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Léa El Khoury, Sehr Naseem-Khan, Karolina Kwapien, Zeina El Hobaika, Richard El Maroun, et al.. Importance of explicit smeared lone-pairs in anisotropic polarizable molecular mechanics. Torture track angular tests for exchange-repulsion and charge transfer contributions. Journal of Computational Chemistry, 2017, 121, pp.3997 - 1920. 10.1021/acs.jpcb.7b01836 . hal-01571959

HAL Id: hal-01571959 https://hal.science/hal-01571959v1

Submitted on 4 Aug 2017

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Importance of explicit smeared lone-pairs in anisotropic polarizable molecular mechanics. Torture track angular tests for exchange-repulsion and charge transfer contributions.

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Summary. A correct representation of the short-range contributions such as exchange-repulsion (E_{rep}) and charge-transfer (E_{ct}) is essential for the soundness of separable, anisotropic polarizable molecular mechanics potentials. Within the context of the SIBFA procedure, this is aimed at by explicit representations of lone pairs in their expressions. It is necessary to account for their anisotropic behaviors upon performing not only in-plane, but also out-of-plane, variations of a probe molecule or cation interacting with a target molecule or molecular fragment. Thus, E_{rep} and E_{ct} have to reproduce satisfactorily the corresponding anisotropies of their quantum chemical (QC) counterparts. A significant improvement of the out-of-plane dependencies was enabled when the sp² and sp localized lone-pairs are, even though to a limited extent, delocalized on both sides of the plane, above and below the atom bearer but at the closely similar angles as the in-plane lone pair. We report calibration and validation tests on a series of monoligated complexes of a probe Zn(II) cation with several biochemically-relevant ligands. Validations are then performed on several polyligated Zn(II) complexes found in the recognition sites of Zn-metalloproteins. Such calibrations and validations are extended to representative mono- and polyligated complexes of Mg(II) and Ca(II). It is emphasized that the calibration of all three cations was for each ΔE contribution done on a small training set bearing on a limited number of representative N-, O-, and S-mono-ligated complexes. Owing to the separable nature of ΔE , a secure transferability is enabled to a diversity

of polyligated complexes. For these the relative errors with respect to the target $\Delta E(QC)$ values are generally < 3%. Overall the paper proposes a full set of benchmarks that could be useful for force field developers.

Introduction.

Anisotropic, polarizable molecular mechanics/dynamics (APMM/APMD) are an appealing alternative to QM/MM calculations for large-scale ligand-macromolecule simulations. Their reliability could be ensured if they enabled to compute the intermolecular interaction energies (ΔE) in the QM recognition site with an accuracy that would match that of quantum chemistry (QC) on which they are grounded. This can be particularly critical regarding the recognition sites of metalloproteins,^[1] in which large polarization and charge-transfer effects come into play. In such sites, furthermore, the binding of the cation(s) to some conjugated ligands such as imidazole, formamide or formate, can occur out of the ligand plane. Out-of-plane binding, with ϕ values of up to 45°, has been observed in some metal binding sites, by both X-ray crystallography and QC optimizations. This can be necessary to enable the cation to optimize its simultaneous interactions with all its ligands, since the latter have limited freedom due to their anchoring to the protein main-chain. It is thus necessary to accordingly account for the ΔE dependencies upon out-of-plane, and not only in-plane, cation binding.

The present work is along the continued enrichments and refinements of the SIBFA procedure.^[2-4] Most of the previous validation studies had resorted to the CEP 4-31G(2d) basis set,^[5] the reliability of which could be shown from parallel ΔE computations with much more extended basis sets,^[4] and Refs. therein. We have subsequently reconstructed a library of fragments with distributed multipoles and polarizabilities derived from QC calculations with the aug-cc-pVTZ(-f) basis set.^[6] The procedure was calibrated,^[7] using the I-NoLLS^[8] procedure and used to study the complexes of the alkali cations $(Li^+ - Cs^+)$,^[9] and the intermolecular interactions of the guanine and cytosine bases.^[10] Its extensions to divalent cations Zn(II), Mg(II) and Ca(II) are reported in the present paper. We consider first the monoligated complexes of Zn(II) with O-, N-, and S-containing ligands. At the optimized cation-ligand distances, we investigate both in- and out-of-plane angular variations. For ligands having localized sp lone-pairs, such as imidazole, or sp² lone-pairs, such as formamide or formate, out-of-plane variations uncovered a shortcoming in E_{rep} . That is, it underwent a regular decrease until the ϕ angle came close to 90°, namely upon entering the zone of the π lone-pairs. By contrast, the QC exchange-repulsion contribution, E_X, had limited amplitudes of variations. Thus, even though the overall value of ΔE favors in-plane binding, the relative destabilization of out-of-plane binding is not sufficient. This could have adverse consequences in MD simulations, in which the poses are populated as a function of the Boltzmann factor. We have found that a very simple, yet efficient, improvement of E_{rep} was obtained upon 'smearing' the sp and sp² lone pairs on both sides of the atom-bearer plane (Figure 1). That is, the in-plane occupation number of 2 is reduced to a value, α , in the range 1.50-1.68, and two additional, out-of-plane, lone-pairs have each half the $(2-\alpha)$ difference. Owing to the separable nature of the SIBFA intermolecular interaction energy, all comparisons with QC will be done by energy-decomposition analyses (EDA), enabling to compare each SIBFA contribution to its QC counterpart regarding non-isotropy in monoligated complexes and non-additivity in polyligated complexes. Because of the

proof-of-principle nature of the present study, we focus on results from Reduced Variational Space (RVS) Energy Decomposition Analyses (EDA),^[11] limited to the Hartree-Fock level, but enabling an operational resolution of ΔE into Coulomb (E_c) and exchange-repulsion (E_x) in first-order, and polarization (E_{pol}) and charge-transfer (E_{ct}) in second-order. Extensions to correlated QC calculations are underway in the context of the SAPT energy decomposition analysis and will be reported separately.

The organization of the paper is as follows. In the Methods section, we shortly recall the leading formulas of E_{rep} and E_{ct} , and the manner the sp and sp² lone-pairs are 'smeared' and probed by an incoming dicationic probe. We next investigate in succession the mono- and polyligated complexes of three divalent metals of upmost importance in all chemistry fields: a 'soft' transition metal, Zn(II), and two hard' alkali-metal cations, Mg(II) and Ca(II).

Monoligated complexes. All selected ligands are encountered as side-chains of proteins, and one (hydroxamate) as a metal-binding moiety of metalloprotein inhibitors. Their complexes with all three divalent cations are first investigated regarding the radial dependencies of ΔE and its individual contributions. In- and out- of plane dependencies are next investigated for one cation, Zn(II), and its complexes with conjugated ligands embodying smeared sp or sp² lone-pairs, namely: formate, formamide, imidazole and hydroxamate. We specify in the Tables which complexes were used for the calibration.

Polyligated complexes. All polyligated complexes were subsequently considered to validate the procedure. The Zn(II) complexes were considered first, with four up to six identical ligands, then with mixed ligands, in arrangements encountered in Zn-metalloprotein binding sites. For Mg(II), three same complexes are considered as with Zn(II), namely those with four and six water molecules, and the one with four imidazoles. This enables to quantify how Zn(II) 'softness' versus Mg(II) 'hardness' are translated in terms of the individual $\Delta E(QC)/\Delta E(SIBFA)$ contributions. A bimetallic Mg(II) binding site is next considered, extracted from the HIV-1 integrase. For Ca(II), which can have coordination numbers in the 6-8 range, three different octahydrate arrangements were considered. This was followed by the study of a Ca(II) polyligated complex with three formate anions, an extreme case of shielding of the dicationic charge.

Regarding the mono-ligated complexes, each ligand is thus probed in a diversity of in- and out-of-plane positions. Considering extreme cases should enable to unravel possible artefacts of the energy potential away from equilibrium, such as in 'vertical' approaches of the probe to the ligand plane, deliberately shortened theta angles, etc. Regarding the polyligated complexes, a diversity of cases is considered, embodying 'hard' as well as 'soft' ligands, while the total charge of the ligands can range from 0 to -3. This should also stretch the possibilities of the potential to reveal possible artefacts, notably regarding non-additivity. This has led us to adopt the somewhat unconventional term 'torture track' overall.

Methods.

1) SIBFA computations.

In the SIBFA procedure,^[4] the intermolecular interaction energy is computed as the sum of five contributions: electrostatic multipolar (E_{MTP*}), short-range repulsion (E_{rep}), polarization (E_{pol}), charge transfer (E_{ct}), and dispersion (E_{disp})

$$\Delta E_{\text{TOT}} = E_{\text{MTP}*} + E_{\text{rep}} + E_{\text{pol}} + E_{\text{ct}} + E_{\text{disp}}$$

 E_{MTP} is computed with distributed multipoles (up to quadrupoles) derived from the QC molecular orbitals precomputed for each individual molecule. They were originally derived from a procedure due to Vigné-Maeder and Claverie^[12] and distributed on the atoms and bond barycenters. The anisotropic polarizabilities are distributed on the centroids of the localized orbitals (heteroatom lone pairs and bond barycenters) using a procedure due to Garmer and Stevens.^[13] The distributed multipoles are presently derived, for large basis sets, using the Generalized Multipole Analysis (GDMA) method by Stone.^[14] E_{MTP*} is augmented with a penetration term.^[15]

 E_{rep} and E_{ct} , the two short-range repulsions, are computed using representations of the molecular orbitals localized on the chemical bonds and on localized lone-pairs.^[2-16] The present work is along continued refinements of the representation of the lone pairs belonging to conjugated atoms which intervene in their expression.

At this stage we wish to recall the starting expressions for the formulation of these two contributions.

Thus E_{rep} is expressed as a sum of bond-bond, bond-lone pair, and lone pair-lone pair interactions.

$$\mathbf{E}_{rep} = C_1 (\sum_{AB} \sum_{CD} rep(AB, CD) + \sum_{AB} \sum_{L_{\gamma}} rep(AB, L_{\gamma}) + \sum_{L\alpha} \sum_{CD} rep(L_{\alpha}, CD) + \sum_{L\alpha} \sum_{L_{\gamma}} rep(L_{\alpha}, L_{\gamma}))$$
(2)

 $\mathbf{rep}(AB,CD) = N_{occ}(AB)N_{occ}(CD)S^{**2}(AB,CD)/(D_{AB,CD})^{n}$

AB and CD denote the chemical bonds belonging to the interacting molecules M_A and M_B , and L_α and L_γ the lone pairs of M_A and M_B . $N_{occ}(AB)$ and $N_{occ}(L_\alpha)$ denote the occupation numbers of bonds AB and CD. The S function is computed as sum of exponentials of the distance between the interacting atoms or lone pair bearers, modulated by coefficients taking into account their type of hybridization and an angular factor, depending upon the angle between the lone pair tip and the direction of approach of the interacting atom (see Refs.^[16-18] for details). The denominator D is the distance between the midpoints of the interacting bonds or lone pair bearers

of MA and those of MB. Each the rep(AB,CD), rep(L_{α} ,CD), and rep(L_{α} ,L_{β}) has two components corresponding to n=1 and 2.

Ect is expressed as:

$$E_{\rm ct} = -2S_{\rm M} \sum_{\alpha} N_{\rm occ}(\alpha) \left((I_{\alpha \rm M^*})^{**2} / \Delta E_{\alpha \rm M^*} \right)$$
(3)

It runs over all lone pairs of electron donating molecules M_A . $N_{occ}(\alpha)$ is the occupation number of lone pair L_{α} belonging to atom A, $I_{\alpha M^*}$ is a function of the overlap between the electron donor, L_{α} , and the anti-bonding orbital of the electron acceptor bond HX or metal cation M. It has dependencies upon the hybridization of L_{α} and the angle centered at A between its direction and the AX, AH, or AM direction. The denominator is the difference between the ionization potential of L_{α} and the electron affinity of the acceptor. Both $I_{\alpha M^*}$ and the denominator are modulated by the electrostatic potentials undergone by both the electron donor and the electron acceptor (see [3-4, 19] for details).

The actual location of the lone pairs has evolved in the course of the SIBFA refinements. On the one hand, those bearing an 's' character, namely sp³, sp² and sp could be oriented along standard hybridization angles such as: for sp³ lp's, θ angles close to 109°.5 and ϕ angles close to 120° and 240°; for sp² lp's, θ angles of 120° and ϕ angles of 0° and 180° ; for sp, a θ angle corresponding to the external bisector of the bearer. On the other hand, the actual distance to the bearer did evolve. Distances corresponding to those given by the Boys localization algorithm,^[20] as listed by the GAMESS software^[21] in the context of the Garmer and Stevens procedure were initially adopted. More recently, upon recalibration of the method with aug-cc-pVTZ multipoles and polarizabilities, we resorted to analyses^[22-23] given by the Electron Localization Function (ELF)^[24-26] giving longer distances of the centroid to the bearer. Thus, for water O sp³ lone pairs, such a distance amounts to 0.59 Å instead of 0.295 Å (Boys), etc. The location of the π lone pairs on aromatic and conjugated fragments had evolved earlier. The initial implementations had them located at the vertical of the π atom bearer and on both sides of the plane, at distances set to 0.5 Å. We were subsequently led to 'probe' the vertical of the conjugated planes of formate, formamide, imidazole, etc., with a Zn(II) cation (Figure 2) approaching each conjugated atom (C, N, or O) and performing EDA to fit E_{rep} in order to reproduce E_X for a range of distances. A 'bent' representation of the lone pairs, with θ angles of 60° and 120° instead of 90°, such that each π lone pair was actually 'shared' between the two atoms of the conjugated bond (Figure 1), enabled a better match to E_X than a vertical representation. 's' containing lone pairs had occupation numbers of 2. The π lone pairs had occupation numbers enabling to complete the valencies of their bearers: 0.5 for each lone pair on conjugated C's on each side of the plane, as well as for N sp and O sp² atoms, 1.0 for conjugated N atoms connected to three bonds, etc.^[27] More recently, we further elaborated on the representation of the π lone pairs owing to their heavy involvement in stacking interactions, such as between nucleic acid bases. This led us to probe by EDA, using a Mg(II) cation or a water molecule, the vertical of each heavy atom of cytosine and guanine, and optimize the distance to the bearer, the increment of effective radii for E_{rep} along the direction of the probed π lone pair, as well as some occupation numbers.^[10] As mentioned in the Introduction, a shortcoming was identified upon monitoring the ϕ dependencies of E_{rep} and E_{ct} . We have thus considered the binding of a dicationic and of a water probe to a sp² O atom, such as in formamide or formate, or to a sp N atom, such as in imidazole. While the θ , inplane dependencies, of these two contributions matched closely those of their EDA counterparts, a progressive loss of their magnitudes took place upon performing at fixed θ out-of-plane variations until the probe reached the perpendicular. By contrast, such decreases did not take place in the case of sp³ lone-pairs. Regarding the latter, in an early investigation on the linear water dimer, we found that upon departing from the direction of one electron-donating sp³ lone pair, and upon moving the interacting HO bond of the electron-accepting water molecule towards the second electron-donating lone pair, the decrease of E_{rep}/E_{et} due to reduced overlap with the second.^[2, 28]

The 'smearing' of the sp and sp² lone pairs was also inspired by the results of ELF analyses. Thus upon decreasing the resolution requested, such lone pairs could be firstly singled out, and then coalesced with the π lone pairs, an indication of partial delocalization. SIBFA resorts to discretized sites to compute E_{rep}/E_{ct} . Smearing thus brings this procedure one step closer to the GEM^[29] and S/G1 (SIBFA/GEM)^[19] procedures, which resort to continuous QC electronic densities to compute both first- and second-order contributions. The computational overhead is not penalizing, being anyway dominated in multimolecular complexes by the iterative computation of E_{pol} . Thus, molecular fragments such as formamide and imidazole total eight and eleven lone pairs respectively. Smearing of the sp imidazole nitrogen and of the sp² formamide oxygen adds up two additional lone pairs (Figures 1 a,d), whence a total of ten and thirteen lone pairs, respectively. For bond-lone pair interactions, the CPU time is increased by ratios of 10/8 and 13/12, resp. For lone pair-lone pair interactions, of which there are N*(N-1)/2, the increases are obviously larger, namely 1.42 and 1.60, nevertheless the CPU time remains outweighted by that of E_{pol} . Finally the analytical formulas for the gradients of E_{rep} and E_{ct} recently coded in Tinker-HP^[30] remain unaltered.

Presently, a single-point SIBFA computation on complexes of app. 300 atoms, as encountered in the recognition sites of protein, is, depending upon the size of the basis set, 10^5 - 10^6 times faster than a QC Hartree-Fock or DFT one. The SIBFA and QC computing times increase as N² and N⁴-N⁶, respectively, N denoting the number of atoms. This fully motivates our aim for $\Delta E(SIBFA)$ to reproduce $\Delta E(QC)$ with relative errors < 2-3%. Nevertheless, the present speed is not sufficient to consider long-duration MD on very large macromolecular complexes. This should be enabled as a next step by Tinker-HP^[30], a massively parallel code in which the SIBFA

potential and its gradients are being ported. This software enables Smooth Particle Mesh Ewald calculations^[31] to compute the long-distance parts of E_{MTP} and E_{pol} , together with Truncated Conjugate Gradient calculation of E_{pol} ^[32] enabling considerable speed-ups in its calculation.

We give in the Appendix the input data concerning the lone pairs of formate, formamide, hydroxamate, and imidazole.

Molecular Dynamics (MD). The one nanosecond MD simulation for Ca(II) in a droplet of 256 waters was done using the Velocity-Verlet algorithm.^[33] The temperature was set at 300 K with the Berendsen algorithm.^[34] The OH bond distances and the HOH angles were constrained to 0.957 Å and 104.5° respectively with the SHAKE algorithm.^[35] The protocol for the MD simulations of the complexes 5PAH with PMI was detailed in.^[36]

2) QC computations. The RVS analyses were done with the GAMESS software.^[21] The QC calculations were performed at the Hartree-Fock level with the aug-cc-pVTZ(-f) basis set. Effective core potentials were used on $Zn(II)^{[37]}$ and Ca(II).^[38] Some polyligated complexes were computed at the cc-pVTZ(-f) basis set because of integral storage limitations. Computations on smaller (30 atoms) polyligated Zn(II) complexes with such a basis gave very close ΔE values to those with the aug-cc-pVTZ(-f) basis set.

Results and discussion.

We will denote throughout by E_1 the sum of E_C and E_X (QC calculations) and of E_{MTP} and E_{rep} (SIBFA calculations), by E_2 the sum of E_{pol} and E_{ct} , and by ΔE (SIBFA) the sum of E_1 and E_2 . The "torture track" set of tests encompasses usual "radial" distance variations but also angular dependencies along with probe interactions with π systems.

The chart given below summarizes the ligands investigated regarding their nature and occurrence in proteins or in metalloprotein-binding drugs.

Ligand	Net charge	Nature	Occurrence
Water	0	hard	
Methanol	0	hard	Ser, Thr side-chain.
Phenol	0	hard	Phe side-chain.
Formate	-1	hard	Asp, Glu side-chain.
Formamide	0	hard	Asn, Gln side-chain, protein main-chain
Imidazole	0	soft	His side-chain.
Methanethiolate	-1	soft	Cys side-chain.
Hydroxamate	-1	hard	Zn-metalloprotein inhibitors.

Monoligated Zn(II) complexes.

1. Distance variations. The complexes of a Zn(II) probe with representative O-, N-, and S-containing ligands were first used to calibrate the procedure. We considered the evolutions of ΔE and its contributions for in-plane distance variations of the approach of the cation along a well-defined angle θ . We list in Appendix 1, along with the Zn(II)-specific parameters, the effective atomic radii used in each energy contribution and the two ligandspecific parameters used to screen the polarizing field in E_{pol}, namely the prefactor of the Gaussian screening factor and its exponent. We then list the internal coordinates used to locate the lone pair centroids with respect to their atom bearer, and the populations and increments of van der Waals effective radii along each lone pair direction. Table Ia reports the comparisons between the QC and SIBFA results at the optimized R distances for O-containing ligands: water, methanol, formamide, phenol, and formate in either in the bridge position or in an external one. These are encountered in the side-chains of the Ser, Thr, Asn, Gln, Tyr, Asp and Glu residues in proteins. Table Ib reports these comparisons for three N-containing ligands: imidazole, pyridine and trimethylamine, and one S- containing one, methanethiolate. The first and the last are encountered in the sidechains of His and Cy- protein residues respectively. The complete radial dependencies are given as Supporting Informations SI. a–i. Tables Ia-b show the calibration to enable a close match of $\Delta E(SIBFA)$ and its contributions to their QC counterparts, which concerns all four contributions to ΔE . It is notable that ΔE (SIBFA) can retain very close values to those of $\Delta E(RVS)$ concerning the binding of Zn(II) to formate in either the bidentate or the external monodentate mode; in the latter, the Zn-O bond is trans to the bond between the C atom and the unbound O. In all Zn(II) complexes with neutral ligands, E_{pol} has larger magnitudes than E₁, a feature which was already noted in 1995 upon calibrating and testing Zn(II) in the context of the CEP 4-31G(2d) basis set.^[3]

2. ϕ and θ dependencies. As a next step, we performed out-of-plane variations of the approach of Zn(II) to its ligating atoms, as denoted by the ϕ angle. For each ligand, we sought to match the evolutions of E_X at the optimized R and for the same θ angle, θ_0 , as for the distance variations, upon performing step-wise variations of the out-of-plane ϕ angle. This enabled to optimize the value of α , along with the increments of effective radii for E_{rep} and E_{ct} along the lone-pair directions. The validation of the calibration was then assessed by parallel SIBFA and RVS computations upon resuming the ϕ variations for several θ angles encompassing θ_0 .

The complete results for Zn(II) complexes are given under the form of Tables as Supporting Information (SII.ad, SIII a-d, SIV a-d, SV a-d and SVI a-d).

Figures 3a-b concern the Zn(II)-formamide complex. They show the evolutions of E_{rep} , and ΔE as a function of θ at two values of ϕ , namely 0° and 90°. The following figures report the corresponding evolutions for Zn(II)-

formate in the 'E' complex (4a-b), Zn(II)-hydroxamate bound to either the carbon-bound O atom (5a-d) or the N-bound O (5a-b), and Zn(II)-imidazole (6a-b).

Formamide-Zn(II) (Figures 3a-b). $E_{rep}(SIBFA)$ matches correctly $E_X(RVS)$ upon stepwise increasing ϕ . This is an essential result for the present study, indicating that upon out-of-plane excursions of the approach of Zn(II) to an amide ligand, as can occur during MD or MC simulations, there should be no imbalances in E_{rep} that could stem from an under-representation of the lone pairs out of the ligand plane and until the perpendicular is reached. If the sp² lone pairs are not smeared, E_{rep} undergoes a strong decrease in magnitude upon increasing ϕ , until ϕ =90° is reached, when the cation starts to undergo the π lone pair repulsions. On the other hand, for each of the four values of ϕ , the directionality of E_X upon varying θ from 105° to 150° is correctly accounted for by E_{rep} . For both $\phi = 0^\circ$ and 30°, the two curves run close to one another. They start to diverge, albeit by small amounts, for $\phi = 60^\circ$ and 90°, which correspond to less favorable ΔE values (Supp. Info. SII a-d). For the least favorable ϕ value of 90°, E_{rep} runs parallel to E_X , and the offset is small (3 kcal/mol) (Figure 3-b). Overall, and since for each ϕ value the evolutions of E_{MTP}^* as a function of θ parallel those of E_C (not shown), E_1 (SIBFA) runs parallel to E_1 (RVS). The other short-range contribution, E_{ct} , has values reasonably close to those of $E_{ct}(RVS)$, as a function of both θ and ϕ . The overlap is less than found for the short-range repulsion, but the differences remain small and do not prevent the close agreement between ΔE (SIBFA) and ΔE (RVS) for all ϕ and θ values investigated.

Formate-Zn(II). (Figures 4a-b). The formate anion is an ubiquitous cation-binding motif, and it is essential to control the anisotropic character of its cation complexes, exemplified here by in- and out-of-plane Zn(II)-binding. $E_{rep}(SIBFA)$ has a much lesser θ dependency than E_X . Thus for $\phi=0^\circ$, it decreases by 2 kcal/mol out of 72 upon increasing θ from 105° to 150°, while the corresponding decrease of E_X is 16 kcal/mol. The same trends are found for the three other ϕ angles. These mismatches are partly absorbed, within E₁, by the large magnitudes of E_{MTP} /E_C, leaving out relative errors of 3% in terms of E₁. We have observed that abolishing the smearing by recovering populations of 2 on the in-plane components of the sp² lone pairs, leaving out zero for the smeared out-of-plane components, restores only a small fraction of the θ dependencies, but at the cost of a severe loss of E_{rep} , of up to 17 kcal/mol upon increasing ϕ from 0° to 60° (Supp. Info. SIII a-d). On the other hand, resorting to larger decrements of the lone pair radii along the two 'internal' oxygen sp² lone pairs enabled for a faster decrease of E_{rep} than displayed here (unpublished). This would correspond to the fact that the two 'inner' sp² O lone pairs would have reduced a radial extension, owing to their mutual Pauli repulsions, than the 'outer' lone pairs. Adopting this would however, require a new cycle of parametrization. It is a clue for further refinements, planned for a subsequent work at the correlated level within the context of SAPT-DFT^[39-40] approaches. E_{ct}(SIBFA) has a different θ dependency than E_{ct}(RVS). However, the numerical differences are small, and they are compensated for by the polarization energy which is shallow in QC, while it decreases in magnitude upon increasing θ . The agreement between $\Delta E(SIBFA)$ and $\Delta E(RVS)$ is to < 3% in relative errors for all ϕ values reported. It is to be

noted that the present tests are done at the equilibrium Zn-O distance of 1.80 Å. In polyligated complexes, the Zn-ligand will increase, clearly enabling for improved SIBFA and QC agreements regarding the individual Zn(II) –formate complexes.

Hydroxamate-Zn(II). Hydroxamate is an essential Zn-binding moiety of several high-affinity Zn-metalloenzyme inhibitors, some of which are clinically promising.^[41-42] and Refs. therein. Therefore, it could be very important to unravel the in- and out-of-plane angular dependencies of ΔE and its contributions, and how well these could be accounted for by PMM. The first parallel QC and PMM study of mono- and polyligated complexes of Zn(II) with hydroxamate was published in 1998,^[43] but did not probe the potential energy surface at the same detailed level as here. As in Ref.^[43] we will focus here on the deprotonated form.

a) Zn(II)-binding to the CO oxygen (Supp. Info. SIV a-d). E_{rep} and E_1 (SIBFA) have a near-parallelism to their RVS counterparts. E_{ct} (SIBFA) presents larger divergences with respect to E_{ct} (RVS) for the smaller θ values, which correspond to less-favored ΔE (RVS) values. Such divergences do not appear to compromise the near-parallelism between ΔE (SIBFA) and ΔE (RVS) found as a function of θ for all four ϕ angles. In fact, the sole exception appears at ϕ =90°, for θ values of 140°-150°, where the two curves actually cross. As shown in the Supp. Info., it is also seen that underestimations of E_{ct} (RVS) can to some extent be recovered by corresponding overestimations of E_{pol} (RVS) by E_{pol} (SIBFA).

b) Zn(II)-binding to the NO oxygen (Figures 5a-b; Supp. Info. SV a-d). The interaction energies are much more favorable than for Zn(II)-binding to the CO oxygen. E_X is much more sensitive to both θ and ϕ variations, and E_{rep} has very close values to it throughout, except for $\phi = 0^{\circ}$, and in a limited 15° range, namely for 105° < θ < 120° for which $\Delta E(RVS)$ and $\Delta E(SIBFA)$ are themselves much less favorable anyway (Figure. 5-a). Since E_{MTP*} closely matches E_C throughout, $E_1(SIBFA)$ parallels $E_1(RVS)$, again with the sole exception of $\phi = 0^{\circ}$ and 105° < $\theta < 120^{\circ}$ reflecting the corresponding behavior of $E_{rep}(SIBFA)$. The angular dependencies of $E_{ct}(SIBFA)$ have only slightly lesser parallelism with those of $E_{ct}(RVS)$ than was the case with the short-range repulsion. There are visible exceptions. The first is again for $\phi = 0^{\circ}$, 105° < $\theta < 120^{\circ}$. The second is for $\phi = 90^{\circ}$, where $E_{ct}(RVS)$ present a minimum for $\theta = 120^{\circ}$, while $E_{ct}(SIBFA)$ has a smoother behavior (Supp. Info. SV. a-d). The range of $\phi = 90^{\circ}$ values nevertheless concerns ΔE values that are significantly (app. 50 kcal/mol) less stable than for $\phi = 0^{\circ}$ and 30°.

For both Zn(II) binding to the CO and NO oxygens, there is a close agreement between $\Delta E(SIBFA)$ and $\Delta E(RVS)$ in all relevant energy-wise θ and ϕ values, the relative errors being always < 3%.

Imidazole-Zn(II) (Figures 6a-b). E_{rep} reproduces closely E_X as a function of both ϕ and θ . It could not account satisfactorily for the ϕ dependency of E_X if the N3 sp lone pair were not smeared. Since E_{MTP}^* has itself closely

similar ϕ and θ dependencies as E_c , the corresponding $E_1(RVS)$ dependencies are accordingly closely accounted for by $E_1(SIBFA)$. $E_{ct}(SIBFA)$ has a lesser parallelism to its RVS counterpart, but the mismatches appear essentially for the $\Delta E(RVS)$ values that are the least favored in terms of ϕ and θ (Supp. Info. SVI. a-d). Thus $\Delta E(SIBFA)$ closely reproduces $\Delta E(RVS)$ for all θ and ϕ values considered. This has an important implication for the modelling of Zn-binding sites in metalloproteins, in which His residues are recurrent Zn-binding ligands. Zn-binding mostly occurs in the imidazole plane and close to the direction of the external bisector of the coordinating nitrogen. Nevertheless out-of-plane binding, corresponding to ϕ values of app. 45°, could also be found by high-resolution X-ray crystallography, possibly due to constraints by the anchoring of the His sidechain to a relatively rigid backbone [see, eg.^[44-45] and refs. therein]. The present calculations quantify the $\Delta E(QC)$ cost incurred by out-of-plane binding, and show that they can be accounted for by anisotropic molecular mechanics.

As a conclusion for the present section, the 'smearing' of the sigma lone pairs (sp² for formamide, formate and the CO oxygen of hydroxamate, sp for the purine-like nitrogen of imidazole) enables E_{rep} to closely parallel both in- and out-of-plane dependencies of E_X in the energy-relevant range. E_{MTP*} and $E_{pol}(SIBFA)$ also stably reproduce the angular behaviors of their RVS counterparts. Owing to the separability of the SIBFA potential, this results into a close reproduction of $\Delta E(RVS)$ by $\Delta E(SIBFA)$. It could also be mentioned that some remaining mismatches of E_{ct} or to E_{pol} should be absorbed in polyligated complexes, on account of non-additivity due to screening of the cation-exerted field on any Zn-binding ligand by the other ones.

3. *Perpendicular Zn(II) binding to imidazole, formamide and phenol.* For completeness, we also performed tests which 'probe' the π electrons of conjugated or aromatic ligands by Zn(II) at the vertical of their atoms. Such binding positions are anticipated to be significantly less favorable than Zn(II)-binding to the σ lone-pairs and *a priori* unlikely to be populated by this cation in the course of MD or MC simulations. They should nevertheless be investigated to ensure that no artificial magnification of $\Delta E(RVS)$ by $\Delta E(SIBFA)$ might occur. As reported for the cytosine and guanine bases,^[10] a divalent cation acts can be used as a probe for E_{rep} and E_{ct} involving the π lone-pairs: comparison to their RVS counterparts enables to calibrate their radial extension and extent of effective radius increments/decrements of their atom bearer. This is necessary in a more general context, whenever the ligand is involved in even partially stacked interactions with other ligands. Furthermore, a divalent cation enables to evaluate how the GDMA multipolar representation fares above the plane of such ligands. Thus it was reported that for such ligands, GDMA multipoles derived from extended basis sets could be very sensitive to the input parameters used to derive them,^[46] namely the effective radii on the H atoms and the value of the switch. A non-optimal choice, while delivering E_{MTP^*} values close to E_C in the in-plane position, could severely overestimate it in over-the-plane positions, due to an imbalance of the charge-quadrupole component [unpublished]. The results are reported in Tables IIa-b. Table IIa reports the results at equilibrium distance for

Zn(II) bound at the vertical of each of the five atoms of imidazole. Table IIb reports the corresponding results for over the plane binding of Zn(II) to relevant atoms of two O-containing ligands, phenol and formamide. For completeness, Table III gives the radial dependencies of $\Delta E(QC)$ and $\Delta E(SIBFA)$ and their contributions for the Zn(II)-formamide complex, the C-O-Zn(II) angle being 135°.

The results for imidazole show that $\Delta E(SIBFA)$ retains a close agreement with $\Delta E(RVS)$, except for binding over C6, which has an error of app. 10%. This position is nevertheless 50 kcal/mol less stabilizing for Zn(II) than in-plane binding to N3, and 17 kcal/mol less so than perpendicular binding to N3. For all five binding positions, there is a lesser term-to-term agreement between the individual SIBFA contributions and their RVS counterparts than was the case with in-plane binding to N3. This is due on the one hand to E_{MTP*} which as mentioned above tends for conjugated systems to overestimate the charge-quadrupole component. As further refinements we consider resorting to Iterative Stockholder Atom (ISA) partition-derived multipoles^[47-48] and GEM-derived multipoles.^[49-50] E_{rep} accordingly overestimates E_X. It is recalled that the prefactor of E_{rep} embodies dependences upon the electronic populations of the pair of interacting atoms, and can thus be sensitive to the stability of the multipolar expansion. The calibration of E_{rep} will thus be readdressed in conjunction with the derivation of a new set of multipoles. E_{pol}(SIBFA) is invariably larger in magnitude than E_{pol}(RVS), but this compensates partly the less stabilizing E_1 (SIBFA) than E_1 (RVS). It is due to a maximal exposure of the bond polarizabilities when a cation binds above the plane at short distances of approach (1.90 Å in the present case). Such overestimations of E_{pol} for the vertical positions of approach are not likely to impair future MD simulations, since as mentioned above, in polyligated complexes, the field exerted by the dipositive Zn(II) charge on any one ligand will be shielded by the fields exerted by the other ligands.

The same observations can be done regarding binding to formamide over the C atom: the acceptable reproduction of $\Delta E(RVS)$ by $\Delta E(SIBFA)$ stems from an overestimated $E_{pol}(SIBFA)$ compensated by an overestimated and repulsive E_1 . In any case the ΔE values are app. 80 kcal/mol less favorable than for in-plane binding to O. For the binding over N, $\Delta E(SIBFA)$ has a closer agreement to $\Delta E(RVS)$ and a better term-to-term match of its contributions to the RVS ones. Here again, $\Delta E(RVS)$ remains 80 kcal/mol less favorable than for in-plane binding of O.

We have considered three perpendicular Zn(II) binding positions over phenol: over O, over the O-connected C atom, and over one *meta* C atom. $\Delta E(SIBFA)$ is underestimated by app. 10% with respect to $\Delta E(RVS)$. This is mostly due to the underestimation of $E_{pol}(RVS)$. As discussed above regarding over the plane Zn-binding to imidazole, a significant reduction of such an error can be anticipated in polyligated complexes, owing to the reduction of the Zn-electrostatic field on any ligand by the fields exerted by the other ligands. The unanticipated preference of over-C2 binding on in-plane binding to O stems from E_{pol} and E_{ct} . While these features could be accounted for by SIBFA, such a preference could be anticipated not to prevail either in polyligated complexes.

The improvement brought to E_{rep} by smearing the lone pairs is illustrated in Figures 7a-c, for the Zn(II) complexes with formamide, formate, and imidazole at their equilibrium distances. For formamide and formate, the CO—Zn θ angles are 135° and 150°. For imidazole, Zn(II) is along the external bisector of the ligating N atom. We perform stepwise variations of the ϕ angle until the perpendicular to the plane is reached. While for formamide and formate, E_{rep} parallels the evolution of EX, E_{rep}^* (without smearing the lone pairs) undergoes 10 kcal/mol drops in magnitude past ϕ =60°. With imidazole, E_{rep}^* undergoes a 10 kcal/mol drop at 40° before rising again until ϕ =90° is reached. Such a drop is in marked contrast with the much smaller decrease of both EX and E_{rep} in the 0-60° range, both contributions increasing again from 60° to 0°.

Polyligated Zn(II) complexes. We have in several preceding studies^[3,27, 51-52] addressed the issue of the nonadditivity of ΔE , δE_{nadd} , and its contributions upon passing from mono- to polyligated complexes. The control of δE_{nadd} is clearly critical in view of reliable MD simulations of Zn complexes in a diversity of problems ranging from biological to material science. Our earliest studies were with the CEP 4-31G(2d) basis set.^[27] and refs. therein. We have been more recently^[7, 9] targeting the results from a much more extended basis set, aug ccpVTZ(-f), where second-order effects could have a larger magnitude than with the CEP 4-31G(2d) set. Thus it is all the more necessary to control δE_{nadd} with this basis, and ensure if the impact of out-of-plane Zn(II) binding is controlled as well. We have considered the following complexes:

-the oligohydrated Zn(II) complexes with four and six water molecules, namely $[Zn(H_2O)_4]^{2+}$ in either the square planar alternating or the pyramidal arrangement, and $[Zn(H_2O)_6]^{2+}$ in the octahedral arrangement (Figures 8a-c and Table IVa);

-the complexes of Zn(II) with three and four imidazole ligands, and with three methanethiolate ligands, denoted as [Zn(Imidazoles)₃]²⁺, [Zn(Imidazoles)₄]²⁺, and [Zn(Methanethiolates)₃]⁻¹ respectively (Figures 8d-f and Table IVb);

-three polyligated complexes, with 'mixed' N, S, and O-containing ligands. Each ligand is denoted with one letter, corresponding to the one-letter code for the protein residue of which it is the end side-chain, namely H for imidazole, C for methanethiolate, and E for glutamate. The first complex, Zn[HHCC], recurs in several Zn-finger proteins.^[53] The second, denoted as Zn[HHE]⁺, is found in the Zn-finger motif of the anthrax lethal factor (LT),^[54] a target for the design of anthrax inhibitors.^[55] The third corresponds to the Zn-binding site of the phosphomannose isomerase (PMI) bacterial enzyme, which catalyzes the reversible interconversion of mannose into glucose, and is a target for the design of antibacterial and antiparasitic diseases.^[56]

The energy-minimized structures of complexes Zn[HHCC] and Zn[HHE]⁺ are represented in Figures 9a-b, and the interaction energy values are given in Table V.

-[$Zn(H_2O)_n$]²⁺ complexes (n=4, 6). Table IVa shows that, in line with our previous results,^[27] $\Delta E(RVS)$ and its contributions can be closely reproduced by $\Delta E(SIBFA)$ and its contributions. For n=4, the preference in favor of the pyramidal arrangement over the planar one (14.7 and 14 kcal/mol in QC and SIBFA respectively), is mostly due to E_{pol} by both RVS and SIBFA, and, to a lesser extent, to E_1 and E_c . These conclusions are consistent with those of a study devoted to the tetrahydrates of the alkali cations in the series Li⁺ -- Cs⁺, although the preference due to E_{pol} was not dominating.^[9] The non-additivities of both E_{pol} and E_{ct} appear properly controlled. Thus, consistent with our earlier results that resorted to the CEP 4-31G(2d)^[3, 27,57] basis set, in both QC and SIBFA, E_{ct} increases by only a factor of app. 2 with respect to the monoligated case upon multiplying the number of ligands by 4 or even 6. Table IVa lists the values of E_{pol} prior to, and following, iterated calculations on the induced dipoles. The QC values are denoted as $E_{pol}(RVS)$ and $E_{pol}(VR)$, respectively, and the corresponding SIBFA values by E_{pol}^* and E_{pol} : the latter is 30% smaller in magnitude than the former, which as previously noted^[52] is a signature for anticooperativity.

-[Zn(imidazole)_n]²⁺ complexes (n=3, 4). For completeness, for both n=3 and 4, we list two sets of distances, as they give rise to closely similar ΔE values.

Table IVb shows that for both tri- and tetraligated complexes of imidazoles, $\Delta E(SIBFA)$ matches $\Delta E(RVS)$ with relative errors of 2%. There is a slight imbalance between short-range repulsion and charge-transfer, an underestimation of $E_{rep}(SIBFA)$ being compensated by a corresponding underestimation of $E_{ct}(SIBFA)$. The close agreements of $E_{pol}(SIBFA)$ and $E_{pol}(RVS)$ both at non-iterated and iterated levels is noteworthy. The trends in E_{ct} upon passing from n=3 to n=4 are correctly accounted for. E_{ct} undergoes closely similar and modest increases with both methods, namely -3.5 and -1.3 kcal/mol. Such increases upon adding up one imidazole ligand are markedly smaller than the corresponding values of E_{ct} in the monoligated Zn(II)-imidazole complexes for $d_{N-Zn}=2.10$ Å, namely -17.6 and -16.3. kcal/mol: this attests again to the strong anticooperative nature of E_{ct} in polyligated Zn(II) complexes.

The last complex reported in Table IVb is that of Zn(II) bound to three methanethiolates, Zn[CCC]⁻. It represents one extreme test for the procedure and its handling of non-additivity, owing to the presence of three anionic, highly polarizable ligands complexing Zn(II) in a compact arrangement. $\Delta E(RVS)$ and $\Delta E(SIBFA)$ undergo large magnitude increases. The relative error now amounts to 4%, the largest one reported in this study for polyligated complexes. $E_1(SIBFA)$ reproduces $E_1(RVS)$ with a relative error of 1%, but this occurs due to a compensation of errors between E_{MTP}^* and E_C on the one hand and E_{rep} and E_X on the other. The main reason for the relative underestimation of $\Delta E(SIBFA)$ resides in E_{ct} , which amounts to -44.8 kcal/mol in SIBFA and to -64.2 kcal/mol with RVS after BSSE correction. -*Zn*[*HHCC*] and *Zn*[*HHE*]⁺ complexes. We next investigate two 'mixed' complexes in which Zn(II) is ligated by two different kinds of ligands: imidazole and methanethiolate for the first complex, and imidazole and formate for the second (Figure 9 and Table V). There is a close agreement between $\Delta E(SIBFA)$ and $\Delta E(RVS)$ for the Zn[HHCC] complex, with a relative error of 2%. The close match of $E_1(SIBFA)$ to $E_1(RVS)$ is due to a smaller E_{rep} than E_X compensating for a smaller magnitude of E_{MTP}^* than E_C . The Zn[HHE] ⁺ complex shows close agreements between $\Delta E(SIBFA)$ and $\Delta E(RVS)$ (less than 1% in relative error) and their individual contributions.

At this stage we can conclude that the Zn(II) polyligated complexes giving rise to the larger relative errors are those with methanethiolates, the largest one being 4%. This appears mostly due to E_{MTP}^* as compared to E_C . It could be due to the derivation of the methanethiolate multipoles using the GDMA procedure. An indication that this could indeed be the case is given by the faster radial decay of E_{MTP}^* than E_C in the monoligated complex Zn-methanethiolate complex (Supp. Info SI.i), in contrast to the near-parallel decay of E_{MTP}^* and E_C in all the other mono-ligated complexes. This could in turn impact E_{rep} , which depends upon E_{MTP}^* by a prefactor which includes the electronic population of the pair of interacting atoms.

The impact of smearing the lone pairs can be also illustrated in Table V. E_{rep}^* is 7-12 kcal/mol smaller in magnitude than E_{rep} . In the extreme case of the Zn[HHCC] complex, this can increase to 25-35 kcal/mol the underestimations of E_X . Because of compensations of errors occurring for the Zn[HHCC] complex owing to an underestimation of E_X by E_{MTP}^* , this actually led to an outwardly better agreement of corresponding energy, ΔE^{**} , ΔE^* than ΔE with $\Delta E(QC)$ for it. But this is not deemed meaningful in the prospect of the planned improvement to E_{MTP}^* discussed below. In the case of Zn[HHE]⁺, ΔE^* has a lesser agreement with $\Delta E(QC)$ than ΔE , the relative error being 3.5% compared to <1%.

-*Phosphomannose isomerase (PMI) binding site.* PMI is a Zn-dependent metalloenzyme, responsible for several bacterial and parasitic diseases,^[58-59] and refs. therein, but to date there are no therapeutically useful PMI inhibitors. Hydroxamate is a recurrent Zn-binding moiety of many Zn-metalloenzyme inhibitors [for a recent review, see].^[41] A submicromolar PMI hydroxamate inhibitor, 5-D-phosphonoarabinohydroxamate (5PAH), was originally designed and synthesized by Hardre et al..^[56] There are no available X-ray crystal structures for complexes of PMI with inhibitors. In a preceding study,^[36] we resorted to the X-ray crystal structure of Zn-bound PMI^[60] as a starting point to search for the most-stably bound structures of the complexes of PMI with 5PAH and three dehydroxylated derivatives. This was done by SIBFA Molecular Dynamics (MD) simulations with the hydroxamate ligated to Zn(II) and the phosphate bound at the entrance of the cavity by a lysine and an arginine residue.^[36] Three competing poses for each of the four inhibitors were unraveled by MD, whence a total of twelve complexes. The three poses differed mostly concerning the Zn-hydroxamate binding modes. To lend credence to their computed ΔE values and their ranking, we had performed parallel QC (HF and DFT-D3) and SIBFA computations on an enlarged recognition site encompassing up to 265 atoms including 28 waters. Agreements

between $\Delta E(QC)$ and $\Delta E(SIBFA)$ with relative errors < 3% were obtained. The present comparisons will be limited to the sole Zn(II)-binding site, each of the seven PMI residues being represented by its end side-chain, and 5PAH being represented by its hydroxamate moiety. These residues are: Asn111, His113, Ly136, Glu138, His285, Tyr287, and Asp300. There is thus a total of nine interacting molecules including hydroxamate and Zn(II), which is amenable to RVS analyses. Which agreements could still be obtained between QC and SIBFA regarding not only ΔE , but also their individual contributions? Comparisons with HF and DFT-D3 on an enlarged binding site with the entirety of the inhibitors and 28 structural waters, totaling up to 264 atoms, are reported separately.^[36] The three PMI poses, denoted A-C, are represented in Figures 10a-c. In A, the hydroxamate is bidentate bound to Zn(II), which is also ligated by Asn111, His113 and monodentate bound to Glu138. His285 has a larger Zn-N distance (3.1 Å) than His113 (2.4 Å). Tyr287 donates its hydroxyl proton to Asp300 and Lys136 donates a proton to the second, non-Zn bound anionic oxygen of Glu138. In B, hydroxamate is bound monodentate, through its N-bound O atom, to Zn(II), and Tyr287 now partakes into Zn-coordination instead of the C-bound hydroxamate oxygen. The other Zn-ligands are Gln111, His285 instead of His113, which has a Zn-N distance of 2.9 Å, and Glu138. As in A, Lys136 and Tyr287 donate a proton to Glu138 and Asp300, respectively. In C, Zn(II) is again bound to hydroxamate monodentate, but no longer by Tyr287. Its other ligands are Asn111, His113, and Glu138: the coordination number is thus 4, and not 5 as in A or B. In addition to the two H-bonded interactions between Lys136 and Glu138 and between Tyr287 and Asp300, an additional H-bond connects another Lys136 proton and the C-bound hydroxamate O atom. Thus, structures A-C highlight a diversity of Zn-binding motives. The varying coordination numbers will impact differently non-additivity. In addition, according to the structure, Zn-binding can for some ligands occur in out-of-plane positions, as observed in particular for the N-bound hydroxamate oxygen and for the Zn-ligating His113 nitrogen. Furthermore, some ligand-ligand interactions can themselves occur over the rings. This is observed: -a) between Asn111 and hydroxamate. The amino H *cis* to the CO bond of Asn is at representative π -H-bond distances to the conjugated ring of hydroxamate: in A, 2.8 and 3.0 Å from the N-bound O atom and the N atom, respectively; in B, 2.7 Å from the C and N atoms; and in C, 2.5, 2.6, 2.9 and 3.1 Å from the N-bound O, N, C, and C-bound O atoms, respectively; -and b) between His285 and Tyr287 in structures A and B. Such observations underline the need for a dependable representation of anisotropy, which was addressed in the first part of this study. The results are reported in Table VI. $\Delta E(SIBFA)$ reproduces closely $\Delta E(QC)$ and its trends along the sequence C > A > B, and has the proper balance between E₁ and E₂. Within E₁, there remains some compensation of errors, E_{MTP} being larger than E_C but E_{rep} being larger than E_X leading to closely similar (within 10 kcal/mol out of > 550) E_1 values. Further improvements will resort to alternative, newly-emerging procedures to derive distributed multipoles, such as GEM^[49-50] or the iterative Stockholder method^[47-48] and GEM-derived multipoles.^[49-50] It will be instructive to evaluate if these could remedy some shortcomings of the GDMA approach regarding conjugated or aromatic ligands which were mentioned in the Introduction. These would in turn impact Erep, which has

prefactors embodying the electronic population of the pair of interacting atoms. Work is in progress along these lines.

Handling of non-additivity effects. At this stage, and in line with a point raised by a Reviewer, it is essential to reevaluate how well the QC non-additivity, δE_{nadd} , can be accounted for. We have previously reported comparisons between QC and SIBFA regarding both anti-cooperativity in polyligated complexes of Zn(II) with N-, O-, and S- containing ligands ^[51-52] and cooperativity effect in multiply H-bonded complexes.^[61-63] These calculations were carried out with the CEP 4-31G(2d) basis set. Which outcome is expectable in the context of the aug-cc-pVTZ(-f) basis, which is more polarizable and has much more diffuse orbitals? We compare in Tables VIIa-c the values of δE_{nadd} in three representative complexes: the binding site of the anthrax lethal factor, Zn[HHE]⁺, with two imidazole and an anionic O ligand (Table VIIa); [Zn(Imidazoles)₄]²⁺, with four neutral Nligands (Table VIIb); and [Zn(HHCC)], with two neutral N-ligands and two anionic S-ligands (Table VIIc). δE_{nadd} (SIBFA) reproduces well its QC counterpart for all three complexes. There are two cases when the δE_{nadd} error exceeds 10%, namely for $E_{pol}(VR)$ versus E_{pol} in the $Zn[HHE]^+$ complex, and for $E_{pol}(RVS)$ versus E_{pol}^* in the $[Zn(Imidazoles)_4]^{2+}$ complex. Yet in both cases the agreements between $E_{pol}(VR)$ and E_{pol} , and between $E_{pol}(RVS)$ versus E_{pol}^* respectively are close, with only 2% relative errors. δE_{nadd} is larger in magnitude for E_{pol}(VR) and E_{pol} than for E_{pol}(RVS) and E_{pol}*. This as earlier mentioned is an additional signature for anticooperativity. $\delta E_{nadd}(QC)$ for E_{ct} is closely accounted for by $\delta E_{nadd}(SIBFA)$. It increases in both approaches by factors of 1.2-1.25 upon passing from Zn[HHE]⁺ to [Zn(Imidazoles)₄]²⁺ even though there is no anionic ligand in the latter. It next increases by factors 2.3-2.5 upon passing to [Zn(HHCC)] which has two 'soft' anionic ligands. Such agreements regarding $\delta E_{nadd}(E_{ct})$ validate its formulation which introduces dependencies upon the potentials and fields undergone by the interacting partners.^[2] While E_C is strictly additive, E_X has small δE_{nadd} magnitude, much smaller than those of the second-order contributions. This is fully consistent with the results from Ref.^[18] It could be accounted for by a three-body overlap formula involving the Zn(II) cation, and all pairs of ligand heavy atoms.^[18] For the present study, it was only included in the three test cases of Table VIIa-c but not at this stage in Tables IV-VI. Refining the calibration of $\delta E_{nadd}(E_{rep})$ and its explicit inclusion within ΔE should await prior refinements of E_{MTP}* with alternatives to GDMA multipoles, considering the dependencies of E_{rep} upon the electronic populations of the interacting partners.^[2, 4, 18]

Monoligated Mg(II) and Ca(II) complexes. The very first PMM calibrations of Mg(II) and Ca(II) with an explicit charge-transfer contribution were from 1985.^[64] Refinements resorting to the RVS procedure and the CEP 4-31G(2d) basis set were subsequently reported.^[57] Upon now resorting to the aug-cc-pVTZ basis set, we performed their calibration as follows. The cation-specific parameters for each contribution were fit in order to reproduce the radial dependency of its QC counterpart in their complexes with water along the water external bisector. The K(M-N) and K(M-S) constants used for E_{rep} were fit in order to match the radial dependency of E_X

in the cation complex with imidazole and methanethiolate. Tables VIIIa and VIIIb report the compared SIBFA-QC values for the complexes of Mg(II) with the same O-containing ligands (VIIIa) and N- and S-containing ligands (VIIIb) as in the corresponding Tables Ia and Ib for Zn(II). Table IX compares the radial dependencies of $\Delta E(QC)$ and $\Delta E(SIBFA)$ and their contributions for the monoligated Mg(II)-formamide complex. Tables Xa and Xb report the results for the same monoligated Ca(II) complexes as those of Tables VIIIa and VIIIb for Mg(II). A close reproduction of $\Delta E(QC)$ and its contributions is found with methanol, phenol, formamide and formate, as well as with pyridine and trimethylamine, which were not used to fit the Mg(II) and Ca(II) parameters. Tables XIa and XIb report the radial dependencies of $\Delta E(QC)$ and $\Delta E(SIBFA)$ and their contributions of the formate-Ca(II) complexes in the external (XIa) and bridge (XIb) complexes. The comparison between Mg(II) and Zn(II), which have similar ionic radii, is instructive. At equilibrium distance, for each monoligated complexes, E₁ has a significantly larger weight with Mg(II) than with Zn(II), while the chargetransfer is much smaller, consistent with the 'harder' character of Mg(II).^[65-67] The comparison between Mg(II) and Ca(II) shows that even though the equilibrium distance is significantly larger for Ca(II), by up to 0.5 Å, E_{ct} has a much larger magnitude in the monoligated Ca(II) complex than in the Mg(II) one. The radial dependencies of $\Delta E(QC)$ and $\Delta E(SIBFA)$ and their contributions are given in Supp. Info SVII a-i and SVIII a-h for the monoligated Mg(II) and Ca(II) complexes respectively.

Polyligated Mg(II) and Ca(II) complexes.

Mg(II). We have investigated the tetra- and hexahydrated complexes of Mg(II) in the same geometries as for Zn(II). The results are reported in Table XIIa. We also investigated the polyligated complexes in the binding site of the HIV-1 integrase.^[68] This complex is represented in Figure 11. The results are reported in Table XIIb. For the $[Mg(H_2O)_4]^{2+}$ and the $[Mg(H_2O)_6]^{2+}$ complexes $\Delta E(SIBFA)$ and its contributions closely reproduce their RVS counterparts, as was the case for the corresponding $[Zn(H_2O)_4]^{2+}$ complexes. Both $\Delta E(SIBFA)$ and $\Delta E(RVS)$ have app. 10 kcal/mol lesser magnitudes than in the corresponding Zn(II) complexes. With both procedures, this is due predominantly to E_{ct} and, to a lesser extent, to E_{pol} as well, which counteract the inverse preferences of E_1 . Within E_1 , E_C/E_{MTP}^* and E_X/E_{rep} have opposite preferences, the former favoring Zn(II) due to penetration and the latter favoring Mg(II) owing to its smaller size, and it is E_X/E_{rep} that imposes the preference of E_1 . This highlights again the need for the separability of the intermolecular potential. The same trends in Mg(II) versus Zn(II) are found in their complexes with three and four imidazoles (compare Tables IVb and XIIb). The $[2Mg(DDE)]^+$ is a more challenging case, owing to the presence of two divalent cations and three anionic ligands. Nevertheless, $\Delta E(SIBFA)$ can still retain a close agreement to $\Delta E(RVS)$, the relative error being 2%, although some compensations between E_1 on the one hand, and E_{pol} and E_{ct} on the other hand, could again be noted.

Ca(II). Ca(II) in water solution can adopt fluctuating coordination numbers, in the 6-8 range. This was shown in a MD simulation using AMOEBA.^[69] In view of long-duration MD simulations using SIBFA and now enabled by the massively parallel Tinker-HP code,^[30] it is essential to evaluate how well the values of $\Delta E(QC)$ and its individual contributions can be matched upon changes of the coordination Figures 12a-c show three structures of Ca(II) bound by eight water molecules, denoted as: a: $[Ca(H_2O)_{6+2}]^{2+}$, *b*: $[Ca(H_2O)_{7+1}]^{2+}$, and *c*: $[Ca(H_2O)_8]^{2+}$ namely with six and two first- and second-shell molecules, seven first and one second-shell water, and eight first-shell water molecules, respectively. These structures were extracted from a one nanosecond SIBFA MD simulation in a droplet of 512 water molecules. Table XIII compares the SIBFA and QC results in these arrangements. The ordering found by SIBFA is *b*: $[Ca(H_2O)_{7+1}]^{2+} > c$: $[Ca(H_2O)_8]^{2+} > a$: $[Ca(H_2O)_{6+2}]^{2+}$. The *b* versus *a* preference is dictated by E₁ overcoming the inverse preferences of both E_{pol} and E_{ct} within E₂. The preference of *b* over *c* is, conversely, now dictated by E₂ and goes in par with a small preference of E₁ as well. E_{pol} has in *b* values that are intermediate between those found in *c* and *a*. All these trends, as well as the numerical values and ΔE energy differences, are fully consistent with those of $\Delta E(RVS)$ and its contributions.

Calmodulin (CaM) is a pivotal Ca(II)-binding protein, relaying the role of Ca(II) as an intracellular messenger [Review in^[70-71]]. It has two Ca(II)-binding loops, denoted I-IV, at each N- and C-terminal ends. Ca(II)-binding loops III and IV have been previously studied by 2D-NMR spectroscopy and SIBFA energy-minimization.^[72] Ca(II) binding triggers CaM conformational changes, which it could be very interesting to monitor by long-duration MD. We are now given the possibility of validating SIBFA by high-level QC EDA. Thus as a first stage toward such simulations, we have considered Ca(II) binding loop III, having up to four formate anionic ligands and a hydroxyl one (Figure 13). The geometry of the complex was extracted from 1CLL.pdb.^[73] There is an excess of two negative charges. Which accuracy could still be expected? Table XIII (last two columns) shows that Δ E(RVS) can be reproduced by Δ E(SIBFA) with a relative error of 3%. The very significant decrease of E_{pol}(QC) and E_{pol}(SIBFA) compared of its values in the Ca(II) polyhydrates is noteworthy. It translates the very strong reduction of the dicationic field by the fields exerted by the four anionic charges.

Conclusions and perspectives.

The requirement for each contribution of an APMM/APMD potential to reproduce both anisotropy and nonadditivity features of its QC counterpart is critical for transferability.^[4] In the context of SIBFA, the anisotropy of the short-range contributions E_{rep} and E_{ct} is predominantly enabled by the angular features of electron spⁿ and π lone pairs. An additional contribution to the anisotropy of E_{rep} is conferred by its S²/Rⁿ formulation,^[2-16] expressing its dependence upon intermolecular bond-bond, bond-lone pair, and lone pair-lone pair overlaps: the S²/Rⁿ bond-bond repulsion should enable to account for anisotropy also in the absence of lone pairs, as in the model case of two hydrogen molecules in linear, parallel, and crossed configurations.^[74-75] Of course, results are not perfect as other effects in metal cation polyligated complexes such as three-body effects which are sometimes significant. ^[18] Such additional corrections as mentioned above exist in SIBFA^[18] and will upon refinements allow for further corrections in the final incoming set of parameters of the force-field. In the first integration of Ect in the context of SIBFA^[2, 28] we considered the linear water dimer and monitored its evolution for out-ofplane variations of the electron acceptor as a function of the HOH (electron donating water)-H angle. Ect(SIBFA) was able to closely parallel the evolution of Ect(QC). Its relatively shallow behavior stemmed from mutual compensations, upon augmenting this angle, between the decreased amplitude of E_{ct} contributed by one sp³ lone pair concomitant with the increased amplitude contributed by the other sp³ lone pair. But such a situation does not apply in out-of-plane complexes which involve sp² or sp lone pairs in conjugated molecules: if for a fixed θ angle an incoming H-bond donor or cationic probe is raised progressively above their plane, both E_{rep} and E_{ct} will undergo a regular decrease in their magnitude due to decreased overlap, until a π lone pair is encountered when the probe approaches or reaches the perpendicular. Such decreases are not observed with E_X and E_{ct}(RVS), and could thus significantly impair the agreement with these contributions. For conjugated molecules, we found that a simple remedy consisted into smearing the sp² and sp lone pairs, by relocating a fraction, α , of the initial population, namely two, in a plane perpendicular to the molecule, at a well-defined angle and at equal distances above and below it. For sp lone pairs, as in imidazole, the plane of the two smeared lone pairs cuts the molecular plane along the external bisector of the atom bearer. For sp² lone pairs, as for carbonyl or carboxylate oxygens, the corresponding plane cuts the molecular plane along the CO bond. We have considered four conjugated fragments: formamide, formate, hydroxamate, and imidazole. For each fragment, and at a given value of the θ angle for in-plane variations, the values of α and of the out-of-plane elevation z were fit so that E_{rep} reproduces the evolution of E_X upon ϕ -dependent out-of-plane evolutions of a probe molecule. Validations were done upon resuming these variations for a range of θ values encompassing the initial one. Such validations showed that both E_{rep} and E_{ct} could reproduce closely the in- and out-of plane dependencies of their QC counterparts for such values. Some differences could be observed concerning Ect, as in the case of Zn-formate or Zn-hydroxamate, but these either bore on small amounts, or were compensated by Epol within E2, and in any case did not impair the agreement between $\Delta E(SIBFA)$ and $\Delta E(RVS)$ in the relevant energy-wise geometries.

Along with these refinements, we integrated in the SIBFA library of fragments the polar or anionic moieties of several protein side-chains, constructed with augcc-pVTZ(-f) distributed multipoles and polarizabilities. We retained the general parameters which were previously optimized by the automated INoLLS procedure^[7] in this context, and performed limited recalibration of the atom-specific parameters owing to the smearing of the lone-pairs. We resorted to Zn(II) as a probe for ΔE (SIBFA) and its contributions to match their QC counterparts upon varying the Zn-ligand distance at fixed angles. The calibration and validation of the Mg(II), and Ca(II) cations were performed subsequently.

In this context, the most stringent tests and criteria for transferability clearly bear on polyligated complexes of the divalent cations. Zn(II) plays a prominent role, catalytic as well as structural, in numerous metalloproteins.^[1] We focused first on a series of polyligated complexes of Zn(II) with O-, N-, and S- containing ligands. We considered first model complexes with four and six water ligands, three and four imidazoles, and three methanethiolates [Tables IVa-b]. Several such complexes had been previously reported by us in the context of the CEP 4-31G(2d) basis set,^[27,52] but it could not be granted that close agreements would still hold with the aug-cc-p-VTZ basis owing to the onset of much larger overlap as well as polarizability effects. We further considered a complex extracted from the X-ray structures of Zn-fingers, [Zn(HHCC)]. A second complex, [Zn(HHE)]⁺, was extracted from the X-ray structure of the anthrax lethal factor. Close numerical agreements were retained between $\Delta E(SIBFA)$ and $\Delta E(RVS)$ in all cases with relative errors in the 1-3% range, with one exception [Zn(CCC)]⁻, in which it amounted to 4%. Limited compensations of errors between contributions, however, could be noted in some cases.

The most stringent test of this study related to the complex of hydroxamate with the binding site of phosphomannoseisomerase (PMI). PMI is a Zn-metalloenzyme responsible for bacterial and parasitic diseases^{[58-} ^{59]} and refs. therein, and hydroxamate is a recurrent Zn-binding moiety of pharmacological inhibitors. There are no known X-ray structures for inhibitor-bound PMI. Structural insight could be provided by PMD simulations. Thus three candidate structures, denoted A-C, were unraveled by SIBFA MD for the complexes of a hydroxamate-based inhibitor, 5PAH, and three dehydroxylated derivatives. Important, although not exclusive, differences between modes A-C concerned the modes of coordination of hydroxamate to Zn(II): bidentate in A, and monodentate in B and C, with five- and four-fold coordinations in A-B and in C, respectively. It was essential to evaluate if the generally close agreements found between $\Delta E(SIBFA)$ and $\Delta E(QC)$ for all twelve complexes in enlarged recognition sites with up to 265 atoms obtained as well upon focusing on the sole Zn-binding site encompassing the hydroxamate moiety and seven end side-chain residues, not only regarding ΔE but their individual contributions as well. Table VI confirmed this to be the case, with even smaller relative errors (1%) than in the other polyligated complexes. Closer consideration of Figures 10a-c showed that Zn-binding to its hydroxamate, formate, formamide and/or imidazole ligands could occur out of the ligand plane, thus not exclusively in the plane; it also showed in some complexes the onset of 'promiscuous' interactions between the Zn-coordinating residues, occurring near, or at, the vertical of the conjugated rings, such as between one formamide H and the hydroxamate or one imidazole CH and the Zn-coordinating N of the other imidazole. These findings, as well as those done recently for the binuclear Zn-binding site of a metallo- β -lactamase bound to a thiazole-thione inhibitor [Kwapien et al., submitted] stress the need for a correct smearing of the sp and sp² lone pairs enabling to account for in- and out-of-plane binding on a balanced level.

We next considered representative O- and N-polyligated complexes of Mg(II) in the same geometries as for Zn(II). This enabled to compare the weights of the energy contributions upon passing from Zn(II) to Mg(II). In the context of SIBFA, the preference in favour of the 'softer' cation, Zn(II), was due to E_{ct} , while E_1 favoured the 'harder' and smaller cation, Mg(II), due to a lesser E_{rep} value within E_1 , overcoming the more favorable E_{MTP}^* value for Zn(II) due to penetration. The results were fully consistent with the RVS ones. E_{disp}/E_{corr} will further stabilize Zn(II) over Mg(II).^[57, 67] These findings reemphasize again the need for separability. A stringent test for polycoordinated Mg(II) concerned the binding site of the HIV-1 integrase, in which two Mg(II) cations are chelated by three Asp/Glu end side-chains. This gave rise to very large electrostatic/polarization effects, yet the relative error remained confined to 2%.

The third cation considered is Ca(II), involved in a wealth of structural and enzymatic processes, and in cellular signaling. Ca(II) is prone to adopt variable, and possibly fluctuating, coordination numbers. This occurs in water, where n can vary between 6 and 8. A stringent test for transferability/nonadditivity bore on three competing octahydrated Ca(II) complexes, extracted as snapshots from MD. We have compared ΔE and its contributions in : *a*: $[Ca(H_2O)_{6+2}]^{2+}$, *b*: $[Ca(H_2O)_{7+1}]^{2+}$, and *c*: $[Ca(H_2O)_8]^{2+}$ with six, seven, and eight first-shell water molecules, respectively. $\Delta E(SIBFA)$ ranked the complexes along the same ranking as $\Delta E(QC)$, namely b > c > a. Its numerical values were close to the QC ones with relative errors < 1%., and the individual contributions of $\Delta E(SIBFA)$ closely followed the same trends as their QC counterparts. $\Delta E(SIBFA)$ also retained a close agreement with $\Delta E(QC)$, with a relative error <2%, in a model of a Ca(II)-binding loop in CaM, which encompasses four formate anions and one hydroxyl group, an extreme case of screening of a dipositive charge by up to four anionic charges. The agreements with QC are a strong incentive for extensions of SIBFA to a diversity of Ca(II)-dependent regulatory proteins, as well as to engineered Ca(II) sensors and transporters [reviewed in ^[76]].

The smeared lone-pair representation has been very recently used to a model bimetallic Zn-binding site of a Zndependent metallo- β -lactamase (MBL). In one monozinc site, a Zn(II) cation is chelated by three imidazoles and by a cyclic N atom of thiazole-thione (TZT), the conjugated anionic Zn-binding group of a novel group of MBL inhibitors. In the second site, the other Zn(II) cation is complexed by two other imidazoles, a formate and the extracyclic S atom of TZT [Kwapien et al., submitted]. $\Delta E(SIBFA)$ in the dizinc site as well as in both monozinc sites, whether TZT-ligated or unligated, was found to reproduce $\Delta E(QC)$ with <2% errors, a highly encouraging outcome given the complex interplay of first- and second-order effects and the magnitudes of the interaction energies. Presently, energy-minimizations on the complexes of a series of TZT-based MBL inhibitors are underway. At the present stage, Zn-coordination appears stable and fully consistent with the available data from X-ray crystallography of the complexes of MBL with such TZT inhibitors [Kwapien et al., work in progress]. It remains to be seen how stable such coordinations will remain upon passing to very long duration MD. This constitutes the next major step of our developments. It will be undertaken as soon as the integration of the full SIBFA potential and its gradients into the massively parallel Tiker-HP code [Tinker HP_team, Tinker-HP, <u>http://www.ip2ct.umpc.fr/tinkerHP</u>, 2016] is completed.

A last note concerns the representation of the localized lone pairs. They are explicitly accounted for in SIBFA in four contributions: along with E_{rep} and E_{ct} , they are represented in E_{disp} as fictitious atoms [see eg., Ref. ^[10]] and, in E_{pol} , through the dipole polarizabilities on the sigma-type lone-pairs. The importance of lone pairs to improve directionality in H-bonded complexes was demonstrated in the context of classical, non-polarizable force-fields by Lii and Allinger [^[77] and Refs. therein]. Lone-pairs are also represented in another polarizable potential, the classical Drude oscillator model, as centers for virtual sites improving the anisotropy of the polarization response^[78-79]

As a next step of this study, we are extending the construction of the SIBFA library with correlated aug-cc-pVTZ multipoles and polarizabilities. ΔE (SIBFA) and its contributions, now integrating correlation effects, as well as E_{disp} , are being recalibrated on the basis of SAPT EDA's. The present proof-of-principle validations pave the way for those performed at the correlated level. Recent are recently available regarding the stacked and H-bonded complexes of cytosine and guanine bases [Ref. 10] and the channeling of alkali cations across two stacked guanine tetramers [Gresh et al.; submitted]. They showed that upon passing from the HF to correlated levels, resorting to correlated, instead of HF, multipoles and polarizabilities could be sufficient to reproduce correlated ΔE results with accuracy. Is this generalizable, and to which extent could correlation impact the parameters (internal coordinates, electron population, increase/decrease of the van der Waals radii) of the in- and smeared out-of plane lone pairs? We are addressing this point for a diversity of ligands upon resorting to Electron Localization Function (ELF) results.^[25] We follow a similar strategy as in [Refs. ^[22-23]]. The results will be reported in a forthcoming study.

Acknowledgments.

We wish to thank the Grand Equipement National de Calcul Intensif (GENCI), Institut du Développement et des Ressources en Informatique Scientifique (IDRIS), Centre Informatique de l'Enseignement Supérieur (CINES, France, project No. x2009-075009), and the Centre de Ressources Informatiques de Haute Normandie (CRIHAN, Rouen, France, project 1998053).

We sincerely acknowledge the Research Council of Saint-Joseph University of Beirut, Lebanon (Project FS71), the Lebanese National Council for Scientific Research, CNRS-L (Project FS80), as well as the French Institute-Campus France au Liban and the French-Lebanese program CEDRE (Project FS30) for their financial support

and, particularly, for funding the PhD thesis of Léa El Khoury. The research by Karolina Kwapien is supported by the ANR 'Anti-MBL'.

Appendix. Internal coordinates of the lone pairs (σ and π) in formate, formamide, hydroxamate, and imidazole. They are denoted as theta, phi and r (first three columns). The following columns give the occupation numbers of the lone pairs, nocc and the increment of effective radii used for E_{rep} and E_{ct} , denoted as dwlp and twlp, respectively. The last column gives the designation of the atom bearer.

formate

phi	r	nocc	dwlp	twlp	bearer
90.00	0.51	0.00	0.00	0.0	<= C2
-90.00	0.51	0.00	0.00	0.0	<= C2
00.00	0.58	1.68	-0.05	0.0	<= 03
180.00	0.58	1.68	-0.05	0.0	<= 03
090.00	0.82	0.32	0.20	0.0	<= 03
270.00	0.82	0.32	0.20	0.0	<= 03
90.00	0.50	1.00	0.25	0.0	<= 03
-90.00	0.50	1.00	0.25	0.0	<= 03
000.00	0.58	1.68	-0.05	0.0	<= 04
180.00	0.58	1.68	-0.05	0.0	<= 04
090.00	0.82	0.32	0.20	0.0	<= 04
270.00	0.82	0.32	0.20	0.0	<= 04
90.00	0.50	1.00	0.25	0.0	<= 04
-90.00	0.50	1.00	0.25	0.0	<= 04
	phi 90.00 -90.00 00.00 180.00 270.00 90.00 -90.00 000.00 180.00 090.00 270.00 90.00 -90.00	phir90.000.51-90.000.5100.000.58180.000.58180.000.82270.000.8290.000.50-90.000.50000.000.58180.000.58180.000.58270.000.8290.000.5890.000.5890.000.5890.000.50-90.000.50-90.000.50	phirnocc90.000.510.00-90.000.510.0000.000.581.68180.000.581.68180.000.820.32270.000.820.3290.000.501.00-90.000.501.00000.000.581.68180.000.581.68180.000.581.68180.000.820.32270.000.820.3290.000.501.00-90.000.501.00	phirnoccdwlp90.000.510.000.00-90.000.510.000.0000.000.581.68-0.05180.000.581.68-0.05090.000.820.320.20270.000.820.320.2090.000.501.000.25-90.000.581.68-0.05180.000.581.68-0.05180.000.581.68-0.05180.000.581.68-0.0590.000.501.000.20270.000.820.320.2090.000.501.000.25-90.000.501.000.25	phirnoccdwlptwlp90.000.510.000.000.0-90.000.510.000.000.000.000.581.68-0.050.0180.000.581.68-0.050.090.000.820.320.200.0270.000.820.320.200.090.000.501.000.250.0-90.000.581.68-0.050.0180.000.581.68-0.050.0180.000.581.68-0.050.0270.000.820.320.200.090.000.501.000.250.090.000.501.000.250.090.000.501.000.250.090.000.501.000.250.090.000.501.000.250.0

formamide

theta	phi	r	nocc	dwlp	twlp	bearer
60.00	90.00	0.50	0.50	-0.25	0.0	<= C2
60.00	-90.00	0.50	0.50	-0.25	0.0	<= C2
113.00	00.00	0.58	1.60	-0.05	0.0	<= 03
113.00	180.00	0.58	1.60	-0.05	0.0	<= 03
113.00	90.00	0.78	0.40	0.10	0.0	<= 03
113.00	-90.00	0.78	0.40	0.10	0.0	<= 03
60.00	90.00	0.50	0.50	0.25	0.0	<= 03
60.00	-90.00	0.50	0.50	0.25	0.0	<= 03
60.00	90.00	0.68	1.00	0.20	0.0	<= N4
60.00	-90.00	0.68	1.00	0.20	0.0	<= N4

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hydroxamate

16						
theta	phi	r	nocc	dwlp	twlp	bearer
90.00	90.00	0.50	0.50	0.00	0.0	<= C2
90.00	-90.00	0.50	0.50	0.00	0.0	<= C2
113.00	000.00	0.58	1.75	0.00	0.0	<= 03
113.00	180.00	0.58	1.75	0.00	0.0	<= 03
113.00	090.00	0.75	0.25	0.05	0.0	<= 03
113.00	-90.00	0.75	0.25	0.05	0.0	<= 03
60.00	90.00	0.75	1.00	-0.05	0.0	<= 03
60.00	-90.00	0.75	1.00	-0.05	0.0	<= 03
60.00	90.00	0.50	0.50	0.00	0.0	<= N4
60.00	-90.00	0.50	0.50	0.00	0.0	<= N4
105.00	000.00	0.60	1.00	-0.05	0.0	<= 05
105.00	060.00	0.60	1.00	-0.05	0.0	<= 05
105.00	120.00	0.60	1.00	-0.05	0.0	<= 05
105.00	180.00	0.60	1.00	-0.05	0.0	<= 05
105.00	240.00	0.60	1.00	-0.05	0.0	<= 05
105.00	300.00	0.60	1.00	-0.05	0.0	<= 05

imidazole

theta	phi	r	nocc	dwlp	twlp	bearer
110.00	131.00	0.97	0.50	0.00	0.0	<= C2
110.00	229.00	0.97	0.50	0.00	0.0	<= C2
127.54	00.00	0.72	1.50	-0.10	0.0	<= N3
120.00	43.00	1.02	0.25	+0.05	0.0	<= N3
120.00	-43.00	1.02	0.25	+0.05	0.0	<= N3
45.00	90.00	0.85	0.50	0.00	0.0	<= N3
45.00	-90.00	0.85	0.50	0.00	0.0	<= N3
45.00	90.00	0.85	0.50	0.00	0.0	<= C4
45.00	-90.00	0.85	0.50	0.00	0.0	<= C4
45.00	90.00	0.85	1.00	0.00	0.0	<= N5
45.00	-90.00	0.85	1.00	0.00	0.0	<= N5
45.00	90.00	0.85	0.50	0.00	0.0	<= C6
45.00	-90.00	0.85	0.50	0.00	0.0	<= C6

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Figure captions.

Figure 1. Representation of the lone-pair locations in monoligands; **a:** formamide, **b:** formate-external, **c:** hydroxamate and **d:** imidazole. sp^{2*} and α represent the smeared lone-pairs and nocc, respectively.

Figure 2. Representation of the variations undergone by Zn (II) probe around monoligands; **a:** Zn²⁺-formamide, **b:** Zn²⁺-formate-external, **c:** Zn²⁺-hydroxamate (Zn²⁺ Probing around O3) and **d:** Zn²⁺-imidazole. Out-of-plane variations of φ are represented in different colors; magenta for (φ =0°), pink for (φ =30°), purple for (φ =60°) and iceblue for (φ =90°).

Figure 3. Zn^{2+} -formamide complex. Compared evolution of ΔE (RVS) and ΔE (SIBFA) and of the repulsion contribution on variation of the Zn2+-O θ angle; a: ($\varphi=0^{\circ}$) and b: ($\varphi=90^{\circ}$). Zn2+-O distance is 1.90 Å. RVS: Reduced Variational Space (RVS). All energy values are represented in Supp. Info. (Table SII a-d).

Figure 4. Zn^{2+} -formate external complex. Compared evolution of ΔE (RVS) and ΔE (SIBFA) and of the repulsion contribution on variation of the Zn^{2+} -O θ angle; **a:** ($\varphi=0^{\circ}$) and **b:** ($\varphi=90^{\circ}$). Zn^{2+} -O distance is 1.80 Å. RVS: Reduced Variational Space (RVS) analysis. All energy values are represented in Supp. Info. (Table SIII a-d).

Figure 5. Zn^{2+} - hydroxamate complex. Compared evolution of ΔE (RVS) and ΔE (SIBFA) and of the repulsion contribution on variation of the Zn²⁺-O5 θ angle; **a:** (φ =0°) and **b:** (φ =90°). Zn²⁺-O distance is 1.80 Å. RVS: Reduced Variational Space (RVS) analysis. All energy values are represented in Supp. Info. (Table SIV a-d).

Figure 6. Zn^{2+} - imidazole complex. Compared evolution of ΔE (RVS) and ΔE (SIBFA) and of the repulsion contribution on variation of the Zn²⁺-O θ angle; **a:** (φ =0°) and **b:** (φ =90°). Zn²⁺-O distance is 1.90 Å. RVS: Reduced Variational Space (RVS) analysis. All energy values are represented in Supp. Info. (Table SV a-d).

Figure 7. a. Zn^{2+} -formamide complex. Compared evolution of the repulsion contribution on variation of the Zn^{2+} -O φ angle with ($\theta = 135^{\circ}$); Zn^{2+} -O distance is 1.90 Å. **b.** Zn^{2+} -formate external complex. Compared evolution of the repulsion contribution on variation of the Zn^{2+} -O φ angle with ($\theta = 150^{\circ}$); Zn^{2+} -O distance is 1.80 Å. **c.** Zn^{2+} -imidazole complex. Compared evolution of the repulsion contribution on variation of the repulsion contribution on variation of the Zn²⁺-O φ angle with ($\theta = 127^{\circ}$); Zn^{2+} -O distance is 1.90 Å. RVS: Reduced Variational Space (RVS) analysis. (*) Energy values when lone-pairs are not smeared.

Figure 8. Polycoordinated complex of Zn(II) with: a) four planar water molecules, b) four pyramidal water molecules, c) six octahedral water molecules, d) three methanethiolates, e) three imidazoles and f) four imidazoles.

Figure 9. Polycoordinated complex of Zn(II) with: a) two imidazoles and two methanethiolates [HHCC], and b) two imidazoles and one formate (Anthrax lethal factor active site [HHE]).

Figure 10. Polycoordinated complex of Zn(II) with phosphomannose isomerase (PMI). Three structures provided by SIBFA MD are represented in a, b and c.

Figure 11. Polycoordinated complex of Mg(II) with three formate in the integrase catalytic site (Catalytic triad [DDE]).

Figure 12. Polycoordinated complex of Ca(II) with 8 water molecules. a) $[Ca(H2O)_8]^{2+}$, b) $[Ca(H2O)_{7+1}]^{2+}$ and c) $[Ca(H2O)_{6+2}]^{2+}$.

Figure 13. Polycoordinated complex of Ca(II) with four formate and one methanol in the calmoduline active site [DDDTE].

Table captions.

Table I. Complexes of Zn(II) with: Ia : -Oxgen ligands:water, methanol, formamide, phenol, formate E, formateB; Ib: -Nitrogen and Sulfur ligands: imidazole, pyridine, trimethylamine, methanethiolate.

Table II. Complexes of Zn(II) with: IIa: imidazole over C2, N3, C4, N5, C6 at equilibrium distances. IIb: Formamide over C and N phenol over C2, C3, O30.

Table III. Monoligated Zn(II) complexes: Formamide. Distance variations of $\Delta E(QC)$, $\Delta E(SIBFA)$ and their contributions.

Table IV. Polyligated Zn(II) complexes with IV. a: pyramidal $[Zn(H_2O)_4]^{2+}$, $planar [Zn(H_2O)_4]^{2+}$, $[Zn(H_2O)_6]^{2+}$; IV.b: $[Zn(Imidazoles)_3]^{2+}$, $[Zn(Imidazoles)_4]^{2+}$, $[Zn(Methanethiolates)_3]^{2+}$

Table V. Polyligated Zn(II) complexes: [Zn(HHCC)] and [Zn(HHE)]⁺.

Table VI. PMI- Zn(II) complexes.

Table VII.a-c Nonadditivity values of VII.a: [Zn(HHE)]⁺; VIIb: [Zn(Imidazoles)₄]²⁺; VIIc: [Zn(HHCC)].

Table VIII. Complexes of Mg(II) with: VIIIa: -O-ligands: water, methanol, formamide, phenol, formate B, formate E; VIIIb: -N and S ligands: imidazole, pyridine, trimethylamine, methanethiolate.

Table IX. Monoligated Mg(II) complexes: Formamide. Distance variations of $\Delta E(QC)$, $\Delta E(SIBFA)$ and their contributions.

Table X. Complexes of Ca(II) with: Xa : -O-ligands: water, methanol, formamide, phenol, formate B, formate E; Xb: -N and S ligands: imidazole, pyridine, trimethylamine, methanethiolate.

Table XI. Monoligated Ca(II) complexes with XIa. formate external; XIb. Formate bridge. Distance variations of $\Delta E(QC)$, $\Delta E(SIBFA)$ and their contributions.

Table XII. Polyligated complexes of Mg(II) with XIIa: pyramidal $[Mg(H2O)_4]^{2+}$, planar $[Mg(H2O)_4]^{2+}$, $[Mg(H2O)_6]^{2-}$: $[Mg(Imidazoles)_3]^{2+}$, $[Mg(Imidazoles)_4]^{2+}$, $[2Mg(DDE)]^+$.

Table XIII. Polyligated complexes of Ca(II): $[Ca(H_2O)_{6+2}]^{2+}$, $[Ca(H_2O)_{7+1}]^{2+}$, $[Ca(H_2O)_8]^{2+}$, $[Ca(DDDTE)]^{2-}$.

Supporting Information.

- Table S.I.a. Monoligated Zn(II) complexes: H₂O.
- Table S.I.b. Monoligated Zn(II) complexes: Methanol.
- Table S.I.c. Monoligated Zn(II) complexes: Formate external.
- **Table S.I.d.** Monoligated Zn(II) complexes: Formate bridge.
- Table S.I.e. Monoligated Zn(II) complexes: Phenol.
- Table S.I.f. Monoligated Zn(II) complexes: Imidazole.
- Table S.I.g. Monoligated Zn(II) complexes: Pyridine.
- Table S.I.h. Monoligated Zn(II) complexes: Trimethylamine.
- Table S.I.i. Monoligated Zn(II) complexes: Methanethiolate.

Table S.II.a. Monoligated Zn(II) complexes: Formamide. Variation of the Zn(II)-O θ angle; $\Phi=0^{\circ}$; d=1.90Å**Table S.II.b.** Monoligated Zn(II) complexes: Formamide. Variation of the Zn(II)-O θ angle; $\Phi=30^{\circ}$; d=1.90Å

Table S.II.d. Monoligated Zn(II) complexes: Formamide. Variation of the Zn(II)-O θ angle; Φ =90°; d=1.90Å

Table S.II.c. Monoligated Zn(II) complexes: Formamide. Variation of the Zn(II)-O θ angle; Φ =60°; d=1.90Å

- **Table S.III.a.** Monoligated Zn(II) complexes: Formate. Variation of the Zn(II)-O θ angle; $\Phi=0^{\circ}$ d=1.80Å
- **Table S.III.b.** Monoligated Zn(II) complexes: Formate. Variation of the Zn(II)-O θ angle; Φ =30° d=1.80Å
- **Table S.III.c.** Monoligated Zn(II) complexes: Formate. Variation of the Zn(II)-O θ angle; Φ =60° d=1.80Å
- **Table S.III.d.** Monoligated Zn(II) complexes: Formate. Variation of the Zn(II)-O θ angle; Φ =90° d=1.80Å
- **Table S.IV.a.** Monoligated Zn(II) complexes: Hydroxamate. Variation of the Zn(II)-O3 θ angle; $\Phi=0^{\circ}$ d=1.80Å
- **Table S.IV.b.** Monoligated Zn(II) complexes: Hydroxamate.Variation of the Zn(II)-O3 θ angle; Φ =30° d=1.80Å

Table S.IV.c. Monoligated Zn(II) complexes: Hydroxamate. Variation of the Zn(II)-O3 θ angle; Φ =60° d=180Å **Table S.IV.d.** Monoligated Zn(II) complexes: Hydroxamate. Variation of the Zn(II)-O3 θ angle; Φ =90° d=1.80 Å

Table S.V.a. Monoligated Zn(II) complexes: Hydroxamate. Variation of the Zn(II)-O5 θ angle; $\Phi=0^{\circ}$ d=1.80Å **Table S.V.b.** Monoligated Zn(II) complexes: Hydroxamate. Variation of the Zn(II)-O5 θ angle; $\Phi=30^{\circ}$ d=1.80Å **Table S.V.c.** Monoligated Zn(II) complexes: Hydroxamate. Variation of the Zn(II)-O5 θ angle; $\Phi=60^{\circ}$ d=1.80Å **Table S.V.d.** Monoligated Zn(II) complexes: Hydroxamate. Variation of the Zn(II)-O5 θ angle; Φ =90° d=1.80 Å

Table S.VI.a. Monoligated Zn(II) complexes: Imidazole. Variation of the Zn(II)-O θ angle; $\Phi=0^{\circ}$ d=1.90Å **Table S.VI.b.** Monoligated Zn(II) complexes: Imidazole. Variation of the Zn(II)-O θ angle; $\Phi=30^{\circ}$ d=1.90Å

Table S.VI.c. Monoligated Zn(II) complexes: Imidazole. Variation of the Zn(II)-O θ angle; Φ =60° d=1.90Å

Table S.VI.d. Monoligated Zn(II) complexes: Imidazole. Variation of the Zn(II)-O θ angle; Φ =90° d=1.90 Å

Table S.VII.a. Monoligated Mg(II) complexes: H₂O.

Table S.VII.b. Monoligated Mg(II) complexes: Methanol.

Table S.VII.c. Monoligated Mg(II) complexes: Formate external.

Table S.VII.d. Monoligated Mg(II) complexes: Formate bridge.

Table S.VII.e. Monoligated Mg(II) complexes: Phenol.

Table S.VII.f. Monoligated Mg(II) complexes: Imidazole.

Table S.VII.g. Monoligated Mg(II) complexes: Pyridine.

Table S.VII.h. Monoligated Mg(II) complexes: Trimethylamine.

Table S.VII.i. Monoligated Mg(II) complexes: Methanethiolate.

Table S.VIII.a. Monoligated Ca(II) complexes: H₂O.

Table S.VIII.b. Monoligated Ca(II) complexes: Methanol.

Table S.VIII.c. Monoligated Ca(II) complexes: Formamide.

Table S.VIII.d. Monoligated Ca(II) complexes: Phenol.

 Table S.VIII.e.
 Monoligated Ca(II) complexes: Imidazole.

Table S.VIII.f. Monoligated Ca(II) complexes: Pyridine.

Table S.VIII.g. Monoligated Ca(II) complexes: Trimethylamine.

Table S.VIII.h. Monoligated Ca(II) complexes: Methanethiolate.

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Figure 1. Representation of the lone-pair locations in monoligands; **a**: formamide, **b**: formate-external, **c**: hydroxamate and **d**: imidazole. sp^{2^*} and α represent the smeared lone-pairs and nocc, respectively.



Figure 2. Representation of the variations undergone by Zn (II) probe around monoligands; **a:** Zn^{2+} -formamide, **b:** Zn^{2+} -formate-external, **c:** Zn^{2+} -hydroxamate (Zn^{2+} Probing aroundO3) and **d:** Zn^{2+} -imidazole. Out-of-plane variations of phi are represented in different colors; magenta for (φ =0°), pink for (φ =30°), purple for (φ =60°) and iceblue for (φ =90°).



Figure 3. Zn^{2+} -formamide complex. Compared evolution of ΔE (RVS) and ΔE (SIBFA) and of the repulsion contribution on variation of the Zn2+-O θ angle; a: (ϕ =0°) and b: (ϕ =90°). Zn2+-O distance is 1.90 Å. RVS: Reduced Variational Space (RVS) analysis, All energy values are represented in Supp. Info. (Table SII a-d).



Figure 4. Zn^{2+} -formate external complex. Compared evolution of ΔE (RVS) and ΔE (SIBFA) and of the repulsion contribution on variation of the Zn^{2+} -O θ angle; **a:** (ϕ =0°) and **b:** (ϕ =90°). Zn^{2+} -O distance is 1.80 Å. RVS: Reduced Variational Space (RVS) analysis. All energy values are represented in Supp. Info. (Table SIII a-d).



Figure 5. Zn^{2+} hydroxamate complex. Compared evolution of ΔE (RVS) and ΔE (SIBFA) and of the repulsion contribution on variation of the Zn^{2+} -O5 θ angle; **a**: (ϕ =0°) and **b**: (ϕ =90°). Zn^{2+} -O distance is 1.80 Å. RVS: Reduced Variational Space (RVS) analysis. All energy values are represented in Supp. Info. (Table SVI a-d).



Figure 6. Zn^{2+} - imidazole complex. Compared evolution of ΔE (RVS) and ΔE (SIBFA) and of the repulsion contribution on variation of te Zn^{2+} -O θ angle; **a**: (ϕ =0°) and **b**: (ϕ =90°). Zn^{2+} -O distance is 1.90 Å. RVS: Reduced Variational Space (RVS) analysis. All energy values are represented in Supp. Info. (Table SV a-d).



Figure 7. a. Zn^{2+} -formamide complex. Compared evolution of the repulsion contribution on variation of the Zn^{2+} -O φ angle with (θ =135°); Zn^{2+} -O distance is 1.90 Å. **b.** Zn^{2+} -formate external complex. Compared evolution of the repulsion contribution on variation of the Zn^{2+} -O φ angle with (θ =150°); Zn^{2+} -O distance is 1.80 Å. **c.** Zn^{2+} -imidazole complex. Compared evolution of the repulsion contribution on variation of the repulsion contribution on variation of the repulsion contribution on variation of the Zn^{2+} -O φ angle with (θ =127°); Zn^{2+} -O distance is 1.90 Å. RVS: Reduced Variational Space (RVS) analysis. (*) Energy values when lone-pairs are not smeared.



Figure 8. Polycoordinated complex of Zn(II) with: a) four planar water molecules, b) four pyramidal water molecules, c) six octahedral water molecules, d) three methanethiolate, e) three imidazoles and f) four imidazoles.



Figure 9. Polycoordinated complex of Zn(II) with: a) two imidazoles and two methanethiolates [HHCC] and b) two imidazoles and one formate (Anthrax letal factor active site [HHE]).



Figure 10. Polycoordinated complex of Zn(II) with phosphomannoseisomerase (PMI). Three structures provided by SIBFA MD are represented in a, b and c.



Figure 11. Polylicoordinated complex of Mg(II) with three formate in the integrase catalytic site (Catalytic triad [DDE]).



Figure 12. Polylicoordinated complex of Ca(II) with 8 water molecules. a) $[Ca(H2O)_8]^{2+}$, b) $[Ca(H2O)_{7+1}]^{2+}$ and c) $[Ca(H2O)_{6+2}]^{2+}$.



Figure 13. Polylicoordinated complex of Ca(II) with four formate and one methanol in the calmoduline active site [DDDTE].

Table I.a.	Monoliga	ted Zn ²⁺ c	omplexes: (Dxygen lig	ands.							
	H	₂ O*	Meth	nanol	Forma	amide**	Fomate e	external**	Formate	bridge**	Ph	enol
	d = 1	.90 Å	<i>d</i> = 1	.90Å	<i>d</i> = 1	.90 Å	<i>d</i> = 1	.90 Å	<i>d</i> = 1	.94Å	<i>d</i> = 1	.80 Å
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA
$\text{Es/E}_{\text{MTP}^{\star}}$	-74.4	-74.8	-75.7	-75.2	-97.9	-96.4	-292.1	-291.2	-360.1	-358.3	-68.1	-67.6
Ex/E_{rep}	35.1	35.1	37.4	35.0	34.1	36.4	45.9	46.8	78.2	78.0	56.1	55.2
E ₁	-39.3	-39.7	-38.4	-40.1	-63.8	-60.0	-246.1	-244.4	-281.9	-280.3	-12.0	-12.4
E _{pol}	-41.6	-41.8	-51.5	-49.5	-61.8	-64.6	-64.8	-61.4	-80.3	-81.5	-86.9	-87.1
E _{ct}	-10.6	-11.9	-12.3	-11.6	-12.3	-15.9	-21.5	-22.4	-24.6	-23.1	-15.1	-13.0
ΔE	-91.3	-93.4	-102.2	-101.3	-137.9	-140.5	-332.6	-328.1	-386.8	-384.9	-114.0	-112.5

Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions

* Used to calibrate the effective vdW radii

** Used to calibrate the smeared lone pair (effective population and increment of the vdW effective radius)

Table I.b. Monoligated Zn ²⁺ complexes: Nitrogen and sulfur ligands.											
	lmida:	zole**	Pyri	dine	Trimeth	ylamine	Methane	ethiolate			
	<i>d</i> = 1.90 Å		<i>d</i> = 1.90Å		<i>d</i> = 1.90 Å		d = 2.	20 Å			
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA			
Es/E _{MTP*}	-112.9	-112.4	-98.9	-98.3	-104.3	-110.2	-323.8	-323.2			
Ex/E _{rep}	57.6	53.8	59.7	59.6	69.0	66.6	69.9	71.0			
E ₁	-55.3	-58.5	-39.2	-38.7	-35.3	-43.6	-254.0	-252.2			
E _{pol}	-78.3	-80.1	-84.6	-88.3	-82.5	-82.3	-91.6	-92.7			
E _{ct}	-22.1	-22.6	-23.6	-21.1	-25.9	-15.6	-50.9	-51.4			
ΔE	-155.6	-161.2	-147.3	-148.1	-143.7	-141.5	-396.4	-396.3			

Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions

** Used to calibrate the smeared lone-pairs (effective population and increment of the vdW effective radius)

Table II.a.	Monoligat	ted Zn ²⁺ con	nplexes: Imida	zole**.							
	С	2	N	3	C	4	Ν	15	C	6	
	<i>d</i> = 1.90 Å		<i>d</i> = 1	<i>d</i> = 1.90Å		<i>d</i> = 1.90 Å		<i>d</i> = 1.90 Å		<i>d</i> = 1.90Å	
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	
$\mathrm{Es/E}_{\mathrm{MTP}^*}$	-49.7	-61.4	-61.2	-66.6	-25.3	-29.8	-20.2	-25.0	-36.3	-46.5	
Ex/E _{rep}	48.0	66.4	42.9	63.2	40.1	60.3	40.9	66.1	49.3	69.9	
E ₁	-1.6	5.0	-18.4	-3.4	14.8	30.5	20.6	41.1	13.0	23.4	
E _{pol}	-94.0	-105.6	-88.9	-104.7	-92.4	-106.0	-87.8	-96.2	-94.2	-98.7	
E _{ct}	-25.4	-20.6	-18.6	-20.5	-22.0	-21.9	-15.3	-22.9	-27.0	-22.6	
ΔE	-121.0	-121.2	-125.9	-128.6	-99.6	-97.4	-82.4	-78.0	-108.2	-97.9	
Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions											
** Used to	** Used to calibrate the smeared lone-pairs (effective population and increment of the vdW effective radius)										

Table II.b.	Table II.b. Monoligated Zn ²⁺ complexes: Formamide** and Phenol.												
	Forma	mide-N	Forma	mide-C		Phen	ol-C2		Phen	ol-C3		Phene	ol-O30
	<i>d</i> = 1.90 Å		d= 1	<i>d</i> = 1.90Å		<i>d</i> = 2.10 Å			<i>d</i> = 2.10Å			d=1	.90 Å
	QC	SIBFA	QC	SIBFA		QC	SIBFA	-	QC	SIBFA	· <u>-</u>	QC	SIBFA
Es/E _{MTP*}	-11.2	-21.9	-2.3	-10.2		-38.8	-42.5		-35.0	-39.2		-45.9	-50.8
Ex/E _{rep}	41.3	50.3	29.3	61.6		36.3	34.1		34.7	37.0		45.0	51.9
E ₁	30.2	28.3	27.0	51.4		-2.5	-8.4		-0.2	-2.2		-0.9	1.1
E _{pol}	-64.9	-69.6	-69.0	-88.4		-103.6	-93.6		-103.6	-98.3		-106.3	-76.9
E _{ct}	-17.4	-13.3	-13.1	-24.5		-26.0	-17.7		-24.4	-19.0		0.5	-15.9
ΔE	-52.1	-54.6	-55.1	-61.5		-132.2	-119.7		-128.3	-119.5		-106.8	-91.6

Values (Kcal/mol) of the QC and SIBFA interaction energies and their contributions

** Used to calibrate the smeared lone-pairs (effective population and increment of the vdW effective radius)

Table III.	Monoligate	ed Zn²⁺ com	plexes: Form	amide**.							
						I		*		a (a 1	
	<i>d</i> = 1.70 A		d= 1	.80A	d= 1	.90 A	d=2	.00 A	d=	2.10 A	
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	
Es/E _{MTP*}	-124.0	-121.3	-109.4	-107.6	-97.9	-96.4	-88.5	-87.1	-80.7	-79.2	
Ex/E _{rep}	79.9	85.7	52.2	55.7	34.1	36.4	22.3	23.9	14.6	15.8	
E,	-44.0	-35.6	-57.3	-51.9	-63.8	-60.0	-66.2	-63.2	-66.2	-63.4	
E _{pol}	-75.7	-76.6	-68.1	-70.5	-61.8	-64.2	-56.2	-58.0	-51.1	-52.0	
E _{ct}	-18.4	-21.5	-15.1	-18.4	-12.3	-15.6	-10.0	-13.2	-8.3	-11.2	
ΔE	-138.1	-133.7	-140.5	-140.8	-137.9	-139.8	-132.4	-134.4	-125.	-126.6	
Values (k	/alues (kcal/mol) of the OC and SIBFA interaction energies and their contributions										

** Used to calibrate the smeared lone-pairs (effective population and increment of the vdW effective radius)

Table IV.a	Table IV.a. Polyligated Zn ²⁺ complexes.										
		[Zn(H	₂ O) ₄] ²⁺		[Zn(H2	2O) ₆] ²⁺					
	Pyramidal,	<i>d</i> = 2.10 Å	Planar, d	/= 2.20 Å	d= 2	<i>d</i> = 2.30 Å					
	QC	SIBFA	QC	SIBFA	QC	SIBFA					
Es/E _{MTP*}	-218.2	-219.7	-201.6	-202.8	-264.4	-265.9					
Ex/E _{rep}	57.5	56.5	43.6	42.2	46.1	43.3					
E ₁	-161.0	-163.2	-158.0	-160.6	-218.3	-222.7					
E _{pol}	-85.7	-83.8	-75.9	-74.4	-75.7	-74.5					
E _{pol*}	-98.9	-95.8	-88.6	-85.2	-95.2	-91.6					
E _{ct}	-16.0	-17.6	-14.1	-15.6	-13.4	-15.6					
E _{ct*}	-15.8	-17.6	-13.9	-15.6	-13.0	-15.6					
ΔE	-262.7	-264.6	-248.5	-250.6	-307.5	-312.8					
ΔE*	-262.5	-264.6	-248.3	-250.6	-307.0	-312.8					
Values (ko	Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions										
ΔE^* BSSE corrected total interaction energy											

Table IV.b	. Polyligated Z	n ²⁺ complexes								
	[Zn(lmida	zoles) ₃] ²⁺	[Zn(Imida	zoles) ₄] ²⁺	[Zn(lmida	zoles) ₄] ²⁺	[Zn(Methan	ethiolates) ₃] ⁻		
	d = 2	.11 Å	d= 2	2.01Å	d= 2	2.11Å	d= 2.37 Å			
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA		
Es/E _{MTP*}	-254.9	-254.2	-370.8	-374.2	-332.5	-332.6	-660.2	-642.9		
Ex/E _{rep}	85.0	74.7	167.3	151.4	116.2	102.6	144.9	128.2		
E ₁	-170.0	-179.5	-203.6	-222.8	-216.3	-230.0	-515.3	-514.7		
E _{pol}	-120.8	-123.4	-139.5	-134.7	-130.5	-131.0	-84.6	-78.4		
E _{pol*}	-148.2	-149.8	-181.5	-186.2	-172.2	-176.2	-110.95	-103.8		
E _{ct}	-31.5	-24.7	-38.1	-29.7	-34.0	-25.9	-64.2	-44.8		
E _{ct*}	-30.9	-24.7	-36.9	-29.7	-33.0	-25.9	-30.0	-44.8		
ΔE	-322.3	-327.6	-381.1	-387.2	-380.8	-386.9	-698.3	-637.9		
ΔE*	-321.7	-327.6	-380.0	-387.2	-379.8	-386.9	-664.1	-637.9		
Values (ko	values (kcal/mol) of the QC and SIBFA interaction energies and their contributions									

 ΔE^* BSSE corrected total interaction energy

Table V. Polyligated Zn ²⁺ complexes.										
	[Zn(H	IHCC)]	[Zn(H	HE)]+						
	QC	SIBFA	QC	SIBFA						
Es/E _{MTP*}	-656.6	-640.1	-498.2	-493.1						
Ex/E _{rep}	172.1	149.1	127.7	113.7						
Ex/E _{rep**}	172.1	137.5	127.7	102.1						
E ₁	-484.5	-491.0	-370.5	-379.3						
E _{pol}	-110.2	-88.9	-108.4	-109.8						
E _{pol*}	-130.7	-123.9	-128.8	-136.4						
E _{ct}	-35.7	-38.6	-33.6	-26.2						
E _{ct*}	-34.4	-38.6	-32.9	-26.2						
ΔE	-630.4	-618.4	-512.5	-515.3						
ΔΕ*	-629.1	-618.4	-511.8	-515.3						
ΔE**	-629.1	-630.6	-511.8	-528.6						
Values (kcal/	mol) of the QC a	and SIBFA interactio	n energies and their o	contributions						
ΔE* BSSE co	ΔE^* BSSE corrected total interaction energy									
**Calculated v	vith no smeared	d lone-pairs								

Table VI. PM	∕II-Zn²+ comp	lexes.				
	/	4	E	В	(0
	QC	SIBFA	QC	SIBFA	QC	SIBFA
Es/E _{MTP*}	-770.5	-785.0	-789.3	-818.0	-790.3	-818.1
Ex/E _{rep}	177.6	201.5	219.0	242.9	209.9	239.8
E ₁	-592.2	-583.6	-570.3	-575.1	-580.5	-578.4
Epol(VR)/Epol	-102.9	-99.6	-100.6	-98.7	-110.3	-102.5
E _{ct}	-39.1	-37.3	-51.2	-41.0	-47.0	-40.8
E _{ct} *	-27.3	-37.3	-39.2	-41.0	-36.2	-40.8
ΔE	-734.2	-720.5	-722.1	-714.8	-737.8	-721.6
ΔΕ*	-723.1	-720.5	-710.1	-714.8	-727.1	-721.6
Values (kca	al/mol) of the	QC and SIBFA	interaction ene	ergies and their	contributions	

 ΔE^* BSSE corrected total interaction energy

Table VII.a. Nonaddi	itivity value	s of [Zn(H	HE)]⁺								
	Es/E	MTP*	Ex/I	Erep	_Epol(RV	′S)/Epol*	Epol(V	'R)/Epol		E	ēct
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA		QC	SIBFA
Σ	-498.2	-493.1	124.8	113.7	-211.5	-222.5	-220.9	-233.6	-	-62.5	-56.6
[Zn(HHE)]⁺	-498.2	-493.1	127.7	113.7	-128.8	-136.5	-108.4	-109.8		-33.6	-26.2
δE _{nonadditivity}	0.0	0.0	2.9	3.6	82.7	80.1	112.5	123.8		28.9	30.4
Values (Kcal/mol) of	the QC an	d SIBFA in	teraction energ	gies and tl	heir contributior	าร					

Table VII.b. Nonadditiv	rity value:	s of [Zn(Im	$nidazoles)_4]^{2+}$							
	Es / E	EMTP*	Ex /	Erep	Epol(RVS	S) / Epol*	Epol(V	R) / Epol	 E	ct
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA
Σ	-332.5	-332.6	112.1	101.9	-262.3	-254.3	-270.5	-266.1	-68.9	-64.4
[Zn(Imidazoles) ₄] ²⁺	-332.5	-332.6	116.2	101.9	-172.2	-176.2	-130.5	-131.0	-34.0	-25.9
δE _{nonadditivity}	0.0	0.0	4.1	3.9	90.1	78.2	140.2	135.1	34.9	38.5
Values (Kcal/mol) of th	ne QC an	d SIBFA in	teraction energ	gies and tl	heir contribution	IS				

Table VII.c. Nonadditivity values of [Zn(HHCC)] ⁻

	Es / E	EMTP*	Ex /	Erep		Epol(RV	S) / Epol*		Epol(V	R) / Epol	Ð	ct
	QC	SIBFA	QC	SIBFA		QC	SIBFA	-	QC	SIBFA	QC	SIBFA
Σ	-656.6	-640.1	164.5	149.1		-315.6	-308.8		-325.0	-324.4	-135.7	-125.5
[Zn(HHCC)] ⁻	-656.6	-640.1	172.1	149.1		-130.7	-123.9		-97.9	-88.9	-48.0	-38.6
δE _{nonadditivity}	0.0	0.0	7.6	6.4		184.8	185.0		227.1	235.5	87.7	86.9
Values (Kcal/mol) o	f the QC an	d SIBFA inte	eraction energ	gies and th	neir co	ontributior	ıs					

Table VIII	Table VIII.a. Monoligated Mg²+ complexes: Oxygen ligands.													
	Н	2 O	Meth	nanol	Form	amide	Fomate	external	Format	e bridge	Phe	enol		
	<i>d</i> = 1	.90 Å	d = 1	I.90Å	<i>d</i> = 1	.80 Å	<i>d</i> = 1	.80 Å	<i>d</i> = 1	.94Å	<i>d</i> = 190 Å			
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA		
Es/E _{MTP*}	-68.9	-69.6	-70.1	-70.0	-101.7	-99.9	-302.8	-297.1	-349.2	-342.0	-52.2	-51.4		
Ex/E _{rep}	26.2	25.1	28.3	24.8	38.4	40.4	54.4	52.2	58.7	55.2	27.3	25.8		
E ₁	-42.7	-44.5	-41.8	-45.2	-63.3	-59.5	-248.4	-244.8	-290.6	-286.7	-24.9	-25.6		
E _{pol}	-36.6	-35.6	-46.0	-42.8	-61.2	-64.4	-60.8	-53.5	-71.8	-66.9	-74.6	-71.8		
E _{ct}	-1.4	-2.2	-1.5	-2.1	-2.2	-3.3	-3.6	-4.0	-4.2	-4.9	-1.8	-2.0		
ΔE	∆E -80.7 -82.3 -89.3 -90.1 -126.6 -127.2 -312.8 -302.3 -366.5 -358.6 -101.3 -99.4													
Values (ko	cal/mol)	of the QC	and SIBFA	interact	ion energies	and the	ir contributio	ons						

Table VIII.b. Mo	able VIII.b. Monoligated Mg ²⁺ complexes: Nitrogen and sulfur ligands.												
	Imida	azole	Pyri	idine	Thrimeth	nylamine	Methane	ethiolate					
	d = 2.	.00 Å	<i>d</i> = 1	.90Å	<i>d</i> = 1.	.90 Å	<i>d</i> = 2.20 Å						
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA					
Es/E _{MTP*}	-97.1	-92.9	-92.3	-90.5	-96.7	-97.3	-318.3	-318.3					
Ex/E _{rep}	34.6	31.2	51.5	52.2	60.3	58.6	67.6	69.9					
E ₁	-62.5	-61.7	-40.8	-38.4	-36.4	-38.7	-250.7	-248.4					
E _{pol}	-63.6	-64.1	-75.9	-77.8	-70.3	-72.1	-83.7	-77.1					
E _{ct}	-2.2	-3.0	-2.4	-3.1	-3.9	-2.2	-11.7	-6.6					
ΔE	-128.3 -128.8 -119.1 -119.3 -110.7 -112.9 -346.0 -332.1												
Values (Kcal/mo	ol) of the QC ar	nd SIBFA intera	ction energies a	nd their contribu	utions								

Table IX.	able IX. Monoligated Mg ²⁺ complexes: Formamide.													
	-d 4	co Å		70 Å	- 4	00 Å	4 - 4	00 Å	4-0	00.4				
	<i>a</i> = 1	.60 A	<i>d</i> = 1	.70A	a= 1.	80 A	<i>a</i> = 1	.90 A	a = 2	.00 A				
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA				
Es/E _{MTP*}	-125.7	-122.5	-112.5	-110.4	-101.7	-99.9	-92.7	-90.9	-85.0	-83.2				
Ex/E _{rep}	89.5	98.6	58.6	62.9	38.4	40.4	25.2	26.1	16.6	16.9				
E ₁	-36.2	-23.9	-53.9	-47.4	-63.3	-59.5	-67.4	-64.8	-68.4	-66.2				
E _{pol}	-73.3	-65.6	-66.8	-62.6	-61.2	-58.9	-56.0	-54.8	-51.0	-50.4				
E _{ct}	-2.7	-3.4	-2.5	-3.4	-2.2	-3.3	-1.9	-3.1	-1.8	-2.9				
ΔE	-112.2	-92.9	-123.2	-113.4	-126.6	-121.7	-125.4	-122.7	-121.2	-119.6				
Values (k	/alues (kcal/mol) of the QC and SIBFA interaction energies and their contributions													

Table X.a.	Monoligat	ed Ca²+ com	plexes: Oxy	ygen ligands	s.							
	н	₂ 0	Meth	nanol	Form	amide	Fomate	external	Formate	e bridge	Phe	enol
	d = 2	.30 Å	d= 2	2.30Å	d= 2	.10 Å	d=2	.30 Å	d= 2	.50Å	<i>d</i> = 2	.20 Å
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA
$\mathrm{Es/E}_{\mathrm{MTP}^{\star}}$	-51.0	-50.6	-48.7	-49.9	-84.0	-83.2	-237.0	-230.9	-279.5	-265.7	-41.6	-39.9
Ex/E _{rep}	18.1	19.8	18.9	19.5	40.4	46.4	23.2	26.4	22.6	22.6	31.4	34.1
E ₁	-33.0	-30.8	-29.9	-30.4	-43.6	-36.8	-213.8	-204.5	-256.9	-243.1	-10.2	-5.8
E _{pol}	-17.3	-20.0	-23.3	-25.4	-35.2	-40.7	-30.6	-33.2	-30.5	-37.2	-47.9	-50.3
E _{ct}	-4.6	-3.8	-6.1	-3.7	-11.7	-10.2	-13.3	-8.5	-13.2	-8.0	-8.8	-4.5
E _{ct*}	-3.5	-3.8	-5.5	-3.7	-10.9	-10.2	-11.1	-8.5	-10.7	-8.0	-8.8	-4.5
ΔE	-54.9	-54.6	-59.3	-59.5	-91.3	-87.7	-257.7	-246.2	-300.6	-288.3	-66.9	-60.7
ΔE*	-53.8	-54.6	-58.6	-59.5	-90.5	-87.7	-255.5	-246.2	-298.0	-288.3	-66.9	-60.7

Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions

 ΔE^* BSSE corrected total interaction energy

Table X.b.	Table X.b. Monoligated Ca ²⁺ complexes: Nitrogen and sulfur ligands.										
	Imida	azole	Pyri	idine	Trimeth	nylamine	I	Methan	ethiolate		
	d = 2	.30 Å	d= 2	2.30Å	d= 2	2.50 Å	_	d=2	.50 Å		
	QC SIBFA		QC	SIBFA	QC	SIBFA		QC	SIBFA		
$\text{Es/E}_{\text{MTP}^*}$	-76.4	-72.5	-63.4	-60.9	-47.5	-50.4	-	295.3	-274.8		
Ex/E _{rep}	37.2	34.4	38.1	35.8	20.0	22.4		79.4	76.2		
E ₁	-39.2	-38.2	-25.3	-25.1	-27.5	-28.0	-	215.9	-198.7		
E _{pol}	-38.2	-45.2	-42.3	-48.6	-34.7	-43.9		-44.3	-68.2		
E _{ct}	-10.3	-5.0	-10.8	-5.0	-7.9	-2.0		-25.5	-18.8		
E _{ct*}	-9.9	-5.0	-10.5	-5.0	-7.7	-2.0		-22.5	-18.8		
ΔE	-87.7	-88.4	-78.4	-78.7	-70.1	-73.9	-	285.6	-285.6		
ΔE*	-87.3	-88.4	-78.1	-78.7	-69.9	-73.9	-	282.7	-285.6		

Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions

 ΔE^* BSSE corrected total interaction energy

Table XI.a	Table XI.a. Monoligated Ca ²⁺ complexes: Formate external.												
	d= 2	.10Å	<i>d</i> = 2	.20Å	<i>d</i> = 2	30 Å	d=	2.40 Å	<i>d</i> = 2	.50 Å			
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA			
Es/E _{MTP*}	-264.5	-258.0	-249.6	-243.5	-237.0	-230.9	-226.3	-219.8	-216.9	-209.9			
Ex/E _{rep}	52.2	59.2	35.0	39.5	23.2	26.4	15.3	17.8	10.0	12.0			
E ₁	-212.3	-198.8	-214.6	-204.0	-213.8	-204.5	-211.0	-202.0	-206.9	-197.9			
E _{pol}	-36.9	-39.0	-33.6	-36.1	-32.8	-33.2	-27.9	-30.5	-25.4	-27.9			
E _{ct}	-18.3	-12.5	-15.3	-10.3	-13.3	-8.5	-12.1	-6.9	-11.4	-5.7			
E _{ct*}	-16.0	-12.5	-13.0	-10.3	-11.1	-8.5	-9.9	-6.9	-9.3	-5.7			
ΔE	-265.2	-250.3	-261.2	-250.4	-257.7	-246.2	-248.8	-239.4	-241.5	-231.5			
Values (ko	alues (kcal/mol) of the QC and SIBFA interaction energies and their contributions												
∆E* BSSE	∆E* BSSE corrected total interaction energy												

Table XI.b. Monoligated Ca ²⁺ complexes: Formate bridge.											
	<i>d</i> = 2.30Å		<i>d</i> = 2.40Å		<i>d</i> = 2	<i>d</i> = 2.50 Å		<i>d</i> = 2.60 Å		<i>d</i> = 2.70 Å	
	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	QC	SIBFA	
Es/E _{MTP*}	-309.1	-293.3	-293.3	-278.8	-279.5	-265.7	-268.7	-256.3	-257.3	-243.9	
Ex/E _{rep}	50.5	50.7	34.0	33.9	22.6	22.6	15.7	16.5	10.1	10.4	
E ₁	-258.6	-242.5	-259.3	-244.9	-256.9	-243.1	-253.0	-239.8	-247.2	-233.5	
E_{pol}	-37.3	-43.7	-33.7	-40.4	-27.9	-37.2	-27.9	-34.6	-25.1	-31.2	
E _{ct}	-16.1	-11.9	-14.3	-9.8	-13.2	-8.0	-12.7	-6.8	-12.4	-5.4	
E _{ct*}	-13.2	-11.9	-11.6	-9.8	-10.7	-8.0	-10.3	-6.8	-10.2	-5.4	
ΔE*	-309.1	-298.2	-304.6	-295.2	-298.0	-288.3	-291.2	-281.2	-282.5	-270.1	
Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions											
ΔE^* BSSE corrected total interaction energy											

Table XII.a. Polyligated Mg ²⁺ complexes.									
		[Mg(H	[Mg(H2O)6] ²⁺						
	Pyramidal,	<i>d</i> = 2.10 Å	Planar, d	= 2.20 Å	<i>d</i> = 2.30 Å				
	QC	SIBFA	QC	SIBFA	QC	SIBFA			
Es/E _{MTP*}	-209.0	-210.0	-195.0	-195.4	-257.9	-258.1			
Ex/E _{rep}	42.2	38.9	32.8	29.4	35.0	31.0			
E ₁	-166.8	-171.1	-162.2	-166.0	-222.9	-227.1			
E _{pol}	-81.2	-76.5	-72.7	-68.6	-73.6	-69.1			
E _{pol*}	-93.0	-97.8	-83.4	-78.6	-90.7	-85.1			
E _{ct}	-3.6	-3.0	-3.6	-3.2	-4.2	-3.4			
E _{ct*}	-3.5	-3.0	-3.4	-3.2	-3.9	-3.4			
ΔE	-251.8	-250.7	-238.7	-237.9	-301.0	-299.6			
ΔE*	-251.6	-250.7	-238.5	-237.9	-300.7	-299.6			
Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions									

 ΔE^* BSSE corrected total interaction energy

Table XII.b. Polyligated Mg ²⁺ complexes.									
	[Mg(Imida	zoles)3] ²⁺	[Mg(Imida	$zoles)_4]^{2+}$					
	d = 2	.11 Å	d= 2	11Å	[2Mg(DDE)] ⁺				
	QC	SIBFA	QC	SIBFA	QC	SIBFA			
Es/E _{MTP*}	-244.7	-241.5	-318.9	-315.7	-849.4	-847.2			
Ex/E _{rep}	73.5	65.2	99.8	89.9	119.0	110.2			
E ₁	-171.2	-176.3	-219.0	-225.8	-730.3	-736.9			
E _{pol}	-117.8	-112.9	-127.9	-121.2	-159.9	-137.2			
E _{pol*}	-140.6	-138.7	-163.5	-162.8	-175.6	-137.2			
E _{ct}	-5.1	-3.0	-6.6	-2.5	-13.6	-9.4			
E _{ct*}	-4.4	-3.0	-5.5	-2.5	-13.1	-9.4			
ΔE	-293.4	-292.2	-352.5	-349.5	-903.3	-883.6			
ΔE*	-292.8	-292.2	-351.4	-349.5	-902.8	-883.6			
Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions									
ΔE^* BSSE corrected total interaction energy									

Table XIII. Polyligated Ca ²⁺ complexes.											
	[Ca(H2O) ₆₊₂] ²⁺		[Ca(H2	[Ca(H2O) ₇₊₁] ²⁺		[Ca(H2O) ₈] ²⁺			[Ca(DDDTE)] ²⁻		
	QC	SIBFA	QC	SIBFA		QC	SIBFA	-	QC	SIBFA	
Es/E _{MTP*}	-262.4	-263.2	-278.1	-282.3		-275.0	-281.6		-558.6	-566.9	
Ex/E _{rep}	103.0	118.2	95.1	109.1		94.6	109.1		87.0	113.5	
E ₁	-159.3	-145.0	-183.0	-173.2		-180.4	-172.5		-471.6	-453.4	
E _{pol}	-59.6	-69.9	-53.1	-58.5		-50.1	-54.4		-35.4	-21.0	
E _{pol*}	-62.3	-84.7	-56.0	-75.4		-53.8	-72.9		-29.8	-24.6	
E _{ct}	-20.0	-14.4	-15.4	-13.2		-14.3	-12.5		-15.3	-19.2	
E _{ct*}	-11.0	-14.4	-6.2	-13.2		-5.1	-12.5		-3.2	-19.2	
ΔE	-238.9	-229.3	-251.4	-244.9		-244.8	-239.4		-522.3	-493.5	
ΔE*	-230.0	-229.3	-242.2	-244.9		-235.6	-239.4		-510.2	-493.5	
Values (kcal/mol) of the QC and SIBFA interaction energies and their contributions											
ΔE^* BSSE corrected total interaction energy											