

Structure and Bonding 150

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Mihai V. Putz

D. Michael P. Mingos *Editors*

Applications of Density Functional Theory to Biological and Bioinorganic Chemistry

 Springer

150

Structure and Bonding

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Aims and Scope

The series *Structure and Bonding* publishes critical reviews on topics of research concerned with chemical structure and bonding. The scope of the series spans the entire Periodic Table and addresses structure and bonding issues associated with all of the elements. It also focuses attention on new and developing areas of modern structural and theoretical chemistry such as nanostructures, molecular electronics, designed molecular solids, surfaces, metal clusters and supramolecular structures. Physical and spectroscopic techniques used to determine, examine and model structures fall within the purview of *Structure and Bonding* to the extent that the focus is on the scientific results obtained and not on specialist information concerning the techniques themselves. Issues associated with the development of bonding models and generalizations that illuminate the reactivity pathways and rates of chemical processes are also relevant.

The individual volumes in the series are thematic. The goal of each volume is to give the reader, whether at a university or in industry, a comprehensive overview of an area where new insights are emerging that are of interest to a larger scientific audience. Thus each review within the volume critically surveys one aspect of that topic and places it within the context of the volume as a whole. The most significant developments of the last 5 to 10 years should be presented using selected examples to illustrate the principles discussed. A description of the physical basis of the experimental techniques that have been used to provide the primary data may also be appropriate, if it has not been covered in detail elsewhere. The coverage need not be exhaustive in data, but should rather be conceptual, concentrating on the new principles being developed that will allow the reader, who is not a specialist in the area covered, to understand the data presented. Discussion of possible future research directions in the area is welcomed.

Review articles for the individual volumes are invited by the volume editors.

In references *Structure and Bonding* is abbreviated *Struct Bond* and is cited as a journal.

Mihai V. Putz • D. Michael P. Mingos
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Applications of Density Functional Theory to Biological and Bioinorganic Chemistry

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Preface

In the early twentieth century following the elucidation of the structure of atoms it became evident that atoms and molecules with even numbers of electrons were far more numerous than those with odd numbers of electrons. In 1916, G. N. Lewis provided the first comprehensive description of ionic and covalent bonds, when he postulated that atoms tend to hold an even number of electrons in their outer shells and a special stability was associated with eight valence electrons, which he speculated were arranged symmetrically at the eight corners of a cube. In 1919, I. Langmuir suggested that the structure of the periodic table could be rationalized using an extension of Lewis' postulates. In 1922, N. Bohr updated his model of the atom by assuming that certain numbers of electrons (for example 2, 8, and 18) corresponded to stable "closed shells." In 1926, Schrödinger established a wave mechanical description of the hydrogen atom which was subsequently extended to polyelectron atoms. Pauli was the first to realize that the complicated numbers of electrons in closed shells can be reduced to the simple rule of *one* per state, if the electron states are defined using four quantum numbers. For this purpose he introduced a new two-valued quantum number, identified by Goudsmit and Uhlenbeck as electron spin. The resulting Pauli Exclusion Principle states that no two electrons in a single atom can have the same four quantum numbers; if n , l , and m_l are the same, m_s must be different such that the electrons have opposite spins.

The idea of shared electron pairs introduced by Lewis provided an effective qualitative picture of covalent bonding and it still forms the basis of the universal notation for chemical communication, but it was Heitler and London who in 1927 developed the first successful quantum mechanical expression for this bonding model. Initially they provided a description of the bonding in molecular hydrogen, but it was subsequently adapted to more complex molecules and its widespread applications were articulated with great conviction by Linus Pauling. An alternative molecular orbital description of chemical bonding originated from Burrau's description of the hydrogen molecule ion and this model was subsequently widely developed by Mulliken and Lennard-Jones. The electrons occupy molecular orbitals which are delocalized over the whole molecule and were filled according to the Aufbau Principle and assigned quantum numbers according to the Pauli

Exclusion Principle. The orbitals are calculated in a self-consistent fashion in a manner analogous to those developed previously for atomic orbitals and are based on linear combination of the atomic orbitals of the individual atoms. The number of molecular orbitals equals the number of atomic orbitals in the atoms being combined to form the molecule. A molecular orbital describes the behavior of one electron in the electric field generated by the nuclei and some average distribution of the other electrons. This approximation proved to be more amenable to computer programming than the valence bond model and was widely developed and used in increasingly less approximate forms from 1960 to 1990.

In the early 1970s, a new electronic structure approach emerged from the physics community and was described as density functional theory (DFT). The total energy of a molecule was expressed as a functional of the total electron density. Hohenburg and Kohn proved the unique relationship between electron density and energy and Kohn and Sham put forward a practical variational DFT approach. Although calculations in solid-state physics had been reported since the 1970s DFT was not considered accurate enough for calculations in quantum chemistry until the 1990s, when the approximations used in the theory were refined to more accurately describe the exchange and correlation interactions. Computational costs for *ab initio* DFT calculations are relatively low when compared to the valence bond and molecular orbital methods. DFT thus began to approach the goals of computational thermochemistry to calculate the energetic properties of chemical processes to an accuracy of 1 kcal mol^{-1} . The widespread acceptance of these methodologies by the chemical community led to Kohn and Pople sharing the Nobel Prize in Chemistry in 1998.

When in 2004 Volumes 112 and 113 of *Structure and Bonding* were devoted to the “Principles and Applications of Density Functional Theory in Inorganic Chemistry” the editors N. Kaltsoyanis and J.E. McGardy noted “It is difficult to overestimate the impact that Density Functional Theory has had on computational quantum chemistry over the last two decades. Indeed, this period has seen it grow from little more than a theoretical curiosity to become a central tool in the computational chemist’s armory.” In these volumes they described recent applications in inorganic and biochemistry and addressed key issues in spectroscopy, mechanistic studies, and magnetism.

As possibly the dominant discipline of the twenty-first century the biological sciences have assimilated analytical, conceptual, and computational techniques from the other natural sciences. The continuing need for interpreting the vast amount of new data from *in vivo* and *in vitro* experiments using causal and deterministic hypothesis requires a wide range of statistical and computational tools and algorithms. As a consequence bioinformatics and mathematical, physical, and chemical biology have flourished and been used to interpret complex natural biological phenomena and pharmaceutical/toxicological effects of chemicals to natural systems.

The universal implications of chemical interactions and more specifically the structure and bonding characteristics of biomolecules suggest that DFT may also play a crucial role *in cerebro* and *in silico* experiments. Establishing the molecular

basis of biological principles by means of quantum mechanical tools has become a realistic possibility given the current accuracy of DFT methods. The present volume opens with an authoritative review of the extensions of DFT (dispersion-corrected functionals, Born–Oppenheimer dynamics, hybrid with molecular mechanics, constrained, and interpretational) from chemical reactions to biochemical systems (containing over a hundred atoms, enzyme kinetics, etc.). The dispersion problem and the development of dispersion-corrected DFT, which may be used accurately to describe weakly bonded biological systems, are further formalized by specific density functional features in the second chapter. Computational models of DFT are used in the next chapter to exemplify the theoretical counterparts of the spectroscopic data to define the binding and activation energies of small molecules with high bioinorganic implications such as water, congeners of molecular oxygen, nitrogen oxides and oxyanions, sulfide, sulfur oxides and oxyanions, carbon dioxide, organic compounds, halogens, molecular hydrogen, and protons. The computational DFT approach as applied to the electronic localization functions and maximum probability domain analyses for modeling metal–porphyrins. These results suggest that the bonding is primarily ionic in porphyrins containing transition and non-transition metals. The last two chapters deal with the important problem of modeling toxicity phenomena using reactivity principles derived from DFT calculations. After introducing the connection between chemical structure and biological information by connecting the chemical reactivity with biological activity within the quantitative structure–activity relationship (QSAR) technique, the possible anticancer activity of two new metal–borane clusters is explored. It is further generalized by the last chapter which describes the full merging of the QSAR with logistic enzyme kinetics. This leads to a description of the mechanisms of chemical–biological interactions in chlorinated-PAHs by means of chemical reactivity principles derived from conceptual DFT.

Overall the volume provides a coherent exposition of the application of DFT to various biological and bioinorganic chemical systems. We hope that it will encourage the DFT community in further refining and extending the electronic models to complex and correlated biological–chemical systems and interactions in the years to come.

We thank the contributors to this volume for the consistent efforts they have made in writing high-class scientific reviews and for providing the readers with a broad perspective which has revealed the widespread uses of DFT in interpreting biological and bioinorganic systems. MVP acknowledges the research and editing facilities provided for the present volume by the Romanian Education and Research Ministry within the project CNCS-UEFISCDI-TE-16/2010-2013. MVP and DMPM sincerely thank the Springer team and in particular Marion Hertel, Ursula Gramm, Elizabeth Hawkins, and Tanja Jaeger for professionally supervising the production of the **Structure and Bonding** series in general and of this volume in particular.

Timișoara, Romania
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Contents

| | |
|---|------------|
| Recent Progress in Density Functional Methodology for Biomolecular Modeling | 1 |
| Dennis R. Salahub, Aurélien de la Lande, Annick Goursot, Rui Zhang, and Yue Zhang | |
| Density Functional Theory and Molecular Interactions: Dispersion Interactions | 65 |
| Eugene S. Kryachko | |
| Redox Activation of Small Molecules at Biological Metal Centers | 97 |
| Radu Silaghi-Dumitrescu | |
| The Bond Analysis Techniques (ELF and Maximum Probability Domains) Application to a Family of Models Relevant to Bio-Inorganic Chemistry | 119 |
| Mauro Causà, Maddalena D'Amore, Carmine Garzillo, Francesco Gentile, and Andreas Savin | |
| Biological Activity and Toxicity: A Conceptual DFT Approach | 143 |
| Arindam Chakraborty, Sudip Pan, and Pratim K. Chattaraj | |
| DFT Chemical Reactivity Driven by Biological Activity: Applications for the Toxicological Fate of Chlorinated PAHs | 181 |
| Mihai V. Putz and Ana-Maria Putz | |
| Index | 233 |

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Recent Progress in Density Functional Methodology for Biomolecular Modeling

Dennis R. Salahub, Aurélien de la Lande, Annick Goursot,
Rui Zhang, and Yue Zhang

Abstract Density Functional Theory (DFT) has become the workhorse of applied computational chemistry. DFT has grown in a number of different directions depending on the applications concerned. In this chapter, we provide a broad review of a number of DFT and DFT-based methods, having in mind the accurate description of biological systems and processes. These range from pure “cluster” DFT studies of the structure, properties, and reactions of biochemical species (such as enzymatic catalysts) using either straight DFT or dispersion-corrected functionals (DFT-D), to Born–Oppenheimer-DFT dynamics of systems containing up to a hundred atoms or more (such as glycerol-lipids), to hybrid DFT/Molecular Mechanical Molecular Dynamics methods which include protein and solvent environments (for enzymes or ion channels, for example), to constrained-DFT (working within the Marcus framework for electron-transfer reactions), to Interpretational-DFT (which provides the interpretational benefits of the Kohn–Sham DFT methodology).

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 • Interpretational DFT • QM/MM methodology

Contents

| | | |
|-----|--|----|
| 1 | Introduction | 2 |
| 2 | DFT and ADFT | 5 |
| 2.1 | Methodology | 5 |
| 2.2 | Applications | 8 |
| 3 | DFT-D | 15 |
| 4 | BODFT-MD | 19 |
| 4.1 | Introduction | 19 |
| 4.2 | Properties of Phosphatidyl Choline Lipids | 20 |
| 4.3 | Activation of Triplet Dioxygen by Bio-inspired Cuprous Complexes | 26 |
| 5 | DFT/MM-MD | 29 |
| 6 | Constrained-DFT | 39 |
| 6.1 | Methodological Background | 39 |
| 6.2 | cDFT and Population Analyses | 43 |
| 6.3 | Modeling Electron Transfer Reactions | 44 |
| 6.4 | Other Applications of cDFT | 48 |
| 7 | Interpretational-DFT | 49 |
| 8 | Conclusions and Perspectives | 51 |
| | References | 52 |

1 Introduction

Density Functional Theory (DFT) has become the workhorse of applied computational chemistry because of its particularly appealing combination of accuracy, speed, and interpretability. Nearly five decades have transpired since the seminal papers of Hohenberg and Kohn [1] and Kohn and Sham [2] and nearly nine since the first use of a density functional in atomic theory by Thomas [3] and by Fermi [4]. Each decade has seen steady, inexorable, progress—more accurate functionals have been developed, better and faster algorithms have been implemented, and new analytical concepts have been devised (see e.g., [5] for a review covering the 1964–2004 period). The new methodologies and techniques have allowed systems of ever growing complexity to be addressed, to the point where, now, DFT is starting to have a real impact on biological questions.

In this review, we will focus on biology and try to capture the state of the art in studies that approach aspects of biological systems and processes from various points of view, all of them involving DFT. We will exclude from our scope the semiempirical Tight-Binding DFT (DFTB) and also Time-Dependent DFT (TDDFT). Although these methods have been used to study systems of biological interest, we choose to discuss here methods and applications that involve “real” ground-state DFT, including hybrid functionals, at the core.

We have chosen a methodological hierarchy that extends from “simple” molecules and cluster models in which the usual tools of quantum chemistry, (geometry optimization, transition state localization, reaction path following), are applicable, to dynamical studies within the Born–Oppenheimer approximation, to systems of greater extent and complexity in which part of the system is treated with a molecular-mechanical force field (the so-called QM/MM approaches) and, finally, to systems involving long-range charge transfer for which the newly developed constrained DFT has considerable merit. Calculations at any level of the hierarchy require an interpretational framework and a section of the chapter is devoted to some of the concepts involved, most notably various population analyses and the Electron Localization Function (ELF).

We have chosen this particular cross section of DFT-centric subjects because of our own contributions to the methodologies and their applications but we will attempt to put our own work in the context of other contributions by reviewing some of the main contributions over the last few years. Our review will not be exhaustive, and we apologize for the omission of any particular works that undoubtedly may be as pertinent as those we have chosen. Our goal is to paint as faithful a picture as possible of the state of the art through inclusion of a representative sampling rather than an exhaustive review. We do, however, aim to give a comprehensive account of the main issues involved in choosing a computational protocol for the types of biological models we have treated, including the strengths and the limitations of the various methodologies.

To the novice (or even expert) biomolecular modeler the choice of a problem and of the appropriate methodology to address it can often be a daunting challenge. In order to put the particular biological processes we have chosen for this review, along with the six types of methodology we describe, within a common organizational framework we offer the following protocol (enzymatic catalysis has been chosen as an exemplar but we think the protocol also applies to other problems (ion channels, lipid conformations, peptide agglomeration, electron transfer, etc., with some changes)).

1. Choose a relevant biological process and system. What are the biological/biochemical/biophysical questions that are to be addressed? This choice will of necessity involve only a small part of biological reality. We are in reductionist mode here; the question of integration into a systems framework will, for the most part, be left to other publications.
2. Choose an appropriate active component of the biological system (in our exemplar, an enzyme active site).
3. Consider a proposed mechanism and build a model, using crystallographic data if available, from the active site outwards. Choose important protein residues, substrates, intermediates and products, water molecules involved in the mechanism, etc., knowing that the validity of the results will depend critically on these choices.
4. In the case of a “simple” cluster model, decide whether to fix any of the peripheral atoms at their crystallographic positions.

5. Decide on the charge state(s) of acidic residues by calculating pK_a 's. If more than one charge state (protonated or unprotonated) seems possible, extend the study to examine both possibilities.
6. Also in the case of a cluster model, decide whether to include the effects of the surrounding protein and solvent through the use of a Polarizable Continuum Model (PCM). Choose the value of the dielectric constant (often chosen to be 4.0).
7. Since in this review we are considering only DFT, decide on the variant of DFT to use (mainly GGA or hybrid functionals, with dispersion corrections for highest accuracy) along with the computational parameters (basis sets, auxiliary basis sets, integration grids, SCF and geometry convergence criteria, etc.). Choose a software package (or write one...).
8. Design the study to check that the cluster model is large enough either through cluster-convergence tests or, if that is not possible, through well-reasoned choices of the residues to include, ideally calibrated against experimental data.
9. Decide whether the methodology is accurate enough to allow the use of calculated energies for all steps of the mechanism or if the use of some empirical data is preferred/necessary.
10. If a simple cluster model is thought or shown to be inadequate, extend the model, most often by using hybrid QM/MM methods to incorporate the effects of the surrounding protein and solvent.
11. Decide whether dynamical effects are of interest. If so, design a Born–Oppenheimer Molecular Dynamics protocol using either a cluster or QM/MM.
12. Decide whether entropic effects are likely to be important (for example if charged species are released to the solvent) and, if so, decide on whether a quantum chemical approach (calculating the partition function within a harmonic-oscillator approximation) may be used or whether a molecular dynamics-based approach (e.g., free-energy perturbation theory) should be used to properly sample phase space.
13. For an MD approach with QM/MM design a protocol (preequilibration with a classical force field, substrate docking, boundary conditions, number and length of sampling “windows,” etc.).
14. In all of the above, one of the most delicate choices is that of a suitable reaction coordinate.
15. For all approaches, analyze the results in terms of structural, energetic, and dynamic aspects and using the tools of interpretational-DFT as appropriate.

Although we do not pretend that the above protocol is unique or complete in all aspects, we will use it as a guide in the following sections, in the hope that it will be helpful to novices as they gain experience in what is, in the end, a complex field of computational chemistry/biology.

2 DFT and ADFT

2.1 Methodology

We take as a starting point the Kohn–Sham equations:

$$\left[-\frac{1}{2}\nabla^2 + v(r) + \int \frac{\rho(r')}{|r-r'|}dr' + v_{xc}(r) \right] \varphi_i(r) = \varepsilon_i \varphi_i(r) \quad (1)$$

with the operators in parentheses being, respectively, the kinetic energy of the noninteracting Kohn–Sham reference system, the external potential (nuclear-electron attraction), the classical coulomb potential felt at the point \mathbf{r} due to the electronic charge distribution (including the self-interaction), and the exchange-correlation potential. φ_i is a Kohn–Sham orbital and ε_i is the corresponding Kohn–Sham orbital energy. The exchange-correlation potential is the functional derivative of the Kohn–Sham exchange-correlation energy with respect to a variation of the density:

$$v_{xc}(\mathbf{r}) = \frac{\delta E_{xc}[\rho(\mathbf{r})]}{\delta \rho(\mathbf{r})}. \quad (2)$$

The Kohn–Sham equations are exact but, of course, for practical calculations approximations have to be made and these will determine the accuracy, the speed, and the interpretability of approximate KS-DFT methods.

The most important choice is that of the exchange-correlation functional. Here there is a wide variety of options. Some standard choices are functionals of the Generalized Gradient Approximation (GGA) (which we favor because of their computational speed when used within the Auxiliary DFT approach and their overall good level of accuracy—see below; PBE [6–8] is a common choice) and the hybrid functionals that involve a component of Hartree–Fock exchange, the B3LYP functional [9, 10] providing the prototypical example:

$$E_{xc}^{B3LYP} = E_{xc}^{VWN} + a_0(E_x^{HF} - E_x^{VWN}) + a_x(E_x^{Becke} - E_x^{VWN}) + a_c(E_c^{VWN} - E_c^{LYP}), \quad (3)$$

where $a_0 = 0.20$, $a_x = 0.72$, and $a_c = 0.81$.

Other functionals have been specially parameterized for various applications, notably by the Truhlar group [11]. GGA functionals of the OPTX type have had considerable success in the area of bio-organic complexes [12–16].

Once the functional has been chosen, one needs to solve the Kohn–Sham equations, typically using some sort of basis set, although numerical approaches have seen some use [17]. Here we will focus on the use of Gaussian basis sets which are used in a number of software packages (Gaussian [18], NWChem [19],

Q-Chem [20], etc.) and also in our own code deMon2k [21]. We further outline the use of fitting functions [22] and the Auxiliary-DFT (ADFT) [23] methodology because it provides a real computational advantage provided (for the moment) that pure, nonhybrid, functionals are employed. Of course one has to address the accuracy question, both as concerns the inherent errors of approximate functionals and the numerical errors associated with the choice of basis sets, auxiliary basis sets, numerical integration grids, etc. These issues will be addressed in the discussion of the applications below, but first we complete the overview of the methodology with a discussion of density fitting and the ADFT methodology.

The use of Gaussian functions in DFT was pioneered by Dunlap, Connolly, and Sabin [22] who, in 1982, formulated the LCGTO-X α method that incorporated a variational fit of the coulomb terms. With the usual LCAO approximation, and using a general form for the exchange-correlation energy rather than X α (an early density functional that uses a local density approximation for exchange, incorporating a parameter α that, arguably, accounts for correlation to a certain extent), one can write the total energy in terms of the density matrix:

$$E = \sum_{\mu, \nu} P_{\mu\nu} H_{\mu\nu} + \frac{1}{2} \sum_{\mu, \nu} \sum_{\sigma, \tau} P_{\mu\nu} P_{\sigma\tau} \langle \mu\nu || \sigma\tau \rangle + E_{xc}[\rho(\mathbf{r})]. \quad (4)$$

Here $P_{\mu\nu}$ and $P_{\sigma\tau}$ denote elements of the density matrix and $H_{\mu\nu}$ an element of the core Hamiltonian (kinetic energy and nuclear attraction). The basis functions that are used to expand the molecular orbitals are indicated by μ, ν, σ and τ . The (up to) four-center electron repulsion integral is represented by:

$$\langle \mu\nu || \sigma\tau \rangle = \iint \frac{\mu(\mathbf{r}') v(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} \sigma(\mathbf{r}) \tau(\mathbf{r}) d\mathbf{r}' d\mathbf{r}. \quad (5)$$

The key development came from the realization that if one could fit the density using an auxiliary basis set then in practice one index could be saved, turning the problem from a basically N^4 scaling, where N is the size of the orbital basis set, to N^2M where M is the size of the auxiliary basis set. Writing

$$\tilde{\rho}(\mathbf{r}) = \sum_k x_k k(\mathbf{r}) \quad (6)$$

along with the expansion of the density using the density matrix:

$$\rho(\mathbf{r}) = \sum_{\mu, \nu} P_{\mu\nu} \mu(\mathbf{r}) v(\mathbf{r}) \quad (7)$$

and minimizing the following error

$$\varepsilon_2 = \frac{1}{2} \iint \frac{[\rho(\mathbf{r}) - \tilde{\rho}(\mathbf{r})][\rho(\mathbf{r}') - \tilde{\rho}(\mathbf{r}')] }{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r} d\mathbf{r}' \quad (8)$$

leads to the following equation:

$$\frac{\partial \varepsilon_2}{\partial x_k} = - \sum_{\mu,v} P_{\mu,v} \langle \mu v || k \rangle + \sum_l x_l \langle l || k \rangle \equiv 0 \forall k. \quad (9)$$

Solution of this equation is often called the Resolution of the Identity and is now also common in wave function methods. A second auxiliary set was introduced for the exchange-correlation potential; however, this used a least-squares procedure on a numerical grid and it was not done variationally. This level of theory is identified by the keyword BASIS in our deMon2k software since the density matrix is used (within a min-max scheme).

Going one step further in terms of potential speed, Köster and coworkers [23] studied the use of the auxiliary density throughout the variational process. The corresponding auxiliary DFT (ADFT) energy expression is the following:

$$E = \sum_{\mu,v} P_{\mu v} H_{\mu v} + \sum_{\mu,v,k} P_{\mu,v} \langle \mu v || k \rangle x_k - \frac{1}{2} \sum_{k,l} x_k x_l \langle k || l \rangle + E_{xc}[\tilde{\rho}(\mathbf{r})], \quad (10)$$

where, in practice, as implemented in deMon2k, the orbital basis sets are contracted Cartesian Gaussians while the auxiliary sets are primitive Hermite Gaussians. In Eq. (10) the first term represents the core energy, the second the coulomb repulsion energy of the electrons using the density matrix once and the fitted density once, the third term complements the second in the variational fitting procedure and involves only the fitted density and the final exchange-correlation term also involves only the fitted density. The derivatives of this energy expression with respect to the density matrix elements define the ADFT Kohn-Sham matrix elements:

$$K_{\mu v} = \frac{\partial E}{\partial P_{\mu v}} = H_{\mu v} + \sum_k \langle \mu v || k \rangle x_k + \frac{\partial E_{xc}[\tilde{\rho}(\mathbf{r})]}{\partial P_{\mu v}}. \quad (11)$$

After some algebra and using the variational property of the density fitting the following equation is derived:

$$K_{\mu v} = \frac{\partial E}{\partial P_{\mu v}} = H_{\mu v} + \sum_k \langle \mu v || k \rangle (x_k + z_k), \quad (12)$$

where z_k are exchange-correlation fitting functions. The important thing about Eq. (12) is that the Kohn-Sham matrix elements are independent of the density matrix elements. As a result, only the fitted density (and, in the case of GGA functionals, the corresponding density derivatives) have to be numerically calculated on a grid. These quantities scale linearly by construction and so the grid work is reduced considerably. In addition, the use of shared auxiliary function exponents between the two auxiliary sets dramatically decreases the number of expensive exponential function evaluations, resulting in very favorable computing times.

2.2 Applications

2.2.1 Cluster Calculations for Histone Lysine Methyltransferase

We start the discussion of biomodeling using straight DFT with a brief overview of some recent work from Per Siegbahn and coworkers using the recent review of Siegbahn and Himo as the leading reference [24]. Siegbahn has been a champion for (properly converged) cluster models for a number of years and his well-chosen applications have shed considerable light on classes of enzymes for which reliable cluster models can be formulated. With recent increases in computer power and program efficiency, cluster models with upwards of 150 atoms are now feasible; for some, but not all, types of reactions, these are able to capture the essence of the relevant free-energy profiles.

Siegbahn's recent work focuses on the B3LYP hybrid functional for which he claims "In spite of numerous attempts, it has been difficult to improve the accuracy beyond that of this functional." Of course, that does not relieve us from the responsibility of addressing the question of accuracy for a given reaction. Siegbahn cites three sources of inaccuracy in DFT methods (1) the self-interaction error, (2) the inherent limitations of a single-determinant approach, and (3) the lack of van der Waals interactions in the usual functionals, including B3LYP. According to Siegbahn, errors (1) and (2) (using a spin-unrestricted formalism) tend to cancel and this can at least partially explain the relatively good performance of B3LYP. He also discusses the effects of varying the amount of exact exchange in the hybrid functionals, leading to the rule of thumb that if the results do not change very much when the amount of exact exchange is decreased from 20 % to 15 % the methodology seems to be reliable. While this seems less than an ideal procedure from an "ab initio" perspective, such procedures are necessary at the present stage of advancement in the search for more accurate and generally applicable functionals. The body of work using B3LYP indicates that it can provide results of useful accuracy if it is applied with due caution. We will discuss the third source of error in the next section on DFT-D methodologies. We only indicate here that the empirical corrections for dispersion-like interactions can now be readily incorporated and that they lead in many cases to significant improvement. They should become the default option.

Two further aspects of Siegbahn's cluster approach are (1) a coordinate-locking scheme and (2) the use of a polarizable continuum method (PCM) to model the electrostatic effects of the surrounding medium (protein and solvent). The coordinate-locking scheme fixes the coordinates of key atoms on the periphery, hence preventing large artificial movements of the active site groups. For very small cluster models this approach can lead to artifacts, but as the cluster grows it behaves better and better (and ultimately becomes unnecessary). The PCM approach assumes that the surroundings can be represented as a homogeneous polarizable medium with a dielectric constant that has to be chosen (often $\epsilon = 4$ gives good results but in some studies ϵ is varied to gauge the sensitivity of the results to this parameter).

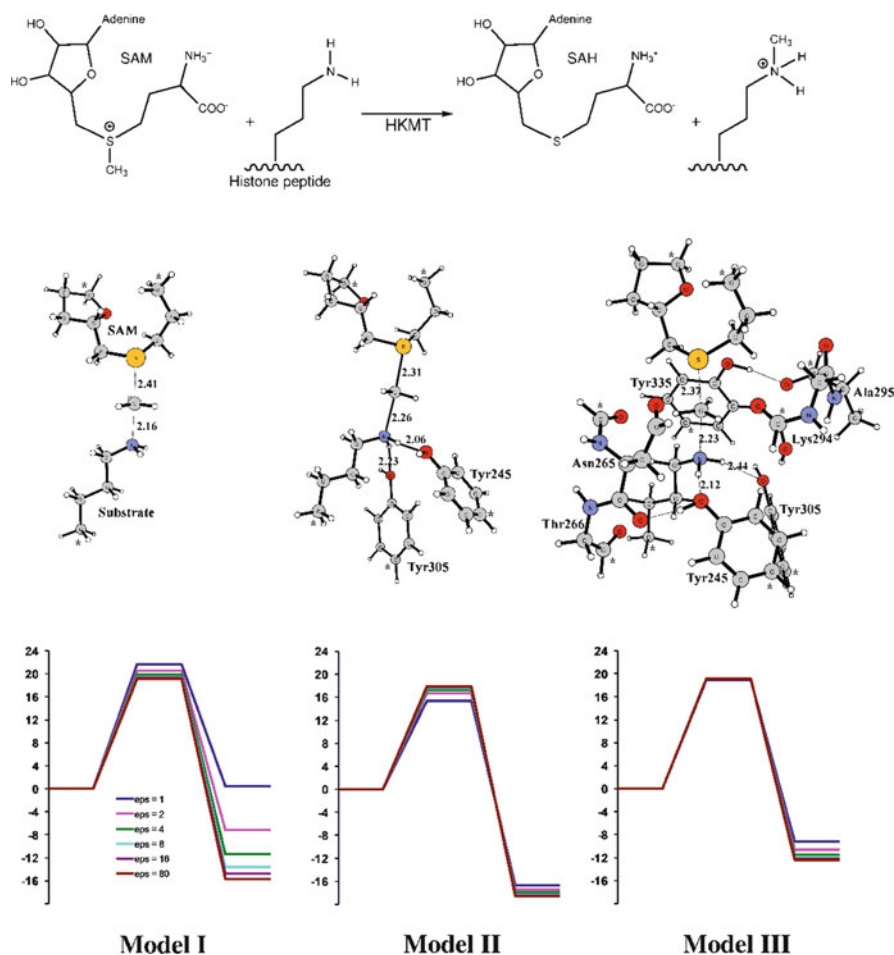


Fig. 1 *Top*: Reaction mechanism for the methylation of the Lysine side chain of histone by *S*-adenosylmethionine using HKMT as the catalyst. *Middle*: The three cluster models used. *Stars* indicate fixed atoms in the coordinate-locking scheme. *Bottom*: Potential energy profiles (kcal/mol). Results for various values of the dielectric constant are shown in *different colors*. Reproduced with permission from [24]

Clearly, the more protein residues and waters of solvation that are included in the cluster model explicitly, the less critical will it be to resort to this somewhat doubtful PCM approximation.

We turn now to histone lysine methyltransferase (HKMT) which catalyzes the methylation of the N-terminal histone tail of chromatin using the *S*-adenosylmethionine (SAM) cofactor as the methylating agent (see Fig. 1).

Models of increasing size were used; Model I (46 atoms) contained only truncated models of SAM and the substrate; Model II (72 atoms) also contained two important tyrosine residues that form hydrogen bonds with the amino group of

the substrate and Model III (132 atoms, Middle part of Fig. 1) contains additional groups that form a ring around the substrate and interact with the transferred methyl group. The total overall charge on the models is +1. Considering only this particular reaction one has a “quantum-chemistry friendly” situation, where all of the reaction steps (reactants, transition state, and products) take place in a reasonably similar environment (provided by the interacting residues of Model III, for example) and a single positive charge is transferred from the cofactor to the substrate. Geometries of the critical points were calculated at the B3LYP/6-31G(d,p) level with single-point calculations of the energies using a larger 6-311 + G(2d,2p) basis. Solvation effects were calculated as single points at the same level as the geometry optimizations using a range of dielectric constants (2, 4, 8, 16, 80). As expected the larger models showed little dependence on the value of ϵ chosen (essentially identical values for the entire range of ϵ for the activation energy from the reactant side and roughly a range of 4 kcal mol⁻¹ for the overall exothermicity of the reaction) (see the bottom part of Fig. 1).

It was found that the transition-state geometries for the three models were very similar which is of practical importance, allowing the TS search to be performed for small models and then only having to refine the structures for the larger models. Moreover, the calculated energy differences are not very different in the different models. In fact all of the barriers are close to the experimental barrier of 20.9 kcal mol⁻¹. The solvation effects saturate quickly; 132 atoms in this case seem to be well converged, a fortunate circumstance for this reaction and a good number of other reactions, because such cluster sizes are well within the range of current possibility. It was emphasized that geometry optimization is crucial; taking raw crystallographic geometries usually leads to wrong energy profiles.

Some final comments on the entropy (free energy) are in order. Because of the coordinate-locking scheme, there are a number of small imaginary frequencies ($<30i$ cm⁻¹) which, although they do not affect the energetics significantly, do render the calculation of the harmonic frequencies and their associated entropy contributions inaccurate. So the reported energies correspond to enthalpies and not free energies. For the present HKMT case, Siegbahn refers to work by Hu and Zhang [25] to show that the entropy effects are quite small. Hu and Zhang used a QM/MM methodology (B3LYP with single-point MP2) with free-energy perturbation theory (see below) for the MM contributions to the entropy and harmonic frequencies for the QM part (now possible because there are no frozen atoms). The QM part contained 66 atoms, SAM and the lysine side chain only. The free energy barriers are found to be only about 1.1 kcal mol⁻¹ lower than the potential energy barriers, confirming the validity of neglecting entropy for this particular reaction, which is well contained within the QM part of the system. We will see below that this is not a general result. For some reactions, such as those involving DNA or RNA polymerase, entropy effects can be very large; they definitely cannot be neglected.

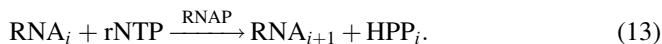
For other examples of the valid use of the finite cluster model, we refer the reader to [26–31].

2.2.2 Cluster Calculations for RNA Polymerase

In the central dogma of biology (“DNA to RNA to proteins”) the first step transcribes the genetic code from DNA to messenger RNA. This task is accomplished by a marvelous nano-molecular machine, the RNA Polymerase (RNAP) enzyme, which, once initiated, processes a DNA template strand, adding successive matching nucleoside triphosphates to a growing chain of m-RNA. RNAP is a complex multidomain protein containing about 3,500 residues and 28,000 nonhydrogen atoms [32]. One of the great triumphs of modern protein crystallography is surely the elucidation of many aspects of the mechanism for transcription for which Roger Kornberg was awarded the 2006 Nobel Prize in Chemistry. In principle, the work of Kornberg and the others who have accomplished this magnificent task is simple—isolate and crystallize proteins that are ligated by various intermediates along the multistate reaction path and derive their structures by X-ray analysis. In this way we get “flash-frozen” snapshots of the reaction mechanism. This has been so successful that a movie of the process has been made [33] which may be viewed at <http://www.lmb.uni-muenchen.de/cramer/pr-materials>. We show some snapshots from the movie in Fig. 2.

This is clearly a more complex reaction than that of HKMT just described. Of the seven steps shown in the movie all but one involve conformational changes of the enzyme that are more in the domain of Molecular Mechanics than in that of quantum chemistry (we are studying these conformational steps, but they are not the topic of the present chapter). The sole exception is step 4, catalytic incorporation, which actually involves a multistep mechanism of chemical reactions. This is where theory and computation have to step in to help elucidate the mechanism. The first steps of the process involve the construction of cluster models for the calculation of relevant portions of the potential energy surface corresponding to proposed reaction steps. Several key choices have to be made for which reactions to consider.

The basic enzymatic function of RNAP is the transfer of the nucleotidyl motif from the rNTP substrates to the hydroxyl at the 3'-end of the nascent RNA transcript. The nucleotidyl transfer reaction can be simplified as



The catalytic center of RNAP includes the binding site for the 3'-end of RNA and the insertion site for the incoming rNTP. In the nucleotidyl transfer reaction, the 3'-OH group in the sugar ring of the RNA primer reacts with the α -phosphorous atom of a ribonucleoside triphosphate by nucleophilic attack, then the $\text{P}_\alpha\text{-O}_{\alpha\beta}$ bond is broken and pyrophosphate (PP_i) is released. Thus, a nucleotidyl addition to the RNA primer is achieved. Structural and biochemical data have shown that the active centers of all polymerases share certain common features: a pair of metal ions (normally divalent magnesium ions Mg^{2+}) and three universally conserved carboxylates. The two-metal-ion mechanism for the nucleotidyl transfer reaction

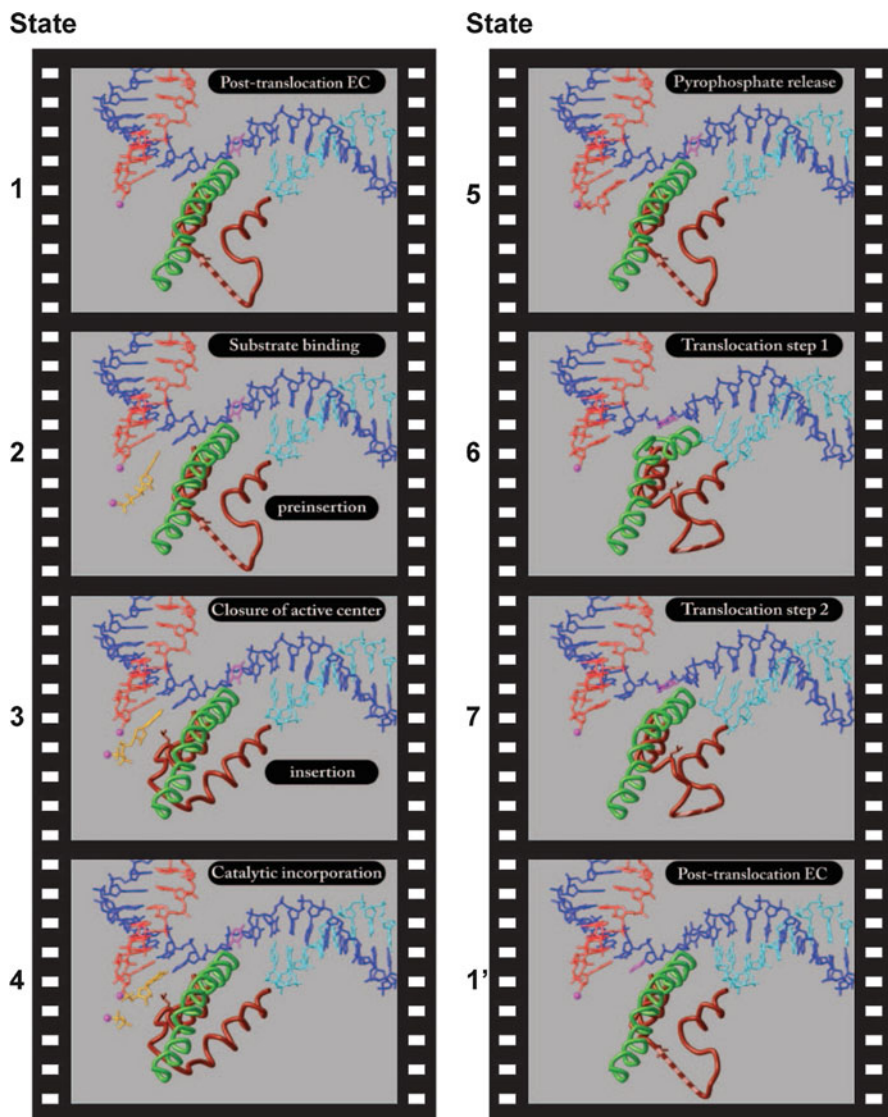


Fig. 2 Snapshots from the NAC (Nucleotide Addition Complex) movie have been depicted that correspond to different functional states of the EC. Reproduced with permission from [33]

was proposed by Steitz [34]. A schematic diagram of the ternary elongation complex for yeast RNA polymerase is shown in Fig. 3.

We have considered several different detailed reaction mechanisms for the nucleotide addition [36]. In all of them, the 3'OH group has to be deprotonated, either by passing its proton to another residue of the enzyme or to a solvent water

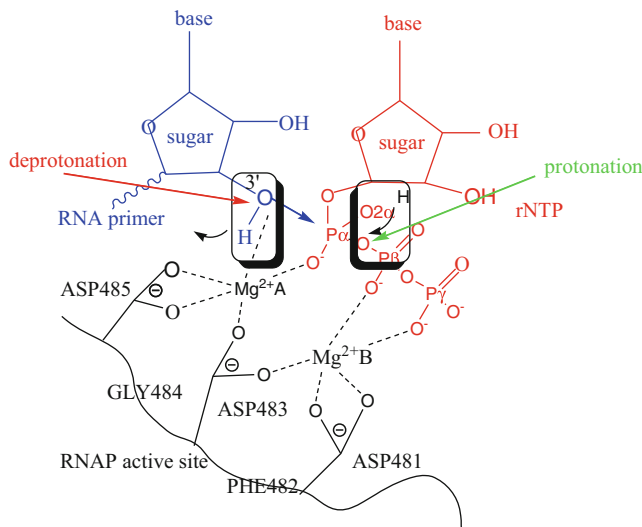


Fig. 3 The two-metal-ion mechanism and two-proton transfer model for the nucleotidyl reaction of yeast RNA polymerase II [34, 35]. The ternary elongation complex consists of three parts: the growing RNA transcript (*blue*), the incoming rNTP (*red*), yeast RNA polymerase active center (*black*) that is mainly composed of two divalent magnesium ions, Mg^{2+} A and B, and three conserved amino acid residues, aspartates. The two protons are directly or indirectly transferred at two sites proposed according to deuterium isotope effect experiments. One site is the primer RNA 3' terminus (*left square*), where the RNA primer 3'OH must be deprotonated by the incoming rNTP, a nearby residue or a mediated water molecule before nucleophilic attack. The other is the α - and β -phosphate bridging oxygen atom (*right square*), where the pyrophosphate should be protonated by a nearby residue or a mediated water molecule before it leaves. The three aspartate residues ASP481, 483, 485 connected by PHE482 and GLY484 conjugate with two magnesium ions to form the active center of yeast RNA polymerase

molecule or directly to the departing pyrophosphate. One of these mechanisms is illustrated in Fig. 4.

Initial molecular dynamics simulations were performed in order to derive a plausible starting structure for subsequent optimization with DFT. The QM model includes a simplified RNA primer (a sugar ring with 2' and 3'OH groups), an incoming nucleotide GTP substrate, two divalent metal cations, Mg^{2+} A and B, a solvent water molecule, and three universally conserved Aspartate residues (Asp481, Asp483, Asp485) where Asp481 and Asp483 are connected by PHE482, Asp483 and Asp485 are connected by Gly484. The three conserved aspartate residues form a lotus-like complex so that they are quite robust during geometry optimizations. Our previous work showed that the simplified aspartate residues (formic acid $HCOO^-$ or acetic acid CH_3COO^-) changed a lot during geometry optimizations [37].

We found that the barrier height of direct proton transfer from the 3'OH of the RNA primer to the $O2\alpha$ of rNTP is higher than that for proton transfer from the 3'OH of RNA primer to water. Thus, the latter model will be discussed in the following.

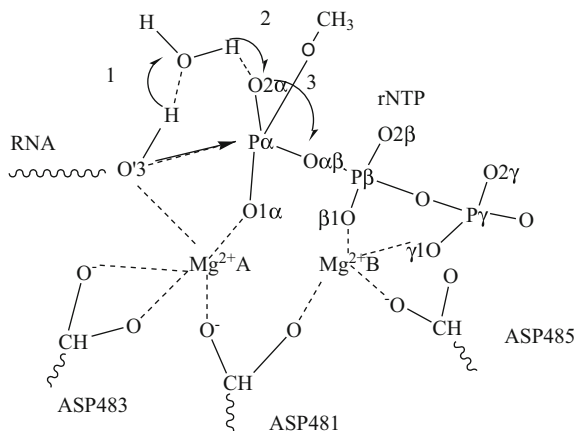


Fig. 4 The proposed nucleotidyl transfer reaction mechanism. Step 1: The proton of the 3'/OH of RNA primer transfers to a solvent water and a proton on water transfers to O2 α of α -phosphate simultaneously; Step 2: the proton of the O2 α atom rotates to the side of β -phosphate; Step 3: the 3'O atom performs a nucleophilic attack at the α -phosphorus atom of the α -phosphate; Step 4: the P α -O3 β bond of the intermediate cleaves to form a phosphodiester bond and the proton on O2 α migrates to O3 β

The model consists of two Mg²⁺, three conserved aspartate residues, one ribose, the simplified RNA primer, a simplified rNTP, and a water molecule that is located between the growing RNA primer and the incoming rNTP and is closest to the 3'/OH of RNA primer and the α -phosphorus atom of the incoming rNTP. The model has 94 atoms with a total charge of -3 . This is due to the three conserved aspartate residues (-3 charge) of the yeast RNA polymerase II active site, the triphosphate (-4 charge) of the incoming rNTP, 2 Mg²⁺ ($+4$ charge) at physiological pH (7.2–7.5). Considering the physiological pH in the experimental condition and the pK_a of triphosphate, the incoming rNTP is thought to be in a deprotonated state in this study.

All density functional (DFT) calculations were performed using the deMon2k (version 2.4.4) program. We carried out full optimizations on all structures, reactant, intermediates, transition states using the PBE exchange-correlation functional with the basis set DZVP-GGA and auxiliary basis set GEN-A2. No constraints were imposed on any atom of these systems.

We proposed the following detailed nucleotidyl transfer reaction mechanism for yeast RNA polymerase II, shown in Fig. 4. The proton of the 3'/OH first transfers to the O2 α of α -phosphate via a solvent water molecule, then one of the water molecule's protons transfers to the bridging phosphate O3 β atom. Note that the water molecule is located by our CHARMM molecular dynamics (MD) simulations.

The calculated potential energy profile for the reaction is shown in Fig. 5.

If taken at face value the analysis of these four steps could provide detailed insight into the various proton transfer and bond making and -breaking steps and the

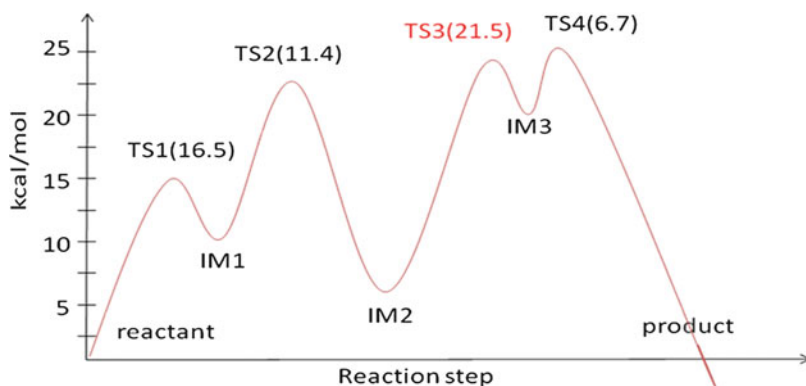


Fig. 5 The calculated potential energy curve for the PBE/DZVP-GGA method

role of water molecules. Indeed, we have performed such an analysis [36] but we have chosen not to publish it yet because of potentially large effects of the protein/solvent environment and, especially, the probably large entropic effects, based on analogous QM/MM calculations for DNA polymerases that are discussed in the QM/MM section below.

3 DFT-D

Biological systems are large, solvated in water and comprise domains with low overlapping density, which are only weakly interacting with each other. They belong to what has generally been called “soft matter.” Simultaneous charge fluctuations in the various nonbonded parts of soft matter systems generate attractive van der Waals interactions that are nonlocal correlation effects. These correlation effects are naturally accounted for in correlated wave function methods including a sufficiently high level of electron correlation (the basic dispersion interaction is already accounted for at the MP2 level), but these methods are relevant only for small systems and they are mainly used for benchmark calculations.

Despite the fact that the exact density functional contains van der Waals correlation, the inability of local (LDA) or generalized gradient approximations (GGA) or even more sophisticated meta-GGA functionals to treat properly long-range interactions is now recognized. Significant progress in making DFT more appropriate for weakly interacting systems has been made with long-range density functional theory, applying to nonoverlapping [38–42] and also overlapping electron densities [43, 44]. Molecular polarizabilities, from which the dispersion interaction energy can be calculated, have been estimated from time-dependent calculations [45] or from the instantaneous dipole moment of the exchange hole [46–48]. New combinations and parameterizations of GGA [49, 50] or meta-GGA [51] exchange-correlation functionals have also been proposed to allow the incorporation of the long-range part of electron correlation.

Based on the treatment of asymptotic van der Waals forces [52], a van der Waals correlation density functional has been proposed by Dion et al. [53] and applied to solid and biosystems [54]. The use of this correlation functional for large systems has been limited due to the double spatial integration for calculating the correlation energy. However, a very recent implementation algorithm has been proposed to overcome this bottleneck [55].

Because it has been known for a long time that van der Waals interaction energies are important for large systems, a more empirical approach was first used by Wu and Yang [56], adding an empirical dispersion energy correction (E_{disp}) to the usual DF energy (DFT-D approach) based on atomic C_6 coefficients, derived from molecular C_6 coefficients. The idea of using a correction term originates from Hartree–Fock-based studies [57–59] and from Elstner et al. [60] for correcting the Self-Consistent-Charge DFT-Tight Binding method. Whereas the nonlocal character of the Hartree–Fock exchange provides a correct description of the long-range intermolecular forces, the DF approaches based on the local density expansion are not strictly applicable and their performance depends on the particular exchange-correlation functional. A damping function has to be used in order to set E_{disp} to zero as the electron densities overlap. Different analytic forms were tested [56, 61]. Based on a test set of small van der Waals systems, proper scaling factors of E_{disp} were proposed for different XC density functionals [62]. Instead of scaling the empirical dispersion energy, Jurečka et al. [63] adopted a global scaling factor of the atomic van der Waals radii, optimized for a training set of noncovalent complexes. More recently, inclusion of higher order correction terms, involving C_8 and C_{10} coefficients associated with an adequate damping function, has been proposed by Johnson and Becke [48], aiming at a better description of π stacked systems.

In recent years more sophisticated DFT-D dispersion corrections have been proposed and benchmarked on large test sets, allowing comparison with a very complete set of XC functionals [64]. Improving the use of atomic C_6 coefficients for dispersion corrections, a density-dependent energy correction for long-range dispersion has been proposed, based on the Becke–Johnson exchange hole dipole moment formalism [65] and an extended Tang–Toennies damping function [66] accounting for charge-overlap effects [67]. This formalism has been benchmarked for different functionals and test sets [68].

In our DFT-D applications, the dispersion term, limited to the dipole–dipole contribution to the dispersion interaction energy,

$$E_{\text{disp}} = - \sum_{i=1}^N \sum_{j=i+1}^{N-1} \frac{C_6^{ij}}{r_{ij}^6} f_{\text{damp}}(r_{ij}) \quad (14)$$

is expressed as the sum of the i,j atom-pair contributions in an N -atomic system with an interatomic distance r_{ij} . In the DF approach including a damped empirical correction for the van der Waals interactions, E_{disp} is simply added to the usual DF energy and ∇E_{disp} is added to the DF energy gradient. The present E_{disp} expression,

limited to the first C_6/R^6 term, contains implicitly most of the physical intermolecular dispersion via the fitting of the atomic C_6 coefficients to molecular C_6 values, obtained from a training set of 44 pairs of molecules including hydrocarbons and other small organic compounds [56].

The dispersion coefficients

$$C_6^{ij} = \frac{2C^i C^j}{(C^i + C^j)} \quad (15)$$

are computed from the atomic C_6^i , as proposed by Wu and Yang [56], but averaged over the possible hybridization states of the atoms, which are 2.845 and 26.360 a.u. for H and C atoms, respectively. The damping function used is

$$f_{\text{damp}}(r_{ij}) = \frac{1}{1 + e^{-\alpha \left(\frac{r_{ij}}{r_0} - 1 \right)}} \quad (16)$$

with $\alpha = 23.0$ [24, 56] and with r_0 being the sum of the atomic van der Waals radii [69, 70].

The dispersion correction energy, its gradient and second derivative were implemented in the deMon2K program [29], allowing geometry optimization, Born–Oppenheimer molecular dynamics, and vibrational frequency analysis. We used a nonscaled empirical dispersion term, unlike previously mentioned DFT-D calculations [62, 63].

In fact, scaling E_{disp} has been used to compensate the erroneous behavior of GGA exchange functionals which either show an unexpected attractive tendency in the van der Waals region (and thus avoids a double counting of the “dispersion”) or, in contrast, have a too strong repulsive slope in this region. The latter trend is mainly displayed by the Becke exchange functional [71] whereas PW91 [72] and PBE [6] exchange functionals are responsible for the former.

Exchange-only calculations of alkane dimers illustrate this problem, as well as the related question of choosing a correlation functional which can compensate the exchange repulsion at nonbonding distances that correspond to short-range van der Waals attractive interactions between hydrophobic alkyl chains [73]. Taking as an example two butane molecules, one can analyze the effect of the exchange functional on their interaction energy as a function of distance: the revised version of PBE, i.e., revPBE [74] and TPSS [75] repulsive curves are close to HF for long distances, i.e., between 4.4 and 5 Å where they reach zero, insuring no double counting of any correction for “dispersion” effects. In contrast, Becke exchange leads to a much stronger repulsion, whereas PBE is attractive for distances beyond 4.4 Å. Similar conclusions have been reported for the benzene–benzene interaction [53].

On the other hand, correlation interaction energies also have to be analyzed in combination with the exchange interaction energy term. As expected, correlation interaction becomes more attractive when decreasing the separation between the two butane monomers. In this test example, PBE exchange and correlation

functionals without the empirical dispersion correction lead to a weakly bound butane dimer ($-0.8 \text{ kcal mol}^{-1}$ at a separation of 4.5 \AA), whereas a PBE-D (including E_{disp} correction) calculation leads to a much too large stabilization of $4.1 \text{ kcal mol}^{-1}$ compared with the MP2 and CCSD(T) interaction energy of $2.7 \text{ kcal mol}^{-1}$. However, combining the PBE correlation with the revPBE exchange yields a large repulsive interaction in the range of $4\text{--}4.4 \text{ \AA}$ which cannot be compensated by the damped dispersion correction. In contrast, combining the “semiempirical” revPBE exchange, adjusted on atomic exchange energies, with the LYP correlation functional [76], self-interaction (SI) free by construction, allows a good balance of repulsive and attractive effects between the two subsystems. In fact, the revPBE exchange leads to less SI error than PBE (5 times less for the H atom). The PBE exchange SI error is mostly compensated by the SI error of the PBE correlation itself. Therefore, combining the revPBE exchange with the PBE correlation is much less appropriate than combining it with the LYP correlation. This strategy is not based on fundamental arguments but on an empirical analysis of XC functional behavior. Different applications on saturated and aromatic hydrocarbon compounds have shown that the revPBE-LYP exchange-correlation functional augmented with the empirical van der Waals correction gives a good description of soft matter containing alkyl chains, without being considered as the panacea for a general and accurate estimate of long-range interactions [73]. In a similar vein, a recent study of exchange-only interaction energies of small molecules led to the proposal that a re-parameterized PW86 exchange functional can be recommended for its performance in yielding no spurious intermolecular binding when augmented by an empirical dispersion term [77].

The interest of taking dispersion interactions into account for soft matter systems can be illustrated by the thermal properties of lipids in cell membranes, which get more rigid at a given temperature when their alkyl chains get longer. This property is exploited by natural mechanisms for maintaining cell membranes fluid but not leaky, either increasing the number of long chain lipids or making them shorter. This very important property is related to the increasing dispersion-like stabilizing interaction of alkane chains when increasing the number of CH_2 units in the chains. The estimated value, corresponding to the most favorable distance of 4.10 \AA between two chains, has been found to be $-0.81 \text{ kcal mol}^{-1}$ from DFT-D, $-0.86 \text{ kcal mol}^{-1}$ from MM, and $-0.88 \text{ kcal mol}^{-1}$ from the G3 (CCSD(T)) results (butane and hexane) [73]. Calculations using MP2 (limit) lead to a slightly overestimated value of $0.97 \text{ kcal mol}^{-1}$, as usually observed [78].

It is worth noting that dispersion energy also contributes to the total energies of the individual n -alkane monomers. This contribution varies from -3.7 for butane to $-14.8 \text{ kcal mol}^{-1}$ for dodecane, with a regular decrease of $-1.4 \text{ kcal mol}^{-1}$ per CH_2 unit. The internal E_{disp} value is thus about 50 % larger than the dispersion contribution to the dimer binding. This shows that, in the alkane monomers, the damped dispersion at mid-range contributes also to the total energies. However, the amount of stabilization provided by the internal dispersion in an alkane molecule is very small with respect to the other energy contributions (0.55 % of the correlation energy for n -butane, 0.57 % for isobutane, 0.70 % for n -octane, and 0.81 % for

isooctane). This explains why this E_{disp} term cannot compete with some other effects, such as those governing the branched/linear alkane relative stabilities [79].

Similar intra- and inter-molecular dispersion effects have been estimated in the case of phosphatidyl choline (PC) lipids, with two acyl chains of 12 (dilauroyl, DL), 14 (dimyristoyl, DM), and 16 (dipalmitoyl, DP) CH_2 units. The conformational analysis of these lipids has shown the existence of two relative orientations of the chains with close energies, having their carbon backbones in parallel and in perpendicular planes [80].

It is worth noting that the dispersion-type energies differ for these two chain arrangements with a larger stabilization (about $4.5 \text{ kcal mol}^{-1}$) for the parallel arrangement. Indeed, the optimum distance for dispersion between two alkane monomers in a dimer is about 4.2 \AA , which is also the distance found between the PC lipid alkane chains in the “parallel” conformers as compared to the distance of 4.8 \AA for “perpendicular” conformers. However, the shorter distance stabilizing effect is compensated by a structural rearrangement in the glycerol backbone and the balance of the above two effects leads to similar total energies for the two types of chain structures. The comparison of the computed empirical dispersion energies for several lipid isomers with 12, 14, and 16 carbons in their alkyl chains and in the two tail arrangements reveals a regular increase of stability of approximately $3.1 \text{ kcal mol}^{-1}$ per CH_2 unit [81]. This increase of stabilization from DLPC to DPPC is more regular for parallel chains. These results allow one to anticipate qualitatively the thermodynamic behavior of these lipids as well as, more generally, the fluidity of lipid bilayers (membranes): long chains are more stable than shorter chains and will gain more stability in assemblies of lipids. As a consequence, the evolution from ordered (more stable) chains to disordered (less stable) will be less easy for longer chains, needing higher temperature, than for shorter chains. In fact, the transition from ordered to disordered chains, called the main phase transition for lipid bilayers, is measured at -2° , 27° , and 41° C for DLPC, DMPC, and DPPC, respectively.

4 BODFT-MD

4.1 Introduction

The concept of motion in science goes back to Newton [82]. But there was a long time gap until 1959 when a general method for the computer simulation of particle dynamics (500 hard spheres) appeared [83]. Molecular dynamics (MD) simulations using atomic parameters started in the 1970s, studying water [84] and very soon protein conformations and folding, with the early mention of “computer simulations” [85]. About 10 years later, dynamics was introduced into the world of quantum chemistry with *ab initio* molecular dynamics (AIMD) studying the dynamical motion of atoms by solving, as exactly as is required, the entire quantum mechanical electronic structure problem and deriving the forces on the atomic

nuclei using the Hellmann–Feynman theorem. These forces are then used to move the atoms as in classical Newton dynamics. The breakthrough in this domain was the method proposed by Car and Parrinello (CPMD), solving coupled equations of motion for both nuclei and electrons based on DFT potential energy calculations [86]. This work opened perspectives in many areas of science. Converging the electronic structure at every time step, i.e., following the Born–Oppenheimer (BO) potential energy surface was proposed later, bringing the advantage of providing fully converged structures all along the trajectory, which is of great interest for the statistical estimate of dynamical properties [87].

Thanks to the evolution of software and computers, CPMD or BOMD dynamics can be applied now to larger systems than in the 1990s. However, these AIMD calculations are still very time consuming due to the repetition of the self-consistent electronic structure and gradient calculations at every time step, that is generally about 1 fs for BOMD and about ten times less for CPMD. Even if parallelization decreases dramatically the computational effort, long trajectories at the ns scale, which is still very short in biochemistry, are only possible for systems with less than about 100 atoms.

Although questions asked in biology and biochemistry are at a much coarser, macroscopic level, experimental tools and studies have brought accumulated information on molecular structures and mechanisms, reducing the gap with theoretical investigations at the microscopic level. BOMD can thus bring knowledge about the role of relatively fast structural time evolution of properties that are revealed at a macroscopic level. In this perspective, one must stress that *ab initio* methods are not avoidable, because of their accuracy in taking into account the correlation between all degrees of freedom of the system, whatever are its elements and size, which is not possible with interatomic pair potentials. This particular point may be the major reason to perform AIMD on model systems rather than classical MD on more extended ones.

We believe that it is also why the results presented here on single lipid molecules may bring fundamental information on properties of lipid assemblies.

4.2 Properties of Phosphatidyl Choline Lipids

All living systems are made up of cells and all cells are limited by membranes which are built from lipid bilayers. Membranes are held together in water by hydrophobic intermolecular interactions that favor self-assembly of lipid molecules and tend to close bilayers, avoiding holes. Cell membranes include proteins and carbohydrates and modulate the flow of ions and polar molecules. The fluid nature of the lipid membranes is of critical importance. The fluidity of the cell membranes, normally in the liquid crystalline state, is precisely regulated because lipids undergo phase changes in response to temperature. Phospholipids, according to their fatty acid compositions, have a specific main phase transition temperature, T_m , also called the melting temperature, between the gel (ordered)

and liquid crystalline (disordered) states. The melting transition is accompanied by enthalpy and volume changes.

It is worth noting that lipids in membranes are not covalently bound and are moving individually, diffusing laterally inside their layer (about one time per 100 ns) and, even, undergoing a flip from one leaflet to the other, although much more rarely. These data show that individual lipids behave as separate entities, even if they are subject to a collective physics. Moreover, one can expect that intramolecular bonding prevails over intermolecular interactions, at least, for neutral lipids.

4.2.1 Structural Studies of Phosphatidyl Choline Lipids

Experimental X-ray and NMR studies of lipids have recognized that the knowledge of the structure and dynamics of a phospholipid monomer within an assembly is essential for the understanding of the functional role of the bilayer in biomembranes [88–94]. The effects of the intra and intermolecular electronic forces on the structure and energetics of these systems make their study nontrivial both experimentally and computationally. However, there is general agreement concerning structural differences between the fluid-like and the crystal conformers. This fact questioned the relevance of using the single crystal atomic positions for fluid phase simulations wherein the internal monomer properties appear to be better preserved, due to the relatively small intermolecular interactions [91, 92]. The use of isotopically labeled atoms in Fourier transform infrared (FTIR) spectroscopy allowed the assignment of lipid functional group vibrations, the estimate of their shift in water, and the presence of blue-shifted CH₂ stretching vibrations for temperatures above T_m which have been associated with the appearance of gauche conformations in the alkyl chains [95–99].

Among the large number of MD studies devoted to phospholipid structural properties, dihedral angle values [100, 101], tail orientations [102], head group flexibility [103, 104], phase changes [105, 106], hydration effects [107, 108], and evaluation of local order parameters [109, 110] have been explored. A thorough exploration of the conformational spaces of the dilauroyl (DLPC), dimyristoyl (DMPC), and dipalmitoyl (DPPC) phosphatidylcholine molecules was performed using DFT-D, showing, for the three lipids, the existence of a large number of quasi isoenergetic conformers [80, 81, 111, 112]. These conformers differ by the relative position of the two alkyl chains and by different combinations of the backbone torsion angles (Fig. 6). For the three lipids, the backbone rotational conformers share a common geometric profile which includes a balance of attractive, repulsive, and constraint forces between and within specific groups of atoms. The definition of this profile fits with most of the structural characteristics deduced from measured NMR properties of DMPC and DPPC solutions. The calculated vibrational spectra are similar and in very good agreement with experimental data obtained for these PC bilayers. These results support the idea that these molecules preserve their individual molecular structures in their various assemblies. In fact, this conclusion is confirmed by our studies of the dynamical behaviors of DMPC and DLPC [111, 112].

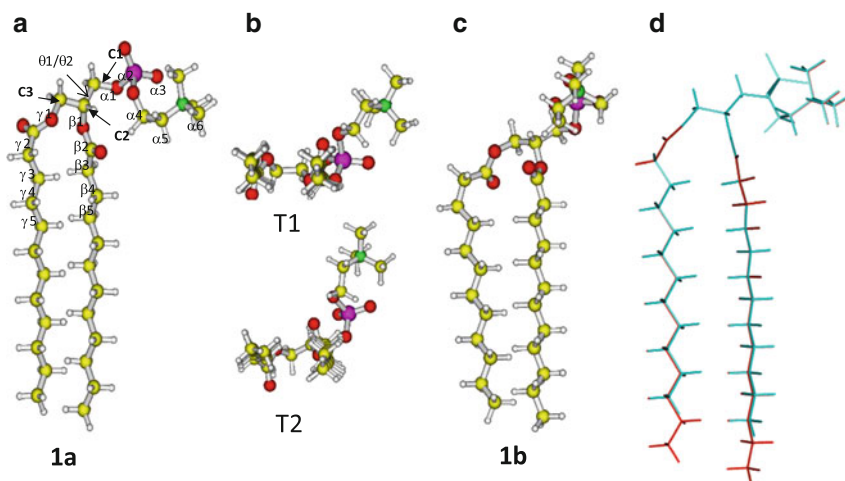


Fig. 6 Structures of phosphatidyl choline lipids: (a) isomer 1a; (b) parallel (T1) and perpendicular (T2) chain conformations; (c) isomer 1b; (d) superposed DLPC (blue) and DMPC (red) isomer 1a optimized structures

4.2.2 Experimental Phase Transitions

Most of the reported structural transformations of PC lipid bilayers with respect to environmental conditions such as temperature, pressure, pH, etc., are associated with isomerizations of the constituent lipid molecules (mostly 14–24 carbon acyl chains). For instance, the transition from the ordered gel to fluid liquid crystalline phase, called the main phase transition, has been related to the melting of the hydrocarbon chains: in the gel phase, phospholipids with all trans alkyl chains are present, whereas in the disordered liquid crystalline phase the most populated conformational states correspond to gauche forms in the alkyl chains.

In the last decade, accurate microcalorimetric and atomic force microscopy experiments have provided detailed information on the lipid melting processes [113–118]. Interestingly, a linear relationship between the isobaric heat capacity and the volume expansion with temperature has been observed for a variety of lipids [115–118] leading to the interpretation that proportional enthalpy and volume changes at the melting transition would be driven by intrinsic structural changes within the lipid molecules, whereas the changes of free volumes and intermolecular interactions could be considered as perturbations [116].

The thermograms of DMPC and longer chain lipid bilayers show a small peak at temperatures well below that of the sharp peak of the main phase transition. The general interpretation is that, when the bilayers are fully hydrated and sufficiently incubated at low temperature, this small endothermic peak (about 1 kcal mol^{-1}) is related to a pretransition between a L_c gel state and a rippled gel state ($P_{\beta'}$), both characterized by all trans chain conformers. This pretransition state is thus not related to molecular structural changes, but to a different packing arrangement of

the molecules in the gel. The pretransition peak is followed by the main phase transition to a liquid crystalline L_α state with predominantly gauche chain conformers of the constituent lipid molecules.

In contrast, the phase pattern of the DLPC thermograms, characterized by one sharp peak followed by a broad band spreading over about 10 K is more complex. This complexity is enhanced by the overlapping of lipids and water solid–liquid phase transitions. Various interpretations of the unusual DLPC thermograms were proposed. From most experiments using various techniques, two phase transitions were assigned to the two peaks [119–123]. These latter studies agreed also in reporting a strong dependence of the position of the first peak on the incubated temperature and the heating rate. Finally, the most recent experiments, using a mixture of water and ethylene glycol and thus avoiding water freezing at the temperature close to the first peak, showed the existence of three peaks [124].

MD simulations of lipid bilayers as a function of temperature have not been extensive. The reliability of force fields is dependent on the accuracy of describing the torsional potential energies of alkanes which prompted improvements of some of the most commonly applied force fields for biomolecules [125] using recent ab initio computations of torsional potential in various trans/gauche *n*-alkanes (up to *n*-decane) [126]. A few recent MD simulations have analyzed the temperature effects on models of PC lipid bilayers, mainly for long chains (16–18 C), with a reasonably successful prediction of the phase transition temperature (error of 12–50 °C) [106, 127–130]. These studies relate the melting temperature to the fast increase of gauche conformers in the alkyl chains.

4.2.3 BODFT-MD of DLPC and DMPC Molecules

In agreement, with the experimental results, the simulated dynamical behaviors of the DMPC [112] and DLPC [111] molecules were found to be quite different. In both cases, BOMD simulations using the DFT-D methodology were performed for a set of temperatures ranging from below to above the experimental T_m for lipid bilayers, i.e., about 295 and 270 K for DMPC and DLPC, respectively.

Quantification of the structural deformations occurring along the molecular dynamics is obtained using the distance-fluctuation criterion initially introduced by Berry et al. [131]. The Berry parameter Δ_B is expressed for a system of N atoms and r_{ij} inter atomic distances between atoms i and j as

$$\Delta_B = \frac{2}{N(N-1)} \sum_{i < j} \frac{\sqrt{\langle \Delta r_{ij}^2 \rangle - \langle r_{ij} \rangle^2}}{\langle r_{ij} \rangle}. \quad (17)$$

The critical value of Δ_B for the solid-to-liquid transition of finite systems has been suggested to be close to 0.1 [132–134].

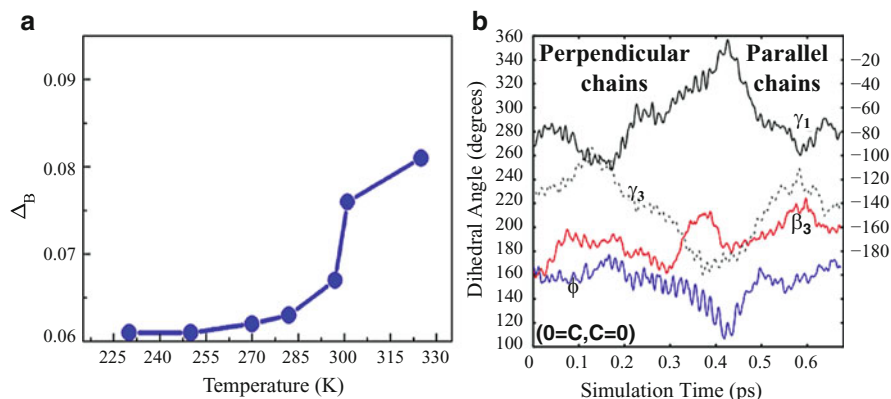


Fig. 7 Dynamics of DMPC: (a) Berry parameter Δ_B evolution with temperature; (b) Short period chain fluctuations at 250 K, showing variation of the four involved dihedral angles. Reproduced with permission from [112]

Focusing on the microscopic structural changes of these molecules as a function of temperature, one can indeed relate them to the existence of only one peak for DMPC and two peaks for DLPC.

The DMPC atomic motions for $T \leq 282$ K correspond essentially to reorientations of the chain CH_2 groups around the chain axes. The planes of the alkyl carbon skeletons are thus fluctuating between parallel and perpendicular (see Fig. 7b). At about 297 K, mostly perpendicular chain orientations remain, fluctuations keeping both chains at a distance comparable to that found at low temperature, but a trans to gauche conformer is formed (kink) at one chain end. Such a transformation also occurs at 301 K in the middle of the chain, increasing the interchain distances and the Δ_B value. Raising the temperature leads to more configurational states introducing more kinks in the chains. In the temperature range explored (230–325 K), no other conformational change was observed than the kink occurrences in the alkyl chains. Figure 7a illustrates the evolution of the DMPC Δ_B parameter with temperature.

The situation is very different in the case of DLPC. It must be noted that DLPC, with 12 C chains is the shortest possible lipid that can assemble in layers. As a consequence of their shortness, DLPC alkyl chains appear much more mobile than those of DMPC, even at low temperature. Moreover, several isomer interchanges occur along the DLPC BOMD trajectories generated at different temperatures, starting from the two isoenergetic isomers illustrated in Fig. 6a, c (1a and 1b). These isomers differ only by the two dihedral angles β_3 and γ_4 . Figure 8a illustrates the concomitant changes of these dihedral angles along the 1a dynamics at $T = 240$ K. When starting from 1b, the first structural change (kink in one chain) occurs at $T = 261$ K. For both 1a and 1b isomers, gauche alkane chain conformers appear at $T \geq 261$ K. It is interesting to note that much larger chain and backbone fluctuations occur at 261 K than at 273 and 285 K, despite the occurrence of a glycerol conformational change in 1b and 1a, at 273

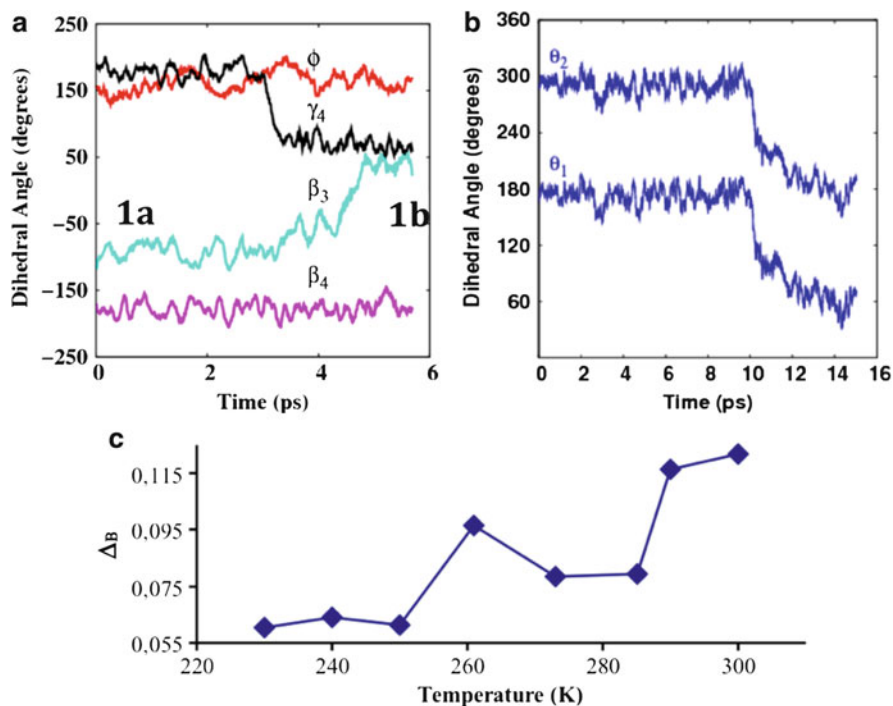


Fig. 8 Dynamics of DLPC: (a) Interchange between isomers 1a and 1b at 240 K; (b) interchange between two glycerol backbone isomers at $T = 273$ K ($\theta_1/\theta_2 = 180^\circ/-60^\circ$) to ($\theta_1/\theta_2 = 60^\circ/180^\circ$); (c) Berry parameter Δ_B evolution with temperature. Reproduced with permission from [111]

(illustrated in Fig. 8b) and 285 K, respectively. At higher temperatures, the number of kinks increases in both chains.

DLPC dynamics thus shows conformational changes associated with the ester groups and the glycerol backbone at various temperatures. This contrast with DMPC is also revealed by the evolution of Δ_B with temperature. The Δ_B value for DLPC (Fig. 8c) is weighted by the populations of isomers 1a and 1b. The Δ_B value jumps to 0.097 at $T = 261$ K, the temperature at which a trans to gauche transformation occurs in the alkyl chains. Increasing the temperature to $T = 273$ and 285 K, an unexpected reduction of Δ_B is observed. Despite the fact that the alkyl chains keep one gauche form as is the case at $T = 261$ K, it appears that the fluctuations of the intra-molecular atomic distances decrease. This peculiar behavior of an isolated DLPC molecule corresponds to the conclusions drawn from both DSC [124] and neutron diffraction measurements [123] attributing the broad area between 272.6 and 281 K to an unusual transition from a gel state to an “intermediate” liquid state (L_x), which shows “a substantial slowing of molecular motions” with respect to the first transition [123].

Using the critical Δ_B value of about 0.1 suggested for finite system order–disorder phase transitions, 261 K can be considered as a DLPC intra-molecular “melting”

temperature. The second intra-molecular transition to a more disordered state occurs at $T = 290$ K as follows from the Δ_B evolution. It is interesting to relate these two peaks of the Berry parameter function to the thermograms of the DLPC bilayers, which also display two phase transitions. The calculated 261 K is about 10 K lower than the experimental temperatures, indicated for the first gel to liquid ($P_{\beta'} \rightarrow L_{\alpha}$ or L_{α}) phase transition. The second peak of the Δ_B function is shifted upward by about 10–15 K when compared to the experimental temperatures of 275–280 K [124].

It is also worth noting that for the DMPC molecule, the distance-fluctuation criterion shows only one abrupt increase at T between 297 K and 301 K related to the main phase transition at $T = 295 \pm 1.5$ K [114, 115]. This confirms that C12 and C14 PC lipids have different dynamics at the molecular level. It is thus most likely to attribute the origins for the “intrinsically different” [123] gel to liquid phase transitions of PC bilayers with chains shorter than C13 to the substantially different dynamical behavior of the constituent lipid molecules.

From these molecular BOMD results, we conclude that the dynamical behaviors of the individual DMPC and DLPC molecules are largely preserved in their assembly properties that are not related to changes of the ensemble topology. This conclusion is confirmed by results from short BOMD trajectories (14 ps) of DLPC in 50 waters that show isomer interchanges occurring at 240 K and 273 K similar to those illustrated in Fig. 8 for the isolated molecule.

4.3 Activation of Triplet Dioxygen by Bio-inspired Cuprous Complexes

We now present a second example of application of the BODFT-MD methodology for the case of dioxygen activation by bio-inspired copper complexes. Despite the fact that dioxygen can lead to many radical oxygen species such as HO^\bullet , $\text{O}_2^{\bullet-}$, or HOO^\bullet , that are harmful for the cells, aerobic organisms massively exploit dioxygen and direct its oxidative power toward many metabolites. Free dioxygen is, however, a triplet molecule while most organic substrates are singlet molecules. This so-called O_2 spin-mismatch implies that (1) reaction of $^3\text{O}_2$ with organic molecules is slow at ambient temperature, and (2) that catalysts must be used to overcome this kinetic barrier and to limit the possible formation of ROS in the cells. Biochemists have identified numerous enzymes allowing O_2 to be activated, most of them containing metal ions (primarily Fe and Cu) at their active sites [135]. In fact biochemical studies have revealed an impressively large panel of strategies employed by different enzymes to efficiently activate O_2 depending on the chemical nature of their substrate. In that regard the living realm provides a formidable source of inspiration for the development of innovative models [136].

We have been interested in the noncoupled copper-monooxygenases comprising Peptidylglycine α -Monooxygenase (PHM), Dopamine β -Monooxygenase (D β M), and Tyramine β -Monooxygenase (T β M) that catalyze the hydroxylation of C–H

bonds of C-terminal glycine-extended peptides of dopamine and of tyramine, respectively. Biochemical studies suggest that activation of the substrate C–H bond is promoted by a mononuclear cupric-superoxo species ($\text{Cu(II)/O}_2^{\bullet-}$) in the singlet state. The understanding of the process by which such a singlet species is formed from the interaction between a triplet O_2 molecule and the singlet cuprous complex is far from being satisfactory.

The kinetics of spin-forbidden reactions can be described by various theoretical models like those [137] derived from Transition State Theory that include nonadiabatic effects [138]. Basically these approaches follow the key concepts of TST (equilibrium between the reactant and the activated complex, semiclassical description of the molecular system) but take into account the probability the system has to hop from the initial quantum state (the triplet state here) to the final quantum state (the singlet) when reaching degeneracy between the two spin states. We have developed an alternative mixed quantum–classical (MQC) expression building on the work of Prezhdho and Rossky [139] and of Jasper and Truhlar [140] who explored the manifestations of decoherence effects in physico-chemical processes. In theoretical physics the term decoherence denotes processes by which a quantum system comes to behave classically upon interactions with its environment [141]. Applying these ideas to the case of chemical reactions involving two quantum states we obtained [142]:

$$k_{\text{SC}}^{\text{MQC}} = v \cdot \frac{8\mu^2}{\hbar^2 + 8\mu^2} \exp\left(-\frac{\Delta G^\ddagger}{k_{\text{B}}T}\right). \quad (18)$$

In this expression μ is the product of the electronic coupling between the two quantum states (including the spin–orbit coupling terms) and of a characteristic decoherence time τ_{dec} . The latter reflects the mixed quantum–classical character of the molecular system: “*although intrinsically obeying quantum mechanical laws, behave semiclassically after a finite but nonzero amount of time (τ_{dec})*” and as such Eq. (18) may be “*regarded as a mixed quantum classical rate constant expression.*” We recall here that this rate constant expression is expected to be valid for fast decoherence times (say <100 fs) that are shorter than the time characterizing the fluctuations of the diabatic energy gap.

Now the question is to devise algorithms for estimating τ_{dec} in molecular systems of significant size like those found in biology. A fully ab initio estimation would require performing simulations of the quantum density matrix of the system [143], which is out of the question for molecular systems composed of even tens of atoms. Coming back to the question of dioxygen activation by the PHM enzyme, we have considered the complexes depicted in Fig. 9 that mimic some of the main geometrical features of the parent enzymatic active site [144]. We followed the methodology proposed by Prezhdho et al. where decoherence is modeled by the decaying overlap between nuclear wave packets evolving on different electronic states. Contrary to these authors we have employed a BODFT-MD approach to perform our simulations. The computational protocol is illustrated in Fig. 10. In a

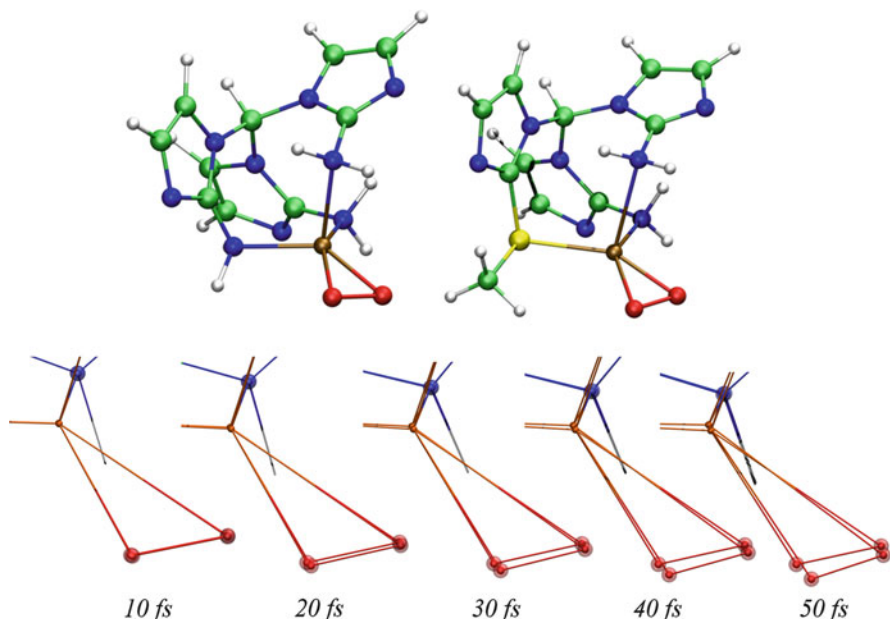


Fig. 9 DFT-based modeling of decoherence within a bio-inspired model of the active site on mononuclear copper enzymes. *Top*: two complexes investigated presenting an N_3 or an N_2S coordination sphere. *Bottom*: diverging motion of the copper and oxygen atoms on the fs timescale for one set of diverging trajectories. The transparent spheres represent the mass-dependent wave packet of each nucleus. *Color code*: Cu in brown, O in red, N in blue, C in green, S in yellow, and H in white

first step we performed a few tens of ps of BODFT-MD simulation in the NVT ensemble (at 300 K) on the singlet or on the triplet PES. In addition since the hopping probabilities between the two states (the quantities we wish to evaluate) reach a maximum in the region of degeneracy of two spin states, the BOMD simulations were biased by application of a harmonic constraint on the energy gap ($V_{\text{bias}} = k_{\text{bias}} (\Delta E_{\text{ST}} - 0)^2$). From this initial trajectory we extracted snapshots providing the starting conditions (positions and momentum of the nuclei and KS determinants) of so-called diverging trajectories which were run independently on the singlet and on the triplet surfaces (Step 2). A final step consists in the postprocessing of the data, that is, the estimation of the characteristic decoherence times and of the hopping probabilities obtained by integration of the Time-Dependant Schrödinger equation along the diverging BOMD [145].

The characteristic decoherence time for this copper complex has been found to be slightly below 10 fs for both complexes, whatever the direction of the reaction ($S \rightarrow T$ or $T \rightarrow S$) Table 1. The computed hopping probabilities entering the transmission coefficient of the rate constant show an interesting feature: the introduction of a sulfur atom within the copper coordination sphere induces an increase of the hopping probability by a factor of more than 3. These effects are related to the

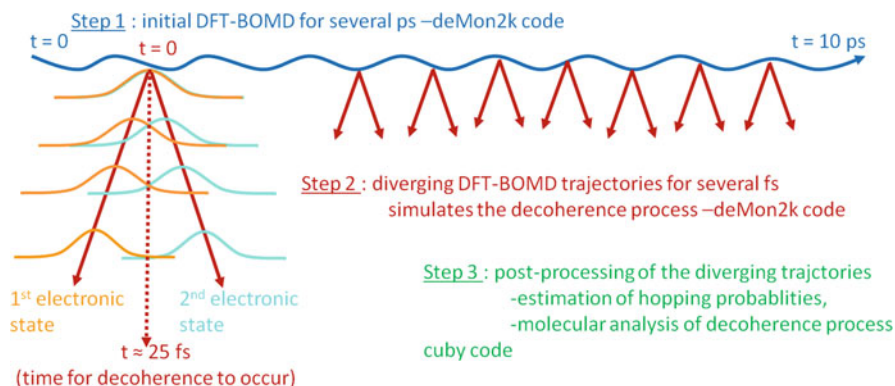


Fig. 10 Three-step computational protocol employed to estimate decoherence times with deMon2k. The orange and cyan Gaussian functions along the left-hand side of diverging trajectory represent one nuclear wave packet on the singlet and triplet PES

Table 1 Decoherence time and average hopping probabilities for the N_2N - and N_2S -based $[CuO_2]^+$ adducts following the computational protocol described in [142] that is based on DFT MD simulations

| | | $[(N_2N)CuO_2]^+$ | $[(N_2S)CuO_2]^+$ |
|-------------------|-------------------|-------------------|-------------------|
| τ_{dec} (fs) | $T \rightarrow S$ | 9.7 | 9.6 |
| | $S \rightarrow T$ | 9.2 | 9.4 |
| P_h | $T \rightarrow S$ | 0.035 (0.004) | 0.115 (0.010) |
| | $S \rightarrow T$ | 0.036 (0.004) | 0.106 (0.018) |

The numbers in parentheses are the standard deviations

higher electronic coupling computed for the $[(N_2S)CuO_2]^+$ adduct (ca. 160 cm^{-1}) compared to $[(N_2N)CuO_2]^+$ (ca. 60 cm^{-1}). It should be remarked, however, that these computations have been carried out in the gas phase for a bio-mimicking model of the real biological enzymatic active sites. New estimates of the above quantities with inclusion of the copper complex environment composed of protein residues and of water molecules are underway in our laboratories with the QM/MM computational approaches described in the next section.

5 DFT/MM-MD

Atomistic simulations of biomolecules remain one of the challenges of present day computational chemistry and biophysics. In such simulations one of the main difficulties is to give an accurate description of the structural and energetic aspects of the biochemical processes, while simulating very large systems for the lengths of time relevant to the studied problems. Particularly if one is concerned with properties that require high accuracy electronic structure, such as bond breaking/making, transition metal sites, magnetic properties, and sufficiently accurate

energetics (for example, relative energies of different conformers or transition states), the choice of methods is restricted to methods based on the calculation of the full electronic structure of the system. These electronic structure, or quantum mechanical (QM), methods are usually based on Density Functional Theory (DFT).

Despite the great development of computer and code performances, the size of the biosystem is still limited to about 1,000 atoms for geometry optimization and 150 atoms for reasonably long dynamics. Moreover, taking into account the solvent, when a global treatment is not sufficient, increases the computational demand even more.

As mentioned in the protocol for studying biomolecular systems in Sect. 1 of this review, if a simple cluster model is thought or shown to be inadequate, the model is most often extended by using hybrid QM/MM methods to incorporate the effects of the surrounding protein and solvent. Hybrid methods where the system is partitioned into a chemically active part, treated by first-principles QM methods and a larger environment, assumed to be chemically inert, which is modeled by classical molecular mechanics (MM), have been introduced in the late 1970s and have incorporated technological improvements over the years [146–152].

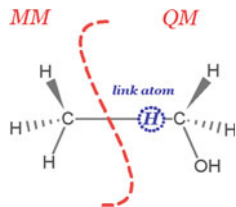
QM/MM has proven to be extremely successful in the study of biochemical, especially enzymatic, reactions and therefore has been widely applied in this field [153–165]. Numerous reviews have been devoted to overviews of QM/MM studies of biochemical reactions over the last 20 years [152, 166–175]. In this section, to introduce the methodology, we will first use the implementation of a QM/MM interface [176] between our computational chemistry program, deMon2k [70] and the widely used CHARMM [177] MM/MD package, as an example. For a more comprehensive treatment, please refer to our previous review on this topic [178].

In a typical QM/MM scheme, a system is usually divided into two subsystems: the QM subsystem treated by “high-level” QM methods and the MM subsystem treated by “low-level” force field-based methods. The boundary between these two subsystems distinguishes the QM region from the MM region. Ideally, partition of the system should not cut any covalent bonds to ensure the completeness of the QM subsystem. However, crossing covalent bonds is often unavoidable for large molecules, such as polymers and proteins. In order for a QM/MM calculation to mimic the real system the QM chemical structure has to be complete, i.e., no dangling bonds are permitted. An intuitive way to remedy a dangling bond is to cap it with an artificial atom, which gives rise to the so-called Link Atom approach [148]. In this approach, an additional atom, as a link between the QM and MM regions, is added to saturate the QM frontier atom at one end of the cut covalent bond. The link atom scheme is illustrated in Fig. 11. This link atom in most cases is a hydrogen atom because of its simplicity and practicality.

To explicitly calculate the QM subsystem (QMS)-MM subsystem (MMS) interaction on the QM level, an additive energy scheme has been widely used. It is formulated as

$$E(S) = E_{\text{MM}}(\text{MMS}) + E_{\text{QM}}(\text{QMS} + L) + E_{\text{QM}}[(\text{QMS} + L) - \text{MMS}], \quad (19)$$

Fig. 11 Illustration of the link atom scheme using ethanol as an example, in which the methyl group is treated as the MM subsystem and the rest is QM. Reproduced with permission from [178]



where L denotes the link atoms. The last term in Eq. (19) accounts for the coupling between QM and MM subsystems. It consists of electrostatic and van der Waals interactions between QM and MM atoms, all of which will be detailed below.

The electrostatic interactions between QM and MM subsystems are often calculated based on the following formula:

$$\hat{H}_{\text{QM-MM}} = - \sum_{i,m} \frac{q_m}{r_{im}} + \sum_{A,m} \frac{Z_A q_m}{R_{Am}}, \quad (20)$$

where q_m are the charges of MM atoms, Z_A the atomic number of QM atoms, i runs over all QM electrons, A over all QM atoms including link atoms, and m over all MM atoms. The first term is a one-electron operator and the second accounts for the nuclei–MM charge interaction. When acting on the QM wave function, Eq. (20) results in the electrostatic interaction between QM and MM subsystems as a portion of $E_{\text{QM}} [(QMS + L)\text{-}MMS]$.

As one can see in Eq. (20), calculation of the first term on the DFT level can significantly slow down the entire QM/MM calculation, especially when there are a large number of MM charges. However, these charges cannot be simply ignored because long-range electrostatic effects can often be of considerable importance. To speed up the calculation of long-range electrostatics, we have introduced a multipole expansion-based approach [179] in the DFT framework. This has resulted in significant savings in time and plausible accuracy compared to full DFT calculation as shown in Figs. 12 and 13 and Table 2.

Besides electrostatic interactions, the QMS–MMS coupling also includes van der Waals (vdW) interactions. The vdW interaction is usually described by a Lennard–Jones 12-6 potential:

$$E_{ij} = \sum_{ij} \left(\frac{A_{ij}}{r_{ij}^{12}} - \frac{B_{ij}}{r_{ij}^6} \right), \quad (21)$$

where i runs over QM atoms and j over MM atoms, and A and B are constants pertaining to atom types. In QM/MM calculations, when electrical embedding is used, the vdW interaction of QMS–MMS could be incorrect as the corresponding electrostatic interaction is not the parameterized point charge–point charge interaction any more. However, Cui and co-workers have tested three sets of vdW

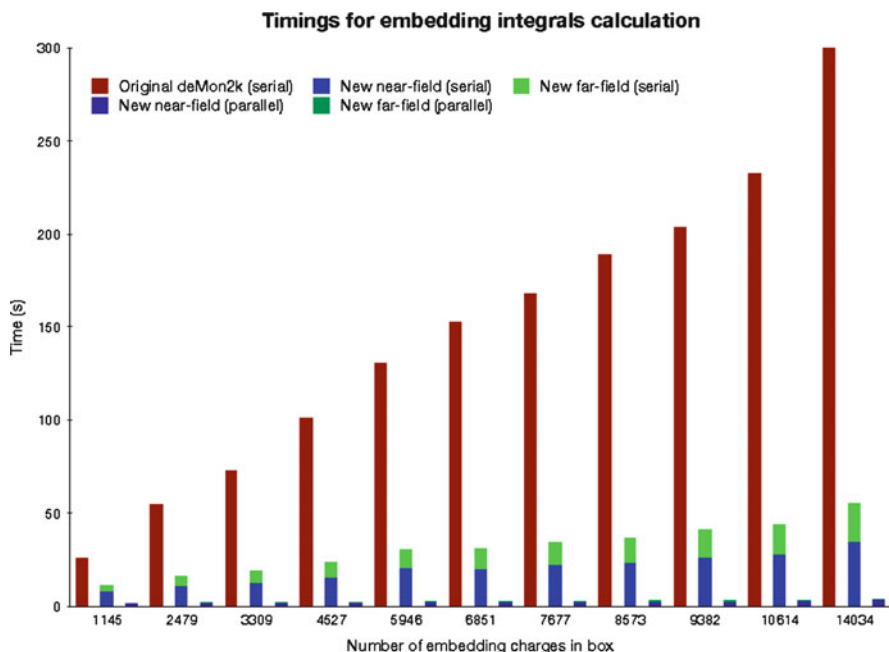


Fig. 12 Comparison of computational timings for the embedding integral calculation in small QM/MM systems

parameters and concluded that the QM/MM energetics were not sensitive to the vdW parameters and efforts to improve QM/MM accuracy should focus elsewhere [180].

Built on the energy scheme, the main focus of QM/MM calculations is often on geometry. In principle, the whole QM/MM system can be simultaneously optimized with a uniform optimizer using the QM/MM potential and gradient. Convergence, however, could be difficult to reach when the starting geometry is far from the minimum. Considering the different natures of QM and MM methods, optimization is sometimes easier to run separately for each subsystem on their respective levels. To this end, a macro/microiterative scheme has been implemented [181–187]. In this scheme, the optimization is driven by the QM optimizer. The MM subsystem is optimized to convergence with the QM part frozen, and this is termed as micro iteration. Thereafter, the QM region is optimized till convergence with the MM part frozen, and this is termed a macro iteration. These two iterations alternate until the whole system is fully optimized. A diagram of this scheme is shown in Fig. 14.

In addition to optimized geometries, which are minima on the potential energy surface, curvature on the free energy surface is also of interest in many QM/MM studies. Therefore, efficient calculations of free energy are required. One of the most widely used techniques to calculate the free energy in QM/MM methods is free energy perturbation (FEP) theory. Originally developed in MM, this FEP

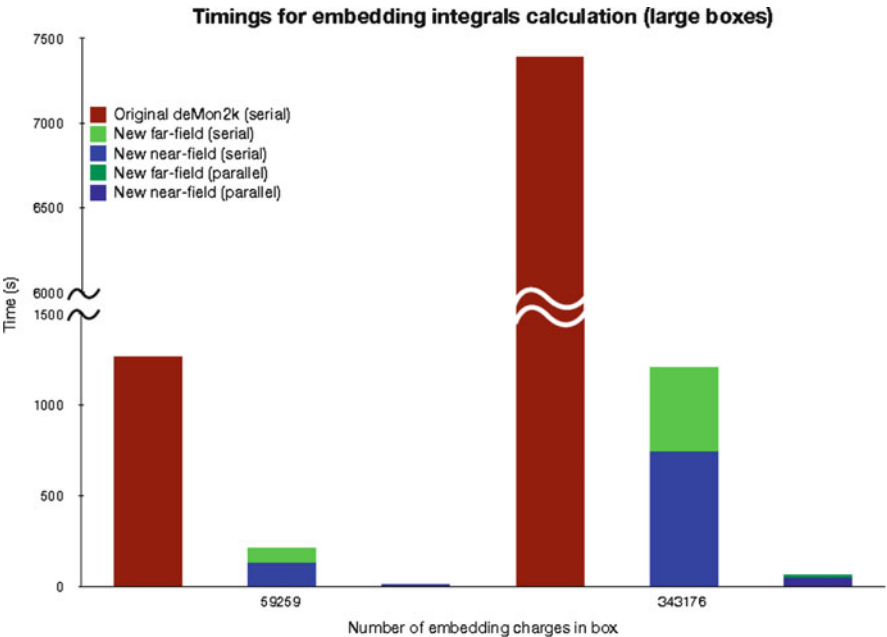


Fig. 13 Comparison of computational timings for the embedding integral calculation in large QM/MM systems

Table 2 Comparison of the total energies (a.u.) of the embedded systems

| Embedding charges in box | Original deMon2k (serial) | New code (serial) | New code (parallel 16 processors) |
|--------------------------|---------------------------|-------------------|-----------------------------------|
| 1145 | −4880.219852928 | −4880.219852922 | −4880.219853263 |
| 2479 | −4880.197609272 | −4880.197609273 | −4880.197609798 |
| 3309 | −4880.151039302 | −4880.151039303 | −4880.151039189 |
| 4527 | −4880.342973267 | −4880.342973268 | −4880.342973155 |
| 5946 | −4880.292634368 | −4880.292634369 | −4880.292633976 |
| 6851 | −4880.359550818 | −4880.359550818 | −4880.359550949 |
| 7677 | −4880.314753835 | −4880.314753835 | −4880.314754031 |
| 8573 | −4880.672278586 | −4880.672278586 | −4880.672278983 |
| 9382 | −4880.728606709 | −4880.728606709 | −4880.728607116 |
| 10614 | −4880.785583226 | −4880.785583226 | −4880.785583612 |
| 14034 | −4880.580437797 | −4880.580437798 | −4880.580438027 |
| 59259 | −4880.428695928 | −4880.428695929 | −4880.428695680 |
| 343176 | −4880.335242269 | −4880.335242272 | −4880.335241941 |

technique relates the free energy difference between state A and state B to the potential energy difference between these two states [188]. If we assume that the perturbation required to transform system A to system B is small (<2 kT), it can be shown that:

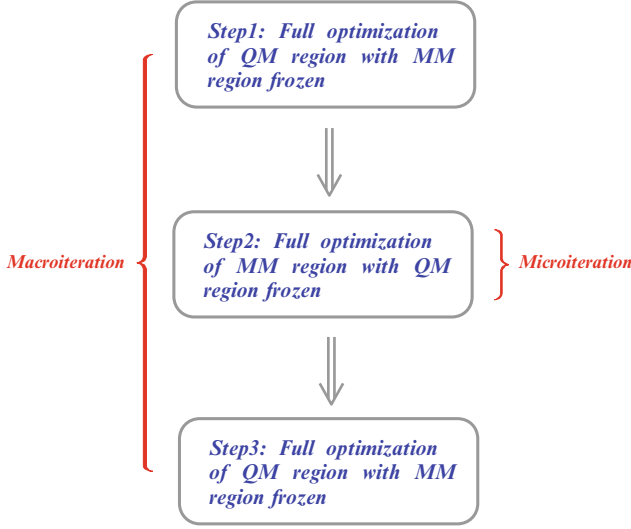


Fig. 14 Schematic of the macro–micro iteration scheme for QM/MM geometry optimization

$$\Delta G_{(A \rightarrow B)} = -k_B T \ln \left\langle \exp \left(-\frac{E_B - E_A}{k_B T} \right) \right\rangle_A, \quad (22)$$

where the potential energy difference between two states is weighted by the energy of the initial state. In case of larger perturbations, one can always use additional windows to connect starting and ending points of this perturbation. The free energy is a path-independent property of the system and can be evaluated regardless of how “alchemical” the perturbation path may be.

In the formalism of QM/MM, a parameter λ is introduced to relate the two states. Thus, the Hamiltonian describing the system which is changing from state A to state B during FEP calculations can be re-written as

$$H(\lambda) = \lambda E_A + (1 - \lambda) E_B, \quad (23)$$

where E_A and E_B represent distributions of states for the two end points of the perturbation. The dynamics of two replicas of the system each corresponding to end points with $\lambda = 0$ and $\lambda = 1$ is treated explicitly and simultaneously and appropriate weighting is applied to reconstitute the Hamiltonian for intermediate windows.

We have applied this method in our QM/MM implementation [176] to the study of ion solvation and the mechanisms of selectivity in ion channels or ion-coupled transporters. Ion selectivity has been extensively studied at the MM level with both classical and polarizable force fields. However, classical simulations may be compromised by their inability to account for charge transfer and electronic polarization, thought to be critical for ion binding to proteins. We have extended QM/MM FEP to studies of Na^+/K^+ solvation and selectivity by water clusters with

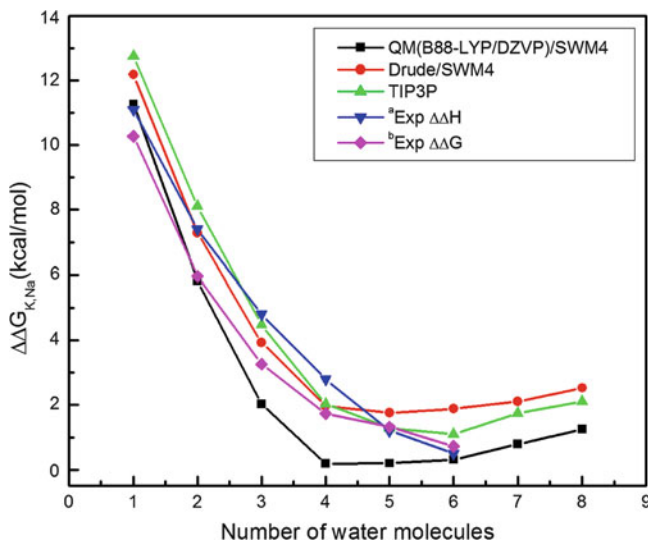


Fig. 15 Relative (to the bulk) free energy of selectivity for Na^+/K^+ in water clusters as a function of cluster size. Reproduced with permission from [176]

variable numbers of ligands, for which results are illustrated in Fig. 15. In [176] QM/MM FEP calculations have been performed for ion-water clusters with different numbers of water molecules, where the ion was treated as the QM region using B88-LYP with the DZVP basis set implemented in deMon2k, and waters were represented by the polarizable Drude force field implemented in CHARMM. The same approach can be used for selectivity calculations in more complex biological systems. Also, the close connection between QM/MM calculations and those with polarizable force fields developed to account for electronic effects is visible. Therefore, the polarizable force fields can be sufficient for some cases.

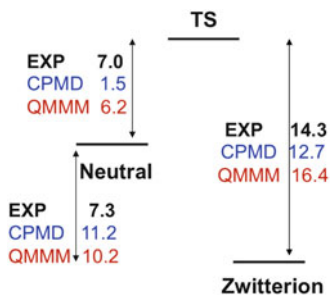
Hybrid QM/MM methodology looks particularly attractive for reproducing solvent effects when a discrete representation of the solvent molecules is needed, as in the example above, and for many other bio-processes, such as microsolvation at enzymatic sites or at metal binding sites in peptides, proton transfer, free-energy calculations. In fact, comparison of full QM and QM/MM results for sufficiently small peptides in water has been performed recently, including an analysis of the full MM results [189, 190]. These comparative studies led to the following main conclusions (1) the solvation pattern around most groups of the dipeptides is similarly described by the three methods (QM, QM/MM, and MM); (2) solvation patterns around terminal $-\text{CO}_2^-$ and $-\text{NH}_3^+$ groups are poorly reproduced in MM simulations; (3) solvation patterns of the full QM results are well reproduced, at a lower cost, by the QM/MM calculation involving MM waters; (4) DFT/MD computed chiroptical properties of a solvated glycine molecule are comparable to those obtained with the much more expensive coupled cluster CC2 method.

The QM/MM methodology is also powerful in providing accurate structural and free energy changes in solution at a moderate computational cost for relatively large models treated with a QM method in the presence of a solvent calculated explicitly using an MM approach. Several simulation techniques based on *ab initio* quantum mechanics have been proposed for this purpose. The quantum mechanical free energy (QM-FE) approach mentioned above consists of optimizing the reaction path in the QM system in the gas phase and of applying free-energy perturbation (FEP) along the preoptimized gas-phase reaction path with the inclusion of the QM/MM interactions that are treated classically [191]. This method may have some inaccuracy if the gas-phase reaction path does not yield a good description of the reaction path in solution. The QM(ai)/MM method developed for the study of condensed phase reaction processes approximates the potential-energy surface from *ab initio* QM with an empirical valence bond (EVB) potential [192]. At the same time, QM/MM-FE was proposed for the simulation of enzymatic reaction processes [193–195]. In this method, the dynamics of the QM and MM systems are assumed to be independent of each other, although the two subsystems interact energetically, and the QM subsystem motions are taken to be harmonic. In order to avoid the dependency of the reaction path on the choice of the initial conformation, the reaction path is thus calculated on the potential of mean force (PMF) surface of the QM/MM system, yielding the QM/MM minimum free-energy path (QM/MM-MFEP). The free-energy gradients are used to perform the path optimization on the free energy surface.

The QM/MM-MFEP method is easily combined with the deMon2k version that includes MM and QM/MM energy and gradient evaluations inside the program core. The QM/MM-MFEP method using this deMon2k version has been tested on glycine in aqueous solutions, namely the path from neutral to zwitterionic glycine in a water droplet (53, 108, and 182 water molecules). The glycine molecule is the QM subset surrounded by SPC MM waters.

Being an amino acid prototype, glycine solvation in water has been studied in the literature using different methods and aiming at different goals. Indeed, neutral glycine only exists in the gas phase, whereas its zwitterionic form is mainly present in water solution [196]. Qualitatively, the larger zwitterion stability is found using a dielectric solvation model [197]. Several studies searching for the minimum number of water molecules that are needed to stabilize the zwitterions concluded to various values ranging from 2 to 7 [197–199]. Statistical averaging from finite-temperature simulations has been performed using MM, QM/MM, full QM methods, and the continuum solvent model [189, 190, 200–204]. In most of these studies, detailed hydration shells and solute–solvent interactions of the zwitterionic form are investigated. The free energy difference, ΔG , between neutral and zwitterionic solvated species has also been studied, despite the fact that neutral aqueous glycine is not observed experimentally. A ΔG value of $8.7 \text{ kcal mol}^{-1}$, close to the experimentally proposed value of about $7.0\text{--}7.5 \text{ kcal mol}^{-1}$ [196], was obtained using the QM gaseous energies of neutral and zwitterionic forms of glycine and its radical (calculated with the accurate CBS-QB3 procedure [205]) and MM solvation energies (free energy cycle) [200]. Full QM simulations using the Car and

Fig. 16 Schematic representation of the free energy proton transfer path from neutral to zwitterionic glycine or vice versa in 53 waters. The QM/MM-MFEP results are presented in *red*, compared with experimental values in *black* [205, 206] and CPMD calculations in *blue* [201]



Parrinello (CPMD) approach [201] showed variable hydration numbers along the proton transfer path and led to a zwitterion–neutral glycine ΔG of $11.2 \text{ kcal mol}^{-1}$ and a proton transfer barrier, ΔG^* , of $12.7 \text{ kcal mol}^{-1}$. The advantage of performing the dynamics of the peptide in a full QM or in a QM/MM calculation is to provide the reaction path and the barrier estimate. Combining the experimental barrier value of $14.3 \text{ kcal mol}^{-1}$ for the zwitterion to neutral glycine reaction [206] and the energy difference between the neutral and zwitterionic glycine, one concludes that the experimental neutral glycine ΔG^* is about 7 kcal mol^{-1} . The full QM CPMD calculations [201] thus lead to an underestimated free energy barrier of $1.5 \text{ kcal mol}^{-1}$ for the neutral glycine.

Figure 16 illustrates the results obtained for our QM/MM simulation of glycine in 53 MM waters using the QM/MM-MFEP methodology.

First, the neutral glycine was placed into a spherical droplet of 53 waters, cut from a drop of 2500 waters equilibrated at 300 K. The glycine molecule BOMD was performed using the DFT methodology (PBE98-LYP-D, DZVP/GEN-A2 bases), whereas the MD of the water molecules used the SPC force field [207].

The system was left to evolve at 300 K using a Nose–Hoover thermostat in the NVT ensemble. After about 100 ps, the proton of the H_a acidic glycine moved to N. As also mentioned in the CPMD study [201], a decrease of the water coordination to glycine was observed before and during the transfer (2 waters bound to the acidic OH and to the N atom moved farther away). The proton transferred directly to the amine N without any intermediate bonding with water.

In order to calculate the free-energy changes following the PMF description of the proton transfer, we used the QM/MM-MFEP method [195]. The reaction path was divided into six steps and the reaction coordinate along this path was chosen to be the distance between the acidic proton H_a to the amine nitrogen N. Step 0 was the neutral and step 5 the zwitterionic glycine. The $H_a \cdots N$ distances of steps 1–4 were taken from snapshots of BOMD-MD dynamics.

The calculated free energy changes from neutral to zwitterionic glycine are obtained from the calculation of the potential of mean force of the proton transfer in water, with the contribution of the water degrees of freedom being ensemble-averaged out.

The optimization of the proton transfer is carried out in a discretized representation, using the free-energy perturbation formula for the chain of conformations (steps)

$$A(r_{\text{QM}}) = A_{\text{ref}} - \frac{1}{\beta} \text{Ln} \langle \exp(-\beta[E(r_{\text{QM}}, r_{\text{MM}}) - E_{\text{ref}}(r_{\text{MM}})]) \rangle_{E_{\text{ref}}, r_{\text{MM}}}, \quad (24)$$

where $E_{\text{ref}}(r_{\text{MM}})$ is the total energy of the system expressed in terms of the coordinates of the QM and MM subsystems, at the QM reference geometry. The QM geometry and the MM ensemble are mutually dependent. The first QM geometry (step 0) is set as the reference point on the PMF. Its free energy is A_{ref} . The $E_{\text{ref}}(r_{\text{MM}})$ values for each step were obtained by optimizing the full QM/MM system keeping the constraint of the $\text{H} \cdots \text{N}$ distance fixed for every step from 1 to 4. Finally, sampling of the MM space was provided by about 40 ps MM water dynamics keeping frozen the full QM subsystem at the reference geometry of the step. It has been verified that doubling this MM sampling time did not change the $A - A_{\text{ref}}$ values.

In order to take into account the sensitivity of the energetic results with respect to the orbital basis set extension, the $E_{\text{ref}}(r_{\text{MM}})$ values obtained from DZVP/GEN-A2 calculations were followed by one single point energy calculation with the cc-pVTZ basis [208], and the auxiliary GEN-A2* set including up to g functions.

Figure 16 shows that both theoretical methods overestimate the neutral–zwitterion free energy difference, whereas the QM/MM methodology leads to a better estimate of the neutral to transition state free energy difference. However, increasing the number of waters from 53 to 108 brings the results into very close agreement with both experimental values. The effects of glycine dilution are presently under study [209].

We conclude this section with some remarks on the RNA polymerase enzyme for which cluster calculations have been discussed in Sect. 2.2.2. Extensions of that work to include more of the protein and solvent environment through QM/MM calculations and free-energy perturbation theory are in progress, but they are not yet ready for publication. Instead, we will give a glimpse of some related work from the literature on DNA polymerases, which show many of the same essential features as their RNA counterparts.

DNA polymerases and RNA polymerases help replicate DNAs and RNAs with considerably high fidelity, selecting the matched deoxynucleoside triphosphate (dNTP) and NTP over the mismatched dNTPs and NTPs, and catalyzing the nucleotidyl transfer reaction. In the QM/MM studies of DNA pol IV and T7 pol, Zhang and coworkers [210, 211] utilized the pseudobond approach to cap the QM subsystem, the micro/macro iteration scheme to optimize the geometry, the reaction coordinate driven method for the reaction path search, and the free energy perturbation method for free energy calculations along the path. They found that the nucleophilic attack was the rate-limiting step and the initial proton transfer was assisted by water.

Figure 17 shows a comparison of potential energy and free energy profiles for this reaction. The potential energy profile is in fact reminiscent of the results for our cluster models of RNA polymerase shown in Fig. 5. The warning about the very large effects of entropy should be clear from Fig. 17.

Finally, we mention that the initial proton transfer has also been a focus in the study of RNA polymerase II. Ramos and coworkers [212] adopted an ONIOM method in their QM/MM study of RNA polymerase II. They tested several proton transfer possibilities and concluded that none of the adjacent residues or the RNA primer could be the deprotonation agent due to the high activation barriers. As an alternative solution, they introduced a hydroxide ion in the reaction site to be the proton acceptor, which resulted in a reasonable activation barrier. However, the source of the hydroxide ion was not discussed and it is not clear to us whether this hydroxide-ion assisted mechanism is in fact plausible.

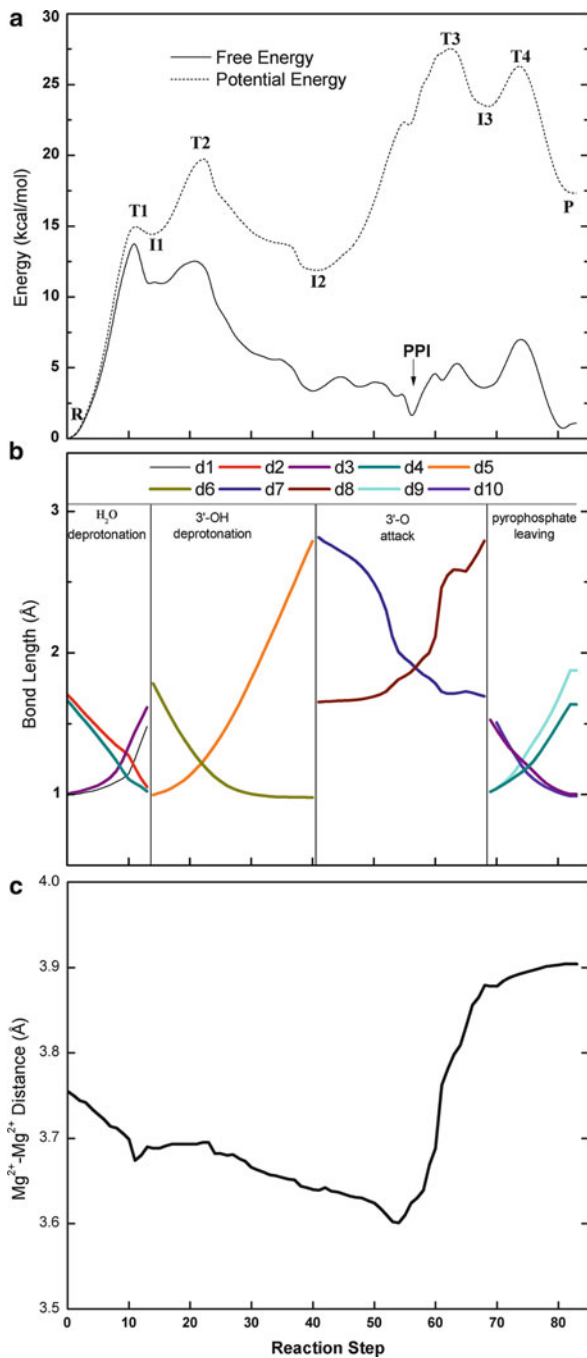
6 Constrained-DFT

6.1 Methodological Background

The resolution of the KS equations gives access to the electronic density of the adiabatic ground state of the molecular system of interest. In other words, one finds by the SCF procedure a relaxed electronic density that corresponds to a minimum of the DFT energy. It is, however, possible that one desires to investigate properties of electronic densities that *do not* correspond to the electronic ground states. The purpose of constrained DFT is precisely to enable the user to impose a particular atomic charge or a spin density while solving the KS equations. Although artificial, this procedure can have considerable interest for the modeling of (bio-)chemical systems. We will provide various examples of applications in the second part of this section but we can already mention the question of electron transfer reactions within the context of the Marcus Theory [213]; this theory relies on the definition of phenomenological states that cannot be obtained by standard DFT (which provides adiabatic states). cDFT is a means to define such ad hoc diabatic states. As another example we remark that a strength of computational chemistry is to give access to systems or situations that are not amenable to direct experimental measurements. Chemical trends like the electronic effects of substituents on a reactive energy profile can be investigated by various methods including cDFT or other methods by imposing artificial constraints on the chemical system of interest. As an example of the usefulness of other procedures we refer to [214] where cDFT is not used but the nuclear charge of nitrogen atoms is varied to modify its electronegativity and indirectly the energy cost for activating an alkyl C–H bond.

In the next paragraphs we explain how this objective is achieved with cDFT and how an efficient implementation within the context of ADFT can be devised. Our aim is not to provide here a delineated mathematical derivation of the

Fig. 17 Potential energy and free energy profiles for the nucleotidyl addition reaction of a DNA polymerase. Reproduced with permission from [211]



formalism, but rather to introduce the key aspects of the method so as to provide the reader with a view of the potential applications of cDFT. For more mathematical details, the reader is referred to recent reviews on the method [215, 216]. The foundations of constrained DFT are to be found in the pioneering works of Gunnarsson and Lundqvist [217] and of Dederich et al. in the early 1980s [218]. Wesolowski, Muller, and Warshel also developed a related, though algorithmically different, constrained DFT approach in the 1990s [219] with remarkable success for the modeling of condensed phase proton transfers, electron transfer reactions [220], and S_N2 reactions [221], in connection with the Empirical Valence Bond approach [222]. We will not, however, cover this methodology in this chapter and the reader is referred to the references for more details. Continuing with the principal works on cDFT formalisms we finally mention that of Wu and Van Voorhis in 2005 [223]. In 1976 Gunnarsson and Lundqvist showed that in addition to the true ground state it was possible to obtain relaxed electronic densities of the particular molecular symmetries. A few years later Dederich et al. proposed a generalization of this idea that used a Lagrange Multiplier to constrain the DFT energy:

$$\varepsilon[\rho, \lambda_c] = \min_{\rho} \max_{\lambda_c} \left(E[\rho] + \lambda_c \left[\int \rho(\mathbf{r}) w(\mathbf{r}) d\mathbf{r} - N_C \right] \right). \quad (25)$$

In this equation E is the nonconstrained DFT energy whose mathematical expression has been given in Sect. 2.1, $w(r)$ is a weight function that defines the constraining property, and N_C is the set-point supplied by the user. For example to constrain N_C electrons to occupy a volume Ω the weight function would equal 1 inside Ω and zero everywhere else. Both $w(r)$ and N_C are thus user-defined terms. We will come back later to the practical definition of $w(r)$ within the LCAO framework. For the moment we focus on the Lagrange multiplier λ_c that needs to be determined. To emphasize the role of this term in the formalism it is useful to write down the set of modified KS equations. These are obtained by differentiation of the cDFT energy with respect to the MO coefficients under the orthonormalization constraint.

$$\left(-\frac{1}{2} \nabla^2 + v_{\text{ext}}(\mathbf{r}) + \int \frac{\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}' + v_{\text{xc}}(\mathbf{r}) + \lambda_c w(\mathbf{r}) \right) \psi_i = \varepsilon_i \psi_i(\mathbf{r}) \forall i. \quad (26)$$

Within the LCGTO formalism employed in deMon2k the elements of the constrained KS matrix are given by

$$K_{\mu\nu} \equiv \left(\frac{\partial \varepsilon_{\text{ADFT}}}{\partial P_{\mu\nu}} \right)_x = H_{\mu\nu} + \sum_k \langle \mu\nu || \bar{k} \rangle (x_{\bar{k}} + z_{\bar{k}}) + \lambda_c W_{\mu\nu}. \quad (27)$$

It is apparent from these equations that the product $\lambda_c W$ acts like a supplementary potential felt by the electrons. Its role is to drive the convergence of the

SCF procedure toward a relaxed density that fulfills the desired constraint. It must be noted, however, that λ_c is not known beforehand (contrary to W or N_c) and must be determined. In the older applications of cDFT, it was customary to scan over possible values of λ_c to find the correct one. This was obviously a cumbersome strategy that probably restricted the application of cDFT to only a few fields of research. In addition, a critical point when solving the cDFT KS equations is related to the fact that the Coulomb and XC potentials on the one hand and the $\lambda_c w$ potential on the other are inter-dependent. This makes the resolution of the constrained KS equations difficult. Wu and Van Voorhis proposed an efficient solution to these difficulties building on Optimized Potential Theory. By examining the stationary conditions of the cDFT energy with respect to λ_c , these authors proved that, for a *given* Coulomb and Exchange Correlation (XC) potential there is a unique λ_c that leads to an electronic density fulfilling the desired constraint. In addition, the correct λ_c corresponds to a maximum of the energy. The first and second derivatives of the energy with respect to λ_c are easily obtained from the KS orbital coefficients so that an automatic algorithm can now be implemented for searching the correct λ_c . In practice this optimization is conducted according to the combination of steepest descents and a Newton–Raphson algorithm for best efficiency [224]. Then to address the difficulty of the coupling between the Coulomb, XC potential, and the constraint, Wu and Van Voorhis proposed a dual-loop strategy whereby the determination of KS orbital energies is decoupled from the determination of λ_c . The algorithm is illustrated in Fig. 18. At every SCF step λ_c is optimized keeping the Coulomb and XC potential constants (“inner loop” in the figure). Once the correct λ_c is determined the constrained density matrix is used to calculate the new Coulomb and XC potentials for the next SCF step. Our experience told us that the convergence of the inner loop is usually not problematic in terms of CPU time, especially if a guess for λ_c from the previous SCF iteration can be provided. At global convergence the SCF procedure provides a converged electronic density fulfilling the desired constraint. The most demanding task in ADFT/cDFT remains the numerical calculation of the XC matrix elements, but since these terms remain constant when converging the inner loop the necessity to optimize λ_c is not lethal for the computational efficiency of the ADFT framework. In other words, cDFT is compatible with ADFT and its implementation with the min–max algorithm [225]. The calculation of the contribution of the constraining term to the energy gradients with respect to the nuclear positions has also been implemented in deMon2k, allowing geometry optimization, frequency analysis, and BOMD with cDFT [226]. However, we identified a computational bottleneck in the calculation that becomes problematic for molecular systems of a few tens of atoms (ca. 70–80 atoms). This is due to the involvement of the orbital density in the constraining term and therefore to the calculation of products of GTOs in the numerical evaluation of the energy derivative terms. We are pursuing our efforts to derive a fully cADFT formalism where the constraint would apply on the auxiliary function density instead of on the orbital density.

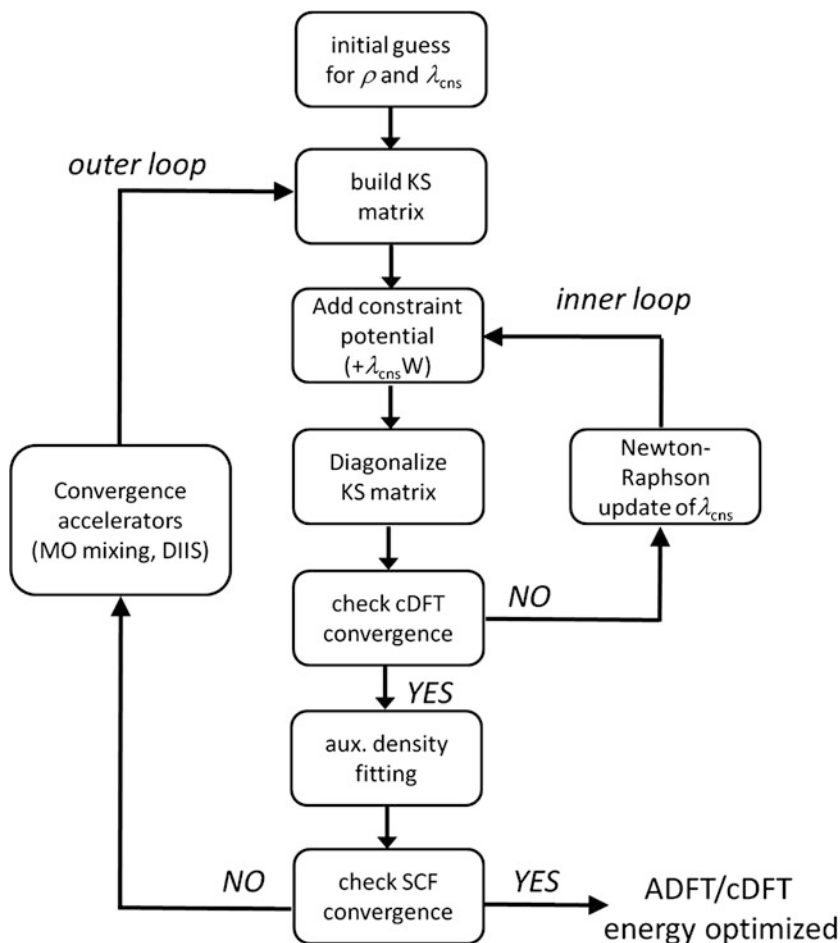


Fig. 18 Dual-loop SCF algorithm for solving the modified ADFT/cDFT KS equations in deMon2k. Reproduced with permission from [225]

6.2 cDFT and Population Analyses

So far we have presented cDFT as a means to impose a charge or a spin density on a group of atoms. However, we have not specified how such charges are calculated in practice. It is well known that atomic charges cannot be strictly defined in quantum mechanics and, as a corollary, that multiple population analysis (PA) approaches can be developed to reach this goal. A central point PAs have to address is to define a criterion for assigning each fraction of the electronic density of every point in space to the individual atoms. Some PAs realize this partition with criteria relying on the KS molecular orbital coefficients. The Mulliken [227], Löwdin [228], and the more sophisticated Natural Bond Orbital [229, 230] approaches are examples of

such methods. Other PAs work on the electronic density itself, a strategy that seems most appropriate to the philosophy of DFT. The Voronoi Deformation Density (VDD) [231], Becke [232], Hirshfeld [233], and the more recent iterative Hirshfeld [234–236] and iterative Stockholder [237] are examples of PAs falling into this second category. For the sake of completeness, we also mention the Atoms-in-Molecules approach developed by Bader that uses a topological analysis of the electron density, but we will come back to this method in Sect. 7. The mathematical expressions for calculating the atomic charges for some of these approaches are given in Table 3.

In this table, P is the density matrix and S is the overlap matrix of the atomic GTOs (μ and ν). Z_A is the nuclear charge of atom A . For the Becke, Hirshfeld, and VDD approaches, the electron density is integrated numerically over a grid of points. The Voronoi cell of atom A is the ensemble of points closer to A than to any other atom. The Becke cell is defined similarly except that a smoothing function is applied at the borders between the cells. ρ^A is the electron density of the neutral atom A .

Coming back to cDFT one needs to make a choice of a particular PA to define the charge of the atoms to be constrained. This choice determines the mathematical definition of the W matrix elements. So far the PAs that have been used for cDFT are the Mulliken, Löwdin, Becke, Hirshfeld, and VDD approaches. Our personal experience has led us to prefer the Hirshfeld scheme that generally provides more reasonable results from a chemical point of view. However, apart from the Mulliken approach that is usually not satisfying, cDFT with the Löwdin, or Becke method is usually satisfactory. A future worthwhile objective would be to devise algorithms for making cDFT compatible with more sophisticated PA approaches such as the iterative Hirshfeld, the iterative Stockholder, or Bader's Atoms-In-Molecules approach. Many of these approaches are available in deMon2k [238], but using them for cDFT is however nontrivial because the mathematical definition of the W matrix would then depend on the density matrix itself (at variance with the cases reported in Table 3). Said otherwise, the W matrix would vary between successive SCF steps like the Coulomb and XC matrices, making the dual-loop procedure of Fig. 18 partially obsolete. We now report on the applications of cDFT to biologically relevant electron-transfer systems. A further example will be given in the section on "interpretational-DFT."

6.3 Modeling Electron Transfer Reactions

Along with hydrogen transfers (in the form of protons, hydrogens, or hydrides), electron transfer is one of the most fundamental chemical processes in biological systems. Electron transfers are found in the respiratory chain, in photosynthesis, and in uncountable metabolic pathways. Enzymes such as the family of oxidoreductases have evolved to catalyze chemical reactions such as the functionalization of C–H bonds. Inner-sphere or outer-sphere (Long Range) electron

Table 3 Five population approaches commonly used in cDFT computations to define the weight matrix elements

| PA | Definition of the charge in the volume Ω | Weight matrix elements | cDFT weight w_i |
|-----------|---|---|---|
| Mulliken | $Q_A^{\text{Mulliken}} = Z_A - \sum_{\mu \in A} (PS)_{\mu\mu}$ | $W_{\mu\nu} = S_{\mu\nu}$ if μ and $\nu \in A$ $W_{\mu\nu} = \frac{1}{2} S_{\mu\nu}$ if μ or $\nu \in A$ $W_{\mu\nu} = 0$ if μ and $\nu \notin A$ | – |
| Löwdin | $Q_A^{\text{Löwdin}} = Z_A - \sum_{\mu \in A} (S^{1/2} P S^{1/2})_{\mu\mu}$ | $W_{\mu\nu} = \sum_{\lambda} S_{\mu\lambda}^{1/2} S_{\lambda\nu}^{1/2}$ | – |
| Becke | $Q_A^{\text{Becke}} = Z_A - \int_{\text{Becke cell of } A} \rho(r) dr$ | $W_{\mu\nu} = \sum_i a_i w_i \mu(r_i) \nu(r_i)$ | $w_i^{\text{Becke}} = 1.0$ inside the cell $w_i^{\text{Becke}} = 0.0$ outside the cell + smoothing function |
| Hirshfeld | $Q_A^{\text{Hirshfeld}} = Z_A - \int \rho(r) \frac{\rho_A^a(r)}{\sum_{\text{all atoms}} \rho^a(r)} dr$ | $W_{\mu\nu} = \sum_i a_i w_i \mu(r_i) \nu(r_i)$ | $w_i = \frac{\sum_{\text{atoms in } A} \rho^a(r_i)}{\sum_{\text{all atoms}} \rho^a(r_i)}$ |
| VDD | $Q_A^{\text{VDD}} = \int_{\text{Voronoi cell of } A} \left[\rho(r) - \sum_{\text{all atoms}} \rho^a(r) \right] dr$ | $W_{\mu\nu} = \sum_i a_i w_i \mu(r_i) \nu(r_i)$ | $w_i^{\text{VDD}} = 1.0$ inside the cell $w_i^{\text{VDD}} = 0.0$ outside the cell |

transfers are usually at play within the catalytic cycles of these enzymes [239]. However, the precise molecular strategies enabling efficient catalysis are often elusive. Computational chemistry, and in particular DFT, provides a valuable means to complement biochemical studies to reach a detailed understanding of these biological processes. From a theoretical point of view, the most popular conceptual framework for dealing with the kinetics of ET is the Marcus Theory (MT) [213]. The theory assumes the definition of two phenomenological electronic states corresponding to situations where the electron to be transferred is localized on the reductant (initial redox state) or on the oxidant (final redox state). These diabatic electronic states correspond to the empirical description familiar to the chemist. In MT two limiting regimes are considered. First when the quantum coupling between the two diabatic states is strong as in the case of inner-sphere ET, an adiabatic rate constant can be employed:

$$k_{\text{ET}}^A = \nu \cdot \exp\left(-\frac{(\Delta G^\circ + \lambda)^2}{4\lambda k_{\text{B}}T}\right), \quad (28)$$

where ΔG° is the free energy of the reaction, λ is the reorganization energy, and ν is the effective frequency along the reaction coordinate. Actually the diabatic energy gap is often a good choice of reaction coordinate [240]. The other terms have their usual meaning. When the coupling between the two diabatic states is weak, as in the case of a long-range ET or of symmetry-forbidden ET, a nonadiabatic expression applies

$$k_{\text{ET}}^{\text{NA}} = \frac{2\pi}{\hbar} \frac{1}{\sqrt{4\pi\lambda k_{\text{B}}T}} |H_{\text{DA}}|^2 \exp\left(-\frac{(\Delta G^\circ + \lambda)^2}{4\lambda k_{\text{B}}T}\right). \quad (29)$$

where H_{DA} is the electronic coupling factor between the initial and final redox states (respectively $|\varphi_{\text{D}}\rangle$ and $|\varphi_{\text{A}}\rangle$):

$$H_{\text{DA}} = \langle \varphi_{\text{D}} | H_{\text{el}} | \varphi_{\text{A}} \rangle, \quad (30)$$

H_{el} being the electronic Hamiltonian.

The cDFT formalism of Dederich et al. and Wu et al. has been employed by various groups to model the two diabatic states and to compute the terms entering the adiabatic and nonadiabatic rate constant expressions [241–244]. In particular, cDFT is one of the few DFT-based approaches that enable the evaluation of the electronic coupling terms between the diabatic states with no need for the adiabatic representation [242, 245]. In addition, the possibility of performing BOMD simulations on cDFT potential energy surfaces makes possible the evaluation of free energy terms. In this sense, cDFT holds great promise for the modeling of ET reactions at the DFT level, especially thanks to the computational advantages of the cADFT framework. We already mentioned the cDFT method of Wesolowski et al. that also has found numerous applications in the field [246, 247].

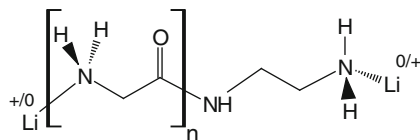


Fig. 19 $\text{Li}_2^{+/0}$ redox pairs separated by polypeptide chains of increasing lengths investigated in [226]. Reproduced with permission from [226]

In [226] we investigated the decay of the electronic coupling as a function of the donor to acceptor distance for the electron transfer between two lithiums separated by a polyglycine peptide of increasing length (Fig. 19). The simulations were run in the gas and aqueous phases thanks to the QM/MM scheme described in Sect. 5. Note that the BOMD simulations were biased so as to sample exclusively the molecular configurations corresponding to the degeneracy of the two diabatic states. Note also that the lithium and its first coordination shell were fixed during the simulations to control the D to A distance. In this work we also reported the first estimates of the characteristic decay times of the Franck–Condon factors by means of cDFT/MM MD simulations. This term reflects the time the molecular system remains in the Franck–Condon region that is active for the ET. It can be computed by the following expression:

$$\tau_{\text{dec}} = \left[\left\langle \sum_n \frac{1}{2a_n\hbar^2} (F_{1n} - F_{2n})^2 \right\rangle_T \right]^{-1/2}, \quad (31)$$

where F_{xn} and a_n , respectively, denote the forces felt by the n th degree of freedom in the electronic state 1 or 2 and the typical width of the wave packet associated with this degree of freedom. Actually Eq. (31) provides a means to evaluate the decoherence time in the context described in Sect. 4.3, but under a short time approximation and within the high temperature limit [139]. Note that this approximate equation enables an estimation of the decoherence time without resorting to computationally expensive diverging trajectories (see Sect. 4.3).

The graphs shown in Fig. 20 are in line with common knowledge of the electronic coupling decaying exponentially with the D to A distance. Characteristic decays of 0.52 and 0.58 \AA^{-1} are found for the gas phase and aqueous phase, respectively, in good agreement with the experimental values reported in the literature [248] for LRET rates through β -strands. The characteristic decoherence times are found to be around 4.5 and 1.5 fs in the gas and aqueous phases, but quite interestingly independent of the bridge length. This constant trend can be understood by looking at Eq. (31). The atoms that contribute the most to decoherence are those which feel the different forces in the two redox states. Therefore, the molecular fragments that are far from the donor and from the acceptor do not appreciably contribute to decoherence. And, similarly, when the bridge length is increased the new intervening atoms contribute less and less to decoherence. Recall, however, that in these simulations the most contributing atoms were fixed

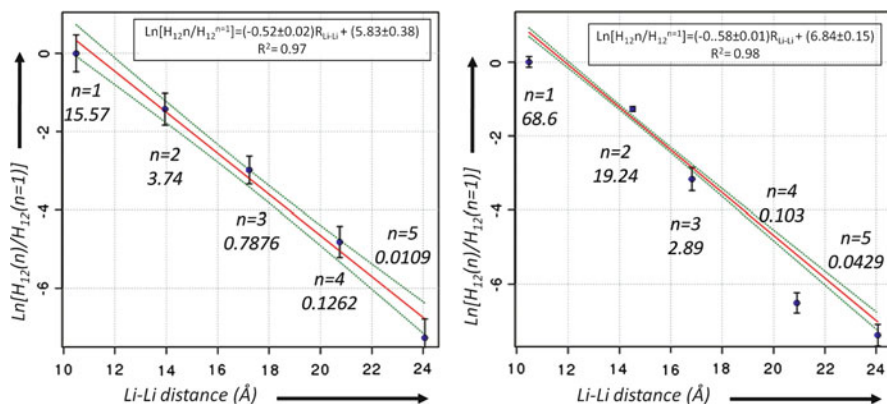


Fig. 20 Linear decay of the electronic coupling logarithm for the inter-lithium ET for different donor-to-acceptor distances in the gas phase (*Left*) and in water (*Right*). Each point represent an average over 1 ps of BOMD simulation in the NVT ensemble ($T = 300$ K). The values of the average H_{DA} on the graphs are given in cm^{-1} . Reproduced with permission from [226]

in space for practical reasons so that the actual characteristic decoherence time for this reaction is probably much below 1 fs. In fact the study reported in [226] is more to be seen as a proof of principles of the use of cDFT/MM MD approaches for the estimation of decoherence times in large molecular systems. Investigations of decoherence times in real enzymatic systems are currently being performed in our laboratories using this methodology and will be reported in due course.

6.4 Other Applications of cDFT

Apart from the modeling of electron transfer reactions, the cDFT formalism has been applied to many other chemical problems (see [215]). So far we have considered constraints applied to the atomic charges only; however, the cDFT formalism also allows one to enforce the spin density on specific atoms. Therefore, the method provides a valuable tool to prepare electronic guesses for (nonconstrained) SCF-DFT computation with desired spin-density properties. Such a possibility is certainly of great added value for the modeling of complex electronic structures like those encountered in bio-inorganic metal complexes. In addition, the ability of cDFT to provide trustworthy values of the magnetic coupling constants within polynuclear clusters such as those found in many metalloenzymes holds great promise for future applications in this field [249].

To conclude this section, and although we do not cover the TD-DFT and Tight-Binding DFT methods in this chapter we mention that extensions of the cDFT ideas of Dederich et al. have been developed for these methods. More generally, there is little doubt that further development of cDFT in the coming years will allow its application to numerous biologically relevant systems.

7 Interpretational-DFT

As seen throughout this chapter, DFT represents a powerful computational tool. The primary outputs of DFT computations are the energies of the molecules. Then, when coupled to geometry optimizers or BOMD engines, the stable conformers or Boltzmann ensembles can be determined, giving access, for example, to the free energy balance of a chemical reaction (or its free energy of activation). In addition, the calculations of spectroscopic properties such as, for example, the infra-red or Raman vibrational spectra provide valuable supplements to understand the chemical systems of interest. Properties such as polarizabilities, hyperpolarizabilities [250], NMR (Nuclear Magnetic Resonance) shieldings [251, 252], or NQR (Nuclear Quadrupole Resonance) [253] energy splittings, to mention but a few, are other quantities that can be efficiently computed with DFT. Having recognized the immense merits of DFT it is also worth noting that the outcomes of these first principles computations are not always directly connected to some of the most fruitful concepts of chemistry like the notion of electronegativity, of hardness, or even the notion of chemical bonding. Many efforts have been spent to fill this gap leading for example to the so-called conceptual DFT [254, 255] framework or the topological analysis.

In this section, we have chosen to focus on the interpretational tools that are based on the topological analysis of well-defined functions like the electronic density, ρ and the Electron Localization Function (ELF) η . The former is a direct output of a DFT computation or of experimental measurements while the latter is a function introduced by Becke and Edgecombe as a mean to measure the electron localization in molecular systems [256]

$$\eta = \frac{1}{\left[1 + \left(\frac{D}{D_h}\right)^2\right]}. \quad (32)$$

with

$$D = \frac{1}{2} \sum_i |\nabla \varphi_i|^2 - \frac{1}{8} \frac{(\nabla \rho)^2}{\rho}, \quad (33)$$

$$D_h = \frac{3}{10} (3\pi^2)^{5/3} \rho^{5/3}, \quad (34)$$

where D and D_h represent the curvature of the electron pair density for electrons with identical spins for the system under study, and a homogeneous electron gas with the same density, respectively. The ELF function can be interpreted as a measure of the Pauli repulsion in the atomic or molecular space and reflects the probability of finding opposite spin electron pairs. The function η is bounded between 0 and 1.

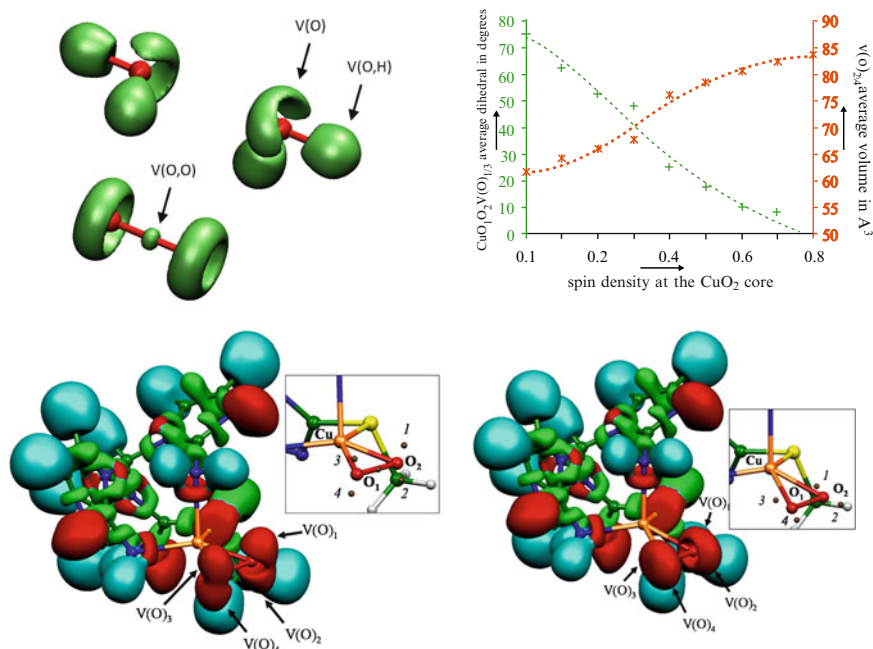


Fig. 21 *Upper left:* ELF isosurface (0.7) for the $O_2(H_2O)_2$ system showing the valence basins within the molecular fragments. *Upper right:* cDFT/ELF analysis showing the rotation of the dioxygen lone pairs and their volumes changes as a function of the spin density imposed at the CuO_2 core. *Bottom:* Orientation of the dioxygen lone pairs revealed by the ELF topological analysis in the singlet (*Left*) and triplet (*Right*) spin states within minimal model of noncoupled monooxygenases. (isosurface for ELF = 0.8). *Insets:* zooms on the CuO_2 core showing the localization of the O_2 valence basin attractors. *Color code:* red for nonbonding (lone pair), green for bonding, and light blue for protonated bond. Reproduced with permission from [259]

Topological analysis, as defined in the context of the theory of dynamical systems, was pioneered by Bader in the 1970s in the case of the electronic density. The approach proposed by Bader performs a partition of the electrons within the molecular space into basins associated with each atom. Therefore, it provides a mathematically grounded way to define an atom within a molecule. Following a similar approach, Savin and Silvi showed that the topological analysis of the ELF leads to the identification of molecular basins that can be connected to the chemical intuitive picture of bonds, lone pairs, π -systems [257]. For example, the topological analysis allows one to distinguish between core basins (labeled $C(X)$, X being a nuclei) encompassing the nuclei and valence regions. These latter then split into bonding basins connected to more than one atomic center (labeled $V(X, Y, \dots)$, X, Y, \dots being nuclei) and nonbonding basins (lone pairs) connected to only one atomic center (labeled $V(X)$). These valence basins match closely the nonbonding (lone pairs) and bonding domains of Gillespie's VSEPR (Valence Shell Electron Pair Repulsion) model [258]. The ELF isosurface ($\eta = 0.7$) is showed in Fig. 21 for a

triplet water molecule surrounded by two water molecules. The dioxygen lone pairs (V(O)) and the covalent O–O bond are clearly visible.

As a further illustration we take here the example of the $[(\text{N}_2\text{S})\text{CuO}_2]^+$ complex which was presented in Sect. 4 (Fig. 10). As already mentioned the transition between the triplet and singlet spin state is a prerequisite in the dioxygen activation process. Recently, we reported a systematic study of the ELF topology on a series of metallated dioxygen complexes (including, Fe, Ni, Pd, and Cu atoms) [259]. We observed that upon spin transition the dioxygen lone pairs experienced a systematic rotation around the metal-O₂ plane making them more or less accessible toward exogenous substrate. In addition, an important variation of the lone pair spatial extension was observed upon this transition. This trend could be related to the molecular orbital diagrams of the complexes. We also reported in this communication, the ELF topological analysis on constrained DFT densities where the spin density of the CuO₂ triad was constrained between 0.1 and 0.8 thereby confirming our initial interpretation, leading to the conclusion that metalloenzymes could use spin state control to tune the regioselectivity of substrate oxidations.

More generally the ELF topological analysis has been found to be a powerful interpretational framework to rationalize the regioselectivity of numerous types of chemical reactions [260, 261]. We already mentioned the approach of Bader that is based on the electronic density itself. Other scalar functions such as the electrostatic potential [262] have also been the subject of topological analysis.

8 Conclusions and Perspectives

In this review we have seen examples representing three main types of biomolecules and biomodels: proteins, nucleic acids, and lipids. Although it is not always classified as a “biomolecule,” water represents a fourth major type and it has been center stage in many of the examples we have treated. For each of these classes of biomolecules, DFT has played a major role in our work and in that of other workers. But this is not a one-man show; we have also shown how DFT can be combined with molecular dynamics, either in the Born–Oppenheimer-MD approach or in hybrid QM/MM methodologies. And we have shown examples that go beyond strictly Kohn–Sham DFT in the use of constraints that allow connections with other theories and concepts, notably the Marcus theory of electron transfer. Finally, we have given a glimpse of some of the tools that can be used to analyze and interpret the DFT-based computations.

While there has been very remarkable recent progress in DFT towards descriptions of ever more complex models for biological processes, we do not view DFT as a panacea. Much work remains to develop both the functionals necessary for quantitative work and the appropriate suite of modeling tools within which DFT can perform. More and more the lines are blurring between DFT (and quantum chemistry in general) and the more conventional molecular dynamics and statistical mechanics methodologies that use empirical force fields. Much more

work remains to develop powerful and efficient hybrid methodologies that will, in the fullness of time, allow more complete and sophisticated models combining all four classes of biomolecules. We find the prospect of combining proteins, lipids, nucleic acids, and water (not to neglect ions and small molecules...) into a multiscale suite of methodologies to be a very exciting one. Much progress has been made and we hope that our review will help to introduce new workers into this vibrant field.

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Density Functional Theory and Molecular Interactions: Dispersion Interactions

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Abstract We are definitely witnessing an ever-increasing need to study dispersion molecular interactions that govern a weakly bound molecular world within the density functional theory. This chapter outlines the basic approaches currently undertaken to resolve this density functional paradigm.

Contents

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|-----|---|----|
| 1 | Density Functional Theory: Background | 65 |
| 2 | Kohn–Sham Density Functional Theory: The Hohenberg–Kohn Theorem | 68 |
| 3 | Dispersion Molecular Forces: Introduction | 73 |
| 4 | Dispersion-Corrected DFT Approaches | 79 |
| 5 | Local-Scaling Transformations’ DFT | 81 |
| 5.1 | Mathematical Preliminaries: Local-Scaling Transformations | 81 |
| 5.2 | One-Electron Densities: Definition | 83 |
| 5.3 | Many-Electron Wavefunctions and Concept of Orbit | 86 |
| 5.4 | Energy Density Functional and Variational Principle | 87 |
| 6 | Overlook | 92 |
| | References | 92 |

1 Density Functional Theory: Background

In the last two decades, since Walther Kohn was awarded the Nobel Prize 1991 in Chemistry for the density functional theory (DFT), the latter has become the most popular and useful computational approach to study many-electron systems in the ground states because of its physically transparent underlying concept and lower computational cost [1–8]. The Kohn–Sham version of DFT [1–3, 7] is the most widely used many-body method for electronic structure calculations of atoms, molecules, solids, and solid surfaces.

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Generally speaking, a molecule \mathbf{M} is a stable quantum Coulomb system that consists of the following two subsystems:

- The electronic—of N electrons of the mass m_e and the charge $-e$ which positions in the spin-configurational space are determined by the corresponding radii vectors $\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_N$ where each $\mathbf{r}_i, i = 1, 2, \dots, N$ belongs to the real three-dimensional space \mathbb{R}^3 and the spins $\sigma_1, \sigma_2, \dots, \sigma_N$ where each $\sigma_i, i = 1, 2, \dots, N$ takes the value from $\mathbb{Z}_2 = \{\pm 1/2\}$, the discrete two-dimensional spin space
- The nuclear—of M nuclei carrying the nuclear charges $\{Z_\alpha\}_{\alpha=1}^M$ and located at $\{\mathbf{R}_\alpha \in \mathbb{R}^3\}_{\alpha=1}^M$.

According to Löwdin's definition [9, 10]: “A system of electrons and atomic nuclei is said to form a molecule if the Coulombic Hamiltonian H' —with the center of mass motion removed—has a discrete ground-state energy E_0 ” (see also [11–13] and references therein) where the total Hamiltonian $H := \hat{H} = \hat{H}_e + \hat{T}_{nn} + \hat{U}_{nn}$ is, respectively, the sum of the electronic Hamiltonian operator, the nuclear kinetic energy operator, and the nuclear–nuclear Coulomb interaction energy operator. Consider, within the Born–Oppenheimer approximation, the electronic Hamiltonian operator (in the atomic units) of \mathbf{M} :

$$\hat{H}_e = \hat{T}_e + \hat{U}_{ee} + \hat{V}_{en} = -\frac{1}{2} \sum_{i=1}^N \nabla_{\mathbf{r}_i}^2 + \sum_{1 \leq i < j}^N \frac{1}{|\mathbf{r}_i - \mathbf{r}_j|} + \sum_{i=1}^N \hat{v}(\mathbf{r}_i), \quad (1)$$

where \hat{T}_e is the nuclear kinetic energy operator, \hat{U}_{ee} the nuclear–nuclear Coulomb interaction energy operator, and the “external,” electron–nuclear potential is defined as

$$\hat{v}(\mathbf{r}_i) := \sum_{\alpha=1}^M \frac{Z_\alpha}{|\mathbf{r}_i - \mathbf{R}_\alpha|}. \quad (2)$$

\hat{H}_e acts on the class \mathcal{L}_N of “admissible” N -electron wavefunctions $\Psi(\mathbf{r}_1, s_1; \dots; \mathbf{r}_N, s_N)$ obeying the following conditions:

(Fi) the wavefunction normalization:

$$\langle \Psi | \Psi \rangle = \sum_{s_1, \dots, s_N} \int d^3\mathbf{r}_1 \dots \int d^3\mathbf{r}_N |\Psi(\mathbf{r}_1, s_1; \dots; \mathbf{r}_N, s_N)|^2 < \infty \quad (3)$$

implying that $\mathcal{L}_N \subset \mathcal{L}_\sigma^2(\mathbb{R}^{3N} \otimes \mathbb{Z}_2^N)$, the Hilbert space of antisymmetric, square-integrable N -electron wavefunctions. Henceforth it is assumed that an arbitrary $\Psi \in \mathcal{L}_N$ is normalized to unity: $\langle \Psi | \Psi \rangle = 1$;

(Fii) the boundness from below of the expectation value $\langle \Psi | \hat{H}_e | \Psi \rangle > -\infty$: In fact, (Fii) results from the aforementioned definition of molecule which lowest energy is finite. If \hat{U}_{ee} and \hat{V}_{en} are of Coulomb type, (Fii) is equivalent to

$$T_e[\Psi] = \langle \Psi | T_e | \Psi \rangle < \infty \quad (4)$$

implying that $\Psi \in \mathcal{L}_N$ is a differentiable function of all spatial coordinates, together with each component of $\nabla_{\mathbf{r}}, \Psi \in \mathcal{L}_N$.

One can prove [4, 14] that the conditions (Fi) and (Fii) fully determine \mathcal{L}_N of “admissible” N -electron wavefunctions where the energy functional

$$E[\Psi] \equiv \langle \Psi | \hat{H}_e | \Psi \rangle \quad (5)$$

is thus well defined. Its lowest energy, the infimum, equal to the ground-state electronic energy E_0 as the lowest eigenenergy of the N -body Schrödinger equation

$$\hat{H}_e \Psi_0 = E_0 \Psi_0, \quad (6)$$

is attained at the ground-state electronic wavefunction Ψ_0 , that is

$$E_0 \equiv \inf_{\Phi \in \mathcal{L}_N} \{E[\Phi]\} = E[\Phi]|_{\Phi=\Psi_0 \in \mathcal{L}_N}. \quad (7)$$

The stationary quantum mechanical variational principle then reads as

$$\delta E[\Phi]|_{\Phi=\Psi_0} = 0. \quad (8)$$

The basic postulate of the many-electron density functional theory [1–8] suggests, first, the existence of the so-called functional

$$\mathcal{E}[\rho(x)] = \begin{cases} \mathcal{E}[\rho(\mathbf{r})] & \text{spin – restricted functional} \\ \mathcal{E}[\rho_{\uparrow}(\mathbf{r}), \rho_{\downarrow}(\mathbf{r})] & \text{spin – polarized functional} \end{cases} \quad (9)$$

that has the meaning of the energy and depends, in some functional manner, on one-electron density $\rho(\mathbf{r})$,

$$\rho_{\Psi}(\mathbf{r}) := N \sum_{s_1, \dots, s_N} \int d^3 \mathbf{r}_2 \dots \int d^3 \mathbf{r}_N |\Psi(\mathbf{r}, s_1; \mathbf{r}_2, s_2; \dots; \mathbf{r}_N, s_N)|^2, \Psi \in \mathcal{L}_N \quad (10)$$

or on its both spin components, $\rho_{\Psi_{\uparrow}}(\mathbf{r})$ and $\rho_{\Psi_{\downarrow}}(\mathbf{r})$,

$$\begin{aligned} \rho_{\Psi_s}(\mathbf{r}) &:= \rho_{\Psi}(\mathbf{r}, s) \\ &= N \sum_{s_1, \dots, s_N} \int d^3 \mathbf{r}_2 \dots \int d^3 \mathbf{r}_N |\Psi(\mathbf{r}, s; \mathbf{r}_2, s_2; \dots; \mathbf{r}_N, s_N)|^2, s = \uparrow, \downarrow. \end{aligned} \quad (11)$$

The latter yield together $\rho_{\Psi}(\mathbf{r}) = \rho_{\Psi_{\uparrow}}(\mathbf{r}) + \rho_{\Psi_{\downarrow}}(\mathbf{r})$. Each $\rho_{\Psi_s}(\mathbf{r})$ is normalized to N_s so that $N_{\uparrow} + N_{\downarrow} = N$. The second suggestion is that the infimum of $\mathcal{E}[\rho(\mathbf{r})]$ does exist and

$$E_0 \equiv \inf_{\Phi \in \mathcal{L}_N} \{E[\Phi]\} = E[\Phi]_{\Phi=\Psi_0} = \inf_{\rho \in \mathcal{P}_N} \{\mathcal{E}[\rho(\mathbf{r})]\} = \mathcal{E}[\rho_\Phi(\mathbf{r})]_{\Phi=\Psi_0} \quad (12)$$

where \mathcal{P}_N is a set of one-electron densities associated with \mathcal{L}_N (see below). Formally, this postulate looks rather strong, however it is widely accepted that it is guaranteed by the Hohenberg–Kohn theorem [1] (for the new proof of the Hohenberg–Kohn theorem see [15, 16]).

Equation (9) assumes the existence of the “Functional mapping”

$$F : E[\Psi] \mapsto \mathcal{E}[\rho_\Psi(\mathbf{r})] \quad (13)$$

that implicitly presumes the existence of the “Variable mapping”

$$\Psi \leftrightarrow \rho_\Psi(\mathbf{r}). \quad (14)$$

Obviously, the mapping (14) is valid if, first, there are defined the sets of “variables” on its left- and right-hand sides. Second, the symbol \leftrightarrow does not mean at all that this is precisely a one-to-one correspondence. The sub-mapping of (14), $V:\Psi \rightarrow \rho_\Psi(\mathbf{r})$, is given by the reduction mapping, either (10) or (11), that is, $\rho_\Psi(\mathbf{r}) = V(\Psi)$ and $\mathcal{P}_N \equiv V\mathcal{L}_N$. Besides, the reduction mapping has another facet—this is a so-called N -representability: any one-electron density obtained via V possesses its own image in \mathcal{L}_N . Generally speaking, the inverse mapping V^{-1} is one-to-many, that is, a given one-electron density has many preimages in \mathcal{L}_N . It is trivial to show this. Let us consider any stable two-electron system which ground-state wavefunction and one-electron density are $\Psi_0(\mathbf{r}_1, \mathbf{r}_2)[\alpha(s_1)\beta(s_2) - \beta(s_1)\alpha(s_2)]$ and $\rho_0(\mathbf{r})$, respectively. The two-electron Slater determinant $\sqrt{\rho_0(\mathbf{r}_1)\rho_0(\mathbf{r}_2)}[\alpha(s_1)\alpha(s_2) - \beta(s_1)\alpha(s_2)]/2$ possesses the same one-electron-density $\rho_0(\mathbf{r})$ as well. Q. E. D. The Hohenberg–Kohn theorem [1] (see also [15, 16]) states however that there exists a one-to-one correspondence between the ground-state wavefunctions and ground-state densities.

2 Kohn–Sham Density Functional Theory: The Hohenberg–Kohn Theorem

The Hohenberg–Kohn theorem [1] underlies the foundation of the density functional theory [2–7], the Kohn–Sham density functional theory. On p. B864 of their work [1], Hohenberg and Kohn state that they “... develop an exact formal variational principle for the ground-state energy, in which the density” $\rho(\mathbf{r})$ (in a widely accepted notation) “is the variable function. Into this principle enters a universal functional” $F[\rho(\mathbf{r})]$, “which applies to all electronic systems in their ground state no matter what the external potential is.” Following Hohenberg and Kohn [1], let us consider “a collection of an arbitrary number of electrons, enclosed in a large box and moving under the influence of an external potential $v(\mathbf{r})$ and

mutual Coulomb repulsion.” The total Hamiltonian H of a given N -electron system is written as Eq. (1), viz.

$$H = T_e + V_{ee} + V, \quad (15)$$

where T_e is the kinetic energy operator of N electrons, V_{ee} is the interelectronic Coulomb operator, and

$$V = \sum_{i=1}^N v(\mathbf{r}_i) \quad (16)$$

is the total external potential. Hohenberg and Kohn [1] further assume (p. B865) that H possesses the least bound-state (ground-state) wavefunction $\Psi_0(\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_N) \in \mathcal{H}_N$ (spins are omitted for simplicity) and the latter is nondegenerate. \mathcal{H}_N is the Hilbert space of square integrable N -electron wavefunctions and $\mathbf{r}_i \in \mathbb{R}^3$, $i = 1, 2, \dots, N$. Let us define the corresponding ground-state one-electron density [3]

$$\rho_0(\mathbf{r}) \equiv \int \prod_{i=2}^N d^3\mathbf{r}_i |\Psi_0(\mathbf{r}, \mathbf{r}_2, \dots, \mathbf{r}_N)|^2, \quad (17)$$

“which is clearly a functional of $v(\mathbf{r})$ ” ([1], p. B865), that is, there exist such mappings

$$v(\mathbf{r}) \Rightarrow \Psi_0(\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_N) \Rightarrow \rho_0(\mathbf{r}). \quad (18)$$

Hohenberg–Kohn Theorem [1]. “ $v(\mathbf{r})$ is a unique functional” of $\rho(\mathbf{r})$, “apart from a trivial additive constant.”

Proof ([1], p. B865). “The proof proceeds by *reductio ad absurdum*.” We assume the existence of two “external” potentials $v_1(\mathbf{r})$ and $v_2(\mathbf{r})$ such that

$$v_1(\mathbf{r}) \neq v_2(\mathbf{r}) + \text{constant}. \quad (19)$$

Via Eqs. (16) and (15), $v_1(\mathbf{r})$ and $v_2(\mathbf{r})$ define the Hamiltonians H_1 and H_2 associated with two different N -electron systems. Let us further assume the existence of the ground-state normalized wavefunctions $\Psi_0^{(1)} \in \mathcal{H}_N$ and $\Psi_0^{(2)} \in \mathcal{H}_N$ of H_1 and H_2 , respectively. By virtue of Eq. (17), $\Psi_0^{(1)}$ and $\Psi_0^{(2)}$ yield the corresponding ground-state one-electron densities $\rho_0^{(1)}$ and $\rho_0^{(2)}$. Hohenberg and Kohn [1] finally assume that

- (i) $\Psi_0^{(1)} \neq \Psi_0^{(2)}$
- (ii) $\rho_0^{(1)}(\mathbf{r}) = \rho_0^{(2)}(\mathbf{r}) = \rho_0(\mathbf{r})$.

Applying the Rayleigh–Ritz variational principle, one obtains

$$\begin{aligned}
 E_0^{(1)} &= \langle \Psi_0^{(1)} | H_1 | \Psi_0^{(1)} \rangle \stackrel{(i)}{<} \langle \Psi_0^{(2)} | H_1 | \Psi_0^{(2)} \rangle \\
 &\stackrel{Eq.(5)}{=} \langle \Psi_0^{(2)} | H_2 | \Psi_0^{(2)} \rangle + \langle \Psi_0^{(2)} | V_1 - V_2 | \Psi_0^{(2)} \rangle \\
 &= E_0^2 + \int d^3r [v_1(\mathbf{r}) - v_2(\mathbf{r})] \rho_0(\mathbf{r})
 \end{aligned} \tag{20}$$

and

$$\begin{aligned}
 E_0^{(2)} &= \langle \Psi_0^{(2)} | H_2 | \Psi_0^{(2)} \rangle \stackrel{(i)}{<} \langle \Psi_0^{(1)} | H_2 | \Psi_0^{(1)} \rangle \\
 &\stackrel{Eq.(5)}{=} \langle \Psi_0^{(1)} | H_1 | \Psi_0^{(1)} \rangle + \langle \Psi_0^{(1)} | V_2 - V_1 | \Psi_0^{(1)} \rangle \\
 &= E_0^1 + \int d^3r [v_1(\mathbf{r}) - v_2(\mathbf{r})] \rho_0(\mathbf{r})
 \end{aligned} \tag{21}$$

where the used formulas are indicated above the signs.

Hohenberg and Kohn then conclude ([1], p. B865) that adding (20) to (7) “leads to the inconsistency”

$$E_0^{(1)} + E_0^{(2)} < E_0^{(1)} + E_0^{(2)}, \tag{22}$$

and therefore, (22) implies that the assumption (ii) fails. “Thus $v(\mathbf{r})$ is (to within a constant) a unique functional of” $\rho(\mathbf{r})$, “since, in turn, $v(\mathbf{r})$ fixes H we see that the full many-particle ground state is unique functional of” $\rho(\mathbf{r})$. Q. E. D.

Examine Eq. (22). It is obviously self-contradictory (see also [15–17]). Equation (22) is deduced under the assumption that (19) is true together with the to-be-refuted assumptions (i) and (ii) both composing the negation of the Hohenberg–Kohn theorem. Equation (22) then appears to be absurd in a sense of being obviously false and therefore the statement of the Hohenberg–Kohn theorem is true. This might, logically speaking, imply that one of the to-be-refuted assumptions, (i) or (ii), or simultaneously both, (i) and (ii), lead to the contradiction with (19) *or* they are a priori invalid in a sense that one of them *or* both are incompatible with (5) and therefore, the statement of the Hohenberg–Kohn theorem is not true unless it is proved in the other way (see below). Explicitly, all these cases are the following:

$$(I) \Psi_0^{(1)} = \Psi_0^{(2)} = \Psi_0.$$

This directly gives $\rho_0^{(1)} = \rho_0^{(2)} = \rho_0$, that is, (ii) does hold. This also yields that

$$V_1 \equiv V_2 \equiv E_0 - \frac{(T_e + V_{ee})\Psi_0}{\Psi_0} \tag{23}$$

if V_1 and V_2 are multiplicative operators, as suggested by Eq. (16). Equation (23) clearly contradicts (19). However, there is no “inconsistency” because the last terms in the last lines of Eqs. (20) and (21) simply vanish.

- (II) $\Psi_0^{(1)} \neq \Psi_0^{(2)}$ and $\rho_0^{(1)} \neq \rho_0^{(2)}$

This is precisely in the line of the original reasoning by Hohenberg and Kohn [1] proving that different external potentials determine different ground-state one-electron densities.

- (III) $\Psi_0^{(1)} = \Psi_0^{(2)}$ and $\rho_0^{(1)} \neq \rho_0^{(2)}$

These two relations contradict to each other due to (17).

- (IV) A self-contradiction (*ad absurdum*) of Eq. (22) might also mean that the to-be-refuted assumptions (i) or/and (ii) of the Hohenberg–Kohn theorem are self-contradictory with Eq. (19) and this is precisely the case of many-electron Coulomb systems with Coulomb-type class of external potentials. In other words, the original *reductio ad absurdum* proof of the Hohenberg–Kohn theorem based on the assumption (19) is incompatible with the *ad absurdum* assumption (ii) since the Kato theorem is valid for such systems [18].

The Kato theorem [18] (see also [19–29]) determines the character of the singularity of the exact N -electron wavefunction at the electron–nucleus coalescence where the external potential $v(\mathbf{r})$ of the Coulomb form (see Eq. (2.2) and the conditions (i) and (ii) on p. 154 and Theorem I on p. 156 of [19]; in atomic units)

$$v(\mathbf{r}) = - \sum_{\alpha=1}^M \frac{Z_{\alpha}}{|\mathbf{r} - \mathbf{R}_{\alpha}|} \quad (24)$$

is singular. In Eq. (24), the α th nucleus with the nuclear charge Z_{α} is placed at $\mathbf{R}_{\alpha} \in \mathbb{R}^3$. Any N -electron eigenwavefunction Ψ of H with $v(\mathbf{r})$ of the form (24) and its one-electron density ρ_{Ψ} then satisfy the electron–nucleus cusp condition, that is, their corresponding nonvanishing radial logarithmic derivatives obey at \mathbf{R}_{α} the following relationships:

$$\begin{aligned} \frac{d}{dr_i} \log \Psi(\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_i, \dots, \mathbf{r}_N) \big|_{\mathbf{r}_i = \mathbf{R}_{\alpha}} &= -Z_{\alpha}, \quad i = 1, 2, \dots, N \\ \frac{d}{dr} \log \rho_{\Psi}(\mathbf{r}) \big|_{\mathbf{r} = \mathbf{R}_{\alpha}} &= -2Z_{\alpha}. \end{aligned} \quad (25)$$

Therefore, the true one-electron density of the given N -electron system exhibits cusps (local maxima) at the positions of the nuclei. Analyzing the topology of the given ground-state one-electron density $\rho_0(\mathbf{r})$ over the whole coordinate space \mathbb{R}^3 , one determines the positions of its cusps and evaluates the lhs of Eq. (25) (the last one) at these points. Altogether, the positions of the electron–nucleus cusps [as being always negative, see Eq. (25)] and the halves of the radial logarithmic derivatives of $\rho_0(\mathbf{r})$, taken with the opposite sign at these points, fully determine the external potential $v(\mathbf{r})$, Eq. (24), of the given system. This constitutes a naïve interpretation of the Hohenberg–Kohn theorem originally proposed by Coleman [30], Bamzai and

Deb [31], Smith [32], and E. Bright Wilson (quoted by Löwdin [33]; for the recent applications of the Kato theorem to the Hohenberg–Kohn theorem see also [34–36]). Therefore, if a given pair of N -electron systems with the Hamiltonians H_1 and H_2 of the type (1) are characterized by the same ground-state one-electron densities (\equiv to-be-refuted assumption (ii)), their external potentials $v_1(\mathbf{r})$ and $v_2(\mathbf{r})$ of the form (24) are identical. The latter contradicts (19) and hence, the assumption (ii) cannot be used in the proof via *reductio ad absurdum* of the Hohenberg–Kohn theorem together with the assumption (19). In other words, they are Kato-type incompatible with each other.

Vice versa, the nuclei of the given N -electron system are isolated 3D point attractors behaving topologically as critical points of rank three and signature minus three [37]. However, there exist some “particular many-electron systems” which show local maxima of their ground-state one-electron density at non-nuclear positions ([38–48] and references therein). These local non-nuclear maxima might be the true ones or might appear as a consequence of an incomplete, inadequate quantum mechanical treatment. Therefore, despite the present conclusion that the original proof of the Hohenberg–Kohn theorem by *reductio ad absurdum* is flawed in a sense that its to-be-refuted assumption (ii) is incompatible, by virtue of the Kato theorem, with the assumption (19) (for a similar proof of the ensemble generalization of the Hohenberg–Kohn theorem see Sect. II of [16]), the Kato theorem itself corroborates the existence of the one-to-one correspondence between the Coulomb-type class of external potentials (10) and the ground-state one-electron densities for nearly all many-electron except probably the aforementioned “particular” ones.

According to the work [1] by Hohenberg and Kohn, the Hohenberg–Kohn theorem implies the existence of the universal energy density functional for any isolated many-electron Coulomb system. This statement has been usually interpreted as the second Hohenberg–Kohn theorem [3]. In the density functional theory, there exist two rigorous constructions of the universal energy density functionals based on their own rigorous proofs of the Hohenberg–Kohn theorem. This is the Levy–Lieb energy density functional [49–54] and the universal energy density functional based on the group of the local-scaling deformations in \mathbb{R}^3 consisting of topological deformations mapping or topologically deforming any pair of one-electron densities to each other [4] (see also Sect. 5). The related Jacobian of such deformation gives rise to the first-order nonlinear differential equation, a so-called “Jacobian equation” [55–63] whose solution, namely, the corresponding deformation, does exist within the present approach and it is unique [4]. Solving the “Jacobian equation” enables to determine the deformation for any pair of well-behaved one-electron densities and to consistently extend the action of the local-scaling deformation group onto \mathcal{H}_N [49]. This larger local-scaling transformation group partitions \mathcal{H}_N into disjoint classes, orbits. All orbits exhaust \mathcal{H}_N and within each orbit, there establishes the one-to-one correspondence between its wavefunctions and the one-electron densities. That is, these orbits are endowed with the characteristic that there are no two wavefunctions belonging to the same orbit that have the same density. Each orbit, say $\mathcal{O}^{[\alpha]}$, is determined by its

generator wavefunction $\Psi_g^{[\alpha]}$. Therefore, for a given orbit $\mathcal{O}^{[\alpha]}$, one defines the energy density functional $E_\alpha[\rho(\mathbf{r})]$ as merely the restriction of the energy functional $E[\Psi]$ on those wavefunctions that belong to the α th orbit, $\Psi \in \mathcal{O}^{[\alpha]}$. It is trivial to prove that, first, there are as many different energy density functionals as the orbits in \mathcal{H}_N , and second, that each density functional $E_\alpha[\rho(\mathbf{r})]$ implicitly depends on the generator wavefunction $\Psi_g^{[\alpha]}$. Evidently, the true ground-state N -electron wavefunction of the given Hamiltonian operator H belongs to a certain orbit, called the Hohenberg–Kohn one, $\mathcal{O}^{[\text{HK}]}$ [4]. Within $\mathcal{O}^{[\text{HK}]}$, the Levy–Lieb energy density functional [49] exactly coincides with $E_{\text{HK}}[\rho(\mathbf{r})]$ that is defined within the local-scaling deformation approach [64–75]. The explicit form of any energy density functional $E_\alpha[\rho(\mathbf{r})]$ for any α has been obtained and the corresponding variational Euler–Lagrange equation has been also derived in [64–75] (see also Chaps. 7 and 8 of [4]). The rigorous mathematical framework of the local-scaling deformation approach to the density functional theory based on the “Jacobian equation” has recently been elaborated in [55, 57–63]. The local-scaling deformation approach to the density functional theory has been also generalized to include spin densities, for the momentum space representation, for excited states, for the fractional occupation numbers, and finally to study nonadiabatic effects. The corresponding analogue of the Kohn–Sham approach has been also formulated in terms of the orbit generators (see Chap. 8 in [4]). A variety of theoretical and computational applications of the local-scaling deformation density functional theory to atoms and molecules has recently been elaborated as well (see [76–87] and references therein).

However, in the computational perspective, the Kohn–Sham DFT remains the most widely used many-body approach to conduct the electronic structure calculations of atoms, molecules, solids, and solid surfaces. In the context of this DFT, only the exchange–correlation energy must be approximated as a functional of the electron density [8]. The simplest approximation is the local spin density (LSD) approximation [88] which uses the local electron spin densities $\rho_\uparrow(\mathbf{r})$ and $\rho_\downarrow(\mathbf{r})$, as in (9), as the only ingredients. In the development of DFT, a significant step was the introduction of the density gradients $\nabla_{\rho_\uparrow}(\mathbf{r})$ and $\nabla_{\rho_\downarrow}(\mathbf{r})$ as additional local ingredients in the generalized gradient approximation or shortly GGA [89–91].

3 Dispersion Molecular Forces: Introduction

Some people think that there is an air between molecules.

A. S. Kompaneets [92]

According to our current knowledge, there exist four basic types of forces: gravitational, electromagnetic, strong, and weak. The electromagnetic force is the force that binds the electrons and the nuclei inside the atom, and the atoms inside a molecule, and that governs the interaction between atoms and molecules. These are referred to as the so-called intermolecular interactions [92–94, 186, 187]. Indeed,

Table 1 Van der Waals radii R_{vdW} (in Å) of atoms

| Atom | R_{vdW} | Atom | R_{vdW} | Atom | R_{vdW} | Atom | R_{vdW} | Atom | R_{vdW} | Atom | R_{vdW} |
|------|------------------|------|------------------|------|------------------|------|------------------|------|------------------|------|------------------|
| H | 1.20 | He | 1.30 | N | 1.50 | O | 1.40 | F | 1.35 | Ne | 1.40 |
| P | 1.90 | S | 1.85 | Cl | 1.80 | Ar | 1.70 | As | 2.00 | Se | 2.00 |
| Br | 1.95 | Kr | 1.80 | Sb | 2.20 | Te | 2.20 | I | 2.15 | Xe | 2.05 |

intermolecular interactions play the important role in the world: it is true that we “touch” them everywhere, at a macroscopic scale, in our everyday life. It is rather hard to imagine what would be the world without them—obviously, the situation would be significantly more dramatic than to have the ideal gases that surround us! Simply imagine that would be no such molecules as the DNA and RNA. Many chemical and biochemical processes involve van der Waals interactions [93].

The intuitively clear idea of that atoms and molecules do interact with each other is a very old indeed and dates back to the ancient times. Recall in particular Democritus, Leukippos, and Lucretius whose philosophical thoughts on the interaction through a direct contact were developed and rationalized by R. Boscovich in “Theory of Natural Science Reduced to the Single Law of Forces Existing in Nature” (1758) and independently by A. C. Clairault in “Théorie de la Figure de la Terre” (1743).

In Nature, atoms are located at different interatomic distances depending on a kind of the forces between them: either by cohesion forces or chemical bonds. The latter prevail at the distances which are smaller or equal to the sum of van der Waals radii of atoms. At such distances atoms form a molecule. By definition, the van der Waals (vdW) radii of a given atom is the half of the shortest distance that is observed in crystals between the nuclei of the same atoms. The vdW radii of atoms are listed in Table 1. At the distances beyond the sum of van der Waals radii of atoms, there exists a specific van der Waals interaction often referred to as the dispersion interaction between atoms, after Johannes Diderik van der Waals who first postulated its existence in his well-known equation of state derived in his PhD thesis in 1837 and which won him the 1910 Nobel Prize in Physics.¹ For the first time van der Waals explained the deviations of gases from the ideal behavior. Let us consider a vessel filled by a gas of atoms. Within this vessel, the pressure exerted by a gas of atoms on its wall is lower compared to that predicted by the ideal gas law since the atoms may collide with the wall and are thus retained by the attraction they undergo from the other atoms in the bulk of the gas that results in the pressure P obeying the equation [94],

$$P = \left(\frac{nRT}{V - nb} \right) - a \left(\frac{n^2}{V^2} \right). \quad (26)$$

Here a is a so-called vdW factor and $b = (16\pi N/3)R_{\text{vdW}}^3$.

¹ <http://www.nobelprize.org/nobelprizes/physics/laureates/1910/waals-bio.html>.

Table 2 Dipole moments d^o of some molecules in D

| \mathcal{M} | d^o | \mathcal{M} | d^o | \mathcal{M} | d^o |
|--------------------|-------|-------------------|-------|--------------------|-------|
| <i>n</i> -Butane | 0.00 | Pyridine | 2.23 | <i>n</i> -Pentane | 0.00 |
| <i>n</i> -Hexane | 0.00 | Cyclohexanone | 2.90 | Acetone | 2.900 |
| Benzene | 0.00 | Propionitrile | 3.20 | Dimethylacetamide | 3.70 |
| Cyclohexane | 0.00 | Nitroethane | 3.2 | Lithium fluoride | 6.40 |
| Toluene | 0.36 | Dimethylsulfoxide | 3.92 | Lithium chloride | 7.09 |
| Triethylamine | 0.78 | Diethylether | 1.21 | Tetrahydrofurane | 1.76 |
| Hydrogen chloride | 1.10 | Methylacetate | 1.80 | Water | 1.84 |
| Propylenecarbonate | 4.98 | Sodium chloride | 9.06 | Potassium chloride | 10.70 |

It merits to notice that the dispersion interaction energy between the ground-state molecules is always negative. Therefore, such molecules always attract each other—in this regard it is appropriate to notice the celebrated work by E. H. Lieb and W. E. Thirring [95] that proves that the attractive van der Waals force between an atom and a molecule exists only at some mutual orientations and was generalized to any orientation in 2006 by M. Lewin [96]. The leading term of the dispersion energy is the dipole–dipole term which is proportional to $1/R^6$ and determined by a change in the zero-point vibrational energy of electric field created by zero-point vibrations of fluctuating dipole moments of interacting species. Since the zero-point fluctuations are the quantum phenomenon, the dispersion interactions have the quantum origin, as though in the beginning of the 1970s, T. H. Boyer [97] derived the London formula for the dispersion interactions within the classical electrodynamics, additionally assuming the existence of the classical electromagnetic zero-point radiation (Table 2). In fact, the van der Waals forces are cohesive attractions between molecules which operate at long intermolecular separations. From a quantitative viewpoint, van der Waals forces between molecules correspond to interactions between electric dipoles. Generally speaking, there exist three types of electric dipoles in molecules. These are permanent, induced, and temporary dipoles. If a molecule \mathcal{M} under study consists of the positive nuclear charges $q_1 = Z_1, \dots, q_M = Z_M$ and negative electron charges e_1, \dots, e_N , its total permanent dipole moment is defined as

$$\mathbf{d}^o = \sum_{\alpha=1}^M q_{\alpha} \mathbf{R}_{\alpha} + \sum_{i=1}^N e_i \mathbf{r}_i \quad (27)$$

d^o is distinct from zero in some state if the center of charge of the nuclei, $\mathbf{R}_n \equiv \sum_{\alpha=1}^M q_{\alpha} \mathbf{R}_{\alpha} / \sum_{\alpha=1}^M q_{\alpha}$, does not coincide with that of the electron subsystem, $\mathbf{r}_n \equiv \sum_{i=1}^N e_i \mathbf{r}_i / \sum_{i=1}^N e_i$. If $d^o \neq 0$, a molecule \mathcal{M} is called polar. Permanent dipole moments of neutral molecules usually vary from 0 to 15 Debye (D) that is reflected in Table 3.²

² Al. McClellan, Tables of Experimental Dipole Moments. Raha Enterprises, El Cerrito, 1974.

Table 3 Intramonomer distances (in Å) and binding energies (in kcal/mol), below, of representative van der Waals complexes, Ne_2 , Ar_2 , $(\text{CH}_4)_2(D_{3d})$, $(\text{C}_2\text{H}_4)_2(D_{2d})$, $\text{T}-(\text{C}_6\text{H}_6)_2$, $\text{SP}-(\text{C}_6\text{H}_6)_2$, $\text{P}-(\text{C}_6\text{H}_6)_2$, taken from Tables 1 and 2 of [98]

| DFT | Ne_2 | Ar_2 | $(\text{CH}_4)_2$ | $(\text{C}_2\text{H}_4)_2$ | $\text{T}-(\text{C}_6\text{H}_6)_2$ | $\text{SP}-(\text{C}_6\text{H}_6)_2$ | $\text{P}-(\text{C}_6\text{H}_6)_2$ |
|-----------|------------------|------------------|-------------------|----------------------------|-------------------------------------|--------------------------------------|-------------------------------------|
| VSXC | 2.47 | 3.59 | 3.21 | 3.35 | 4.65 | 3.47 | 3.41 |
| | 0.49 | 1.15 | 4.33 | 9.96 | 8.28 | 16.95 | 11.23 |
| PW91 | 3.00 | 4.0 | 4.0 | 4.0 | 5.15 | 5.76 | 4.22 |
| | 0.97 | 0.36 | 0.52 | 1.56 | 1.65 | 1.55 | 0.24 |
| HCTH407 | 2.58 | 3.84 | 3.89 | 4.05 | 5.45 | 5.84 | 4.22 |
| | 0.76 | 0.48 | 0.71 | 1.53 | 1.41 | 1.63 | 0.67 |
| PBE | 2.52 | 3.88 | 3.86 | 3.76 | 5.19 | 5.76 | 4.31 |
| | 0.69 | 0.19 | 0.29 | 1.30 | 1.33 | 1.27 | −0.22 |
| PBE1PBE | 2.54 | 3.93 | 3.90 | 3.75 | 5.14 | 5.76 | 4.40 |
| | 0.53 | 0.13 | 0.20 | 1.16 | 1.32 | 1.15 | −0.42 |
| BHANDHLYP | 2.48 | 3.93 | 3.99 | 3.76 | 5.20 | 5.77 | U |
| | 0.81 | 0.03 | 0.02 | 0.91 | 1.00 | 0.90 | |
| MPW1K | 2.62 | 4.31 | 4.35 | 3.82 | 5.19 | 5.81 | U |
| | 0.34 | 0.06 | 0.08 | 0.69 | 1.00 | 0.80 | |
| MPW1PW91 | 2.61 | 4.30 | 4.37 | 3.88 | 5.27 | 5.83 | U |
| | 0.40 | 0.07 | 0.10 | 0.62 | 0.89 | 0.75 | |
| B3LYP | 2.51 | U | U | 3.85 | 5.37 | 5.81 | U |
| | 0.63 | | | 0.55 | 0.63 | 0.60 | |
| B3P86 | 2.64 | U | U | 3.76 | 5.21 | 5.78 | U |
| | 0.04 | | | 0.47 | 0.64 | 0.48 | |
| CCSD(T) | 3.2 ^a | 3.8 ^a | 3.8 ^a | 3.8 ^a | 4.89 ^b | 3.8 ^b | 3.7 ^b |
| | 0.07 | 0.26 | 0.495 | 1.15 | 2.74 | 2.78 | 1.81 |

U implies that at this computational level, the studied complex is unbound. The notations ^a[99] where the computational level CCSD(T) and the aug-cc-pV5Z basis set are used, and ^b[100], and U for unbounded. The experimental value of the bound distance in Ne_2 dimer is equal to 3.09 Å. The benzene dimer exists in three isomeric structures *T*—*T*-shaped, *P*—parallel, and *SP*—slipped parallel

Any pair of polar molecules, \mathcal{M}_1 and \mathcal{M}_2 , separated by a distance R_{12} , in the states n and m , respectively, interact with each other by their dipoles, $\mathbf{d}_n^{(1)}$ and $\mathbf{d}_m^{(2)}$, via the dipole–dipole interaction read as

$$E_{\text{dd}}^{(1)} = \frac{\mathbf{d}_n^{(1)} \mathbf{d}_m^{(2)}}{R_{12}^3} - \frac{3(\mathbf{d}_n^{(1)} \mathbf{R}_{12})(\mathbf{d}_m^{(2)} \mathbf{R}_{12})}{R_{12}^5}. \quad (28)$$

Structurally, $E_{\text{dd}}^{(1)}$ consists of two terms. A polar molecule \mathcal{M}_1 interacts with the electric field $\mathbf{E}^{(2)}(\mathbf{R}_{12})$ created by another molecule \mathcal{M}_2 at the position of the first molecule. As known from electrostatics [101–103], molecule \mathcal{M}_1 gains the energy $-\mathbf{d}_n^{(1)} \mathbf{E}^{(2)}(\mathbf{R}_{12})$. Expressing the electric field

$$\mathbf{E}^{(2)}(\mathbf{R}_{12}) = \nabla[(\mathbf{d}_m^{(2)} \mathbf{R}_{12})/R_{12}^3] \quad (29)$$

results then in Eq. (28).

Let us consider another, so-called, second-order effect of an external electric field \mathbf{E} on a given molecule \mathcal{M}_2 . This field influences the molecular charges, electrons, and nuclei, causing their displacements, and as a result, there appears the induced dipole moment \mathbf{d}^{ind} ,

$$\mathbf{d}^{\text{ind}} = \alpha \mathbf{E}, \quad (30)$$

where the proportionality coefficient is merely a polarizability α of a given molecule. Assuming that this electric field is generated by the presence of the second molecule \mathcal{M}_1 , one obtains

$$\mathbf{d}_{2,m}^{\text{ind}} = \alpha \nabla[(\mathbf{d}_m^{(1)} \mathbf{R}_{12})/R_{12}^3]. \quad (31)$$

Therefore, the interaction of the permanent dipole $\mathbf{d}_n^{(1)}$ of \mathcal{M}_1 with the dipole $\mathbf{d}_{2,n}^{\text{ind}}$ that is induced on \mathcal{M}_2 by \mathcal{M}_1 takes the following expression:

$$E_{\text{dd}}^{(2)} = \frac{\mathbf{d}_n^{(1)} \mathbf{d}_{2,n}^{\text{ind}}}{R_{12}^3} - \frac{3(\mathbf{d}_n^{(1)} \mathbf{R}_{12})(\mathbf{d}_{2,n}^{\text{ind}} \mathbf{R}_{12})}{R_{12}^5} \quad (32)$$

that is known as the Keesom dipole–dipole interaction [104].

By a straightforward analogy, the dipole–dipole interaction of two mutually induced dipoles on \mathcal{M}_1 and \mathcal{M}_2 is described by the expression

$$E_{\text{dd}}^{(2)} = \frac{\mathbf{d}_{1,n}^{\text{ind}} \mathbf{d}_{2,m}^{\text{ind}}}{R_{12}^3} - \frac{3(\mathbf{d}_{1,n}^{\text{ind}} \mathbf{R}_{12})(\mathbf{d}_{2,m}^{\text{ind}} \mathbf{R}_{12})}{R_{12}^5}. \quad (33)$$

If the distance R_{12} between dipoles is small enough compared to the wavelength λ , corresponding to transitions between the ground and excited states, within the second- and higher-order Rayleigh–Schrödinger perturbation theory as [92, 105, 106] there appear, as first shown by Fritz London [105, 106], the dispersion interaction [105, 106]

$$E_{\text{disp}}^{(2)} = - \sum_{m,n \neq 0} \frac{\left| \langle \Psi_n^{(1)} \Psi_m^{(2)} | V_{12} | \Psi_0^{(1)} \Psi_0^{(2)} \rangle \right|^2}{(E_n^{(1)} - E_0^{(1)}) + (E_m^{(2)} - E_0^{(2)})} \quad (34)$$

where $\Psi_n^{(i)}$ is the n -state eigenfunction of \mathcal{M}_i , $i = 1, 2$ and V_{12} is the electrostatic interaction between molecules \mathcal{M}_1 and \mathcal{M}_2 .

The dispersion energy is traditionally represented by means of the multipole expansion [94]

$$E_{\text{disp}}^{(2)} = - \sum_{n=6}^{\infty} \frac{C_n}{R^n}, \quad (35)$$

where C_n are dispersion constants among which C_6 corresponds to dipole–dipole interaction, C_8 to dipole–quadrupole and C_{10} to dipole–octupole and dipole–quadrupole interactions.

Dispersion interactions play a role of the attractive interaction between rare gas atoms and are also one of the important intermolecular interactions that govern the molecular organic world [93, 107]. Dispersion interactions are mostly responsible for the heats of sublimation of hydrocarbon molecules, make significant contributions to the solvent properties of polar and apolar neutral compounds [108, 109] and are also important for crystal packing of organic molecules [110] as well as for the stacking of nucleic acids in DNA [93, 111]. The world of dispersion interaction is rich (see [112] and references therein), despite the fact that it is a weaker form of intermolecular attractions. Dispersion forces as one of the two types of van der Waals force are also known as “London forces,” named after Fritz London [105, 106].

Density functional theory [113–116] as one of the approaches to evaluate electron correlation is considerably less demanding on computational resources than the MP2 or CCSD(T) methods. DFT might therefore be considered as a powerful computational tool, if it can adequately describe and accurately evaluate intermolecular interactions. The suitability of DFT for the evaluation of dispersion interaction has been an important issue in the recent literature [117–125] that is mirrored in Table 3. While DFT calculations with local exchange–correlation functionals lead to overestimate binding energies of weakly bound systems, it was reported that non-local exchange–correlation functionals very often underestimate the attraction [118–120]. The DFT calculations with Becke’s exchange and Lee, Yang, and Parr’s correlation functionals, BLYP [113, 114], and Becke’s three-parameter functional combined with Lee, Yang, and Parr’s correlation functional, B3LYP [114, 116], also fail to evaluate the attractive dispersion interaction between hydrocarbon molecules [120–122]. Recently Zhang et al. and Wesolowski et al. reported that the Becke exchange functional due to its erroneous asymptotic behavior at low density is responsible for the failure in the evaluation of the attraction between weakly bound systems [123–125]. It was shown, however, that other non-local exchange–correlation functionals such as Perdew and Wang’s exchange and correlation functionals, PW91 [115], are possible alternatives to describe the binding between rare gas dimers or other systems. The performance of some exchange–correlation functionals and the PW91 one in particular for the representative van der Waals systems is demonstrated in Table 3. Notice that the PW91 functional is a general functional, i.e., it is not biased towards the description of intermolecular interaction. In this investigation we will examine the basis set dependence of the interaction energies, and benchmark those against the results obtained from MP2 [188] and CCSD(T) theory. For comparison, the results with the BLYP and B3LYP are also presented. Interestingly, DFT adequately describes, on the one hand, atoms and molecules as stable many-electron systems and on the other hand, the molecules formed under interaction of its composing molecules. However, its description of those molecular interactions is not always perfect. The simplest DFT approximation widely used in computational practice is the local

density approximation (LDA) [2], based on the properties of the uniform electron gas. In principle, DFT yields the exact ground-state energy, including long-range van der Waals energies, very important in organic chemistry and elsewhere. However, the commonly used LDA and GGA, designed for nonuniform electron gases, fail to capture the essence of vdW energies. The latter reflect correlated motions of electrons due to the Coulomb interactions between distant, even non-overlapping atoms, molecules, and solids. In [4] Kohn and coauthors propose a first-principles approach, which contains the following ingredients (i) The density distribution, $\rho(\mathbf{r})$, is approximated by the LDA or GGA. (ii) The Coulomb interaction is divided into short- and long-range parts, of which only the latter contributes to vdW energies. (iii) The contribution of the long-range interactions to the energy is expressed by the so-called adiabatic connection formula. (iv) This expression is transformed into the time domain, avoiding the need to solve a self-consistent equation for the density–density response function.

4 Dispersion-Corrected DFT Approaches

DFT methods with currently available functionals failed completely for London-type clusters for which no minimum was found at the potential energy surfaces.

P. Hobza, J. Reschel and J. Sponer (1995) J Comput Chem 16:1315

Density functional theory is often the preferred electronic structure method to study moderate and large systems. This preference reflects the efficiency of DFT compared to correlated wavefunction theories such as coupled cluster theory even though accuracy, and more importantly, predictability (as in systematic convergence to the right answer) are sacrificed. DFT which incorporates currently accepted exchange-correlation functionals can be used with reasonable reliability on chemically bound systems around the equilibrium geometry but inevitably fail when applied to systems which are bound by weak van der Waals forces [126–130] and to a lesser degree for chemically bound systems away from equilibrium, like transition states [131]. These failings of density functional theory are well known [132]. Here, we address weak interactions. Attempts to compute weak intermolecular forces using DFT fall into two categories. Some would simply modify functionals until reasonable results are obtained (see [133] and references within). Others would focus on an add-on correction that explicitly introduces the van der Waals C_6 coefficient. This can be made to work, but it is unsatisfactory that the weak interactions do not occur naturally as they would in wavefunction methods. This is the experimental or computational fact which has not been still proved. Though, the problem of description of London dispersion in density functional theory (DFT) using (semi) local exchange-correlation functionals is a well-known problem [134, 135] since the first diagnostic in 1994 [134].

A step in the right direction was made by Engel et al. [136] who obtained reasonable results for the helium and neon dimers. In [137], Bartlett and

co-workers proposed *ab initio* density functional theory has been applied for the weakly interacting, He_2 , $[\text{He}-\text{Be}]^{2+}$, Ne_2 and Be_2 that results in fair agreement with the highly accurate coupled-cluster method. Generally, one assumes that the cause lies in the local character of the widely used correlation functionals, which, in contrast to the correlation contribution in post-Hartree–Fock methods such as Møller–Plesset or coupled cluster, only utilize information on the density of the system at one point and are therefore unsuitable for the description of a non-local phenomenon such as dispersion. Attempts to introduce non-local correlation to DFT, such as the random phase approximation (RPA) [138, 139] or the non-local van der Waals functionals [140–143], are being investigated, but unfortunately the improvement comes with a significant increase in the computational cost. Since the relatively low computational cost of DFT is one of the major factors responsible for its status as the most widely used quantum chemical method today, a range of more pragmatic approaches has been developed to correct the performance of DFT for dispersion interactions. Part of these methods rely on reparametrization of existing local correlation functionals [144–146], motivated by the fact that dispersion is partially included in many functionals and that a suitable reparametrization will allow one to achieve the aspired results more consistently. The drawback of such an approach is that the strong empirical character decreases the reliability. For instance, the performance of the reparametrized functionals often decreases for properties other than the electronic energy. Other attempts are based on adding a correction term, representing the dispersion energy, to the energy calculated using standard DFT methods. Also in this category, one can find highly empirical but computationally attractive methods [147] based on parameters fitted to reproduce high-level results, as well as the methods with deeper theoretical foundation but computationally more expensive, where *ab initio* information of the systems is used to evaluate the dispersion energy, such as the static or frequency dependent polarizabilities [148–154] or the dipole moment of the exchange-correlation hole (XDM) [156–159]. Another noteworthy approach is the adaptation of the symmetry adapted perturbation theory [160] to the framework of DFT, i.e., SAPT(DFT) [161–164]. SAPT(DFT) has a significant computational advantage against the highly scaling SAPT as the contribution of intramonomer correlation, already embedded within the Kohn–Sham orbitals, does not need to be evaluated. Although possible to use for the correction of DFT dispersion energies [165], SAPT(DFT) is mostly meant for an evaluation of the total interaction energy. The explicit expression for the repulsive contribution of electron-exchange to the dispersion energy within SAPT(DFT), though rarely calculated fully due to the computational expense, offers a more theoretically attractive alternative to the empirical damping functions used in other methods. SAPT(DFT) does have the disadvantage of requiring explicit separation of the system in two parts, which makes it impossible for application on intramolecular dispersion interactions, such as those occurring, for example, in biomolecules. Table 4 reports some developed dispersion-corrected DFT functionals and their performance for the representatives of the van der Waals dimers.

Table 4 Dispersion-corrected DFT functionals

| Method | He ₂ | He-Ne | Ne ₂ | He-Ar | Ar ₂ | (CH ₄) ₂ | (C ₂ H ₄) ₂ | T-(C ₆ H ₆) ₂ |
|---------------------------|-----------------|-------|-----------------|-------|-----------------|---------------------------------|---|---|
| CCSD(T) | -0.02 | -0.04 | -0.07 | -0.06 | -0.27 | -0.53 | -1.51 | -2.74 |
| D-B3LYP/A ^a | 0.04 | 0.04 | 0.05 | 0.07 | 0.17 | 0.38 | 0.49 | 0.98 |
| D-BH-B3LYP/A ^a | -0.00 | -0.02 | -0.05 | -0.02 | -0.19 | -0.49 | -1.68 | -2.64 |
| D-B3LYP/B ^a | 0.05 | 0.04 | 0.03 | 0.08 | 0.07 | 0.76 | 0.40 | 0.96 |
| D-BH-B3LYP/B ^a | 0.03 | 0.02 | -0.03 | 0.03 | -0.08 | -1.12 | -0.87 | -1.89 |
| DFT-B97-D/C ^b | | | -0.17 | | -0.26 | | | -2.99 |

Basis sets A \equiv aug-cc-pVTZ, B \equiv 6-311++G(2d,f,p), and C \equiv TZV2P

^a[166].

^b[167].

One of the most promising new DFT methods is the non-local van der Waals correlation functional vdW-DF-04 in [168, 169], which was derived from first principles, describes dispersion interactions in a seamless fashion, and yields the correct asymptotic form of intermolecular van der Waals forces. Recently we reported a self-consistent implementation of vdWDF-04 with Gaussian basis functions [168, 169]. The code includes analytic gradients of the energy with respect to nuclear displacements, enabling efficient geometry optimizations. The alternative and consistent approach to the dispersion-corrected DFT can be formulated, as we believe, within the so-called local-scaling transformations' (LST') DFT whose key features are given in the next section.

5 Local-Scaling Transformations' DFT

In order to properly assess the local-scaling transformation formulation of the density functional theory, we first consider the concept of local-scaling transformation and second, apply it to the topological features of atomic and molecular one-electron densities.

5.1 Mathematical Preliminaries: Local-Scaling Transformations

Define on the Euclidean \mathbb{R}^3 the following mapping: $\mathbb{R}^3 \xrightarrow{f} \mathbb{R}^3$ such that $r \in \mathbb{R}^3$ is mapped into

$$\mathbf{f}(\mathbf{r}) = f(\mathbf{r})\mathbf{e}_r = f(r; \mathbf{e}_r)\mathbf{e}_r, \quad (36)$$

where $\mathbf{e}_r \equiv \mathbf{r}/r \equiv \mathbf{e}(\Omega)$ is a unit vector, specified in \mathbb{R}^3 and defined by the spherical angles $\Omega = (\theta, \phi)$ and $r = |\mathbf{r}|$. For a given $\mathbf{e}_r(\Omega)$, the transformation (36) that deforms \mathbb{R}^3 onto itself, nonuniformly in general, is referred to as a local-scaling transformation or LST for short [4] and is the special class of point transformation

[170, 171]. LSTs satisfy all axioms of group and hence form the group \mathcal{F} of local-scaling transformations. A scalar function $f(\mathbf{r})$ in Eq. (36) can be arbitrary, though often it belongs to \mathcal{C}^1 or higher. In the former, \mathbf{f} is a \mathcal{C}^1 -diffeomorphism on \mathbb{R}^3 .

Equation (36) nontrivially generalizes the well-known scaling: $\mathbf{f}_\lambda(\mathbf{r}) := \lambda \mathbf{r}$ which Fock [172] used in 1930 to prove the virial theorem. $\lambda \neq 0$ is a constant that means that all vectors $\mathbf{r} \in \mathbb{R}^3$ are scaled uniformly. Bearing in mind that an arbitrary vector \mathbf{r} is uniquely determined by its Cartesian coordinates $\mathbf{r} = (x, y, z)$, the equivalent representation of (36) is the following:

$$\mathbf{r} \equiv \begin{pmatrix} x \\ y \\ z \end{pmatrix} \xrightarrow{f} \mathbf{f}(\mathbf{r}) \equiv \begin{pmatrix} \frac{x}{r}f(x, y, z) \\ \frac{y}{r}f(x, y, z) \\ \frac{z}{r}f(x, y, z) \end{pmatrix} \equiv \begin{pmatrix} f_x(\mathbf{r}) \\ f_y(\mathbf{r}) \\ f_z(\mathbf{r}) \end{pmatrix} \equiv \begin{pmatrix} x\sigma(\mathbf{r}) \\ y\sigma(\mathbf{r}) \\ z\sigma(\mathbf{r}) \end{pmatrix}, \quad (37)$$

where $\mathbf{f}(\mathbf{r}) = \sigma(\mathbf{r})\mathbf{r}$.

The Jacobian of (1) is defined as

$$J\{\mathbf{f}(\mathbf{r}); \mathbf{r}\} \equiv J\{\mathbf{f}; \mathbf{r}\} = \begin{vmatrix} \frac{1}{r}f - \frac{x^2}{r^3}f + \frac{x}{r}\frac{\partial f}{\partial x} & -\frac{xy}{r^3}f + \frac{x}{r}\frac{\partial f}{\partial y} & -\frac{xz}{r^3}f + \frac{x}{r}\frac{\partial f}{\partial z} \\ -\frac{xy}{r^3}f + \frac{y}{r}\frac{\partial f}{\partial x} & \frac{1}{r}f - \frac{y^2}{r^3}f + \frac{y}{r}\frac{\partial f}{\partial y} & -\frac{yz}{r^3}f + \frac{y}{r}\frac{\partial f}{\partial z} \\ -\frac{xz}{r^3}f + \frac{z}{r}\frac{\partial f}{\partial x} & -\frac{yz}{r^3}f + \frac{z}{r}\frac{\partial f}{\partial y} & \frac{1}{r}f - \frac{z^2}{r^3}f + \frac{z}{r}\frac{\partial f}{\partial z} \end{vmatrix} \\ = \frac{f^2}{r^3} \left(x \frac{\partial f}{\partial x} + y \frac{\partial f}{\partial y} + z \frac{\partial f}{\partial z} \right) = \frac{1}{3r^3} \mathbf{r} \cdot \nabla f^3. \quad (38)$$

In terms of $\sigma(\mathbf{r})$, the Jacobian (38) has the form $J\{\mathbf{f}(\mathbf{r}); \mathbf{r}\} = \sigma(\mathbf{r})[1 + \mathbf{r} \cdot \nabla \ln \sigma(\mathbf{r})]$ [171]. For the uniform scaling $\mathbf{f}_\lambda :=$, the corresponding Jacobian is equal to

$$J\{\mathbf{f}_\lambda; \mathbf{r}\} = \begin{vmatrix} \lambda & 0 & 0 \\ 0 & \lambda & 0 \\ 0 & 0 & \lambda \end{vmatrix} = \lambda^3. \quad (39)$$

It is trivial to generalize a three-dimensional local-scaling transformation (1) on other dimensions, say \mathbb{R}^D , simply by considering a vector \mathbf{r} as a D -dimensional one. If $D = 1$, $f(r)$ is a function of a single variable r . The corresponding Jacobian $J\{f(r); r\} = df(r)/dr$. Let us consider some examples of local-scaling transformations \mathbf{f}_{LST} :

- (1) $f^{[1]} = \left[\left(\frac{1}{r} \right)^m + \left(\frac{\delta}{\sqrt{r}} \right)^m \right]^{-1/m}$, where δ is the constant [173].
- (2) $f^{[2]} = \begin{cases} r(1 + ar^2)^{1/3} & \text{if } r \leq R \\ \sqrt{\frac{d_{-2}}{r^2} + \frac{d_{-1}}{r}} + d_o + d_1 r + d_L \ln r & \text{otherwise.} \end{cases}$

This form results from the asymptotes at small and large r [174].

- (3) $f^{[3]} = \begin{cases} r & \text{if } r \leq R \\ a\sqrt{\frac{8r}{a} - \frac{8a}{r} + \frac{a^2}{r^2} - 12 \ln\left(\frac{r}{a}\right)} & \text{otherwise} \end{cases}$ [8].

- (4) Let $\Omega :=]-L/2, L/2[^3 \subseteq \mathbb{R}^3$ be a cube with volume $|\Omega| = L^3$. $\mathbf{f}^{[4]}$ is defined as a periodic deformation on the cube Ω if it is a C^1 -diffeomorphism on \mathbb{R}^3 that leaves Ω invariant: $\mathbf{f}^{[4]}(\Omega) = \Omega$ and if $\mathbf{f}^{[4]}(\mathbf{r} + L\mathbf{m}) = \mathbf{f}^{[4]}(\mathbf{r}) + L\mathbf{m}$ for any $\mathbf{m} \in \mathbb{Z}^3$ [175].
- (5) $f_{p,q,r}^{[5]}$ is defined by the inverse function $r\left(f_{p,q,r}^{[5]}\right) = \left[f_{p,q,r}^{[5]}\right]^p \left(1 + \alpha \left[f_{p,q,r}^{[5]}\right]^q\right)^r$ where α, p, q , and r are variational parameters. If $p = q = r = 1$, $r\left(f_{p,q,r}^{[5]}\right)$ refers to Hall's transformation [176]. The other $r\left(f_{p,q,r}^{[5]}\right)$ with $q = r = 1, p = r = 1$ and $p = q = 1$ were defined in [177, 178]. Hall's local-scaling transformation is then $f_{1,1,1}^{[5]} = [(1 + 4\alpha r)^{1/2} - 1]/(2\alpha)$.

Let $\phi(\mathbf{r})$ be an arbitrary function given on domain $\Sigma \subset \mathbb{R}^3$. A local-scaling transformation (16) transforms $\phi(\mathbf{r})$, generally speaking, into another function

$$\psi(\mathbf{r}) := \phi(\mathbf{f}(\mathbf{r})) \quad (40)$$

within the Jacobian (3), depending on the normalization of $\phi(\mathbf{r})$ on Σ if any. If $\phi(\mathbf{r}) = \exp(-r)$ is a simple exponential orbital, under Hall's local-scaling transformation it converts to

$$\psi(\mathbf{r}) = \frac{(1 + 4\alpha r)^{1/2} - 1}{2\alpha r(1 + 4\alpha r)^{1/4}} \exp\left(-[(1 + 4\alpha r)^{1/2} - 1]/(2\alpha)\right) \quad (41)$$

that was used to approximate the 1s orbital.

5.2 One-Electron Densities: Definition

A function $\rho(\mathbf{r}) : \mathbb{R}^3 \rightarrow \mathbb{R}_+^1$ is defined as a one-electron density associated with some system of N electrons iff:

- (Di) $\rho(\mathbf{r})$ is non-negative everywhere in \mathbb{R}^3 ;
- (Dii) $\rho(\mathbf{r})$ is normalized to the total number N of electrons,

$$\int_{\mathbb{R}^3} d^3\mathbf{r} \rho(\mathbf{r}) = N. \quad (42)$$

Here \mathbb{R}_+^1 stands for the non-negative semi-axis of \mathbb{R}^1 . Equation (42) merely implies that the square root of $\rho(\mathbf{r})$ is a square-integrable function, i.e., $[\rho(\mathbf{r})]^{1/2} \in L^2(\mathbb{R}^3)$;

- (Diii) $\rho(\mathbf{r})$ is a continuously differentiable function of \mathbf{r} almost everywhere in \mathbb{R}^3 .
It is a well-behavedness of densities.

Let \mathcal{D}_N be the class of the one-electron densities associated with a Coulomb system of N electrons and obeying the conditions **(Di)**–**(Diii)**. Obviously, $V\mathcal{L}_N \subset \mathcal{D}_N$. The fact that the condition **(Diii)** is valid for $V\mathcal{L}_N$ is the consequence of the following

Proposal 1. For any $\rho_\Psi(\mathbf{r}) = V\Psi$ where $\Psi \in \mathcal{L}_N$, $\nabla_{\mathbf{r}}\rho_\Psi(\mathbf{r}) \in \mathcal{L}^2(\mathbb{R}^3)$.

Proof. [88]: According to the Schwarz inequality, it follows from Eq. (11) that

$$[\nabla_{\mathbf{r}}\rho_\Psi(\mathbf{r})]^2 \leq 4N\rho_\Psi(\mathbf{r}) \sum_{s_1, \dots, s_N} \int d^3\mathbf{r}_2 \cdots \int d^3\mathbf{r}_N |\Psi(\mathbf{r}, s_1; \mathbf{r}_2, s_2; \dots; \mathbf{r}_N, s_N)|^2. \quad (43)$$

□.

Corollary 1.1. $\nabla_{\mathbf{r}}[\rho_\Psi(\mathbf{r})]^{1/2} \in \mathcal{L}^2(\mathbb{R}^3)$.

Proof. [10]:

$$\int d^3\mathbf{r} (\nabla_{\mathbf{r}}[\rho_\Psi(\mathbf{r})]^{1/2})^2 = \frac{1}{4} \int d^3\mathbf{r} \frac{[\nabla_{\mathbf{r}}\rho_\Psi(\mathbf{r})]^2}{\rho_\Psi(\mathbf{r})} \leq T_e[\Psi]. \quad (44)$$

□.

The term $[\nabla_{\mathbf{r}}\rho_\Psi(\mathbf{r})]^2/\rho_\Psi(\mathbf{r})$ is known as the von Weizsäcker kinetic energy $t_W[\rho_\Psi(\mathbf{r})]$ [4]. Hence, in the other words, **Corollary 1.1** tells that $t_W^{1/2}[\rho_\Psi(\mathbf{r})]$ is square-integrable. Usually, the von Weizsäcker term is only a part of the total many-electron kinetic energy [4]. The exception is the Hartree–Fock 2-electron model systems for which $t_W[\rho_\Psi(\mathbf{r})]$ is the exact kinetic energy.³ We further have

Corollary 1.2. Thomas–Fermi [172, 179] one-electron density $\rho_{\text{TF}}(\mathbf{r})$ is not N -representable.

Proof. According to [180, 181], $t_W^{1/2}[\rho_{\text{TF}}(\mathbf{r})]$ is not square-integrable. □.

Furthermore, the Thomas–Fermi energy density functional cannot be inserted in the density functional philosophy presented by the mappings (13) and (14) for all $\rho(\mathbf{r}) \in \mathcal{D}_N$ since the ground-state energies of many Thomas–Fermi atoms and ions⁴ lie below the exact ones.⁵

³ Except H^- which is unstable within the Hartree–Fock picture since its Hartree–Fock ground-state energy is equal to -0.488 hartree and placed above $E_0[H] = -0.5$ hartree. Note that the exact ground-state energy of H^- is -0.5278 hartree.

⁴ Thomas–Fermi molecules are unstable (see [189] and e.g., [4] and references therein).

⁵ Some of the widely used density functionals predict the ground-state energies below the experimental ones. For example, the B3LYP density functional in conjunction with the 6-31+G(d, p) basis set yields the energy -0.500273 hartree $< E_0[H]$ [182]. The B3LYP and B3PW91 show a similar trend for the atoms Li, C, O, F, Na, and Mg, and diatomics O_2 , F_2 , and LiF [183]. This implies that the corresponding ground-state wavefunctions, if do exist, are not square-integrable.

Consider an N -electron atom or ion with the nucleus centered at the origin of the Cartesian coordinate system. Let $\rho(\mathbf{r}) \in \mathcal{D}_N$ be one-electron density associated with a given atom and $\rho(\mathbf{r}) = \{\rho(r, \mathbf{e}_r) | r \in \mathbb{R}_+, \Omega \equiv (\theta, \phi), 0 \leq \theta \leq \pi, 0 \leq \phi \leq \pi\}$ is merely a bundle of one-dimensional curves. $\tilde{\rho}_2(r)$

Let two densities $\rho_1(\mathbf{r})$ and $\rho_2(\mathbf{r})$ from \mathcal{D}_N be given. Both of them are represented by the corresponding bundles of curves. Let us choose the unit vector \mathbf{e}_r and in these bundles, the projections of $\rho_1(\mathbf{r})$ and $\rho_2(\mathbf{r})$ onto \mathbf{e}_r ; the curves $\tilde{\rho}_1(r)$ and $\tilde{\rho}_2(r)$ which are, according to (Diii), continuously differentiable functions of $r = |\mathbf{r}|$. Hence, they are homotopically equivalent, or equivalently, there does exist such topological deformation that maps or deforms $\tilde{\rho}_1(r)$ into $\tilde{\rho}_2(r)$. Formally,

$$\tilde{\rho}_2(r) = J\{\mathbf{f}(\mathbf{r}); \mathbf{r}\} \tilde{\rho}_1(f(r, \mathbf{e}_r)). \quad (45)$$

The Jacobian in (38) ensures the normalization (Dii) for both densities ρ_1 and ρ_2 . Generalizing Eq. (24) over all directions in \mathbb{R}^3 results in that [7]

$$\rho_2(\mathbf{r}) = \frac{1}{3r^3} \mathbf{r} \cdot \nabla f^3 \rho_1(\mathbf{f}(\mathbf{r})). \quad (46)$$

To hold the electron–nuclear Kato cusp, the nuclear position is invariant of f . If f is a uniform scaling f_λ , the latter equation takes the form

$$\rho_\lambda(\mathbf{r}) = \lambda^3 \rho_1(\lambda \mathbf{r}). \quad (47)$$

Given \mathbf{e}_r , combining Eqs. (36) and (38) yields

$$\frac{df(r, \mathbf{e}_r)}{dr} = \frac{r^2 \rho_2(r, \mathbf{e}_r)}{f^2 \rho_1(f(r, \mathbf{e}_r), \mathbf{e}_r)} \quad (48)$$

or in spherical coordinates, along a chosen unit vector \mathbf{e}_r determined by $\Omega = (\theta_0, \phi_0)$,

$$\frac{df(r, \theta_0, \phi_0)}{dr} = \frac{r^2 \rho_2(r, \theta_0, \phi_0)}{f^2 \rho_1(f(r, \theta_0, \phi_0), \theta_0, \phi_0)}. \quad (49)$$

Equation (49), or (48), is the first-order nonlinear differential equation for deformation $f(\mathbf{r})$ for given densities ρ_1 and ρ_2 . Due to (Diii), its solution exists and it is unique (see e.g., [4] and references therein). Therefore, for any pair of well-behaved densities, one enables to determine the deformation that transforms one of them into another. This means that F acts on \mathcal{D}_N transitively, that is, in algebraic terminology, \mathcal{D}_N is a single orbit with respect to \mathcal{F} . For a given and fixed density $\rho_1(\mathbf{r})$, defined hereafter as the generator density $\rho_g(\mathbf{r})$, Eq. (46) then implies:

Proposal 2. There exists the one-to-one correspondence between \mathcal{F} and \mathcal{D}_N that is explicitly expressed as $f \in \mathcal{F} \Leftrightarrow \rho_f^{[g]}(\mathbf{r}) := J\{\mathbf{f}, \mathbf{r}\} \rho_g(\mathbf{f}(\mathbf{r}))$.

In the integral form, Eq. (28) is as follows:

$$f(r, \theta_0, \phi_0) = \left[3 \int_{r_0}^r dr \frac{r^2 \rho_2(r, \theta_0, \phi_0)}{\rho_1(f(r, \theta_0, \phi_0), \theta_0, \phi_0)} \right]^{1/3}. \quad (50)$$

Note that the rhs of (50) contains a cubic root that reflects that the group \mathcal{F} of local-scaling transformations acts on \mathbb{R}^3 . It is shown in [184] that the dimensionality D of \mathbb{R}^D enters the corresponding Jacobian in the power D and, respectively, the corresponding integral form as $1/D$, this is on the one hand. On the other, there exists another remarkable facet of Eq. (28). This equation is well known in mathematics as the “Jacobian problem” ([55, 57–63, 190], see also [56]).

5.3 Many-Electron Wavefunctions and Concept of Orbit

To build the “variable mapping” (14), let us generalize the concept of the local-scaling transformations on \mathcal{L}_N . This is rather simple and straightforward. For this purpose, let us choose an arbitrary “reference” or generator wavefunction $\Phi_g(\{\mathbf{r}_i, \sigma_i\}_{i=1}^{i=N})$ where σ_i is the spin of the i -th electron and $\rho_g(\mathbf{r}) \in \mathcal{D}_N$ is the associated one-electron density. Then define a new wavefunction

$$\Phi_f(\{\mathbf{r}_i, \sigma_i\}) = \Phi_\rho(\{\mathbf{r}_i, \sigma_i\}) \equiv \left[\prod_{i=1}^N J(\mathbf{f}(\mathbf{r}_i); \mathbf{r}_i) \right]^{1/2} \Phi_g(\{\mathbf{f}(\mathbf{r}_i), \sigma_i\}) \quad (51)$$

with the density $\rho(\mathbf{r}) \equiv \rho_g(\mathbf{f}(\mathbf{r}))$ casting in **Proposal 2**. Φ_f is a locally scaled image of the “reference” wavefunction. Formally, $\Phi_f \equiv F\Phi_g$ where $F \in \mathcal{F}^{\times N} := [\times]^N \mathcal{F}$ and $F = (f, f, \dots, f) := f^{\times N}$ and (51) is nothing then else as the definition of the action of the group $\mathcal{F}^{\times N}$ on \mathcal{L}_N . Arbitrariness in choosing Φ_g ensures the validity of the definition (30) on the entire \mathcal{L}_N . Due to the isomorphism of the groups \mathcal{F} and $\mathcal{F}^{\times N}$, it is obvious that a local-scaling transformation that maps a given pair of N -electron wavefunctions into each other matches unambiguously the local scaling that transforms the corresponding one-electron densities into each other. However, although any pair of densities are locally scaled, this property no longer holds for an arbitrary pair of N -electron wavefunctions. Hence, \mathcal{L}_N is nontrivially partitioned, with respect to the group $\mathcal{F}^{\times N}$ of local-scaling transformations, into the orbits

$$\mathcal{L}_N = \bigcup_i \mathcal{O}^{[i]}. \quad (52)$$

In this sense, the group \mathcal{F} entangles \mathcal{D}_N and \mathcal{L}_N . By construction, an arbitrary orbit $\mathcal{O}^{[i]}$ is closed with respect to $\mathcal{F}^{\times N}$, that is, for any pair Φ_1 and Φ_2 in $\mathcal{O}^{[i]}$, there exists such local-scaling transformation $F_{1 \Rightarrow 2}$ that $\Phi_2 = F_{1 \Rightarrow 2} \Phi_1$. In the other words, if Φ_1 is the generator wavefunction of $\mathcal{O}^{[i]}$, for all $F \in \mathcal{F}^{\times N}$, $F_{1 \Rightarrow 2} \Phi_1 \in \mathcal{O}^{[i]}$. We thus prove:

Proposal 3. There exists a one-to-one map of variables on any orbit in \mathcal{L}_N .

Corollary 3.1. Orbit $\mathcal{O}^{[i]}$ is invariant relative to generator wavefunction.

Corollary 3.2. On each orbit $\mathcal{O}^{[i]} \subset \mathcal{L}_N$, there exists one and only one N -electron wavefunction which one-electron density is $\rho(\mathbf{r}) \in \mathcal{D}_N$.

Corollary 3.3. For any given orbit $\mathcal{O}^{[k]} \subset \mathcal{L}_N$ generated by $\Phi_g^{[k]}$ and the latter one-electron density $\rho_g^{[k]}, \mathcal{F}_{\rho_g^{[k]}}^{[k]}$ exhausts the whole \mathcal{D}_N .

Remark 1: Corollary 3.3 implies that any density $\rho(\mathbf{r}) \in \mathcal{D}_N$ is N -representable. In other words, the group \mathcal{F} of local-scaling transformations and its actions on \mathcal{D}_N and \mathcal{L}_N defined above ensures the N -representability of \mathcal{D}_N .

The uniqueness of the local-scaling transformation as the solution of Eq. (48) guarantees that the transformed wavefunction $\Phi_\rho^{[i]}$ is also unique. Thus, for any $\rho(\mathbf{r}) \in \mathcal{D}_N$ there exists a unique wavefunction $\Phi_\rho^{[i]}$ generated by means of local-scaling transformation from the arbitrary generator wavefunction $\Phi_g^{[k]}$. The orbit in \mathcal{L}_N is actually the set of all the wavefunctions thus generated which yield one-electron densities $\rho(\mathbf{r})$ in \mathcal{D}_N :

$$\mathcal{O}^{[i]} \equiv \left\{ \Phi_\rho^{[i]} | \Phi_\rho^{[i]} \rightarrow \rho(\vec{r}); \Phi_\rho^{[i]} \in \mathcal{L}_N; \rho(\vec{r}) \in \mathcal{D}_N \right\}. \quad (53)$$

Therefore, the orbit patterns in \mathcal{L}_N predetermine the inverse “variable mapping” V that was the premise in (13) and (14) and that naturally generalizes the Hohenberg–Kohn theorem on the entire set \mathcal{D}_N .

Note that \mathcal{L}_N includes N -electron Slater determinants which are structurally invariant with respect to $\mathcal{F}^{\times N}$. Define \mathcal{S}_N as the proper subset of \mathcal{L}_N consisting of Slater determinants. Since $\mathcal{F}^{\times N} \mathcal{S}_N \subseteq \mathcal{S}_N$, then $\mathcal{S}_N = \cup_i \mathcal{O}_S^{[i]}$ over all Slater orbits. Thus, we have:

Corollary 3.4. An arbitrary one-electron density $\rho(\mathbf{r}) \in \mathcal{D}_N$ is N -representable in \mathcal{S}_N .

5.4 Energy Density Functional and Variational Principle

5.4.1 Energy Density Functional: Definition

Proposal 3 definitely allows to propose the following rigorous definition of the energy density functional

$$\mathcal{E}_i[\rho(\mathbf{r})] \equiv \mathcal{E}_i[\rho(\mathbf{r}); \Phi_g^{[i]}] := \mathbf{E}[\Phi] |_{\Phi \in \mathcal{O}^{[i]} \subset \mathcal{L}_N} \quad (54)$$

and hence express the “functional mapping” (13) in the explicit way. It is evident that this mapping is one-to-many and there exist as many density functionals as there are orbits in \mathcal{L}_N . To derive $\mathcal{E}_i[\rho(\mathbf{r})]$ that is defined on the orbit $\mathcal{O}^{[i]} \subset \mathcal{L}_N$

explicitly, let us first write down the explicit expression for the energy functional $E[\Phi_g^{[i]}]$ of the orbit-generating wavefunction $\Phi_g^{[i]}$, in terms of its 1- and 2-matrices, $D_g^{1[i]}(x_1, x'_1)$ and $D_g^{2[i]}(x_1, x_2; x_1, x_2)$, respectively, and its one-electron density $\rho_g(x)$:

$$E[\Phi_g^{[i]}] = \frac{1}{2} \int d^4x_1 \nabla_{x_1} \nabla_{x'_1} D_g^{1[i]}(x_1; x'_1)|_{x'_1=x_1} + \int d^4x \rho(x) \hat{v}(\mathbf{r}) + \int d^4x_1 \int d^4x_2 \frac{D_g^{2[i]}(x_1, x_2; x_1, x_2)}{|\mathbf{r}_1 - \mathbf{r}_2|}, \quad (55)$$

where $\int d^4x \equiv \sum_s \int d^3\mathbf{r}$. Let us apply the local-scaling transformation that casts in **Proposal 3** to the wavefunction $\Phi_g^{[i]}$, precisely to its 1- and 2-matrices and its density. This yields

$$D_\rho^{1[i]}(\mathbf{r}'_1, s_1; \mathbf{r}'_1, s'_1) = [J(\mathbf{f}(\mathbf{r}_1); \mathbf{r}_1) J(\mathbf{f}(\mathbf{r}'_1); \mathbf{r}'_1)]^{1/2} D_g^{1[i]}(\mathbf{f}(\mathbf{r}_1), s_1; \mathbf{f}(\mathbf{r}'_1), s'_1), \quad (56)$$

$$D_\rho^{2[i]}(\mathbf{r}_1, s_1, \mathbf{r}_2, s_2; \mathbf{r}_1, s_1, \mathbf{r}_2, s_2) = J(\mathbf{f}(\mathbf{r}_1); \mathbf{r}_1) J(\mathbf{f}(\mathbf{r}_2); \mathbf{r}_2) D_g^{2[i]}(\mathbf{f}(\mathbf{r}_1), s_1, \mathbf{f}(\mathbf{r}_2), s_2; \mathbf{f}(\mathbf{r}_1), s_1, \mathbf{f}(\mathbf{r}_2), s_2), \quad (57)$$

and

$$\rho(\mathbf{r}, s) = J(\mathbf{f}(\mathbf{r}); \mathbf{r}) \rho_g(\mathbf{f}(\mathbf{r}), s). \quad (58)$$

Partitioning the 1- and 2-matrices, appearing in the rhs of Eqs. (56) and (57), into their local and non-local components:

$$D_g^{1[i]}(\mathbf{f}(\mathbf{r}_1), s_1; \mathbf{f}(\mathbf{r}'_1), s'_1) = [\rho_g(\mathbf{f}(\mathbf{r}_1), s_1) \rho_g(\mathbf{f}(\mathbf{r}'_1), s'_1)]^{1/2} \tilde{D}_g^{1[i]}(\mathbf{f}(\mathbf{r}_1), s_1; \mathbf{f}(\mathbf{r}'_1), s'_1), \quad (59)$$

$$D_g^{2[i]}(\mathbf{f}(\mathbf{r}_1), s_1; \mathbf{f}(\mathbf{r}_2), s_2; \mathbf{f}(\mathbf{r}_1), s_1, \mathbf{f}(\mathbf{r}_2), s_2) = \frac{1}{2} \rho_g(\mathbf{f}(\mathbf{r}_1), s_1) \rho_g(\mathbf{f}(\mathbf{r}_2), s_2) \left(1 + \mathcal{F}_{\text{xc}, g}^{[i]}(\mathbf{f}(\mathbf{r}_1), s_1, \mathbf{f}(\mathbf{r}_2), s_2) \right) \quad (60)$$

where $\tilde{D}_g^{1[i]}$ is the non-local part of the 1-matrix and $\mathcal{F}_{\text{xc}, g}^{[i]}$ is the non-local exchange-correlation factor. Therefore, the 1- and 2-matrices of (51) take the appearance:

$$D_\rho^{1[i]}(\mathbf{r}_1, s_1; \mathbf{r}'_1, s'_1) = [\rho(\mathbf{r}_1, s_1) \rho(\mathbf{r}'_1, s'_1)]^{1/2} \tilde{D}_g^{1[i]}(\mathbf{f}(\mathbf{r}_1), s_1; \mathbf{f}(\mathbf{r}'_1), s'_1), \quad (61)$$

and

$$D_{\rho}^{2[i]}(\mathbf{r}_1, s_1, \mathbf{r}_2, s_2; \mathbf{r}_1, s_1, \mathbf{r}_2, s_2) = \frac{1}{2} \rho(\mathbf{r}_1, s_1) \rho(\mathbf{r}_2, s_2) \times \left(1 + \mathcal{F}_{\text{xc}, \text{g}}^{[i]}(\mathbf{f}(\mathbf{r}_1), s_1, \mathbf{f}(\mathbf{r}_2), s_2) \right). \quad (62)$$

Finally, we obtain [4]

$$\begin{aligned} E[\Phi_{\rho}^{[i]}] &\equiv \mathcal{E}[\rho(x); \Phi_{\text{g}}^{[i]}] \\ &= \frac{1}{8} \int d^4x \frac{[\nabla_{\mathbf{r}} \rho(x)]^2}{\rho(x)} + \frac{1}{2} \int d^4x \rho(x) \nabla_{\mathbf{r}} \nabla_{\mathbf{r}'} \bar{D}_{\text{g}}^{1[i]}(\mathbf{f}(\mathbf{r}), s; \mathbf{f}(\mathbf{r}'), s')|_{x'=x} \\ &\quad + \int d^4x \rho(x) v(\mathbf{r}) \\ &\quad + \frac{1}{2} \int d^4x_1 d^4x_2 \frac{\rho(x_1) \rho(x_2) \left(1 + \mathcal{F}_{\text{xc}, \text{g}}^{[i]}(\mathbf{f}(\mathbf{r}_1), s_1, \mathbf{f}(\mathbf{r}_2), s_2) \right)}{|\mathbf{r}_1 - \mathbf{r}_2|}. \end{aligned} \quad (63)$$

Few statements can be drawn from Eq. (63):

- (i) The kinetic energy density functional is composed of two components. The first, the von Weizsäcker term, is local and orbit-invariant. The second is non-local, orbit-dependent, and due to the one-third power in Eq. (50), transformed to the modified Thomas–Fermi term within the local density approximation;
- (ii) The exchange-correlation energy density functional is explicitly expressed as

$$\begin{aligned} E_{\text{xc}}[\Phi_{\rho}^{[i]}] &\equiv \mathcal{E}_{\text{xc}}[\rho(x); \Phi_{\text{g}}^{[i]}] : \\ &= \frac{1}{2} \int d^4x_1 d^4x_2 \frac{\rho(x_1) \rho(x_2) \mathcal{F}_{\text{xc}, \text{g}}^{[i]}(\mathbf{f}(\mathbf{r}_1), s_1, \mathbf{f}(\mathbf{r}_2), s_2)}{|\mathbf{r}_1 - \mathbf{r}_2|}; \end{aligned} \quad (64)$$

- (iii) In fact, each density functional $\mathcal{E}[\rho(x); \Phi_{\text{g}}^{[i]}]$ depends on two basic variables: the one-electron density $\rho(x)$ and the generator wavefunction $\Phi_{\text{g}}^{[i]}$. Equation (42) expresses the energy as a functional of the one-electron density $\rho(x)$ within the orbit $\mathcal{O}^{[i]}$. True, Eq. (63) satisfies the condition of N-representability; (iv) One of the orbits in the decomposition (52) of \mathcal{L}_N is actually the orbit that contains the exact ground-state wavefunction. Refer it as the Hohenberg–Kohn orbit $\mathcal{O}^{[\text{HK}]}$ $\subset \mathcal{L}_N$. If a generator wavefunction is chosen to belong to $\mathcal{O}^{[\text{HK}]}$, Eq. (63) then determines the Hohenberg–Kohn energy density functional in the explicit manner.

5.4.2 Orbit Variational Principle and Euler–Lagrange Equation

The variational principle of the energy density functional theory based on the definition (33) is a straightforward consequence of the quantum mechanical variational principle (8) and the “functional mapping” (13). It is clearly orbit-dependent

or, equivalently, it is of the intra-orbit type. Let us consider the energy density functional $\mathcal{E}[\rho(\mathbf{r}, s); \Phi_g^{[i]}]$ given by Eq. (63) and defined within the $\mathcal{O}^{[i]}$ only. In this functional, $\rho(\mathbf{r}, s)$ stands for the density variable resulted from the initial density $\rho_g(\mathbf{r}, s)$ associated with the generator wavefunction $\Phi_g^{[i]}$. The extremum of $\mathcal{E}[\rho(\mathbf{r}, s); \Phi_g^{[i]}]$ on \mathcal{D}_N is attained at the i -th-optimal density $\rho_{\text{opt}}^{[i]}(\mathbf{r}, s)$ which is obtained by varying the following auxiliary functional:

$$\mathcal{E}[\rho(\mathbf{r}, s); \Phi_g^{[i]}] - \mu^{[i]} \left(\int d^4x \rho(\mathbf{r}, s) - N \right), \quad (65)$$

where $\mu^{[i]}$ is the Lagrange multiplier that accounts for the normalization of the density and that actually plays the role of a chemical potential on the orbit $\mathcal{O}^{[i]}$. Therefore, the stationary ground-state variational principle for the energy density functional $\mathcal{E}[\rho(\mathbf{r}, s); \Phi_g^{[i]}]$ is given by

$$\frac{\delta}{\delta \rho(\mathbf{r}, s)} \left\{ \mathcal{E}[\rho(\mathbf{r}, s); \Phi_g^{[i]}] - \mu^{[i]} \left(\int d^4x \rho(\mathbf{r}, s) - N \right) \right\} = 0, \rho(\mathbf{r}, s) \in \mathcal{D}_N, \quad (66)$$

we obtain the following integro-differential equation for the one-electron density [4]:

$$\begin{aligned} \frac{1}{8} \left[\frac{\nabla \rho(\mathbf{r}, s)}{\rho(\mathbf{r}, s)} \right]^2 - \frac{1}{r} \frac{\nabla^2 \rho(\mathbf{r}, s)}{\rho(\mathbf{r}, s)} + v_{\text{T},g}^{[i]}([\rho(\mathbf{r}, s)]; \mathbf{r}, s) + v(\mathbf{r}) + v_{\text{H}}([\rho(\mathbf{r}, s)]; \mathbf{r}) + \\ v_{\text{xc},g}^{[i]}([\rho(\mathbf{r}, s)]; \mathbf{r}, s) = \mu^{[i]}, \end{aligned} \quad (67)$$

where $v_{\text{H}}([\rho(\mathbf{r}, s)]; \mathbf{r}) = \int d^4x \rho(\mathbf{r}, s) |\mathbf{r} - \mathbf{r}'|^{-1}$ is the Hartree potential,

$$\begin{aligned} v_{\text{T},g}^{[i]}([\rho(\mathbf{r}, s)]; \mathbf{r}, s) = \frac{1}{2} \left\{ [\nabla_{\mathbf{r}} \nabla_{\mathbf{r}'}, \tilde{D}_g^{1[i]}(\mathbf{f}(\mathbf{r}), s; \mathbf{f}(\mathbf{r}'), s')]_{\mathbf{r}'=\mathbf{r}, s'=s} \right. \\ \left. + \rho(\mathbf{r}, s) \frac{\delta}{\delta \rho(\mathbf{r}, s)} ([\nabla_{\mathbf{r}} \nabla_{\mathbf{r}'}, \tilde{D}_g^{1[i]}(\mathbf{f}(\mathbf{r}), s; \mathbf{f}(\mathbf{r}'), s')]_{\mathbf{r}'=\mathbf{r}, s'=s}) \right\} \end{aligned} \quad (68)$$

is the potential originated from the non-local component of the kinetic energy in (63), and,

$$v_{\text{xc},g}^{[i]}([\rho(\mathbf{r}, s)]; \mathbf{r}, s) = \mathcal{E}_{\text{xc},g}^{[i]}([\rho(\mathbf{r}, s); \Phi_g^{[i]}]; \mathbf{r}, s) + \rho(\mathbf{r}, s) \frac{d\mathcal{E}_{\text{xc},g}^{[i]}([\rho(\mathbf{r}, s); \Phi_g^{[i]}]; \mathbf{r}, s)}{\delta \rho(\mathbf{r}, s)} \quad (69)$$

the exchange-correlation potential resulted from the non-local part of the electron–electron interaction where

$$\mathcal{E}_{\text{xc},g}^{[i]}([\rho(\mathbf{r}_1, s_1); \Phi_g^{[i]}; \mathbf{r}_1, s_1) = \frac{1}{2} \int d^4x_2 \frac{\rho(\mathbf{r}_2, s_2) \mathcal{F}_{\text{xc},g}^{[i]}(\mathbf{f}(\mathbf{r}_1), s_1; \mathbf{f}(\mathbf{r}_2), s_2)}{|\mathbf{r}_1 - \mathbf{r}|}. \quad (70)$$

Solving Eq. (67) for the given generator wavefunction $\Phi_g^{[i]}$, we obtain the i -th optimal or i -th approximate ground-state density $\rho_0^{[i]}(\mathbf{r}, s) \in \mathcal{D}_N$ and the i -th optimal or i -th ground-state energy

$$E_0^{[i]} \equiv \mathcal{E}_i[\rho_0(\mathbf{r})] \quad (71)$$

that simply casts as the i -th orbit variational principle:

$$E_0^{[i]} \equiv \inf_{\Phi \in \mathcal{O}^{[i]} \subset \mathcal{L}_N} \{E[\Phi]\} = E[\Phi]_{\Phi=\Psi_0^{[i]} \in \mathcal{O}^{[i]} \subset \mathcal{L}_N} = \inf_{\rho_\Phi \rightarrow \Phi \in \mathcal{O}^{[i]}} \{\mathcal{E}_i[\rho_\Phi]\}. \quad (72)$$

The next step is to substitute the densities $\rho_1(\mathbf{r})$ and $\rho_2(\mathbf{r})$ by $\rho_g^{[i]}(\mathbf{r})$ and $\rho_0^{[i]}(\mathbf{r})$ in Eq. (46) correspondingly and to solve the latter. The solution is the i -th optimal local-scaling transformation $\mathbf{f}_0^{[i]}(\mathbf{r}) \in \mathcal{F}$ which is further applied to $\Phi_g^{[i]}$ to get, via Eq. (51), the i -th optimal, ground-state wavefunction $\Phi_0^{[i]} \in \mathcal{L}_N$. True, generally speaking, the latter is the approximate ground-state wavefunction that yields an upper bound to the exact ground-state energy E_0 which is attained, by definition, only at the Hohenberg–Kohn orbit $\mathcal{O}^{[\text{HK}]}$, that is, $E_0^{[\text{HK}]} = E_0$.

5.4.3 Global Variational Principle: The Concept of Local-Scaling Self-Consistent Field

The orbit variational principle (65) deduced in Sect. 5.4.2 is solely defined on a particular orbit. The reason is trivial: this is precisely that orbit where the energy density functional is defined according to Eq. (54). In contrast, the global ground-state quantum mechanical variational principle (8) is carried out over the whole Hilbert space \mathcal{L}_N . Within the local-scaling formulation of the density functional approach is achieved due to the fact that the energy density functional in fact depends on two basic variables of theory: one—the one-electron density—is the key variable of the density functional theory and the other is the generator wavefunction that determines an orbit. Hence, the orbit partitioning (52) of \mathcal{L}_N is governed by the orbit generators. Therefore,

$$E_0 = \inf_{\substack{\text{over} \\ \text{all orbits} \\ \mathcal{O}^{[i]} \subset \mathcal{L}_N}} \left\{ \inf_{\rho(\mathbf{r}) \in \mathcal{D}_N} \{\mathcal{E}[\rho(\mathbf{r}); \Phi_g^{[i]}]\} \right\}. \quad (73)$$

Equation (73) implies that the search for the exact ground-state wavefunction must be carried out by a combined intra-orbit and inter-orbit minimization [7]. The former reflects the charge consistency variational principle, whereas the latter the

inter-orbit one, the orbit consistency. The latter is actually the variational principle of the “inter-orbit” self-consistent field that resembles the Kohn–Sham self-consistent field approach and results in inter-orbit “jumps” that finally leads to the exact, Hohenberg–Kohn orbit.

6 Overlook

All of what is scientific in chemistry is physics—the rest is cooking.
L. D. Landau [187]

I have a feeling that a success in the adequate description of dispersion interactions within the DFT has been achieved. Again what we have at our current disposal is a number of D-DFT functionals among which choosing the best still remains a challenge that leaves a feeling of certain yet-imperfectness in this DFT area and, on the other hand, shows the ways to improve it. This was in fact the key goal undertaken in the present review which has definitely been achieved, I believe.

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Redox Activation of Small Molecules at Biological Metal Centers

Radu Silaghi-Dumitrescu

Abstract Some recent applications of density functional theory (DFT) are described and specifically the review addresses the activation of small molecules such as water, molecular oxygen and its reduced congeners, nitrogen oxides and oxyanions, sulfide, sulfur oxides and oxyanions, carbon dioxide, organic compounds, halogens, molecular hydrogen, and protons. A range of predictions on geometry, electronic structure, and spectroscopic properties are made and binding energies and activation energies are critically reviewed.

Keywords DFT • Metalloenzyme • Small molecule activation

Contents

| | | |
|-----|---|-----|
| 1 | Introduction | 98 |
| 2 | Molecular Oxygen, Superoxide, Peroxide, Water | 99 |
| 2.1 | Dioxygen Activation | 99 |
| 2.2 | Superoxide Reductases and Dismutases | 104 |
| 2.3 | Redox Activation of Water/Hydroxide | 106 |
| 3 | Nitrogen Oxides and Oxyanions | 106 |
| 3.1 | Nitric Oxide | 106 |
| 3.2 | Nitrite and Nitrate | 107 |
| 4 | Sulfur-Based Compounds | 107 |
| 5 | Carbon-Based Compounds | 109 |
| 6 | Halides and Their Derivatives | 111 |
| 7 | Hydrogen/Protons | 113 |
| 8 | Summary and Outlook | 114 |
| | References | 114 |

Abbreviations

| | |
|---------|----------------------------------|
| CANH | Carbonic anhydrase |
| CPOI | Chloroperoxidase |
| DFT | Density functional theory |
| Fe-SOD | Iron-containing SOD |
| MM | Molecular mechanics |
| MPO | Myeloperoxidase |
| Ni-SOD | Nickel-containing SOD |
| NOR | Nitric oxide reductase |
| P450NOR | P450-like nitric oxide reductase |
| P450 | Cytochrome P450 |
| QM | Quantum mechanics |
| SOD | Superoxide dismutase |
| SOR | Superoxide reductase |
| TD-DFT | Time-dependent DFT |
| TSR | Two-state reactivity |
| VPO | Vanadium-containing peroxidase |

1 Introduction

Activation of small molecules by metal centers may occur via mechanisms involving changes in redox state, spin state, bond breaking and formation, or acid–base properties [1, 2]. Density functional methods (DFT) are particularly suited for providing insights into such aspects [3]. Biological metal centers may be deemed to have evolved so that the biomolecule part would selectively favor one (or at least a small number) of the many reactions and mechanisms of which the metal center is intrinsically capable. On one hand, this selectivity implies a less complicated array of reactions connecting reactants and products (presumably a single shortest path allowed by chemical rules), compared to what would be observed for simpler synthetic metal complexes (multiple uncontrolled parallel reactions with multiple products). Such presumed simplicity is a remarkable advantage for experimentalists seeking to investigate these pathways. On the other hand, precisely the specialization brought about by the biomolecules around the metal, often entails not only particular sensitivity to reaction conditions (e.g., proteins stable only in biological membranes or with very labile metal-binding sites) but also very efficient catalysis—meaning that the lifetimes of the investigated reaction intermediates will often seriously challenge the performance of most cutting-edge experimental techniques. In such cases, DFT methods have the advantage of allowing extensive examination not only of fleeting/unstable reaction intermediates but also of otherwise unobservable transition states.

In the following pages, some recent applications of DFT are discussed for activation of small molecules such as water, molecular oxygen and its reduced congeners, nitrogen oxides and oxyanions, sulfide, sulfur oxides and oxyanions, carbon dioxide, organic compounds, halogens, molecular hydrogen, and protons. A range of predictions on geometry, electronic structure, and spectroscopic properties are made and binding energies and activation energies are critically reviewed.

2 Molecular Oxygen, Superoxide, Peroxide, Water

Oxygen-based small molecules are among the longest known substrates for bioinorganic chemistry and are pervasive throughout processes essential for life—respiration, photosynthesis, defense, or selective biosynthesis of essential building blocks within the human organisms [1, 2]. The constant progress made by theoretical chemists in these areas, with DFT being a key tool, is therefore not surprising.

2.1 Dioxygen Activation

In heme-containing monooxygenases and peroxidases, a general theme is cleavage of the oxygen–oxygen bond, leading to formally high oxidation states at the metal (Fe(IV), Fe(V), also referred to in the literature as “high-valent”) [4–7]. Figure 1 illustrates the main reaction pathways accepted to be involved in these processes and the key players. It may be seen that small changes in the first coordination sphere, as well as changes in the second coordination sphere, can significantly affect the choice of reaction pathway. Globins normally oscillate between the “oxy” and the Fe(II) forms, with side reactions leading to Fe(III) (“autooxidation”) and then possibly to the ferric-hydroperoxo → Compound I route. Peroxidases normally use the path between the ferric, ferric hydroperoxo, Compound I/II, and ferric-hydroxo states. Cytochrome oxidases close a catalytic cycle that connects the Fe(II) state with the “oxy,” then all the way to Compound II and Fe(III)-OH, and then back to Fe(II). Heme oxygenases, similarly to cytochrome oxidases, start out at Fe(II) and to the ferric-hydroperoxo state via the oxy; however, at the ferric-hydroperoxo stage the site self-destructs, as the heme itself is the substrate of this enzyme. Notably, in all these enzymes of such different reactivities, the axial ligand (*X* in Fig. 1) is histidine—illustrating how important the role of the second coordination sphere may be.

Binding of molecular oxygen (often referred to as “dioxygen”) to heme centers has been described computationally for several proteins, with emphasis on the electronic structure within the Fe–O₂ moiety, on the reversibility of the process and on the importance of spin state preferences [8–11]. An interesting question to

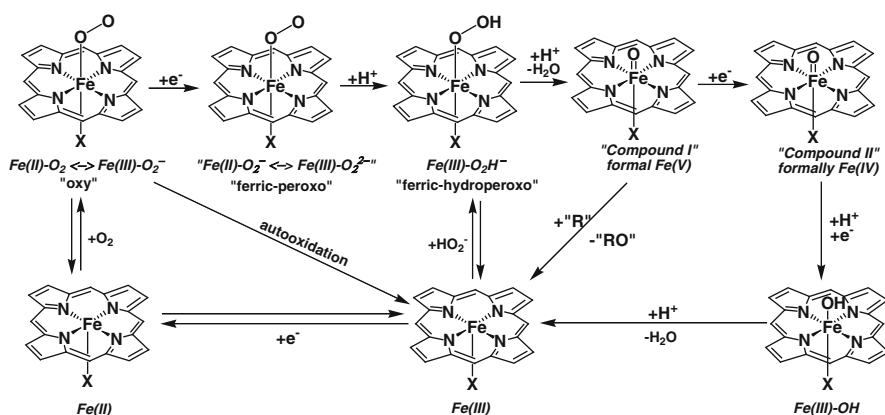


Fig. 1 Reactions involving molecular oxygen and its reduced congeners in heme proteins. Reactions where equilibria are manifest are indicated as such. In globins, peroxidases, cytochrome oxidases, and heme oxygenases, the axial ligand X is a protein-derived histidine; in catalases, X is a tyrosine phenolate; in chloroperoxidases and cytochromes P450, X is a cysteine thiolate

investigate is also the reason why globins and heme oxygenase (or indeed cytochrome oxidases) make such drastically different choices with respect to the stability of the ferrous–dioxygen adduct towards reduction: stable in one case, easily reduced in the other cases. The computed thermodynamics indicate that the one-electron reduction of the ferrous–dioxygen adduct is indeed reasonably allowed—certainly more allowed than the reduction of ferrous–dioxygen adducts of cytochrome P450, where the axial ligand is not a neutral imidazole ligand but rather an anionic thiolate from a cysteine [8, 9]. Paradoxically, precisely this cysteinate has been considered a key element in *facilitating* dioxygen activation—an effect that was dubbed *thiolate push* on the grounds that the extra charge density at a sulfur would, by being transferred onto the dioxygen via the iron, help occupy the antibonding orbitals of the O₂ ligand and thus weaken the O–O bond as well as increase its proton affinity (protonation being, as seen in Fig. 1, a key prerequisite for O–O bond cleavage) [9, 10]. The *thermodynamic* obstacle presented by the thiolate ligand in the early step of dioxygen reduction has in this context been dubbed *thiolate obstruction*, and, as detailed in the next sections, is seen at work in more than one instance in small molecule activation [9, 12, 13].

The nature of the higher oxidation state species in hemoproteins, especially the Compound I and Compound II depicted in Fig. 1, has been thoroughly revisited in recent years. DFT-based predictions, based on proton affinities and electronic structures [14], have argued that many of these high oxidation states would in fact occur in a protonated form (Fe(IV)–hydroxo, as opposed to Fe(IV)=oxo)—and these predictions were soon conformed experimentally [4, 7, 15–20]. A debate is nevertheless still open about the extent of these protonations, with arguments supporting either one of two theories (1) that only thiolate-ligated Compound II species may be anionic (and, possibly, other anion-ligated species) [19] and (2) that all Compound II species are protonated [4]. Essentially, this debate revolves around

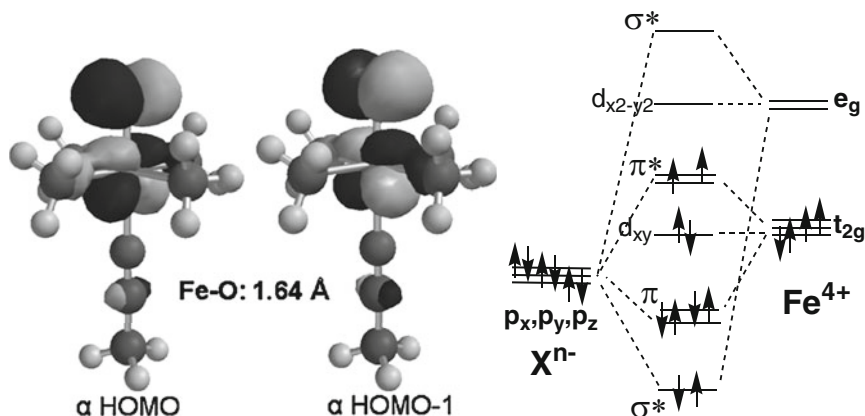


Fig. 2 Molecular orbital diagram and iron–oxygen π^* orbitals for an octahedral ferryl model, with four equatorial amine ligands and an axial acetonitrile, illustrating a high degree of covalence and a triplet dioxygen-like electronic structure [22]

the pK_a of the formally Fe(IV)=oxo moiety in the histidine-ligated Compound II. Estimates based on experiment have placed this pK_a at ~ 4.7 [21], which being close to the pH range where most experiments are conducted, and being subject to temperature dependence in low-temperature crystallography and spectroscopy measurements, may partly explain why different groups looking at the same experiments appear to obtain conflicting results [4, 19, 21]. On the other hand, for Compound I species where the extra oxidizing equivalent compared to Compound II is still located on the heme-oxo moiety, the protonated state is predicted by DFT to be too acidic to be observed under physiological conditions [21].

Bonding within the Fe(IV)=oxo moiety has been described by several groups, highlighting a strongly covalent interaction with a preferred dioxygen-like triplet ground state, as illustrated in Fig. 2 [22–24]. Nevertheless, Fig. 3 illustrates that according to non-DFT computations (Hartree-Fock, MP2 [22]) an alternative description of these centers may be proposed, where the two unpaired electrons reside one on the oxygen and one on the iron—making for an Fe(III)-oxyl unit (formally, a deprotonated hydroxyl radical). Such a description would fit well with the fact that “ Fe(IV)=oxo ” complexes are capable of hydrogen atom abstraction, much like hydroxyl radical. In models where the oxo ligand was replaced by sulfido or by nitride ligands, DFT has allowed the “covalent deadlock” to be broken: population analyses and molecular orbitals have revealed that with sulfide and nitride the interaction is no longer a “triplet dioxygen-like,” and that the monoatomic ligands are in fact partially oxidized, in the same manner previously predicted by MP2 calculations [24, 25]. Under these circumstances, one may argue that many biological Fe(IV)=oxo complexes are in fact better described as *not* containing iron in a high oxidation state—but rather an oxyl ligand bound to Fe (III) [24]. Evidence for an oxyl-type structure as well as for an oxyl-type reactivity with high-oxidation state centers of other metals, such as manganese, is also available [26, 27].

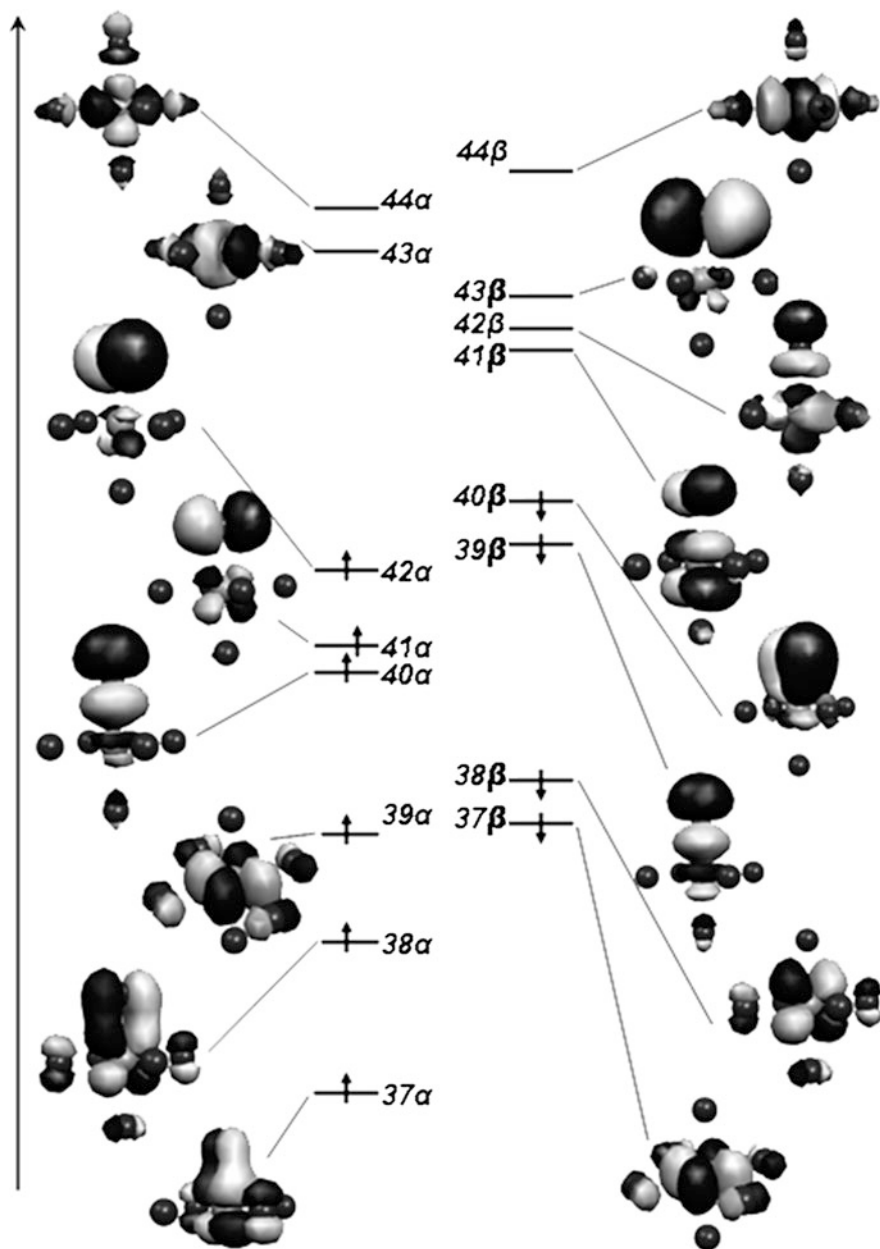


Fig. 3 UMP2/6-311 + G** molecular orbital diagram for a $S = 1$ $[\text{FeO}]^{2+}$ complex in an octahedral environment [22]. The π^* orbitals are 41 and 42 on the α manifold, and 41 and 43 on the β manifold; 43 β illustrates a distinct “hole” in the oxygen p orbitals, and implicitly 38 β illustrates one extra electron in the Fe orbitals, compared to the expected Fe(IV) d^4 description—hence, a d^5 Fe(III) iron bound to an O^{-1} oxyl ligand

A complication of the oxo-containing Compound I species is the presence of one extra unpaired electron, which may couple to the triplet iron-oxo moiety either ferromagnetically (to yield a net doublet) or antiferromagnetically (yielding a quartet). According to DFT data, these two spin states display slightly different reactivities in terms of kinetics and of mechanism. Such differences were proposed to explain experimental observations according to which some cytochrome P450-catalyzed reactions would incur two different oxidants. This concept is now known as *two-state reactivity* [23]. On the other hand, with more recent data now suggesting that protonated oxo groups may also be effective oxidants (*vide infra*), and with other data supporting the involvement of other intermediates in the catalytic cycle as direct oxygenation agents, or the fact that in nonheme enzymes the ferric-hydroperoxo rather than a ferryl species is accepted as the active oxygenating agent [28], the concept of *mechanistic promiscuity* was proposed—describing a situation where a substrate may be converted into a single product via more than one mechanism [27, 29]. Examples have been given of other enzyme mechanisms where this concept may apply—pointing out that mechanistic promiscuity would be an evolutionary advantage for an enzyme [30–32].

Comparisons between the reactivities of the protonated and nonprotonated high-oxidation state Fe(IV)-hydroxo/Fe(IV)=oxo states towards hydrocarbons have been attempted, with hydrogen abstraction processes being taken as representative for enzymatic P450 chemistry. In some of the studies, the oxo and hydroxo ligands were in fact placed as ligands to the same metal in the same model, in order to allow more straightforward comparisons [27, 33, 34]. Although the free-radical character assigned to the oxo ligand (both in the dioxygen-like description and in the MP2-like description) would suggest that the oxo atom would be distinctly more adept than a hydroxo unit at activating hydrocarbons, it was surprisingly found that the barriers for hydrogen atom abstraction from model hydrocarbons by oxo vs. hydroxo ligands are remarkably close to each other [27]. As such, the thermodynamics of the process (and not the kinetics) would be the ones dictating the preferred reaction pathways.

Nonheme iron proteins deal with molecular oxygen on the same principles as illustrated in Fig. 1. Differences arise from the increased mobility of the coordination sphere, which often allows dioxygen, superoxide and peroxide to bind in a bidentate fashion to the metal. Additionally, the absence of the porphyrin “charge reservoir” may disfavor oxygen–oxygen bond cleavage thermodynamically. The case of Rieske dioxygenases, where two different DFT functionals, coupled with two slightly different choices of model, yielded two apparently conflicting results, is particularly instructive. Thus, on one hand B3LYP data suggest that indeed substrate oxygenation occurs reasonably well directly from the ferric-hydroperoxo state, with an activation barrier higher by a few kcal/mol than the barrier for O–O bond cleavage [28]. On the other hand, BP86 results suggest the opposite: O–O bond cleavage favored by a few kcal/mol over the direct oxygenation of substrate by a ferric-hydroperoxo adduct; then, such a high oxidation state would very easily oxygenate the substrate [35, 36]. This dependence of the outcome on the choice of the functional is important to remember whenever one examines conclusions based

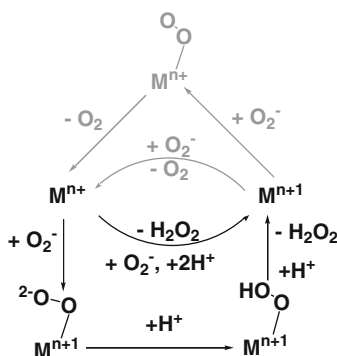


Fig. 4 Reaction pathways for superoxide dismutases and superoxide reductases. Shown in *black* are the two possible pathways leading to oxidation of the active site by superoxide, producing peroxide; this half of the catalytic cycle is common to SOR and SOD. Shown in *gray* is the re-reductive path specific to SOD, and which in SOR is replaced by a single-electron transfer from the specialized redox protein, rubredoxin. M may be Fe ($n = 2$, in SOR, or in SOD), Ni, Mn ($n = 2$, in SOD), or Cu ($n = 1$, in SOD)

on energy differences of less than ~ 10 kcal/mol: two barriers computed to be different by that much, may, for all we know, be perfectly equal in reality—or even display the opposite trend compared to what our single DFT method has predicted.

2.2 Superoxide Reductases and Dismutases

Figure 4 shows proposed catalytic mechanisms for superoxide dismutases (SOD) and superoxide reductases (SOR) [37, 38]. A single redox metal is present at all these enzyme active sites, oscillating between two oxidation states (+2 and +3 for Fe, Ni, or Mn, +1 and +2 for Cu).

A common theme of the SODs and SORs is the question of whether the reduction and/or oxidation of superoxide occurs inner-sphere or outer-sphere. Although there appears to be a consensus, strongly supported computationally, that on both pathways binding of the superoxide to the metal is feasible [38–40], the issue of whether this binding is a prerequisite for completing the electron transfer, or a side effect of it, is still open for discussion. Thus, for nickel-containing superoxide dismutases (NiSOD) the nonprotonated Ni(II)–superoxide complex was predicted by DFT to be unfeasible, featuring the superoxide noncovalently associated with the nickel ligands, rather than coordinated at the Ni itself; at this point, contrary to the formal description from Fig. 4, a Ni(II)–superoxo rather than a Ni(III)–peroxo description was given based on the population analyses and on the O–O bond length [38]. However, protonation of the superoxide led to a state where the OOH moiety was clearly describable as a hydroperoxide, and was shown to be able to coordinate to Ni—hence, a Ni(III)–hydroperoxo state [38]. The one-electron

reduction of the superoxide to (hydro)peroxide appears to be mainly provoked by the protonation event and not by coordination to the metal: indeed, even at Ni–OOH distances excluding coordinative interaction, the OOH moiety retained its hydroperoxo character, suggesting that the reduction of superoxide in NiSOD operates via a protonation-induced outer-sphere electron transfer [38]. Under these conditions, while it is entirely possible for the hydroperoxide to bind to the metal, this event would happen only *after the reduction of superoxide was completed*—and may thus be regarded as a form of substrate inhibition rather than as a key step within the catalytic cycle. Interestingly, for Fe–SOD [38] or for copper-containing SOD [40] such a clear-cut description is not possible: the M^{n+} -superoxo adduct is computed to feature reasonably short Fe–OO bonds (and, incidentally, nonnegligible M^{i+1} -peroxo character). Remarkably, while the NiSOD active site is completely solvent exposed, the iron, the manganese, and the copper SODs have their metal sites further buried inside the protein, as if to preclude superoxide–metal coordination as a form of a nonproductive/inhibitory interaction [38].

For the putative ferrous–superoxide complexes in SOD as well as in SOR, the alternative electromeric state, resulted from a one-electron transfer (ferric-peroxo), has been favored [38], although the O–O bond length, computed at ~ 1.36 Å with the BP86 functional and slightly shorter with hybrid functionals, was intermediate between those computed to free hydrogen peroxide and free superoxide [37]. Nevertheless, the most recent computational data for SOR suggest a high-spin ferrous-superoxo state, as characterized by \sim one unit of unpaired electron density at the OO ligand, and by O–O bond lengths closer to free superoxide than to free hydrogen peroxide [37, 41]. Furthermore, TD-DFT UV–vis computations suggest that the electronic absorption spectrum of such a species may even feature a band at ~ 600 nm, in the same region where the resting ferric state features a ligand-to-metal transition (sulfur-to-ferric); spectacularly, in the ferrous-superoxo adduct this band would now arise from *ligand-to-ligand* and *metal-to-ligand* charge transfers [41]. An ongoing debate on whether such a nonprotonated reaction intermediate would be observable is fuelled by reports according to which only one reaction intermediate is observed experimentally for SOR [42]. Based on DFT-derived thermodynamics, correlated with crystallographic information on the O–O bond length [39], this intermediate would most likely be the ferric-hydroperoxo, rather than the nonprotonated ferrous-superoxo—especially as the active site is water exposed and decorated with two potential proton-donating amino acids [37, 39].

The re-reductive pathway, shown in gray in Fig. 4 and specific to SOD but not to SOR, appears more likely to prefer the outer-sphere pathway. Thus, according to DFT data, superoxide coordination to Ni(III)–SOD would be disfavored thermodynamically and unnecessary: the electron transfer from superoxide to Ni, to yield a neutral dioxygen molecule loosely associated with Ni(II)–SOD, occurs even at nonbonding Ni–OO distances [38]. For Fe–SOD, although the diamagnetic ferrous-dioxygen/ferric-superoxo state appeared to feature bond lengths reminiscent of the isoelectronic stable oxy form of globins, the higher spin states, including an essentially nonbonding $S = 3$ situation, were thermodynamically favored, suggesting a drive to avoid proper metal–superoxide adducts on this pathway [38].

2.3 *Redox Activation of Water/Hydroxide*

As detailed above in the discussion on metal centers in high oxidation states, water may in principle be oxidized upon binding to a metal center, to yield oxyl units or even electrically neutral oxygen atoms. The photosynthetic pathways in plants use this reaction as a convenient source of electrons—with a specialized four-manganese cluster at the active site of the enzyme responsible for the reaction, in an interesting parallel with the Mn-oxyl evidence cited above for mononuclear model compounds. Recent progress in X-ray diffraction determinations has allowed intimate attempts of correlating geometrical and EPR (electron paramagnetic resonance) data with DFT-derived data, attempting to assign oxidation and protonation states in the reactive intermediates en route from water to dioxygen. Paradoxically to some extent, much more appears to remain uncovered on this pathway than on the reverse pathway—dioxygen activation discussed earlier in this chapter [43–46].

3 Nitrogen Oxides and Oxyanions

The nitrogen cycle in nature involves complex transformations, via at least three separate pathways, between nitrate and organic nitrogen/ammonia. Every single step in these processes is catalyzed by metalloenzymes [47]—some of which (e.g., nitrogenase, nitrous oxide reductase) have offered remarkable findings in recent years and continue to pose interesting challenges to computational chemists [48]. Among other enzymes, nitrogenase continues to tantalize computational chemists with new structural features of the iron–sulfur–molybdenum cofactor at active site, especially the mysterious central atom within this cluster; while a few years ago it appeared that computational methods would correctly predict not only structural but even spectroscopic parameters going as far as Mössbauer, recent data suggests that in the center of the cluster is a previously unsuspected carbon atom—hence a new member of the bioorganometallic family [49].

3.1 *Nitric Oxide*

Most nitric oxide reductases (NOR) fall into two categories: fungal P450-type NOR and bacterial heme/nonheme NOR. P450NOR illustrate a unique mechanism wherein a hydride atom is transferred directly from NAD(P)H to an NO moiety bound at the single heme active site [12]. By contrast, bacterial NOR feature a binuclear active site with a heme and a nonheme iron facing each other; here, the electrons required for reducing nitric oxide are delivered in sequential one-electron steps. Interestingly, while in P450NOR the axial ligand at the heme is a thiolate, in

the bacterial NOR the axial ligand is a histidine. DFT calculations have shown that a sequence of two successive one-electron reductions of an NO molecule bound to an Fe(III) heme would be thermodynamically feasible with a neutral ligand (e.g., histidine), whereas with a thiolate axial ligand the reaction would be blocked at the Fe(II)–NO stage, after the first electron transfer (an example of thiolate obstruction, similar to those discussed above for dioxygen activation) [9, 12]. The thiolate-bound active site of P450NOR manages to bypass the Fe(II)–NO state, by transferring both electrons at once, in the form of a hydride, to the initial Fe(III)–NO adduct—yielding an Fe(II)–HNO state.

Oxidation of nitric oxide by ferrous–dioxygen adducts of hemoproteins occurs accidentally in blood (with hemoglobin) but also as physiological function for certain bacterial globins (e.g., flavohemoglobin); the mechanism of this reaction appears to entail ferric–peroxynitrite adducts at the heme, whose extremely short lifetimes were predicted by DFT calculations [50, 51].

3.2 Nitrite and Nitrate

Nitrite binding to metals is well known in traditional inorganic chemistry as an example of linkage isomerism—occurring either via the nitrogen or via the oxygen atom, cf. Fig. 5. In the past few years it has been recognized that the same linkage isomerism is also at work in metalloproteins [30, 52–56]. An initial DFT prediction was made despite a wealth of experimental data showing nitrite binding to hemoproteins only via its nitrogen atom, binding via the oxygen should also be possible [30]. Soon afterwards, the crystal structures of nitrite-bound myoglobin and hemoglobin were reported, showing indeed the DFT-predicted isomer [52–56]. Subsequent detailed studies, with DFT, *ab initio* dynamics, and QM/MM approaches [57] have also revealed that nitrite reduction is almost equally feasible via the two isomers, suggesting perhaps one more case of mechanistic promiscuity, as illustrated in Fig. 5 [57]. Also based on DFT data, linkage isomerism was proposed to be feasible in copper-containing nitrite reductases [58], as well as in molybdenum-containing nitrate reductases—as the latter enzymes can reduce nitrite at competition with nitrate [32]. In the case of nitrate reductase, nitrate and nitrite were computed to be reducible at very similar rates [32].

4 Sulfur-Based Compounds

Sulfur, its oxides and its oxyanions are placed on metabolic pathways involved in energy conservation/extraction in bacteria, and are more generally important as basic reagents in biochemical experiments (with dithionite a prime example) [59, 60].

$\text{Fe(III)}\text{--SO}_2^0$ species where the ligand binds to iron via an *oxygen* atom, in an example of linkage isomerism reminiscent of the nitrite case and unprecedented for sulfur compounds in enzymes. A facile one-electron reduction to a $\text{Fe(III)}\text{--SO}_2^-$ adduct ensues. This adduct would then undergo two subsequent protonation events and eliminate a water molecule to yield a $\text{Fe(III)}\text{--SO}^+$ species. The latter was computed to easily undergo three successive one-electron reduction events, to $\text{Fe(II)}\text{--SO}^-$. From here, two successive protonation events were computed to lead to a Compound I-type species, with a $\text{Fe(IV)}\text{--sulfide}$ moiety coupled to a cation radical delocalized onto the axial thiolate and onto the porphyrin. A one-electron reduction and a subsequent protonation would lead to an $\text{Fe(IV)}\text{--SH}$ state akin to the $\text{Fe(IV)}\text{--OH}$ Compound II species seen in thiolate-ligated hemoproteins from the chloroperoxidase family. This protonated sulfide Compound II would then return to the ferric state by a further one-electron reduction, liberating hydrogen sulfide in a protonation-dependent final step of the catalytic cycle. The sulfide adducts are likely to provide further computational interest, in light of hydrogen sulfide's recently discovered implications in cellular signaling; [62] sulfur oxyanions also appear to offer surprises in coordination chemistry not only in heme proteins but also in systems such as cobalamin [63].

5 Carbon-Based Compounds

Carbon-based compounds such as carbon dioxide or aliphatic hydrocarbons pose challenges of reactivity to metalloenzymes insofar as they are generally reluctant to bind to metal active sites; a general rule is that such substrates would usually be attacked by metal-activated nucleophiles. However, in selected cases, even binding to metal may be feasible, as detailed below.

One class of carbon dioxide-activation enzymes, the carbonic anhydrases (CANH), employ a metal-bound hydroxide as a nucleophile with (bi)carbonate as a final product [64]. As illustrated in Fig. 7, the metal employed in CANH is zinc; recent DFT data shows that the choice of zinc in this enzyme (as opposed to other transition metals, such as cobalt, iron, or copper) is not so much related to differences imparted on the nucleophilicity of the metal-bound hydroxide by the various metals: indeed, hydroxides in other enzyme models (including well-known species such as met-hemoglobin, lipoxxygenase, or a superoxide reductase mutant), as well as with other metal centers, were all computed to require reasonably small activation energies when reacting with carbon dioxide [64]. Nevertheless, in most cases the coordination environment around the metal was too crowded, thus hindering an efficient binding of the product to the metal—eventually resulting in an endergonic process. Therefore zinc, with its preferred tetrahedral coordination and reasonable kinetic lability (arguably, eventually linked to other intrinsic electronic properties [65, 66]), is ideally suited for the CANH reaction.

Carbon dioxide may also be activated via direct redox reaction with metals—possibly involving metal–carbon bonds, and necessarily involving low oxidation

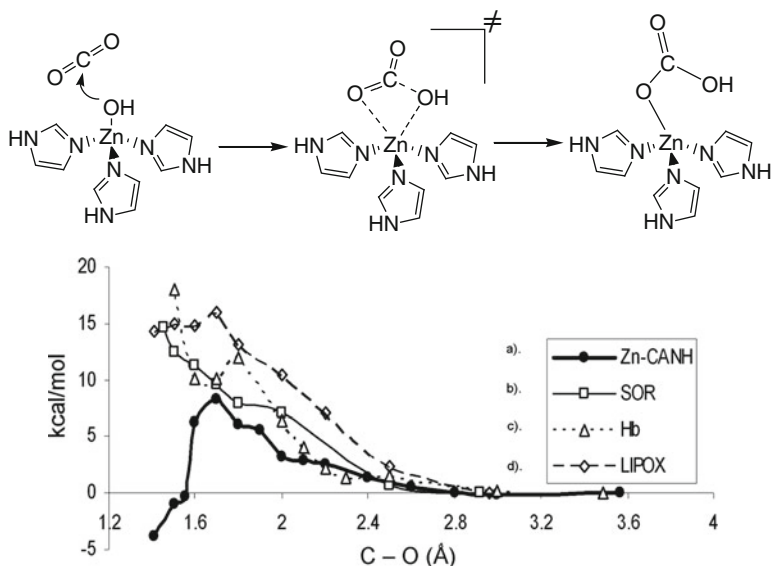
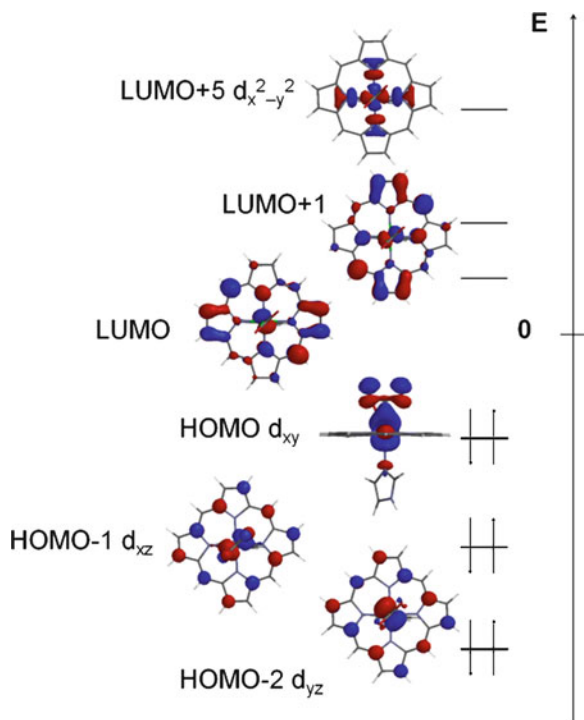


Fig. 7 Upper panel: currently accepted mechanism for CO₂ activation by carbonic anhydrases. Lower panel: comparison of reaction profiles for CANH versus three other enzyme active sites, not known to catalyze carbon dioxide hydration (SOR superoxide reductase, Hb hemoglobin, LIPOX lipoxigenase)

states at the metal [67]. A recent survey of low-oxidation state heme and related systems, with Fe(0) and Fe(I) states [67, 68], has revealed that although in general the extra oxidizing equivalents in such “super-reduced” states will tend to be localized on the ligands rather than on the metal itself, in one case an Fe–CO₂ bond could be established, with an electronic structure suggestive of a true low oxidation state at the metal. Thus, Fig. 8 illustrates the frontier molecular orbitals for an Fe(0) heme system ligated axially by CO₂ and imidazole; it may be seen that only the $d_{x^2-y^2}$ orbital at the metal is empty—hence a formally d^8 description, suggestive of an Fe(I) center bound to an activated carbon dioxide radical. This is relevant in the context where such low oxidation states are known experimentally to be involved in catalytic reduction of carbon dioxide [69, 70]. Interestingly, the CO adducts of Fe(0) or Fe(I) heme-like systems were computed to in fact consist of Fe(II) coupled to macrocyclic anion radicals [67].

The oxygenation of organic compounds at metal centers is often accomplished by metal-bound oxygen atoms—be it a hydroxide as in lipoxigenase, or a peroxide-derived atom as in cytochromes P450 or Rieske dioxygenases discussed above [25, 27, 29, 71]. An interesting alternative, not seen in enzymes but seen in model compounds, are the metal-bound nitrites: these are known experimentally to act as reasonable oxygenation agents; their reactivity in this respect has been tested with DFT methods, revealing that despite an apparently drastic difference in electronic structure, the oxyl-like ligand in P450 Compound I (as discussed above) and an

Fig. 8 Metal-localized frontier orbitals for a formally Fe(0) heme bound to carbon dioxide, with an imidazole ligand trans to the CO₂



oxygen atom within a ferric-nitro adduct, feature very similar energy profiles [71]. A key role was therefore inferred the redox potential in dictating oxygenation ability, in good agreement with a previous proposal of Green and coworkers based on P450 models alone [7, 71] (Fig. 9).

6 Halides and Their Derivatives

Heme enzymes such as chloroperoxidase (CPO) and myeloperoxidase (MPO) perform a two-electron oxidation of halides to formally generate hypohalous acids, HOX. These acids can either be released or used for halogenation reactions. Most peroxidases oxidize iodide, some oxidize bromide, and very few (of which MPO and CPO are classical examples) are known to oxidize chloride. Using UV-vis spectroscopy, intermediates arising from the reaction of Compound I with chloride and proposed to consist of Fe(III) adducts with HOCl and OCl⁻ were detected with model compounds as well with CPO. Recently, the structure of these intermediates was explored with DFT. The oxygen-halogen bonds were computed to be particularly weak (cf. Fig. 10, broken exergonically with a barrier of only ~10 kcal/mol), to the extent where ferric hypohalous acid complexes with intact oxygen-halide bonds should be difficult to observe in heme peroxidases. Instead, a “caged” adduct, featuring a ferric species in close (but not covalent) contact with the oxidized halide, appears

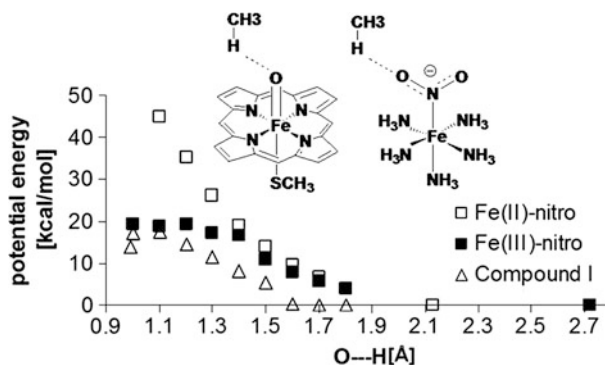


Fig. 9 Potential energy as a function of the O...H distance for hydrogen atom abstraction from methane by three different agents: a thiolate-ligated Compound I, or a ferric or ferrous-nitro complex (cf. structures shown). For each model, the energy of the reactants (*far right side* of the plot) is taken as reference [71]

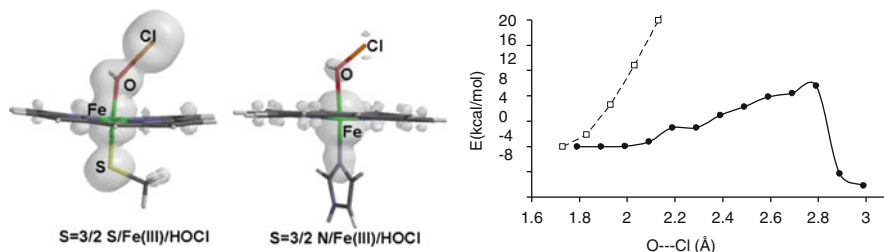


Fig. 10 *Left*: optimized geometries and spin density distributions in models of ferric heme-containing haloperoxidases bound to hypochlorous acid, with axial thiolate or imidazole ligands. *Right*: Potential energy surface for O-Cl bond elongation (*black circles*) in the thiolate-ligated Fe-O-Cl model. *White squares*: equivalent potential surface in an isolated HOCl molecule

as a likely alternative for an observable intermediate in CPO and MPO. The presence of an axial thiolate (cysteinate in CPO) or imidazole (histidine in MPO) ligated to the iron trans to the hypohalous moiety was calculated to have a distinct effect on halide activation, including a spin-state preference with relevance to the TSR (two-state reactivity) concept discussed above [72]. Indeed, Fig. 10 illustrates optimized geometries and computed spin densities for ferric-HOCl models of CPO and MPO. Notably, the O-Cl bond is almost completely broken in the thiolate-ligated model (and consequently significant spin density accumulates to the chlorine), whereas the same bond is essentially intact in the imidazole-ligated model (with the Fe-O bonds following opposite trends—i.e., weaker in the imidazole-ligated model). Furthermore, the thiolate-ligated model, but not the imidazole-ligated one, has a low-spin ground state, where the O-Cl bond is even further activated. Remarkably, in good correlation with these differences, the thiolate-ligated CPO specializes in immediate cleavage of the O-Cl bond so that it can insert the resulted chlorine atom into a hydrocarbon, whereas the imidazole-ligated MPO specializes in liberating intact HOCl.

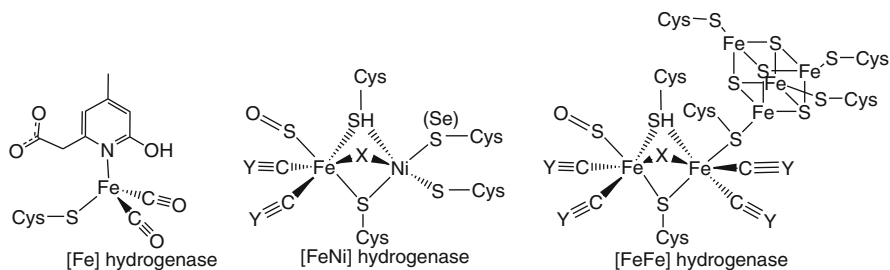


Fig. 11 Hydrogenase active sites. *Y* can be nitrogen, oxygen or sulfur; *X* is most likely sulfur

Vanadium haloperoxidases (VPO) offer a perplexing alternative to heme-containing CPO or MPO. While VPOs also use hydrogen peroxide to activate halides, their active site contain a vanadyl unit already in a high oxidation state—as opposed to CPO and MPO where the role of the hydrogen peroxide was precisely to mount the metal into a higher oxidation state [73–77]. Experimental and DFT data appear to concur on the fact that in VPO the halide is directly activated via a side-on peroxo adduct at vanadium; the protonation states of the ligands, as well as the influence of the protein on the reactivity and spectroscopic properties, have been thoroughly indexed with DFT, TD-DFT, and QM/MM calculations [73–77].

7 Hydrogen/Protons

Hydrogenases catalyze the reversible transformation of molecular hydrogen into protons + electrons; both of the partners in this reaction—but especially the protons—are particularly reluctant to ligate to most biological metal centers. Hydrogenases feature three types of active sites: mononuclear iron, diiron, and binuclear iron–nickel, cf. Fig. 11.

In all cases, unusual ligands are found at the metals—with carbon monoxide or cyanide being the least expected; together with the sulfur-rich ligands, this makes for an unusual coordination environment, where, among others, lower oxidation states at the metal are possible—a clear advantage in the attempt to convince protons to bind to what may otherwise be regarded as an ordinary transition metal in aqueous phase. The small size of the substrate implies serious difficulties in applying some of the most accurate and informative experimental techniques for trapping reaction intermediates—X-ray crystallography and X-ray absorption spectroscopy. The complete details of the mechanisms of hydrogenases are therefore still a matter of investigation. This is thus a field where the reliability of DFT applications is of particular importance; not surprisingly, a significant number of reports have covered the area in recent years [78–91]. The *qualitative* dependence of the results on the size of the model as well as on the functional employed [87], imply a significant need for reconsideration of the performance of the DFT

methods, as different functionals may yield opposing conclusions on the same problem (as also discussed above for e.g., Rieske dioxygenases). In this respect, the progress made by newer parameterized methods, where selective targets are set for performance (e.g., only weak interactions or only (organo)metallic compounds) [3] offers significant hope. Nevertheless, the concept of such target-dependent functionals reignites, in parallel, the interest in true *ab initio* methods—whose performance would be less dependent on the type of chemical system examined.

8 Summary and Outlook

Recent progress in methodology with DFT methods has allowed tackling of increasingly challenging tasks in terms of understanding mechanisms and making predictions on small molecule activation by biological metal centers. The potential practical applications of such investigations cannot be overstated: fields like proton activation for molecular hydrogen production, water splitting, molecular nitrogen activation, or carbon dioxide fixation still have unexplored facets, with computational methods able to provide key input for generating robust economically feasible catalysts. On a different note, species that may be too short lived to be observed experimentally are facile targets for DFT predictions. Also, recent years have seen a focus on allowing predictions on spectroscopic parameters [92–95]. One important component is, however, the responsibility of the researcher to properly assess the predictive value of the DFT results. The times where a single method would be applied on a small set of models, concluding that “theory confirms experiment,” have long passed. We have given here examples of how some of the mainstream functionals can easily lead to opposing conclusions on key aspects of the topic investigated; in our experience these differences between functionals may amount to as high as 30 kcal/mol in terms of spin state preferences, and not much lower for activation barrier differences; these values are much larger than the 1–5 kcal/mol commonly assumed in past years. From this point of view, methodological contributions such as reference [3] will in our opinion prove to be by far among the most useful for the future development of the field.

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The Bond Analysis Techniques (ELF and Maximum Probability Domains) Application to a Family of Models Relevant to Bio-Inorganic Chemistry

Mauro Causà, Maddalena D'Amore, Carmine Garzillo, Francesco Gentile, and Andreas Savin

Abstract Electron Localization Function (ELF) and Maximum Probability Domain (MPD) analyses have been applied to model metal–porphyrins and show compatible and complementary results. ELF basins are quite different from MPDs, but are a necessary starting point for optimizing them. The analyses of the bond between the metal and porphyrin do not show significant differences between non-transition and transition metals. In all the cases considered, we find signatures characteristic of essentially ionic bonds.

Keywords Density functional theory • Electron localization function • Maximum probability domains • Molecular orbitals

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Contents

| | | |
|-----|--|-----|
| 1 | Introduction | 120 |
| 2 | Methods | 123 |
| 2.1 | Algorithms for Managing and Optimizing Maximum Probability Domains | 123 |
| 2.2 | Computational Details | 126 |
| 3 | Results | 127 |
| 3.1 | Bader's "Atoms in molecules" Analysis | 127 |
| 3.2 | MPDs Analysis of Pure Covalent and Ionic Systems | 127 |
| 3.3 | ELF and MPDs Analysis of Metal Porphyrins | 132 |
| 4 | Conclusions and Outlook | 139 |
| | References | 139 |

Abbreviations

| | |
|------|-----------------------------------|
| AIM | Atoms in molecule |
| AO | Atomic orbital |
| DFT | Density functional theory |
| ELF | Electron localization function |
| ELI | Electron localizability indicator |
| LOL | Localized orbital locator |
| MO | Molecular orbital |
| MPDs | Maximum probability domains |

1 Introduction

In the recent years, the traditional discussion of chemical bonding in terms of the canonical orbitals has been challenged [1–3]. In fact, in general SCF theory, the definition of orbitals remains ambiguous [4–6] and several localization procedures have been introduced by suitable unitary transformations of the set of canonical orbitals in order to provide a more rigorous mathematical meaning to chemical concepts such as bonded atoms, reactants, chemical bonds, electron shells, and lone electron pairs [7–9]. Yet, the realization that theoretical partitionings are not unique “casts a shadow of mistrust over certain definitions” [10].

Alternative approaches to the many-electron problem, working in real space rather than in Hilbert space and with the electron density playing the major role, are provided by Bader's “atoms in molecule” [11, 12], which partitions the molecular space into basins associated with each atom and density-functional methods [3, 13]. These latter are based on a modified Kohn–Sham form of the one-electron effective Hamiltonian, differing from the Hartree–Fock operator for the inclusion of a “correlation potential.” In these methods, it is possible to mimic correlated natural orbitals, as eigenvectors of the first-order reduced density operator, directly

from one-electron eigenvalue equations, bypassing the calculation of multi-configurational Möller–Plesset or CI wave functions [14]. Such a localized “Lewis-like” description of bonding and reactivity, which uses a natural bond orbital analysis (associating bonds with localized two-center two-electron wave functions), has recently been expanded to bulk materials and/or periodic surface models [15].

As a matter of fact, there are systems that cannot be easily characterized in terms of localized molecular orbitals, and thus not easily understood using localized bonds. An extreme example is provided by the fluxional behavior of CH_5^+ , where quantum-mechanical fluctuations induce delocalization phenomena and thus fluctuating bonds [16–20].

Most importantly, these systems are amenable to the Electron Localization Function (ELF) method [21]. This is a local measure based on the reduced second-order density matrix, which as pioneered by Lennard-Jones [22] should retain the chemical significance and at the same time reduce the complexity of the information contained in the square of the wave function Ψ . ELF is defined in terms of the excess of local kinetic energy density due to the Pauli exclusion principle, $T(r)$, and the Thomas–Fermi kinetic energy density, $T_h(r)$:

$$\text{ELF} = \left[1 + \left(\frac{T(r)}{T_h(r)} \right)^2 \right]^{-1}. \quad (1)$$

In the seminal paper of Becke and Edgecombe [20] it was proposed that a topological analysis of the ELF [23–25] permitted a more quantitative analysis of the three-dimensional function. The latter is normalized to the interval between 0 and 1, and is large where the Pauli repulsion is small (two electrons with anti-parallel spin are paired), and is small in the regions between electron pairs. To identify a region around a maximum (an attractor), which forms a basin, one can consider all the points in the space with $\text{ELF} \geq \zeta$, which defines the ζ localization domain and with ζ being a positive constant smaller than the value of the maximum. This region in the space can be visualized by showing the iso-surface $\text{ELF} = \zeta$. All points leading to the same attractor belong to the same basin.

The ELF has been extensively applied to a large number of systems and has also been used to quantify chemical concepts like the strength of the hydrogen bond [26] and aromaticity [27, 28]. Moreover, other interesting ELF-like scalar fields have been developed, such as the localized orbital locator (LOL) [29], based on the comparison of the local noninteracting kinetic energy density with that of the uniform electron gas, and the electron localizability indicator (ELI) [30–32], derived directly from the electron pair density without any reference to the uniform electron gas.

In order to get out of the somewhat confusing definition and interpretation of ELF, a more thorough investigation of the competition of kinetic terms $T(r)$ and $T_h(r)$ of Eq. (1), between regions in which the inhomogeneous or the homogeneous behavior dominates, was carried out by Putz [33, 34]. Employing a path integral

Markovian pair conditional probability density with the basic concepts of the catastrophe theory, he succeeded in introducing new Markovian ELF classes which generalize the previous Becke–Edgecombe definition. Going beyond the actual interpretation of ELF as the error in electron localization, this new approach provides a quantum step-function indicating where the electrons are trapped rather than where they have peaks of spatial density.

From a computational point of view, it should be stressed that the computational tool of Francisco et al. [35] results in obtaining the electron number probability distribution functions of an n -electron molecule through an exhaustive partitioning of the real space into arbitrary regions. From the computed probabilities, several magnitudes relevant to chemical bonding theory are obtained, such as average electronic populations and localization/delocalization indices.

Recently, Corminboeuf et al. [36] have compared the ELFs of both the canonical wave functions and electron-localized states with those of ELI and LOL in order to discriminate between enhanced or reduced electron (de)localization within cyclic π -conjugated systems. As a result, the authors say, the simplest LOL function gives a more appealing and intuitive picture of the π -bond, whereas the most popular ELF fails to capture subtle contrasting local electronic properties and suffers from the arbitrariness of the σ/π dissection. Some ELF limitations have been thoroughly discussed in recent years [37] and attempts to search for new tools, capable of providing solutions to still unsolved difficulties in chemical understanding, have been done. A promising methodology has been obtained through a topological analysis by means of Maximum Probability Domains (MPDs) [38–41, 53] and this is discussed in detail below.

In this paper, the chemical bonding of some metal porphyrins is extensively analyzed using both ELF and MPDs, the latter revealing itself to be necessary to gain further chemical information and allowing us to identify and discuss the subtle variations of the electron density with respect to the various macrocycle/metal junctions. Macrocycles such as porphyrins and their analogues [42, 43] have attracted much attention because of their remarkable role in biological photoredox processes and in potentially semiconducting polymers; more recently, they have found applications in the detection and photodynamic therapy of tumors [44]. Therefore, attempts to clearly establish the structure–property relationships in this class of molecules promise to be extremely useful for a better understanding of the mechanisms of those processes. A preferential way to achieve that goal consists in trying to identify the structural features most likely to be responsible for electron donor/acceptor properties and eventually to compare the properties of porphyrin macrocycles to those of other macrocycles with different features, where the features in question are absent. In the present study, for the first time, we explore the performances of the MPDs method when applied to metal porphyrin compounds.

2 Methods

2.1 *Algorithms for Managing and Optimizing Maximum Probability Domains*

The optimization algorithm of the Domains in accordance with MPDs approach has been described in references [26, 38]. An atomic or ELF basin, defined on a grid, or another domain chosen by the user (a sphere, an ellipsoid, a cube, or a previously obtained MPD) can be a first guess for the MPDs.

2.1.1 Calculation of Surface and Integration

First a regular cubic grid of points $G = \{g_i\}$ is established with a given increment dx . The classical value of dx is 0.05 bohrs. A volume $dV = dx^3$ is associated to the points g_i .

Definition of Domains

A subset $D = \{d_i\}$ of G is defined As constituted by grid points belonging to the domain. The number of the d_i points is n_v . The volume of the domain is: $V_1[D] = n_v dV$.

Definition of D for AIM and ELF Basin

The scalar field $s(\mathbf{x})$ (where s is the electronic density for AIM or ELF) is calculated for all the points $\{g_i\} = G$. Afterwards, the set of attractors (maxima) is calculated, first through the use of the numerical gradient of $s(\mathbf{x})$ on the grid G , and next the positions of the attractors are refined using the analytical gradient of s . The g_i points are attributed to each attractor using the numerical gradient. $V_1[D]$ is the most accurate method of integration of the volume for ELF and AIM basins.

2.1.2 Definition of D for MPD

A first guess D can be defined by:

- Using a D set of points taken from AIM or ELF analysis.
- Searching for a set of grid points interior to a guess surface (sphere, ellipse, cube, or a previous MPD surface).

The set D of points d_i is maintained and redefined during the surface optimization for keeping the information about the interior–exterior space of the surface.

2.1.3 Construction of the Surface

Given the set of domain of grid points D , a subset $S = \{s_i\}$ of D is defined as constituting points that have less than six neighbors belonging to D . This is the surface set of grid points. The set S is then made independent of G and D , because the position of s_i may be changed.

2.1.4 Elaboration of the Surface S

The set S is then multiply screened and smoothed (the positions of s_i are a little modified to set triangularization).

2.1.5 Triangularization

The set of points S is triangularized. A set of triangles $T = \{t_i\}$ having corners s_i is defined. The process is a non-Delaunay one to allow for concavities. The triangles are defining walking in the set of nearest neighbors s_i for obtaining locally the most regular triangles. The set $B = \{\mathbf{b}_i\}$ of barycenter of the triangles t_i is defined. The set $N = \{\mathbf{n}_i\}$ of normal vectors to each t_i is defined, pointing toward the external space of the domain: the set of domain grid points D is used for defining the versor of \mathbf{n}_i . The information about the domain is then represented by D , B , and N .

2.1.6 Variation and Optimization of the Surface

The set B is changed during optimization, \mathbf{b}_i are moved along \mathbf{n}_i . The shape derivative is calculated in the points \mathbf{b}_i and helps defining the step and sign of variation of \mathbf{b}_i along \mathbf{n}_i .

For optimization, the barycenter of the triangles (of the surface S) is moved along the normals. The displacements are proportional to the shape derivatives computed at the barycenter [40]. They are larger at the start of the optimization, smaller towards the end. During the optimization process, certain regions of domain can collapse to a surface, or even points. These low-coordinated grid points are then eliminated.

2.1.7 Redefinition of the Surface

After a controlled number of variations of B , the set B is no more coherent with D . So the domain and the surface are redefined: a new set of grid points $D = \{d_i\}$ is defined internally to the surface represented by B and N . Then, the sets S , T , B , and N are redefined on the base of the newly defined D .

2.1.8 Integration Method for Probability Evaluation

The overlap matrix $S[D] = \{S_{mn}\}$ between AO m and n is calculated; s_{mn} is the overlap limited to the domain D , whereas S_{mn} is the integral extended to all the space. A Becke atomic partition method is adopted for defining the quadrature set of points $X = \{x_i\}$ and the weights w_i :

$$S_{mn} = \chi_m(\mathbf{x}_i)\chi_n(\mathbf{x}_i)w_i$$

$$s_{mn} = \chi_m(\mathbf{x}_i)\chi_n(\mathbf{x}_i)w_i f(\mathbf{x}_i).$$

$f(\mathbf{x}_i)$ is the domain weight function for limiting the overlap to D , χ_m is the m -th atomic orbital of the molecule model or of the cluster chosen as a representation of the periodic system in terms of a set of local Wannier functions. The volume of the domain D can be calculated as:

$$V_2[D] = \sum_i w_i f(\mathbf{x}_i).$$

The average number of electrons in the domain $\langle N \rangle[D]$ can be calculated in two ways:

1.

$$\langle N \rangle[D] = \sum_{n=0}^{n_e} n p_n$$

where p_n are the probabilities of finding exactly n electrons in the domain D and n_e is the number of electrons in the molecule or cluster. The set of probabilities $P = \{p_n\}$ is a functional of the domain overlap matrix $S[D]$.

2.

$$\langle N \rangle[D] = \sum_i w_i \rho(x_i)$$

where $\rho(x_i)$ is the electronic density calculated in grid point x_i . This method is used only for checking the accuracy of p_i .

There are two methods for calculating the domain function $f(\mathbf{x}_i)$

1. Given the quadrature point \mathbf{x}_i , the closest point of the grid g_i is found: $f(\mathbf{x}_i)$ is equal to 1 if g_i belongs to the set of domain points D ; otherwise $f(\mathbf{x}_i)$ is 0.
2. Given the quadrature point \mathbf{x}_i , the closest barycenter b_i of the surface triangles is found. The scalar product ps is calculated: $ps = (x_i - b_i) \cdot n_i$. A "Fermi type" step function is calculated: $f(x_i) = \frac{1}{1 - \exp[sf \cdot ps]}$ where sf has been optimized to a value of 50 atomic units as a compromise between a sharp step and the accuracy of the integration accuracy.

2.1.9 The Adopted Integration Methods

p_n and $\langle N \rangle[D]$ for MP Domains

The probabilities P are calculated on the basis of $S[D]$ evaluated by the method described in Sect. 2.1.8. The $f(\mathbf{x}_i)$ domain function is calculated by method 2 described above. $\langle N \rangle[D]$ is calculated using P . The volume V of MPD is calculated using the method $V_2[D]$, described in Sect. 2.1.8.

AIM and ELF Analysis

The probabilities P are evaluated on the base of $S[D]$ matrix, calculated using the method reported in Sect. 2.1.8; in this case, the $f(\mathbf{x}_i)$ domain function is calculated using method 1 described in Sect. 2.1.8. $\langle N \rangle[D]$ is calculated using P . The volume V of AIM and ELF basins are computed using the method $V_1[D]$ as reported in section “Definition of Domains.”

2.2 Computational Details

As starting point for our calculations, we have used the following PDB files for structures: Fe-protoporphyrin PDB Code: HEM, Co-protoporphyrin PDB Code: COH, Mg-protoporphyrin PDB Code: HEG, Ni-protoporphyrin PDB Code: HNI, Zn-protoporphyrin PDB Code: ZNH [45] and Cambridge Crystallographic Data Centre for Ca-protoporphyrin [46].

All the calculations have been performed within the Density Functional Hybrid Approximation, using the B3LYP parametrization [47]. A double zeta plus polarization Gaussian Type Orbital basis set has been adopted (6-31g(dp) [48]).

A 3D extension of MPDs analysis, implemented in a development version of CRYSTAL code [49], able to deal with molecules and crystalline systems in the Hartree–Fock and DFT approximation, has been adopted. The MPD code can study closed and open shell systems.

In MPDs analysis, our first objective has been to identify spatial subsystems $D \in R^3$ in the metal porphyrin compounds by specifying a given electron number n ; for example, we considered +2 cationic domains specifying the number of electrons diminished by two (Fe ion with 24 electrons).

We searched two electrons domains in the metal–ligand regions of the macrocycle.

The next choice is the arbitrary definition of a volume D where we are going to find a given number of electrons n with a certain probability p ; in fact, according to quantum mechanics, for a given state Ψ , we can find a certain number of electrons n in D with a certain probability. We computed the probability of finding exactly n

electrons in D for one Slater determinant wave functions and define the regions of space for which the probability to find n electrons becomes a maximum; to find out the local maxima for a certain probability with respect to the shape of D , the optimization of D for a given n (indicated as D_n) has been performed by making use of shape derivatives.

As in all distributions, information about the probability distribution $p_{\downarrow}(0)(D)$, $[(p)]_{\downarrow}(1)(D)$, $[(p)]_{\downarrow}(2)(D)$, \dots , $p_{\downarrow}(N)(D)$ is obtained by means of some parameters, in particular the mean $\langle N \rangle$ has been computed for this system.

For an N electrons system, the average number of electrons in D is given by:

$$\langle N \rangle [D] = \sum_{n=0}^N n p_n(D). \quad (2)$$

We have obtained it by integrating the electron density $\rho(r)$ over the domain D as indicated in Eq. (3), representing the population of:

$$\int_D \rho(r) d^3(r) = \int_D \langle \psi | \hat{\rho}(r) | \psi \rangle d^3(r) = \left\langle \psi \left| \int_D \hat{\rho}(r) d^3(r) \right| \psi \right\rangle. \quad (3)$$

3 Results

3.1 Bader's "Atoms in molecules" Analysis

The reference analysis of the electronic structure of molecules and crystals, as coming from experiment or calculation, is due to Richard Bader [11], who showed how chemical concepts related to Atoms can be recognized from a pure physical observable, like the electronic density, its derivatives, and its critical points. In the present study, Bader's analysis has been applied to porphyrin and metal porphyrins, mainly for calculating atomic charges. Moreover, we have explored how the electronic density analysis describes the metal–ligand bond (Figs. 1, 2 and 3; Table 1).

3.2 MPDs Analysis of Pure Covalent and Ionic Systems

For illustrating the MPDs analyses to the metal–ligand bonding in metal-protoporphyrins, we report some significant results of MPDs analysis of extreme ionic and covalent situations. We refer to crystalline magnesium oxide as a prototype of pure ionic situation and crystalline silicon, as a pure covalent situation. We select

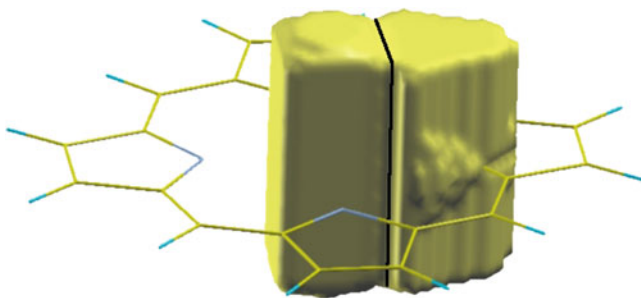


Fig. 1 Atom in molecules (AIM) basins, corresponding to the metal (*center*) and to the ligand (*right*), for the Mg porphyrin. In principle, perpendicular to the molecular plane, the AIM basins extend up to infinite. All the plots of molecules, basins and domains are obtained by XcrysDen computer program [54]

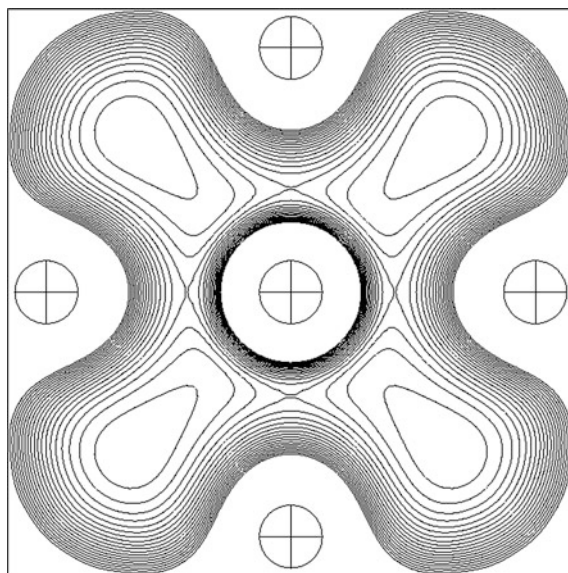


Fig. 2 Electronic charge density map for Mg porphyrin along the molecular plane. The atomic positions are marked by *crossed circles*. The metal is in the center of the map, while the four N ligands are the remaining *circles*. Neighbor iso-lines correspond to values that differ by 0.01 atomic units (electrons per cubic bohr). The lower value iso-lines (0.01 atomic units) lie in the diagonals of the map, out of the metal–ligand directions. The highest reported value iso-lines (0.20 atomic units) lie in the direction of the nuclei, both metal and N. Close to the middle of the metal–ligand segment, a saddle point lies

crystalline situation because the extremely ionic bond exists only in the solid state. Even for the perfect homo-nuclear covalent situation, the crystalline status is a perfect paradigm.

Figures 4, 5 and 6 show that ELF reaches very high values in valence regions; it means that the electronic structure is very localized both in perfect ionic and perfect covalent systems; in terms of ELF, the main difference between ionic and covalent

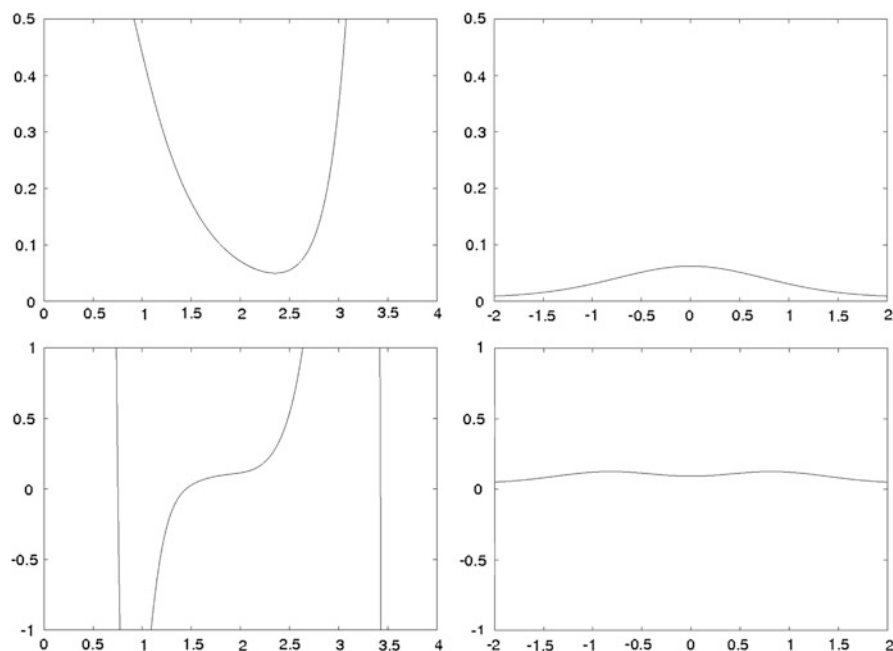


Fig. 3 Electronic charge density (*upper plots*) and Laplacian of the charge density profiles (*lower plots*). *Left plots* refer to the metal–ligand direction; cationic position is at the origin while N position is 3.88 bohrs, close to the end of the axis. *Right plots* refer to a direction perpendicular to the “bond” direction, lying on the molecular plane, passing through the minimum of the electronic density (*upper left*). The *upper pictures* show the $(3,-1)$ critical point in the charge density: a saddle point in a scalar field has two negative and one positive hessian eigenvalues, so is defined $(3,-1)$. The *bottom plots* show that the Laplacian of the charge density is positive in the critical point

Table 1 Atomic net charges for metal porphyrins (phy) following Bader’s analysis

| | $Q(M)$ | $V(M)$ | $Q(L)$ | $V(L)$ |
|--------------------------------------|--------|--------|--------|--------|
| Be phy | +1.32 | 81.2 | −1.25 | 169.1 |
| Mg phy | +1.45 | 136.9 | −1.24 | 163.6 |
| Ca phy | +1.47 | 205.7 | −1.17 | 154.3 |
| Fe phy | +1.39 | 158.9 | −0.94 | 152.7 |
| Co phy | +1.13 | 158.9 | −0.89 | 151.6 |
| Ni phy | +0.99 | 158.1 | −1.02 | 153.7 |
| Zn phy | +1.26 | 158.0 | −0.87 | 155.6 |
| Fe phy (CO) ₂ | +1.43 | 46.5 | −0.88 | 92.0 |
| [Fe(CO) ₆] ²⁺ | +1.03 | 97.5 | +1.63 | 120.7 |

Basin charges and volumes are reported for the metal (M) and the nitrogen ligand (L); L is Nitrogen in all cases but in the last two lines, where the atomic charge of Carbon in CO is reported. The absolute values of the volumes has no meaning for the square planar coordinated molecules (the first seven systems), because the basins will extend to infinite perpendicularly to the molecular plane. In those cases, the reported volumes refer to basins truncated to a distance of 6 bohrs from the molecular plane

Fig. 4 Electron localization function (ELF) plot for crystalline magnesium oxide. The iso-surface corresponding to $\text{ELF} = 0.3$ is reported. A magnesium ion is at the center of the cube and it appears like a medium-sized spheroid. Oxygen ions appear like large quasi cubes. The *small spheres* represent the oxygen cores

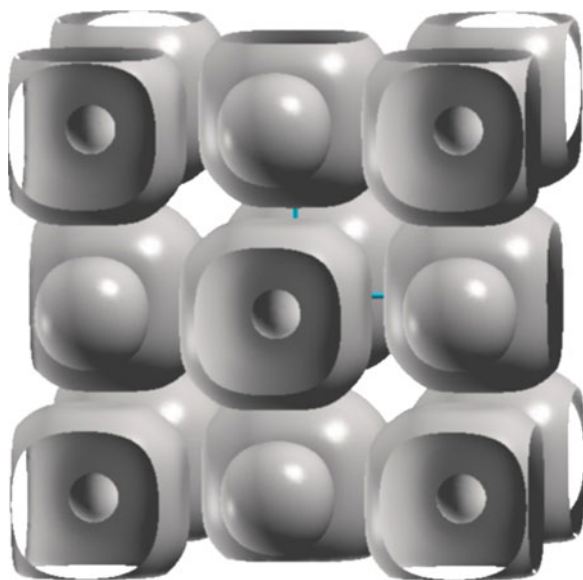
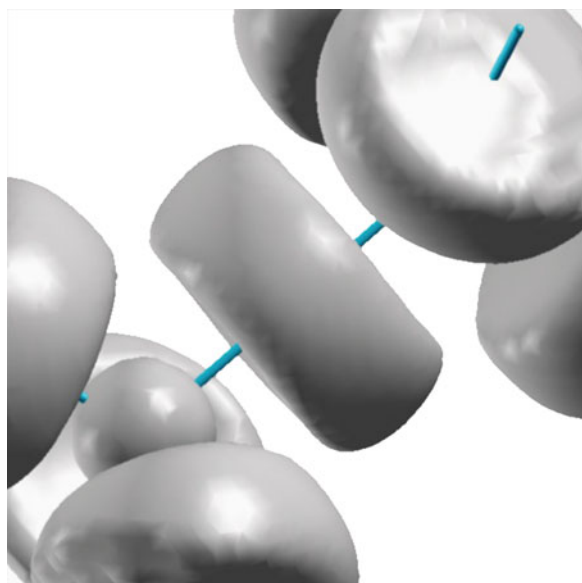


Fig. 5 ELF iso-surface, corresponding to the value of 0.66, for crystalline silicon. The *little spheres* underline the silicon cores; the large cylinders, coaxial with the Si–Si direction, underline the Si–Si covalent bond



lies out of the bond regions: in ionics, ELF drops to very low values, below 0.05, while in covalents, it remains higher, amounting at about 0.1 in bond direction and it is even higher in “out of bond” directions. As a consequence, the ELF basins remain well separated even at low level, like the 0.3 iso surface shown in Fig. 4. In covalent situations, the chemical basins which correspond to cores and covalent bonds easily melt together: the covalent picture is clearly restituted by 0.6 iso-surface reported for silicon in Fig. 5.

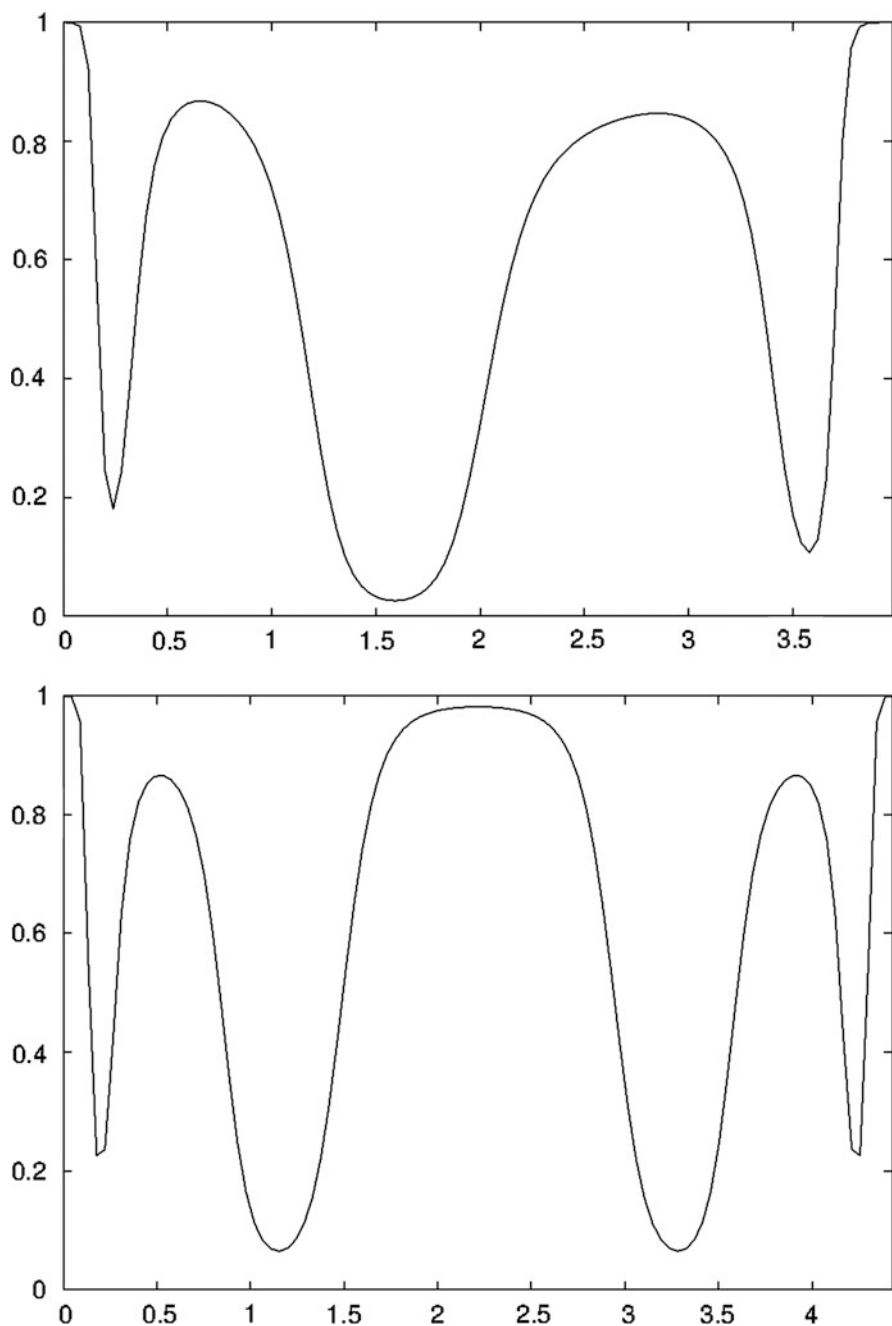


Fig. 6 ELF profile between the closest neighbors of crystalline magnesium oxide (*top*) and silicon (*bottom*). In both cases, the atomic core-valence shell structure is evident. In the MgO, *top* plot Mg position is on the *extreme left* whereas Oxygen is on the *extreme right*. For Si two symmetry equivalent Si nuclei are located at the extremes of *horizontal axis*. ELF reaches maximum values that can be considered as maximum pairing of electrons in the 1 s cores and in the middle of Si-Si segment that we commonly attribute to a Lewis pair covalent bond

Fig. 7 Cationic (*left*) and anionic (*right*) maximum probability domains for crystalline MgO. Both the domains are optimized by maximizing the probability of containing ten electrons

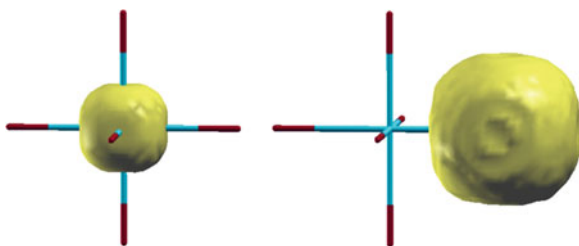
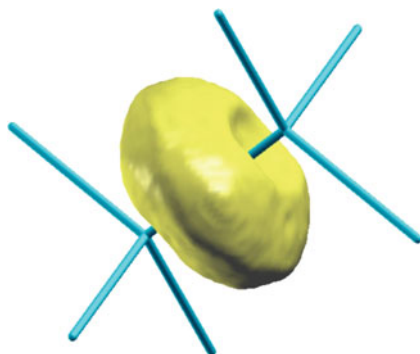


Fig. 8 Two-electron MPD for Crystalline silicon, representing the Si–Si covalent bond



The chemically important Maximum Probability Domains for the same systems are shown in Figs. 7 and 8. For magnesium oxide, the cation and anion 10 electrons MPD have high probability: 0.85 and 0.57 for cation and anion, respectively; the volumes of two ions are very different, 19.4 and 90.5 cubic bohrs; the MPD average net charges are, respectively, +1.85 and -1.85 and these are very close to the formal ones. The two-electron MPD reported in Fig. 8 has a probability equal to 0.42, a volume of 56.0 cubic bohrs, and an average population of 2.04 electrons.

In all our previous MPD studies, the probability lies in the range (0.30–0.45) for covalent bonds, whereas the probability for ionic MPD is higher in fact it amounts to (0.70–0.90) for cations and to (0.5–0.7) for anions.

3.3 *ELF and MPDs Analysis of Metal Porphyrins*

In the present study, we consider the bond analysis in metal porphyrins. In the following discussion, the porphyrin will be often indicated as “phy.” We apply Electron Localization Function (ELF) analysis and Maximum Probability Domains (MPDs) analysis for understanding the eventual differences between transition metal and non-transition metals inserted into the porphyrin ring. As an example of non-transition metal, we consider Be, Mg, and Zn (a d^{10} metal, so d orbitals can be considered core-like). For transition metals, we consider Fe, Co, and Ni. Due to the difficulties in finding experimental examples of pure square planar coordination

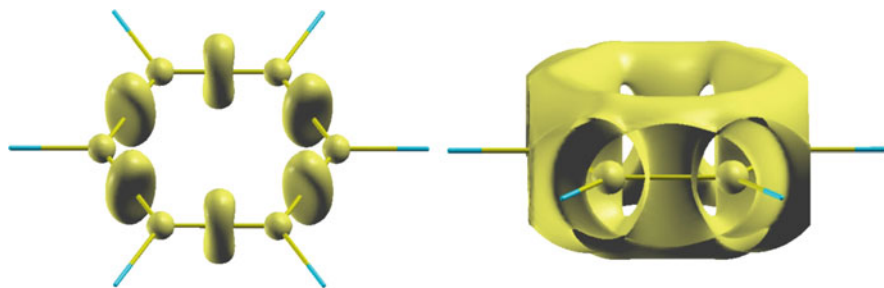


Fig. 9 Electronic localization function (ELF) for benzene. *Left*: ELF = 0.85 iso-surface. *Right*: ELF = 0.66 iso-surfaces. The right iso-surface is cut on the front side for showing the complex topology: the Carbon core basins sharply separate from the scalar ELF field. The aromatic ELF basin is clearly defined. Both in the high or lower ELF value iso-surface, no separation between Sigma and Pi basins can be found, which is a characteristic of delocalized bonds, similar to that shown in Fig. 11. All the molecular plots reporting ELF iso-surfaces or MPD are performed using the Xcrysden program [44]

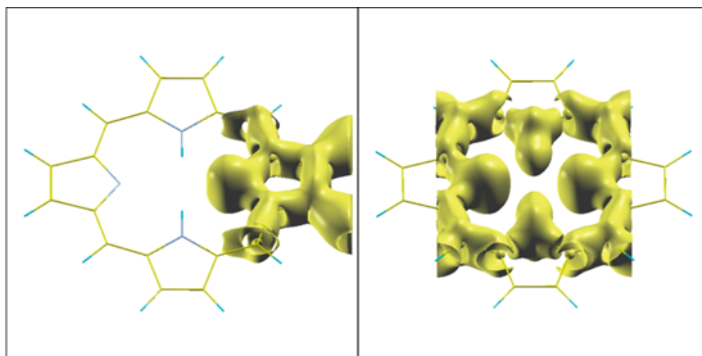


Fig. 10 ELF for porphyrin. In both parts, portions of ELF = 0.66 iso-surface are shown to highlight different features. The ELF shows the same delocalization features as benzene in Fig. 9. The nitrogen lone pairs show a slightly larger localization basin than NH bonds. NH induces a small interruption in the delocalized ring between N and neighbor C(H); this small interruption disappears in deprotonated porphyrin and in metal-porphyrin

for Fe, because the d^6 electronic structure for Fe(II) does not favor this coordination geometry, we also consider the penta-coordinate Fe porphyrin CO and the quasi octahedral Feporphyrin $(CO)_2$ examples. As a classical reference, we also study with the same methods and approximations $[Fe(CO)_6]^{2+}$, as a prototype for the CO-metal bond.

In Figs. 9 and 10, we show the bond delocalization in benzene and porphyrin. In Fig. 9, left, the iso-surface corresponding to ELF equal to 0.85 is presented. At high values, the maxima of ELF relative to atom cores and bonds are shown. Lowering the ELF value to 0.66, the iso-surfaces melt together in a classical hexagonal domain, as shown in Fig. 9, right. Due to the relatively high value of ELF that

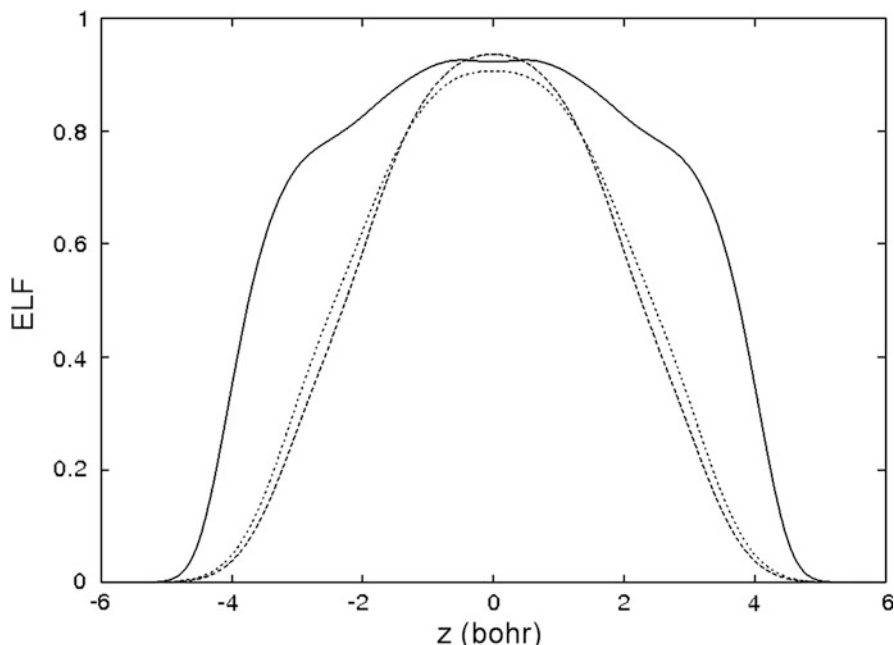


Fig. 11 Vertical ELF profiles for ethylene, benzene, and porphyrin. The z coordinate is perpendicular to the molecular plane. The $z = 0$ value corresponds to the middle of the C–C bond. The *continuous line* refers to ethylene, the *long-dotted one* to benzene and the *short dotted one* refers to porphyrin (the external C–C bond of the ring has been selected)

corresponds to this “aromatic” topology, we interpret as the existence of a wide spatial region where the electrons localize (are paired), without the possibility of distinguishing them in separated pairs. The valence electrons n in the porphyrin ring shows an analogous behavior as the one shown in Fig. 10, where the iso-surface corresponding to ELF equal to 0.66 is shown.

From the comparison with ELF of benzene, we can assume a perfect aromaticity of the porphyrin ring. Further, in Fig. 11 we analyze vertical profiles of ELF.

The ethylene ELF profile is on average higher than the others because the bond is localized; the π bond local maxima are clearly distinguishable. Out of the molecular plane, in the benzene and porphyrin case, ELF is rapidly decaying and only a central maximum is present: we interpret this behavior of benzene and porphyrin as an indication of electron delocalization that corresponds to an objective difficulty in distinguishing between σ and π bonds.

It is noteworthy that Coulson et al. [50], calculating the electron density in benzene in the plane 0.35 above the molecular plane, found that there is only a very small region directly above each of the six carbon nuclei where the charge density of the π electrons is as great as that of the σ electrons, meaning that the π electron charge cloud does overlap considerably with that of σ electrons. Thus “this calculation threw a considerable doubt on the validity of the σ – π separation” [51],

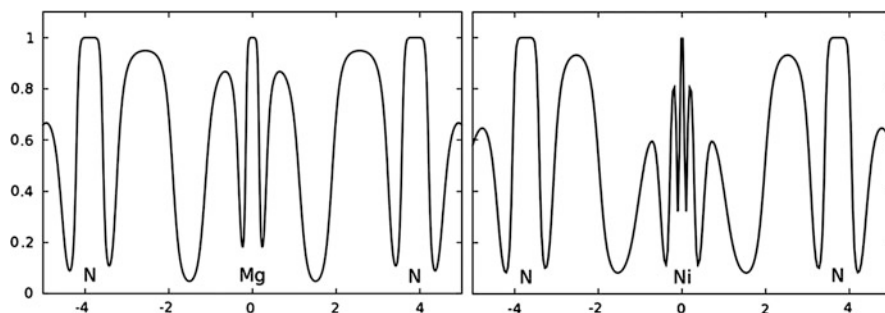


Fig. 12 ELF profiles along the N–M–N segments. *Left plot* refers to Mg phthalocyanine (left); *right plot* refers to Ni phthalocyanine (right). *From left to right* of each plot, the following ELF localization domains are distinguishable: N core, N lone pair, Metal outer and inner core, N lone pair and N core. The ELF analysis provides the same picture for the non-transition and the transition metal compound

becoming also more doubtful for larger molecules when σ and π configuration interaction would become more important with the increasing number of atoms.

The ELF analysis of metal porphyrins as reported in Figs. 12, 13, and 14 shows no relevant difference between non-transition and transition metals. There are low values in ELF in the space region between metal and ligands, both nitrogen, or CO. The ELF profiles reported in Fig. 12 show very clearly the localization of the electronic structure in the ligand–metal segment: The higher values of ELF are in the N and Metal 1s cores. Then are clearly distinguishable the 2sp outer core of Mg^{2+} ion and the 2sp and 3spd outer cores of Ni^{2+} . The nitrogen “lone pairs” also show high value of localization. The low value of ELF in d-electron systems has been already remarked upon [29]. At moderately high ELF level (0.66), the porphyrin ring and the central metal regions remain well separated, as shown in Fig. 4. In addition, at the low 0.3 ELF level, the metal and ligand domains are still separated, as shown in Figs. 12 and 13. The ELF level equal to 0.3 is usually adopted for analyzing very ionic systems, like alkaline halides, where the cationic and anionic ELF domains stay distinct also at 0.05 ELF level. In the metal–porphyrin complex, metal and ligand ELF basin remain separated for ELF values larger than 0.12. The same behavior can be noted in $[\text{FeCO}_6]^{2+}$, where the metal and ligand (C of CO molecule) ELF basins melt at 0.17 level. From the ELF analysis, as shown in Figs. 12 and 13, there is no difference between non-transition and transition metal porphyrins.

In Fig. 15 two Maximum Probability Domains for metal porphyrins are shown: the central cation MPD and the two-electron domain in the metal–ligand space region. These MPDs are well defined and do not overlap. Following previous applications of MPD analyses [33–36], the non-overlap of MPD is a clue of a good chemical hypothesis. Figure 15 shows that there are no significant topological differences between non-transition and transition metals. This confirms the ELF analysis. Figure 16 shows that in the case of a metal–ligand bond, there is a small difference between the ELF basin and the Maximum Probability Domain. On the other hand, the ELF basin is an essential starting point for optimizing the MPD.

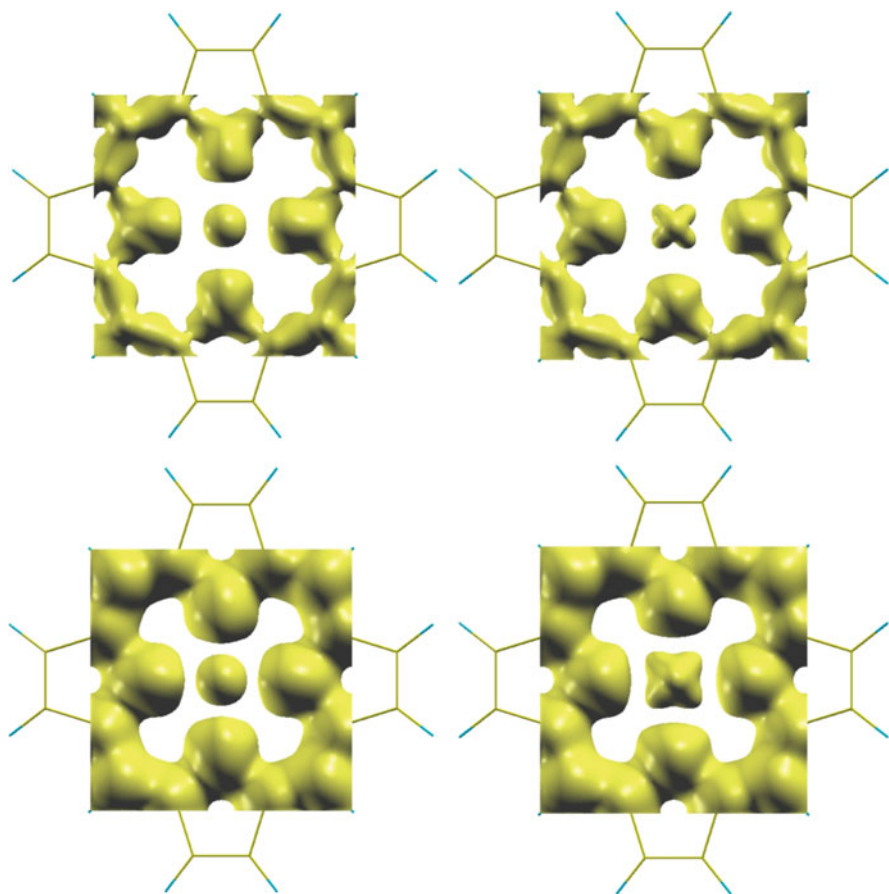


Fig. 13 ELF iso-surfaces: *upper row* corresponds to $\text{ELF} = 0.66$, *lower row* corresponds to $\text{ELF} = 0.30$; *left column* refer to Mg phy, *right column* refers to Ni phy

The numerical data relative to the MPD optimization are reported in Table 2. The trend of $p(\text{M}^{2+})$, the maximal probability of the +2 cationic domains, and the trend of their volumes, is chemically understandable. $p(\text{M}^{2+})$ monotonically decreases in the alkaline-earth series. Almost iso-electronic Ca phy and Zn phy have the same values. The first transition series Fe phy to Ni phy has almost the same cationic probability $p(\text{M}^{2+})$. The cationic probability $p(\text{M}^{2+})$ is the only difference between non-transition and transition metal porphyrins. The non-transition cations, including the $3d^{10} \text{Zn}^{2+}$ in Zn phy model, are obviously more defined physical systems. The average electronic population $\langle N \rangle$ is always close to the formal number of electrons.

The range of variation of the maximal two-electron probability of metal–ligand domain $P(2, \text{M-L})$ is very small, spanning the interval (0.36–0.41) for porphyrins: a little lower for metal–CO domains where $P(2, \text{M-L})$ is about 0.32. The average

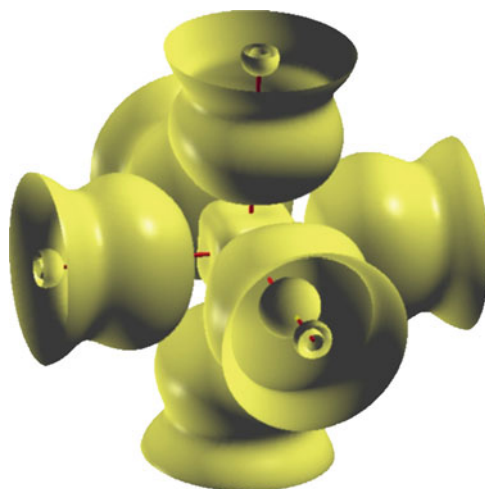


Fig. 14 Iso-surface for $\text{ELF} = 0.30$ of $[\text{Fe}(\text{CO})_6]^{2+}$

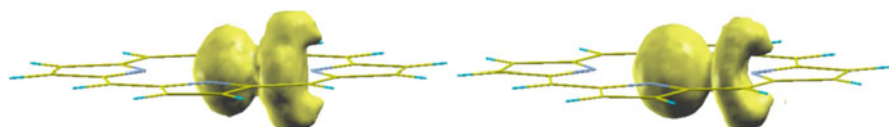


Fig. 15 Metal cation MPD and 2-electron metal–ligand MPDs for Mg phy (*left*) and Ni phy (*right*)

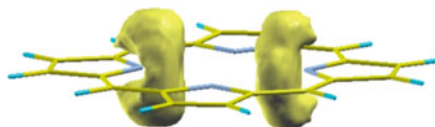


Fig. 16 Difference between ELF Basin (*left*) and MPD domain (*right*) for the metal–ligand bond in Mg-phy. The average number of electrons $\langle N \rangle$ is, respectively, 2.5 for ELF basin and 2.0 for MPD

number of electrons $\langle N \rangle(\text{M-L})$ is every time close to 2.0. The volumes are very similar, about 20 cubic bohr, but in the Mg phy model the volume is 40.5 cubic bohr. In Fig. 16, we show the ELF M–L basin and the MPD, obtained optimizing $P(2, \text{M-L})$ starting from the ELF basin; even if topologically equivalent, ELF basin and MPD have a very different dimension in fact: for Mg phy the volume goes from 23.5 to 40.5 cubic bohr upon optimization, whereas $P(2, \text{M-L})$ goes from 0.23 to 0.34.

Table 3 reports some energetic and structural data. The formation energy of the metal porphyrin models is defined as the energy of the reaction:

Table 2 Numerical results about maximum probability domains in metal–porphyrins

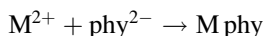
| | $p(M^{2+})$ | $V(M^{2+})$ | $\langle N \rangle(M^{2+})$ | $P(2,M-L)$ | $V(M-L)$ | $\langle N \rangle(M-L)$ |
|--------------------------------------|-------------|-------------|-----------------------------|------------|-----------|--------------------------|
| Be phy | 0.88 | 4.7 | 2.0 | 0.39 | 24.1 | 2.0 |
| Mg phy | 0.77 | 21.6 | 10.1 | 0.40 | 26.3 | 2.0 |
| Ca phy | 0.55 | 42.6 | 17.9 | 0.41 | 22.6 | 2.0 |
| Fe phy | 0.39 | 34.8 | 24.1 | 0.36 | 20.3 | 1.9 |
| Co phy | 0.36 | 34.4 | 25.2 | 0.36 | 19.5 | 1.9 |
| Ni phy | 0.35 | 32.3 | 26.2 | 0.36 | 19.4 | 1.9 |
| Zn phy | 0.50 | 33.6 | 28.1 | 0.37 | 20.7 | 1.9 |
| Fe phy (CO) ₂ | 0.29 | 31.3 | 24.0 | 0.36,0.38 | 19.7,25.6 | 1.9,1.9 |
| [Fe(CO) ₆] ²⁺ | 0.27 | 30.9 | 24.0 | 0.32 | 17.1 | 1.9 |

$p(M^{2+})$ is the maximal probability of a domain centered in the metallic position, relative to a number of electron equal to the atomic number minus two, that is exactly the situation of a +2 ion; V is the volume of the domain in atomic units (cubic bohr); $\langle N \rangle$ is the average number of electrons in the domain, that is the integral value of the electronic density; $P(2,M-L)$ is the maximal probability having two electron in a domain located in the space between the M^{2+} ion and the ligand L core. In metal porphyrins L is Nitrogen. In [Fe(CO)₆]²⁺ L is Carbon of CO molecule. In Fe phy (CO)₂, the two values refer to N and C respectively. $V(M-L)$ and $\langle N \rangle(M-L)$ are volumes and average numbers of electrons of M–L domains

Table 3 Relevant geometric parameters and formation energy of metal–porphyrins

| | Formation energy (Kcal/mol) | M–L distance (Angstrom) |
|--------------------------------------|-----------------------------|-------------------------|
| Q phy | –843.0 | |
| Be phy | –824.7 | 1.94 |
| Mg phy | –704.1 | 2.05 |
| Ca phy | –573.8 | 2.19 |
| Fe phy | –703.9 | 2.00 |
| Co phy | –805.8 | 1.98 |
| Ni phy | –803.0 | 1.96 |
| Zn phy | –726.7 | 2.05 |
| Fe phy (CO) ₂ | –763.4 | 2.02, 1.86 |
| [Fe(CO) ₆] ²⁺ | –259.3 | 1.93 |

The formation energy refers to the reaction: $M^{2+} + \text{phy}^{2-} \rightarrow \text{Mphy}$. The M–L bond distance refers to N ligand in all cases, but for [Fe(CO)₆]²⁺ and for the second value for Fe phy (CO)₂, where it refers to M–C. The Q phy compound has the equilibrium geometry of Be phy, but the central ion is substituted by a +2 point charge



where M^{2+} and phy^{2-} are, respectively, the cation and the de-protonated porphyrin. This model reaction is highly exoergonic. As a reference, we calculated the formation energy of the complex Q phy, having a +2 point charge in the cation position. We adopted for Q phy the equilibrium geometry of Be phy, the complex with the smallest cation. The formation energy of Q phy clearly depends on the geometry of the porphyrin ring: we calculated it adopting the equilibrium geometry of all the considered complexes, and we obtained a strict correlation between the

formation energy of Q phy and M phy. The extremes of the set of Q phy formation energies are -843 and -731 kcal/mol for the Be phy and Ca phy geometry, respectively, that corresponds to -825 and -574 kcal/mol, that are the extremes of the set of M phy formation energies, for the same compounds. This feature underlines the role of electrostatic in determining M-phy bonds. The formation energy monotonically decreases in absolute value in the series Q-Be-Mg-Ca, as the M-L distance increases, and the dimension of the cation increases (volume of the M^{2+} MPD, reported in Table 2). This trend indicates a strong electrostatic character in the metal porphyrin bond for non-transition metal models. In the model containing first transition row metals, the formation energy is in the same range. Between models containing first transition row metals, the formation energy relative to Fe is lower than in Co phy and Ni phy, perhaps this could be attributed to the disadvantageous square planar coordination of the $3d^6$ Fe^{2+} cation. The quasi octahedral Fe phy $(CO)_2$ shows additivity between Fe-N bond energy and Fe CO bond energy, as calculated for $[Fe(CO)_6]^{2+}$. The metal-ligand equilibrium distances anti-correlate with formation energies.

4 Conclusions and Outlook

We have analyzed the metal-ligand bond in metal porphyrin models, using Electron Localization Function (ELF) and Maximum Probability Domains Analysis (MPDs). We do not find any relevant difference between non-transition and transition metal models. Our MPDs analysis shows that the metal-ligand bond is essentially ionic for all models, similar to the “electride” (charge separated structures) behavior of metal ions within cages [52]. In our MPDs analysis, we do not find clues of stabilization of bonds due to covalent interactions between the central metals and the porphyrin ring both in non-transition and in transition metal porphyrins.

On the basis of results collected hitherto, the Maximum Probability Domains Analysis has been shown as a valuable tool for performing further analysis of bonds in other coordination compounds, without biases coming from historical concepts.

Acknowledgments We, as all the researchers in the field of physical interpretation of chemical bond, are deeply indebted to Richard Bader for his pioneering work [11].

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Biological Activity and Toxicity: A Conceptual DFT Approach

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Abstract Quantitative structure–activity relationship (QSAR) models are generated for biological activity and toxicity in terms of global and local reactivity descriptors within a conceptual density functional theory framework. Possible anticancer activity of two new metal–borane clusters is analyzed.

Keywords Biological activity • Conceptual DFT • Structure–activity relationships • Toxicity

Contents

| | | |
|-----|--|-----|
| 1 | Introduction | 145 |
| 2 | Quantitative Structure–Activity/Structure–Property-Based Methods | 146 |
| 2.1 | Role of Different Theoretical Methods Towards Constructing Effective QSAR Models | 149 |
| 3 | Conceptual Density Functional Theory | 151 |
| 3.1 | Computational Details | 160 |
| 3.2 | Applications of Conceptual DFT in QSAR/QSTR | 160 |
| 3.3 | Application of Conceptual DFT in Designing Anticancer Drugs | 163 |
| 4 | Conclusions | 169 |
| | References | 170 |

List of Abbreviations

| | |
|-----------|---|
| 3D-MEDNEs | Three-dimensional Markovian electron delocalization negentropies |
| 3D-QSAR | Three-dimensional quantitative structure activity relationship |
| ADME | Allied toxicological physicochemical and absorption Disposition Metabolism and Excretion |
| AHH | Aryl hydrocarbon hydroxylase |
| Amax | Maximum acceptor superdelocalizability |
| ArHs | Aryl-hydrocarbon receptors |
| CDFT | Conceptual density functional theory |
| CoMFA | Comparative molecular field analysis |
| CPs | Chlorophenols |
| DFT | Density functional theory |
| DNA | Deoxyribonucleic acid |
| EPR | Electron paramagnetic resonance |
| ESIP | Element specific influence parameter |
| HCV | Hepatitis C Virus |
| HIV | Human Immunodeficiency Virus |
| HPA | Hirschfield population analysis |
| MD | Molecular dynamics |
| MEP | Minimum electrophilicity principle |
| MMP | Minimum magnetizability principle |
| MPA | Mulliken population analysis |
| MPP | Minimum polarizability principle |
| MTD | Minimum topological difference |
| NICS | Nucleus independent chemical shift |
| NMR | Nuclear magnetic resonance |
| NPA | Natural population analysis |
| NQR | Nuclear quadruple resonance |
| PCA | Principal component analysis |
| PCBs | Polychlorinated biphenyls |
| PCDFs | Polychlorinated dibenzofurans |
| PMH | Principle of maximum hardness |
| QC | Quantum chemistry |
| QM/MM | Quantum mechanical/molecular mechanics |
| QSAR | Quantitative structure–activity relationship |
| QSPR | Quantitative structure–property relationship |
| QSRR | Quantitative structure–retention relationships |
| QSTR | Quantitative structure–toxicity relationship |
| QTMS | Quantum topological molecular similarity |
| SAR | Structure–activity relationship |
| SCC-DFTB | Self-consistent charge density functional tight binding |
| SCF-MO | Self-consistent field molecular orbital |
| STR | Structure toxicity relationship |
| TQSI | Topological quantum similarity index |
| VSPM | Variable selection and modeling method based on the prediction |

1 Introduction

Chemists, the experimentalists, or theoreticians although sharing a different world of knowledge are very much aware of the fact that our mother earth is plundered with millions of molecules that may be segregated into two broad classes, namely organic and inorganic. In order to study the chemical behavior of molecules in such large numbers, an effective correlation technique needs to be built up, which can further categorize them into several classes based on closely related structures or properties. Although both the theorists and experimentalists share the same motivation towards classifying molecules into various categories, leading to a better understanding of reactivity patterns, yet they share a somewhat different platform. While the experimental chemists carry out reactions on a large number of molecules to set up a correlation, the theoreticians at the same time use mathematical algorithms to obtain powerful regression models, which usually help to develop effective structure–activity or structure–property relationships. Nevertheless, the development of fruitful activity or property relationships related to changes in molecular structure is what we need in order to predict the reactivity of an unknown compound. Moreover, hundreds and thousands of reactions are attempted and subsequently published, which eventually enrich the chemical database every day. Such increasing data bank needs to be managed skillfully to bring out a logistic correlation among analogous chemical moieties with a view of acquiring superior prediction towards molecular behavior. In this regard, the ideas developed by both experimental and theoretical chemists must be considered together to offer a multidisciplinary approach in dealing with and to envisage chemical reactivity. Again, the various organic and inorganic molecules upon consumption in any form (diet or medicine) are found to affect the general metabolism in human beings and other bio-organisms, which may even lead to toxicity upon an overdose. The mode of reactivity of the several compounds on the living organisms can be nicely predicted in a quantitative approach through suitable structure–activity relationship (SAR) or structure–toxicity relationship (STR) models that are built from chemical intuition. Once a meaningful rationale is established, the structure and property of an unknown molecule can be predicted from this standardized algorithm. Thus, the extent of the lethal effect of an unknown molecule towards a given living organism can be predicted with respect to the behavior of some structurally analogous molecules chosen as a reference, whose toxic action is already known. So quantitative structure–activity relationship (QSAR) and quantitative structure–property relationship (QSPR) techniques help to build powerful mathematical regression models that efficiently predict the reactivity of an unknown molecule in the backdrop of the behavior of an analogous set of chemically or structurally similar compounds. The possibility of an overdose and hence the development of a toxicity criterion of a particular molecule towards a biological entity can also be analyzed through suitable model equations that relate toxicity with molecular structure. Such mathematical equations are also called quantitative structure–toxicity relationship (QSTR) models and have been widely used to assess bio-toxicity. Consequently, the discovery of suitable antidotes, called drugs, to combat such

toxic effects in bio-organisms has also been made possible by utilizing the fruitful quantitative techniques. So, in addition to providing a reliable rationale towards predicting the chemical reactivity and toxicity of molecules, the QSAR, QSPR, and QSTR techniques also play a useful role in drug discovery and design.

In this article, we first of all outline the various theoretical tools and their role in deciphering the biological activity and toxicity patterns in several molecules. Section 2, in this regard, describes the development of QSAR and various allied quantitative relationship-based models and their applications in predicting the reactivity and toxicity of several classes of molecules. Section 2.1 delineates other theoretical techniques like quantum mechanical density functional theory (DFT), molecular dynamics (MD) simulations, nuclear quadrupole resonance (NQR) spectroscopy, their role in building suitable structure–activity relationship (SAR) based models towards analyzing the biological activity of molecules and subsequent applications in medicinal chemistry and drug design. The penultimate section (Sect. 3) outlines a brief overview of conceptual density functional theory (CDFT) and the role of various CDFT-based global and local reactivity descriptors towards developing suitable regression models to correlate and quantify the biological activity and toxicity of molecules. Section 4 finally concludes our discussion with some queries and remarks.

2 Quantitative Structure–Activity/Structure–Property-Based Methods

The trends in the reactivity pattern for a given set of structurally related compounds can be mathematically predicted by building effective structure–activity oriented models. A known set of structurally analogous molecules is chosen and the variation in their reactivity and/or toxicity as a function of the changes in associated molecular framework is studied. The trends obtained as such are then hailed as a benchmark index and further extrapolated to another set of unknown but structurally similar compounds with an aim to correlate their reactivity patterns. Thus, the structure–activity relationship (SAR) based models become useful pathfinders towards predicting chemical reactivity and toxicity of molecules that usually give a nice correlation between theoretical predictions and experimental findings. When such SAR-based models are applied to derive a quantitative relationship between the activity and toxicity trends for a series of chemical compounds, we, hereby, obtain a quantitative structure–activity (or toxicity) relationship, QSAR (or QSTR). QSAR, unlike being conceived merely as an academic tool to predict the reactivity and toxicity of molecules through rigorous data analysis, should be better looked upon as a novel technique that helps to build meaningful models correlating structure, activity, and toxicity with ample judgment and reasoning. The success of such QSAR-based models towards an accurate prediction is sometimes a matter of fortune but, nevertheless a proper judicious rationale and the choice of appropriate

parameters always hold the key. QSAR-based studies, as mentioned above, have got widespread applications in many fields like eco-toxicology, drug discovery, antitumor treatment, molecular modeling, which are nonetheless our direct concern. Schultz et al. [1, 2] in a couple of articles have elaborated the role of QSAR in chemical toxicology. The success beyond any such quantitative prediction lies in the availability of accurate experimental data. An enriched databank renders a given structure–activity relationship (SAR) to be more defined towards an accurate prediction and also helps interpreting the plausible reaction pathway. Some relevant works [3–7] critically highlight the development of the QSAR methodology and its application in predicting bio-toxicity. Insights into the basics of QSAR regarding a nice predictive algorithm can also be obtained from the literature [8–10]. In fact, a hint of the effect of structural changes on reaction mechanisms has been earlier enunciated by Hammett [11, 12] in the well known linear free-energy relationship-based equations applicable to organic reactions involving substituted benzoic acid derivatives. Hammett's idea is also further extended by Taft to understand the varying substituent effects in aliphatic compounds [13]. Based on Hammett's idea, Hansch and his colleagues [14, 15] discussed the role of QSAR, especially the application of biological QSAR containing the terms σ and σ^- . Other studies establishing the idea of constructing strong QSAR models towards elucidating chemico-biological interactions [16–27] and drug design [28–37] are well known. Applications of QSAR in designing biomolecules, drug discovery through a quite accurate in silico toxic data analysis and constructing more effective models by introducing new, improved molecular descriptors are also reviewed [38–40]. The roles of QSAR in biochemistry and several allied toxicological, physicochemical, absorption, disposition, metabolism, and excretion (ADME) processes related to the discovery and subsequent development of various bioactive molecules distinguishing drug-like from nondrug-like molecules [41], drug resistance [42], toxicity prediction [43–48], prediction of various physico-chemical parameters [49], gastro-intestinal absorption [50], activity of peptides [51], data mining [52], drug metabolism [53], and prediction of other pharmacokinetic and ADME properties [54, 55] are quite evident in the relevant literature. Some pertinent articles [56–66] and book [67] provide additional insights into the applicability of QSAR in these fields. Now, in order to build an appropriate structure-reactivity model towards assessing toxicity, a strong chemical database is always needed as a backup. Such databases are usually built by experimentally studying several reaction mechanisms for a large number of toxic assays in vitro or in vivo. In this regard, with an aim towards acquiring a more informative database for an even better predictive correlation, various computational approaches are also proposed [68–71]. Some other studies [72, 73] emphasize on the reaction between an electron rich (nucleophile) and electron poor (electrophile) species. Such triggering reactions are considered as a benchmark that provides a detailed in chemico reactivity database towards toxicity prediction.

Parallel to the development of the QSAR and several models based on the given phenomenon, the advent of an analogous quantitative structure–property relationship (QSPR) rationale is also in the cards. Several research groups have aimed to strengthen QSPR on a theoretical basis and some novel contributions [74–83] in

this regard are worth-mentioning. Katritzky et al. [84] further describe some significant inroads that establish QSPR as a reliable tool in correlating chemical structure with various molecular (physical, chemical, biological, and technological) properties. Application of QSPR in the field of medicinal chemistry through the generation of whole molecular descriptors is also reported [85]. QSPR equations constructed on an empirical basis have been found to maintain a good regime between structure and property for even a large number of data sets. It is shown [56] that for a large number of molecular structures, data reduction methods, like principal component analysis (PCA) of a matrix formed by the assembly of related properties, provide insight into how these related properties correspond to each other quantitatively. Further applications of this method are also discussed. The development and applications of QSPR in pharmaceutical science engrossing a detailed study of the various pharmacokinetic aspects towards drug modeling is also provided in a two-part review by Grover et al. [86, 87].

In recent years, the development of three-dimensional correlation models for analyzing the structure–property relationships has also become quite popular. The three-dimensional quantitative structure–activity relationships (3D-QSAR) algorithm actually analyzes the quantitative relationship between the biological activity of a set of compounds and their three-dimensional properties using statistical correlation methods. 3D QSAR involves the application of force field calculations requiring three-dimensional structures and is concerned with the overall molecule rather than a single substituent. It examines the steric fields (shape of the molecule) and the electrostatic fields based on the applied energy function [88, 89]. Comparative molecular field analysis (CoMFA) [90], a well known three-dimensional quantitative structure–activity relationship (3D-QSAR) method, has been successfully invoked as a powerful tool towards drug design and other allied applications [91–98]. In the CoMFA approach, the molecules are structurally and pharmacologically oriented on the basis of the assumption that each compound acts via a common binding mode [90, 92, 95, 99, 100]. 3D-QSAR CoMFA in association with QSAR methods have been implemented to build useful structure–activity relationship models towards designing effective enzyme inhibitors [101–104], which are also sometimes guided by molecular docking [105–114]. The above phenomenon is also applied for studying the interactions between different organic polychlorinated derivatives with aryl-hydrocarbon receptors (ArHs) [115–118].

QSAR technique is also found to be effective in the simultaneous designing and study of inhibitors to combat the lethal growth of Human Immunodeficiency Virus (HIV) and Hepatitis C Virus (HCV) within living cells [119–121]. Quantitative structure-relationship-based modelings of different series of compounds acting as anti-HIV enzyme inhibitors are broadly studied [122–132]. QSAR, as an *in silico* methodology, has also become very relevant towards modeling antitumor enzyme inhibitors [133] and anticancer drugs [134, 135]. Tan et al. [136], in a recent article, have discussed about the use of coinage metals towards modeling antitumor drugs. In this regard, several other studies [137–149] have also reported that the metal-based compounds are indeed effective in tumor therapy.

Applications of other in silico tools like Three-Dimensional Markovian Electron Delocalization Negentropies (3D-MEDNEs) [150, 151] for toxicity prediction are also reported. QSAR technique is also applied to study the toxicity of phenols [152], alkaloids [153], aliphatic esters [154], and modeling anti-tubercular compounds [155–157].

QSAR studies have become extremely useful towards understanding the toxic levels of bio-organisms. In this context, various topological descriptors [158–160] are introduced to quantitatively correlate the hazardous effect of several chemical compounds on the ecosystem. The toxic effects of various compounds on fishes are reported in detail through fruitful SAR-based regression analyses [161–173].

2.1 Role of Different Theoretical Methods Towards Constructing Effective QSAR Models

Several sophisticated theoretical tools have turned out to be quite efficient towards strengthening the basis of a given structure–activity relationship, which, in turn, evokes a more approximate quantitative rationale. Quantum chemistry (QC) based methods are supposed to be quite significant in this aspect for applications in biological systems [174]. QC-based SAR models provide a much better understanding of the electronic correlation effects than empirical methods [175]. The various QC-based molecular descriptors serve as potential determinants to predict the biochemistry of various enzyme-mediated processes, hallucinogenic activity [176–179] as well as the reactivity of several organic molecules, derivation of partition coefficients, and correlation of other physical parameters [180–184]. The influence of quantum chemical methods and the role of the various allied QC-based descriptors in the structure–activity/toxicity studies are also discussed in detail elsewhere [185]. Applications of the SCF-MO [186] and various high level quantum chemical techniques towards modeling biomolecules and correlating their chemical reactivity with molecular structure are a vastly cultivated topic. Sophisticated ab initio methods like localized perturbation theory and density functional theory (DFT) are employed to understand the catalytic activity of enzymatic reactions, conformational energetics, and nonbonded interactions [187]. Biological activity measures of large molecules like proteins, DNA, etc. from first principles quantum chemical approaches sometimes pose problems due to size and complexity in their structures. Still ab initio DFT and DFT-based molecular dynamics (MD) simulation studies [188, 189] are found to be quite fruitful towards understanding the bond cleavage and bond formation processes of various biomolecular interactions. The roles of ab initio DFT and hybrid quantum mechanical/molecular mechanics (QM/MM) methods towards elucidating the biological modeling and reactivity of various complex motifs [190, 191] including the cytochrome P450 family of enzymes [192, 193] are also reported. DFT-based studies in the field of medicinal chemistry [194] as well as the prediction of toxicity measures for molecules acting as anticancer

agents or as drugs are also well documented [195, 196]. Quantum chemistry-based QSAR/QSPR models are also employed towards studying the bio-toxicity of *cis*-platin complexes as effective anticancer drugs [197, 198]. Sarmah and Deka [199] have further made a DFT-based 2D QSAR study on the anticancer activities of various nucleosides.

Superior QSAR-based regression models explaining the predictive nature of enzyme activity and the toxicity of various organic molecules like substituted aromatic compounds are studied vividly using the DFT protocol. Density functional theory as a powerful mathematical algorithm is exploited as a useful tool to provide sets of quantum chemical descriptors that help finding suitable correlations for enzymatic activities and toxicity of various classes of organic compounds and their derivatives [200–217]. Again Elstner and coworkers [218, 219] have elaborated that the self-consistent charge density functional tight binding (SCC-DFTB) formalism, compared to DFT may appear as a better mathematical rationale for large biological systems, as the former method considers the phenomena of nonbonded interactions and charge transfer processes in a better way.

In another approach, with an aim to offer a realistic motive towards handling millions of databases and hundreds of descriptors for a fruitful structure–activity relationship (SAR), Putz and coworkers have proposed a unique QSAR model called spectral-SAR (S-SAR) [220], which considers the spectral norm in quantifying toxicity and reactivity with molecular structure. A handful of applications of the S-SAR algorithm in dealing with ecotoxicity, enzyme activity, and anticancer bioactivity are well established [221–227]. The S-SAR model coupled with Element Specific Influence Parameter (ESIP) formulations [228] are also utilized for predicting ecotoxicity measures. QSAR studies on the anti-HIV-1 activity of HEPT (1-[(2-hydroxyethoxy)methyl]-6-(phenylthio)thymine) [229] and further studies involving the minimum topological difference (MTD) method [230, 231] are also reported [232].

A number of other novel methodologies providing good correlations and meaningful structure–activity-based regression models are also available. Latosińska [233, 234] delineates the application of nuclear quadrupole resonance (NQR) spectroscopy in studying drug molecules. Applications of other spectroscopic methods like NMR, EPR, UV, and IR towards investigating the biological activity of molecules are presented earlier [235–237]. The variable selection and modeling method based on the prediction (VSMP) [238], quantitative structure–(chromatographic) retention relationships (QSRR) [239], topological quantum similarity index (TQSI) [240, 241] models, and quantum topological molecular similarity (QTMS) descriptors [242, 243] also have ample contributions towards ensuring refinements in the predictive nature of quantitative structure–activity relationships. Modeling of soft electrophilicity and hydrophobicity [68, 244–246] as descriptors for toxicology-based QSAR predictions has also become a quite effective analytical algorithm.

3 Conceptual Density Functional Theory

Chemistry is all about the formation of new molecules from some parent motifs. All chemical processes follow the basic philosophy of rupture of old inter-atomic linkages (bonds) followed by the formation of newer ones upon electronic interactions between two reacting species. Chemical reactions can be experimentally studied as well as theoretically monitored. The advent of quantum chemistry and its subsequent developments with time have boasted off some exceptionally sophisticated mathematical paradigms, which are proven to be very unique in studying several classes of chemical reactions along with accurately screening the corresponding reaction pathways. Applications of a few such theoretical techniques towards elucidating the toxicity trends of various molecules and building of fruitful SAR-based regression models as a useful benchmark for further prediction are already mentioned in the preceding section. In this section, we discuss a very popular theoretical model, conceptual density functional theory (CDFT) and its utility towards setting up more refined mathematical relationships to quantitatively correlate biological activity and toxicity with molecular structure.

Conceptual density functional theory (CDFT) [247–251], after its inception a few decades ago, has proved itself to be a very powerful theoretical paradigm to understand molecular reactivity. Conceptual DFT basically owes its origin from the research group of Prof. Robert G. Parr, which is later made more coherent and is developed with an aim to offer a logistic idea about electronic interactions. In this effort, several mathematical response functions, also known as reactivity descriptors are introduced, which nicely explain the mode of chemical reactivity between a given set of interacting molecules. Conceptual DFT in association with its various global reactivity descriptors like electronegativity [252, 253] (χ), hardness [254, 255] (η), and electrophilicity [256–259] (ω) originating from the same popular qualitative concepts, respectively, along with the corresponding local variants like atomic charges [260] (Q_k), Fukui functions [261] $f(\mathbf{r})$, and their condensed-to-atom variants [262] (f_k) develops intuitive reactivity trends for a molecule as a whole and for each and every active site of the same as well.

The term electronegativity (χ) was initially coined by Pauling [263, 264], which virtually meant the “power” of an atom in a molecule to attract bonded electrons towards itself. Mulliken [265, 266] further defined the term electronegativity (χ) as a function of the two main experimentally determined parameters of a system, viz., the first ionization energy (IP) and the electron affinity (EA) and showed that χ may be expressed as an arithmetic mean of the respective IP and EA values. Thus, according to Mulliken:

$$\chi = \frac{IP + EA}{2}. \quad (1)$$

The chemical potential (μ) of an atom/molecule may be quantitatively assessed by the theory of statistical ensembles. Thus, if an atom or a molecule be considered

as a member of a grand canonical ensemble where the energy (E) and the number of electrons (N) are continuous functions and vary independently, the chemical potential of the ensemble may be expressed as

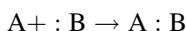
$$\mu = \frac{\partial E}{\partial N}, \text{ at constant entropy.} \quad (2)$$

Gyftopoulos and Hatsopoulos [267] proposed a quantum thermodynamic definition of electronegativity (χ) of a system where the electronic chemical potential (μ) is being considered as the negative of the same.

Based on Mulliken's [265, 266] prescription for electronegativity, the Mulliken chemical potential can be similarly expressed as a finite-difference approximation of the derivative of the electronic energy with respect to the number of electrons and can be expressed as

$$\mu_{\text{Mulliken}} = -\chi_{\text{Mulliken}} = -\frac{\text{IP} + \text{EA}}{2} = \left[\frac{\partial E(N)}{\partial N} \right]_{N=N_0}. \quad (3)$$

The concept of “hardness” or “softness” in chemistry was introduced by Ralph G. Pearson [268–272] in connection with the study of generalized Lewis acid–base reactions



where A, the electron acceptor is the acid while the electron-pair donor B is the base. Studies on the above donor–acceptor interaction (between B and A) provided a splendid qualitative rationale to conceive global hardness (or softness). Global hardness (η) is simply attributed to the degree of compactness of the electron cloud encompassing the nucleus/nuclei of an atomic/molecular system. Global softness (S) on the other hand being the reciprocal of global hardness describes the extent to which the electronic environment surrounding the nucleus/nuclei of an atomic/molecular species tends to loosen itself. The qualitative concept of molecular hardness was put forward quantitatively in the form of a mathematical descriptor by Parr and Pearson [254, 255], who defined that the chemical hardness (η) for a molecular system can be identified as the first derivative of the chemical potential (μ) or the second derivative of the energy (E) as a function of the number of electrons, N , at a fixed external potential $v(\mathbf{r})$:

$$\eta = \left(\frac{\partial \mu}{\partial N} \right)_{v(\mathbf{r})} = \left(\frac{\partial^2 E}{\partial N^2} \right)_{v(\mathbf{r})}. \quad (4)$$

The convexity in the E vs. N curve renders the value of η to remain positive and from the finite-difference approach the corresponding curvature equals $\text{IP} - \text{EA}$, which actually signifies hardness. Therefore:

$$\eta = \left(\frac{\partial \mu}{\partial N} \right)_{v(\mathbf{r})} = \left(\frac{\partial^2 E}{\partial N^2} \right)_{v(\mathbf{r})} \approx \text{IP} - \text{EA}. \quad (5)$$

Thus, for a normal charge-transfer interaction between a donor–acceptor couple like a Lewis acid–base pair, the chemical hardness (η) of a given species signifies the measure of its reluctance towards further electron shift to the other species, thereby upholding its qualitative concept of compactness.

Global softness [273] (S), as a reciprocal of the global hardness (η) for a molecular system, is expressed as

$$S = \frac{1}{\eta} = \left(\frac{\partial N}{\partial \mu} \right)_{v(\mathbf{r})}. \quad (6)$$

Among the various reactivity descriptors, electrophilicity [256–259] (ω) and its recently reported mathematical refinement, net electrophilicity ($\Delta\omega^\pm$) [274] deserve a special mention from the viewpoint of constructing useful QSAR models.

The term “electrophile” in chemistry is coined to those species, which have got a “special” kind of affinity for electrons. Electrophiles are in general electron-deficient systems, sometimes bearing positive charge(s) or may be a free radical that has got a propensity towards attracting electron-rich motifs called nucleophiles. Electrophiles and nucleophiles together play a vital role in determining the mechanistic courses for several types of organic and inorganic chemical reactions like acid–base, redox, addition, substitution, elimination, molecular rearrangement, etc. An electrophile upon interaction with an electron rich species (nucleophile) will strongly attract the electron density of the latter and get stabilized with the gradual lowering of energy and formation of a stable covalent bond. A clear quantitative estimate of this simple qualitative idea as to which extent an electrophilic system would be energetically stabilized upon transfer of electrons from the adjacent nucleophile was, however, lacking. In this context, Maynard et al. [275], upon experimenting with the human immunodeficiency virus type 1 (HIV-1) nucleocapsid protein p7 (NCp7) with a variety of electrophilic agents, showed that the fluorescence decay rates vary almost linearly with the ratio of the square of electronegativity (χ) to hardness (η), χ^2/η , which can further be designated as the capacity of an electrophile to attract electrons from a nucleophile to create a covalent bond. Such an innovative idea developed by Maynard et al. [275] inspired Parr and coworkers [256] to put forward a more precise mathematical rationale for correlating the electron-attracting power of a species. Subsequently, a new descriptor called electrophilicity index (ω) is introduced. Parr’s postulate of electrophilicity index (ω) unlike Maynard et al. (which was developed from a kinetic consideration by studying reaction rates) was based on a thermodynamic background and thus ω became a measure of the favorable change in energy upon saturation of a system with electrons. The electrophilicity index (ω) as prescribed by Parr et al. [256] is

$$\omega = \frac{\mu^2}{2\eta} = \frac{\chi^2}{2\eta}. \quad (7)$$

The above expression of electrophilicity (ω) is comparable with the equation of electrical power (W) in classical physics where $W = V^2/R$, where V and R attribute to the voltage and electrical resistance, respectively. Therefore, ω represents the “electrophilic power” of a species. Comprehensive reviews on electrophilicity index (ω) [256–259] regarding its genesis and rigorous applications towards an understanding of chemical reactivity are available.

In an attempt to further correlate the associated energy changes between the interacting acceptors and donors in a charge-transfer process, Gazquez et al. [276], based on the second-order Taylor series energy expansion formula as a function of the number of electrons (N) in the intervals between $N - 1$ and N , and N and $N + 1$, showed that the electrodonating (ω^-) and the electroaccepting (ω^+) powers may be defined as

$$\omega^- = \frac{(\mu^-)^2}{2\eta^-}; \quad \omega^+ = \frac{(\mu^+)^2}{2\eta^+}, \quad (8)$$

where μ^- and μ^+ denote the chemical potential for electron donation and electron acceptance, respectively. η^- and η^+ signify the hardness for electron donation and electron acceptance, respectively. Further evidences [276] reveal that μ^- and μ^+ can be equalized to μ so that: $\mu^- = \mu^+ = \mu$ and, in the same spirit, $\eta^- = \eta^+ = \eta$. Thus, the electrodonating (ω^-) and the electroaccepting (ω^+) powers become equivalent with the concept of electrophilicity (ω) owing to which ω , ω^- , or ω^+ may be expressed in terms of chemical potential (μ) and hardness (η) as $\omega^- = \omega^+ = \omega = \mu^2/2\eta$. Gazquez et al. [276] proposed two sets of definitions for ω^- or ω^+ , which were based on two different approaches; one exploiting the original formula as above, which is expressed as

$$\omega^+ = \frac{EA^2}{2(IP - EA)}, \quad (9)$$

$$\omega^- = \frac{IP^2}{2(IP - EA)} \quad (10)$$

and the other by using an alternative expression for energy as

$$\omega^+ = \frac{(IP + 3EA)^2}{16(IP - EA)}, \quad (11)$$

$$\omega^- = \frac{(3IP + EA)^2}{16(IP - EA)}, \quad (12)$$

where IP and EA correspond to the first ionization energy and electron affinity of the system, respectively. It [276] was also shown that the alternative approach of expressing ω^- or ω^+ yielded better correlations than the other one.

The concept of net electrophilicity ($\Delta\omega^\pm$), proposed recently by Chattaraj et al. [274] for a system, is an attempt to assess the resultant electron-accepting power exhibited by a molecule upon chemical response due to the combined attractive and repulsive effects arising out of the presence of both electrons and nuclei. This new dual descriptor ($\Delta\omega^\pm$), therefore, seems to be an appraisal of the electrophilicity of a system relative to its own nucleophilicity and serves better to provide a more meaningful physical rationale towards understanding the “electrophilic power” of a system. Energy considerations, however, reveal that a larger value of ω^+ for a system presupposes its enhanced capability to accept charge whereas a smaller value of ω^- necessitates the same to serve as a better donor. The mathematical basis of $\Delta\omega^\pm$ thus arises out of a parity between ω^+ and ω^- where the negative (or reciprocal) of ω^- is compared with ω^+ . Thus, net electrophilicity ($\Delta\omega^\pm$) in terms of ω^+ and ω^- is formulated as

$$\begin{aligned}\Delta\omega^\pm &= \{\omega^+ - (-\omega^-)\} = (\omega^+ + \omega^-) \\ &\text{or} \\ \Delta\omega^\pm &= \left\{ \omega^+ - \left(\frac{1}{\omega^-} \right) \right\}.\end{aligned}\quad (13)$$

The polarizability (α) of an atom or a molecule means the lowest order response of its electron cloud to an external weak electric field [277, 278]. The static dipole polarizability (α), a linear response property, is defined as the second derivative of the total electronic energy (E) with respect to the external homogeneous electric field as

$$\alpha_{\alpha\beta} = - \left(\frac{\partial^2 E}{\partial F_\alpha \partial F_\beta} \right)_{F=0}, \quad (14)$$

where F_α and F_β are the electric field components for a fixed coordinate system with $\alpha, \beta, \gamma = x, y, z$. The polarizability (α) is very sensitive to basis set, electron correlation and relativistic effects, and to the vibrational structure in case of a molecule. Qualitatively, polarizability (α) varies inversely with global hardness [279–284] (η) and is proportional to the global softness (S) as expected. Hence, an interesting linear correlation is established between the static dipole polarizability (α) and the third power of global softness [285–287].

The magnetizability (ξ) for a chemical system is a measure of the linear response of its electron cloud to an externally applied magnetic field and is expressed as

$$\xi = - \left(\frac{\partial^2 \varepsilon(B)}{\partial B^2} \right)_{B=0}, \quad (15)$$

where B is the external magnetic field.

The magnetizability (ξ) of a chemical species varies proportionately with its softness (S) and polarizability (α). A softer species is, therefore, more polarizable and hence more magnetizable [288].

In addition to the various conceptual DFT-based global reactivity descriptors, the local reactivity descriptors also provide valuable insights in determining chemical reactivity patterns. The various local reactivity indices like electron density ($\rho(\mathbf{r})$), Fukui function ($f(\mathbf{r})$), local softness ($s(\mathbf{r})$), local hardness ($\eta(\mathbf{r})$), and philicity ($\omega(\mathbf{r})$) help to assess the response of a particular atomic site in a molecule during a chemical attack.

The electron density ($\rho(\mathbf{r})$) provides useful information regarding site-reactivity for a molecule and is assessed in terms of the first-order variation of the energy as a function of the external potential [247, 289] ($v(\mathbf{r})$).

$$\rho(\mathbf{r}) = \left(\frac{\delta E}{\delta v(\mathbf{r})} \right)_N. \quad (16)$$

The Fukui function [261, 290] ($f(\mathbf{r})$) is usually applied to quantitatively evaluate the extent of local response of a particular active site of a chemical system. It is widely applied as a popular local variant and is defined as the differential change in electron density ($\rho(\mathbf{r})$) due to an infinitesimal change in the number of electrons (N).

$$f(\mathbf{r}) = \left(\frac{\partial \rho(\mathbf{r})}{\partial N} \right)_{v(\mathbf{r})} = \left(\frac{\delta \mu}{\delta v(\mathbf{r})} \right)_N. \quad (17)$$

Owing to a discontinuity in the derivative in Eq. (17) for integral values of N , three different types of Fukui functions can be defined by applying the finite difference and frozen core approximations as follows [261, 290]:

For nucleophilic attack

$$f^+(\mathbf{r}) = \left(\frac{\partial \rho(\mathbf{r})}{\partial N} \right)_{v(\mathbf{r})}^+ \approx \rho_{N+1}(\mathbf{r}) - \rho_N(\mathbf{r}) \approx \rho_{\text{LUMO}}(\mathbf{r}) \quad (18a)$$

For electrophilic attack

$$f^-(\mathbf{r}) = \left(\frac{\partial \rho(\mathbf{r})}{\partial N} \right)_{v(\mathbf{r})}^- \approx \rho_N(\mathbf{r}) - \rho_{N-1}(\mathbf{r}) \approx \rho_{\text{HOMO}}(\mathbf{r}) \quad (18b)$$

For radical attack

$$f^0(\mathbf{r}) = \frac{1}{2} [f^+(\mathbf{r}) + f^-(\mathbf{r})]. \quad (18c)$$

Thus, $f(\mathbf{r})$, on the basis of the frontier orbital theory proposed by Fukui et al. [291–294], physically correlates the tendency of a local atomic site in a molecule to interact upon chemical response. The Fukui functions are, therefore, referred to as the DFT analogue of the frontier orbitals [295].

A coarse-grained atom by atom representation of the Fukui function, called condensed-to-atom Fukui function, was proposed by Yang and Mortier [262] and based on a finite-difference approach in terms of the Mulliken population analysis (MPA) scheme, they may be represented as

$$f_k^+ = q_k(N+1) - q_k(N) \quad \text{for nucleophilic attack,} \quad (19a)$$

$$f_k^- = q_k(N) - q_k(N-1) \quad \text{for electrophilic attack,} \quad (19b)$$

$$f_k^o = [q_k(N+1) - q_k(N-1)]/2 \quad \text{for radical attack,} \quad (19c)$$

where q_k refers to the electron population at a particular atomic site k in a molecule. The Eqs. (19a)–(19c) can be easily evaluated through population analysis data that helps in further quantum chemical calculations.

The global softness (S) mathematically means an integration of the individual local softness ($s(\mathbf{r})$) values computed for all the atomic sites of an entire molecule. A normalized condition between S and $s(\mathbf{r})$ is thus expressed as

$$S = \int s(\mathbf{r}) d\mathbf{r}, \quad (20)$$

which is quite at par with the condition as $f(r)$ is normalized to unity.

The local softness $s(\mathbf{r})$ is related to the Fukui functions $f(\mathbf{r})$ through a chain rule as

$$s(\mathbf{r}) = \left(\frac{\partial \rho(\mathbf{r})}{\partial \mu} \right)_{v(\mathbf{r})} = \left(\frac{\partial \rho(\mathbf{r})}{\partial N} \right)_{v(\mathbf{r})} \cdot \left(\frac{\partial N}{\partial \mu} \right)_{v(\mathbf{r})} = f(\mathbf{r}) \cdot S. \quad (21)$$

The Fukui function ($f(\mathbf{r})$) makes an assessment of the tendency towards reactivity of the different local sites of the same molecule and thus serves as an intramolecular reactivity descriptor. The local softness ($s(\mathbf{r})$), on the other hand, compares and correlates the propensity of a pair of interacting molecular neighbors during chemical response and unlike $f(\mathbf{r})$ serves as an intermolecular reactivity descriptor.

Local hardness ($\eta(\mathbf{r})$) [296, 297] is introduced to quantify the reluctance of a particular atom or group in a molecule towards chemical attack. Unlike a normalized condition between $s(\mathbf{r})$ and S , however, $\eta(\mathbf{r})$ cannot be normalized to η . $\eta(\mathbf{r})$ is thus differently expressed as

$$\eta(\mathbf{r}) = \frac{1}{N} \int \frac{\delta^2 F[\rho(\mathbf{r})]}{\delta \rho(\mathbf{r}) \delta \rho(\mathbf{r}')} \rho(\mathbf{r}') d\mathbf{r}', \quad (22)$$

where $F[\rho(\mathbf{r})]$ defines the Hohenberg–Kohn universal functional. Local hardness is also expressed [296, 297] differently at par with global hardness (η) by replacing the number of electrons (N) with electron density ($\rho(\mathbf{r})$) as

$$\eta(\mathbf{r}) = \left(\frac{\delta\mu}{\delta\rho(\mathbf{r})} \right)_{v(\mathbf{r})}. \quad (23)$$

But Eq. (23) is ambiguous due to an inter-dependence of $\rho(\mathbf{r})$ and $v(\mathbf{r})$ and thus cannot be considered as a basic definition for $\eta(\mathbf{r})$. Thus, unlike η and S , $\eta(\mathbf{r})$ and $s(\mathbf{r})$ do not even hold a reciprocal relationship and are interconnected as

$$\int \eta(\mathbf{r})s(\mathbf{r})d\mathbf{r} = 1. \quad (24)$$

An acceptable definition of local hardness and subsequent identification of the “hard” sites in a molecule is, therefore, in demand [298–304]. The “minimum Fukui function rule” [301] asserts that hard reactions, unlike softer ones, would prefer sites with minimum Fukui function values. Although there are applications [305–307] of this rule, subsequent criticisms [303, 308] are also reported. The “minimum Fukui function rule” is unable to correlate the electrostatic hard–hard interactions that are predominantly charge-controlled with hardly any relevant effect from the associated frontier orbitals. The given rule cannot justify the inadequacy of the frontier orbitals and also misses the role of electrostatic interactions in hard–hard interactions [303, 304, 308].

The local hardness ($\eta(\mathbf{r})$) and the Fukui function ($f(\mathbf{r})$) can be used [309] to obtain the global hardness, which can be expressed as

$$\eta = \int \eta(\mathbf{r})f(\mathbf{r})d\mathbf{r}. \quad (25)$$

The concept of philicity ($\omega^{\alpha}(r)$) introduced by Chattaraj et al. [310, 311] signifies the aptitude towards reactivity (electrophilicity or nucleophilicity) of a local atomic site in a molecule. In other words, an increase (or decrease) in the reactivity of any local site of a molecule does not affect the electrophilicity (ω) of the entire species as a whole—the global reactivity parameter, ω remains conserved. The transformation of the local electrophilicity ($\omega(\mathbf{r})$) into global electrophilicity (ω) is expressed as

$$\omega = \omega \int f(\mathbf{r})d\mathbf{r}$$

or,

$$\omega = \int \omega f(\mathbf{r})d\mathbf{r} = \int \omega(\mathbf{r})d\mathbf{r}$$

thus,

$$\omega(\mathbf{r}) = \omega f(\mathbf{r}). \quad (26)$$

Although $\omega(\mathbf{r})$ contains information about both ω and $f(\mathbf{r})$ but $f(\mathbf{r})$ alone cannot provide any input regarding $\omega(\mathbf{r})$ without the knowledge of ω . The concept of philicity is widely applicable to diverse types of chemical reactions, which, therefore, presupposes $\omega(\mathbf{r})$ to be represented as $\omega^\alpha(\mathbf{r})$ —the symbol of “philicity.” The sign of α varies with the different categories of reactions, i.e. $\alpha = +, -, \text{ and } 0$ represents nucleophilic, electrophilic, and radical attacks, respectively. Thus:

$$\begin{aligned} \omega^\alpha(\mathbf{r}) &= \omega \cdot f^\alpha(\mathbf{r}) \\ \text{or, } \omega_k^\alpha &= \omega \cdot f_k^\alpha \end{aligned} \quad (27)$$

where ω_k^α refers to the condensed-to-atom local philicity variant for the k th atomic site in a molecule. The philicity (ω_k^α) seems to be a better intermolecular local reactivity index than the Fukui function (f_k^α).

The group philicity concept [312] (ω_g^α) gains importance when we consider the reactivity of an atomic assembly or a group and is usually expressed as a summation of the individual condensed-to-atom philicities (ω_k^α) over all the relevant atoms. Thus

$$\omega_g^\alpha = \sum_{k=1}^n \omega_k^\alpha \quad (28)$$

where n denotes the number of atoms present in the reacting group and $\alpha = +, -, \text{ and } 0$ refers to nucleophilic, electrophilic, and radical attacks, respectively.

The CDFT-based global and local reactivity descriptors and their modes of influencing chemical reactivity are further appreciated in terms of the various allied molecular electronic structure principles like the principle of maximum hardness [273, 313–319] (PMH), minimum polarizability principle [279, 320–322] (MPP), minimum electrophilicity principle [323, 324] (MEP), minimum magnetizability principle [288, 325] (MMP), etc. The various reactivity descriptors theoretically justify the observed chemical behavior of a molecule. Widespread applications of conceptual DFT towards modeling a plethora of molecular cluster assemblies accompanied by a rigorous analysis of their bonding, stability, and reactivity trends are adequately reported in the literature. The variation of the conceptual DFT-based reactivity indices as a function of structural changes and/or substituent effects convey invaluable insights into a quantitative correlation between chemical reactivity and toxicity. It [326] is, however, pointed out that the magnitudes of the various reactivity descriptors are quite dependent on the level of theory and the type of the basis set used. Thus, for an effective theoretical benchmarking to provide a structure–activity/toxicity correlation at par with experimental values, proper knowledge regarding the selection of level of theory and/or molecular basis set(s) is necessary. An appropriate choice indeed becomes new pathfinders towards building fruitful regression models for QSAR/QSPR/QSTR analyses.

3.1 Computational Details

The molecular geometries of the different classes of compounds under study are optimized at various levels (semi-empirical and DFT) of theory using the Gaussian 98 [327], Gaussian 03 [328] program packages. The stability of the optimized structures is characterized by their harmonic vibrational frequency values. The number of imaginary frequencies (NIMAG) is zero in every case, which corresponds to the existence of the given molecular geometry at a minimum on the potential energy landscape. The ionization potential (IP) and electron affinity (EA) are computed through the Koopmans' theorem [329] or by exploiting the Δ SCF technique. The atomic charges [260] (Q_k) and Fukui functions [261] ($f(\bar{r})$) are computed from the Mulliken population analysis (MPA) [262], natural population analysis (NPA), or Hirschfield population analysis (HPA) [330] schemes. The Hirschfield charges are computed with the BLYP/DND method using the DMOL³ package [331].

3.2 Applications of Conceptual DFT in QSAR/QSTR

Various conceptual DFT-based reactivity indices in association with some new parameters are successfully employed in the development of stronger QSAR/QSTR models [332]. Deeper correlations of the toxicity of different classes of organic compounds like chlorinated benzenes [333], polychlorinated biphenyls [312, 334–336], and benzidine [337] at DFT level of theory are reported. The toxicity of the polychlorinated biphenyls as well as benzidine is influenced by its electron affinity and planarity. The interactions of the chlorinated benzo-derivatives and benzidine with other biomolecules like nucleic acid/base pairs or aryl hydrocarbon hydroxylase (AHH) receptors are primarily of charge-transfer type, which can be quantitatively assessed from Parr to Pearson formula [254] and can be given as

$$\Delta N = \frac{\mu_B - \mu_A}{2(\eta_A + \eta_B)}, \quad (29)$$

where, ΔN represents the fractional number of electrons transferred from system A to system B during a charge-transfer process occurring between two interacting systems A and B , which eventually produce an adduct $A:B$. μ_A , η_A and μ_B , η_B denote the chemical potential and hardness of systems A and B , respectively.

In these cases, the chlorobenzenes act as electron acceptors while the benzidine molecule behaves as an electron donor. The effect of chlorine substitution on the aromaticity of the planar benzene ring of the chlorobenzene derivatives is assessed from the nucleus independent chemical shift (NICS) criterion proposed by Schleyer et al. [338]. Among the various global and local molecular descriptors, electrophilicity (ω) is found to be the most appropriate reactivity parameter regarding toxicity

assessments for the entire molecule as well as for a particular local reactive site. Toxicity-based regression models developed by correlating the experimental biological activities (pIC_{50}) of polychlorinated dibenzofurans (PCDFs) with their corresponding activity (pIC_{50}) values computed by using the electrophilicity and the local electrophilic power illustrate a fine predictive relationship [339]. Padmanabhan et al. [340], in the line of a contemporary article by Toro-Labbé and coworkers [341], utilized the global philicity concept to propose a new multiphilic descriptor ($\Delta\omega_k$), which considers both the nucleophilicity and electrophilicity of a local atomic site in a molecule. The $\Delta\omega_k$ values computed for an entire set of chlorinated benzenes in both the gas and solvent media give a fair potential of its predictive ability regarding toxic effects on aquatic species. The group philicity concept as a possible descriptor in association with electrophilicity is also applied to build structure–toxicity-based relationship models for chlorophenols (CPs) [342] with high internal predictive ability. Effective QSAR models for the toxicity of CPs upon *Daphnia magna* are presented by utilizing the group philicities. Group philicity and electrophilicity together also provide useful structure–activity-based predictions for the CPs against *Brachydanio rerio* and *Bacillus*. Cronin et al. [343], in an earlier study, showed that electrophilicity can be successfully implemented to assess the toxic effects of a large number of aromatic compounds on *Tetrahymena pyriformis* and in that case, the maximum acceptor superdelocalizability (A_{max}) proved to be a better descriptor than the energy of the lowest unoccupied molecular orbital (E_{LUMO}) in parametrizing ω . Electrophilicity (global and local) as well as its modified variants are, therefore, quite effective in presenting useful correlation of ecotoxicological trends of various organic molecules with bio-organisms.

Electrophilicity, as a cardinal reactivity index, is quite successfully implemented to study the biological activities of male and female sex hormones like testosterone and estrogen derivatives [344]. Another subsequent article [345] delineates that the number of atoms in a molecule (N_A) can be used as a valid molecular descriptor in the QSAR parlance towards explaining the activity of the sex hormones. A recent QSAR-based modeling [346] on the ecotoxicological effects of a large set of aromatic compounds on micro-organisms like *Tetrahymena pyriformis*, *Daphnia magna*, *Brachydanio rerio*, and *Bacillus* also reflects the usefulness of the atom counting (number of non-hydrogenic atoms) criterion as a simple and effective descriptor in association with ω and the ground state energy (E_{GS}). The electrophilicity protocol is also quite perfect in providing an impeccable prediction of toxicity, which builds a considerable data bank upon interaction of a large number of aliphatic compounds with *Tetrahymena pyriformis* [347]. In most cases, excellent correlations are obtained between toxicity and electrophilicity and/or its local variant, thereby giving rise to a rationale for the toxic action. Quantitative assessments of the logarithm of *n*-octanol/water partition coefficient ($\log K_{\text{ow}}$) (an important index for toxicological and pharmacological studies) on a large set of polychlorinated biphenyls (PCBs) [348] using electrophilicity as a descriptor yields reasonably good coefficients and internal predictive ability values. A recent article [349] describes the relative efficacy of electrophilicity, energy of the lowest unoccupied molecular orbital (E_{LUMO}), and $\log P$ (1-octanol/water partition coefficient) as global descriptors in predicting the toxicity (pIGC_{50}) of a large number of

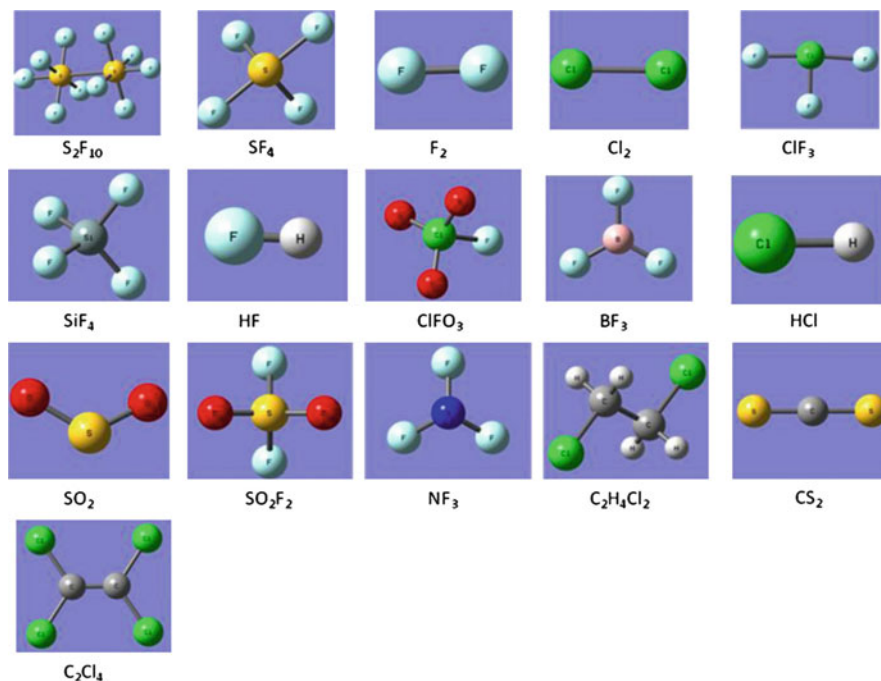


Fig. 1 Optimized ground state geometries (B3LYP/6-311 + G(d)) of some selected sulfur and halogen compounds (Reprinted with permission from [350]. Copyright 2011 IGI Global)

aliphatic compounds towards *Tetrahymena pyriformis*. It further transpires that electrophilicity marginally supersedes over E_{LUMO} to provide better regression analyses in most cases, refinements also being dependent on the units of the IGC_{50} magnitudes.

QSTR-based one-parameter and multi-parameter regression models involving the conceptual DFT-based global reactivity indices and the newly proposed net electrophilicity [274] ($\Delta\omega^\pm$) are developed to assess the toxic effects of some halogen, sulfur, and chlorinated aromatic compounds [350]. Two sets of inorganic compounds, containing mainly halogen and sulfur in the first set, displayed in Fig. 1, and chlorinated aromatic compounds in the second set, displayed in Fig. 2, are considered for investigation.

The R^2 , R^2_{CV} , and R^2_{adj} values given in Table 1 and the experimental vs. calculated Log (LC_{50}) plots utilizing the one-parameter regression equations for sets 1 and 2 in Figs. 3 and 4, respectively, reveal that in case of the first set, the newly proposed net electrophilicity descriptor [274] ($\Delta\omega^\pm$) provides the best result, whereas, for the second set, both electrophilicity index (ω) as well as net electrophilicity index ($\Delta\omega^\pm$) put up a comparably decent performance. The newly proposed net electrophilicity index ($\Delta\omega^\pm$) as a refinement of electrophilicity (ω) has got a fair potential in constructing effective regression models suitable for QSAR/QSTR studies. Appropriate QSAR-based regression models are quite capable of explaining the toxicity trends of several alkali metal ions and various

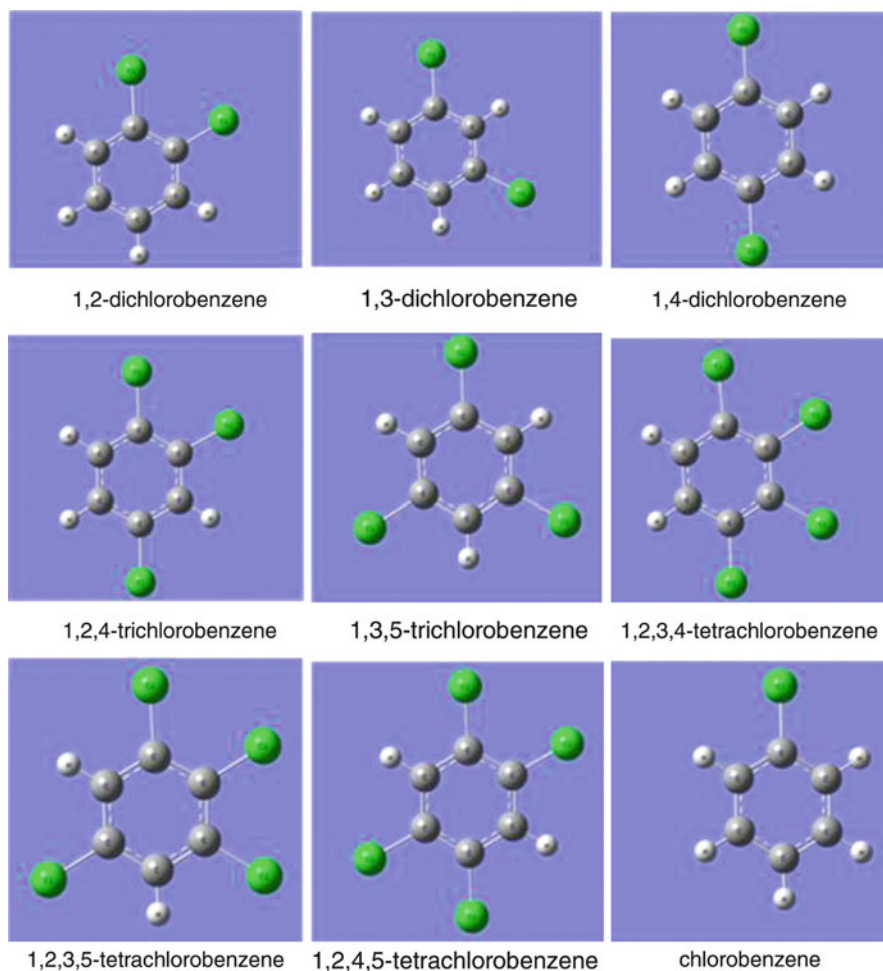


Fig. 2 Optimized ground state geometries (B3LYP/6-311 + G(d)) of chlorinated aromatic compounds (Reprinted with permission from [350]. Copyright 2011 IGI Global)

arsenic compounds [351]. A suitable three-parameter-based regression model consisting of electrophilicity (ω), philicity (ω_{As}^+), and atomic charge (Q_{As}) is found to be quite effective in predicting the toxicity of the arsenic compounds.

3.3 Application of Conceptual DFT in Designing Anticancer Drugs

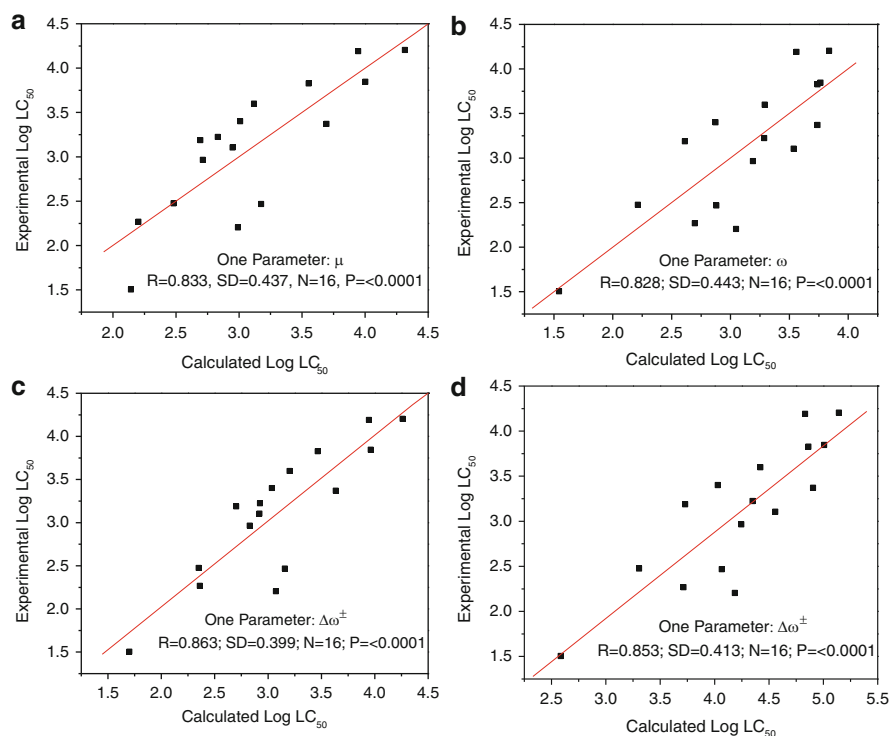
Boron in many cases is now-a-days considered as a “new carbon” in drug modeling [352]. A conceptual DFT-based study is made to assess the anticancer properties of

Table 1 Regression models and various coefficients of determination with the various combinations of μ , ω , $\Delta\omega^{\pm a}$, and $\Delta\omega^{\pm b}$ for the sets 1 and 2

| Regression model | R^2 | R_{CV}^2 | R_{adj}^2 |
|---|-------|------------|-------------|
| Set 1 (Halogen and Sulfur compounds) | | | |
| $\text{Log LC}_{50} = (0.521 \times \mu) + 6.60$ | 0.694 | 0.618 | 0.673 |
| $\text{Log LC}_{50} = (-0.417 \times \omega) + 4.64$ | 0.686 | 0.618 | 0.662 |
| $\text{Log LC}_{50} = (-0.2341 \times \Delta\omega^{\pm a}) + 6.33$ | 0.745 | 0.713 | 0.742 |
| $\text{Log LC}_{50} = (-0.233 \times \Delta\omega^{\pm b}) + 6.33$ | 0.728 | 0.696 | 0.727 |
| Set 2 (Chlorinated aromatic compounds) | | | |
| $\text{Log LC}_{50} = (-2.95 \times \mu) - 7.89$ | 0.729 | 0.549 | 0.691 |
| $\text{Log LC}_{50} = (2.66 \times \omega) - 0.4$ | 0.872 | 0.798 | 0.854 |
| $\text{Log LC}_{50} = (1.74 \times \Delta\omega^{\pm a}) - 10.3$ | 0.686 | 0.449 | 0.641 |
| $\text{Log LC}_{50} = (1.43 \times \Delta\omega^{\pm b}) - 2.50$ | 0.844 | 0.751 | 0.822 |

^aCalculated as per Eqs. (9), (10).^bCalculated as per Eqs. (11), (12).

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**Fig. 3** (a) Experimental versus calculated toxicity (Log (LC₅₀)) values as obtained using one-parameter chemical potential (μ) regression model for the complete set of Fluorine and Sulfur compounds; (b) same for electrophilicity index (ω); (c) same for net electrophilicity ($\Delta\omega^{\pm}$) from Eqs. (9), (10) and (13); (d) same for net electrophilicity ($\Delta\omega^{\pm}$) from Eqs. (11), (12) and (13) (Reprinted with permission from [350]. Copyright 2011 IGI Global)

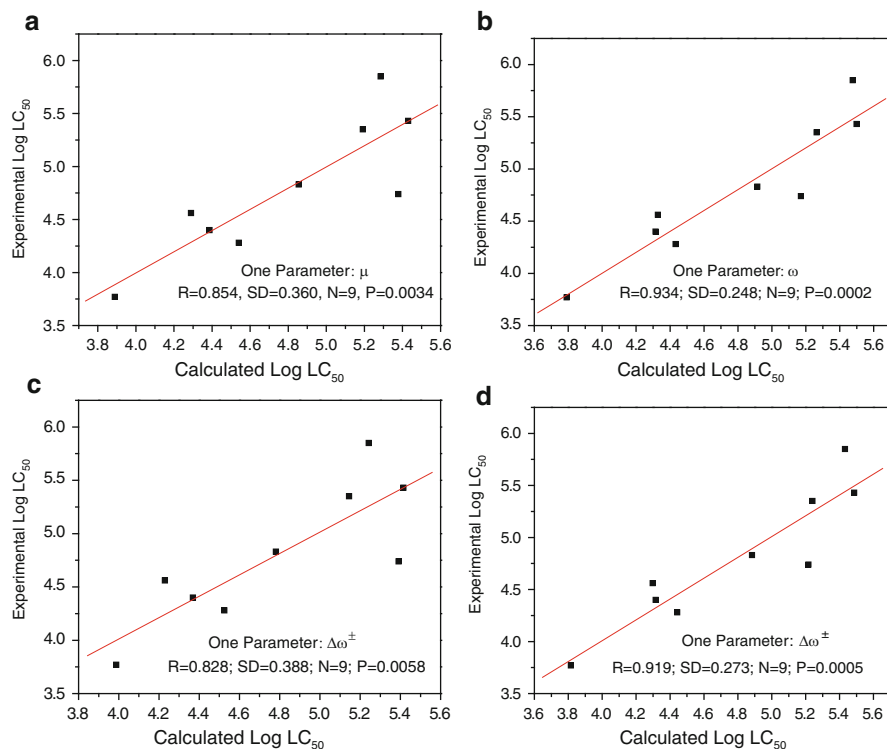


Fig. 4 (a) Experimental versus calculated toxicity (Log (LC₅₀)) values as obtained using one-parameter chemical potential (μ) regression model for the complete set of chlorinated aromatic compounds; (b) same for electrophilicity (ω) index; (c) same for net electrophilicity ($\Delta\omega^{\pm}$) from Eqs. (9), (10) and (13); (d) same for net electrophilicity ($\Delta\omega^{\pm}$) from Eqs. (11), (12) and (13) (Reprinted with permission from [350]. Copyright 2011 IGI Global)

two newly proposed transition metal–borane clusters namely (B₃H₃)₂Ti and (B₃H₃)₂V. In this regard the interaction of these metalloboranes with adenine (dAMP), guanine (dGMP), cytosine (dCMP), and thymine (dTMP) nucleotides are explored here. Figure 5 depicts the optimized geometries of four nucleotides, (B₃H₃)₂Ti, (B₃H₃)₂V, and their corresponding adducts computed at the B3LYP/6-311 + G(d) level of theory.

The interaction energy (ΔE), Gibbs free-energy change (ΔG), reaction enthalpy (ΔH), and reaction electrophilicity ($\Delta\omega$) of all the complexation reactions between the metalloboranes and the nucleotides, provided in Table 2, are negative, which ensures thermodynamic feasibility. The given metalloboranes thus effectively bind with nucleotide moieties and can be conceived as suitable anticancer drugs.

The reaction schemes to derive (B₃H₃)₂Ti and (B₃H₃)₂V from Cp₂Ti²⁺ and Cp₂V²⁺, respectively, (Table 3) also justify the thermodynamic spontaneity of the given processes. Therefore, (B₃H₃)₂Ti and (B₃H₃)₂V, considered as new boron-based anticancer drugs, are supposed to be a possible alternative of Cp₂Ti²⁺ and Cp₂V²⁺, respectively.

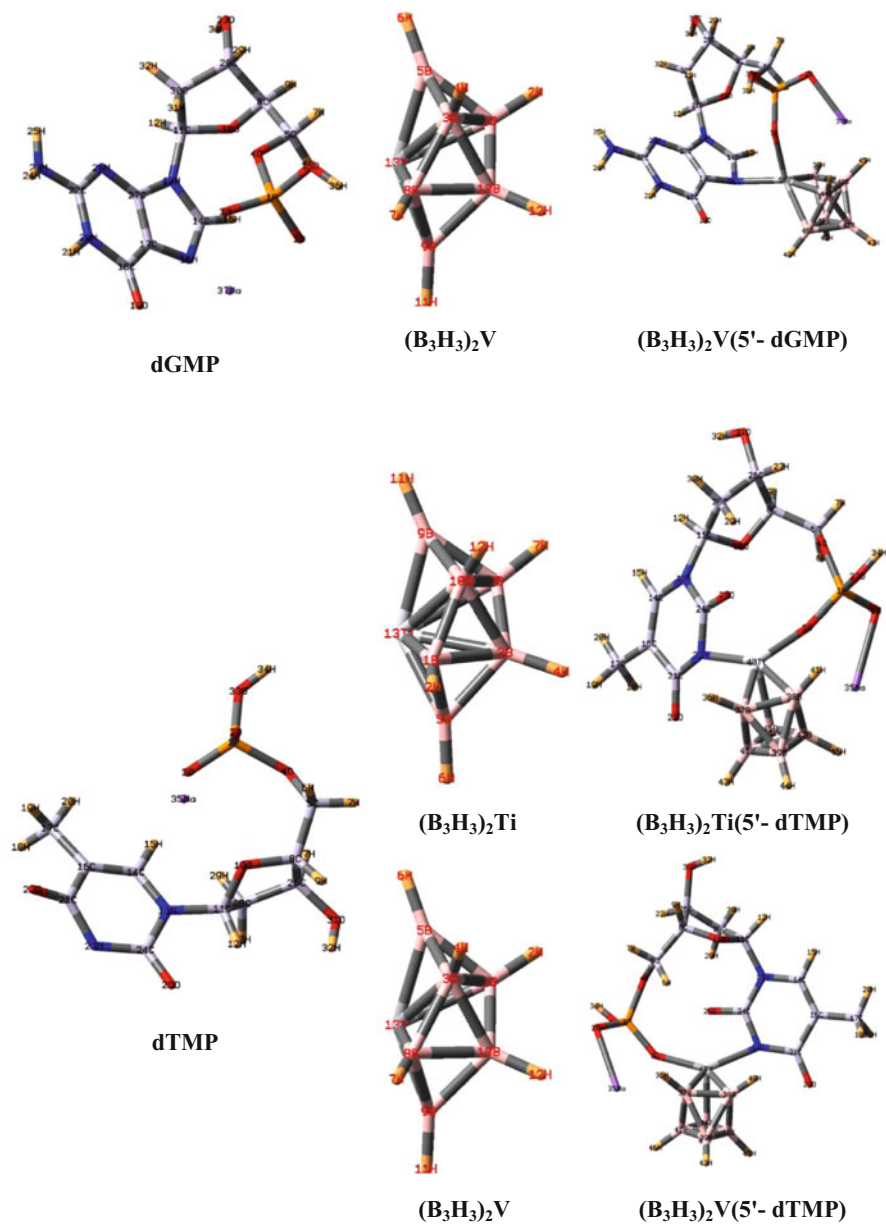


Fig. 5 (continued)

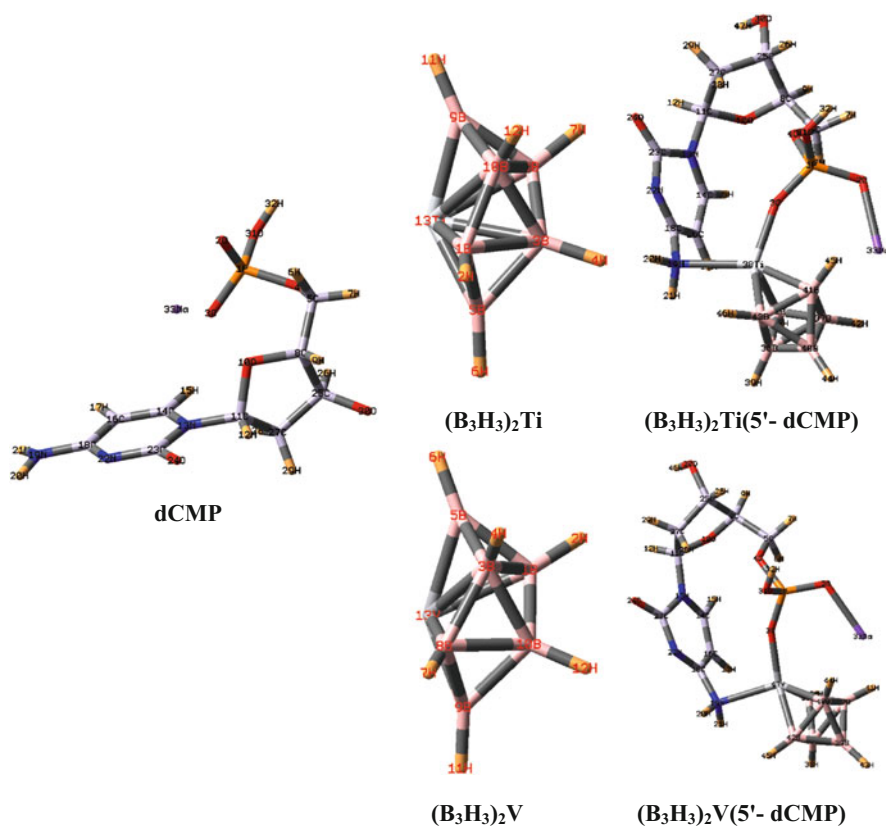


Fig. 5 The optimized geometries of the four nucleotides, $(\text{B}_3\text{H}_3)_2\text{Ti}$, $(\text{B}_3\text{H}_3)_2\text{V}$, and their corresponding adducts at B3LYP/6-311 + G(d) level of theory

Table 2 Interaction energy (ΔE , kcal/mol), Gibbs free-energy change (ΔG , kcal/mol), reaction enthalpy (ΔH , kcal/mol), hardness (η , eV), electrophilicity (ω , eV), and reaction electrophilicity ($\Delta\omega$, eV) of $(\text{B}_3\text{H}_3)_2\text{Ti}$, $(\text{B}_3\text{H}_3)_2\text{V}$, and their corresponding adducts with the four nucleotides at B3LYP/6-311 + G(d) level of theory

| Systems | ΔE | ΔG | ΔH | η | ω | $\Delta\omega$ |
|---|------------|------------|------------|--------|----------|----------------|
| $(\text{B}_3\text{H}_3)_2\text{Ti}$ | | | | 2.865 | 3.787 | |
| $(\text{B}_3\text{H}_3)_2\text{V}$ | | | | 2.810 | 4.290 | |
| $(\text{B}_3\text{H}_3)_2\text{Ti}(5'\text{-dAMP})$ | −89.1 | −72.8 | −86.5 | 2.762 | 2.523 | −2.816 |
| $(\text{B}_3\text{H}_3)_2\text{Ti}(5'\text{-dAMP})$ | −89.7 | −72.5 | −86.9 | 3.218 | 2.465 | −3.375 |
| $(\text{B}_3\text{H}_3)_2\text{Ti}(5'\text{-dGMP})$ | −82.1 | −66.2 | −79.3 | 2.776 | 2.006 | −3.055 |
| $(\text{B}_3\text{H}_3)_2\text{V}(5'\text{-dGMP})$ | −80.9 | −64.5 | −78.1 | 3.211 | 2.008 | −3.554 |
| $(\text{B}_3\text{H}_3)_2\text{Ti}(5'\text{-dTMP})$ | −152.3 | −133.3 | −148.3 | 3.023 | 2.851 | −5.307 |
| $(\text{B}_3\text{H}_3)_2\text{V}(5'\text{-dTMP})$ | −141.7 | −119.3 | −137.1 | 2.906 | 3.754 | −4.906 |
| $(\text{B}_3\text{H}_3)_2\text{Ti}(5'\text{-dCMP})$ | −85.3 | −66.6 | −82.3 | 2.744 | 2.583 | −3.060 |
| $(\text{B}_3\text{H}_3)_2\text{V}(5'\text{-dCMP})$ | −86.9 | −67.3 | −83.8 | 3.107 | 2.607 | −3.537 |

Table 3 Interaction energy (ΔE , kcal/mol), Gibbs free energy (ΔG , kcal/mol), reaction enthalpy (ΔH , kcal/mol), and reaction electrophilicity ($\Delta\omega$, eV) of the studied reaction schemes at B3LYP/6-311 + G(d) level of theory

| Scheme | ΔE | ΔG | ΔH | $\Delta\omega$ |
|---|------------|------------|------------|----------------|
| $\text{Cp}_2\text{Ti}^{2+} + \text{B}_3\text{H}_3^{2-} = (\text{B}_3\text{H}_3)\text{CpTi}^+ + \text{Cp}^-$ | -304.3 | -306.6 | -304.0 | -25.960 |
| $(\text{B}_3\text{H}_3)\text{CpTi}^+ + \text{B}_3\text{H}_3^{2-} = (\text{B}_3\text{H}_3)_2\text{Ti} + \text{Cp}^-$ | -270.7 | -270.6 | -270.4 | -21.008 |
| $\text{Cp}_2\text{V}^{2+} + \text{B}_3\text{H}_3^{2-} = (\text{B}_3\text{H}_3)\text{CpV}^+ + \text{Cp}^-$ | -311.2 | -314.6 | -312.1 | -25.672 |
| $(\text{B}_3\text{H}_3)\text{CpV}^+ + \text{B}_3\text{H}_3^{2-} = (\text{B}_3\text{H}_3)_2\text{V} + \text{Cp}^-$ | -271.7 | -269.9 | -270.8 | -18.095 |

4 Conclusions

A quantitative prediction of molecular behavior based on some known and tested standards is often found to give quite close correlations with allied experimental results. The precise theoretical assessment of a plausible practical thought (molecular property) indeed becomes a useful pathfinder to understand chemical reactivity. The application of this idea to chemistry is abbreviated in many ways, viz. QSAR, QSPR, QSTR, and QSRR. Such methods correlate the changes in molecular structure with chemical behavior like reactivity, toxicity, or any other property to build suitable one or multi-parameter-based regression models for predicting the corresponding parameters of unknown compounds. A comprehensive analysis of the growth of QSAR as a popular predictive algorithm is reported. The roles of various molecular parameters, chemical descriptors in building suitable regression models, and their useful applications towards explaining the biological activity and toxicity of several classes of compounds are mentioned. The crucial roles of QSAR methodology in designing drugs, tumor inhibitors, and in assessment of aquatic toxicity deserve a special mention. A conceptual DFT-based QSAR/QSTR treatment for biomolecules further adds some invaluable insights. Among the various conceptual DFT-based reactivity descriptors, the role of electrophilicity (ω), net electrophilicity ($\Delta\omega^\pm$), and philicity (ω_g^α) concepts are very useful in predicting molecular toxicity. Atom counting as a simple descriptor gives beautiful regression results in some cases. Novel boron-based transition metal clusters as potential anticancer drugs can be theoretically modeled and their further complexations with nucleotide moieties are studied. Boron can be treated as a “new carbon” in drug design. The search for better variables as well as methodologies for an even better prediction of molecular parameters is on.

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DFT Chemical Reactivity Driven by Biological Activity: Applications for the Toxicological Fate of Chlorinated PAHs

Mihai V. Putz and Ana-Maria Putz

Abstract The logistic kinetics model of quantitative reactivity–activity relationships is presented as a modern tool for modeling chemical–biological interactions using catalytic progress curves for the chemical ligand species that bind biological cell receptors. Chemical reactivity principles of electronegativity and chemical hardness, their necessity, the associated scenario of chemical bonding, the critical points in *ab initio* quantum chemical computations of the molecular structures, and the QSAR–OECD principles are reviewed. Illustrative applications are described for chlorinated polycyclic aromatic hydrocarbons (Cl-PAHs) bound to aryl hydrocarbon receptors (AhRs) in human breast cancer MCF-7 cells, *Pimephales promelas* and rats by recognizing key features of *Biological Activity driving Chemical Reactivity* (BioAct-ChemReact) interaction mechanisms. The specific feedback behavior of biological sites against chemical attacks unveil the manner in which chemical reactivity indices and principles of electronegativity and chemical hardness act successively or synergistically (in chemical power and electrophilicity) to regulate chemical reactivity hierarchies in ligand–receptor mechanisms that reflect observed (or *in silico* computed) biological or toxicological effects.

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Contents

| | | |
|-----|--|-----|
| 1 | Introduction | 182 |
| 2 | Survey of Chemical Reactivity Principles | 186 |
| 2.1 | Electronegativity and Chemical Hardness Principles | 186 |
| 2.2 | On the Necessity of the Charge-Parabola for Chemical Energy | 189 |
| 3 | Survey on Computational Quantum Chemistry | 193 |
| 3.1 | Quantum Chemical Theories and Approximations | 193 |
| 3.2 | Self-Consistent Mono-Electronic Orbitals' Equations | 195 |
| 3.3 | Basics of Ab Initio Methods | 198 |
| 3.4 | Hartree-Fock Implementation | 199 |
| 3.5 | Density Functional Theory Implementation | 199 |
| 4 | Logistic Modeling of Chemical-Biological Interactions | 202 |
| 5 | Quantitative Reactivity-Activity Modeling of Chemical-Biological Interactions | 209 |
| 5.1 | On QSAR Principles | 209 |
| 5.2 | In silico-Based Biological Activities | 210 |
| 5.3 | Chemical Reactivity Principles Hierarchy According to the Biological Activity of Cl-PAHs | 211 |
| 6 | Conclusions | 222 |
| | References | 226 |

1 Introduction

In present-day chemistry, and especially in biological chemistry, there is a move toward replacing preindustrial *in vivo* experiments (expensive in materials and time) and *in vitro* experiments (expensive in equipment costs and reproducibility efforts across labs) with *in cerebro* and *in silico* experiments (i.e., evaluation and computation/simulation studies) [1]. The latter experiment types are reasonably inexpensive because they [2]

- Are based on physico-mathematical laws and software implementation
- Provide results on a controlled time scale that are dependent only on the researcher and the computation's theoretical and informatics capabilities
- Are more highly reproducible
- Minimize errors, and most importantly
- Are predictable and reproducible (specific to qualitative and quantitative treatments), considerably reducing the cost and time of validation, especially for newly synthesized compounds, by improving toxicity studies and calculating their impact on biological, ecological, medical, and social environments.

In this context, the quantitative structure-activity relationship (*QSAR*) technique has become the standard European Commission-approved computational/*in silico* method for estimating *in vivo*/*in vitro* experiments regarding chemical toxicity for

environmental hazards in the alimentary, pharmaceutical, somatic, and agricultural industries [3–6]. For this aim to be feasible, conceptual and computational methods for studying both the chemical structure and biological activity aspects of a given compound should be used. Researchers should employ chemical structural information that furthers reactivity-to-activity conceptual–computational knowledge. Fortunately, the current chemical indices developed within density functional theory (DFT) respond to the *conceptual* need because they describe chemical reactivity in terms of observable quantities such as total (or valence) energy and number of electrons (see [7–12]):

- Electronegativity, viewed as an instantaneous variation of total (or valence) energy for the equilibrium (neutral) charged system [13–27]

$$\chi \equiv -\left(\frac{\partial E_N}{\partial N}\right)_{V(\mathbf{r})} \quad (1)$$

- Chemical hardness, viewed as the instantaneous electronegativity change around atoms in a molecule [28–45]

$$\eta \equiv -\frac{1}{2}\left(\frac{\partial \chi}{\partial N}\right)_{V(\mathbf{r})} \quad (2)$$

- Chemical power, providing the *dynamic charge* of atoms in a molecule

$$\pi = \frac{\chi}{2\eta} \quad (3)$$

- Electrophilicity, relating the *promotion energy* of atoms in a molecule [46–51]

$$\omega = \frac{\chi^2}{2\eta} \quad (4)$$

Electronegativity (EL) and chemical hardness (HD) comprise an orthogonal 2D chemical space in which chemical reactivity can be described analogous to physical phenomena in the kinetic space of velocity and acceleration

$$v = \partial r / \partial t \leftrightarrow \partial E / \partial N = -\chi \quad (5)$$

$$a = \partial^2 r / \partial t^2 \leftrightarrow \partial^2 E / \partial N^2 = 2\eta \quad (6)$$

while recognizing the phenomenological equivalence

$$E(\text{valence energy}) \leftrightarrow r(\text{coordinate}) \quad (7)$$

$$N(\text{no. valence } e^-) \leftrightarrow t(\text{time}) \quad (8)$$

Such conceptual orthogonality is apparent when the chemical principles of EL and HD are discussed in the finite difference (gauge ionic species) framework in Sect. 2. It is worth noting that the numerical evaluation of χ , η , ω , and π actually depends on the computational DFT framework of the various energetic states computed. In this way, the *computational* approach is utilized for the chemical reactivity indices aspect, and it will be surveyed in Sect. 3.

The chemical–biological species interaction, on the other hand, currently requires *further conceptual* tools for modeling. The present approach parallels the substrate–enzyme reaction for the consecrated Michaelis–Menten mechanism [52–62] to provide the ligand–receptor (L–R) logistic kinetics in general and the ligand progress curve $L(t)$ in particular, depending on the 50 %-effective concentration (EC_{50}) dose for the recorded activity (A) for a given species:

$$A = \ln\left(\frac{1}{EC_{50}}\right). \quad (9)$$

Because $L(t)$ usually also depends on kinetic parameters such as the initial chemical concentration L_0 and the maximum biological uptake β_{\max} , eventually under the natural exponential form, i.e., the nuclear radioactivity equation—here adapted as

$$L(t) \cong L_0 \exp\left(-\frac{\beta_{\max} t}{EC_{50}}\right) \quad (10)$$

its complete unfolding requires *further computational* ligand–receptor activity modeling. This may be performed by using (QSAR) methodology [63–74], which can be achieved by correlating the observed activity with the above reactivity indices within multilinear regressions yielding computed or predicted activity (A^*). However, for reasons that will be immediately revealed, only linear equations will be considered:

$$A^* = A^*(\chi \vee \eta \vee \pi \vee \omega). \quad (11)$$

Following (9), one can immediately make the correspondence between the initial ligand concentration and computed activity

$$L_0 = \exp(-\|A^*\|) \quad (12a)$$

with $\|\bullet\|$ accounting for the algebraic (Banach) norm in the chemical space (with the dimension equal to the cardinal of the set of chemicals considered in the QSAR), while for the EC_{50} parameter, a similar relationship holds at the level of recorded activities, namely

$$EC_{50} = \exp(-\|A\|). \quad (12b)$$

For maximum biological uptake (via interaction with a ligand), one can employ the working definition

$$\beta = -\frac{d}{dt}L(t) \rightarrow -\frac{L_t - L_0}{t}. \quad (13)$$

With complete consumption of the ligand ($L_t \rightarrow 0$), a maximum value is achieved.

$$\beta_{\max} = \frac{L_0}{\Delta t_{\infty}}. \quad (14)$$

The “infinite” time interval may be shaped by considering the re-scaling first,

$$t = e^{\frac{1}{1-\tau}} - e \rightarrow \begin{cases} 0 \dots \tau \rightarrow 0 \\ \infty \dots \tau \rightarrow 1 \end{cases} \quad (15)$$

followed by differentiation

$$dt = \frac{d\tau}{(1-\tau)^2} \exp\left(\frac{1}{1-\tau}\right). \quad (16)$$

For practical considerations, the interval $(0, \infty)$ projected into $(0, 1)$ can safely use the setting $\tau = 1/2$ as a sufficient condition for maximal biological uptake, so that the associated working time interval (the so-called “*receptor time*”) is $\Delta t_{\infty} = dt(\tau = 1/2, \Delta\tau = 1) = 4e^2$, using Eqs. (12a, 12b) and (14) as the working parameters:

$$\beta_{\max} = \frac{1}{4} \exp(-\|A^*\| - 2) = \frac{L_0}{4e^2}. \quad (17)$$

The working ligand progress curve (10) directly depends on recorded and computed activity, which in turn depend on the chemical reactivity indices considered, employing what can be called *quantitative reactivity–activity relationships* (QRAR). For each DFT framework and each reactivity index, a different progress curve for a given ligand–receptor interaction is obtained. The immediate inference can be made that faster consumption of $L(t)$ is involved for the more reactive index, and therefore, a more preeminent reactivity principle is associated. This allows the researcher

- To formulate chemical–biological interactions with the help of conceptual–computational reactivity indices of electronegativity, chemical hardness, electrophilicity, and chemical power.

- To obtain a hierarchy of the allied chemical principles (electronegativity, chemical hardness, electrophilicity, and chemical power) for a given pool of molecules of certain species and to check whether they are maintained across many species.

This algorithm is schematically presented in Fig. 1. It will be detailed in the next sections by reviewing the basic conceptual and computational concepts invoked here and then illustrated by modeling chlorinated polycyclic aromatic hydrocarbons (Cl-PAHs) on various species (*Pimephales promelas* and *Rattus norvegicus*).

2 Survey of Chemical Reactivity Principles

2.1 Electronegativity and Chemical Hardness Principles

Because the principles of quantum mechanics do not suggest any operator whose eigenvalue is electronegativity, years after Pauling's electronegativity, several definitions and interpretations of electronegativity have been formulated [13–27]. One of most preminent of these was created by Mulliken in 1934 as the average of the ionization potential (*IP*) and electron affinity (*EA*) for the valence state of an atom [14]. This empirical spectroscopic definition dominated chemistry for almost a half-century until its quantitative counterpart was introduced by Parr [7–10, 18, 19] as the additive inverse of the chemical potential of a multi-electronic system. The link between these two definitions is clear if the finite difference approximation is performed on the ground state energy, E_N , around the reference integer total number of electrons N_0 in Eq. (1) to obtain [7–10]:

$$\begin{aligned}\chi &\equiv -\left(\frac{\partial E_N}{\partial N}\right)_{V(\mathbf{r})} \cong -\frac{E_{N_0+1} - E_{N_0-1}}{2} = \frac{(E_{N_0-1} - E_{N_0}) + (E_{N_0} - E_{N_0+1})}{2} \\ &= \frac{\text{IP} + \text{EA}}{2} \cong -\frac{\varepsilon_{\text{LUMO}} + \varepsilon_{\text{HOMO}}}{2}.\end{aligned}\quad (18)$$

The problem with equivalent EN forms in Eq. (18) is with the mixed potential conditions that they imply. Because the EN definition on the left side of the chain equation (18) involves *ground state* energy in a nonzero constant potential $V(\mathbf{r})$, it assumes almost vertical values for the energies when electrons change with environment. On the other hand, the values at the right extreme of Eq. (18), the so-called finite-difference EN, correspond to the *valence state* and are thus characterized by the almost adiabatic case $V(\mathbf{r}) = 0$ because no further electrons are attached to the system. The intriguing problem of treating chemical reactivity by indices defined with ground state vs. valence state has continuously challenged the conceptual DFT community. There is not always clear insight with respect to the chemical phenomenology covered or the viewpoint that requires such clarification, especially with respect to equalization reactivity principles (starting from electronegativity

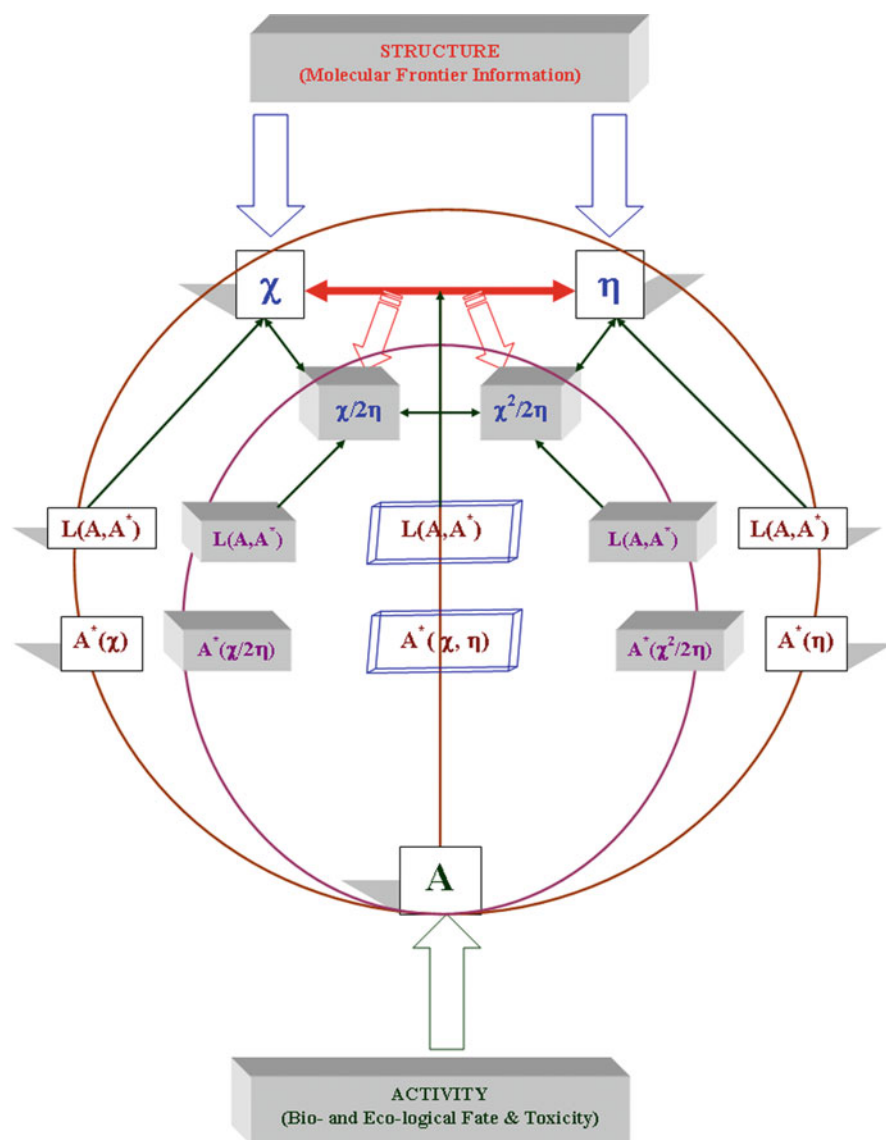


Fig. 1 The workflow of structure–activity relationships involving conceptual and computational density functional theory through molecular frontier information (e.g., HOMO and LUMO), which are primarily employed as electronegativity (χ) and chemical hardness (η) indices and are then combined to form chemical power ($\chi/2\eta$) and electrophilicity ($\chi^2/2\eta$) reactivity measures. These values are correlated with observed biological and ecological activity (A) to provide the QSAR models (A^*) to finally produce ligand progress curves (L) that provide a hierarchy of chemical reactivity principles involved in biological activity within a given DFT computational framework and species of interest

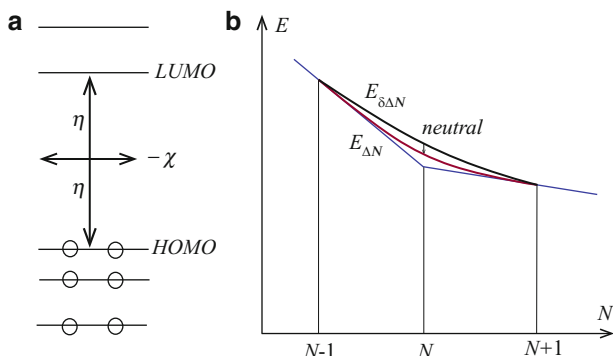


Fig. 2 (a) Orbital energy diagram for a molecule [30, 44] showing electronegativity and hardness according to Eqs. (18) and (19), on which basis the equalization of electronegativity and hardness principles, EE and HSAB, follow (see Table 1). (b) Plot of the electronic energy vs. electrons for a molecule, for which the electronegativity inequality and maximum hardness principles of chemical reactivity, EL and MH, follow from the parabola of energies of the *valence* states (see Table 1)

equalization principles through to charge transfer between adjacent atoms in forming a bond). However, assuming electronegativity is well defined, its natural companion, chemical hardness, can be immediately introduced and developed in the same finite-difference way, as abstracted from Eq. (2) [28]

$$\begin{aligned}\eta &\equiv -\frac{1}{2} \left(\frac{\partial \chi}{\partial N} \right)_{V(r)} = \frac{1}{2} \left(\frac{\partial^2 E_N}{\partial N^2} \right)_{V(r)} \cong \frac{E_{N_0+1} - 2E_{N_0} + E_{N_0-1}}{2} \\ &= \frac{IP - EA}{2} \cong \frac{\varepsilon_{LUMO} - \varepsilon_{HOMO}}{2}.\end{aligned}\quad (19)$$

Like electronegativity [75–80], chemical hardness supports a second-order equalization principle that equalizes the frontier orbitals such that charge transfer is equilibrated between acids and bases in bonding [29, 33, 35, 37]. How this qualitative description of hard-and-soft acids-and-bases (HSAB) [81–87] yields global maximum hardness (MH) is still under debate [88–90], especially when the relationship to electronegativity is invoked. Note that the factor “1/2” in Eq. (19), although considered optional [11, 91], is rooted in parabolic total/valence energy expansion in terms of electronegativity and chemical hardness and describes the fractional quantum distribution of charges in bonding. Only upon averaging do the two adducts with opposite reactivity perspectives compute to unity, as in the classical picture of electronic transfer. Note that the same quantum undulatory information is present (by the appearance of factor “1/2”) in the electrophilicity and chemical power definitions (3) and (4).

In order to form a more intuitive (although not complete) idea of how the above-nominated reactivity principles act at the energetic level, Fig. 2 depicts the equality and inequality variants in the left (a) and right (b) parts, respectively. In Fig. 2a, there is a clear difference during a reaction or bonding process in the electronegativities of the partners first encountered: the main effect of reactivity will be the

adjustment of the middle of the HOMO-LUMO gap so that an easier frontier transfer of the electrons is realized. This behavior illustrates the EE principle on a relative energetic scale. Hard molecules are characterized by a large HOMO-LUMO gap, and soft molecules are characterized by a small gap. Therefore, as a second effect of reactivity, hard-hard and soft-soft interactions are favored over hard-soft and soft-hard interactions because the exchange of the charge induced by the ionic character of bonding is much more easily performed through the first two cases due to nearby energies of the frontier orbitals. This justifies the HSAB principle on a relative energetic scale. Instead, from Fig. 2b, one determines the energetic curve of interaction with the paradigmatic parabolic form, whose justification is based on electronegativity/chemical potential equalization. While approaching the optimum reaction path/parabola achieves a minimum for the neutral collection of nuclei, the electronegativity (slope) at that point should be at a minimum and chemical hardness (the curvature) should be at a maximum to assure the stability of the system in the ground state. A summary of these phenomenological considerations is presented in Table 1, which allows for an understanding of chemical reactivity or chemical bonding as the succession of equal and unequal realizations of chemical indices involved in atoms-in-molecules modeling.

Nothing is specified regarding electrophilicity or chemical power equalization for atoms-in-molecules. Because their basic indices in (3) and (4) are defined as a combination of electronegativity and chemical hardness, it is unclear how to a priori combine their associated principles, i.e., equalization of EL with equalization of HD is one option from Table 1 but not the only one, and the same is true for equalization of EL with the maximum of HD, etc. Therefore, this work explores in an a posteriori (applicative) way the degree to which the electrophilicity principle, for instance, derives (via QRAR modeling technique) from electronegativity- and chemical hardness-based reactivity-activity results. In the systematic affirmative cases (for many species, computation frameworks of structures, etc.), electrophilicity cannot be assessed as carrying the new equalization principle because it is a natural consequence of the EL and HD equalizations (most preeminent in driving chemical reactivity/bonding to its equilibrium with atoms-in-molecules; in all other cases, electrophilicity may provide new insight as an equal stabilization energy for atoms in a molecule). It remains to be seen whether there is an intermediate stage either between the EL and HD actions, or before both of them, in which case it will play a unifying role for the allied principles of electronegativity and chemical hardness, as recently challenged [47–51].

2.2 On the Necessity of the Charge-Parabola for Chemical Energy

Here, we explore the reciprocal case usually treated in chemical reactivity, namely, to derive the energetic shape for chemical valence behavior starting from the equalization of chemical potentials of atoms in molecules [92–95],

Table 1 Synopsis of the basic principles of reactivity at the chemical potential (or electronegativity) and chemical force (or hardness) levels, with both equalization and inequality stages on valence states of atomic, pro-molecule, and atoms-in-molecule systems

| Reactivity/ bonding index | General principle | Special principle | Principle of chemical reactivity/bonding |
|------------------------------|----------------------|----------------------|--|
| $\mu = -\chi$ | $\Delta\mu \geq 0$ | $\Delta\mu = 0$ | <i>Chemical potential (or electronegativity) equality (EE)</i> : “the chemical potential of all constituent atoms in a bond or molecule have the same value” [76] |
| | | $\Delta\mu > 0$ | <i>Chemical potential (or electronegativity) inequality (EI)</i> : “the constancy of the chemical potential is perturbed by the electrons of bonds bringing about a finite difference in regional chemical potential even after chemical equilibrium is attained globally” [77] |
| η | $\Delta\eta \geq 0$ | $\Delta\eta = 0$ | <i>Hard-and-soft acids and bases (HSAB)</i> : “hard likes hard and soft likes soft” [81–83] |
| | | $\Delta\eta > 0$ | <i>Maximum hardness (MH)</i> : “molecules arranges themselves as to be as hard as possible” [88, 89] |

$$\mu_i = \mu_j = \cdots = \mu, \quad (20)$$

as rooted in a well-established thermodynamic (Gibbs) principle of charge flowing toward equilibrium between two charged regions.

When systems are treated quantum-mechanically, especially within DFT, one may assume that the uni-electronic orbitals $\varphi_i, i = \overline{1, N}$ for the N -electronic system relate to the electronic density via spin–orbital occupancies

$$\rho(\mathbf{r}) = \sum_i^N n_i |\varphi_i(\mathbf{r})|^2, \quad 0 \leq n_i \leq 1, \quad \sum_i n_i = N \quad (21)$$

such that within a continuous limit, one recovers the DFT normalization

$$N = \int \rho(\mathbf{r}) d\mathbf{r} = \int \varphi_{\text{KS}}^*(\mathbf{r}) \varphi_{\text{KS}}(\mathbf{r}) d\mathbf{r} \quad (22)$$

involving the Kohn–Sham wave function $\varphi_{\text{KS}}(\mathbf{r})$. It is this wave function that, in equilibrium (dynamic or valence state), corresponds to the global (equalized) chemical potential through the Schrodinger-like equation:

$$\left[-\frac{1}{2} \nabla^2 + V_{\text{eff}}^{\text{DFT}}(\mathbf{r}) \right] \varphi_{\text{KS}}(\mathbf{r}) = \mu \varphi_{\text{KS}}(\mathbf{r}). \quad (23)$$

Equation (23) allows for the expression of chemical potential as an eigenvalue of the N -electronic system

$$\mu = \frac{\int \varphi_{\text{KS}}^*(\mathbf{r}) \left[-\frac{1}{2} \nabla^2 + V_{\text{eff}}^{\text{DFT}} \right] \varphi_{\text{KS}}(\mathbf{r}) d\mathbf{r}}{\int \varphi_{\text{KS}}^*(\mathbf{r}) \varphi_{\text{KS}}(\mathbf{r}) d\mathbf{r}} \equiv \frac{\int \varphi_{\text{KS}}^*(\mathbf{r}) \hat{H} \varphi_{\text{KS}}(\mathbf{r}) d\mathbf{r}}{\int \varphi_{\text{KS}}^*(\mathbf{r}) \varphi_{\text{KS}}(\mathbf{r}) d\mathbf{r}} = \frac{E}{N} \quad (24)$$

in close agreement with Gibbs (grand canonical) formalization. Until now, we have equated chemical potential compatible with Kohn–Sham DFT with the Gibbs definition. If the energy in Eq. (24) is merely activation or free Gibbs energy, it places this phenomenon in the dynamic (and not really ground-state) framework of chemical reactivity. One may call it chemical energy to differentiate it from physical energy and incorporate the effective potential (the electronic correlation) and activation (valence state) information. This allows for consideration of the chemical force expression, analogous to its classical definition, as the minus of the potential gradient while modeling the driving of charge transfer along the reaction path [92–95]:

$$F_\mu = -\frac{d}{dN}(\mu) = \frac{E}{N^2} - \frac{1}{N} \left(\frac{dE}{dN} \right)_V. \quad (25)$$

At this point, by recognizing the electronegativity derivative definition of Eq. (1) as the additive inverse of chemical potential

$$\mu = -\chi = \left(\frac{dE}{dN} \right)_V \quad (26)$$

and also with chemical force

$$F_\chi = -F_\mu = -\frac{d}{dN}(\chi) = \eta^* \quad (27)$$

one immediately arrives at the working equation ($E^* = -E$)

$$E^* = -\chi N + \eta^* N^2 \quad (28)$$

eventually supporting the energy-charge variation form

$$\Delta E = -\chi \Delta N + \eta^* (\Delta N)^2. \quad (29)$$

Note that Eq. (29) differs from conceptual DFT

$$E[N + \Delta N] = E_0[N] + \left(\frac{\partial E}{\partial N} \right)_V \Delta N + \frac{1}{2} \left(\frac{\partial^2 E}{\partial N^2} \right)_V (\Delta N)^2 + \dots \quad (30)$$

by a factor of “1/2” that comes from the undulatory nature of the charges themselves, a feature not taken into account when considering the classical-to-quantum analogy in Eq. (27). The correct classical-to-quantum correspondence should

account for the factor “1/2” on the quantum side due to the electronic self-interaction that should be smeared out in analytical considerations.

In this context, the interaction energy $E_{\Delta N}$ of an electronic system that has a change in charge ΔN assumes the paradigmatic parabolic analytical form:

$$E_{\Delta N} = E_0 + \mu_1 \Delta N + \frac{1}{2} \eta_1 (\Delta N)^2. \quad (31)$$

Let us now assume that energy in Eq. (31) is associated with the minimum perturbation to produce the chemical reaction or molecular (trans-)formation. If further perturbation is considered with respect to the change in charge ($\delta \Delta N$ with the small quantity $\delta \in [0, 1]$), another reaction path unfolds through expansion [44]:

$$E_{\delta \Delta N} = E_{0/v} + \mu_1 \delta \Delta N + \frac{1}{2} \eta_1 (\delta \Delta N)^2 \equiv E_{0/v} + \mu_2 \Delta N + \frac{1}{2} \eta_2 (\Delta N)^2, \quad (32)$$

where $\mu_2 = \mu_1 \delta$ and $\eta_2 = \eta_1 \delta^2$ are the new driving chemical potential and force, respectively. Searching for the naturally selected conditions for optimum reactivity, we can immediately observe that as the variational principle is applied to chemical reactivity, it demands a restoring path from interaction energy [Eq. (32)] back to [Eq. (31)] (see also Fig. 2b)

$$E_{\delta \Delta N} \rightarrow E_{\Delta N} \rightarrow \text{minimum} \quad (33)$$

as the differences in their slope and curvature achieve maximum values

$$\Delta \mu = \Delta \mu_{\delta \Delta N \rightarrow \Delta N} = \mu_1 - \mu_2 = \mu_1 (1 - \delta) \geq 0, \quad (34)$$

$$\Delta \eta = \Delta \eta_{\delta \Delta N \rightarrow \Delta N} = \eta_1 - \eta_2 = \eta_1 (1 - \delta^2) \geq 0 \quad (35)$$

with the specialization and chemical reactivity rules listed in Table 1. The major conclusion is that all principles of chemical reactivity, especially those of chemical hardness, are rooted in the chemical potential equalization principle, though they are not necessarily identical nor do they describe the same stage of bonding. This becomes clear when one translates Table 1 into chemical bonding scenarios through the following four stages [45]:

1. *Encountering stage*: dominated by the difference in electronegativity between reactants and consumed when the electronegativity equalization principle is fulfilled among all product constituents. This stage is associated with charge flow from the more electronegative regions to the lower electronegativity regions in a molecular formation covering the *covalent* binding step.
2. *Promolecule*: even after chemical equilibrium is attained globally, the electrons involved in bonds act as foreign objects between pairs of regions at any level of the molecular partitioning procedure, inducing the appearance of a finite

difference in adjacent electronegativity of neighbor regions in the molecule. This appearance is due to quantum fluctuations associated with the quantum nature of the bonding electrons and corresponds to ionic bond character.

3. *Atoms-in-molecule*: the induced ionic character of bonds is partially compensated by chemical forces through hardness equalization between the pair regions in a molecule. HSAB principles are therefore involved as a second-order effect in charge transfer. See expansion (30) for an example driven by ionic interactions through bonds.
4. *Ω -molecule*: quantum fluctuations provide a further amount of finite difference, this time in attained global hardness that is transposed onto relaxation effects among the nuclear and electronic distributions so that the remaining unsaturated chemical forces are dispersed by stabilization of the molecular structure.

In this way, we have conceptually surveyed how chemical reactivity in general and chemical bonding in particular may be modeled through a minimum set of electronic indices of electronegativity and chemical hardness, which are complementary in their role in promoting and stabilizing electronic structures in chemical combinations.

3 Survey on Computational Quantum Chemistry

3.1 *Quantum Chemical Theories and Approximations* [45]

Quantum chemistry originates in five levels of quantum approximations imposed on many-electron-many-nuclei systems, either in isolated or interacting states. They are summarized below with mentions of the current limitations, controversies and prospects.

1. *The Born–Oppenheimer approximation* [96], created to simplify the electronic calculus for frozen nuclei approximation, breaks down when computing, for instance, the magnetic dipole moment and its derivative with respect to the nuclear velocities or momenta for assessing the molecular properties of surfaces [97].
2. *The single Slater determinant representation of the ground electronic state* [98] elegantly solves the exchange behavior of electrons by incorporating Pauli repulsion in antisymmetric determinants [99] and was conceptually extended to the configuration interaction by the seminal works of Löwdin [100]. Further generalization was later noted in the valuable concept of the so-called “complete active space” (CAS) [101, 102] that, when combined with other quantum chemical methods such as the self-consistent field (SCF) and density functional theory (DFT), becomes very productive in accounting for *all* electronic states that contribute to the *reactive* space, whether they are electronic states of species

(reactants, intermediates, and products) that are involved in chemical reactions or thermally or photo-induced states [103–105].

3. *Simple Hückel* [106, 107] and *molecular orbital theories* [108–115], which are viewed as the next natural step over the paradigmatic Heitler–London theory of homopolar chemical bonding [116], have been making possible the development of self-consistent field Hartree–Fock–Slater theories [117–122] and associated semi-empirical formulations [123–126] for treating a plethora of chemical systems and phenomena on the basis of their internal symmetry, while remarkably agreeing with (and sometimes predicting) observed spectra and reactivity. Pericyclic reactions [127, 128] and the Woodward–Hoffman rules [129, 130] are eminent examples. This method is met, however, with the so-called quantum correlation problem (i.e., modeling the electronic movement in the dynamical field of the other electrons present in the system) that remains intractable within the Slater (or even with configurational interaction) framework. The solution to obtaining accurate correlations arrives with the advent of Density Functional Theory (DFT) but with the price of modifying the overall wave-function of the system and its spectra (see next).
4. *Thomas–Fermi* [131–135] and *Hohenberg–Kohn–Sham* [136, 137] theories, which merged into the celebrated Density Functional Theory [7–9, 11, 12, 138–144], are conceptually exact, i.e., they perform *ab initio* analysis of the electronic spectra relying only upon universal constants of electronic charge, mass, and Planck constant. Their combinations of bare and effective potential, while providing an approximate set of orbitals (called Kohn–Sham orbitals) correctly resemble the observed electronic density of the system along the measured energies (with correlation effects included) but with less significance than the classical wave-function concept. In fact, the current DFT uses the so-called basis function like a mathematical tool that can be adapted or optimized depending on the accuracy needed in relation to the optimized effective potential [145], adding dispersion effects, etc. The computational implementation of DFT becomes a parameterized procedure that makes it a sort of semi-empirical DFT [146] that can be extended to include time-dependency excited state effects [147–152] or even model Bose–Einstein condensates [12]. The great merit and paradox of DFT is that the theory provides the recipe to compute two- or multi-body interactions as exchange and correlations, respectively, by approximations to single body (density) behavior. From a theoretical physics point of view, the picture is flawed, yet the approximation works very well. This landmark achievement was perhaps awarded the Nobel Prize in Chemistry (in 1998) rather than Physics because it involves no new physics but represents a useful reformulation for Chemistry. Recent DFT work further reduces the limits of DFT, formulating various approximations for exchange-correlation functionals [92] that are more or less in agreement with the fundamental theorems and limits at asymptotic and nuclei ranges [153]; this holds until attempting to formulate expectation values of various physical observables based only on density, similar to those based on the expectation value of quantum mechanical wave functions [154]. True *ab initio* DFT is far

from being fully engaged [155, 156]; *semi-empirical DFT* seems to prevail in computational implementation. Still, conceptual DFT [157] is of upmost importance in formulating chemical reactivity and the indices that help in understanding and modeling the chemical systems [11, 91].

5. *Solvent effects* must almost always be taken into account when using quantum chemical treatment to describe the reactivity of chemical systems. The environment, including any strongly interacting solvent molecules (e.g., water molecules in the case of biomolecules, such as amino acids, peptides, nucleic acids and their complexes) must be considered in any modeling study of open chemical systems in order to fully understand and interpret the experimental results, such as vibrational, NMR and electronic spectra, and the chiral analogues [158]. Overall, the interaction of the system with the environment stands in the foreground of quantum theory when predicting additional quantum fluctuations upon the concerned system due to coupling with the media/observer/solvent [159, 160]. Such interactions can be nevertheless implemented by accounting for additional reactions and the stability of the investigated chemical systems, while also having the quantum statistical tools for treating the macro-canonical samples in a correct physical way. DFT is well equipped from its basic definition of density (associated with the total number of electrons in the system) that can be easily extended to include environmental effects [161–163].

Overall, the quantum many-body theory (i.e., the second quantization of fields) and its “chemical” version as DFT may be regarded as the appropriate quantum tools. Quantum chemistry should be used in formulating and implementing the specific indicators of stability and reactivity, among which electronegativity and chemical hardness are the most preminent and versatile.

3.2 Self-Consistent Mono-Electronic Orbitals’ Equations [164]

Following Dirac’s quote, once the Schrodinger equation:

$$H\Psi = E\Psi \quad (36)$$

is established “the underlying physical laws necessary for the mathematical theory of a large part of physics and the whole of chemistry are thus completely known” [165].

Unfortunately, the molecular spectra based on the eigen-problem (36) are neither directly nor completely solved without specific atoms-in-molecule and/or symmetry constraints and approximation. As such, at the mono-electronic level of approximation, the Schrodinger equation (36) is rewritten under the so-called “independent-electron problem”:

$$H_i^{\text{eff}}\psi_i = E_i\psi_i \quad (37)$$

with the aid of effective electron Hamiltonian partitioning:

$$H = \sum_i H_i^{\text{eff}} \quad (38)$$

and the corresponding molecular mono-electronic wave-functions (orbitals) fulfilling the conservation rule of probability:

$$\int \psi_i^2(\mathbf{r}) d\mathbf{r} = 1. \quad (39)$$

When written as a linear combination over the atomic orbitals, the resulting MO-LCAO wave-function:

$$\psi_i = \sum_v C_{vi} \phi_v \quad (40)$$

is replaced in Eq. (37), and following integration over the electronic space, it allows for matrix version of Eq. (37):

$$(H^{\text{eff}})(C) = (S)(C)(E). \quad (41)$$

With the diagonal energy-matrix elements as the eigen-solution

$$(E)_{ij} = E_{ij} = E_i \delta_{ij} = \begin{cases} E_i \dots i = j \\ 0 \dots i \neq j \end{cases} \quad (42)$$

to be found in terms of the expansion coefficients matrix (C), the matrix of the Hamiltonian elements:

$$H_{\mu\nu} = \int \phi_\mu H^{\text{eff}} \phi_\nu d\tau \quad (43)$$

and the matrix of the (atomic) overlapping integrals:

$$S_{\mu\nu} = \int \phi_\mu \phi_\nu d\tau \quad (44)$$

where all indices in Eqs. (42)–(44) refer to matrix elements because the additional reference to the “ i ” electron was skipped to avoid confusion.

The solution of the matrix equation (41) may be unfolded through the Löwdin orthogonalization procedure [166, 167], involving the diagonalization of the overlap matrix via a given unitary matrix (U), $(U)^\dagger(U) = (1)$, using the following procedure:

$$(s) = (U)^+(S)(U), \quad (45)$$

$$\left(s^{-1/2}\right)_{ii} = [(s)_{ii}]^{-1/2}, \quad (46)$$

$$\left(S^{-1/2}\right) = (U)\left(s^{-1/2}\right)(U)^+, \quad (47)$$

$$\left(\left(S^{1/2}\right)(C)\right)^+ \left(\left(S^{-1/2}\right)(H^{\text{eff}})\left(S^{-1/2}\right)\right) \left(\left(S^{1/2}\right)(C)\right) = (E). \quad (48)$$

However, the solution given by Eq. (48) is based on the form of effective independent-electron Hamiltonians that can be empirically constructed as in Extended Hückel Theory [168]. Such “arbitrariness” can be nevertheless avoided by the so-called *self-consistent field* (SCF), in which the one-electron effective Hamiltonian is considered to depend on the solution of Eq. (40) itself, i.e., the matrix of coefficients (C). The resulting “Hamiltonian” is called the Fock operator, while the associated eigen-problem is the Hartree–Fock equation:

$$F\psi_i = E_i\psi_i. \quad (49)$$

In matrix representation, Eq. (49) looks like:

$$(F((C)))(C) = (S)(C)(E) \quad (50)$$

and may be solved iteratively through a diagonalization procedure starting from an input (C) matrix or from a starting electronic distribution quantified by the density matrix:

$$P_{\mu\nu} = \sum_i^{\text{occ}} C_{\mu i} C_{i\nu} \quad (51)$$

with a major influence on the Fock matrix elements:

$$F_{\mu\nu} = H_{\mu\nu} + \sum_{\lambda\sigma} P_{\lambda\sigma} \left[(\mu\nu|\lambda\sigma) - \frac{1}{2}(\mu\lambda|\nu\sigma) \right]. \quad (52)$$

Note that the one-electron Hamiltonian effective matrix components $H_{\mu\nu}$ differ from those of Eq. (43) in what they truly represent. In this form, it represents the kinetic energy plus the interaction of a single electron with the core electrons around all nuclei present. The other integrals appearing in Eq. (52) are generally called two-electron-multi-centers integrals and are written as:

$$(\mu\nu|\lambda\sigma) = \int \phi_\mu^A(\mathbf{r}_1)\phi_\nu^B(\mathbf{r}_1)\frac{1}{r_{12}}\phi_\lambda^C(\mathbf{r}_2)\phi_\sigma^D(\mathbf{r}_2)d\mathbf{r}_1d\mathbf{r}_2. \quad (53)$$

From definition (53), there is immediate recognition of the special integral $J = (\mu\mu|\nu\nu)$ as the Coulomb integral describing repulsion between two electrons with probabilities ϕ_μ^2 and ϕ_ν^2 .

Moreover, the Hartree–Fock Eqs. (50) implemented in Eqs. (51) and (52) are known as Roothaan equations [109] and constitute the basis for closed-shell (or restricted Hartree–Fock, RHF) molecular orbitals calculations. Their extension to spin effects provides the equations for the open shell (or unrestricted Hartree–Fock, UHF), which are also known as the Pople–Nesbet Unrestricted equations [118].

3.3 Basics of *Ab Initio* Methods [164]

The alternative to semi-empirical methods is the full self-consistent calculation, the so-called *ab initio* approach. It is based on computing all integrals in Eq. (52), with the atomic Slater-type orbitals (STO), $\exp(-\alpha r)$, replaced by Gaussian-type orbitals (GTO) [169]:

$$\phi_A^{\text{GTO}} = x_A^l y_A^m z_A^n \exp(-\alpha r_A^2) \quad (54)$$

in molecular orbitals expansion, a procedure allowing for much simplification in multi-center integrals computation. Nevertheless, in turn, each GTO may be generalized to a contracted expression constructed upon the primitive expressions of Eq. (54):

$$\phi_\mu^{\text{CGTO}}(r_A) = \sum_p d_{p\mu} \phi_p^{\text{GTO}}(\alpha_p, r_A) \quad (55)$$

where α_A and $d_{p\mu}$ are the exponents and the contraction coefficients of the primitives, respectively. Note that the primitive Gaussians involved may be chosen as approximate Slater functions [170], Hartree–Fock atomic orbitals [171], or any other set of functions desired to speed up computation. Under these conditions, a minimal basis set may be constructed with one function for H and He, five functions for Li and Ne, nine functions for Na and Ar, 13 functions for K and Ca, 18 functions for Sc to Kr, etc., to describe the core and valence occupancies of atoms [172–174]. Although such a basis does not generally provide accurate results because of its small cardinal, it contains the essential information regarding the chemical bond and may be useful for qualitative studies, as is the present case for aromaticity scales where the comparative trend is studied.

3.4 Hartree–Fock Implementation [164]

When simple ab initio methods are referred to, it means that the Hartree–Fock equation (49) with full Fock matrix elements [117, 175–177] of Eqs. (51) and (52) is solved for a Gaussian-contracted basis [Eq. (55)]. In fact, the method iteratively evaluates the kinetic energy and nuclear–electron attraction energy integrals for the effective Hamiltonian along the overlap and electron–electron repulsion energy integrals (for both the Coulomb and exchange terms), which are written as:

$$T_{\mu\nu} = \langle \mu | \left(-\frac{1}{2} \nabla^2 \right) | \nu \rangle, \quad (56)$$

$$V_{\mu\nu} = \langle \mu | \frac{Z_A}{r_A} | \nu \rangle, \quad (57)$$

$$S_{\mu\nu} = \langle \mu | \nu \rangle, \quad (58)$$

$$(\mu\nu|\lambda\sigma) = \left(\mu\nu | \frac{1}{r_{12}} | \lambda\sigma \right), \quad (59)$$

until consistency in the electronic population of Eq. (51) between two consecutive steps is achieved.

Note that such calculations assume the total wave function as a single Slater determinant, while the resultant molecular orbital is described as a linear combination of the atomic orbital basis functions (MO-LCAO). Multiple Slater determinants in MO description project the configurational and post-HF methods and will not be discussed here.

3.5 Density Functional Theory Implementation [164]

The main weakness of the Hartree–Fock method, namely the lack in correlation energy, is ingeniously restored by the Density Functional method through the introduction of the so-called “effective one-electron exchange-correlation potential.” Unfortunately, this means that the analytical form will not be determined. However, the working equations have the simplicity of the HF versions, while replacing the exchange term in Eq. (52) by the exchange-correlation (“XC”) contribution; this results in the (general) unrestricted matrix form of the Kohn–Sham equations [137]:

$$F_{\mu\nu}^{\dagger} = H_{\mu\nu}^{\dagger} + \sum_{\lambda\sigma} P_{\lambda\sigma}^T (\mu\nu|\lambda\sigma) + F_{\mu\nu}^{\text{XC}\dagger}, \quad (60)$$

$$F_{\mu\nu}^{\downarrow} = H_{\mu\nu}^{\downarrow} + \sum_{\lambda\sigma} P_{\lambda\sigma}^T (\mu\nu|\lambda\sigma) + F_{\mu\nu}^{\text{XC}\downarrow}, \quad (61)$$

$$P^T \equiv P^{\uparrow\downarrow} = P^{\uparrow} + P^{\downarrow} \quad (62)$$

in a similar fashion as with the Pople–Nesbet equations of Hartree–Fock theory. The restricted (closed-shell) variant resembles the density constraint:

$$\rho^{\uparrow} = \rho^{\downarrow} \quad (63)$$

in which case the Roothaan analogous equations (for exchange–correlation potential) are obtained.

Either Eq. (60) or (61) fulfills the general matrix equation of type (50) for the energy solution

$$E = \sum_{\mu\nu} P_{\mu\nu} H_{\mu\nu} + \frac{1}{2} \sum_{\mu\nu\lambda\sigma} P_{\mu\nu} P_{\lambda\sigma} (\mu\nu|\lambda\sigma) + E_{\text{XC}}, \quad (64)$$

which can be regarded as the solution of the Kohn–Sham equations. The apparent exchange–correlation energy E_{XC} may be conveniently expressed through energy density (per unit volume) using the integral formulation:

$$E_{\text{XC}} = E_{\text{XC}}[\rho^{\uparrow}, \rho^{\downarrow}] = \int f(\rho^{\uparrow}, \rho^{\downarrow}) d\tau \quad (65)$$

once the Fock elements of exchange–correlation are recognized to be of a density gradient form [178]:

$$F_{\mu\nu}^{\text{XC}\uparrow(1)} = \int \frac{\partial f}{\partial \rho^{\uparrow(1)}} \phi_{\mu} \phi_{\nu} d\tau. \quad (66)$$

The quest for various approximations for the exchange–correlation energy density $f(\rho)$ has spanned decades in quantum chemistry and was recently reviewed [92]. Here, we will present the “red line” of its implementation, as it will be further used for the current applications. The benchmark density functional stands as the Slater exchange approximation, derived within the so-called “X α theory” [179]:

$$f^{\text{X}\alpha} = -\frac{9}{4}\alpha \left(\frac{3}{4\pi}\right)^{1/3} \left(\rho^{\uparrow 4/3} + \rho^{\downarrow 4/3}\right) \quad (67)$$

with α taking the values:

$$\alpha = \begin{cases} 1 \dots \text{Slater} \\ 2/3 \dots \text{uniform electronic gas} \end{cases} \quad (68)$$

With Eq. (67) in Eq. (66), the resulting Kohn–Sham “exchange-correlation” matrix elements (although rooted only in the exchange) yield the integral representation:

$$F_{\mu\nu}^{\text{XC}\uparrow} = F_{\mu\nu}^{\text{X}\uparrow} = -\frac{9}{4\pi} \alpha \int \rho^{\uparrow 1/3} \phi_{\mu} \phi_{\nu} d\tau. \quad (69)$$

Considerable improvement of molecular calculation was given by Becke’s density gradient correction of the local spin density (or Slater exchange) approximation of the exchange energy [180]:

$$E_{\text{X}} = \sum_{\sigma=\uparrow,\downarrow} \int e_{\text{X}\sigma}^{\text{LSDA}} g_{\text{X}\sigma} d\tau \quad (70)$$

where

$$g_{\text{X}\sigma} \left(\frac{|\nabla \rho^{\sigma}|}{\rho^{\sigma 4/3}} \right) = g_{\text{X}\sigma}(x) = \begin{cases} 1 \dots \text{Slater} \\ 1 + \frac{bx^2}{a(1 + 6bx \sinh^{-1} x)} \dots \text{Becke88} \end{cases} \quad (71)$$

with the parameters $a = (3/2)(3/4\pi)^{1/3}$ and $b = 0.0042$ chosen to fit the experiment. Other exchange functionals were developed along the same line, having different realization of the gradient functions [Eq. (71)], most notably those of Perdew and collaborators (e.g., Perdew–Wang-91, PW91) [181].

The correlation contribution was developed using a somewhat different algorithm, namely, employing its definition as the difference between the exact and Hartree–Fock (HF) total energy of a poly-electronic system [182]. Without reproducing the results (more details are provided in the dedicated review of [92]), we mention only the Lee–Yang–Parr (LYP) correlation functional [183–185] along the Vosko–Wilk–Nusair (VWN) local correlation density functional [186].

The exchange and correlation density functionals combine into so-called “hybrid functionals.” Those used in the present study refer to

- Becke-97: a hybrid exchange-correlation functional created by extending the $g(x)$ of Equation (71) as a power series containing three terms with an admixture of 19.43 % HF exchange [187].
- Becke88-VWN: combines Becke88 exchange of Eq. (71) with VWN correlation.
- B3-PW91: also developed by Becke with PW91 correlation instead of LYP.
- B3-LYP: advanced by Becke by empirical comparisons against very accurate results; it contains the exchange contribution (20 % HF + 80 % Slater + 72 % Becke88) added to the correlation energy (81 % LYP + 19 % VWN) [188].

These are the main methods, at both conceptual and computational levels, used to assess the structure of molecules involved as ligands in various chemical–biological interactions.

4 Logistic Modeling of Chemical–Biological Interactions

In the post-genomic era, the development of kinetic models that allow simulation of complicated metabolic pathways and protein interactions is becoming increasingly important [189, 190]. Unfortunately, the difference between an in vivo biological system and homogeneous in vitro conditions is large, as shown by Schnell and Turner [191]. Mathematical treatments of biochemical kinetics have been developed from the law of mass action in vitro, but the modifications required to bring them in line with stochastic in vivo situations are still under development [192–194].

The mechanism of the biological activity produced by a substance usually involves the combination of interactions between the molecules of that substance, called the effector or ligand (L), with a receptor (R), a protein, a biologically macromolecule, or a complex of macromolecules within the cell. The intensity of the biological action is illustrated in an ordinary way as the logarithm of the inverse of the concentration C to produce a specific biological answer (see Eq. (9)). Because the C_{50} concentration (the molar concentration that produces 50 % of the maximum biological activity) is used in many situations, it can be shown that the biological activity A is proportional to the affinity of the molecule or ligand L_i for the receptor R , which is at the basis of the explicated biological action. However, the most rational hypothesis concerning the mode of action of bioactive substances presumes that biological activity produced by a ligand L is proportional to the complexation degree of the receptor R with L . In this situation, the presumed biological activity of $\alpha\%$ (compared to a maximum of 100 %) is produced by the concentration $[L]$ of ligand, which closely agrees with substrate–enzyme kinetics modeling (see Table 2).

Therefore, we may safely generalize the chemical–biological kinetics in terms of the general rate of biological uptake β while respecting the chemical concentration $[L]$

$$\beta = \frac{\beta_{\max}[L]}{[L] + EC_{50}} \quad (72)$$

while recognizing that the temporal link in Eq. (72) between the biological activity and chemical concentration is represented as

$$\beta = -\frac{d}{dt}[L](t). \quad (73)$$

The full temporal version can be widely formulated as [59–62]:

$$-\frac{d}{dt}[L](t) = \frac{\beta_{\max}[L](t)}{[L](t) + EC_{50}}. \quad (74)$$

The main problem with Eq. (74) is that it accounts only for the velocity of the initial time of the reaction. The information outside the first moments of the inherent progress curve is virtually lost or neglected. Another complication of Eq. (74) is that, even when describing a generalized kinetic, it differs from ordinary chemical curves in its rectangular hyperbola shape instead of the expected exponential form. A further generalized kinetic may instead be assumed, which can be applied to the enzymatic Michaelis-Menten case [196–200] as follows.

We use a probabilistic approach [197, 198], based on the law of mass action, to characterize in vitro ligand-receptor interaction as quoted in Table 2:

$$1 = P_{\text{REACT}}([L]_{\text{bind}}) + P_{\text{UNREACT}}([L]_{\text{bind}}). \quad (75)$$

In Eq. (75), $P_{\text{REACT}}([L]_{\text{bind}})$ is the probability that the ligand–receptor interaction of Table 1 proceeds at a certain concentration of ligand binding to the receptor $[L]_{\text{bind}}$. The limits are:

$$P_{\text{REACT}}([L]_{\text{bind}}) = \begin{cases} 0, & [L]_{\text{bind}} \rightarrow 0 \\ 1, & [L]_{\text{bind}} \gg 0. \end{cases} \quad (76)$$

Note that $P_{\text{REACT}}([L]_{\text{bind}}) = 0$ when the enzymatic reaction does not proceed or when it stops because the ligand fails to bind or is entirely metabolized.

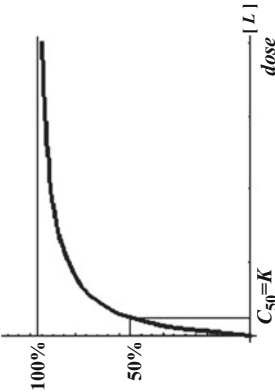
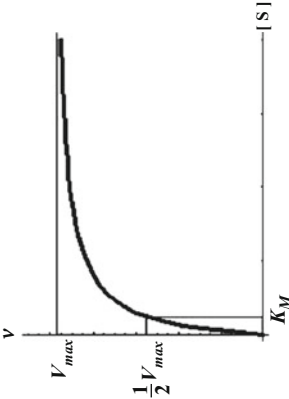
Conversely, $P_{\text{REACT}}([L]_{\text{bind}}) = 1$ when the ligand–receptor interaction proceeds and is related to the standard quasi-steady-states approximation (QSSA). The probability of the occurrence of products in L–R reactions lies between these limits. Similarly, in the case where specific interactions do not take place, $P_{\text{UNREACT}}([L]_{\text{bind}})$, the limits are:

$$P_{\text{UNREACT}}([L]_{\text{bind}}) = \begin{cases} 1, & [L]_{\text{bind}} \rightarrow 0 \\ 0, & [L]_{\text{bind}} \gg 0. \end{cases} \quad (77)$$

This probabilistic treatment of enzyme kinetics is based on the chemical bonding behavior of enzymes that act upon substrate molecules through diverse mechanisms and may offer the key to the quantitative treatment of different types of enzyme catalysis.

To unpack the terms of Eq. (75) to analyze L–R reactions, we first recognize that the bound ligand concentration can be treated as the instantaneous concentration, i.e., $[L]_{\text{bind}} = [L](t)$.

Table 2 The face-to-face ligand–receptor (L–R) and substrate–enzyme (S–E) kinetics [195]

| Property | L–R kinetics | S–E kinetics |
|-----------|--|--|
| Reaction | $R + L \overset{K}{\rightleftharpoons} C^*$ | $E + S \overset{k_1}{\underset{k_{-1}}{\rightleftharpoons}} ES \overset{k_2}{\rightarrow} E + P$ |
| Species | R | E |
| Chemicals | L | S |
| Equation | $\frac{\alpha}{100} = \frac{[L]}{[L] + K}$ | $v = \frac{[S]}{[S] + K_M}$ |
| Constant | $K = \frac{[R][L]}{[C^*]}$ | $K_M = \frac{k_{-1} + k_2}{k_1}$ |
| Graph |  |  |

Maintaining quasi-steady-state conditions for in vitro systems, we may assume constant association–dissociation rates so that probability of interaction/reaction is written as the rate of consumption of the ligand [see Eq. (73)], to saturation:

$$P_{\text{REACT}}([L](t)) = \frac{\beta(t)}{\beta_{\text{max}}} = -\frac{1}{\beta_{\text{max}}} \frac{d}{dt}[L](t) \quad (78)$$

after the initial transient of the ligand–receptor adduct-complex interchanges.

We know only that expression (78) behaves like a probability function, with values in the realm [0,1]. Given expressions (75), (78) and the general Michaelis-Menten equation (74), we derive an expression for the unreacted probability term, $P_{\text{UNREACT}}([S](t))$. As such, the expression:

$$P_{\text{UNREACT}}([L](t))^{\text{MM}} = \frac{EC_{50}}{[L](t) + EC_{50}} \quad (79)$$

satisfies all probability requirements, including the limits in Eq. (77). When combined with equations (75) and (78), the equation gives the instantaneous version of the classical Michaelis-Menten equation (72). Remarkably, expression (79) can be seen as a generalization of the efficiency of the Michaelis-Menten reaction under steady-state conditions. The efficiency depends on two parameters: EC_{50} , which embodies the toxicological conditions of the ligand–receptor reaction, and the initial ligand concentration $[S_0]$; these determine the ratio of free to total ligand concentration in the L–R interaction. That is, when efficiency is equal to one, we do not expect to find free ligand in the reaction; the L–R reactions are all consumed so that the first branch of the limit (77) is fulfilled and no further binding will occur. It is clear that the Michaelis-Menten term [Eq. (79)] is just a particular choice for a probabilistic ligand kinetic model of the conservation law [Eq. (75)]. However, a more generalized version of equation (79) that preserves all of the above probabilistic features may look like

$$P_{\text{UNREACT}}([L](t))^* = e^{-\frac{[L](t)}{EC_{50}}} \quad (80)$$

from which the Michaelis-Menten term [Eq. (79)] is returned by performing the $[L](t)$ first-order expansion for the case where the bound ligand approaches zero:

$$P_{\text{UNREACT}}([L](t))^* = \frac{1}{e^{\frac{[L](t)}{EC_{50}}}} \stackrel{[L](t) \rightarrow 0}{\cong} \frac{1}{1 + \frac{[L](t)}{EC_{50}}} = P_{\text{UNREACT}}([L](t))^{\text{MM}}. \quad (81)$$

It is worth noting that there is no monotonic form between 0 and 1 other than that of Eq. (80) to reproduce the basic Michaelis-Menten term [Eq. (79)] when approximated for a small $x = [L](t)/K_M$. For instance, if one decides to use $\exp(-x^2)$, the unreactive probability will give $1/(1+x^2)$ as the approximation for a small x , which is definitely different than expected for basic Michaelis-Menten treatment [Eq. (79)].

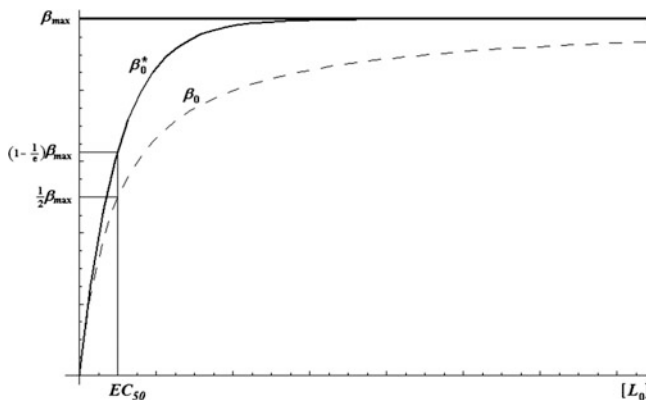


Fig. 3 Initial Michaelis–Menten and logistic velocities plotted against initial ligand concentration for the L–R direct interaction/binding. The *dashed curve* corresponds to the Michaelis–Menten equation (72), while the continuous *thick curve* represents its logistic generalization from (82) (see [198])

The physico-chemical meaning of equation (80) is that the Michaelis-Menten term [Eq. (79)] and its associated kinetics apply to fast ligand–receptor reactions/metabolization, i.e., for fast consumption of $[L](t)$. However, by using Eq. (80) instead of Eq. (79), the range of reaction rates is expanded and provides a new kinetic equation, in the form of the logistic expression

$$-\frac{1}{\beta_{\max}} \frac{d}{dt}[L](t) = 1 - e^{-\frac{[L](t)}{EC_{50}}} \quad (82)$$

by incorporating Eqs. (75) and (78). Under initial conditions, the logistic equation (82) gives an initial velocity of reaction (β_0^*) that is uniformly higher than that calculated by Michaelis-Menten (79) at all initial concentrations of the ligand, except for the case where $[L_0] \rightarrow 0$, when both are zero (see Fig. 3).

To test whether the logistic kinetic equation (82), which is a natural generalization of the Michaelis-Menten equation, may provide a workable analytical solution in an elementary form, we first integrate it under the form

$$\int_{[L_0]}^{[L](t)} \frac{d[L](t)}{\exp(-[L](t)/EC_{50}) - 1} = \int_0^t \beta_{\max} dt \quad (83)$$

generating the new equation to be solved:

$$[L_0] - [L](t) + EC_{50} \ln\left(e^{-\frac{[L_0]}{EC_{50}}} - 1\right) - EC_{50} \ln\left(e^{-\frac{[L](t)}{EC_{50}}} - 1\right) = \beta_{\max} t. \quad (84)$$

This can be solved exactly by substituting

$$\varphi([L](t)) = \frac{[L](t)}{EC_{50}} \quad (85)$$

into Eq. (84) to obtain the simple equation:

$$-\varphi([L](t)) - \ln\left(e^{-\varphi([L](t))} - 1\right) = \Psi(t) \quad (86)$$

where we have also introduced the functional notation:

$$\Psi(t) = \frac{1}{EC_{50}} (\beta_{\max} t - [L_0]) - \ln\left(e^{-\frac{[L_0]}{EC_{50}}} - 1\right). \quad (87)$$

Now, the exact solution of Eq. (86) takes the logistic expression:

$$\varphi([L](t)) = \ln\left(1 - e^{-\Psi(t)}\right). \quad (88)$$

Finally, substituting function (87) into expression (88) gives the logistic progress curve for ligand consumption in an analytically elementary form [198]:

$$[L]_L(t) = EC_{50} \ln\left(1 + e^{-\frac{\beta_{\max} t}{EC_{50}}}\left(e^{\frac{[L_0]}{EC_{50}}} - 1\right)\right). \quad (89)$$

This time-dependent solution (89) substitutes an elementary logarithmic dependency for the *W*-Lambert function. It is nevertheless remarkable that the solution of a generalized logistic kinetic version of the Michaelis-Menten instantaneous equation provides an analytically exact solution. It clearly reduces to the above Eq. (74) in the first order expansion of the chemical concentration time evolution with respect to the 50-effect concentration (EC_{50}) observed.

The original chemical–biological-kinetics [Eq. (74)] gave no analytical solution for the actual working kinetics [Eq. (75)] that provides the logistical solution (89) whose reliability was previously tested on various enzyme kinetics mechanisms. This testing produced remarkable results [196, 197, 199, 200] that constituted a trusted background for employing it in the currently envisaged ecotoxicological studies.

Remarkably, rearranging the logistic solution of the chemical–biological interaction [Eq. (89)] under the equivalent form

$$e^{\frac{[L]_L(t)}{EC_{50}}} - 1 = e^{-\frac{\beta_{\max} t}{EC_{50}}}\left(e^{\frac{[L_0]}{EC_{50}}} - 1\right) \quad (90)$$

provides for a relatively higher concentration $EC_{50} \gg$ (specific to environmental toxicological fate studies) for the working equation employed in Eq. (10) as the basis for advancing quantitative reactivity–activity relationships.

Table 3 The vectorial descriptors in a Spectral-SAR analysis

| Activity | Structural predictor variables | | | | | |
|--------------------------|--------------------------------|---------------|----------|---------------|----------|---------------|
| $ A_{OBS(ERVED)}\rangle$ | $ X_0\rangle$ | $ X_1\rangle$ | \dots | $ X_k\rangle$ | \dots | $ X_M\rangle$ |
| $A_1\text{-OBS}$ | 1 | x_{11} | \dots | x_{1k} | \dots | x_{1M} |
| $A_2\text{-OBS}$ | 1 | x_{21} | \dots | x_{2k} | \dots | x_{2M} |
| \vdots | \vdots | \vdots | \vdots | \vdots | \vdots | \vdots |
| $A_N\text{-OBS}$ | 1 | x_{N1} | \dots | x_{Nk} | \dots | x_{NM} |

As such, for a *logistic-spectral analysis*, based on the molecular M -data for N -chemical species and toxicity activities of Table 3, the next steps are considered in producing the chemical–biological progress curves according to which chemical species “dissolve” in biological/environmental receptors.

1. For the activities of Table 3, the Spectral-SAR [74, 201, 202] relationships are formulated

$$|A^*\rangle^{\text{ENDPOINT}} = B_0|X_0\rangle + B_1|X_1\rangle + \dots + B_M|X_M\rangle \quad (91)$$

for each envisaged molecular-set or model of structural parameters or computational framework.

2. The predicted spectral norm is computed in the N -chemical space

$$\begin{aligned} |||A^*||| &= \sqrt{\langle A^*|A^* \rangle} = \sqrt{\sum_{k=0}^M B_k^2 \langle X_k|X_k \rangle} = \sqrt{\sum_{k=0}^M B_k^2 \left(\sum_{j=1}^N x_{kj}^2 \right)} \\ &= \sqrt{\sum_{j=1}^N (A_j^*)^2}. \end{aligned} \quad (92)$$

3. The initial chemical concentration within the logistical chemical–biological progress picture is related to the predicted S-SAR activity norms (92), according to Eq. (12a, 12b).
4. In the same mechanistic line of the chemical–biological interaction framework, the maximum biological effect is seen as the decrease of the initial chemical concentration in the effector time [see Eq. (17)].
5. Along the EC_{50} parameter computed following the generalization of Eq. (9) toward the algebraic version Eq. (12b), one has all the “ingredients” for progress curves that represent the logistical consumption/metabolization/fate of the chemical species following interaction with the biological species/ecological environment; these are written in the working form, first with the aim of Eqs. (17) and (89), as

$$[L]_L(t) = EC_{50} \ln \left(1 + e^{-\frac{1}{4e^2} \frac{[L_0]}{EC_{50}} t} \left(e^{\frac{[L_0]}{EC_{50}}} - 1 \right) \right). \quad (93)$$

Finally, by replacing the Eqs. (12a) and (12b) relationships, the (observed) activity–(predicted)activity form of the ligand progress curve

$$[L]_L(t) = e^{-\| |A\rangle \|} \ln \left(1 + e^{-\frac{\exp(\| |A\rangle \| - \| |A^*\rangle \|)}{4e^2}} t \left(e^{\exp(\| |A\rangle \| - \| |A^*\rangle \|)} - 1 \right) \right) \quad (94a)$$

or toxicity–activity expression

$$[L]_L(t) = EC_{50} \ln \left(1 + e^{-\frac{\exp(-\| |A^*\rangle \|)}{4e^2 EC_{50}}} t \left(e^{\frac{\exp(-\| |A^*\rangle \|)}{EC_{50}}} - 1 \right) \right) \quad (94b)$$

are created for each structural parameter (reactivity indices considered) computed for each DFT framework considered and are correlated with each set of recorded species activities. Through this comparison, one may draw conclusions based on the chemical reactivity principles governing the specific chemical–biological interaction under study.

Next, a pilot application will be carried out to determine the activity of selected Cl-PAHs for various endpoints and biological species.

5 Quantitative Reactivity–Activity Modeling of Chemical–Biological Interactions

5.1 On QSAR Principles

Quantitative structure–activity (or property) relationship (QSA(P)R) methods seem best for unifying the chemical (and biological) interaction into a single model for researchers aiming to quantitatively organize the huge amount of experimental information in comprehensive equations with a predictive value [71, 203]. The QSA(R)R equation is justified by the quantum superposition principle written in the multilinear form of causes ($|X\rangle$ in the Dirac bracket notation of quantum states), resulting in Eq. (91) [204] and providing the appropriate framework in searching for new “natural laws” by various statistical means for computing the coefficients of these expansions (B ’s), such that the error of the predicted to recorded effects $|A^*\rangle$ is minimized. *Classical QSA(P)R* [67–70] assumes as descriptors the structural indices that directly reflect the electronic structures of the tested chemical compounds, e.g., factors describing the lipophilicity (e.g., LogP, surfaces), electronic effects (e.g., Hammett constants, polarization, localization of charges), and steric effects (e.g., Taft indices, Verloop indices, topological indices, molecular mass, total energy at optimized molecular geometry). Therefore, the optimization is centered on the molecular structure and various descriptors for the system of interest.

The driving QSAR principles have been established for the *in silico* validation of a compound. When completely fulfilled, a QSAR model can give, if not a general natural law [4], a working quantitative model for a given pool of chemical–biological interactions. These principles meet *European normative* regulatory

requirements for a given class of compounds, in accordance with the *Organization for Economic Cooperation and Development* (OECD). These QSAR-OECD principles are [5, 205]

- i. **QSAR-OECD-1:** defining the (biological, ecological, or pharmacological) *activity* of a given chemical compound (the so-called *endpoint* A).
- ii. **QSAR-OECD-2:** using a non-ambiguous algorithm in the quantitative attribution of activity (endpoint) for a chemical series based on their structure or physico-chemical properties and indices (X_1, X_2, \dots, X_M in QSAR equation).
- iii. **QSAR-OECD-3:** defining the applicability domain relative to compounds and structural diversity considered in correlation with the QSAR model of the envisaged activity with the physico-chemical selected indices for the *in silico* tests (determination of parameters $B_0, B_1, B_2, \dots, B_M$ in QSAR equation).
- iv. **QSAR-OECD-4:** the quality of the QSAR model measured using the regression factor, robustness and predictability.
- v. **QSAR-OECD-5:** the possibility of formulation for a mechanistic model of the physico-chemical interactions that yield the assumed activity (endpoint) and QSAR equation in general.

QSAR modeling is crucial for understanding and predicting toxicity-specific mechanisms but is faced with two fundamental problems [6]:

- (a) A priori establishment of the physico-chemical factors to be considered in the QSAR equation to produce the most reliable model (i.e., assurance of the QSAR-OECD I, II, and IV principles).
- (b) Assuring that activity A^* depends on physico-chemical factors that are *independent* and associated with the *primary causes* that generate the observable effect of the activity (endpoint) (i.e., the assurance of the QSAR-OECD III and V principles).

The present approach, considering structural parameters characterizing the frontier information of the molecular structures (electronegativity χ , chemical hardness η , electrophilicity ω , and chemical power π) that drive the chemical reactivity principles, provides individual associate QSARs for a given biological action/species. This generates the actual quantitative reactivity–activity relationships that have a major role in modeling the chemical–biological interaction along the logistic ligand–receptor progress curves, as previously described.

5.2 *In silico*-Based Biological Activities

Biological activity for a given molecule (available or newly synthesized) interacting with a certain species is not always known or easily produced;

determining these interactions can require extensive laboratory efforts and adherence to many environmental safety conditions.

Instead, at the *in silico* level, the *fill-in-the-data-gaps* technique may be applied (see Fig. 4), featuring the main algorithm [206, 207]:

- Choosing a *target molecule* provides existing analogues whose experimental data are available for a studied “end-point” (activity to be modeled).
- Providing data that are required for the target molecule either by read-across (analogue approach) or by trend analysis (molecular similarity approach).
- All these imply inner QSAR modeling specific for the profiled activity (ligand-receptor) that is hypothesized or a known binding mechanism, involving all available data (analogues and their endpoint measured effects).

The mechanism of toxic action involved in the algorithm loop of Fig. 4 is associated with the critical biological effect of the toxicant at the molecular or cellular level. The main classes of toxic action mechanisms are as follows: nonpolar narcosis, polar narcosis, weak acid respiratory uncoupling, formation of free radicals, electrophilic reactions, and toxic action by specific (receptor-mediated) mechanisms.

However, identification of the mechanism of toxic action is often difficult due to the complex nature of toxic activity [208].

As a general rule, the narcotic mechanism of toxic action is a result of nonspecific noncovalent reversible interactions with cell membranes [208]. Note that nonpolar and polar narcosis can be included in the narcotic mechanism category: nonpolar narcotics are neutral nonreactive compounds (aliphatic alcohols, ketones, and ethers), while polar narcotics are less inert and often possess a hydrogen donor moiety (phenols, anilines) (see [209]). Alternatively, compounds that undergo direct electrophilic interaction may have covalent interactions with biological macromolecules [210]. However, compounds may also undergo metabolic reactions resulting in more toxic forms; other chemicals produce their toxic effects by forming free radicals. For a general guideline, Table 4 presents a summary of structural criteria that can be used to assign mechanisms of toxic action to compounds.

5.3 Chemical Reactivity Principles Hierarchy According to the Biological Activity of Cl-PAHs

Polycyclic aromatic hydrocarbons (PAHs) are a class of more than 100 chemicals composed of up to six benzene rings fused together, such that any two adjacent benzene rings share two carbon bonds (e.g., phenanthrenes, naphthalene, and pyrene). They are generally produced during the incomplete burning of organic materials, including coal, oil, gas, wood, garbage, and tobacco. Coal tar ointments containing PAHs are used to treat several inflammatory skin conditions. PAHs are most often generated from motor vehicle exhaust, residential and industrial heating sources, coal, crude oil and natural gas processing, waste incineration, and tobacco smoke.

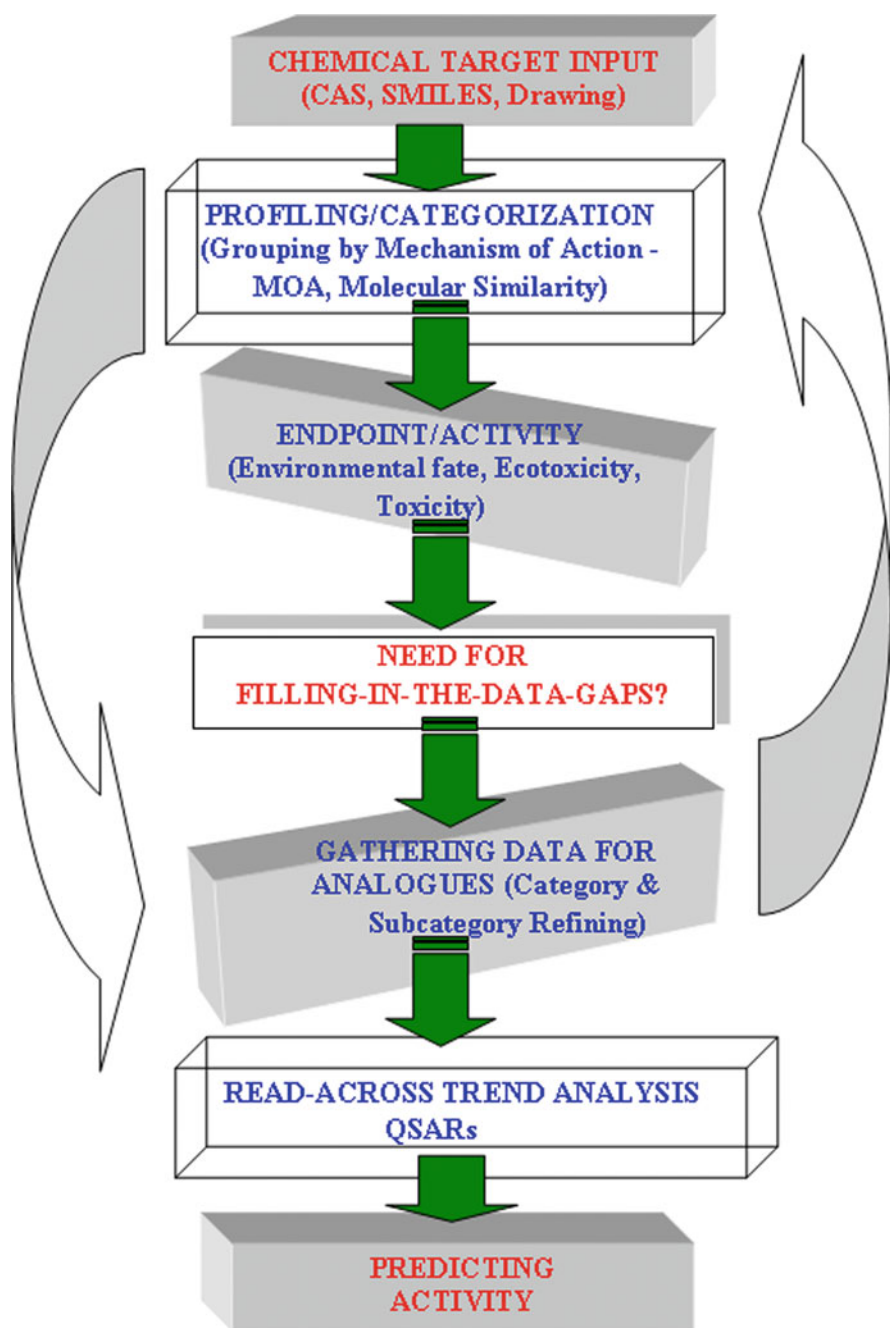


Fig. 4 The *in cerebro* scheme for *in silico* evaluation of environmental or toxicological activities of a given target chemical (of interest, or newly designed or synthesized), following and explicating the Toolbox QSAR computational facility [207]

Table 4 Summary of structural criteria used for classifying compounds according to the mechanism of toxic action [210]

| Mechanism of action (MOA) | Structural features |
|---------------------------------|---|
| Nonpolar narcosis | Saturated alkanes with, e.g., halogen and/or alkoxy substituents (aliphatic alcohols, ketones, ether, amines); halogens and alkyl substituted benzenes |
| Polar narcosis | Phenols with a pK_a greater than or equal to 6.0; phenols and anilines with three or fewer halogen atoms and/or alkyl substituents |
| Weak acid respiratory uncouples | Phenols and anilines with four or more halogen substituents or more than one nitro group or a single nitro group and more than one halogen group |
| Formation of free radicals | Phenol or aniline substituted with an electron-releasing group (alkoxy, hydroxyl, more than one alkyl group) |
| Electrophile/proelectrophile | Activated unsaturated compounds; benzene rings without aniline or phenol substructures that have two nitro groups on one ring; phenols with a single nitro group but not more than one halogen group; aromatic compounds with two or more hydroxy groups in the <i>ortho</i> or <i>para</i> position and at least one unsubstituted aromatic carbon atom; quinines; aldehydes; compounds with halogens at the α -position of an aromatic bond; ketenes; epoxides |
| Specific mechanism | Chemicals that interact with specific biological macromolecules. For example, acetylcholinesterase inhibitors with an organophosphate group |

Emitted PAHs can bind to particles in the air. Particle size depends in part on the source of the PAHs, while ambient air PAH concentrations show seasonal variation [213, 214]. PAHs are found in meat and in other foods as a result of smoking, grilling, broiling, or other high-temperature processing. Uncooked foods and vegetables also contain low levels of PAHs but can be contaminated by airborne particle deposition or growth in contaminated soil. Humans are usually exposed to PAH mixtures rather than to individual chemicals, and PAH mixture composition varies with the combustion source and temperature [215]. CIPAHs are hybrids of dioxins and PAHs suspected of having similar toxicities [216] and are generally known to be carcinogenic, mutagenic, and teratogenic, with greater mutagenicity, aryl-hydrocarbon receptor activity, and dioxin-like toxicity than the corresponding parent PAHs [217]. Especially at the DNA interaction level, CIPAHs have the ability to bind to and activate the aryl hydrocarbon receptor (AhR), a cytosolic, ligand-activated transcription receptor. The biological pathway involves translocation of the activated AhR to the nucleus. In the nucleus, the AhR binds to the AhR nuclear translocator protein to form a heterodimer, leading to transcriptional modulation of genes and causing adverse changes in cellular processes and function [218].

Several CIPAHs have been determined to be AhR-active. AhR-mediated toxicity (Fig. 5) is activated by all embryotoxic HAH and PAH congeners [212, 219, 220] toward nuclear translocation, where the AhR heterodimerizes with the AhR nuclear translocator (ARNT). The resulting ligand-AhR-ARNT complex further combines with various coactivators and promotes their expression through interacting in the

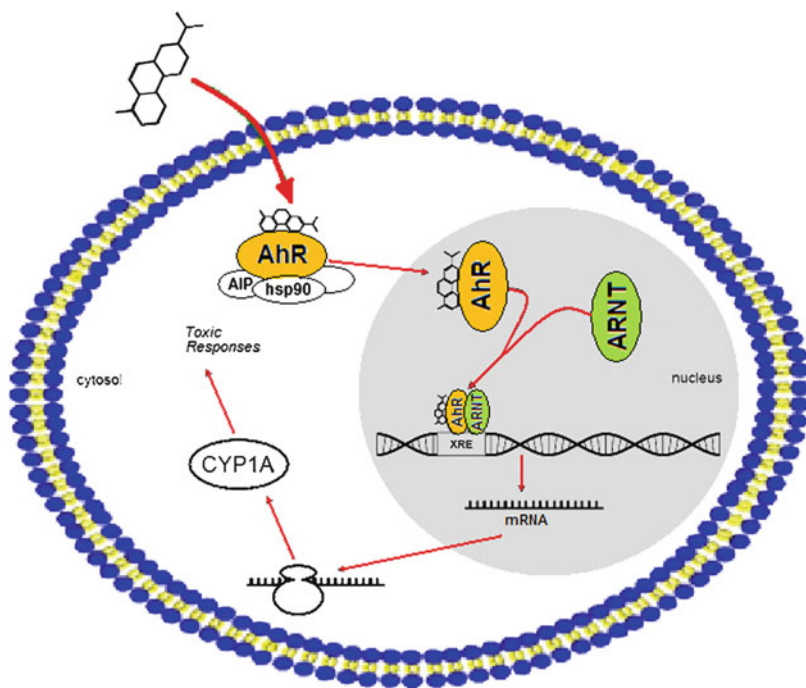


Fig. 5 Generic mechanism of AhR-mediated toxicity: AhR mediates signal transduction by dioxin-like ligands, which form a transcription factor complex with an aryl hydrocarbon nuclear translocator protein (ARNT). This heterodimer recognizes specific DNA sequences, namely dioxin responsive elements (DREs), and leads to induction of several genes forming the so-called Ah gene battery. In this process, the elevated levels of the protein products are assumed to be involved in the toxic action of AhR ligands. *AIP* AhR inhibitory protein, *hsp90* 90-kDa heat shock protein, *ARNT* AhR nuclear translocator, *XRE* xenobiotic response element, *CYP1A* cytochrome P450 1A gene/protein (adapted from [211, 212])

promoter region of AhR-regulated genes with xenobiotic responsive elements (XREs) [221]. This activity plays a role in cell proliferation and differentiation [222] and contributes to the biotransformation of xenobiotics, while also having a functional role in normal development and homeostasis [223–225]. Furthermore, the role of the AhR in TCDD (2,3,7,8-Tetrachlorodibenzo-*p*-Dioxin) and unalkylated PAH toxicity has been assessed [226, 227], but its role in alkylated PAH toxicity has not [211].

CIPAHs may be toxic to humans, and they have an equally important impact on the environment because several CIPAHs have also been found to exhibit mutagenic activity in *Salmonella typhimurium* in the Ames assay [229]. To comprehensively estimate reactivity–activity for representative CIPAHs (see Fig. 6) on human and environmental species, Table 5 presents the ethoxyresorufin-*O*-deethylase (EROD) activities for binding substrates of chlorinated polycyclic aromatic hydrocarbons (CIPAHs) with aryl hydrocarbon receptors (AhRs) in the cytochrome

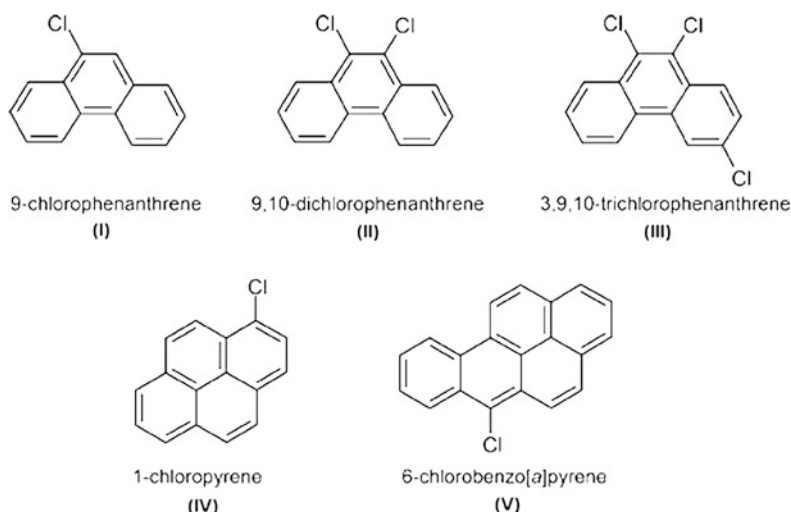


Fig. 6 Chemical structures of CIPAHs of interest [228]

Table 5 Chlorinated polycyclic aromatic hydrocarbons (CIPAH) activity expressed as (a) EROD (ethoxyresorufin-*O*-deethylase) activity as the relative intensity of CIPAH-induced cytochrome P450 (CYP) activity in human breast cancer MCF-7 cells [228]; (b) environmental fate: [Bioaccumulation aquatic] in *Pimephales promelas* over 96 h [10^3 L/kg wet]; (c) ecotoxicological information: [Aquatic Toxicity] LC50 for *Pimephales promelas* after 96 h [10^{-1} mg/L]; and (d) carcinogenicity in rats, TD50 [10^{-3} mol/kg]

| Cl-PAH | CAS | Activity/toxicity | | | |
|--------|-------------|---------------------|------------------------|---------------------------|-----------------------------|
| | | EROD ^(a) | Fate/Pp ^(b) | AquaTox/Pp ^(c) | Carcino/Rats ^(d) |
| (I) | 947-72-8 | 1.2 | 1.90 | 3.41 | 4.72 |
| (II) | 17219-94-2 | 1.4 | 1.74 | 1.3 | 4.05 |
| (III) | 800409-57-8 | 4.4 | 4.22 | 0.828 | 3.84 |
| (IV) | 34244-14-9 | 1.3 | 4.08 | 1.3 | 0.00511 |
| (V) | 21248-01-1 | 9 | 4.22 | 0.33 | 3.46 |

The values of (b)–(d) are computed using the Filling-in-the Data-Gap Toolbox OECD facility [v.1.1.01/2009] with OASIS baseline surface narcosis through DNA binding of PAHs in Fig. 6 [206, 207]

P450 (CYP) 1 family (CYP1A1 and 1B1) and expression in human breast cancer MCF-7 cells [228]. Also included are ecotoxicities on fish (*Pimephales promelas*) and rats, as computed by the previously described fill-in-data gaps method. On the molecular-ligand side, each compound in Fig. 6 is a single point in its symmetrical state computed within no-exchange-no-correlation (X0C0) Hartree–Fock (HF), and specific DFT Becke's exchange-correlation forms (Becke97, Becke88-VWN, B3-PW91, B3-LYP) with large Gaussian basis function (6-31G^{**}) schemes within HyperChem software [230] using the HOMO and LUMO frontier information; they were then employed to produce the reactive indices of electronegativity [Eq. (18)],

Table 6 The values of electronegativity (in eV) for the CIPAHs of Fig. 6 computed from the frontier-like formula (18) with the HOMO and LUMO frontier orbital energies that were evaluated in various quantum mechanical frameworks: no-exchange-no-correlation (X0C0) to Hartree-Fock (HF), and specific DFT Becke's exchange-correlation forms (Becke97, Becke88-VWN, B3-PW91, B3-LYP) with large Gaussian basis functions (6-31G**) within HyperChem software [230]

| χ | Quantum chemical framework | | | | | |
|--------|----------------------------|----------|----------|-------------|----------|----------|
| CI-PAH | X0C0 | HF | Becke97 | Becke88-VWN | B3-PW91 | B3-LYP |
| (I) | 251.1668 | 161.6397 | 190.1459 | 191.0285 | 190.8256 | 190.7167 |
| (II) | 161.12 | 122.6308 | 155.8952 | 129.6772 | 155.9582 | 176.9095 |
| (III) | 16.58217 | 156.6598 | 78.37052 | 30.02489 | 233.2496 | 78.76642 |
| (IV) | 115.2805 | 62.72652 | 199.2066 | 220.7729 | 192.2411 | 194.9907 |
| (V) | 163.9368 | 256.0867 | 88.00236 | 98.01646 | 215.249 | 215.0238 |

Table 7 The values of chemical hardness (in eV) for the CIPAHs of Fig. 6 computed from the frontier-like formula (19) with HOMO and LUMO frontier orbitals' energies evaluated as in Table 6

| η | Quantum chemical framework | | | | | |
|--------|----------------------------|----------|----------|-------------|----------|----------|
| CI-PAH | X0C0 | HF | Becke97 | Becke88-VWN | B3-PW91 | B3-LYP |
| (I) | 1.183525 | 0.28125 | 2.195176 | 2.39061 | 2.329193 | 2.317177 |
| (II) | 2.44944 | 0.231994 | 0.448074 | 0.690163 | 0.385002 | 0.286087 |
| (III) | 0.065345 | 1.273979 | 1.074822 | 1.177084 | 0.214798 | 0.864036 |
| (IV) | 0.065396 | 0.389016 | 0.510063 | 0.298729 | 0.235435 | 0.381012 |
| (V) | 0.183769 | 0.43338 | 0.284748 | 0.567326 | 2.196144 | 1.941712 |

Table 8 The values of chemical power for the CIPAHs of Fig. 6 computed using definition (3) applied to the electronegativity and chemical hardness values from Tables 6 and 7

| π | Quantum chemical framework | | | | | |
|--------|----------------------------|----------|----------|-------------|----------|----------|
| CI-PAH | X0C0 | HF | Becke97 | Becke88-VWN | B3-PW91 | B3-LYP |
| (I) | 106.1096 | 287.3595 | 43.30996 | 39.95393 | 40.96389 | 41.15281 |
| (II) | 32.88916 | 264.2969 | 173.9615 | 93.9468 | 202.542 | 309.1883 |
| (III) | 126.8827 | 61.48444 | 36.45746 | 12.75393 | 542.9511 | 45.58049 |
| (IV) | 881.4097 | 80.62214 | 195.2765 | 369.5204 | 408.2671 | 255.8852 |
| (V) | 446.0405 | 295.4529 | 154.5267 | 86.38468 | 49.00613 | 55.36966 |

Table 9 The values of electrophilicity for CIPAHs of Fig. 6 computed using definition (4) applied to the electronegativity and chemical hardness values from Tables 6 and 7

| ω | Quantum chemical framework | | | | | |
|----------|----------------------------|-----------|-----------|-------------|-----------|-----------|
| CI-PAH | X0C0 | HF | Becke97 | Becke88-VWN | B3-PW91 | B3-LYP |
| (I) | 26,651.22 | 46,448.71 | 8,235.211 | 7,632.341 | 7,816.958 | 7,848.526 |
| (II) | 5,299.102 | 32,410.95 | 27,119.76 | 12,182.76 | 31,588.08 | 54,698.35 |
| (III) | 2,103.989 | 9,632.138 | 2,857.19 | 382.9353 | 126,643.2 | 3,590.212 |
| (IV) | 101,609.3 | 5,057.147 | 38,900.38 | 81,580.09 | 78,485.72 | 49,895.23 |
| (V) | 73,122.46 | 75,661.56 | 13,598.72 | 8,467.12 | 10,548.52 | 11,905.79 |

chemical hardness [Eq. (19)], chemical power [Eq. (3)] and electrophilicity [Eq. (4)], with the results reported in Tables 6, 7, 8, and 9.

The connection between the activity data of Table 5 and the molecular structural-frontier information from Tables 6, 7, 8, and 9 is made based on the *biological activity-driven chemical reactivity* algorithm, which is qualitatively presented in the scheme of Fig. 1 and quantitatively represented by the logistic ligand progress curves of Eq. (94b). It is clear that the present chemical–biological interaction (CIPAH molecule–AhR-mediated toxicity, see Fig. 5) is a specific realization of the generic ligand–receptor kinetics, which is modeled quantum mechanically and mostly using DFT methods. The interaction involves the predicted norm of the respective chemical structure–biological activity correlation through the presence of the predicted initial (in time evolution of ligand–receptor kinetics) bound ligand to the receptor site [see Eq. (12a)], as well as the algebraic norm evaluation of the specific EC_{50} [see Eq. (12b)], for each observed or recorded (experimentally or computationally by filling in the data gaps—see Fig. 4) set of activities for the molecules of interest. At this point, one should note that the employed activities for bioaccumulation in *Pimephales promelas* (P.p.), ecotoxicology of P.p., and carcinogenicity in rats shown in Table 5 were in fact the correspondent 50 % read-across concentrations for aimed effects obtained using ToolBox in an *in silico* environment. However, it turns out that when considering the EC_{50} and then extracting the activity relationships from the logarithmic forms, in each case, no significant distinction between the influences of the reactivity indices on the bio- and eco-toxicology activity correlation were recorded. Instead, the relationship cancels out all chemical information or mechanisms in producing biological effects. This should be avoided (see QSAR-OECD-5 principle of Sect. 5.1), so we consider the 50 % read-across concentrations as the aimed effects and note this as peculiar *in silico* behavior for ToolBox that should be improved in the future. The biological-driving-chemical interaction results for CIPAHs–AhR-mediated toxicity for human breast cancer MCF-7 cells, aquatic bioaccumulation for *P.p.*, aquatic toxicity for *P.p.* and carcinogenicity for rats are summarized in Tables 10, 11, 12 and 13 and in Figs. 7, 8, 9 and 10.

To correctly interpret the results, one can set the following mechanistic rule for hierarchical biological activity-driving chemical reactivity principles: *the higher the QRAR correlating factor is, the closer the predicted initial bound ligand concentration will regulate the toxicity of the in-set EC_{50} concentration*. Thereby, the overall bio- or eco-toxicological effect is modeled. Nevertheless, one notes that in all cases in Tables 10, 11, 12 and 13 and Figs. 7, 8, 9, and 10, the higher Pearson correlation factor associates with the closer L-bound concentrations for the working EC_{50} . In this way, one can systematically identify which reactivity index (and correspondent principle thereof) is dominant in which quantum/DFT computational environment for the biological or ecological system. As such, for the considered systems one finds the following:

- For modeling the interaction of CIPAH ligands that bind human breast cancer MCF-7 cells (Table 10; Fig. 7), it appears that the B3-PW91 DFT exchange-correlation scheme recovers the consecrate chemical reactivity scheme

Table 10 The synopsis of the quantitative (χ , η , π , ω) reactivity indices–EROD activity relationships (EROD-QRAR), including the correlation factor (R) and the initial predicted concentration of Eq. (12a), $L_0 = \exp(-||A^*||)$ [μM], for each index/quantum chemical method considered in Tables 6, 7, 8 and 9 against the EROD activity of Table 5

| Quantum-method | Index | $ A^*\rangle\text{EROD} - \text{QSAR} - \text{EQUATION}$ | R | L_0 |
|----------------|----------|--|-------|---------|
| X0C0 | χ | $4.55 1\rangle - 7.7 \times 10^{-3} \chi\rangle$ | 0.195 | 390.622 |
| | η | $4.7 1\rangle - 1.57 \eta\rangle$ | 0.483 | 225.796 |
| | π | $3.22 1\rangle + 7.63 \times 10^{-4} \pi\rangle$ | 0.08 | 428.397 |
| | ω | $2.83 1\rangle + 1.51 \times 10^{-5} \omega\rangle$ | 0.197 | 389.811 |
| HF | χ | $-2.78 1\rangle + 0.041 \chi\rangle$ | 0.856 | 64.0455 |
| | η | $2.289 1\rangle + 2.243 \eta\rangle$ | 0.284 | 345.096 |
| | π | $2.27 1\rangle + 6.02 \times 10^{-3} \pi\rangle$ | 0.208 | 384.824 |
| | ω | $0.96 1\rangle + 7.39 \times 10^{-5} \omega\rangle$ | 0.631 | 146.062 |
| Becke97 | χ | $10.38 1\rangle - 0.049 \chi\rangle$ | 0.813 | 75.8095 |
| | η | $5.08 1\rangle - 1.80 \eta\rangle$ | 0.417 | 265.837 |
| | π | $3.39 1\rangle + 5.44 \times 10^{-4} \pi\rangle$ | 0.012 | 436.279 |
| | ω | $5.15 1\rangle - 9.32 \times 10^{-5} \omega\rangle$ | 0.406 | 272.73 |
| Becke88-VWN | χ | $6.99 1\rangle - 0.026 \chi\rangle$ | 0.592 | 165.435 |
| | η | $4.67 1\rangle - 1.18 \eta\rangle$ | 0.288 | 342.959 |
| | π | $4.33 1\rangle - 7.21 \times 10^{-3} \pi\rangle$ | 0.306 | 332.855 |
| | ω | $4.299 1\rangle - 3.8 \times 10^{-5} \omega\rangle$ | 0.378 | 289.6 |
| B3-PW91 | χ | $-10.75 1\rangle + 0.072 \chi\rangle$ | 0.62 | 151.004 |
| | η | $2.15 1\rangle + 1.22 \eta\rangle$ | 0.394 | 279.627 |
| | π | $4.23 1\rangle - 3.1 \times 10^{-3} \pi\rangle$ | 0.203 | 386.758 |
| | ω | $3.83 1\rangle - 7.22 \times 10^{-6} \omega\rangle$ | 0.109 | 421.53 |
| B3-LYP | χ | $2.73 1\rangle + 4.25 \times 10^{-3} \chi\rangle$ | 0.067 | 430.681 |
| | η | $1.79 1\rangle + 1.44 \eta\rangle$ | 0.395 | 279.377 |
| | π | $5.40 1\rangle - 0.014 \pi\rangle$ | 0.53 | 198.381 |
| | ω | $5.24 1\rangle - 6.97 \times 10^{-5} \omega\rangle$ | 0.508 | 211.162 |

(electronegativity > chemical hardness > chemical power > electrophilicity), which may be called the *I-BioAct-ChemReact* scheme of biological activity-driven chemical reactivity realization. Almost the same situation appears for Becke97 computations, except for an inversion of chemical power with electrophilicity influence; in both cases, the prescribed reactivity hierarchy affirms the electronegativity equalization first, followed by chemical hardness principles (see Table 1), while their combined equalization is naturally released with electrophilicity and then with chemical power equalizations at the level of ligand–receptor site molecular interactions, *I-BioAct-ChemReact*: $(\chi \wedge \eta) > (\pi \wedge \omega)$. The second scheme of biological–chemical interaction appears when either chemical power or electrophilicity indices and principles dominate those of electronegativity and chemical hardness, as is the case for B3-LYP computation, *II-BioAct-ChemReact*: $(\pi \vee \omega) > (\chi \vee \eta)$, while the third scheme is recorded when one of the basic chemical reactivity (electronegativity or chemical hardness) indices and principles acts in the first instance and is followed by one of the combined reactivity indices (chemical power or electrophilicity), *III-BioAct-ChemReact*: $(\chi \vee \eta) > (\pi \vee \omega)$. In this last case, although one of the

Table 11 The same synopsis as in Table 10 but for the FATE/Pp activity of Table 5

| Quantum-method | Index | $ A^*\rangle FATE - QSAR - EQUATION$ | R | L_0 |
|----------------|----------|--|-------|---------|
| X0C0 | χ | $4.68 1\rangle - 0.01 \chi\rangle$ | 0.677 | 589.933 |
| | η | $4.132 1\rangle - 1.139 \eta\rangle$ | 0.918 | 497.636 |
| | π | $2.511 1\rangle + 2.26 \times 10^{-3} \pi\rangle$ | 0.617 | 610.868 |
| | ω | $2.597 1\rangle + 1.52 \times 10^{-5} \omega\rangle$ | 0.517 | 643.122 |
| HF | χ | $2.75 1\rangle + 3.17 \times 10^{-3} \chi\rangle$ | 0.173 | 716.799 |
| | η | $2.31 1\rangle + 1.77 \eta\rangle$ | 0.586 | 621.244 |
| | π | $4.53 1\rangle - 6.56 \times 10^{-3} \pi\rangle$ | 0.592 | 619.391 |
| | ω | $3.45 1\rangle - 6.52 \times 10^{-6} \omega\rangle$ | 0.145 | 719.657 |
| Becke97 | χ | $4.94 1\rangle - 0.012 \chi\rangle$ | 0.523 | 641.171 |
| | η | $3.9 1\rangle - 0.744 \eta\rangle$ | 0.450 | 662.262 |
| | π | $3.03 1\rangle + 1.64 \times 10^{-3} \pi\rangle$ | 0.095 | 723.676 |
| | ω | $3.28 1\rangle - 2.53 \times 10^{-6} \omega\rangle$ | 0.029 | 726.44 |
| Becke88-VWN | χ | $4.02 1\rangle - 5.86 \times 10^{-3} \chi\rangle$ | 0.343 | 688.379 |
| | η | $4.07 1\rangle - 0.817 \eta\rangle$ | 0.524 | 641.053 |
| | π | $2.91 1\rangle + 2.66 \times 10^{-3} \pi\rangle$ | 0.295 | 698.247 |
| | ω | $2.99 1\rangle + 1.1 \times 10^{-5} \omega\rangle$ | 0.286 | 699.851 |
| B3-PW91 | χ | $-3.71 1\rangle + 0.035 \chi\rangle$ | 0.793 | 546.71 |
| | η | $3.48 1\rangle - 0.231 \eta\rangle$ | 0.195 | 714.148 |
| | π | $2.51 1\rangle + 2.91 \times 10^{-3} \pi\rangle$ | 0.501 | 647.901 |
| | ω | $2.52 1\rangle + 1.4 \times 10^{-5} \omega\rangle$ | 0.550 | 632.856 |
| B3-LYP | χ | $4.165 1\rangle - 5.45 \times 10^{-3} \chi\rangle$ | 0.226 | 709.856 |
| | η | $3.37 1\rangle - 0.123 \eta\rangle$ | 0.088 | 724.144 |
| | π | $3.65 1\rangle - 2.94 \times 10^{-3} \pi\rangle$ | 0.296 | 697.942 |
| | ω | $3.596 1\rangle - 1.42 \times 10^{-5} \omega\rangle$ | 0.271 | 702.58 |

chemical reactivity principles acts first, it is then integrated into the combined reactivity as chemical power or electrophilicity, where implicitly, the other equalization principle for the remaining reactivity index naturally proceeds, as is the case for X0C0 (non exchange and no correlation), HF and Becke-VWN computations. Note that the last two cases, Schemes II and III of the BioAct-ChemReact correlations, indicate a synergism between electronegativity and chemical hardness in the combined reactivity of chemical power and electrophilicity indices that, in this way, promote both indices as global equalization principles, so that electronegativity and chemical hardness are automatically fulfilled as well. For the EROD activity for human breast cancer MCF-7 cells, B3-PW91 and Becke 97 are in accord with an in-chain reactivity hierarchy as prescribed by Table 1, while the other two DFT computational frameworks (Becke88-VWN and B3-LYP) prescribe synergetics for these principles in combined indices and equalization principles of chemical power and electrophilicity.

- For the effect of the bioaccumulation of CIPAHs on the environmental fate of *Pimphales promelas* (Table 11; Fig. 8), a case is made for the ***I-BioAct-ChemReact*** for X0C0 along Becke97 and Becke88-VWN computational schemes, or for ***II-BioAct-ChemReact*** for HF and B3-LYP computations, while the third case ***III-BioAct-ChemReact*** is found only for the B3-PW91 *in silico* approach.

Table 12 The same synopsis as in Table 10 but for the aqua-toxicity/Pp activity of Table 5

| Quantum-method | Index | $[A^*]AQUATOX - QSAR - EQUATION$ | R | L_0 |
|----------------|----------|---|-------|----------|
| X0C0 | χ | $0.132 1\rangle + 9.19 \times 10^{-3} \chi\rangle$ | 0.668 | 28,156.8 |
| | η | $1.11 1\rangle + 0.407 \eta\rangle$ | 0.360 | 36,315.3 |
| | π | $1.74 1\rangle - 9.7 \times 10^{-4} \pi\rangle$ | 0.291 | 37,716.6 |
| | ω | $1.67 1\rangle - 5.66 \times 10^{-6} \omega\rangle$ | 0.211 | 39,011.7 |
| HF | χ | $2.07 1\rangle - 4.19 \times 10^{-3} \chi\rangle$ | 0.250 | 38,418.2 |
| | η | $1.99 1\rangle - 1.06 \eta\rangle$ | 0.386 | 35,736.7 |
| | π | $0.856 1\rangle + 2.92 \times 10^{-3} \pi\rangle$ | 0.289 | 37,739.1 |
| | ω | $1.46 1\rangle - 7.63 \times 10^{-7} \omega\rangle$ | 0.019 | 40,521.2 |
| Becke97 | χ | $-0.62 1\rangle + 0.014 \chi\rangle$ | 0.692 | 27,454.6 |
| | η | $0.234 1\rangle + 1.329 \eta\rangle$ | 0.883 | 21,941.5 |
| | π | $2.268 1\rangle - 6.91 \times 10^{-3} \pi\rangle$ | 0.442 | 34,392.8 |
| | ω | $1.63 1\rangle - 1.07 \times 10^{-5} \omega\rangle$ | 0.134 | 39,909.4 |
| Becke88-VWN | χ | $0.239 1\rangle + 8.92 \times 10^{-3} \chi\rangle$ | 0.574 | 30,855.6 |
| | η | $0.215 1\rangle + 1.189 \eta\rangle$ | 0.837 | 23,264.0 |
| | π | $1.56 1\rangle - 1.05 \times 10^{-3} \pi\rangle$ | 0.128 | 39,968.1 |
| | ω | $1.47 1\rangle - 1.56 \times 10^{-6} \omega\rangle$ | 0.044 | 40,464.4 |
| B3-PW91 | χ | $4.24 1\rangle - 0.014 \chi\rangle$ | 0.352 | 36,491.9 |
| | η | $0.99 1\rangle + 0.414 \eta\rangle$ | 0.384 | 35,785.8 |
| | π | $1.92 1\rangle - 1.94 \times 10^{-3} \pi\rangle$ | 0.367 | 36,155.2 |
| | ω | $1.87 1\rangle - 8.46 \times 10^{-6} \omega\rangle$ | 0.366 | 36,182.4 |
| B3-LYP | χ | $0.801 1\rangle + 3.69 \times 10^{-3} \chi\rangle$ | 0.168 | 39,563.0 |
| | η | $0.814 1\rangle + 0.535 \eta\rangle$ | 0.420 | 34,927.0 |
| | π | $1.606 1\rangle - 1.22 \times 10^{-3} \pi\rangle$ | 0.135 | 39,900.5 |
| | ω | $1.579 1\rangle - 5.683 \times 10^{-6} \omega\rangle$ | 0.119 | 40,042.5 |

- When the ecotoxicology of CIPAHs on the environmental fate of *Pimphales promelas* (Table 12; Fig. 9) is considered to be the driving scheme, **I-BioAct-ChemReact** dominates the computational quantum frameworks because it is specific to Becke97, Becke88-VWN, B3-LYP, and even X0C0, while the schemes HF and B3-PW91 appear specific to the **III-BioAct-ChemReact** mechanism of the ligand–receptor interaction. Therefore, this chemical–biological system clearly favors chemical reactivity principles as triggering the chemical biological specific interactions.
- Finally, when CIPAH ligands mediated AhR toxicity in rats, carcinogenic effects are produced (Table 13; Fig. 10). No specific **I-BioAct-ChemReact** mechanism of action was recorded because the **II-BioAct-ChemReact** hierarchy dominates this system, covering the X0C0, Becke97, Becke88-VWN, and B3-LYP computations, followed by the **III-BioAct-ChemReact** hierarchies for the HF and B3-PW91 *in silico* methods. Consequently, the synergistic mechanism of joint electronegativity and chemical hardness equalization in chemical power and electrophilicity is the driving mechanism specific for carcinogenesis.

The present applications illustrate the reliability of the **BioAct-ChemReact** logistical-QSAR algorithm in modeling specific biological–chemical interactions for various species, particularly its ability to discriminate between tumor promotion, bio-accumulation, ecotoxicity, and carcinogenicity, as presented and analyzed

Table 13 The same synopsis as in Table 10 but for the carcinogenicity/rats activity of Table 5

| Quantum-method | Index | $ A^*\rangle \text{CARCINO} - \text{QSAR} - \text{EQUATION}$ | R | L_0 |
|----------------|----------|--|-------|---------|
| XOC0 | χ | $2.279 1\rangle + 0.61 \times 10^{-3} \chi\rangle$ | 0.305 | 691.13 |
| | η | $2.539 1\rangle + 0.856 \eta\rangle$ | 0.481 | 607.539 |
| | π | $4.79 1\rangle - 4.94 \times 10^{-3} \pi\rangle$ | 0.940 | 339.57 |
| | ω | $4.639 1\rangle - 3.41 \times 10^{-5} \omega\rangle$ | 0.808 | 415.336 |
| HF | χ | $0.872 1\rangle + 0.015 \chi\rangle$ | 0.585 | 548.196 |
| | η | $3.022 1\rangle + 0.369 \eta\rangle$ | 0.085 | 749.612 |
| | π | $1.37 1\rangle + 9.3 \times 10^{-3} \pi\rangle$ | 0.585 | 548.231 |
| | ω | $2.143 1\rangle + 3.17 \times 10^{-5} \omega\rangle$ | 0.493 | 600.768 |
| Becke97 | χ | $4.918 1\rangle - 0.012 \chi\rangle$ | 0.365 | 665.551 |
| | η | $2.187 1\rangle + 1.129 \eta\rangle$ | 0.481 | 607.539 |
| | π | $5.113 1\rangle - 0.016 \pi\rangle$ | 0.639 | 516.73 |
| | ω | $4.994 1\rangle - 9.81 \times 10^{-5} \omega\rangle$ | 0.778 | 432.813 |
| Becke88-VWN | χ | $4.79 1\rangle - 0.012 \chi\rangle$ | 0.482 | 606.849 |
| | η | $1.68 1\rangle + 1.49 \eta\rangle$ | 0.667 | 499.366 |
| | π | $4.712 1\rangle - 0.012 \pi\rangle$ | 0.960 | 328.542 |
| | ω | $4.38 1\rangle - 5.28 \times 10^{-5} \omega\rangle$ | 0.957 | 330.275 |
| B3-PW91 | χ | $3.28 1\rangle - 3.42 \times 10^{-4} \chi\rangle$ | 0.005 | 754.817 |
| | η | $2.37 1\rangle + 0.79 \eta\rangle$ | 0.464 | 616.591 |
| | π | $4.134 1\rangle - 3.69 \times 10^{-3} \pi\rangle$ | 0.443 | 627.51 |
| | ω | $3.866 1\rangle - 1.28 \times 10^{-5} \omega\rangle$ | 0.350 | 672.034 |
| B3-LYP | χ | $4.55 1\rangle - 7.8 \times 10^{-3} \chi\rangle$ | 0.225 | 719.275 |
| | η | $2.019 1\rangle + 1.03 \eta\rangle$ | 0.515 | 588.817 |
| | π | $4.172 1\rangle - 6.76 \times 10^{-3} \pi\rangle$ | 0.476 | 610.099 |
| | ω | $4.244 1\rangle - 4.02 \times 10^{-5} \omega\rangle$ | 0.535 | 577.469 |

in Tables 10, 11, 12 and 13 and Figs. 7, 8, 9 and 10 for the biological systems of Table 5. As a general output, the chemical reactivity principles of Table 1 act according to the general scenario of chemical bonding (see Sect. 2.2), here extended to chemical–biological interactions (see Sect. 4). Some amendments are required when using the Becke97 exchange–correlation scheme as the main *in silico* DFT modeling paradigm for such a binding mechanism (the hierarchy **I-BioAct-ChemReact**). Carcinogenicity seems to be regulated by the **II-BioAct-ChemReact** hierarchy, with Becke97 and Becke88-VWN as the proper computational frameworks. Further studies are necessary to further assess the synergetics of the chemical reactivity principles that work toward biological activity using bi- and multi-dependent quantitative reactivity–activity relationships (QRARs) with various species and toxicants, as well as by considering the biological–chemical hierarchies through the asymptotic times by which the ligand progress curves [Eq. (94b)] first vanish from each computational–conceptual framework (see Figs. 7, 8, 9 and 10). Systematic studies will seek a general formulation of the chemical–biological interactions using quantum catalytic (logistic) activities.

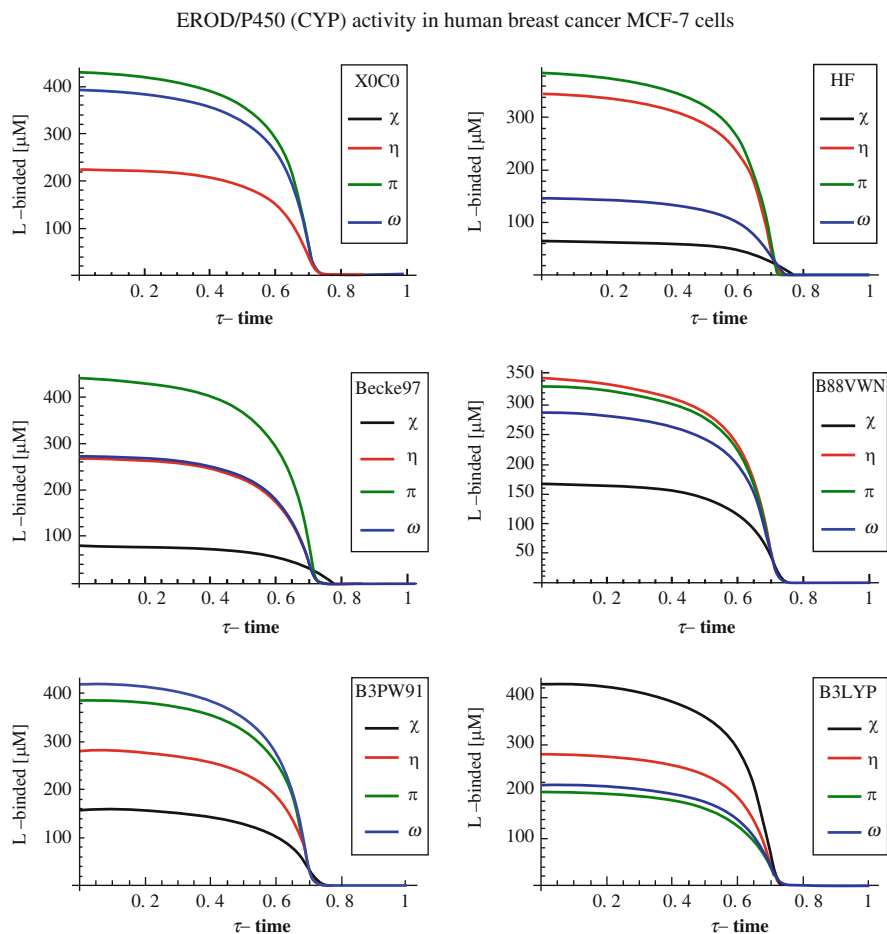


Fig. 7 Comparative representation of the progress curves for the CIPAH ligands of Fig. 6 bound to human breast cancer MCF-7 cells, employing the recorded EROD/human-QRAR reactivity–activity information from Table 10 into logistic chemical–biological interactions modeled by Eq. (89), on the mapped unitary time scale of Eq. (15), for each index/quantum chemical method considered and for an EROD $\text{EC}_{50} = 34.696 \mu\text{M}$ norm parameter as computed with algebraic definition (12b) and the EROD/human data of Table 5

6 Conclusions

Continuous increases in the synthesis of chemical compounds, accompanied by their disposal in the environment, have made hazard toxicity a global issue, especially at the level of the chemical–biological interaction between toxicants and species. In this context, the marriage between structural molecular computing and catalytic activity should be accomplished in a systematic, yet universal, way. In quantitative terms, structural information is today represented by ab initio quantum

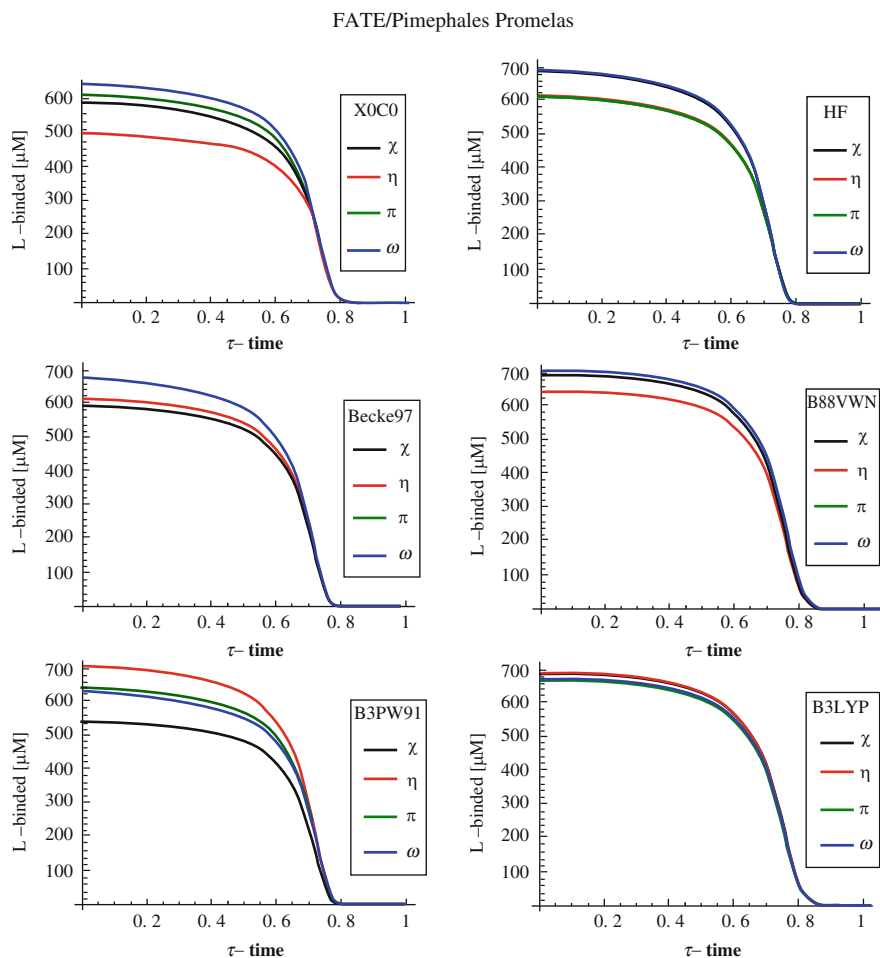


Fig. 8 The same type of representations as in Fig. 7 but for the CIPAH ligand-FATE/Pp QRAR reactivity-activity information from Table 11, with an FATE/Pp $EC_{50} = 464.437 \mu\text{M}$ norm parameter as computed using the algebraic definition (12b) with the FATE/Pp data of Table 5

chemical methods including the advanced methods of Density Functional Theory (DFT). For biological uptake, the preferred paradigms are less obvious, and thus, the present work reviews logistic enzyme-substrate kinetics and generalizes it in terms of ligand-receptor interactions. It is worth noting that the resulting logistic progress curves for ligand receptors, themselves a generalization of the consecrated Michaelis-Menten catalytic paradigm, involve the algebraic employment of activity predicted from a quantitative structure-activity relationship (QSAR), Eq. (94). This method is therefore affirmed as the analytical realization of the chemical-biological interaction, which can be employed for various chemical toxicants and biological species. However, the present approach also includes a broader conceptual

AQUATOX/Pimephales Promelas

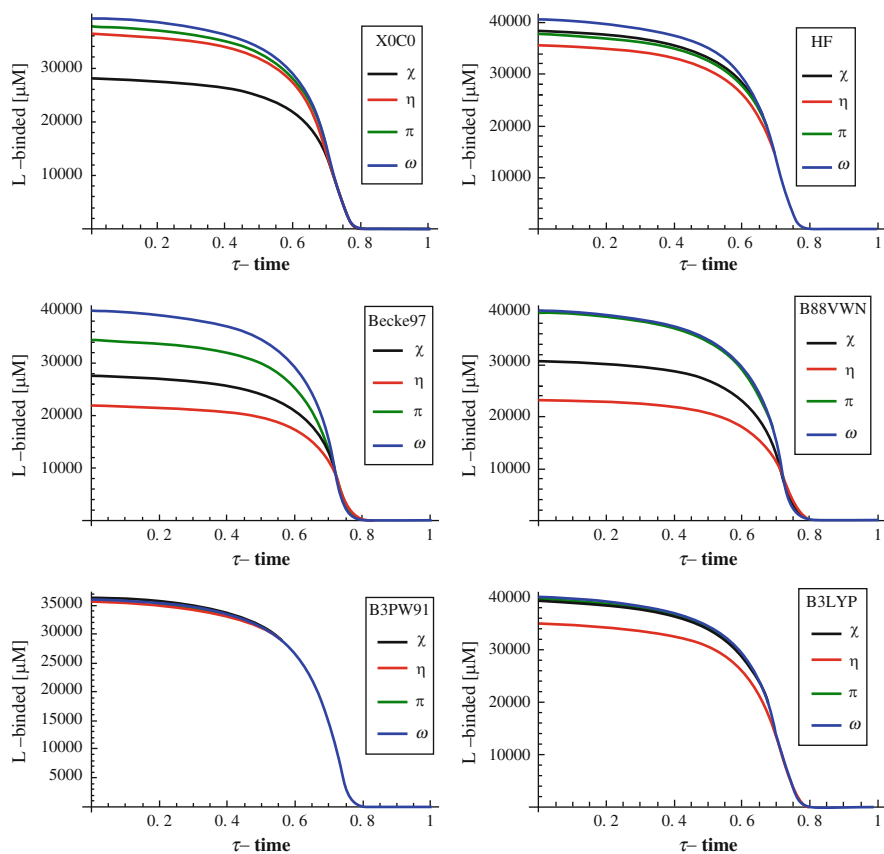


Fig. 9 The same type of representation as in Fig. 7 but for the CIPAH ligand–LC50/Pp QRAR reactivity–activity information from Table 12, with an $\text{EC}_{50} = 18774.7 \mu\text{M}$ norm parameter as computed using the algebraic definition (12b) with the LC50/Pp data of Table 5

consequence: namely, it allows testing of the hierarchy of chemical reactivity principles of electronegativity and chemical hardness.

Recent advances in conceptual DFT have spurred debate over whether these two concepts are manifestly distinct or act synergistically in unified electrophilicity and chemical power indices. Accordingly, chemical reactivity principles are also reviewed, qualitatively and quantitatively, and tested for their hierarchy or synergistic appearance through the “eyes” of recorded biological or computed *in silico* activity (by, for example, the celebrated Toolbox utility). Calculations were carried out on a pilot set of chlorinated polycyclic aromatic hydrocarbons (CIPAHs) using various species (human breast cancer cells), *Pimephales promelas* and rats (*Rattus norvegicus*) according to the aryl hydrocarbon receptor (AhR)-mediated toxicity, as reflected in EROD (ethoxyresorufin-*O*-deethylase) activity, environmental

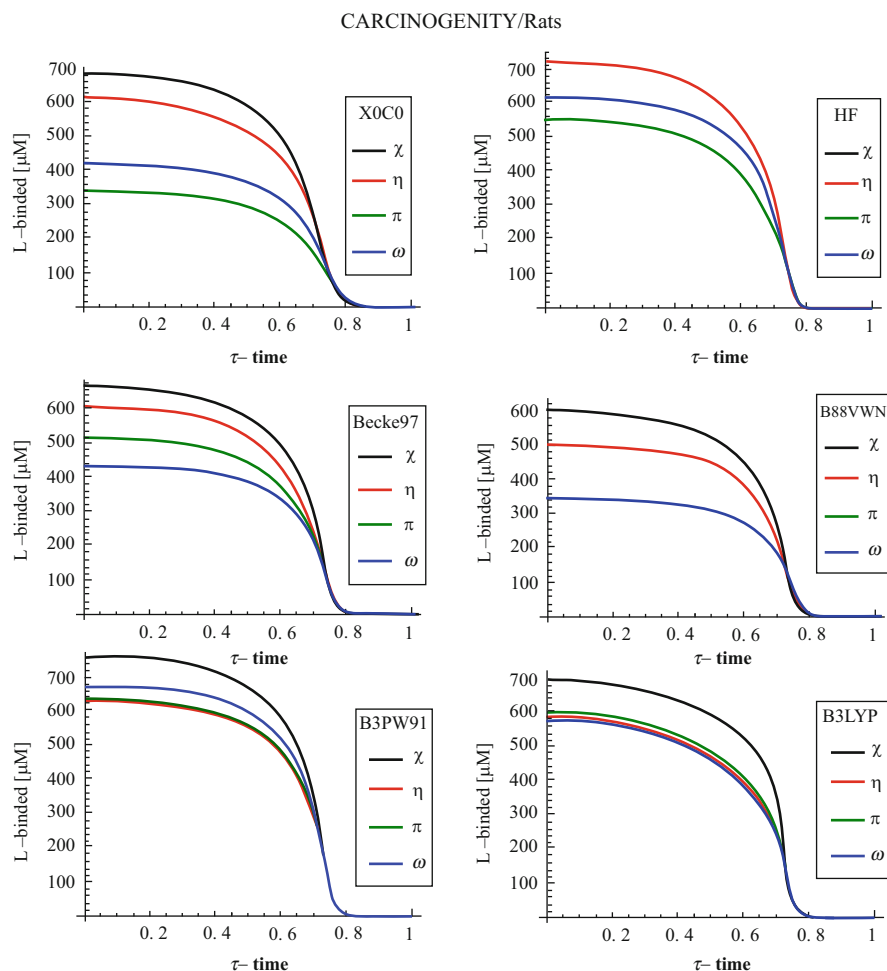


Fig. 10 The same type of representation as in Fig. 7 but for the CIPAH ligand–TD50/rats QRAR reactivity–activity information from Table 13, with an $\text{EC}_{50} = 307.538 \mu\text{M}$ norm parameter as computed using the algebraic definition (12b) with the TD50/Pp data of Table 5

bioaccumulation and toxicity, and carcinogenicity. The results were revelatory because the DFT exchange and correlation methods widely accompanied the chemical bonding scenario (of Table 1) at the level of the biological receptor interactions for most of the environmental hazard toxicity, while modulating the synergistic behavior of the reactivity indices of electronegativity and chemical hardness in the unified chemical bonding measures as the electrophilicity and chemical power, Eqs. (3) and (4), for carcinogenesis modeling. This approach identifies the specific chemical biological interaction for a given toxicant–organism system through chemical reactivity mechanisms, based on logistic ligand–receptor progress-curve-analysis-based QSAR, thus providing the so-called “Biological

Activity driving Chemical Reactivity” realization of the fifth QSAR-OECD principle. The prediction of the initial bound ligand concentration (here in micro-Molar) that triggers the concerned biological uptake is also provided (Figs. 7, 8, 9 and 10). Further studies are envisaged in which quantitative reactivity–activity relationships (QRARs) are revealed for bi- and multi-linear (or even polynomial) correlations between observed or computed activity and frontier chemical information toward systems to determine sequential or synergistic behaviors of chemical reactivity principles and to provide contributions to the biological realm.

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Index

A

Ab initio computations, 181
 Ab initio molecular dynamics (AIMD), 19
 Algebraic norm, 181
 Anticancer activity, 143
 Anticancer drugs, metal–borane clusters, 163
 Aryl hydrocarbon hydroxylase (AHH)
 receptors, 160
 Aryl hydrocarbon receptors (AhRs), 181, 213
 Atoms in molecules, 127, 193
 Auxiliary-DFT (ADFT), 5

B

Becke-97, 201
 Becke, density gradient correlation, 201
 Becke–Johnson exchange hole dipole moment
 formalism, 16
 Becke-88-VWN, 201
 Benzenes, 75, 213
 chlorinated, 160
 electronic localization function, 133
 interactions, 17
 Benzidine, 160
 Biological activity, 143
 Biomolecular modeling, 1
 B3-LYP, 5, 8, 76, 78, 103, 126, 201, 215
 BODFT-MD, 19
 Bond analysis techniques, 119
 Born–Oppenheimer approximation/molecular
 dynamics, 1, 3, 193
 Boron-based transition metal clusters, 165
 B3-PW91, 201, 217

C

Carbon dioxide, 97, 109
 Carbonic anhydrases (CANH), 109

Carcinogenity, 181, 217
 Catalases, 100
 Charge-overlap effects, 16
 Charge-parabola, 189
 Chemical hardness, 181, 183
 Chlorinated polycyclic aromatic hydrocarbons
 (Cl-PAHs), 181, 186, 214
 Chlorobenzenes, 160
 Chloroperoxidases, 100, 111
 Chlorophenols (CPs), 161
 Coal tar ointments, 211
 Comparative molecular field analysis
 (CoMFA), 148
 Complete active space (CAS), 193
 Constrained DFT, 1, 39
 Cupric-superoxo species, 27
 Cysteine thiolate, 100
 Cytochrome oxidases, 99
 Cytochromes P450, 100, 110, 214

D

Density functional theory (DFT), 1, 194
 auxiliary (ADFT), 5
 conceptual, 143, 151, 181
 constrained, 1, 39
 dispersion-corrected (DFT-D), 1, 79
 interpretational, 1, 49
 Kohn–Sham, 1, 65
 local-scaling transformations, 81
 tight-binding (DFTB), 2
 time-dependent (TDDFT), 2
 DFT/molecular mechanical molecular
 dynamics, 1
 Dilauroyl phosphatidylcholine (DLPC), 21
 Dimyristoyl phosphatidylcholine (DMPC), 21
 Dioxin responsive elements (DREs), 214

Dioxygen, activation, 99
 metalled, 51
 triplet, 26
 Dipalmitoyl phosphatidylcholine (DPPC), 21
 Dipole moments, 75
 Dispersion energy, 77
 interactions, 65, 78
 molecular forces, 73
 Dispersion-corrected DFT (DFT-D), 1, 15
 Dithionite, 107
 Dopamine β -monooxygenase (DbM), 26

E

Effective one-electron exchange-correlation potential, 199
 Electric field, 76
 Electron affinity (EA), 186
 Electronegativity, 181, 183
 Electron localizability indicator (ELI), 121
 Electron localization function (ELF), 3, 119, 121
 Electron–nuclear potential, 66
 Electron transfer reactions, 44
 Energy density functional, 87
 EROL activity, 181
 Estrogens, 161
 Ethoxyresorufin-*O*-deethylase (EROD), 214
 Ethylene, ELF, 134
 Euler–Lagrange, 89
 Exchange correlation, 5, 194, 199, 201
 Becke, 215
 energy, 5, 6, 73, 89, 200
 functional, 78, 194
 hole (XDM), 80
 Extended Tang–Toennies damping function, 16

F

Fatty acids, 20
 Ferric-hydroperoxo state, 99
 Ferric hypohalous acid, 111
 Ferrous–dioxygen, 100
 Free energy perturbation (FEP), 32
 Functional mapping, 87

G

Gaussian-type orbitals (GTO), 198
 Generalized gradient approximation (GGA), 5
 Gibbs energy, 165, 191
 Global variational principle, 91

Globins, 99
 Glycine solvation, 36
 Ground state, 186

H

Halides, 111
 Halogenations, 111
 Halogens, 97
 Hartree–Fock–Slater theory, 194, 199
 Heitler–London theory, 194
 Heme oxygenases, 99
 Hemoproteins, 107
 Hilbert space, 66, 69, 91, 120
 Hirschfield population analysis (HPA), 160
 Histone lysine methyltransferase (HKMT), cluster calculations, 8
 Hohenberg–Kohn theorem, 68, 194
 Hückel theory, 194
 extended, 197
 Hydrogenases, 113
 Hydrogen, molecular, 97
 transfers, 44
 Hypohalous acids, 111

I

In silico, 127, 147, 181, 210, 224
 Interpretational DFT, 1, 49
 Ionic bonds, 119
 Ionization potential (IP), 186

J

Jacobian equation, 72

K

Kato theorem, 71
 Keesom dipole–dipole interaction, 77
 Kohn–Sham DFT, 1, 6, 65, 68
 Kohn–Sham orbitals, 194

L

LCGTO- $X\alpha$ method, 6
 Lee–Yang–Parr (LYP) correlation functional, 78, 201
 Levy–Lieb energy density functional, 72
 Ligand-to-ligand, 105
 Lipids, 19, 20
 Lipxygenase, 110
 Local density approximation (LDA), 73, 79

Local spin density (LSD) approximation, 73
Localized orbital locator (LOL), 121
Local-scaling self-consistent field, 91
Local-scaling transformations, 81
Local spin density (Slater exchange), 201
Logistic enzyme kinetics, 181
London forces, 78
Long-range interactions, 15
Lysine, methylation, 9

M

Magnesium oxide, 127, 130
Many-electron wavefunctions, 86
Marcus theory (MT), 45
Maximum probability domains (MPDs), 119, 122, 132
Mechanistic promiscuity, 103
Metal–borane clusters, 143
Metal–ligand bonding, 127
Metal-to-ligand charge transfer, 105
Metalloboranes, 165
Metalloenzymes, 97, 109
Metal–porphyrins, 119, 122, 132
Michaelis–Menten, 205
Minimum electrophilicity principle (MEP), 159
Minimum magnetizability principle (MMP), 159
Minimum polarizability principle (MPP), 159
MO-LCAO, 199
Molecular dynamics (MD) simulations, 19
Molecular orbitals, 119
Monooxygenases, 99
Mulliken population analysis (MPA), 160
Multipole expansion, 77
Myeloperoxidase (MPO), 111

N

Narcotics, 211
Nickel-containing superoxide dismutases (NiSOD), 104
Ni(III)–hydroperoxo state, 104
Ni(II)–superoxide, 104
Nitrate, 107
Nitrate reductase, 107
Nitric oxide reductases (NOR), 106
Nitrite, 107
Nitrite reductases, 107
Nitrogen oxides, 97, 106
No-exchange-no-correlation (X0C0) Hartree–Fock (HF), 215

Nuclear–nuclear Coulomb interaction, 66
Nucleotides, metalloboranes, 165
Nucleotidyl transfer reaction, 13
Nucleus independent chemical shift (NICS), 160

O

One-electron density, 71, 83
Orbit, concept, 86
 variational principle, 89
Organization for Economic Cooperation and Development (OECD), 210
Oxidoreductases, 44
Oxyanions, 97, 106
Oxygen, molecular, 97

P

Peptidylglycine α -monooxygenase (PHM), 26
Peroxidases, 99
Peroxides, 99
Phase transitions, 22
Phosphatidyl choline (PC) lipids, 19, 20
Phospholipids, 20
Pimephales promelas, 181, 217
P450NOR, 106
Polarizable continuum model (PCM), 4, 8
Polychlorinated biphenyls (PCBs), 160, 161
Polychlorinated dibenzofurans (PCDFs), 161
Polycyclic aromatic hydrocarbons (PAHs), 211
 chlorinated (Cl-PAHs), 181, 186, 214
Population analyses (PA), cDFT, 43
Porphyrin, 119
 ELF, 133
Principle of maximum hardness (PMH), 159
Promolecule, 192
Protons, 97, 113
PW91, 78

Q

QM/MM, 1, 4, 30
Quantitative reactivity–activity relationships (QRAR), 185
Quantitative structure–(chromatographic) retention relationships (QSRR), 150
Quantitative structure–activity relationship (QSAR), 143, 145, 181
 OECD, 210
Quantitative structure–property relationship (QSPR), 145
Quantum topological molecular similarity (QTMS), 150

R

Random phase approximation (RPA), 80
Rats (*Rattus norvegicus*), 181, 186
Redox activation, 97
Reductio ad absurdum, 71
Resolution of the identity, 7
Rieske dioxygenases, 103, 110
RNA polymerase, cluster calculations, 11

S

SAPT(DFT), 80
Self-consistent field (SCF), 193, 197
Slater-type orbitals (STO), 198
Small molecule activation, 97
Solvent effects, 195
Structure–activity relationships, 143, 147
Sulfides, 97
Sulfite reductase, 108
Sulfur oxides, 97
Sulfur-based compounds, 107
Superoxide 99
Superoxide dismutases, 104
Superoxide reductases, 104
Surface, 124

T

TCDD, 214
Testosterone, 161
Thiolate obstruction, 100
Thiolate push, 100
Thomas–Fermi theory, 194
Tight-binding DFT (DFTB), 2
Time-dependent DFT (TDDFT), 2
Tobacco smoke, 211
Topological quantum similarity index (TQSI), 150

Toxicity, 143, 181

AhR-mediated, 181
aquatic, 215, 220
environment, 225
fish, 215
PAHs, 214

Two-state reactivity, 103

Tyramine β -monooxygenase (TbM), 26

Tyrosine phenolate, 100

V

Valence shell electron pair repulsion (VSEPR), 50

Valence state, 186

Vanadium haloperoxidases (VPO), 113

van der Waals, 74

forces, 75, 79
interactions, 8, 15, 16, 31
radii, 74

vdW-DF-04, 81

Variable mapping, 86

Variable selection and modeling method based on the prediction (VSMP), 150

Voronoi deformation density (VDD), 44

Vosko–Wilk–Nusair (VWN) local correlation density functional, 201

W

Water, 19, 97

hydroxide, redox activation, 106

X

Xenobiotic responsive elements (XREs), 214

X α theory, 200