

Local perturbations of periodic systems. Chemisorption and point defects by GoGreenGo

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Abstract

We present a software package GoGREENGO—an overlay aimed to model local perturbations of periodic systems due to either chemisorption or point defects. The electronic structure of an ideal crystal is obtained by worldwide-distributed standard quantum physics/chemistry codes, and then processed by various tools performing projection to atomic orbital basis sets. Starting from this, the perturbation is addressed by GoGREENGO with use of the Green's functions formalism, which allows evaluating its effect on the electronic structure, density matrix, and energy of the system. In the present contribution, the main accent is made on processes of chemical nature, such as chemisorption or doping. We address a general theory and its computational implementation supported by a series of test calculations of the electronic structure perturbations for benchmark model solids: simple, face-centered, and body-centered cubium systems. In addition, more realistic problems of local perturbations in graphene lattice, such as lattice substitution, vacancy, and "on-top" chemisorption, are considered.

KEYWORDS

chemisorption, electronic structure, Green's functions, point defects in crystals, substitutional defects

1 | INTRODUCTION

A great number of elementary processes of significant importance in solid-state chemistry are local in nature,¹ although they take place in a "matrix" of crystal (periodic) system. Currently, local processes/effects in solids are addressed by using standard computational chemistry codes pushing towards the limits of their applicability with a considerable loss of efficiency. Indeed, applying "molecular" codes to study a point feature in a crystal requires tricky setting up of boundary conditions in the cluster models of eventually infinite systems.² Any way, one has to consider rather large clusters to compensate boundary effects. At the same time, the required embedding procedures are sometimes awkward and not unique, particularly in the case of clusters cut from ionic crystals or metals. A comprehensive review of the related problems is given in Ref. [3].

An alternative and nowadays predominant approach is to accommodate periodic wave function⁴ or $PAW-DFT^5$ based methods to

study local effects. In order to do so, one needs to increase the size of the unit cell to ensure correct proportion of the point defects and to minimize lateral interactions between them (see, for example, References [6, 7] and references therein). The unit cell enlargement increases the computational costs drastically with the scaling up to $O(N^3)$, where N is the "size" of the unit cell. The entire study consists of full calculations of the original (ideal) periodic system and repeating them for as well periodic models which include intrinsically local features. The effect of the local perturbations is then derived by comparing results of these calculations.

Apparently, the state of the locally perturbed crystal cannot differ too much from that of the ideal one. The mentioned standard approaches do not consider this fact, although it seems to be profitable to avoid repetitive recalculations from the scratch and to make a better use of the eventually more precise information about the electronic structure of the ideal crystal. Based on this idea, further embedding schemes had been developed (for example References [8–11]) which employ the information about the ideal crystal for analysis of the defect ones. In addition, these schemes are not free from numerous ad hoc assumptions leading to an uncontrollable loss of precision and predictive capacity. Embedding methods based on the optimized effective potential method^{8,9} require density inversion procedures to determine non-additive kinetic potential term, which can be ill conditioned.¹¹ Projection-based embedding schemes, employing the idea of localized molecular orbitals in different manifestations,^{10,11} substantially rely on the degree of localization in the system: the quality of calculations decreases significantly for the systems with intrinsically delocalized electronic system. Because of that, embedding in metals presents a considerable challenge for the projection-based methods, and the problem seems to be conceptual, since it is hard to expect, that metals will allow any required localization of one-electronic states. Therefore, the projection-based embedding can hardly be considered as a universal approach to calculate electronic structure of local defects in all solids. At the same time, one can expect them to be a very useful tool to study local defects in insulators.

Remarkably, a detailed theory of local perturbations of periodic systems using the Green's function (GF) formalism had been developed yet in 60s¹²⁻¹⁵ (based on an earlier work¹⁶) and was applied to chemisorption in numerous works.¹⁷⁻²⁸ The most characteristic feature of this approach is that starting from a solution of the ideal periodic problem and from its local perturbation, one obtains the answer as a correction to the unperturbed solution. By this, (1) the highly inefficient step of solving the perturbed problem from the scratch is avoided and (2) the result is a pure effect of the perturbation. The theory is highly pedagogically explained in Reference [29], where it is used to provide pictorial description of the electronic structure perturbations occurring throughout chemisorption. Otherwise more mathematical, but still suitable for a chemistry theory student exposition of the required apparatus is given in Ref. [30].

Being actively used in 70ies to study local defects in rather simple model solids, the GF theory was practically abandoned later and replaced by supercell periodic and finite clusters calculations.³¹ Undeservedly forgotten, this approach has been getting much less attention since then, especially in the community of solid-state chemistry.* Although, it suits better for discussing chemical problems, it has never enjoyed any generalization to more realistic systems described by rigorously defined Hamiltonians. As well, there is no reported software implementing the general theory, which would be suitable for a broad range of solid state and surface chemistry problems (with different kinds of defects in crystals and/or chemisorption). The purpose of the present work, and of the proposed GoGREENGO software package, is to incorporate the mentioned approach into a context of modern theoretical methodology used in solid-state chemistry. Since we mainly focus on the chemical interpretation of electronic structure calculations (described in "chemical" terms such as atomic charges or bond orders), it is implemented for basis of local atomic orbitals. In the present contribution we are concentrated on perturbations of the electronic structure caused by the local defects of different kinds, while structural deformations, which may arise in the defected crystal, are going to be addressed in future works.

CHEMISTRY WILEY 2353

This paper is organized as follows. In the Theory and Implementation Sections we sketch the generalized theory and implementation details of the programs included in the GoGREENGO package. After that we provide test results for various benchmarks by considering substitutional and interstitial defects in the model cubic lattices. Finally, GoGREENGO package is applied to tackle more realistic problems devoted to graphene. One can find more detailed specifications of the GoGREENGO by following the link.³³ The Supporting information collects details of the theory and analytical results used for the testing.

2 | THEORY SKETCH

2.1 | Structure of the electronic problem

Theoretical basis of the proposed software development is the selfconsistent perturbation theory of many-electron systems.³⁴ It starts from representing the electronic structure either by a single Slater determinant formed by one-electron (spin-)orbitals for the wave function based methods (as exemplified by References [4,35,36]) or by one-electron density in the DFT context. In the latter case oneelectron orbitals reappear as Kohn-Sham ones, so that either wavefunction or (Kohn-Sham) DFT procedures can be represented as an iterative eigenvalue-eigenvector problem with some generalized density dependent Fockian matrix $\mathbf{F}[\mathbf{P}]$ in the functional space spanned either by exclusively atomic states or by an assembly of plane waves and atomic local states augmenting (PAW) the former:

$$(\varepsilon \mathbf{I} - \mathbf{F}[\mathbf{P}])|\lambda\rangle = \mathbf{0}.$$
 (1)

Solutions of Equation (1) are the eigenvalue-eigenvector pairs ε_{λ} , $|\lambda\rangle$ numbered by assemblies of quantum numbers λ and satisfying the well-known relations:

$$\mathbf{F}[\mathbf{P}]|\lambda\rangle = \varepsilon_{\lambda}|\lambda\rangle. \tag{2}$$

The density **P** is determined by the occupied eigenvectors $|\lambda\rangle$ whose eigenvalues are subject to the condition $\varepsilon_{\lambda} \leq \varepsilon_{\rm F}$, where $\varepsilon_{\rm F}$ is the Fermi energy selected so that the number of occupied one-electron states equals to that of electrons (Fermi statistics). The eigenvalue-eigenvector problems Equations (1), (2) are sequentially solved until the convergence for **P** is achieved.

2.2 | Green's function's representation

The eigenvalue-eigenvector problem can be alternatively formulated in terms of the quantity[†]:

$$G(z) = (zI - F)^{-1},$$
 (3)

the famous Green's function of a complex argument z. Since the Fockian F (hereinafter, we omit its P dependence for brevity) is a

2354

WILEY -

Hermitian operator, its eigenvalues ε_{λ} are always real. Thus, for an arbitrary complex z unequal to any of ε_{λ} the matrix (zI - F) is nondegenerate and can be inverted producing a z-dependent quantity Equation (3). Its closest relation to the eigenvalue-eigenvector problem stems from the spectral representation:

$$\mathbf{G}(z) = \sum_{\lambda} \frac{|\lambda\rangle \langle \lambda|}{z - \varepsilon_{\lambda}},\tag{4}$$

which immediately derives from the expressions for the identity matrix and the Fockian in the basis of its eigenvectors:

$$\mathbf{I} = \sum_{\lambda} |\lambda\rangle \langle \lambda|; \mathbf{F} = \sum_{\lambda} \varepsilon_{\lambda} |\lambda\rangle \langle \lambda|.$$

As soon as the GF is known in the basis of the eigenvalues of **F**, where it is diagonal, it is known in whatever basis. E.g. in the basis of local atomic spin-orbitals $|a\rangle$, $|b\rangle$,... a Greenian matrix is formed by the elements

$$G_{ab}(z) = \sum_{\lambda} \frac{\langle a | \lambda \rangle \langle \lambda | b \rangle}{z - \varepsilon_{\lambda}}, \qquad (5)$$

where $\langle a|\lambda\rangle$ are expansion coefficients of the eigenvector $|\lambda\rangle$ over atomic basis.[‡]

Being defined as a function of complex variable, GF appears in the expressions for the physical quantities under the integral over the real axis only. Since GF has many poles on the real axis, it should be considered there as a distribution (or a generalized function). Such distribution is defined as a limit:

$$\mathbf{G}(\varepsilon) = \lim_{\nu \to 0^+} \mathbf{G}(\varepsilon + i\nu), \tag{6}$$

where z is set to be $\varepsilon + i\nu$ with real ε and ν . As described in Supporting information section 1, evaluation of the limit entering the later equation leads to the following GF on the real axis

$$\mathbf{G}(\varepsilon) = \Re \mathbf{G}(\varepsilon) + i\Im \mathbf{G}(\varepsilon),$$

$$\Re \mathbf{G}(\varepsilon) = \sum_{\lambda} |\lambda\rangle \langle \lambda | \mathcal{P}\left(\frac{1}{\varepsilon - \varepsilon_{\lambda}}\right),$$

$$\Im \mathbf{G}(\varepsilon) = -\pi \sum_{\lambda} |\lambda\rangle \langle \lambda | \delta(\varepsilon - \varepsilon_{\lambda}),$$

(7)

where $\delta(\varepsilon)$ is the Dirac δ -"function" and \mathcal{P} indicates that the integral of a function $f(\varepsilon)$, multiplied by $(\varepsilon - \varepsilon_{\lambda})^{-1}$, with respect to ε must be understood as the Cauchy principal value. In terms of the GF of a real argument the general expression for the density operator (Equation (3) of Supporting information) takes the form:

$$\mathbf{P} = -\pi^{-1} \int_{-\infty}^{\varepsilon_{\rm F}} \Im \mathbf{G}(\varepsilon) d\varepsilon. \tag{8}$$

Perturbations in terms of Green's function

The GF formalism as sketched in Section 2.2 does not add too much to the usual treatment of the eigenvalue-eigenvector problem. Its power manifests itself when the perturbations are addressed. For the GF of the Fockian F, being a sum of an unperturbed one $\mathbf{F}^{(0)}$ and a perturbation \mathbf{F}' :

 $F = F^{(0)} + F'$,

the Dyson equation

2.3

$$\mathbf{G}(z) = \mathbf{G}^{(0)}(z) + \mathbf{G}^{(0)}(z)\mathbf{F}'\mathbf{G}(z), \tag{9}$$

holds³⁸ valid in the whole complex plane. Being solved for $\mathbf{G}(z)$, it gives:

$$\mathbf{G}(z) = \left(\mathbf{I} - \mathbf{G}^{(0)}(z)\mathbf{F}'\right)^{-1}\mathbf{G}^{(0)}(z),$$
(10)

which generates the perturbation series if one expands the inverse matrix in the geometric series:

$$\begin{split} & \left(\mathbf{I} - \mathbf{G}^{(0)}(z)\mathbf{F}'\right)^{-1} = \mathbf{I} + \mathbf{G}^{(0)}(z)\mathbf{F}' + \mathbf{G}^{(0)}(z)\mathbf{F}'\mathbf{G}^{(0)}(z)\mathbf{F}' + \dots \\ & \mathbf{G}(z) = \mathbf{G}^{(0)}(z) + \mathbf{G}^{(0)}(z)\mathbf{F}'\mathbf{G}^{(0)}(z) + \mathbf{G}^{(0)}(z)\mathbf{F}'\mathbf{G}^{(0)}(z) + \dots \\ \end{split}$$

Thus, the general perturbative treatment rewrites in terms of the Green's functions. Formally, in the case of a point defect in an infinite crystal the solution of the Dyson equation would require inversion of a matrix of infinite dimension. However, switching to a local atomic orbital representation allows to reduce the problem to a finite one, since in this case a point (local) perturbation acts on a relatively low-dimensional subspace (*P*) of the entire space of one-electronic states. In GOGREENGO we employ this possibility and consider the perturbation matrices of a form:

$$\mathbf{F}' = \begin{pmatrix} \mathbf{F}'_{pp} & 0\\ 0 & 0 \end{pmatrix} = \begin{pmatrix} \mathbf{V} & 0\\ 0 & 0 \end{pmatrix}.$$
 (11)

The Greenian matrix is then split in blocks:

$$\mathbf{G}^{(0)} = \begin{pmatrix} \mathbf{G}_{PP}^{(0)} & \mathbf{G}_{PQ}^{(0)} \\ \mathbf{G}_{QP}^{(0)} & \mathbf{G}_{QQ}^{(0)} \end{pmatrix},$$
(12)

where Q refers to the orthogonal complement of the subspace P (argument z is omitted for brevity). Introducing a dimP × dimP matrix $M^{\$}$:

$$\mathbf{M} = \mathbf{V} \left(\mathbf{I}_{PP} - \mathbf{G}_{PP}^{(0)} \mathbf{V} \right)^{-1}, \tag{13}$$

and following the procedure given in Supporting information section 2, one obtains the corrections to the matrix blocks:

$$\begin{split} \mathbf{G}_{PP} &- \mathbf{G}_{PP}^{(0)} = \mathbf{G}_{PP}^{(0)} \, \mathbf{M} \mathbf{G}_{PP}^{(0)}, \\ \mathbf{G}_{QQ} &- \mathbf{G}_{QQ}^{(0)} = \mathbf{G}_{QP}^{(0)} \, \mathbf{M} \mathbf{G}_{PQ}^{(0)}, \\ \mathbf{G}_{PQ} &- \mathbf{G}_{PQ}^{(0)} = \mathbf{G}_{PP}^{(0)} \, \mathbf{M} \mathbf{G}_{PQ}^{(0)}, \ \mathbf{G}_{QP} - \mathbf{G}_{QP}^{(0)} = \mathbf{G}_{QP}^{(0)} \, \mathbf{M} \mathbf{G}_{PP}^{(0)}, \end{split} \tag{14}$$

which, respectively, express the effect of the perturbation on the Green's function in the subspace P itself, in the subspace where the perturbation is absent (Q) and on the coupling between the perturbed and unperturbed subspaces. Density matrix elements of the perturbed system are calculated with Equation (8) once the Dyson equation is solved. If one takes into account the density dependence of the Fockian, the calculated density serves as an input for a next step of iterative solution. This setting is referred below as the self-consistent one.

2.4 Specific of the problems at hand

So far reviewed greenistic representation of the eigenvalueeigenvector problem is fairly general. Here, we apply it to the crystal point defects of different kinds. We consider two types of such defects: (1) point-wise perturbation of the crystalline matrix itself—the lattice substitution and the vacancy; and (2) interaction of the crystal with an extra "particle"—dubbed in different contexts as an interstitial defect or an adsorbate. In either case we shall employ the generic term defect. The unperturbed solution described by GF $G^{(0)}$ of the ideal crystal (and of the finite adsorbate in the case (2)) is assumed to be known. Below we briefly review the specific of the greeninstic formalism as applied to infinite periodic systems with point defects.

2.4.1 | Green's functions of crystals

When it comes to solids, the solution of the eigenvalue-eigenvector problem has specifics described in handbooks on solid-state physics and chemistry³⁹ (see chapters devoted to tight binding approximation). Due to translation invariance of an ideal infinite crystal, its Fockian accepts the block-diagonal form in the basis of the Bloch sums:

$$|ak\rangle = \frac{1}{\sqrt{K}} \sum_{r} \exp(ikr) |ar\rangle$$

of A atomic states. Here A is the number of atomic spin-orbitals $a = 1 \div A$, *r* stands for a unit cell index and K is the number of *k*-points involved in the calculation.^{||} The blocks are numbered by the wave vectors *k* from the first Brillouin zone and respective eigenvalues form A functions ε_{ak} of k ($\alpha = 1 \div A$)–(electronic) bands with the eigenvector expansion coefficients forming *k*-dependent $A \times A$ matrices with elements $\langle \alpha k | ak \rangle$. In this context, the generalized quantum number λ , labeling the eigenvalues of the Fockian, splits in the pair of the wave vector *k* and the band index α : $\lambda = (\alpha, k)$. Thus, the GF of the ideal crystal with A bands reads:

HEMISTRY

$$\mathbf{G}(z) = \sum_{\alpha k} \frac{|\alpha k\rangle \langle \alpha k|}{z - \varepsilon_{\alpha k}}.$$
(15)

It is as well block-diagonal with $A \times A$ blocks numbered by k. A remarkable feature of the "theoretical" GF of the crystal is that the poles coalesce in (quasi-) continuous segments being the allowed energy bands of electrons (see Reference [39]). Considering the GF on the real axis according to Equation (7) produces the electronic density of states (DOS):

$$DOS(\varepsilon) = -\frac{1}{\pi K} Sp \Im \mathbf{G}(\varepsilon) = \sum_{ak} \delta(\varepsilon - \varepsilon_{ak}), \qquad (16)$$

familiar to the workers of the field.

Applying the same trick to the diagonal elements of the Greenian matrix Equation (5) in the basis of local atomic orbitals

$$\mathsf{G}_{aa}(\mathsf{z}) = \sum_{\alpha k} \frac{\langle ak | \alpha k \rangle \langle \alpha k | ak \rangle}{\mathsf{z} - \varepsilon_{\alpha k}},$$

we arrive to

$$\mathsf{DOS}_{a}(\varepsilon) = -\frac{1}{\pi \mathsf{K}} \Im \mathsf{G}_{aa}(\varepsilon) = \sum_{ak} \langle ak | ak \rangle \langle ak | ak \rangle \delta(\varepsilon - \varepsilon_{ak})$$

- the projection of the DOS upon the atomic state *a*, as well, familiar from numerous packages (e.g. LOBSTER⁴¹ or WANNIER90⁴²) performing analysis of numerical data derived from PAW-DFT or whatever computer experiments on solids (see e.g.^{41,43}). Treating similarly off-diagonal elements:

$$G_{ar,br'}(z) = \sum_{ak} \frac{\langle ak | ak \rangle \langle ak | bk \rangle \exp(ik(r-r'))}{z - \varepsilon_{ak}}$$

yields:

$$-\frac{1}{\pi K}\Im G_{ar,br'}(\varepsilon) = \sum_{\alpha k} \Re[\langle ak | \alpha k \rangle \langle \alpha k | bk \rangle \exp(ik(r-r'))]\delta(\varepsilon - \varepsilon_{\alpha k}) \quad (17)$$

which is a close relative of the crystal orbital overlap and crystal orbital Hamilton populations (respectively, COOP and COHP⁴⁴) as well widely available in the solid-state packages. In these expressions $|ar\rangle$ corresponds to the AO *a* in the unit cell *r*, so, the latter formula allows to calculate the density matrix element for the pair of orbitals from different unit cells.

2.4.2 | Green's functions of a finite system

The definition of the GF and all related quantities in the case of the finite system remains the same as sketched in the Subsection devoted to the GF representation of eigenstates problem. The Greenian matrix element has a general form Equation (5), which for the real values of argument reduces to:

$$G_{ab}(\varepsilon) = \sum_{\lambda} \left[\frac{\langle a | \lambda \rangle \langle \lambda | b \rangle}{\varepsilon - \varepsilon_{\lambda}} - i\pi \langle a | \lambda \rangle \langle \lambda | b \rangle \delta(\varepsilon - \varepsilon_{\lambda}) \right],$$
(18)

where the summation goes over discrete levels λ . The imaginary part of the GF consists of discrete signals located at energies ε_{i} and proportional to the Dirac's δ -"function" and has the dimension of inverse energy, which becomes apparent from the Lorentzian approximation of the δ -function. The real part is a continuous function except simple poles at $\varepsilon = \varepsilon_{\lambda}$.

In the general case it is not possible to find an analytical solution for the perturbed GF, therefore one has to treat the initial Greenian matrix of the finite system numerically. Such treatment requires approximating the δ -function in Equation (18) by a Lorentzian of a (small) width ν as explained in Supporting information section 1. Theoretically, such approximation approaches genuine result in the limit $\nu \rightarrow 0$. In practical calculations the value of this parameter has to be finite. It is advised to set ν equal to the step of the energy grid and to choose the latter small enough to guarantee required accuracy.

IMPLEMENTATION AND 3 COMPUTATIONAL DETAILS

3.1 General construction

The programmatic implementation of the theory described in the Theory Section is provided by GoGREENGO package being a set of procedures written in fortran 2010 using the system of high-level objects provided by the cartesius fort library.⁴⁵ The component procedures exchange data through intermediate files packed in the hdf5 archive.⁴⁶ The flow-chart illustrating the relations between different programs of the package and the paths of data transfer between them is shown on the Figure 1. Specifically,

- 1. program GET_GREEN evaluates Greenian matrix elements for a given set of local atomic orbitals from the band structure of the ideal crystal. The program can use eigenvalues derived by VASP,47 ABINIT.48 The projections of eigenvectors produced by these packages to the basis of local orbitals can be obtained by the lobster software⁴¹ whose output format is compatible with get_green. As well $\Theta \Phi$ (TetaPhi)⁴⁹ produces eigenvalues-eigenvectors in the format readable by GoGREENGO. This combination allows extracting Greenian matrix of the pure solid in an atomic basis from the most popular plane wave DFT codes and using it for impurity calculations. One can use the band structure of the solid from any other source, as well, transforming it to the required format. Imaginary parts of G_{ab} are calculated in get_green for each point of the energy grid by applying tetrahedron algorithm⁵⁰ of integration over the Brillouin zone. Since the real parts of GF elements are related with the imaginary parts by the Kramers-Kronig relations⁵¹ they are obtained numerically from the latter as described in Reference [52]. The user has to define the interval of the energy, where the Greenian matrix has to be calculated, and the step of the energy grid. In addition, there is an option to obtain elements of the Greenian matrix between different unit cells r and r' as in Equation (17) with no calculations on extended unit cells.
- 2. HGEN calculates the crystal-defect hopping operator and an array of two-electron Coulomb integrals (required for the self-consistent setting). Using hgen is not mandatory: one can also use any externally prepared hopping operator and Coulomb integrals in the required format. Hgen calculates the interaction within a semiempirical NDDO approximation including different parameterizations such as MNDO,53 AM154 and PM355 which have been recently shown to be compatible with the PAW-DFT setting.56 Other options of calculating atomic integrals will be added in the future releases. To run the hgen utility one has to provide geometries of the defect and crystal, define which atoms interact (by giving a cut-off distance or listing them explicitly) and type of



the AOs basis set to be used (single STO, MAP,⁵⁷ Bunge,⁵⁸ Koga⁵⁹ are available so far). Optionally, one can also change default values of semi-empirical hopping parameters in order to parameterize the Hamiltonian for one's needs.

- 3. Program DYSON reads unperturbed Greenian matrix of the ideal crystal (which comes from GET_GREEN), that of a defect (also comes from GET_GREEN or can be calculated directly in dyson if one provides molecular orbital energies and MO LCAO coefficients), initial perturbation operator and a table of two-electronic integrals, if the self-consistent version is used. The program finds a solution of the Dyson equation and returns perturbed Greenian matrix together with the new density matrix and energy correction caused by perturbation. In the case of the self-consistent setting it performs iterative solution of the Dyson's equation taking into account an adjustment of the self-energy at each step. In this case the user has to specify a convergence threshold. The damping ensuring better convergence is supported and can be used if necessary.
- 4. Program GREEN_OPT performs either gradient or simplex optimization of the defect position and (internal) geometry. Hgen can be used to generate perturbation matrix on each step. In hgen no gradients are available so far, so it can only be used for the simplex optimization. Gradients will be added to hgen in future versions. Regarding geometry optimization we note, that strong local perturbations may cause considerable deformations of the crystal geometry, having significant impact on the electronic structure and energetics of the defect formation. Although, in the present paper we only focus on perturbations of the electronic structure, typically, local defects cause local deformations, which formally translate into additional terms in the perturbation operator \mathbf{F}' . Their impact on the electronic structure can be treated by the proposed approach as of any other perturbation. Of course, consistent evaluation of the atomic forces and search for a deformation, minimizing the total energy, requires special attention. Authors plan to address this issue in future works by introducing electron-phonon interactions into the model.

4 | SOME SPECIFIC FEATURES

4.1 | Self-consistent perturbation theory

As mentioned in the Theory Section, the perturbation of a oneelectron part of the Fockian produces a correction to the density matrix of the same order as the perturbation itself. Then, due to the mean field treatment of the electron-electron interaction either in the wave function or DFT setting, the corresponding Fockian receives further corrections proportional to those of the density matrix elements. They form so called "dressing" which needs to be added to the original one-electron ("bare") perturbation. The solution of the resulting Dyson equation yields further changes of density matrix so that one has to repeat the calculation until the convergence is reached. This option is implemented in the GOGREENGO package.

In metals the effect of perturbation decays with the distance from the defect as $R^{-2\kappa}$ with some $\kappa > 0$ dependent on the form of the Fermi surface and dimensionality of the crystal structure (κ is typically CHEMISTRY WILEY 2357

higher for higher dimension).^{28**} This allows one to restrict the range of the action of the self-consistent (dressed) perturbation by a finite number of unit cells close to the defect. Of course, the corrections to the two-electron part of the Fockian involve additional orbitals, and the dimension of the subspace *P* has to be increased. The amount of this augmentation is system dependent and it is advised to look for an optimal size by a series of convergence tests.

4.1.1 | Position of the Fermi level

Another important aspect is the position of the Fermi level in the perturbed system. In general, it does not remain constant although changes by a small value. The general reason is that the perturbed GF $\mathbf{G}(\varepsilon)$ differs from unperturbed one $\mathbf{G}^{(0)}(\varepsilon)$ and thus the integral of $\mathsf{Sp}\mathbf{G}(\varepsilon)$ from $-\infty$ up to ε_{F} of the unperturbed crystal not necessarily yields the same number of electrons as does the integration of $Sp\mathbf{G}^{(0)}(z)$. The defect of electron's number Δn is to be eliminated by shifting the Fermi energy by $\delta \varepsilon_{\rm F}$. As it is explained in Supporting information section 3 the value of $\delta \varepsilon_{\rm F}$ is determined by the value of $\Delta n/{\rm K}$ and it, evidently, becomes infinitesimally small for the limit $K \rightarrow \infty$. In real calculations K is finite and $\delta \varepsilon_{\rm F}$ is non-vanishing in this case. Evaluation of $\delta \varepsilon_{\rm F}$ required to keep constant the number of electrons in the system with K unit cells is implemented in the GoGREENGO. Practically, in most our calculations the value of $\Delta n/K$ is rather small (we use $31 \times 31 \times 31$ meshes of k-points for three-dimensional models and 51×51 one for graphene) and affects density matrix elements only in fourth-fifth decimal place. However, in all cases it is advised to thoroughly check its impact on the density matrix elements and final results, especially if the number of k-points K used for the band structure calculations of the crystal is relatively small as it sometimes happens in PAW-DFT calculations.

4.1.2 | Energy correction

Even an infinitesimally small shift of the Fermi level causes a finite correction to the perturbation energy, since a summation over infinite number of unit cells is implied in calculations. It may be shown²⁹ that the perturbation energy is:

$$\delta E = -\frac{1}{\pi} \int_{-\infty}^{\varepsilon_{\rm F}} (\varepsilon - \varepsilon_{\rm F}) \delta \Im {\rm SpG}(\varepsilon) d\varepsilon = -\frac{1}{\pi} \int_{-\infty}^{\varepsilon_{\rm F}} \varepsilon \delta \Im {\rm SpG}(\varepsilon) d\varepsilon - \Delta n \varepsilon_{\rm F}, \quad (19)$$

where Δn is a difference in number of electrons in the perturbed system (calculated with the original Fermi level of the unperturbed system) and the initial one. In the self-consistent version of the calculations the modified Equation (25):

$$\delta E = -\frac{1}{\pi} \int_{-\infty}^{\varepsilon_F} (\varepsilon - \varepsilon_F) \delta \Im \mathsf{SpG}(\varepsilon) d\varepsilon - \frac{1}{2} \sum_{i,j,k,l} \left(\mathsf{P}_{ij} \mathsf{P}_{kl} \left(ij \| kl \right) - \left(\mathsf{P}_{ij}^{(0)} \right)_0 \left(\mathsf{P}_{kl}^{(0)} \right)_0 \left(ij \| kl \right)_0 \right),$$
⁽²⁰⁾

taking into account changes in the self-energy needs to be used. The summation in the last term goes over all orbitals involved in the

perturbed subspace *P*. Due to the decaying effect of the perturbation, it is a finite set as explained above. Two-center two-electronic integrals in this equation change in the perturbed system only if the geometry gets distorted.

4.1.3 | Local and virtual states

2358

As it was stressed yet in works,^{12,20} the defects may produce additional poles of the perturbed GF of two types. Either so-called local or virtual states may arise depending on the strength of the perturbation as related to the energy spectrum of the unperturbed system. The local states correspond to the poles on the real axis and, therefore, appear as narrow peaks of the perturbed DOS in the energy ranges, where the unperturbed DOS vanishes—outside the allowed energy bands of the ideal crystal. By contrast, the virtual states are related with the poles in the complex plane¹⁹ and manifest themselves on the real axis as wide Lorentzian peaks of the perturbed DOS inside the allowed energy band (see Supporting information section 4). Both features are perfectly reproduced by the GoGREENGO package as demonstrated in Figures 2 and 3. For more details about handling the pole structure of the perturbed GF in GoGREENGO see Supporting information section 4.

5 | TEST RESULTS AND DISCUSSION

Since the described approach, although, well established theoretically (analytically), did not so far enjoy full scale program implementation, it requires a thorough testing against analytically solvable models, even looking out oversimplified. Below, we present such tests and round up with an intermediate test—adsorption on graphene, which on one hand can be traced analytically far enough to provide necessary reference and on the other hand provides, although a simple, but realistic, example eventually suitable for experimental check. In the main text we mostly concentrate on the numerical results obtained by GoGREENGO, while analytical solutions, used for control, are collected in the Supporting information. Even in the case of simple models not



FIGURE 2 Perturbed densities of states $DOS_0(\varepsilon)$ (blue lines) in comparison with the initial ones (red lines) for the cubia lattices and schematic representations of atomic charges distributions induced by the substitutional defects. Dashed vertical lines show the Fermi level. Individual panels correspond to the following systems: (A) sc v = -1; (B) sc v = 1; (C) bcc v = -1; (D) bcc v = 1; (E) fcc v = -1; (F) fcc v = 1. Radii of the spheres are proportional to the charge of the atom (notice the different scales of Q_0 for different lattices in Table 2). Red spheres correspond to the positive charge, blue—to the negative. The biggest sphere in all cases is located on the defect site ("0"). On the plots red filling between the curves corresponds to the increase of the diagonal electronic density on the site and blue one—to decrease

CHEMISTRY WILEY 2359



FIGURE 3 Perturbed densities of states $DOS_0(\varepsilon)$ (blue lines) compared with the ideal ones (red lines) for the substitution defect in *p*-cubium: (A) v = -0.5; (B) v = 0.5. Color scheme is the same as in Figure 2

TABLE 1 Dispersion relations, Fermi level and the nearest neighbor Coulson bond-order for cubia with single (one electron per site) occupation. Indices k_i correspond to projection of wave vector k to orthogonal basis vectors of reciprocal space chosen so that in all cases the cubic Brillouin zone is defined as $-\pi < k_k \le \pi$

	ϵ_k	W	ε _F	В
Sc	$-2t(\cos k_x + \cos k_y + \cos k_z)$	12t	0	0.3324
Bcc	-8tcosk _x cosk _y cosk _z	16t	0	0.2605
Fcc	$-4t(cosk_xcosk_y+cosk_xcosk_z+cosk_ycosk_z)$	16t	0.915t	0.2184

all functionality of the package can be tested against analytical solutions, since the later are available only for the simplest local perturbations and are inaccessible in the frame of the self-consistent approach. Therefore, we test GoGREENGO against very simple benchmarks and then demonstrate its capabilities for more realistic and comprehensive setting.

5.1 | GoGreenGo for perturbation of cubia

Cubia (see e.g. Reference [60]) are simplest thinkable models of 3D metals. They are formed by s-orbitals centered at the vertices of (simple—sc, body centered—bcc, and face centered—fcc) cubic lattices with one-electron hopping *t* between the nearest neighbors of a given node of a lattice. The dispersion laws of electronic bands for such models⁶¹ allowing for analytic solutions as given in Table 1. Correspondingly, the eigenvectors (Bloch states) related to these eigenvalues are known and, consequently, the Green's functions. Specifically, diagonal GFs for the systems listed in Table 1 are known from References [61–64] and their plots are presented in Supporting information section 6. One can also find the corresponding graphs of the electronic DOS at the website.⁶⁵ The off-diagonal elements of the

respective Greenian matrices are as well accessible through GoGREENGO.

It is as well possible to find analytical solutions for the ideal cubia in the framework of the extended Hubbard model taking into account electron–electron interactions as described in Supporting information section 5. In our subsequent consideration we will use the extended Hubbard model as a starting point for the self-consistent calculations. All energy parameters are given in units of *t*, that is t = 1 everywhere below. The diagonal matrix elements of the unperturbed Fockian are set to zero being by this the energy reference. In further Subsections we present as coherence tests the results of numerical treatment of various local perturbations of cubia with use of GOGREENGO package.

5.1.1 | Lattice substitutions in cubia

First, we consider lattice substitution defects in cubia, where one atom of the crystal (denoted as "O") is replaced by a different atom. In general, the substitute can be a many-electron atom, but here we restrict our tests by single-electron impurities. Chemically this corresponds to substitution defects in alkali metals.

Complexity of the model can be gradually increased by including different terms into the perturbation operator so that different program features are tested independently. In the simplest possible setting we neglect electron-electron interactions and only take into account one-center perturbation: the difference v between diagonal matrix elements of the Fockian over impurity AO and the AO's of the unperturbed crystal. In this case an analytical solution is available (Supporting information section 8). Comparing numerical results to this solution shows that GoGREENGO produces the perturbed Greenian matrix identical to the analytical one up to eighth decimal sign. In addition, this setting allows making qualitative sketch of the features of the perturbed system, which remain valid for more involved crystals and perturbations of this kind. As one can conclude from the analytical form of the mass operator (see Supporting information section 4) and of the cubia GFs $G_{00}^{(0)}$ (Supporting information section 6), the additional poles in the perturbed system appear for: (1) sc |v| > 4.01; (2) bcc |v| > 6.10; (3) fcc v < -9.38 in the case of

WILEY-

2360

occupied local state and v > 0.83 in the case of vacant local state. In all these cases except for the local state above the band in the fcc lattice, the required value of v is unreasonably high and it is difficult to expect any local state to appear if, say, indeed, one alkali atom is exchanged by another (the typical difference in core attractions lies in the range of 0.4-0.8 eV⁶⁶-much smaller than the typical bandwidth). The only option is an appearance of the vacant local state in the fcc lattice, but it presents a minor interest since it does not contribute to the electronic density and the energy of the system. Another conclusion, which can be drawn from the analytical solution, is the behavior of the perturbed function in the vicinity of the pole of the initial GF in bcc. For bcc $\Im G_{00}^{(0)}$ is even function of ε and $\Re G_{00}^{(0)}$ is odd. In addition, due to the presence of the pole at $\varepsilon = 0$, $\Re G_{00}^{(0)}$ has a discontinuity there. Therefore, one would observe a discontinuity of the perturbed function $DOS_0(\epsilon)$ at $\epsilon = 0$, which is indeed observed in our numerical results described below (see Figure 4).



FIGURE 4 Perturbed diagonal densities of states $DOS_0(\varepsilon)$ in comparison with the initial ones and schematic representation of the charge distributions in graphene with the substitution defects: (A) boron, (B) nitrogen. Color code and other legend is the same as in Figure 2. Narrow peaks on the plots above the band in (A) and below the band in (B) correspond to the local states formed predominantly of the defect orbital

Although described model provides a simple and pictorial solution, it is rather far from realistic description of the substitution defects, requiring more involved interaction operator and the self-energy corrections. GoGREENGO supports such description and to test this, we included corrections to one-center two-electron and two-center hopping integrals, Coulomb interactions, and applied the self-consistent procedure. We performed calculations within this setting for cubia lattices for $v = \pm 1(t)$ complemented by a set of further parameters: variation of the hopping between the impurity and its neighbors $(\delta t = -0.2)$, one-center Coulomb repulsion in the unperturbed system $(\gamma_0 = 0.6)$, two-center (nearest neighbor) Coulomb repulsion in the unperturbed system ($\gamma_1 = 0.3$) and a variation of the Coulomb integrals in the defect ($\delta \gamma_0 = -0.1$ and $\delta \gamma_1 = 0.1$). The resulting perturbed $DOS_0(\varepsilon)$ together with obtained charge distributions are depicted in Figure 4; numerical values of charges and the electronic^{††} energy variations due to the defect formation are collected in Table 2.

In all cases the charge distributions induced by the defect have a similar oscillating-decaying behavior. The defect site ("0") carries the highest (by absolute value) net-charge, negative (electron density accumulation) for v = -1 and positive (electron density depletion) for v = 1. Absolute values of the net-charges on other sites decay with the separation from the defect and have alternating signs so that for any site its charge has the opposite sign to the charges of its neighbors. The decay rate is rather high as expected for 3D metals.²⁸ The fastest one is observed for fcc, where the induced charges are insignificant (<0.01) beyond the 2nd neighbors. For bcc the corrections become negligible beyond 3rd neighbors and for sc beyond the 4th ones. In all cases the Fermi level shifts as described in the Implementation Section and Supporting information section 3, but the absolute values of $\delta \varepsilon_{\rm F}$ do not exceed 7×10^{-6} . For v = -1 it shifts downwards and for v = 1—upwards. Such a small shift insignificantly affects individual density matrix elements. Maximal correction of the diagonal density matrix elements due to the Fermi level shift equals to $\Delta n/K$. For $31 \times 31 \times 31$ k-mesh used in our calculations and values of Δn from Table 2 they never exceed 5×10^{-5} for cubia, which is fairly negligible.

The (Coulson) bond orders—the off-diagonal elements of the oneelectron density matrix—are affected much less than the diagonal matrix elements of the density. Again, these corrections decay rapidly with the distance from the defect. In all cases, the bonds formed by the defect site are weaker than those of the innate atom, but even this correction occurs only at the third decimal place. Nevertheless, the sum of all corrections to the bond orders $\sum_{i \sim J} \delta B_{ij}$ can be fairly noticeable due to the large number of bonds in the cubia lattices. These values are presented in Table 2. As one can see, in all cases the total variation of the bond orders is negative, meaning that the system loses a part of the bonding energy because of the defect.

Electronic energy of the defect formation can be evaluated by Equation (20). An alternative expression for it is given in Supporting information section 5. As it can be seen from Table 2, in all cases the total energies are negative and the absolute values are much higher for the systems with v = -1, which is obviously explained by the sign of the one-center contribution of the impurity AO. Further significant contributions are the Coulomb attraction of the atomic charges, which is always negative due to the oscillatory behavior of the charge distribution, and the correction to the bonding energy, which is always positive as we saw above.

5.1.2 | Substitution defects in *p*-cubium

In the previous Subsection, we have tested the main functionality of GOGREENGO for cubia-single-band crystals. To demonstrate package's capabilities in treating multi-band solids we use a p-cubium (pc) model, which is formed by three p-orbitals located in the vertices of the simple cubic lattice. Due to symmetry, each orbital overlaps only with its own nearest translation images in the frame of the tightbinding approximation (e.g., p_x orbital overlaps only with neighboring p_x orbitals and orthogonal p_y and p_z). Two of such overlaps correspond to σ - σ interaction and four others to π - π interaction. In addition we assume the following for the two-electron Coulomb integrals $(ab | cd) = (aa | cc) \delta_{ab} \delta_{cd}$, which allows keeping the same number of two-electron parameters as previously. The analytical band-structure the ideal pc contains three degenerate bands for (see e.g. Reference [60]):

$$\varepsilon_{\alpha k} = -2t \left(-\cos k_{\alpha} + \kappa \sum_{\beta \neq \alpha} \cos k_{\beta} \right)$$
(21)

where $\alpha, \beta = x, y, z$ numerates bands and projections of *k*-vector, κ is a ratio between $\pi - \pi$ and $\sigma - \sigma$ hoping parameters and *t* stands for the $\sigma - \sigma$ one. Minus in front of the first cosine in the brackets occurs because the $\sigma - \sigma$ overlap of two *p*-orbitals is negative (and hopping is positive), if they are aligned in the same direction. In our further consideration we set $\kappa = 0.40$ and t = 1. The elements of the Greenian matrix for pc are accessible through GOGREENGO and can be found in Supporting information section 1. We consider pc with one electron

TABLE 2 Parameters of charge distribution in cubia with the substitutional defect and electronic energy of the defect formation. $Q_i = 1 - n_i$ corresponds to the atomic charge of *i*th neighbor of the defect site "O" with n_i being the electronic population of the *i*th site in the perturbed system

	v	Q ₀	Q1	Q ₂	Q ₃	$\sum_{\mathbf{i}\sim\mathbf{j}}\delta\mathbf{B}_{\mathbf{i}\mathbf{j}}$	Δn	δ Ε
sc	-1	-0.3478	0.1216	-0.0712	0.0555	-0.1620	0.0882	-1.9964
	1	0.3478	-0.1217	0.0711	-0.0555	-0.1594	-0.0899	-0.0029
bc	c –1	-0.4067	0.1058	-0.0835	-0.0389	-0.0600	0.1488	-2.0900
	1	0.4043	-0.1088	0.0819	0.0379	-0.0592	-0.0121	-0.0866
fco	c –1	-0.1984	0.0243	-0.0115	-0.0025	-0.0532	0.0294	-2.1583
	1	0.2903	-0.0293	0.0116	0.0025	-0.0413	-0.0766	-0.2671

per unit cell, yielding the Fermi level to be $\varepsilon_{\rm F} = -1.753t$. Coulson bond-order B^{σ} for the σ -bond is -0.2753 and for the π -bond is $B^{\pi} = 0.1474$. Note, that B^{σ} is negative and being combined with positive σ - σ hopping produces a negative contribution to the energy. In our further discussion of overall change in the bond orders it is convenient, therefore, to use values of $|B^{\sigma}|$.

To test the whole functionality of the program in the case of multi-band crystal we consider substitutional defects in the most rigorous setting by taking into account the self-energy correction and changes of one- and two-center integrals as it was done in Section 5.1.1. The set of parameters used for calculations: $v = \pm 0.5$, $\delta t = -0.2$, $\gamma_0 = 0.6$, $\gamma_1 = 0.3$, $\delta \gamma_0 = -0.1$ and $\delta \gamma_1 = 0.1$.

Perturbed DOS's projected on one of the *p*-orbitals of the defect site "O"are plotted on Figure 5 for the negative and positive perturbation. Resulting charge distributions exhibit similar behavior as in sc with some specific variations. First of all, the charges decay faster than in sc: they become negligible beyond 3rd neighbors for v = -0.5and beyond 4th for v = 0.5. Moreover, charge signs follow a different pattern: (1) for v = -0.5 the charges are $Q_0 = -0.4152$, $Q_1 = 0.0195$, $Q_2 = 0.0102$, $Q_3 = -0.0042$; (2) for v = 0.5 they are $Q_0 = 0.3039$, $Q_1 = -0.0245$, $Q_2 = -0.0044$, $Q_3 = 0.0217$. That is, the charges change their signs every two neighbors from the defect. Also, it can be seen that the negative perturbation causes a larger charge than the positive one of the same magnitude. This is a consequence of a nonsymmetric location of the Fermi level in the band (like in fcc cubium). Values of Δn are 0.1523 for the negative perturbation and -0.2524for the positive one. As previously, such small values do not cause significant shift of the Fermi level for our $31 \times 31 \times 31 \times 31$ k-mesh.

As for corrections to the bond orders, we see that they change significantly only for the bonds incident to the "O"site (σ -bond becomes weaker and π -one stronger in all cases). The total changes in bond orders are $\sum_{i\sim j} \delta B_{ij} = -0.1541$ for the negative perturbation and $\sum_{i\sim j} \delta B_{ij} = -0.2414$ for the positive perturbation, so the system loses bonding energy, upon formation of the defect.

5.1.3 | One-orbital interstitial impurity in simple cubium

Further class of problems accessible for GoGREENGO is that of interactions of solids with some extra additive termed as an interstitial defect. The simplest example of such defect is an interstitial (impurity) atom represented by a single s-orbital, which may be considered as a model of a hydrogen atom. We considered the effect of such impurity placed in a cubic void of the sc lattice being a nontrivial example illustrating general features of the methodology. In the cubic void, the defect is surrounded by eight identical atoms of the crystal system. The unperturbed Fockian of such a system is one of the crystal augmented by an extra row and column having the energy of the impurity orbital on their intersection and filled by off-diagonal zeroes signifying no interaction between the impurity and the crystal. The perturbation consists, in the first approximation, of one-electron hopping between the defect (s-)orbital and those of its immediate neighbors in the lattice (eight neighbors). These nine orbitals form the perturbation subspace (the P-subspace of Equation (11)) hereinafter addressed as the " impurity cluster".

The position of the defect *s*-orbital relative to the cubium Fermi level is given by a difference between ionization potentials of the interstitial atom and crystal atom of the lattice. For all metals, atomic ionization potentials are smaller than for hydrogen, therefore we assume that the diagonal matrix element over the impurity orbital is negative. In the cubia models the nearest neighbor hopping parameter *t* of the crystal provides the natural energy scale. We, at first, considered numerically the effect of interaction on the atomic charges and bond orders in the " impurity cluster" in the frame of the Anderson model (see Supporting information section 8.2). Analytical solutions for the Anderson model used for testing are given in Supporting information section 8.2.

Perturbed GFs obtained numerically with GoGREENGO for a "hydrogen" atom in a cubic void (for the first step of the self-



FIGURE 5 Perturbed diagonal DOS for the site nearest to the vacancy (left) and schematic illustration of the charge distribution caused by the vacancy defect in graphene. Legend for the plot and color scheme are the same as in previous figures

TABLE 3 GoGREENGO results obtained for the hydrogen impurity atom in the cubic void with the Anderson Hamiltonian (supporting information ssection 8.2). Parameter $\gamma = 0.40$ for all cases

En	-2				0	0				
ť	0.20	0.70	0.83	0.95	0.20	0.70	0.83	0.95		
na	1.9910	1.6823	1.4164	1.2164	0.6872	0.6023	0.6319	0.6557		
ni	0.9987	0.9608	0.9347	0.9130	0.9887	0.9534	0.9474	0.9418		
В	0.3311	0.2932	0.2672	0.2454	0.3211	0.2858	0.2798	0.2742		
Β'	0.0108	0.1286	0.2072	0.2446	0.1381	0.2772	0.2909	0.2995		
δΕ	-1.8278	-2.2008	-2.3433	-2.0104	-0.2086	-2.0233	-2.6101	-3.0670		

TABLE 4 GoGREENGO results obtained for the hydrogen impurity atom in the cubic void with taking into account inter-electronic interactions in the crystal. Parameters $\gamma = \gamma_0 = 0.4$, $\gamma_1 = \gamma' = 0.2$ for all cases (see supporting information section 8.2 for the equations)

ε _a t'	-2				0	0				
	0.20	0.70	0.83	0.95	0.20	0.70	0.83	0.95		
na	1.9914	1.6814	1.4592	1.2321	0.6242	0.6079	0.6387	0.6654		
ni	0.9470	0.9306	0.9330	0.8935	1.0098	0.9816	0.9748	0.9701		
В	0.3296	0.2887	0.2779	0.2313	0.3198	0.2833	0.2770	0.2728		
Β'	0.0116	0.1336	0.2417	0.2511	0.1450	0.2791	0.2923	0.3027		
δΕ	-1.8720	-2.2704	-4.5067	-2.0715	-0.2247	-2.0645	-2.6345	-3.3018		

consistent procedure) were compared with corresponding analytical equations from Supporting information section 8.2. The values coincide up to 6th decimal place. Corrections to density matrix elements are not accessible analytically due to a very involved functional form of $\mathbf{G}^{(0)}$ even for the sc cubium. Numerical values are collected in the Table 3. As one can conclude, if the adsorbate level is positioned below the Fermi level the impurity atom acquires a negative charge, while for $\varepsilon_a = \varepsilon_F = 0$ the charge is positive. The absolute value of charge depends on the impurity-matrix hopping parameter t'-as it increases, the charge decays which is a result of changing in coupling patterns (see the four types of perturbed adsorbate levels in Supporting information section 8.2). Atoms of the crystal matrix in all cases acquire a small positive charge.

Bond orders *B* between the crystal atoms of the " impurity cluster" become smaller (the bonds become weaker) and this energetically unfavorable change is compensated by formation of new bonds with the order *B*′ between impurity atom and lattice atoms. With increasing hopping parameter *t*′ the orders of new bonds increase and the orders of original bonds decrease. For $\varepsilon_a = 0$ the bond orders *B*′ are larger than for $\varepsilon_a = -2$.

In a more extended setting, we turn on the Coulomb interactions in the crystal. In this case, non-vanishing matrix elements between the sites outside the impurity cluster appear, which makes an analytical treatment unfeasible. The point to be checked here, is the extension of the number of atoms in the impurity cluster for accounting of the perturbation dressing through electron-electron interactions. According to the tests, it is sufficient to include first and second neighbors of the lattice atoms directly interacting with impurity. As it can be concluded from Table 4 including two-electron Coulomb terms and the self-energy correction does not have major influence on the resulting density matrices and energies; all trends remain fairly the same with slight numerical modifications. The only exception is the case of $\varepsilon_a = -2$ and t = 0.83, where one observes more significant differences in density matrix elements n_a , B and B' between two settings, which translates into quite considerable difference in energies.

5.2 | GoGreenGo for perturbations of graphene

Two-dimensional graphene sheets are widely studied experimentally and theoretically being nature models for two dimensional solids and, respectively, surfaces. Either lattice impurities or adsorbates in/on graphene are of considerable interest from the experimental and theoretical points of view.^{67,68}

The honeycomb lattice of graphene has a primitive cell containing two carbon atoms. The π -system is formed by p_{π} -orbitals, one for each site of the lattice. Dispersion law in the approximation of the nearest neighbor hopping is:

$$\varepsilon(k) = \pm t \sqrt{3 + 2\cos k_x + 2\cos k_x + 2\cos (k_x - k_y)}, \qquad (22)$$

where "-" corresponds to the filled band and "+" to the empty one. Equation (22) allows analytical evaluation of the diagonal DOS⁶⁹ (see also Supporting information section 7). The Fermi level $\varepsilon_{\rm F} = 0$ and the Coulson bond order B = 0.525 between nearest neighbors. 2364 WILEY- COMPUTATION

In the present Section the Greenian matrix of pure graphene serves as a starting point for the study of several types of local defects. We concentrate on the effect of the local perturbations on the π -system of graphene and for the time being ignore the σ -system.

5.2.1 | Substitution defects in graphene

Substitution defects in graphene can be described in the same manner as in cubia. Here we aim to study defects, which are closer to the experimental situation. Specifically, we consider the graphene layer, where one of the carbon atoms is replaced by nitrogen or boron. This is modeled by an appropriate choice of parameters describing oneand two-center interactions as described in Supporting information section 9. The parameters used to study boron and nitrogen impurities in graphene lattice are collected in Table 5.

The bare perturbed subspace is spanned by four π -orbitals, one of boron/nitrogen and three more of their nearest neighbors. The perturbation itself touches the diagonal matrix element of the substituted atom and the hopping integrals with its nearest neighbors.

For 2D graphene being a poor metal (Fermi surface degenerates to two–Dirac–points) one can expect a slower decay of perturbation effect with the separation from the defect. To ensure we do not miss any significant corrections to the density matrix elements we performed self-consistent calculations of substitutional defects using an impurity cluster formed by 7×7 original cells, whose unperturbed Greenian matrix was derived from the band structure of the unit cell with two atoms Equation (22). As previously we denote the defect site as "0".

Diagonal DOS on the "O"th site together with schematic representation of the charge distribution in the perturbed systems is given in Figure 2. The charge distribution parameters and electronic energies of the defect formation are collected in Table 6. As one can see from the plots of DOS₀(ε) the local states outside the band appear, respectively, below the band for nitrogen ($\varepsilon_1 = -3.053$) and above it for boron ($\varepsilon_1 = 3.773$). For nitrogen the local state contributes 0.38 to the diagonal density matrix element which is 26.95% of the total

TABLE 5 Atomic parameters used in this work to describe boron and nitrogen impurities in graphene lattice. All values are in units of *t*

	v	β	γ ₀	γ1
С	-	0.792	2.100	0.792
В	2.488	0.914	1.228	0.699
N	-1.251	0.904	3.573	0.862

value. In the case of boron the local state is vacant and does not contribute. The Fermi level in both cases shifts since nitrogen brings one more electron to the system (hence $\Delta n = 0.91$) and boron withdraws one electron (hence $\Delta n = -1.10$). However, the value of $\Delta n/K$ is again very small and does not affect the individual density matrix elements significantly (we used 51×51 k-mesh).

The qualitative behavior of the atomic charge distribution is fairly similar to that observed in cubia with the only exception that values of the charges decay much slower here, as anticipated. In both cases the corrections to density become negligible (<0.01) beyond 6th neighbor. This is rather reasonable response to very strong perturbation induced by substitutions. Spectacular alternation of signs of the charges induced by substitution seen in Figure 2 is nothing, but the manifestation of the Coulson's "law of alternating polarity",⁷⁰ which had been known already to Hückel⁷¹ and served to explain the rules of ortho-meta-para orientation in the electrophilic substitution in benzene derivatives known in organic chemistry⁷².

In complete analogy with cubia, the bond orders are much less affected by the predominantly diagonal perturbation. This effect manifests only for the bonds closest to the defect and become negligible as from the bonds between 3rd and 4th neighbors. The order of the bond closest to the defect decreases by 0.048 for nitrogen and 0.165 for boron. The total change in bond orders $\sum_{i\sim j} \delta B_{ij}$ is, however, quite significant due to the number of bonds involved (see Table 6) and comprises the contributions of the bond orders up to those between 3rd and 4th neighbors, therefore being more local than the variation of diagonal densities (charges).

Energies of the defect formation have different signs—negative for nitrogen and positive for boron. In the latter case this is the result of the huge positive one-center contribution.

5.2.2 | Vacancy in graphene

Vacancies in graphene lattice are interesting from two viewpoints. First, they can appear in the material during synthesis and, therefore, can have an impact on the properties of available samples. Second, vacancies in the π -system are tentative models of a carbon atom forming an extra bond with some unsaturated particle, which acquiring the sp3 hybridization, leaves the π -system. In the case of not self-consistent setting analytical GFs can be found, which are given in Supporting information section 9. Comparison of GOGREENGO numerical GFs to analytical ones showed they coincide up to eight decimal place. To study vacancy in a more realistic setting we again apply the

TABLE 6 Parameters of charge distribution in the graphene with the substitutional defects and electronic energy of the defect formation. Q_i corresponds to the atomic charge of *i*th neighbor of the defect site "O"

	Q ₀	Q 1	Q ₂	Q_3	Q 4	Q ₅	Q ₆	$\sum_{i\sim j} \delta \mathbf{B}_{ij}$	Δn	δ Ε
В	0.74	-0.39	0.24	-0.22	-0.16	0.14	-0.09	-0.76	-1.10	1.04
Ν	-0.42	0.27	-0.17	0.15	0.12	-0.10	0.07	-0.42	0.91	-1.42

CHEMISTRY WILEY 2365



FIGURE 6 Diagonal AO-projected DOS for the atomic orbital of adsorbate (A) and for the orbital of graphene's site interacting with the H atom (B). On the right plot (B) red curve corresponds to the initial (unperturbed) DOS and the blue one—to the perturbed DOS. Narrow peaks on the plots above the band corresponds to the local state. On the plot a) one can also observe a virtual state being wide peak centered at the point ca. -0.5

self-consistent option of the package with one- and two-center parameters of carbon being the same as in the previous Subection.

As before the perturbation cluster involves four orbitals - one excluded from the π -system and three its nearest neighbors. The perturbation itself reduces to nullifying the hopping matrix elements between the excluded site and its neighbors. The resulting charge distribution as well as AO-projected DOS for the site nearest to the vacancy are shown in Figure 6. Remarkably, the situation is guite different from that of the previous Subsection. For the predominantly offdiagonal perturbation, corrections to the diagonal densities are not that large ($Q_1 = 0.096$, $Q_2 = -0.050$, $Q_3 = 0.045$, $Q_4 = -0.032$) and become negligible beyond 4th neighbors from the vacancy. On the contrary, off-diagonal densities (bond orders) are affected much stronger: $\delta B_{01} = -0.525$ (quite obvious since the bond between the vacant site and its neighbor disappears), $\delta B_{12} = 0.085$, $\delta B_{23} = -0.013$, $\delta B_{34} = 0.012$, $\delta B_{45} = -0.020$ and all other changes occur only at a third decimal place. With a good accuracy (up to third decimal place) the following holds:

$$\sum_{i \sim j} \delta B_{ij} \approx 3\delta B_{01} + 6\delta B_{12}, \qquad (23)$$

so that, all other oscillations of bond orders compensate each other and the total change of the bonding energy is a sum of two local terms – energy of three broken bonds and energies of six nearest bonds, which become stronger.

5.2.3 | Adsorption of hydrogen/alkali metal atom on graphene

On-top adsorption of atomic hydrogen on graphene is a process of considerable interest in material science because it leads to a formation of sp³ defects, which usually present in synthetic graphene and affect its properties. It is known⁷³ that chemisorption of hydrogen forces carbon atom to rise above the plane by 0.4 Å and to form a σ -bond with the adsorbate. This causes a change of hybridization state of the carbon atom (from sp² to sp³) and consequent reorganization of both π - and σ -systems-three π -bonds break and one new σ -bond appears along with distortion of three C–C σ -bonds. Complete treatment of such process can be performed in $\sigma - \pi$ approximation by considering π -bonds breaking as a vacancy forms (as described above) and a rigorous evaluation of the σ -system reorganization energy. Although theoretical basis for that has been already established in our previous works, 57,74-76 full analysis of the problem goes beyond the scope of the present paper. Here, we restrict ourselves by testing the effect of interaction of "hydrogen" atom only with the graphene π -system, neglecting possible distortion of graphene geometry and not touching the σ -core. Treatment of the on-top adsorption on graphene in the framework of the standard Anderson impurity model was given in References [78-83]. Here we employ more advanced self-consistent model taking into account electron-electron interactions at different atomic sites.

Parameters required to describe interactions are calculated within the MNDO setting which has shown its validity for description of carbon allotropes.⁵⁷ We take R(C-H) = 1.1 Å The adsorbate level lies below the Fermi level by $\varepsilon_a = -2.150$ and one-center electron-electron repulsion on the adsorbate *s*-orbital is $\gamma'_0 = 3.952$. Two-center parameters for the C-H pair have the following values $\beta' = 1.095$ and $\gamma'_1 = 1.41$. All energies are in the units of *t*, which is known to be 2.4 eV in graphene.⁷⁷ We denote the graphene site, interacting with the H atom, as "0". The bare perturbation acts in the two-dimensional space (0th π -AO of graphene and adsorbate *s*-orbital):

TABLE 7 Atomic charges induced by the on-top chemisorption of hydrogen atom on graphene calculated within the self-consistent and non-self-consistent settings. Q_i corresponds to the atomic charge of *i*th neighbor of the carbon atom "0" interacting with adsorbate. Q_a stands for the charge of the hydrogen atom. "-" corresponds to a negligibly small charge

	Q _a	Q ₀	Q ₁	Q ₂	Q ₃	Q ₄	Q ₅	Q ₆
self	-0.7516	0.4607	-0.3137	0.1942	-0.1355	0.1174	-0.0775	0.0538
non self	-0.8372	0.1103	-0.0187	0.0081	-	-	-	-

$$\delta F = \begin{pmatrix} 0 & -\beta' \\ -\beta' & 0 \end{pmatrix}, \tag{24}$$

but the size of the cluster used for the self-consistent calculations is, of course, larger due to Coulomb interactions. To ensure we do not miss any significant changes in the density matrix elements, we, as previously, use the impurity cluster of 7×7 original graphene cells, whose unperturbed Greenian matrix was derived from the band structure calculated with the original unit cell of two atoms.

2366 WILEY-

Perturbed AO-projected DOS for adsorbate: $DOS_a(\varepsilon)$ and for the Oth π -AO of graphene: $DOS_0(\varepsilon)$ are plotted in Figure 3. The adsorbate DOS below the Fermi level consists of two broad peaks. The first one lies in the interval [-3, -1] with the maximum at $\varepsilon = \varepsilon_a$. It is obviously the adsorbate level broadened due to interaction with the graphene p-band. The second peak is much smaller than the first one and lies in the interval [-1,0] with the maximum at $\varepsilon = -0.630$. At the point $\varepsilon = -1$ the DOS drops to zero^{‡‡} due to the logarithmic singularity of the pure graphene DOS (GF). There is a local state above the band at $\varepsilon = 3.090$, which does not contribute to the electronic density. No local states show up below the band in this case.

Adsorbate *s*-AO acquires a negative charge $Q_a = -0.7516$ and the site "0" a positive one of $Q_0 = 0.4607$. The charge distribution in graphene lattice generally follows the same pattern as in the impurity problem—the values of the net-charges decay with the distance from the defect with alternating signs. The decay is rather slow and the charges become negligible (<0.01) only beyond 6th neighbors of the adsorption site (values are collected in Table 7). Due to the alternation of atomic charges, the contribution of two-center Coulomb interactions is negative, favoring the perturbed state.

Corrections to the bond orders decay much faster with the distance than atomic charges. Coulson bond order for the C-H bond formed upon adsorption is $B_{a0} = 0.5802$. Three bonds of the adsorption site with its neighbors in the layer turn weaker ($\delta B_{01} = -0.1672$), six bonds between 1st and 2nd neighbors turn stronger ($\delta B_{12} = 0.0173$) and 12 bonds between 2nd and 3rd neighbors again weaker ($\delta B_{23} = -0.01131$). All further corrections to the bond orders are negligible (<0.01) and do not contribute significantly to the adsorption energy. Due to the formation of the C-H bond the system gains the energy:

$$\delta E_{\rm b}^{\rm C-H} = -\left(2\beta' + \frac{\gamma'_1}{2}B_{\rm a0}\right)B_{\rm a0} = -1.5080. \tag{25}$$

At the same time, it loses the energy due to the weakening of three C-C bonds connected with the site "0":

$$\delta E_{\rm b}^{\rm C-C} = -6t \delta B_{01} = 1.0032. \tag{26}$$

In total, the change in bonding energy is negative.

Electronic energy of chemisorption can be calculated by Equation (20) or alternatively one can adapt Equation (31) from Supporting information section 5, which in both cases gives the value of $\delta E =$ -1.392 and favors the adsorption in agreement with DFT calculations.⁷³ However, numerical comparison of binding energies is not possible at the moment, since the calculated electronic energy does not include significant positive terms of core-core repulsion, which will be added in the next release. We, also, did not take into account rehybridization effects. The most significant negative contributions to the binding energy come from one-center term (-1.62), two-center Coulomb and bonding terms mentioned above. The most significant positive contribution comes from one-center repulsion of adsorbate electrons with different spin-projections which equals to $\gamma'_0 n_2^2/4 =$ 1.7306 (note, that we employ a non-magnetic solution for the perturbed system, but take into account that unperturbed hydrogen atom bears one unpaired electron).

In conclusion of this discussion, we mention that omitting the self-consistency procedure in the above model calculations leads to the charge distribution drastically differing from that obtained above. In the non-self-consistent setting with the bare perturbation Equation (24) the fast-decaying charge distribution is obtained as shown in Table 7. The local state below the band at $\varepsilon = -3.05$ appears, which is not there in the self-consistent calculations. This example demonstrates that in some cases the self-consistency is important, when considering point defects, not only quantitatively perspective, but also qualitatively.

6 | CONCLUSION

We present the GoGREENGO package intended for description of point defects in crystals as well as for analysis of adsorption processes on surfaces. The package employs the Green's functions formalism in order to obtain the densities of states of the perturbed crystal containing defects. It builds the Green's function of the unperturbed crystal from the output produced by major ab initio solid-state quantum chemistry codes and solves the Dyson equation for the perturbed GF. Results of this calculation are processed so that the perturbed densities of states, charge distributions and the off-diagonal matrix elements of the density are available in the atomic basis. The package is tested for the electronic structure perturbations caused by various local defects in model cubia crystals, in more realistic graphene and

simple multiband model of p-cubium. In all cases, when the analytic solutions were available, the package manifested perfect agreement with the former.

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DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available either in Supplementary Materials or from the corresponding author upon reasonable request.

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ENDNOTES

- * Relatively modern application of GFs to the problem of point defects in crystals was developed in the framework of Korringa-Kohn-Rostoker method (KKR-GF),³² which is based on ideas of multiple-scattering theory and, consequently, relies on rather different formulation of the GF theory compared to the one used in above cited references and in the present paper (see below).
- [†] Numerous sources are available, in the computational chemistry context Refs. [29,30] can be recommended.
- [‡] The bra-ket notation is used (see, e.g. Ref. [37]).
- [§] Sometimes called a mass operator.
- $^{\parallel}$ It is equal to the number of unit cells used in a periodic model of the infinite crystal.40
- ** This is the most unfavorable situation the decay in insulators is even faster.
- **††** Correction to core-core repulsion term caused by the defect is not included
- ^{‡‡} Since in practical calculations we do not have a genuine pole at $\varepsilon = -1$ and it is approximated by a peak with small vet finite width, the numerical DOS does not drop to zero exactly. However, this fact does not influence an integral of DOS.

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