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Charge-displacement analysis as a tool to study chalcogen bonded adducts and predict their association constants in solution†

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The secondary interaction between a polarized chalcogen atom and different Lewis bases, either anionic or neutral, has been studied by charge displacement analysis. Using charge displacement analysis, the charge rearrangement in the adduct upon the formation of the interaction has been quantified and described in great detail. By comparing the theoretical results with the experimental association constants, two linear correlations can be found for anionic and neutral bases. Such correlations can be used to reliably predict the association constants of adducts for which experimental data are not available yet.

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Introduction

Secondary bonding interactions (SBIs)¹ play a crucial role in chemistry and biochemistry. During the last decades special attention has been paid to the intramolecular and intermolecular contacts between di- and tetracoordinate sulfur, selenium, and tellurium atoms and oxygen, nitrogen, and sulfur atoms² and even π systems.³ Any time these interatomic contacts are longer than a covalent single bond but shorter than the sum of the corresponding van der Waals radii, the presence of a secondary bond is supposed to exist (the interaction is often referred to as a chalcogen bond,⁴ ChB), often determining a peculiar conformation of the molecules and influencing their reactivity.⁵ The presence of this interaction has been demonstrated by solid state experimental data^{2,6} and confirmed in solution by spectroscopic evidence.^{7,8}

SBIs are fundamental in determining the tridimensional structure of inorganic materials and macromolecules, like the secondary and tertiary structures of proteins and biopolymers. Besides the most conventional hydrogen bonding, the investigation of some biopolymers and organic conductors evidenced the role of less conventional interactions, such as those between chalcogen centers, demonstrating their ability in promoting conformational stabilization. ¹⁰

Furthermore, close contacts between nucleophilic sites and electron-deficient covalent-bonded chalcogens have been supposed to be responsible for the enantioselectivity shown by some chalcogen-containing electrophilic reagents in a number of addition reactions to carbon–carbon double bonds, both in inter- and intramolecular processes. It was unequivocally demonstrated that the presence of an intramolecular S–Se interaction is mandatory to achieve a high facial selectivity in methoxyselenenylation reactions as well as in cyclofunctionalization reactions.⁸

SBIs are also responsible for the activation of the catalytic site of glutathione peroxidase, a crucial selenoenzyme responsible for destroying lipid-damaging peroxides in mammals, protecting biomembranes from oxidative stress and, consequently, from a number of pathologies. Recent studies demonstrated that intramolecular Se···O or Se···N interactions play an important role in the catalytic antioxidant activity of these compounds. As a direct consequence of SBIs, the electrophilicity of the chalcogen atom in an organic scaffold is modified, with effects in a series of other biological processes.

All of these examples make the deep understanding of the ChB of fundamental importance and, in recent years, intense research activity was aimed at this point. From the computational point of view, the secondary interactions of chalcogenadiazoles have already been studied with different methods, such as the Quantum Theory of Atoms in Molecules (QTAIM), Natural Bond Orbital (NBO) and Voronoi Deformation Density (VDD).

Recently, Taylor and co-workers synthesized a series of benzotelluradiazoles, with the precise aim to study the ChB between tellurium and Lewis bases in solution. The association constants (K_a) were determined by UV-Vis and H NMR titrations in different organic solvents, varying the substituents

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R₁
$$R_4$$
 R_4 R_4 R_4 R_5 R_5

Scheme 1 Formation of chalcogen-bonded adducts and numbering of the species.

on the aromatic ring to tune the electronic properties of tellurium (compounds 2–5,¹⁷ Scheme 1), and different bases, either anionic (chloride, bromide, iodine and nitrate anions) or neutral (quinuclidine, Q, Scheme 1).

The results demonstrated that the substituents R_{1-4} can finely tune the value of K_a , which can reach considerable values in organic solvents (130 000 M⁻¹ for 5Cl in THF). Substituting the phenyl moiety with two electrowithdrawing cyanide groups (compounds **6a** and **6b**, Scheme 1) and using thiophenolate as the Lewis base, the group of Beckmann, Gritsan and Zibarev obtained even higher values, reaching 10^8 in THF. 15a

In this paper, we study the same adducts using Charge Displacement (CD) analysis, ¹⁸ which has already demonstrated its potential for characterization of secondary interactions (frustrated Lewis pairs, ¹⁹ hydrogen ²⁰ and halogen bonding ²¹) and coordination bonds. ²² Such an approach relies on the integration along a given direction z (eqn (1)) of the difference in electronic density $[\Delta \rho(x,y,z)]$ between the adduct and its non-interacting fragments (in our case the benzotelluradiazole and the Lewis base), frozen in the same positions they occupy in the adduct.

$$\Delta q(z') = \int_{-\infty}^{\infty} \mathrm{d}x \int_{-\infty}^{\infty} \mathrm{d}y \int_{-\infty}^{z'} \mathrm{d}z \Delta \rho(x, y, z) \tag{1}$$

The value of $\Delta q(z')$ defines the amount of the electronic charge that, upon formation of the adduct, has moved across a plane perpendicular to the axis through the coordinate z'. At the boundary between the fragments (see Computational details for the choice of the boundary) the value of Δq is represented by the symbol ω .

To the best of our knowledge, CD analysis has never been used for the characterization of ChB, but its density-based

nature, its stability toward the Computational details (in particular the choice of the basis set size and XC functional, see Computational details) and its propensity to study linear adducts makes it a potentially useful tool.

The results presented here give a detailed and quantitative picture of the ChB between benzotelluradiazoles and Lewis bases, and also provide a reliable methodology to predict the value of the free energy of interaction of systems for which experimental data are not available yet.

Results and discussion

In the optimized structure of 3a, the molecule shows a C_{2v} symmetry (two mirror planes and one C_2 axis) and the two N–Te distances are both 1.994 Å. The formation of the adduct with the chloride (3aCl) lowers the symmetry to C_s (only one mirror plane in the whole molecule) because the chloride approaches the tellurium on the opposite side to one of the nitrogen atoms (hereafter called N¹), where the σ -hole²³ is located, ¹6 forming a N¹–Te–Cl angle of 172.9°. In the adduct 3aCl, the N¹–Te distance is considerably longer than in 3a (2.066 Å), while the N²–Te bond, roughly perpendicular to the Te···Cl axis, is less affected (2.022 Å) by the interaction with the base.

Visualizing the 3D plot of $\Delta \rho$, it is possible to describe in detail the electronic re-organization within the adduct upon the formation of the chalcogen bond (Fig. 1).

On the chloride a large depletion of electron density, especially located at the opposite side with respect to the tellurium, and a small ring-shaped accumulation are visible. Between the chloride and the tellurium there is a small electron accumulation, while two large lobes are present on the tellurium. The lobe facing the chloride indicates a marked electron depletion, the other one an electron accumulation.

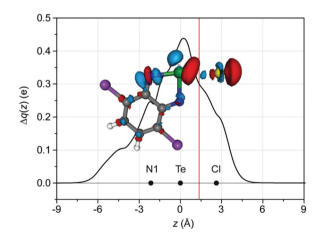


Fig. 1 Charge displacement function for 3aCl. The black dots represent the z coordinate of the atoms. The red vertical line identifies a suitable boundary between the two fragments. Overprinted: 3D contour plot of the change of electronic density upon formation of the adduct 3aCl. Blue (red) isosurfaces identify regions in which the electron density increases (decreases). Density value at the isosurfaces: ± 0.003 au.

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Furthermore, N¹ also shows a remarkable electron accumulation, with a ring-shaped electron depletion around the nucleus. The electron accumulation on N2 is similar but considerably smaller than that on N¹.

The accumulation/depletion pattern described above depicts a net charge transfer from the chloride to the benzotelluradiazole, as denoted by the electronic accumulation visible in the inter-fragment region. The tellurium atom results in being strongly polarized (as indicated by the two opposite lobes), with the electrons that are repelled by the charge of the anion accumulating on the other side. But the tellurium is not the ultimate electron density recipient, as evidenced from the accumulation regions on the two nitrogen atoms (on N¹ more than on N²). Since the N-Te bonds can be described as dative bonds between an imide anion (=N-) and a dicationic tellurium atom (Te2+), the electron density accumulation region on the nitrogen weakens the bond, which is actually longer than in the case of the isolated molecule.

In order to quantify the electronic flux, $\Delta \rho$ has been integrated along the axis connecting Te and Cl (z) giving the CD function depicted in Fig. 1. The CD function is positive throughout the whole molecule, indicating a net charge transfer from the right (chloride) to the left (tellurium) side of the adduct. It can be noted that the flux peaks on the tellurium, but it does not stop on the tellurium, continuing to be positive until N^1 , where it decays to zero. The value of Δq at the interfragment boundary (ω) is 0.3e. As the 3D plot of $\Delta \rho$ revealed, charge transfer and tellurium polarization coexist in the interfragment region, therefore the ω value sums up both contributions. At this stage, the disentanglement between the two is not possible (see later).²³

Fig. 2 shows the same calculations for a neutral adduct, 3aQ. As can be seen, the 3D plot is qualitatively similar to that

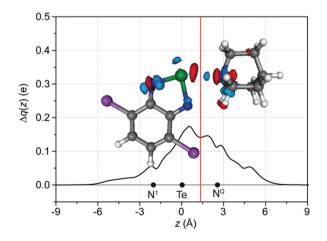


Fig. 2 Charge displacement function for 3aQ. The black dots represent the z coordinate of the atoms. The red vertical line identifies a suitable boundary between the two fragments. Overprinted: 3D contour plot of the change of electronic density upon formation of the adduct 3aQ. Blue (red) isosurfaces identify regions in which the electron density increases (decreases). Density value at the isosurfaces: ±0.003 au.

for 3aCl (depletion on NQ, accumulation between NQ and Te, polarization on Te and accumulation on N1), with the only difference that the polarization on Te is much smaller with respect to the previous case. This can be explained by the charge of the chloride that repels the electrons of the tellurium to a larger degree than a neutral base.

The corresponding CD function (Fig. 2) is again similar in shape to that for 3aCl, but weaker in intensity ($\omega = 0.141e$). This could be due to the smaller donation properties of Q and/ or the smaller polarization induced by Q with respect to Cl-. Incidentally, the same difference can be found by analyzing the CD curves relative to the halogen bonded adducts $C_6F_5I\cdots Cl^-$ and $C_6F_5I\cdots Q$ (Fig. S8†), for which similar values of ω (0.305 and 0.126e, respectively), were obtained.

It is also interesting to note that in the geometry of 3aQ one of the bromine atoms bound to the aromatic moiety is located in the inter-fragment region (z coordinate = 0.966 Å). 3aQ is the only adduct in which an atom of the benzotelluradiazole falls in the inter-fragment region and this is because the N¹-Te-N^Q angle in this case is sensibly lower than 180° (162.6°). The consequence can be appreciated in Fig. 3, where $\Delta \rho$ is plotted at a lower density value than in Fig. 2 (±0.0005 instead of ± 0.003 au).

It is evident that the bromine in the inter-fragment region undergoes a small electronic re-organization, with the electron flowing from the carbon (electron density depletion) to the bromine (accumulation), therefore in the opposite direction with respect to the flux between the chalcogen and the base. Since ω is the electronic charge that, upon formation of the adduct, has moved across a plane perpendicular to the axis, in the case of 3aQ ω contains contributions from both the Te \leftarrow N and $C \rightarrow Br$ fluxes.

In all of the other adducts the N¹-Te-X angle is closer to 180° and none of the atoms of the benzotelluradiazoles fall in the inter-fragment region (all of the atoms of the other benzotelluradiazoles show a z coordinate smaller than 0.6 Å).

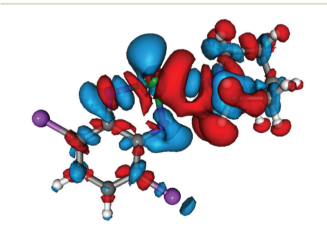


Fig. 3 3D contour plot of the change of electronic density upon formation of the adduct 3aQ. Blue (red) isosurfaces identify regions in which the electron density increases (decreases). Density value at the isosurfaces: ±0.0005 au.

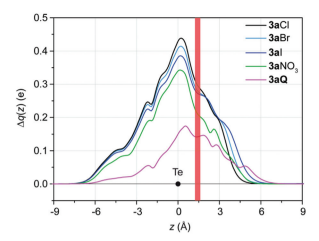


Fig. 4 Charge displacement functions for the 3aX adducts. The black dot represents the z coordinate of the tellurium, always placed at the origin. The red vertical band identifies the range of the inter-fragment boundaries.

Consequently, 3aQ is the only one in which the inter-fragment region contains more than one flux.

Expanding the methodology to other Lewis bases, the CD function has been computed for all of the 3aX adducts experimentally studied by Taylor¹⁶ (Fig. 4). All of them have the same shape, indicating that the nature of the interaction is similar for all of the bases, but there is a trend in the intensities of the functions, which depend on the bond acceptor properties of the base and the magnitude of the polarization.

The ω values are 0.300, 0.277, 0.269, 0.225 and 0.141e for 3aCl, 3aBr, 3aI, 3aNO₃ and 3aQ, respectively. The big gap between the ω value of 3aQ and the others seems to be an indication of the poor bond acceptor properties of Q, but nitrogensubstituted adamantane species (such as Q or 1,4-diazabicyclo-[2.2.2]octane) showed excellent bond acceptor properties when involved in halogen-bonded adducts.²⁴ More likely, most of the difference is due to the smaller contribution of the polarization (compare Fig. 1 and 2).

In order to take into account the difference between tellurium and selenium adducts, the adducts formed by the structurally similar 3,4-dicyano-1,2,5-chalcogenadiazole and the anion SPh⁻ (Scheme 1), as experimentally and theoretically characterized by Zibarev and co-workers, ^{15a} can be compared. The CD curves for the adducts **6a**SPh and **6b**SPh are shown in Fig. 5.

Also, in this case the shape of the curve is similar to that for 3aCl (Fig. 1). The values of ω are 0.376 and 0.330e for 6aSPh and 6bSPh, respectively. As expected, the selenium can accept less electronic density with respect to the tellurium, as already stated in the original paper using different methods. ^{15a} To the best of our knowledge, experimental data are not available for similar systems containing sulfur.

Calculating the value of ω for the adducts for which the K_a has been experimentally evaluated, ²⁵ allows the comparison of ω and the logarithm of the association constants, which is related to the free energy of association (Fig. 6). In particular,

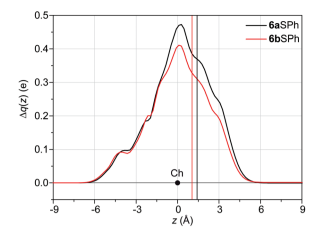


Fig. 5 Charge displacement functions for the 6aSPh and 6bSPh adducts, containing tellurium and selenium, respectively. The black dot represents the *z* coordinate of the chalcogen atom, always placed at the origin, while the anion is always placed on the +*z* semiaxis. The black and red vertical lines identify the inter-fragment boundaries for 6aSPh and 6bSPh, respectively.

all of the K_a values reported here have been measured in THF by Taylor and co-workers (adducts 2–5X)¹⁶ or Zibarev and co-workers (6a–6bSPh).^{15a}

The plot of ω vs. $\ln K_{\rm a}$ reveals that if ionic and neutral adducts are treated separately, the two quantities are linearly correlated, with good correlation coefficients ($r^2=0.9447$ and 0.9999 for ionic and neutral adducts, respectively) and similar slopes (m=0.011 and 0.017 for ionic and neutral adducts, respectively). 3aQ, for which, as shown before, the ω values are lowered by the C \rightarrow Br electronic flux, has been excluded from the fitting procedure.

An alternative way to estimate the charge transfer is the Hirshfeld charge analysis. ²⁶ This allows estimation of the charge on each arbitrarily defined fragment of an adduct, computed as the integral of the adduct electron density over space, weighted at each point by the ratio between the isolated fragment density and the total density of the non-interacting fragments. Using the same fragmentation scheme as that used for the CD method, the values for the charge transferred from $\bf Q$ to 2–5 ($q_{\rm H}$) correlate perfectly with $\ln K_{\rm a}$ (Fig. 6, $r^2=0.9916$), even when including $\bf 3aQ$ in the fitting procedure. For anionic bases, the fitting is slightly poorer ($r^2=0.9177$) but still acceptable. In general, $q_{\rm H}$ shows very good agreement with the values of ω (Table S1†).

It is interesting to note that 2Q and $2NO_3$ have the same value of K_a but different values of ω . In addition to this, the polarization contribution in 2Q is smaller than in $2NO_3$ (see Fig. S5 and S6†). Therefore, combining experimental and theoretical data, it seems that the main contribution in determining the K_a value comes from the charge transfer rather than from the polarization. Stretching further the hypothesis, it can be assumed that, in first approximation, only the effective charge transfer (CT_{eff}) is important for the value of K_a . Under this hypothesis, we can consider the values of the

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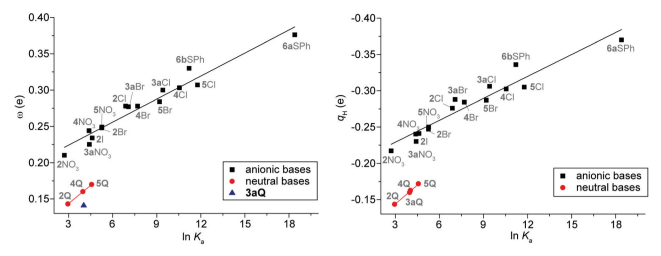


Fig. 6 (Left) Linear correlation between ω and the logarithm of the experimental values of K_a . For the exclusion of 3aQ from the fitting procedure, see the text. (Right) Linear correlation between q_H and the logarithm of the experimental values of K_a .

intercepts (ω_0) found in the fitting procedure previously described (Fig. 6, ω_0 = 0.192 and 0.094e for ionic and neutral adducts, respectively) as the polarization contributions for neutral and ionic adducts. Now, $CT_{\rm eff}$ can be roughly estimated by subtracting the intercept values from ω , which varies from 0.02 (2NO₃) to 0.18 (6aSPh) e, giving the order of magnitude of the "covalent" character of the ChB for these systems.

As shown in Fig. 1 and 2, the N¹-Te bond is also affected by the Te···X interaction. Actually, by plotting the lengthening of the N¹-Te bond with respect to the isolated benzotelluradiazole *versus* ω , a linear correlation can be appreciated (Fig. S7†). In this case there is no differentiation between ionic and neutral adducts, which is likely because N¹, being far from the Lewis base, is unaffected by the charge of X and because the polarization on Te does not involve the N¹-Te axis (Fig. 1).

The goodness of the fitting in Fig. 6 suggests that charge displacement analysis can be used as a reliable method to predict the free interaction energy involved in chalcogen bonding for the benzotelluradiazole class of compounds. In fact, the prediction of the association free energy (ΔG^0) computed by B97-D3 calculations (the best XC functional according to the benchmark conducted in ref. 16 and 15a; the solvent effect was taken into account through the PCM model for THF), even if generally acceptable, fails in some cases (see the case of **2Br** or **3a**NO₃ in Fig. 7), with a deviation standard of 1.2 kcal mol⁻¹ and a maximum error of 3.3 kcal mol⁻¹. Computing ΔG^0 from the ω value using the fitting equations shown in Fig. 6 gives more reliable values (standard deviation = 0.50 kcal mol⁻¹, maximum error = 1.0 kcal mol⁻¹).²⁷ Such reliability can be used for the prediction of ΔG^0 of a chalcogen bonding donor/acceptor couple for which experimental data are missing.

These results open the way to a more reliable theoretical pre-screening for choosing the optimal donor/acceptor couple. In fact, even if no experimental data were available, the trend of the ΔG^0 values of the adducts would be correctly predicted.

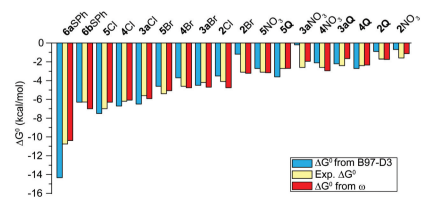


Fig. 7 Free energies of chalcogen bonding interactions of 2X-6X. Experimental and B97-D3-computed values are taken from ref. 16 (2X-5X) and 15a (6aSPh and 6bSPh), ω-computed data are calculated using the proper fitting equation shown in Fig. 6.

Conclusions

In this paper, it has been shown that coupling the visual inspection of $\Delta \rho(x,y,z)$ and charge displacement analysis is very useful for describing the interaction between a chalcogen atom and a Lewis base, from both a qualitative and quantitative point of view. Taking into account a series of adducts between substituted benzochalcogenadiazoles and a series of Lewis bases, either charged or neutral, the corresponding CD functions always have the same general shape, indicating that the nature of the Ch...X interaction does not depend on the exact nature of the base. The only differences rely on the donor properties of the chalcogen, the acceptor properties of the base and, consequently, the exact value of ω . Moreover, the values of the latter were markedly different for neutral and charged bases, since ω also contains a polarization contribution, which is larger for anionic bases. Assuming that the main driving force for the establishment of the ChB is the charge transfer rather than the polarization, the "covalent" character of the interaction can be roughly estimated between 0.02 and 0.18 electrons.

 ω values and experimentally-measured ΔG values were found to be linearly correlated, confirming once again the physical likeliness of the CD analysis in describing the formation of adducts bound by weak interactions. Furthermore, once calibrated with experimental data, ω is more reliable than standard DFT to predict the ΔG of a benzochalcogenadiazoles/ base couple for which experimental data are not available. In general, even if the prediction of the numeric value of ΔG is only possible if a certain number of experimental values are available, the trend of the values can be efficiently predicted even without any experimental values, for instance for new classes of ChB donors, opening the way to a reliable theoretical pre-screening for choosing the optimal donor/acceptor couple. When the inter-fragment region contains atoms other than those involved in the SBI, as in the case of 3aQ, the analysis of the Hirshfeld charges can satisfactorily replace the CD method, even without the details that the analysis of the CD functions throughout the whole molecule allows.

Finally, this study demonstrates how the synergistic coupling of theoretical and experimental data can be fruitfully applied to chalcogen bonding, allowing studies in the near future to detail the nature and the effects of weak interactions in chemically and biologically relevant systems.

Computational details

All thermodynamic calculations were performed using the Gaussian 09 program. ²⁸ All of the optimizations ($C_6F_5I\cdots Cl^-$ and $C_6F_5I\cdots Q$) were carried out using density functional theory employing the B97D3²⁹ functional, the def2-TZVP basis set and the ECP pseudopotential for iodine, ³⁰ as stored in the Basis Set Exchange portal. ³¹ All of the optimized adducts show only positive vibrations. The geometries for 2–5X, and **6a**SPh and **6b**SPh have been taken from ref. 16 and 15a, respectively,

and used without further reoptimization. When necessary, the geometries have been re-oriented to have the z axis passing between the chalcogen atom (origin) and the Lewis base (positive values of z).

All of the charge displacement calculations were carried out using density functional theory employing the BP86³² corrected for the dispersion, using the Grimme D3-parametrized XC functionals,³³ the def2-TZVP basis set and the ECP pseudopotential for tellurium and iodine. The CD method has already been shown to be remarkably stable with respect to the exchange–correlation functional employed, the basis set and the level of theory to account for the relativistic effects (scalar or full four-component hamiltonian).^{22b,c,34} Here, the CD function has been calculated for 3aCl with different XC functionals, giving very similar curves in all of the cases (Fig. S1†).

In order to characterize the interaction with only one value of Δq , it is useful to fix a plausible boundary separating the fragments in the adducts. We chose the isodensity value representing the point on the z axis at which equal-valued isodensity surfaces of the isolated fragments are at a tangent. At this point the value of Δq is represented by the symbol ω .²²

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