Lligadas, B. M. Rosen, C. A. Bell, M. J. Monteiro and V. Percec, Effect of Cu(0) particle size on the kinetics of SET-LRP in DMSO and Cu-mediated radical polymerization in MeCN at 25 °C, *Macromolecules*, 2008, **41**, 8365–8371.
- 65 M. C. R. González, others, Multicomponent Covalent Chemical Patterning of Graphene, *ACS Nano*, 2021, **15**, 10618–10627.
- 66 D. Hahm, others, Direct patterning of colloidal quantum dots with adaptable dual-ligand surface, *Nat. Nanotechnol.*, 2022, **17**, 952–958.

