

Cite this: *Chem. Sci.*, 2022, 13, 9927

All publication charges for this article have been paid for by the Royal Society of Chemistry

# Radical defects modulate the photocatalytic response in 2D-graphitic carbon nitride†

Edoardo Raciti,<sup>ab</sup> Sai Manoj Gali,<sup>a</sup> Michele Melchionna,<sup>b</sup> Giacomo Filippini,<sup>b</sup> Arianna Actis,<sup>c</sup> Mario Chiesa,<sup>c</sup> Manuela Bevilacqua,<sup>de</sup> Paolo Fornasiero,<sup>be</sup> Maurizio Prato,<sup>bfg</sup> David Beljonne<sup>ga</sup> and Roberto Lazzaroni<sup>ga</sup>

Graphitic carbon nitride (gCN) is an important heterogeneous metal-free catalytic material. Thermally induced post-synthetic modifications, such as amorphization and/or reduction, were recently used to enhance the photocatalytic response of these materials for certain classes of organic transformations, with structural defects possibly playing an important role. The knowledge of how these surface modifications modulate the photocatalytic response of gCN is therefore not only interesting from a fundamental point of view, but also necessary for the development and/or tuning of metal-free gCN systems with superior photo-catalytic properties. Herein, employing density functional theory calculations and combining both the periodic and molecular approaches, in conjunction with experimental EPR measurements, we demonstrate that different structural defects on the gCN surface generate distinctive radical defect states localized within the electronic bandgap, with only those correlated with amorphous and reduced gCN structures being photo-active. To this end, we (i) model defective gCN surfaces containing radical defect states; (ii) assess the interactions of these defects with the radical precursors involved in the photo-driven alkylation of electron-rich aromatic compounds (namely perfluoroalkyl iodides); and (iii) describe the photo-chemical processes triggering the initial step of that reaction at the gCN surface. We provide a coherent structure/photo-catalytic property relationship on defective gCN surfaces, elaborating how only specific defect types act as binding sites for the perfluoroalkyl iodide reagent and can favor a photo-induced charge transfer from the gCN surface to the molecule, thus triggering the perfluoroalkylation reaction.

Received 15th July 2022  
Accepted 1st August 2022

DOI: 10.1039/d2sc03964h

rsc.li/chemical-science

## Introduction

Photocatalysis with metal-free catalysts is gaining scientific and industrial attention in view of the growing demand for new sustainable synthetic schemes to access chemicals of extensive use.<sup>1</sup> In comparison to homogeneous systems, heterogeneous metal-free catalysts offer an edge due to their recyclability, low cost, good compatibility with specific reaction environments and sustained reactivity over time. Graphitic carbon nitride (hitherto abbreviated as gCN), a relatively wide band-gap semiconductor, with easy availability, low cost, good stability and excellent electronic properties<sup>2,3</sup> is being extensively used in photocatalytic applications such as H<sub>2</sub> production, water oxidation and CO<sub>2</sub> reduction.<sup>4-6</sup> Recent studies have extended the use of gCN to a wider range of industry-relevant chemical reactions which span from photocatalytic oxidation of small molecules<sup>7,8</sup> to C-C and C-heteroatom coupling reactions,<sup>9-12</sup> including functionalization of arenes and heteroarenes with fluoroalkylated groups.<sup>12-14</sup> The heterogeneous forms of gCN such as mesoporous carbon nitride, polymeric carbon nitride, potassium poly(heptazine imides) were also successfully

<sup>a</sup>Laboratory for Chemistry of Novel Materials, Materials Research Institute, University of Mons, Place du Parc 20, Mons 7000, Belgium. E-mail: david.beljonne@umons.ac.be; roberto.lazzaroni@umons.ac.be

<sup>b</sup>Department of Chemical and Pharmaceutical Sciences, INSTM, University of Trieste, Via L. Giorgieri 1, Trieste 34127, Italy

<sup>c</sup>Department of Chemistry, University of Torino, NIS Centre of Excellence, Via Giuria 9, Torino 10125, Italy

<sup>d</sup>Institute of Chemistry of OrganoMetallic Compounds (ICCOM-CNR), via Madonna del Piano 10, Sesto Fiorentino 50019, Italy

<sup>e</sup>Center for Energy, Environment and Transport Giacomo Ciamician and ICCOM-CNR Trieste Research Unit, University of Trieste, via L. Giorgieri 1, I-34127 Trieste, Italy

<sup>f</sup>Center for Cooperative Research in Biomaterials (CIC biomaGUNE), Basque Research and Technology Alliance (BRTA), Paseo de Miramón 182, Donostia San Sebastián 20014, Spain

<sup>g</sup>Basque Foundation for Science, Ikerbasque, Bilbao 48013, Spain

† Electronic supplementary information (ESI) available: Protocols for the EPR, EIS and H<sub>2</sub> evolution measurements; molecular models for defective gCNs; electronic structure of defective gCNs; electronic structure and adsorption energies of C<sub>4</sub>F<sub>9</sub>-I on defective gCNs; NTO analysis of C<sub>4</sub>F<sub>9</sub>-I/gCN systems; CIF files for the defective gCN structures. See <https://doi.org/10.1039/d2sc03964h>



employed for a plethora of applications such as the photocatalytic/photoredox sulfonylation of alkenes, ketalization of different ketones, aerobic oxidative coupling of resveratrol, and aerobic oxidation of benzylamines.<sup>15</sup>

Structural modifications by simple and scalable approaches were also proposed<sup>16</sup> to fine-tune the properties and reactivity of gCN surfaces. For instance, protonation and doping of gCN can be employed to fine-tune the electronic properties and also modulate the available surface area,<sup>6,16</sup> whereas inclusion of metallic ions *via* intercalation can decrease the band gap, enhance charge carrier mobility and provide additional catalytic sites.<sup>17,18</sup> More recently, carbon nitride surfaces, modified to include electron-rich phenyl moieties, were employed for the metal-free, visible light-responsive photocatalytic synthesis of methyl esters from benzylic alcohols, photocatalytic oxidative esterification by the inclusion of surface acidic sites<sup>19</sup> and for the photocatalytic aerobic oxidation of alcohols, under irradiation using an extended visible-light wavelength of 620 nm.<sup>20</sup> Some of us have recently demonstrated that the visible-light photocatalytic activity of gCN for the perfluoroalkylation of electron-rich organic substrates can be substantially enhanced by diverse surface modifications.<sup>21</sup> Briefly, this strategy includes post-synthetic modifications of melamine-derived gCN (Fig. 1a) to introduce structural and chemical variations into pristine gCN by adopting specific protocols for reduction (red-gCN), oxidation (oxy-gCN) and thermal amorphization (am-gCN). The photocatalytic response of these materials was then investigated for the grafting of perfluorinated groups on aromatic cores, which is a reaction of major interest in medicinal chemistry, agro-chemistry and materials science.<sup>14,22–24</sup> The test case that has been described in ref. 21 is the reaction between 1,3,5-trimethoxy-benzene (**1a**), a model electron-rich aromatic

molecule, and perfluorobutyl iodide (**2a**, C<sub>4</sub>F<sub>9</sub>-I) as the radical precursor to give the fluorinated product **3a** (Fig. 1b). It has been proposed that the key step in the reaction between C<sub>4</sub>F<sub>9</sub>-I and 1,3,5-trimethoxy-benzene is a photo-induced electron transfer from the gCN catalyst surface to the C<sub>4</sub>F<sub>9</sub>-I molecule.

Interestingly, it was found that the photo-catalytic activity of the reduced and amorphous gCN forms is much higher ( $\geq 90\%$ ) compared to that of the gCN surface in its pristine (20%) and oxidized (10%) forms. However, all these post-synthetically modified gCN structures show: (i) a similar C/N ratio, as indicated by the XPS data, with the exception of am-gCN and red-gCN forms that show a comparatively lower N-content attributed to the presence of nitrogen vacancies, consistent with previous observations;<sup>15,25–28</sup> (ii) a similar optical bandgap, as obtained by diffuse reflectance spectroscopy (DRS); and (iii) the presence of native and photo-induced radical species in all samples, as observed by electron paramagnetic resonance (EPR) measurements.<sup>29</sup>

Therefore, despite the structural and spectroscopic data at hand, we are still missing a detailed fundamental understanding of: (i) the nature of the interaction between the perfluoroalkyl iodide and the (modified) gCN surface(s), (ii) the excited-state dynamics triggering the reaction and (iii) the influence of the gCN surface composition on the photocatalytic activity. Such an understanding is crucial to establish a solid structure/property/activity relationship for gCN photocatalysis and to further broaden the spectrum of the reactions that could be catalyzed by gCN, aiding the process of rational function-driven catalyst design for future applications.

Here, we present a detailed theoretical study of model perfluoroalkyl iodide/gCN systems with the aim of providing a comprehensive understanding of the recently reported NMR,

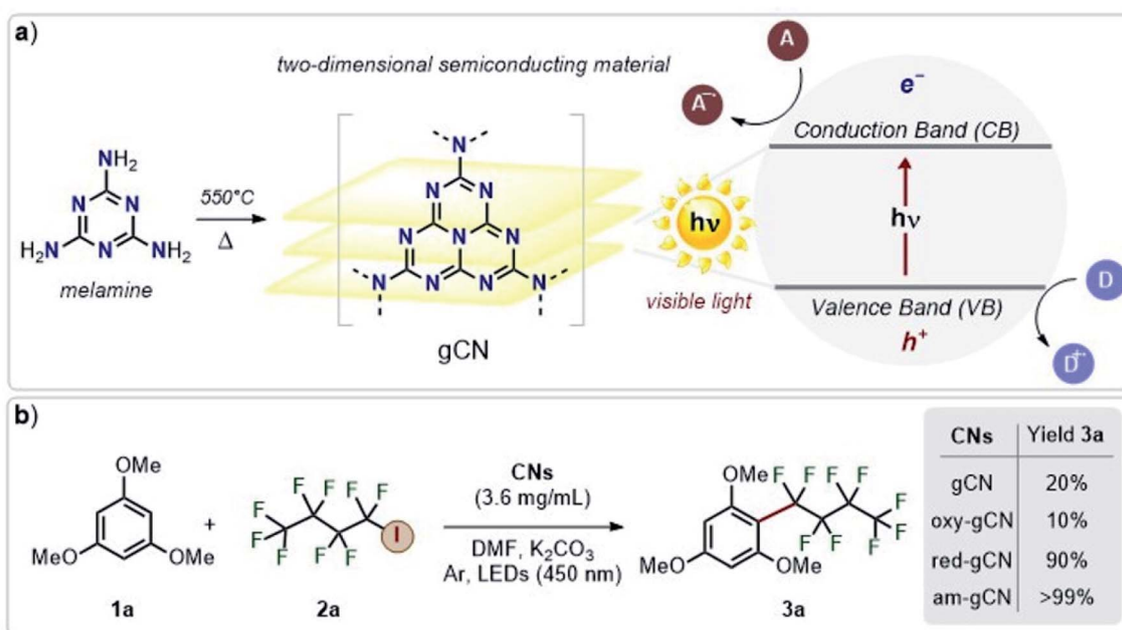


Fig. 1 Schematic representation of (a) graphitic carbon nitride obtained by the pyrolysis of melamine and its conduction and valence bands (CB and VB); (b) the reaction under study and its yield, highlighting the difference in the catalytic efficiency of the various types of gCN. A: electron acceptor; D: electron donor.



XPS and EPR spectroscopic data and photocatalytic activities.<sup>21,29</sup> More specifically, we focus on the specific interaction between different gCN surfaces and the C<sub>4</sub>F<sub>9</sub>-I molecule, which is considered to be the crucial first step governing the experimental photo-catalytic efficiency, quantified only by the global yield of 2-nonafluorobutyl-1,3,5-trimethoxybenzene (product **3a** in Fig. 1).<sup>21</sup> The subsequent reaction steps leading to the formation of product **3a** correspond to a classical case of homolytic aromatic substitution.

For that purpose, we apply a two-fold modelling strategy to: (i) evaluate the ground-state interaction between the pristine, red-gCN, oxy-gCN and am-gCN surfaces and the C<sub>4</sub>F<sub>9</sub>-I molecule to quantify the adsorption profiles and the electronic structure of the combined gCN/C<sub>4</sub>F<sub>9</sub>-I systems and (ii) to determine, with time-dependent DFT (TD-DFT) calculations on the gCN/C<sub>4</sub>F<sub>9</sub>-I systems, the excited-state electronic structure, in order to shed light on the photo-chemical response of these gCN surfaces in the presence of the C<sub>4</sub>F<sub>9</sub>-I molecule. A detailed description of the model gCN structures, the computational protocol employed, and the results pertaining to the ground-state and excited-state gCN/C<sub>4</sub>F<sub>9</sub>-I interactions are presented in the following sections. We find a clear correlation between the nature of the defects on the gCN surface, their electronic structure, and the viability of a photo-induced electron transfer from gCN to the C<sub>4</sub>F<sub>9</sub>-I molecule, which is most probably the key step that triggers its coupling with the electron-rich aromatic molecule.

## Modelling methodology

### Modelling protocol

The following modelling protocol was employed to evaluate the ground-state and excited-state interactions between defective gCN surfaces and the C<sub>4</sub>F<sub>9</sub>-I molecule:

First, starting from pristine gCN, we select model systems including structural point defects to mimic the reduced, amorphous and oxidized forms and we assess how these defects perturb the gCN electronic structure.

We then compute the ground-state interaction between the different defective gCN surfaces and the C<sub>4</sub>F<sub>9</sub>-I molecule to: (i) extract the electronic structure of the defective gCN surfaces and the C<sub>4</sub>F<sub>9</sub>-I molecule adsorbed on those gCN surfaces (hitherto referred as gCN/C<sub>4</sub>F<sub>9</sub>-I for simplicity), (ii) evaluate the adsorption energies, (iii) quantify the ground state (partial) charge transfer between the gCN surface and the molecule, and (iv) evaluate the bond order parameters in relation to the <sup>19</sup>F-NMR experimental results<sup>21,29</sup> for a qualitative assessment of halogen bonding.

Finally, we perform Time-Dependent DFT (TD-DFT) calculations on the gCN/C<sub>4</sub>F<sub>9</sub>-I systems to determine the excited-state electronic structure, in order to shed light on the photo-chemical response of different gCN surfaces in the presence of the C<sub>4</sub>F<sub>9</sub>-I molecule.

These steps, which form the core of the gCN/C<sub>4</sub>F<sub>9</sub>-I interaction, represent the ground-state and excited-state interactions between a given defective form of the gCN surface and the C<sub>4</sub>F<sub>9</sub>-I molecule. More specifically, we investigate the ground-state

electronic interactions between gCN surfaces and the C<sub>4</sub>F<sub>9</sub>-I molecule using both the 2D periodic approach and a molecular approach. For the latter, we use a model triangular heptazine trimer representing a fragment of the (defective) gCN surface (see Fig. SI-1†) that is made to interact with C<sub>4</sub>F<sub>9</sub>-I, as this offers computationally easy access to excited-state calculations. The choice of using a molecular (heptazine trimer) model for excited-state studies is mainly motivated by the fact that: (i) the molecular representation of gCN, as often considered in

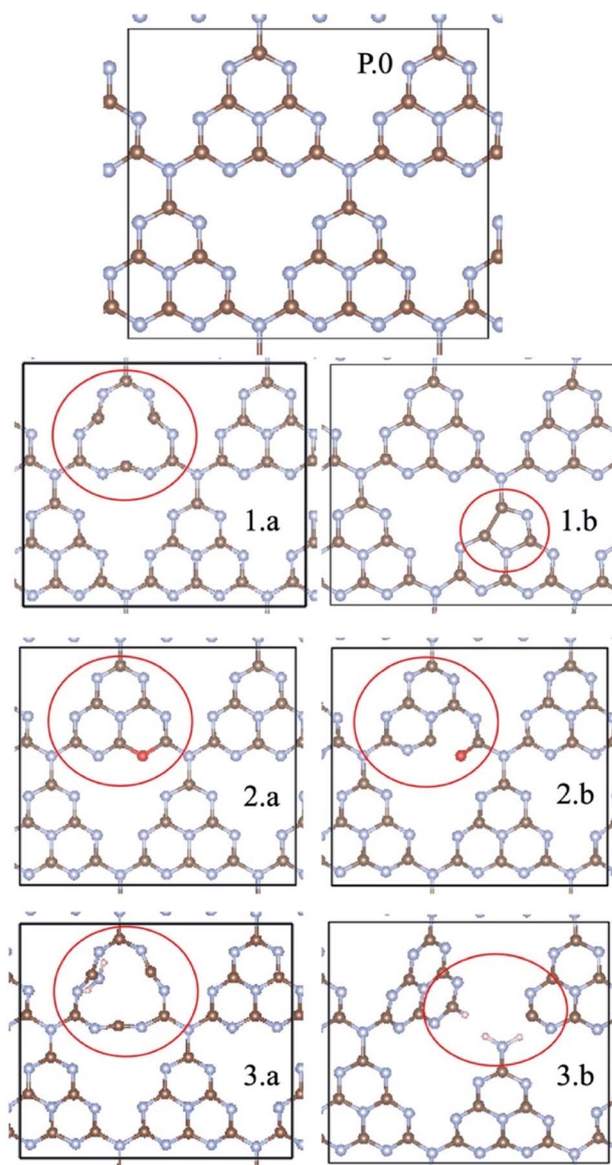


Fig. 2 gCN surfaces in their pristine (P.0) and defective forms (1.a/b, 2.a/b, and 3.a/b) considered in this work (carbon: brown; nitrogen: blue; oxygen: red; hydrogen: pink). am-gCN surfaces are represented to contain nitrogen vacancies either at the centre or at the edge of a heptazine unit, as shown in 1.a to 1.b; oxy-gCN surfaces in their pyran-like (2.a) and carbonyl-like (2.b) forms were created by substituting the heptazine edge nitrogen vacancy. Structural defects with disconnected bonds between nitrogen and carbon atoms, either at the heptazine centre or the edge (3.a and 3.b, respectively) represent the red-gCN forms.



previous studies,<sup>30–32</sup> opens up a wide range of hybrid functionals to evaluate the optical properties, which would be computationally prohibitive within the periodic approaches, and (ii) as the gCN systems are characterized by limited band dispersion, the interaction of the C<sub>4</sub>F<sub>9</sub>-I molecule with a molecular fragment considered to represent the periodic equivalent, turns out to be computationally appropriate, as detailed in the subsequent sections.

### Computational details

All periodic DFT calculations were performed using the VASP code and the projector-augmented wave (PAW) basis set.<sup>33,34</sup> Exchange and correlation effects are treated at the Perdew–Burke–Ernzerhof (PBE) level with the dispersion forces treated by using Grimme correction (PBE + D2),<sup>35,36</sup> with a kinetic energy cut-off of 500 eV and using a Monkhorst–Pack sampling of  $3 \times 3 \times 1$  for the Brillouin zone (BZ) integration on all the gCN unit cells replicated  $4 \times 4 \times 1$  times, with the vacuum spacing set to be 30 Å to avoid the interaction with periodic images. Dipole moment correction was employed along the 'c' axis (Z direction & perpendicular to the gCN surface). The geometries of pristine and defective gCN surfaces along with C<sub>4</sub>F<sub>9</sub>-I adsorbed were fully optimized at the PBE/GGA level of theory prior to the calculation of the electronic properties at the hybrid (HSE06)<sup>37</sup> level of theory. Bond orders and ground state partial charge transfers were computed using the DDEC06 method.<sup>38</sup> DFT and TD-DFT calculations of the molecular equivalent gCN units with and without C<sub>4</sub>F<sub>9</sub>-I adsorption were performed using the Gaussian16 program<sup>39</sup> at the B3LYP/6-311G(d,p) level of theory for molecular optimizations. The subsequent electronic structure and UV-vis absorption spectra calculations, along with the corresponding NTO<sup>40</sup> and attachment-detachment density<sup>41,42</sup> analysis were performed on the molecular systems at the hybrid level of theory (HSE1HSE), to maintain consistency with the electronic structure obtained with the periodic DFT calculations. Prior to TD-DFT calculations on the molecular systems, the consistency between the periodic and molecular gCN/C<sub>4</sub>F<sub>9</sub>-I descriptions was ensured by

comparing the electronic structure at the hybrid level (HSE06) of theory.

### gCN structural models

For the periodic approach, we started from a monolayer of pristine gCN surface (P.0 in Fig. 2), and incorporated a large variety of structural defects on the gCN surface, classified based on the post-synthetic experimental treatment,<sup>21</sup> chemical intuition and literature data, and we calculated their electronic structure and stability. The most representative gCN systems in their pristine (gCN), red-gCN, oxy-gCN and red-gCN forms are shown in Fig. 2. To simulate the formation of amorphous gCN (am-gCN), for which some nitrogen loss has been observed experimentally,<sup>21,29</sup> defects were created by removing one nitrogen atom from the surface of the gCN unit cell. The nitrogen atom is removed from either the core of a heptazine unit or from the edge of a heptazine unit (Fig. 2(1.a and 1.b), respectively). Both these forms of nitrogen vacancies are thermodynamically stable. Here we find that the defect formation energy is 4.71 eV (108.6 kcal mol<sup>-1</sup>) and 4.80 eV (110.7 kcal mol<sup>-1</sup>) respectively, consistent with previous results.<sup>43</sup>

To represent gCN treated under oxidative conditions, oxygenated gCN models (oxy-gCN), considered with respect to recent investigations,<sup>32</sup> were obtained by substituting the nitrogen atoms at the edge of a heptazine unit, to obtain either a pyran-like unit (see Fig. 2(2.a)) or a carbonyl-containing gCN form (see Fig. 2(2.b)).

Since the reductive treatment is carried out in the presence of H<sub>2</sub>, we hypothesize that structural defects appearing upon bond cleavage during reductive treatment are likely to be at least partly saturated by hydrogen atoms. To model the situation of reduced gCN forms (red-gCN), two disconnected gCN forms were considered wherein the bonds between the nitrogen atom and the surrounding carbon atoms were cleaved, either at the center of the heptazine unit (see Fig. 2(3.a)) or at the edge of the heptazine unit (see Fig. 2(3.b)) and were partly saturated with hydrogen atoms.

**Table 1** Descriptive nomenclature of the various gCN defective forms used in this work along with their reference numbers. The reference numbers correspond to the respective gCN structures from Fig. 2 and SI-1 and are hitherto used to indicate the specific gCN surface. The nature of the defect type present on the gCN surface is also shown for clarity

Ref.	Type	Description of the gCN surface
P.0	Pristine	gCN-pristine
1.a	am-gCN	gCN-N vacancy at the center of a heptazine unit Nature of the defect: radical with one unpaired electron
1.b		gCN-N vacancy at the edge of a heptazine unit Nature of the defect: radical with one unpaired electron
2.a	oxy-gCN	Oxygen substitution at the edge of a heptazine unit: pyran-like form Nature of the defect: radical with one unpaired electron
2.b		Oxygen substitution at the edge of a heptazine unit: carbonyl form Nature of the defect: radical with one unpaired electron
3.a	red-gCN	Disconnected gCN network, with N-C bonds cleaved at the center of a heptazine unit Nature of the defect: radical with two unpaired electrons (bi-radical form)
3.b		Disconnected gCN network, with N-C cleaved at the edge of a melamine unit Nature of the defect: radical with one unpaired electron



It must be noted that all these structures, as shown in Fig. 2(1.a/b–2.3.a/b), are open-shell systems with at least one unpaired electron per unit cell, consistent with the experimental EPR results that indicate the presence of radicals on all the gCN surfaces, irrespective of their treatment (oxidation, reduction, and amorphization). The defect formation energies of the oxy-gCN forms were found to be in the 0.6 eV to 1.0 eV range (1.08 eV and 0.66 eV for structures 2.a and 2.b, respectively) whereas that of red-gCN (structure 3.a) is  $\approx 4.1$  eV. See ESI, Fig. SI-3,† for the defect formation energies of these defective gCN forms, along with the corresponding schematics of the reaction mechanism considered.

The molecular equivalents to the periodic structures for pristine and defective gCN forms were built by suitably modifying the pristine heptazine trimer; those model defective molecules are shown in Fig. SI-2.† The specific nomenclature of the different gCN structures pertaining to periodic and molecular gCN forms (related to Fig. 2 and SI-2,† respectively) along with their reference numbers are listed in Table 1. Hitherto, to ease the description, the reference number from Table 1 is used to refer to any given gCN surface/form that is discussed in this work.

## Results and discussion

### Electronic structure of the defective gCN forms

The electronic structure of the different gCN derivatives from Fig. 2, obtained at the HSE06 level of theory, is shown in Fig. 3. While pristine gCN shows an electronic band gap of  $\approx 2.8$  eV, the radical gCN forms show additional defect levels within the electronic band gap. More specifically, for the radical am-gCN form with a nitrogen vacancy at the center of the heptazine unit (see structure 1.a and Fig. 3(1.a)), a localized occupied defect level (*viz.* the radical state with one unpaired ' $\alpha$ ' electron) appears at around mid-gap, *i.e.*  $\approx 1.2$  eV from the conduction band (CB) edge and its corresponding unoccupied electronic level (*viz.* the ' $\beta$ ' state pertaining to the occupied ' $\alpha$ ' state at  $\approx 1.2$  eV) merges with the CB edge. When the nitrogen vacancy is present at the edge of the heptazine unit in am-gCN (see structure 1.b and Fig. 3(1.b)), an occupied defect level (' $\alpha$ ' state) is positioned at  $\approx 1.8$  eV from the CB edge, while a new, corresponding unoccupied (empty ' $\beta$ ' state) electronic level, is positioned within the band gap at  $\approx 0.6$  eV from the CB edge.

For the oxygenated gCN surface with a pyran-like defect (see structure 2.a and Fig. 3(2.a)), the occupied radical defect level is positioned within the bandgap at  $\approx 0.9$  eV from the CB edge. For the carbonyl-containing oxy-gCN form (see structure 2.b and Fig. 3(2.b)), the occupied radical defect level basically merges with the top of the valence band (VB) edge while its corresponding empty level is positioned within the band gap at  $\approx 0.5$  eV from the CB edge.

For the reduced gCN system with a disconnection in the C–N bonding at the center of the heptazine unit (see structure 3.a and Fig. 3(3.a)), the occupied radical defect level is positioned within the bandgap at  $\approx 1.7$  eV from the CB edge. This gCN form (structure 3.a) is a bi-radical, as confirmed by the ground-state broken symmetry calculations, with an energy difference

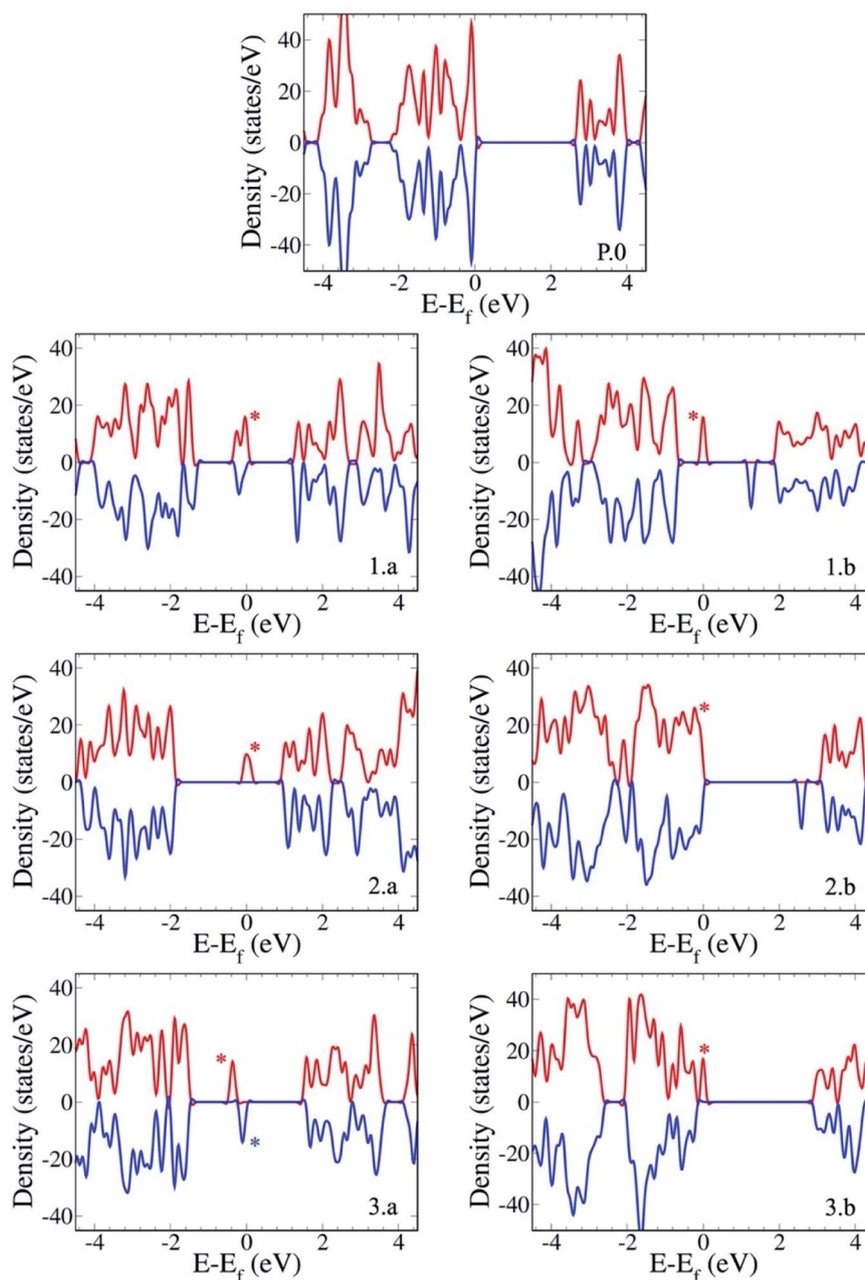
of  $\approx 0.5$  eV between the partly occupied  $\alpha$  and  $\beta$  defect states (see the ESI† for details). For the red-gCN form with a C–N disconnection at the heptazine edge (see structure 3.b and Fig. 3(3.b)), the occupied defect level is positioned within the band gap but very close (0.2 eV) to the VB edge.

The electronic structure of the pristine and defective molecular gCN models, obtained with pristine and modified heptazine units, shows a very good quantitative agreement with the periodic gCN forms in terms of the electronic structure, energy gap and orbital symmetry of the respective gCN forms, as shown in the ESI (Fig. SI-4 and SI-5†), implying the transferability between the periodic and the molecular approaches. In addition, all the intra-gap levels in the radical gCN forms are localized in the vicinity of the defect for both periodic and molecular equivalents, as can be seen in the crystalline/molecular orbitals shown in the ESI (Fig. SI-5†). Indeed, it is the nature of the localized radical defect states present on the gCN surface that generate a significant and specific interaction with the perfluoroalkyl iodide reagent, which subsequently modulate the photo-induced charge transfer from the gCN surface to the  $C_4F_9-I$  molecule (*vide infra*).

It is to be noted that gCN samples prepared by direct thermocondensation of precursors could form a melon-based structure, as reported previously.<sup>44</sup> A melon-based carbon nitride principally represents a one-dimensional (1D) class wherein some of the heptazine sub-units form 1D polymer strands and the other heptazine sub-units align in a zigzag manner due to a hydrogen bonding network with the nitrogen atoms in the 1D strands, resulting in an interconnected network of heptazine units over a flat basal plane. The periodic and the molecular models considered in this work represent the two limit structures of a CN surface, namely, the periodic monolayer represents the result of the complete crosslinking of a melon model into a two-dimensional (2D) structure whereas the molecular model is the zero-dimensional (0D) representation of independent heptazine units, which are also present in the melon-like 1D structure. As both the defective periodic (2D) and molecular (0D) models show a similar electronic structure (see Fig. SI-4 and SI-5† for details), we believe that the conclusions drawn from this work could be extended to other gCN forms, in particular the melon-based 1D carbon nitride structures.

It is also of interest to analyze the computed spin-density distributions on various defective gCN forms. These systematically show larger spin densities on carbon atoms, along with smaller contributions on the nitrogen atoms (see the case of the am-gCN models in Fig. 4(1.a/b)). This result concurs with EPR experiments shown in Fig. 4c, which indicate that all the gCN structures (including pristine gCN) contain radical carbon species, with the radical concentration being the largest in the am-gCN form ( $4.5 \times 10^{16}$  spins/g *vs.*  $2.5 \times 10^{16}$  in pristine gCN). Importantly,  $^{14}N$  2D-EPR experiments (see ESI, section S1†) performed on am-gCN (Fig. 4d) provide evidence for a minute  $^{14}N$  hyperfine coupling ( $A_{\max} \approx 1$  MHz), consistent with the delocalization of the unpaired electron wave function over a number of nitrogen atoms, which further confirms the relevance of the model systems chosen to represent the actual surface-active sites.





**Fig. 3** Electronic structure of the gCN surfaces in their pristine (P.0) and defective forms considered in this study (with am-gCN represented by 1.a/b, oxy-gCN represented by 2.a/b and red-gCN represented by 3.a-b). The positive (red) and negative (blue) density-of-states curves correspond to the  $\alpha$  and  $\beta$  electron densities, respectively. The zero of the energy scale, *i.e.*, the Fermi level, corresponds to the highest-occupied electronic state. Electronic energy levels corresponding to the localized occupied defect states at the Fermi level are indicated by an asterisk.

In addition, the  $\text{H}_2$  evolution reaction by photo-reformation of ethanol in aqueous solution was performed using all four gCN derivatives, employing Pt as a co-catalyst to decrease the recombination rates of the photo-generated electrons and holes. Both red-CN and ox-CN forms achieve higher maximum  $\text{H}_2$  formation rates when compared to the g-CN or am-CN forms, confirming that specific tuning of the CN structure is instrumental for the specific catalytic application (see ESI, section S2† for details). Furthermore, electrochemical impedance spectroscopy (EIS) measurements were performed on the

pristine and am-gCN forms, both in the dark and with a light source, to investigate the effect on the charge transfer mechanism at the electrode–electrolyte interface (see ESI, section S3† for details). According to the data summarized in Table SI-1,† a decrease in the charge transfer resistance ( $R_{ct}$ ) in the presence of a light source was found with respect to the dark condition for both am-CN and g-CN materials. The differences in the absolute  $R_{ct}$  values can be attributed to differences in the homogeneity of the catalyst deposition on the GC support for the different materials. Nevertheless, the evolution of the  $R_{ct}$



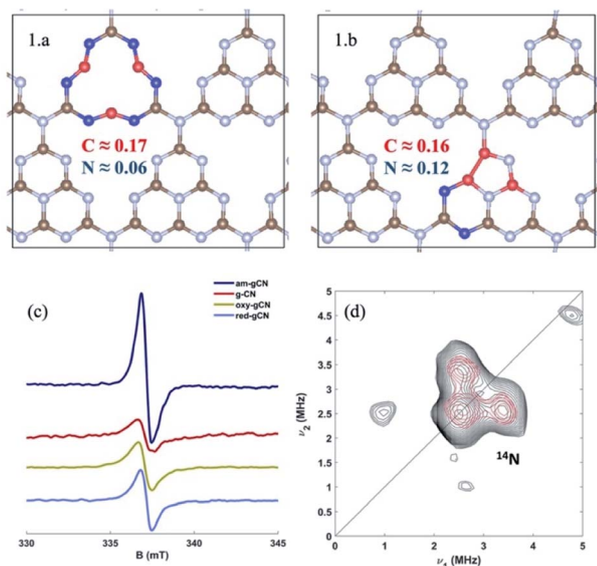


Fig. 4 Atomic spin density distribution in the models for amorphous gCN (structures 1.a and 1.b), extracted using the DDEC6 methodology, and bottom: left (c) continuous wave X-band EPR spectra of the four gCN forms considered in this work recorded in the dark at 77 K; bottom: right (d) Q-band  $^{14}\text{N}$  HYSCORE spectrum of the amorphous gCN system (black experiment, red simulation). The spectrum was recorded at 110 K. Experimental details are shown in the ESI (Section S1†), along with the simulation parameters.

values clearly shows that the charge transfer at the electrode-electrolyte interface, already occurring under the dark condition, has been further improved in the presence of light, clearly confirming the photo-induced nature of the processes taking place at the CN-solution interface.

#### gCN/ $\text{C}_4\text{F}_9\text{-I}$ systems: adsorption energy and bond order analysis

To better quantify the interaction between pristine/defective gCN surfaces and the  $\text{C}_4\text{F}_9\text{-I}$  molecule and to get further

insights, we calculated the adsorption energies between the model gCN surfaces and the  $\text{C}_4\text{F}_9\text{-I}$  molecule, along with estimations of the bond order (focusing on the iodine atom and the neighboring fluorine atoms) and the ground-state partial charge transfer between the partners, as shown in Table 2. Herein, we describe only the lowest-energy conformation of the  $\text{C}_4\text{F}_9\text{-I}$  molecule adsorbed on pristine/defective gCN surfaces, which corresponds to the long axis of the molecule being parallel to the basal plane of the gCN surface, as presented in Fig. 5 and SI-7.† The alternate conformation, wherein the long axis of the molecule is perpendicular to the gCN surface, is  $\approx 2 \text{ kcal mol}^{-1}$  higher in energy, irrespective of the gCN form.

The adsorption energy of  $\text{C}_4\text{F}_9\text{-I}$  is  $\approx 6 \text{ kcal mol}^{-1}$  on the pristine gCN surface (structure P.0) but it has significantly higher values when radical defects are present. More specifically, for the am-gCN forms (structures 1.a and 1.b), the adsorption energy of  $\text{C}_4\text{F}_9\text{-I}$  ranges from  $\approx 11 \text{ kcal mol}^{-1}$  to  $\approx 17 \text{ kcal mol}^{-1}$ , the highest value being found when a nitrogen vacancy is present in the middle of a heptazine unit (structure 1.a). In contrast, when oxygenated defects (pyran/carbonyl-like) are present on the gCN surface, the adsorption energies are reduced to  $\approx 8 \text{ kcal mol}^{-1}$ . The adsorption energies of  $\text{C}_4\text{F}_9\text{-I}$  on the molecular gCN models are in good quantitative agreement with the periodic models (see Table SI-2†). This decrease in adsorption energy for oxygenated defects, despite the radical nature of these defects, could be attributed to the electrostatic repulsion between the oxygen and iodine atoms. In addition, the strong adsorption of  $\text{C}_4\text{F}_9\text{-I}$  on most red/am-gCN surfaces (structures 1.a/b and 3.a) is also associated with a: (i) non-negligible ground-state partial charge transfer from the gCN surface to the  $\text{C}_4\text{F}_9\text{-I}$  molecule ( $\approx 0.1q$ ) and (ii) a small elongation of the carbon-iodine (C-I) bond of the  $\text{C}_4\text{F}_9\text{-I}$  molecule, of  $\approx 0.06 \text{ \AA}$ .

Interestingly, the adsorption energies also reflect a variation in the net bond order of the iodine atom: the highest bond order on iodine is observed when the  $\text{C}_4\text{F}_9\text{-I}$  molecule is adsorbed on gCN surfaces containing nitrogen vacancies at the

**Table 2** Adsorption energies (in  $\text{kcal mol}^{-1}$ ) of the  $\text{C}_4\text{F}_9\text{-I}$  molecule adsorbed on the gCN surface in its pristine and defective forms, along with the respective DDEC bond order and partial charge transfer analysis. The bond order for iodine corresponds to the total bond order on the iodine atom of the molecule adsorbed on the pristine/defective gCN surface.  $\Delta F$  corresponds to the relative change in the bond order on the fluorine atoms in the vicinity of the iodine atom with respect to its value when the  $\text{C}_4\text{F}_9\text{-I}$  molecule is adsorbed on the pristine gCN surface.  $\Delta d$  corresponds to the relative change in the bond length between the iodine atom and the connected carbon atom of the  $\text{C}_4\text{F}_9\text{-I}$  molecule with respect to its value when the  $\text{C}_4\text{F}_9\text{-I}$  molecule is adsorbed on the pristine gCN surface. A negative value of partial charge surface area ( $\partial q/\text{\AA}$ ) corresponds to a charge transfer from the gCN surface to the  $\text{C}_4\text{F}_9\text{-I}$  molecule, while a positive value represents the charge transfer from the  $\text{C}_4\text{F}_9\text{-I}$  molecule to the gCN surface

Ref.	System	$E_{\text{Ads}}$ ( $\text{kcal mol}^{-1}$ )	Bond order		
			I	Bond length ( $\Delta d$ , $\text{\AA}$ )	Partial charge transfer ( $\partial q/\text{\AA}$ )
	P.0	6.1	1.28	-na-	+0.03
	1.a	17.4	1.57	0.20	-0.10
	1.b	11.2	1.38	0.05	-0.07
	2.a	8.1	1.35	0.04	-0.01
	2.b	7.0	1.21	0.01	-0.01
	3.a	12.5	1.51	0.15	-0.10
	3.b	11.4	1.35	0.05	-0.03



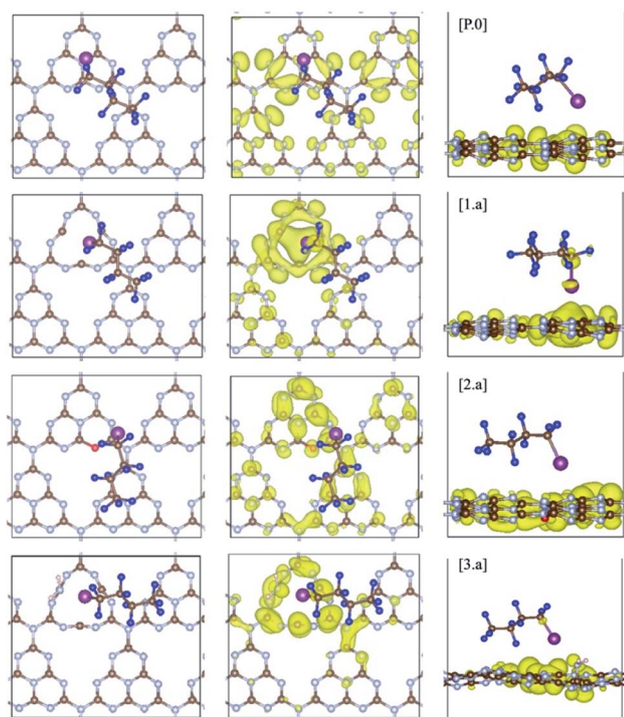


Fig. 5 C<sub>4</sub>F<sub>9</sub>-I adsorbed on pristine (P.0) and defective gCN surfaces (am-gCN: 1.a, oxy-gCN: 2.a and red-gCN: 3.a), in their lowest-energy conformations wherein the long axis of the molecule is parallel to the basal plane of the given gCN surface (carbon: brown; nitrogen: blue; oxygen: red; hydrogen: pink, fluorine: navy and iodine: magenta). Wavefunction plots at the fermi level (highest-occupied electronic state) for gCN/C<sub>4</sub>F<sub>9</sub>-I systems are also reported for gCN surfaces in its (P.0) pristine and (1.a, 2.a and 3.a) defective forms. The wavefunction is found to be partially localized on the C<sub>4</sub>F<sub>9</sub>-I molecule for structures 1.a and 3.a. Corresponding data for gCN structures 1.b, 2.b and 3.b are shown in Fig. SI-6.†

center of the heptazine unit (structures 1a and 3a, and Table 2). This is an indication of the stronger interaction of the iodine atom with the surface defects in these systems. In addition, a variation of net bond order is also observed on the two fluorine atoms that are in proximity to the iodine atom. The relative change in the bond order of these fluorine atoms for different defect types compared to pristine gCN is also the highest for the gCN surfaces containing a nitrogen vacancy at the center of the heptazine unit (see Table 2).

The matching between the increased adsorption energy of the C<sub>4</sub>F<sub>9</sub>-I molecule on defective surfaces representing the amorphous and reduced gCN and the relative variation in the bond orders of iodine and fluorine atoms correlates well with the variation in the experimental <sup>19</sup>F-NMR  $T_1/T_2$  ratios,<sup>21,29</sup> with the highest  $T_1/T_2$  ratios observed for the amorphous/reduced gCN forms. More specifically, these earlier studies,<sup>21,29</sup> using perfluorohexyl iodide as a probe molecule, reported higher <sup>19</sup>F-NMR  $T_1/T_2$  ratios for am-gCN and red-gCN forms compared to pristine/oxy-gCN forms, pointing to stronger molecule/surface interactions in the former systems. These strong interactions between the defective/modified gCN surface and the perfluorohexyl iodide molecule, were hypothetically ascribed to the

formation of halogen bonding (N⋯I) between the freely accessible nitrogen (N) atoms from the defective amorphized gCN surfaces and the iodine(I) atom of the probe molecule.

To validate this hypothesis, we considered C<sub>4</sub>F<sub>9</sub>-I, C<sub>4</sub>F<sub>10</sub> and C<sub>4</sub>H<sub>10</sub> as probe molecules and computed their interactions on the pristine gCN surface (structure P.0) and on the am-gCN surface containing a nitrogen vacancy at the center of the heptazine unit (structure 1.a), as shown in the ESI (Fig. SI-8†). On the pristine gCN surface, all the three probe molecules show a similar adsorption energy in the order of  $\approx 4.5$  to  $6$  kcal mol<sup>-1</sup>, but on the am-gCN surface, their adsorption energies are larger, ranging from  $\approx 11.5$  kcal mol<sup>-1</sup> to  $\approx 17.5$  kcal mol<sup>-1</sup>. Of the three probe molecules adsorbed on the am-gCN surface, C<sub>4</sub>F<sub>9</sub>-I has the largest adsorption energy (17.4 kcal mol<sup>-1</sup>) while both C<sub>4</sub>F<sub>10</sub> and C<sub>4</sub>H<sub>10</sub> molecules show comparatively smaller adsorption energies of 12.7 and 11.4 kcal mol<sup>-1</sup>, respectively. In addition, both C<sub>4</sub>F<sub>10</sub> and C<sub>4</sub>H<sub>10</sub> molecules adsorbed on am-gCN surfaces are characterized by a negligible partial charge transfer between the surface and molecule ( $\approx 0.01$   $\partial q/A$ ) and no variations are observed: (i) in the bond orders of fluorine or hydrogen atoms; (ii) in the C-F/C-H bond lengths. In contrast, C<sub>4</sub>F<sub>9</sub>-I adsorbed on the am-gCN surface shows larger variations in terms of partial ground-state charge transfer ( $\partial q/A$ ), a relative change in the bond order of fluorine atoms ( $\Delta F$ ) and a relative change in the C-I bond length ( $\Delta d$ ), as shown in Table 2, and consequently shows the highest adsorption energy, which is indicative of the strong interaction between the C<sub>4</sub>F<sub>9</sub>-I molecule and the defective gCN surface.

It therefore follows that the increased interaction between the defective gCN surface and C<sub>4</sub>F<sub>9</sub>-I does not appear to be related to a conventional halogen bonding between the freely available nitrogen atoms from the defective gCN surface and the iodine atom of the molecule, as hypothesized previously based on the increased experimental <sup>19</sup>F-NMR  $T_1/T_2$  ratios. Instead, the calculations indicate that it results from a more complex interaction mechanism involving a reorganization of the electronic density on the terminal iodine, which is characterized by: (i) a partial ground state charge transfer from the defective gCN surface to the molecule; (ii) an increase in the ground state C-I bond length; and (iii) a consequential increase (decrease) in the bond order (length) between carbon and fluorine atoms neighboring the iodine atom. These specific interactions between the C<sub>4</sub>F<sub>9</sub>-I molecule and the defects present on the gCN surface could indeed be manifested into an increase in <sup>19</sup>F-NMR  $T_1/T_2$  ratios, observed experimentally.

### Electronic structure of the gCN/C<sub>4</sub>F<sub>9</sub>-I systems

The ground-state electronic structure of the gCN/C<sub>4</sub>F<sub>9</sub>-I systems is analyzed here with a specific focus on the energy separation between the highest-occupied crystalline states of gCN surfaces and the lowest unoccupied molecular states of C<sub>4</sub>F<sub>9</sub>-I, to get insights into the possibility of electron transfer between the partners upon photoexcitation. The electronic structure shown in Fig. 6 is extracted from the partial density of states (pDOS) obtained at the HSE06 level of theory, as shown in the ESI (Fig. SI-6 and SI-7†).



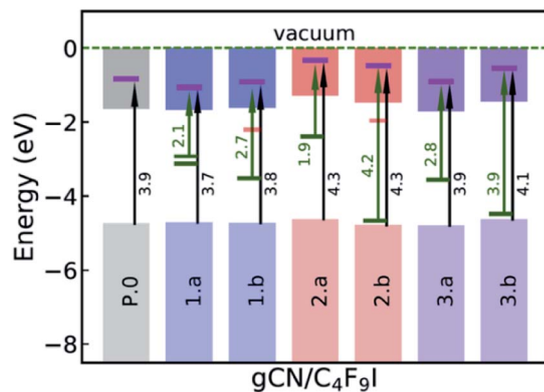


Fig. 6 Summarized electronic structure of the gCN/C<sub>4</sub>F<sub>9</sub>-I systems. The occupied (unoccupied) localized mid-gap defect states within the electronic band gap are indicated with green (red) bars while the iodine-centered electronic energy levels of the C<sub>4</sub>F<sub>9</sub>-I molecule within the conduction band are indicated with a magenta bar. Green (black) arrows correspond to the energy difference between the occupied localized mid-gap state (valence band edge) and the iodine-centered electronic level.

On the pristine gCN surface (P.0), the iodine-localized electronic level of the C<sub>4</sub>F<sub>9</sub>-I molecule lies deep in the conduction band of the system, at  $\approx 3.9$  eV from the VB edge (see Fig. 6). However, when radical defects are present on the gCN surface, gCN/C<sub>4</sub>F<sub>9</sub>-I systems show localized defect levels within the electronic band gap, similar to the description in Fig. 3. In the model systems for the amorphous gCN, structure 1.a, two mid-gap levels are positioned at  $\approx 1.5$  eV from the CB edge. The iodine centered C<sub>4</sub>F<sub>9</sub>-I electronic energy level within the conduction band is positioned at  $\approx 2.1$  eV ( $\approx 2.3$  eV) from the singly (doubly) occupied defect state and at  $\approx 3.7$  eV from the VB edge. For structure 1.b, the singly occupied mid-gap level is positioned at  $\approx 1.9$  eV from the CB edge, while the iodine-centered C<sub>4</sub>F<sub>9</sub>-I electronic energy level within the conduction band is positioned at  $\approx 2.7$  eV from this defect state and  $\approx 3.8$  eV from the VB edge.

For defective surfaces modeling oxidized gCN, the pyran-like structure (2.a) shows a singly occupied defect level at  $\approx 0.9$  eV from the CB edge, which is  $\approx 1.9$  eV from the iodine-centered C<sub>4</sub>F<sub>9</sub>-I electronic energy level within the conduction band. In

contrast, when a carbonyl-like defect (2.b) is present on the gCN surface, the singly occupied defect level is very close to the VB edge positioned at  $\approx 4.2$  eV from the iodine-centered C<sub>4</sub>F<sub>9</sub>-I electronic energy level. For both the oxy-gCN structures (2.a/b) the iodine centered C<sub>4</sub>F<sub>9</sub>-I electronic energy level within the conduction band is positioned at  $\approx 4.3$  eV from the VB edge. In the model systems representing reduced gCN, the localized defect state is  $\approx 1.8$  eV from the CB edge and is positioned at  $\approx 2.8$  eV from the iodine-centered electronic energy level in system 3a, whereas the corresponding values are  $\approx 3.0$  eV and  $\approx 3.9$  eV in system 3b.

Interestingly, in the amorphous and reduced systems showing mid-gap states (structures 1.a and 3.a), the wavefunction at the mid-gap defect level is partially localized on C<sub>4</sub>F<sub>9</sub>-I, as shown in Fig. 5. This indicates strong hybridization between the molecule and the defective surface (as also suggested by the ground-state partial electron transfer in Table 2), which is expected to ease a photoinduced electron transfer reaction from the substrate to the molecule. It is also to be noted that the adsorption energies of the C<sub>4</sub>F<sub>9</sub>-I molecule atop am-gCN (1.a) and red-gCN (3.a) surfaces, as well as the electronic structures of the gCN/C<sub>4</sub>F<sub>9</sub>-I systems at the localized mid-gap defect state, remain unaltered irrespective of the size of the replicated gCN surface unit cell (see ESI Fig. SI-9† for details). This further indicates that it is the localized radical nature of the defects on the gCN surface, rather than the density of defects or the closeness of the molecules, that governs the interaction of the gCN surface with the C<sub>4</sub>F<sub>9</sub>-I molecule.

### Photo-physical properties of the gCN/C<sub>4</sub>F<sub>9</sub>-I systems

The excited-state electronic structure of the gCN/C<sub>4</sub>F<sub>9</sub>-I systems is analyzed by considering the representative heptazine trimers (see Fig. SI-2†). The theoretical UV-vis absorption spectra for the gCN/C<sub>4</sub>F<sub>9</sub>-I systems involving the pristine, am-gCN, and oxy-gCN model surfaces are shown in Fig. 7, while those involving the red-gCN models are shown in Fig. SI-14.† The calculated absorption profile of the pristine gCN/C<sub>4</sub>F<sub>9</sub>-I system shows a very intense peak with a maximum at  $\approx 300$  nm and a second band centered at  $\approx 360$  nm, along with a very weak tail extending into the visible domain. This second band can be related to the broad absorption peak in the 350–400 nm region

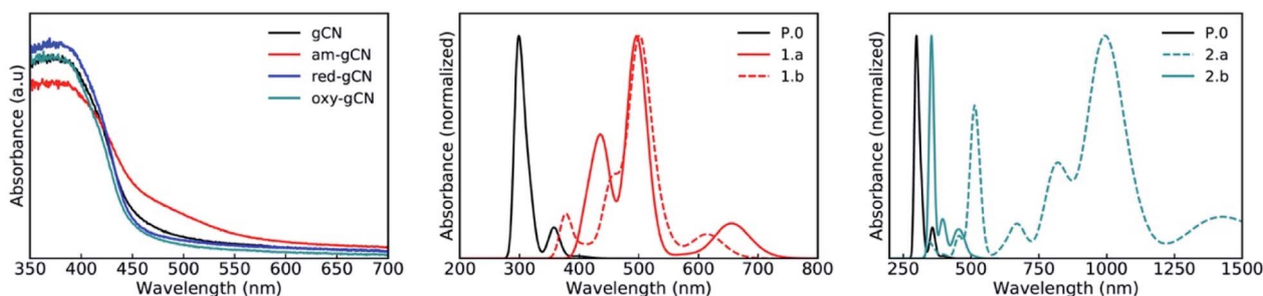


Fig. 7 Experimental (left panel) and theoretical absorption spectra (normalized) of gCN/C<sub>4</sub>F<sub>9</sub>-I systems for the gCN surface in its pristine (black curves) and defective forms; with (middle panel) C<sub>4</sub>F<sub>9</sub>-I on top of the 1.a (full red line) and 1.b (dashed red line) gCN forms and (right) C<sub>4</sub>F<sub>9</sub>-I on top of the 2.a (dashed green line) and 2.b (full green line) gCN forms.



in the experimental spectrum of pristine gCN (see Fig. 7, left panel).

A major change in the calculated optical properties occurs upon the formation of defects in amorphous gCN (structures 1.a and 1.b; Fig. 7, middle panel). Namely, new absorption features appear in the visible domain (an absorption band peak at  $\approx 500$  nm, with additional transitions at higher wavelengths). These changes in the optical absorption predicted by the modelling are in line with the appearance of a shoulder near 500 nm in the experimental absorption spectrum of amorphous gCN. However, it is to be noted that a direct comparison between the calculated and measured intensities should be considered with care, as the defect density considered in the calculations could be different from those present experimentally.

The optical absorption spectra of the oxygen-containing defects are quite contrasting (Fig. 7, right panel): on one hand, the presence of pyran-like oxygenated defects (structure 2.a) on the gCN surface is expected to red-shift the absorption very strongly, up to  $\approx 1000$  nm. On the other hand, the absorption of carbonyl-like oxygenated defects (structure 2.b) shows a peak centered at  $\approx 350$  nm and only a weak response in the visible domain, around 500 nm. The fact that the experimental spectrum of the oxidized gCN shows no intensity at low energy clearly indicates that pyran-like defects are not likely to be present in real samples. Also, the carbonyl-like oxygenated gCN structure is lower in energy by  $\approx 0.6$  eV, indicating that the carbonyl-like form is more stable in comparison. Probably, the low concentration of carbonyl defects and their low expected absorption explain the absence of any new features in the experimental spectrum of oxidized gCN.

To get further insight into the nature of the excited states, we performed the attachment-detachment density (ADD) analysis<sup>41,42</sup> on all the pristine/defective gCN/ $C_4F_9-I$  systems, wherein the detachment density corresponds to the electron density associated with single electronic levels vacated in an electronic transition and the attachment density corresponds to the electron density associated with the single electron levels occupied in the transition. The difference between these densities is called the difference density matrix; it is shown in Fig. 8, wherein the cyan color represents the electronic density of the vacated transition electronic level (detachment density) and the yellow color represents the electronic density of the occupied transition electronic level (attachment density). We also performed the complementary natural transition orbital (NTO) analysis on all the pristine/defective gCN/ $C_4F_9-I$  systems and the representative lowest unoccupied transition orbitals (LUTOs) are shown in the ESI (see Fig. SI-12 to SI-14† for details).

In the case of pristine gCN, the ADD difference corresponding to low-energy excitations ( $\approx 350$  nm) is confined on the semiconductor with both the attachment (in yellow) and detachment (in cyan) transition densities observed only on the surface. Only with excitation at high energy ( $\approx 300$  nm) the transition density partly spreads over the  $C_4F_9-I$  molecule (see Fig. SI-13†). Interestingly, for the models representing the amorphous gCN surface (structures 1.a/b), the ADD difference

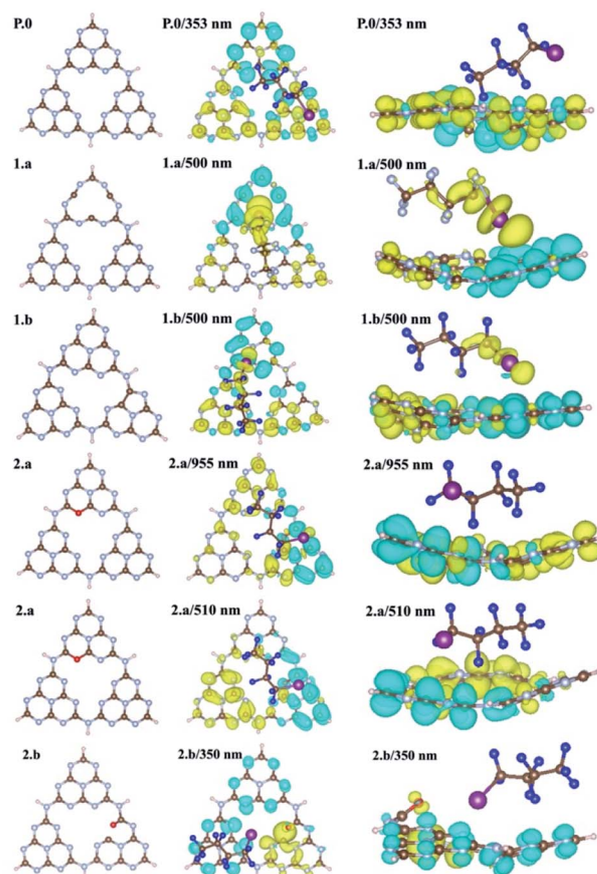


Fig. 8 Top (center) and side (right) views of the attachment-detachment density (ADD) differences of gCN/ $C_4F_9-I$  systems for the gCN surface in its pristine (P.0) and amorphous forms (1.a/b) and pyran-like and carbonyl-like oxygenated forms (2.a/b). Iodine and fluorine atoms of the  $C_4F_9-I$  molecule adsorbed on pristine/defective gCN units are represented with magenta and blue colors, for clarity. The attachment density is represented in yellow and the detachment density in cyan.

calculated at 500 nm now shows major contributions from both the gCN surface and the  $C_4F_9-I$  molecule. While the detachment density is on the am-gCN surface, the attachment density is delocalized on the  $C_4F_9-I$  molecule, in particular on the iodine atom (with a  $d_{z^2}$ -like character). Such transitions are prompted by the molecule-gCN hybridization described above and involve a direct electron excitation from the localized defect state of gCN to the  $C_4F_9-I$  molecule.

In contrast, the oxygenated gCN forms do not show any electronic transition involving the unoccupied  $C_4F_9-I$  molecular orbital as landing levels (Fig. 8, structures 2.a/b) but feature electron transitions occurring exclusively within the gCN surface even for the high-energy excitations ( $\approx 350$  nm), irrespective of the pyran-like or carbonyl-like oxygenated system. In comparison, for models representing the reduced gCN surface (structures 3.a and 3.b), a small delocalization of transition density is observed on the  $C_4F_9-I$  molecule adsorbed on structure 3.a at the lowest energy excitation wavelength of  $\approx 500$  nm, whereas for structure 3.b, the transition density is fully localized only on the gCN surface (see Fig. SI-14† for details).



These results convey the following message: (i) band-edge optical absorption in pristine gCN is not expected to induce an electron transfer to the C<sub>4</sub>F<sub>9</sub>-I molecule; (ii) oxygenated defects on gCN surfaces tend to suppress any photo-induced electron transfer from the gCN surface to C<sub>4</sub>F<sub>9</sub>-I; and (iii) optical absorption in the visible domain for amorphous and reduced gCN forms promotes a photo-induced electron transfer reaction from the semiconductor to the C<sub>4</sub>F<sub>9</sub>-I molecule, thus likely prompting the initial reaction step in the addition of perfluoroalkyl groups to aromatic compounds.

To further corroborate the possibility of electron transfer from the pristine and defective gCN surfaces to the C<sub>4</sub>F<sub>9</sub>-I molecule, we calculated the reaction enthalpies ( $\Delta H$ ) considering a schematic ground-state electron transfer from the pristine/defective gCN surface to the C<sub>4</sub>F<sub>9</sub>-I molecule: C<sub>4</sub>F<sub>9</sub>-I + gCN  $\rightarrow$  C<sub>4</sub>F<sub>9</sub>-I<sup>-</sup> + gCN<sup>+</sup> (see Fig. SI-15<sup>†</sup> for details). Electron transfer to the molecule from the amorphous and reduced gCN forms (structures 1.a and 3.a) is associated with a much lower  $\Delta H$  value of  $\approx 25$  kcal mol<sup>-1</sup>, compared to that of the pristine gCN form ( $\Delta H$  of  $\approx 90$  kcal mol<sup>-1</sup>, ESI Table SI-3<sup>†</sup> for details). Implicitly, it follows that a photo-induced electron transfer from the am-/red-gCN surface to the C<sub>4</sub>F<sub>9</sub>-I molecule is favorable on excitations in the visible domain, consistent with the electronic and excited energy levels shown in Fig. 6 and 8, whereas the electron transfer would happen at much higher energies when the pristine gCN surface is considered.

Very interestingly, upon electron transfer from the gCN surface, the C<sub>4</sub>F<sub>9</sub>-I molecule undergoes an elongation of the C-I bond length by  $\approx 1$  Å, with  $\approx 80\%$  of the charge localized on the iodine atom, while the unpaired electron is strongly localized ( $\approx 65\%$ ) on the neighboring carbon atom (see Fig. SI-16<sup>†</sup>). This is fully consistent with the mechanism proposed for gCN-assisted perfluoroalkylation of electron-rich aromatic compounds, in which the photoexcitation is thought to generate a perfluoroalkyl radical along with the release of iodide.<sup>17</sup> Importantly, the modelling results suggest that the photo-induced electron transfer and C-I bond breaking reactions occur in a concerted way. We further note that the amorphous gCN forms considered in this work show an atomic spin density that is centered on the carbon atoms along with some delocalization on nitrogen atoms, consistent with the EPR results shown in Fig. 4.

These computational results, when compared to the experimental observations in terms of the enhanced photo-catalytic response of am-gCN and red-gCN surfaces, combined with their lower nitrogen contents recorded from the XPS data and higher radical densities observed from the EPR measurements, consistently and strongly imply that specific amorphous/reductive treatments of gCN promotes the formation of more nitrogen vacancies on the gCN surface. These nitrogen vacancies not only favor the adsorption of the C<sub>4</sub>F<sub>9</sub>-I molecule at the defective sites but also promote the photo-induced electron transfer from the radical defect states present on the defective gCN surfaces to the C<sub>4</sub>F<sub>9</sub>-I molecule, which happens at lower excitation energies (at  $\approx 500$  nm), and consequently fully rationalize the superior photo-catalytic response of amorphous and reduced gCN surfaces.

## Conclusions

In conclusion, we employ comprehensive density functional theory (DFT) calculations, using both periodic and molecular approaches, to investigate the nature of radical defects present on the gCN surfaces, in order to provide a detailed structure/photo-physical property relationship between the nature of these defects, their electronic structures, their interactions with a perfluoroalkyl iodide compound in the ground- and excited states and the resulting photo-catalytic efficiencies of the corresponding materials measured experimentally for the perfluoroalkylation of electron-rich organic substrates. It was found that, when radical defect states are created on gCN samples, either by amorphization or reductive or oxidative treatments, electronic defect states appear within the electronic band gap. But only amorphous/reduced gCN forms containing nitrogen vacancies act as strong binding sites for the perfluoroalkyl iodide molecule, *via* partial ground-state charge transfer from these defect states to the substrate molecule and the associated change in bond orders. These defect states are also photo-active in the visible domain: upon photo-excitation electron transfer occurs from these defect states to the C<sub>4</sub>F<sub>9</sub>-I molecule, which is the crucial, initial step in the addition of perfluoroalkyl groups to aromatic compounds. The presence of photo-active defect states within the electronic band gap can therefore directly enhance the resulting photo-catalytic efficiencies of defective gCN samples. This study, therefore, shows the potential of determining the nature and properties of electronic defects in gCN materials for the rational design of gCN-based photocatalysts for other organic substrates.

## Data availability

Data for this paper, including the input and output files of the calculations, are available at the UMONS Institutional Repository (<https://orbi.umons.ac.be>).

## Author contributions

E. R. and S. M. G. performed all the quantum chemical calculations. G. F. synthesized and characterized the experimental gCN samples, under the supervision of M. M. and P. F. A. A. performed the EPR measurements under the supervision of M. C. M. B. performed the EIS measurements. M. P., D. B., and R. L., conceived and designed the study. All authors discussed the results and contributed to the interpretation of the data as well as to the writing and editing of the manuscript.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

This work was supported by the Energy Transition Fund of the Belgian Federal Government (FPS Economy) within the T-REX project, the Belgian National Fund for Scientific Research



(FRS-FNRS) within the Consortium des Équipements de Calcul Intensif – CÉCI, under Grant 2.5020.11, and by the Walloon Region (ZENOBE Tier-1 supercomputer, under grant 1117545). E. R. is grateful to the University of Trieste and the University of Mons for a joint doctoral fellowship. M. M. and G. F. kindly acknowledge FRA2021 funded by the University of Trieste. G. F. also acknowledges Microgrants 2021 funded by Region FVG (LR 2/2011, ART. 4). This work was supported by the University of Trieste, INSTM, and the Italian Ministry of Education MIUR (cofin Prot. 2017PBXPN4). Part of this work was performed under the Maria de Maeztu Units of Excellence Program from the Spanish State Research Agency Grant No. MDM-2017-0720. We acknowledge Prof. E. Salvadori for useful discussions on the EPR results and Dr G. Londi for his assistance in the attachment/detachment density analysis. D. B. is Research Director of FNRS. M. P. is the AXA Chair for Bionanotechnology (2016–2023).

## Notes and references

- C. Rosso, G. Filippini, A. Criado, M. Melchionna, P. Fornasiero and M. Prato, *ACS Nano*, 2021, **15**, 3621–3630.
- X. Wang, K. Maeda, A. Thomas, K. Takane, G. Xin, J. M. Carlsson, K. Domen and M. Antonietti, *Nat. Mater.*, 2009, **8**, 76–80.
- Y. Wang, X. Wang and M. Antonietti, *Angew. Chem., Int. Ed.*, 2012, **51**, 68–89.
- J. Zhang, Y. Chen and X. Wang, *Energy Environ. Sci.*, 2015, **8**, 3092–3108.
- S. Cao, J. Low, J. Yu and M. Jaroniec, *Adv. Mater.*, 2015, **27**, 2150–2176.
- J. Liu, H. Wang and M. Antonietti, *Chem. Soc. Rev.*, 2016, **45**, 2308–2326.
- X. Ye, Y. Cui, X. Qiu and X. Wang, *Appl. Catal., B*, 2014, **152–153**, 383–389.
- X. Chen, J. Zhang, X. Fu, M. Antonietti and X. Wang, *J. Am. Chem. Soc.*, 2009, **131**, 11658–11659.
- L. Möhlmann, M. Baar, J. Rieß, M. Antonietti, X. Wang and S. Blechert, *Adv. Synth. Catal.*, 2012, **354**, 1909–1913.
- C. Cavedon, A. Madani, P. H. Seeberger and B. Pieber, *Org. Lett.*, 2019, **21**, 5331–5334.
- B. Pieber, J. A. Malik, C. Cavedon, S. Gisbertz, A. Savateev, D. Cruz, T. Heil, G. Zhang and P. H. Seeberger, *Angew. Chem., Int. Ed.*, 2019, **58**, 9575–9580.
- I. Ghosh, J. Khamrai, A. Savateev, N. Shlapakov, M. Antonietti and B. König, *Science*, 2019, **365**, 360–366.
- K. Müller, C. Faeh and F. Diederich, *Science*, 2007, **317**, 1881–1886.
- Y. Zhou, J. Wang, Z. Gu, S. Wang, W. Zhu, J. L. Aceña, V. A. Soloshonok, K. Izawa and H. Liu, *Chem. Rev.*, 2016, **116**, 422–518.
- A. Savateev, I. Ghosh, B. König and M. Antonietti, *Angew. Chem., Int. Ed.*, 2018, **57**, 15936–15947.
- Y. Zhang, A. Thomas, M. Antonietti and X. Wang, *J. Am. Chem. Soc.*, 2009, **131**, 50–51.
- H. Pan, Y.-W. Zhang, V. B. Shenoy and H. Gao, *ACS Catal.*, 2011, **1**, 99–104.
- X. Wang, X. Chen, A. Thomas, X. Fu and M. Antonietti, *Adv. Mater.*, 2009, **21**, 1609–1612.
- C. Wang, Q. Wan, J. Cheng, S. Lin, A. Savateev, M. Antonietti and X. Wang, *J. Catal.*, 2021, **393**, 116–125.
- C. Wang, Y. Hou, J. Cheng, M.-J. Lin and X. Wang, *Appl. Catal., B*, 2021, **294**, 120259.
- G. Filippini, F. Longobardo, L. Forster, A. Criado, G. Di Carmine, L. Nasi, C. D'Agostino, M. Melchionna, P. Fornasiero and M. Prato, *Sci. Adv.*, 2020, **6**, eabc9923.
- R. Berger, G. Resnati, P. Metrangolo, E. Weber and J. Hulliger, *Chem. Soc. Rev.*, 2011, **40**, 3496.
- C.-J. Wallentin, J. D. Nguyen, P. Finkbeiner and C. R. J. Stephenson, *J. Am. Chem. Soc.*, 2012, **134**, 8875–8884.
- S. Barata-Vallejo, M. V. Cooke and A. Postigo, *ACS Catal.*, 2018, **8**, 7287–7307.
- N. Robinson, C. Robertson, L. F. Gladden, S. J. Jenkins and C. D'Agostino, *ChemPhysChem*, 2018, **19**, 2472–2479.
- P. Politzer, P. Lane, M. C. Concha, Y. Ma and J. S. Murray, *J. Mol. Model.*, 2007, **13**, 305–311.
- P. Niu, L.-C. Yin, Y.-Q. Yang, G. Liu and H.-M. Cheng, *Adv. Mater.*, 2014, **26**, 8046–8052.
- Y. Kang, Y. Yang, L.-C. Yin, X. Kang, G. Liu and H.-M. Cheng, *Adv. Mater.*, 2015, **27**, 4572–4577.
- F. Longobardo, G. Gentile, A. Criado, A. Actis, S. Colussi, V. Dal Santo, M. Chiesa, G. Filippini, P. Fornasiero, M. Prato and M. Melchionna, *Mater. Chem. Front.*, 2021, **5**, 7267–7275.
- T. Suter, V. Brázdová, K. McColl, T. S. Miller, H. Nagashima, E. Salvadori, A. Sella, C. A. Howard, C. W. M. Kay, F. Corà and P. F. McMillan, *J. Phys. Chem. C*, 2018, **122**, 25183–25194.
- A. Ugolotti and C. Di Valentin, *Nanomaterials*, 2021, **11**, 1863.
- X. Song, X. Li, X. Zhang, Y. Wu, C. Ma, P. Huo and Y. Yan, *Appl. Catal., B*, 2020, **268**, 118736.
- G. Kresse and J. Furthmüller, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1996, **54**, 11169–11186.
- G. Kresse and D. Joubert, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1999, **59**, 1758–1775.
- J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, **77**, 3865–3868.
- S. Grimme, *J. Comput. Chem.*, 2006, **27**, 1787–1799.
- A. V. Krukau, O. A. Vydrov, A. F. Izmaylov and G. E. Scuseria, *J. Chem. Phys.*, 2006, **125**, 224106.
- N. G. Limas and T. A. Manz, *RSC Adv.*, 2018, **8**, 2678–2707.
- M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. a. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. a. Petersson, H. Nakatsuji, X. Li, M. Caricato, a. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, a. F. Izmaylov, J. L. Sonnenberg, D.-Y. Williams, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. a. Montgomery Jr, J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. a. Keith, R. Kobayashi, J. Normand,



- K. Raghavachari, a. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, 2016, *Gaussian 16, Revision C.01*, Gaussian, Inc., Wallin.
- 40 R. L. Martin, *J. Chem. Phys.*, 2003, **118**, 4775–4777.
- 41 T. Etienne, X. Assfeld and A. Monari, *J. Chem. Theory Comput.*, 2014, **10**, 3906–3914.
- 42 A. J. Gillett, C. Tonnelé, G. Londi, G. Ricci, M. Catherin, D. M. L. Unson, D. Casanova, F. Castet, Y. Olivier, W. M. Chen, E. Zaborova, E. W. Evans, B. H. Drummond, P. J. Conaghan, L.-S. Cui, N. C. Greenham, Y. Puttisong, F. Fages, D. Beljonne and R. H. Friend, *Nat. Commun.*, 2021, **12**, 6640.
- 43 O. Faye, U. Eduok and J. A. Szpunar, *J. Phys. Chem. C*, 2019, **123**, 29513–29523.
- 44 F. K. Kessler, Y. Zheng, D. Schwarz, C. Merschjann, W. Schnick, X. Wang and M. J. Bojdys, *Nat. Rev. Mater.*, 2017, **2**, 17030.

