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J. Klein, H. Khartabil, J.C. Boisson, J. Contreras-Garcia, Jean-Philip Piquemal, et al.. A New Way for Probing Bond Strength. *Journal of Physical Chemistry A*, 2020, 124 (9), pp.1850-1860. 10.1021/acs.jpca.9b09845 . hal-02377737

HAL Id: hal-02377737

<https://hal.science/hal-02377737>

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A New Way for Probing Bond Strength

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Abstract

The covalent chemical bond is intimately linked to electron sharing between atoms. The recent Independent Gradient Model (IGM) and its δg descriptor provide a way to quantify locally this electron density interpenetration from wavefunction calculations. Each bond has its own IGM- δg^{pair} signature. The present work establishes for the first time a strong link between this bond signature and the physically grounded bond force constant concept. Analyzing a large set of compounds and bonds, the Intrinsic Bond Strength Index (IBSI) emerges from the IGM formulation. Our study shows that the IBSI does not belong to the class of conventional bond orders (like Mulliken, Wiberg, Mayer, delocalization index or ELF), but is rather a new index complementary thereto, related to the bond force. A fundamental outcome of this research is a novel index allowing to range all two-centre chemical bonds by their intrinsic strength in molecular situation. We believe that the IBSI is a powerful and robust tool for interpretation accessible to a wide community of chemists (organic, inorganic chemistry, including transition metal complexes and reaction mechanisms).

Introduction

The chemical bond¹ is a model commonly used by chemists to describe molecules as made of atoms linked to one another. Although lacking a clear physical basis (the bond is not an observable quantity) it is a quite useful concept frequently employed to help chemists to give explanations and make predictions about physical and chemical properties of substances. In theoretical chemistry, the atom connectivity concept is explicitly involved in models such as force fields used to perform molecular dynamic simulations. In chemical education, the chemical bond, very useful to represent molecular structures and to illustrate chemical processes, has acquired a central place and remains fundamental.² The physical interactions that establish cohesion in molecules can be tackled by the use of quantum chemistry.¹ The use of the electron density (ED) topological analysis for the characterization of bonds and

more generally of interactions in molecular systems has led to a variety of investigative methods based on local descriptors. Examples of such approaches are: the electron localization function ELF,^{3,4} the localized electron detector LED,⁵ the Density Overlap Region Indicator DORI⁶ or the Non-Covalent-Interaction approach NCI.⁷ As complementary tools of the Atoms In Molecules approach (QTAIM^{8,9}), these methods allow for producing insightful three-dimensional isosurfaces localizing the interactions within the system from theoretically derived electron density. Recently, in the continuation of the NCI⁷ analysis originally proposed in 2010, we introduced a new local approach for probing strong and weak interactions in molecules. Based on the Independent Gradient Model (IGM^{10,11}) that represents a non-interacting reference system, the δg descriptor has been defined. It measures locally the electron sharing between interacting fragments and reveals regions of space where the ED between fragments clashes. Compared to the original formulation of NCI, which has a semi-quantitative value, the IGM- δg approach is able to quantify the interaction caused by electron sharing. It provides us with a molecular δg signature as illustrated in Figure 1 (top right panel), by plotting δg versus the ED oriented with the sign of the second density hessian eigenvalue λ_2 . It covers a large range of interactions, associated or not with bond critical point (BCP), including covalent bonds, bonding in transition metal compounds, hydrogen-bonding and van der Waals interactions. It is not dimensionless and good correlations have been found between peak heights in the IGM- δg signature of intermolecular complexes and the calculated stabilization energy for different hydrogen bonds.¹⁰ A critical and appealing feature of the IGM- δg computational scheme is its ability to separate and extract the signature of selected atom pairs from the overall interaction signal.¹¹ Thereby, a bond-by-bond IGM- δg^{pair} picture can be obtained from a wavefunction as illustrated in Figure 1. Each chemical bond has its signature. To the best of our knowledge, this possibility is not available within any other descriptor-based approaches.

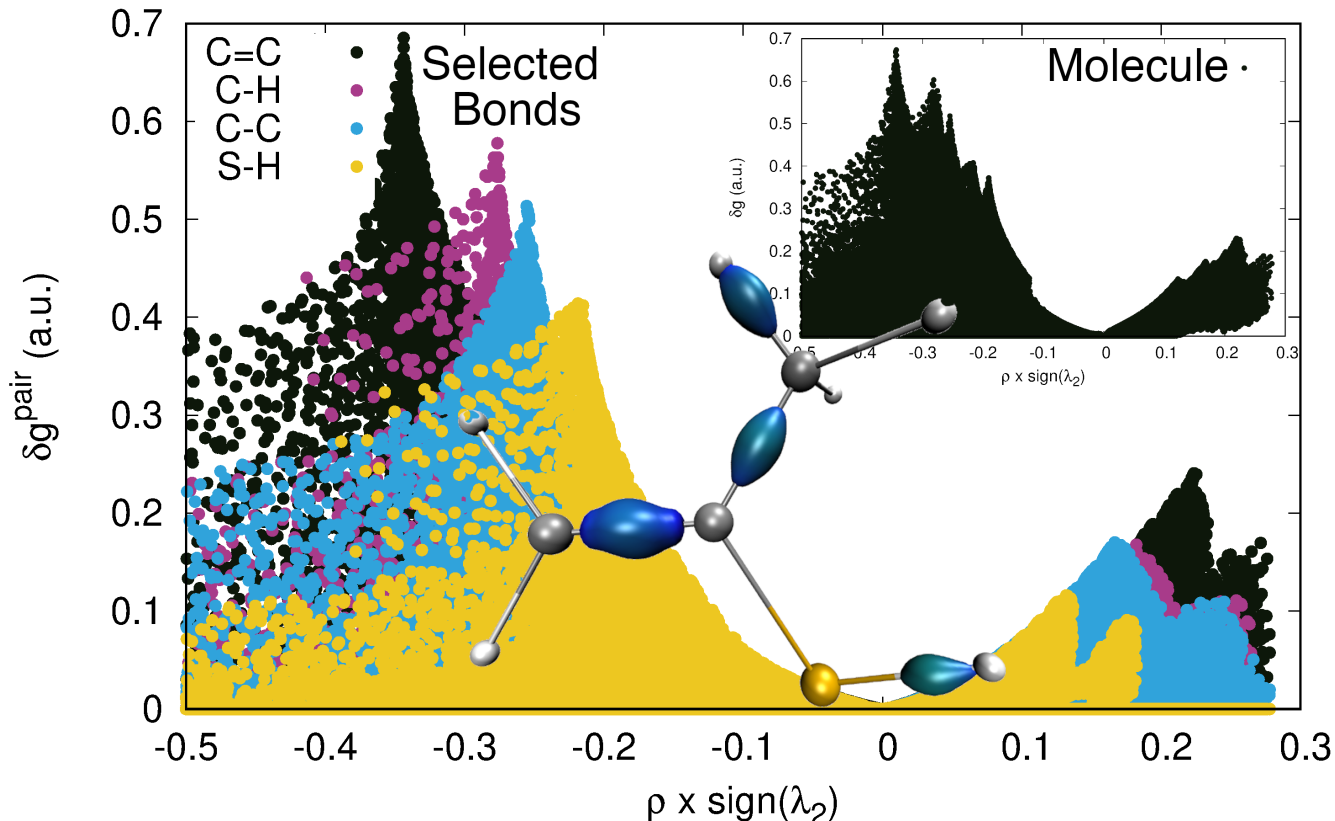


Figure 1: Molecular IGM- δg signature, selected IGM- δg^{pair} bond signatures in prop-1-ene-2-thiol and $\delta g = 0.3$ a.u. isosurfaces colored according to the BGR scheme over the range $-0.2 < \text{sign}(\lambda_2)\rho < 0.2$ a.u. (the sign of the second density hessian eigenvalue λ_2 serves to distinguish between attractive and repulsive situations).

At this stage, it might be tempting to directly associate the strength of the bond with the peak height of the IGM- δg^{pair} bond profile. In the example shown in Figure 1 (prop-1-ene-2-thiol), the $\text{C} = \text{C}$ double bond results in a larger δg^{pair} peak height (0.68) than a single bond $\text{C} - \text{C}$ (0.51). Further, the δg^{pair} peaks for $\text{CH}_2 - \text{H}$ (0.57) and $\text{S} - \text{H}$ (0.41) fairly reflect their homolytic bond dissociation enthalpy ordering (given by Y-R. Luo¹² around 420 and 350-370 $\text{kJ}\cdot\text{mol}^{-1}$, respectively, at 298K). This apparent interrelationship between bond strength and δg^{pair} peak height makes sense since bonds are achieved by electron sharing¹ measured to some extent by δg^{pair} at a point in space. However, each δg^{pair} bond signature has its own individual profile, which suggests that attempts to correlate the δg^{pair} descriptor to energetics should not be limited to the use of a local information (at BCP for instance)

but should consider an integration scheme. Such integration schemes have been explored in the past few years. In their study of the integration of charges within the NCI region,¹³ J. Contreras-García and W. Yang recovered the position of the intermolecular potential minimum for 4 hydrogen-bond complexes. However, although mimicking relatively well the shape of the binding energy curve for the water dimer, the correct curvature around the minimum was not obtained for the other dimers. In a study conducted for a series of 30 weakly bound complexes, Presti¹⁴ found a good correlation between the stabilization energy and the kinetic energy density integrated over the volume enclosed by the NCI isosurface. Unfortunately, this correlation breaks down for complexes of different nature. Moreover, in this previous work based upon the NCI approach, the integration domain was defined in an arbitrary manner (domain below $\text{RDG} = 0.5 \text{ a.u.}$) and ignored some grid points in the interaction region. More recently, for a set of 53 systems, Ananyev¹⁵ presented a strong correlation between effective force constants of diatomic molecules and the potential energy density integrated over the AIM surface between two Bader’s atomic basins. [Very recently, using an integration scheme, the IGM- \$\delta g\$ approach has been employed to derive a score indicating the role played by each atom in the formation of host-guest assemblies.](#)¹⁶ The present study falls in the class of integration schemes that have emerged in the past few years.

The basic aim of this work is to integrate the $\text{IGM-}\delta g^{\text{pair}}$ bond signature to characterize a bond by a unique score. The result is a global bond descriptor Δg^{pair} strongly connected to the physically grounded bond strength concept. In this study, we present results obtained using a set of 235 molecules spanning a broad range of different chemical bonds. Potential relationships between Δg^{pair} and several bond orders as well as with bond stretching force constants have been sought for. A novel Intrinsic Bond Strength Index (IBSI) stems from these attempts. Since the IGM- δg approach is a relatively new concept, we will start by reintroducing it in the Methods section. Also, details on the integration scheme and the design of the new index IBSI will be sketched out. This will be followed by computational

details, results and discussion before to conclude.

Methods

IGM- δg approach

The IGM- δg approach has been designed to reveal and quantify molecular interaction from the ED topology, more precisely from its gradient $\nabla\rho$. All details can be found in our previous works.^{10,11} Its implementation involves the use of a three-dimensional grid built to encompass the chemical system. A prerequisite to apply this local approach is to express each molecular ED gradient component $\frac{\partial\rho}{\partial x}$ as a sum of individual terms, one for each interacting component (atom, fragment, molecule), for instance for two fragments A and B : $\frac{\partial\rho}{\partial x} = \left| \frac{\partial\rho_A}{\partial x} + \frac{\partial\rho_B}{\partial x} \right|$. Then, the idea behind the Independent Gradient Model is to define a non-interacting reference system that cancels the effect of electron delocalization (electron sharing) occurring between the two fragments upon bringing them closer to each other. This is achieved by using absolute values upon summing the gradient contributions of ED sources (fragments) in the expression of the total ED gradient, leading to the non-interacting reference $\frac{\partial\rho}{\partial x}^{IGM} = \left| \frac{\partial\rho_A}{\partial x} \right| + \left| \frac{\partial\rho_B}{\partial x} \right|$. Thereby, the resulting ED gradient attenuation normally expected between bonding fragments in the so-called "interaction corridor" (see Figure 2) and due to individual terms having opposite signs in the gradient sum $\frac{\partial\rho}{\partial x}$, vanishes. The resulting norm $|\nabla\rho^{IGM}|$ is a virtual upper limit of the real total ED gradient. Hence, the descriptor $\delta g = |\nabla\rho^{IGM}| - |\nabla\rho|$ measures locally the electron sharing, hereafter also referred to as ED contragradience. Two very important points must be emphasized for the rest of this study: (1) only the contragradience between A and B is captured by δg and (2) the same ED is considered for the non-interacting system (IGM) and the real system. This makes the δg descriptor intrinsically linked to the AB interaction in molecular situation. In other words, δg quantifies the AB interaction without resorting to calculations performed on separated fragments.

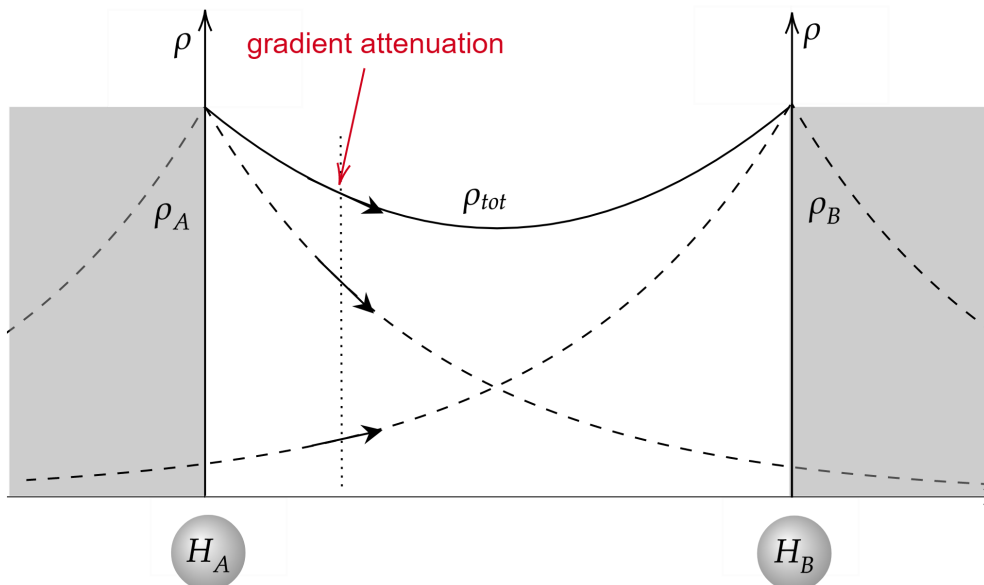


Figure 2: Schematic representation of electron density interpenetration (contragradience) between two fragment sources.

Once δg computed at any point of the three-dimensional grid, a 2D plot can be constructed as illustrated on Figure 1. The first concern in implementing the IGM approach is to choose a gradient decomposition scheme in which each contribution is associated to an ED source to be isolated. For ED coming from quantum-mechanical calculations, this can be done thanks to the Gradient Based Partition (GBP) that we have previously described.¹¹ This partition has been shown to be hardly dependent on the basis set (beyond the small STO-3G basis set). Within the GBP, the most basic ED gradient source that can be considered is an atomic orbital. Regrouping them by atom leads to an atomic partition of the ED gradient. Using this atomic partition yields the δg descriptor extracting the interaction between all atoms of the system. Further, grouping these atomic gradients into two fragments allows for extracting just the intermolecular interaction between them (δg^{inter} , not used in this work). Finally, formulating a gradient partition truncated to two atoms leads to the δg^{pair} bond signature we are interested in, as illustrated in Figure 1. It opens the way to

probe specific bonds in a molecule by using the ED derived from quantum calculations. The local δg^{pair} descriptor was computed using a modified version of NCIPLOT.¹⁷ Such grid computational scheme could leverage the massively parallel GPU technology for the acceleration of δg^{pair} calculation, as demonstrated with NCI in a previous study.¹⁸

Just like in the NCI approach, this δg^{pair} signature can be transposed in the real space taking the form of isosurfaces colored according to the value of signed ED ($sign(\lambda_2)\rho$) (see Figure 1). Clearly, δg^{pair} accounts for the tendency of electrons to be shared between the two atoms. It measures to some extent (not in a direct manner) the kinetic energy lowering resulting from electrons being able to roam over a larger area in the interacting system than in separate fragments. This sharing can take place either in bonding regions or nonbonding regions. Hence, the δg^{pair} bond signature offers two complementary interpretations: (1) the δg^{pair} peak height tells us how much the electron delocalization is important from a kinetic point of view and (2) the ED curvature where the δg^{pair} occurs tells us if we are dealing with an attractive ($\lambda_2 < 0$) or repulsive ($\lambda_2 > 0$) local region from a potential point of view.

IGM bond index IBSI

In order to get a global score for the studied bond, we have decided to carry out the integration of the δg^{pair} signature over the interaction volume V : $\Delta g^{pair} = \int_V \delta g^{pair} dV$. An outstanding advantage of the IGM- δg descriptor in achieving this integration lies in that there is no need to delimit the interaction region. Actually, non-zero values of the δg descriptor exclusively correspond to interaction situations. Thus, the numerical integration is computed as the summation extending over all the grid points lying in the grid volume. No attempt has been made to integrate the ED ρ as others have done. Actually, the 2D δg^{pair} signature seems to hold all the ingredients to characterize bonding patterns and it appears to be a natural candidate for this integration scheme.

The first attempts to compare Δg^{pair} to other properties like bond indices or force constants have shown the need to divide Δg^{pair} by the square of the internuclear distance. The

reason is that ED contragradience only occurs in the space between the two atoms (the interaction corridor) and hence the corresponding integration volume in itself influences the resulting sum Δg^{pair} . For instance, the H_2 bond, with a very short bond length (0.74 Å), is characterized by a narrow interaction corridor. As a result, it is definitely characterized by a too small value of Δg^{pair} compared to other bonds with larger separation distance. Our experience led us to conclude that it is important to make Δg^{pair} independent of the bond size in order to compare different bonds. The resulting IGM bond index, which measures the amount of electron sharing per square unit length, is then: $\Delta g^{pair} = \int_V \frac{\delta g^{pair}}{d^2} dV$. Moreover, to compare between bond indices, the IBSI (Δg^{pair}) has been normalized to 1 for the H_2 molecule:

$$\Delta g^{pair} = \frac{\int_V \frac{\delta g^{pair}}{d^2} dV}{\int_V \frac{\delta g^{H_2}}{d_{H_2}^2} dV} \quad (1)$$

As a consequence, the IBSI is dimensionless. The influence of the level of theory on the gross value of $\int_V \delta g^{H_2} dV$ has been studied and shown to be hardly dependent on both the basis set and level of theory (see ESI). Although the IBSI value for H_2 is rather stable, it is recommended to employ the same level of theory to compare IBSI values across different molecules.

Based on case studies (see supporting information), we have shown that the IBSI is generally not significantly basis set dependent (beyond the STO-3G basis). The extent of variability of the IBSI measured by the relative standard deviation is generally in the range 2-6% for the studied organometallic compounds, generally more homogenous compared to Mayer, Wiberg and Mulliken bond orders. We strongly advise not using the STO-3G basis set, but we recommend the use of 6-31G**, [although adding a diffuse function could be more relevant for weakly bonded systems](#). For atoms beyond Kr, the Def2-TZVP^{19,20} basis set can be used (available for all elements up to Radon) or an effective core potential calculation can be performed (with LANL2DZ²¹⁻²³ basis set). In extreme cases, in very polar bonds like LiH, AlO, AlCl, MgC, a larger dependence has been observed (around 10-20%).

A comparative case study of ab initio (HF, MP2) and DFT (B3LYP, SVWN, BP86 and M06-2X) methods indicates that the IBSI hardly depends on the method used (coefficient of variation of at most 7% in extreme cases, but more generally around 2%). We advise not using the HF method. We therefore recommend the DFT/6-31G** level of theory for its performance/price ratio and using the same method for comparative studies.

Molecule test set

This work aims at finding a physically well-founded characterization of the descriptor δg^{pair} and its integrated form Δg^{pair} for a given atom pair. To this end, a 235-molecule test set has been considered, both organic and inorganic, covering typical functional groups (see ESI) and leading to the study of 677 bonds. It is not limited to known molecules but also includes uncommon compounds. Neutral, charged and radical species have been studied. This large set of molecules covers a full range of bonding interactions, including covalent bonding, metal coordination complexes (with Cr, Fe, Co, Ni, Ru, Re, Pt), weakly hydrogen-bond and halogen-bond complexes.

Quantum-mechanical calculations

In practice, the electron density (ED) is obtained from a quantum-mechanical calculation. In this study, the Gaussian16 package²⁴ was employed. If not otherwise stated, full molecular geometry optimizations were conducted at the DFT(M06-2X/6-31G**) ²⁵ level of theory in the gas phase. Some species with heavy atoms required using the Los Alamos effective core potential through the LANL2DZ basis set specification in gaussian. The integral SuperFineGrid option has been employed in numerical differentiations. Preliminary case studies employed additional levels of theory (HF, MP2) and several other DFT functionals (B3LYP, SVWN, BP86). The harmonic frequency analysis was systematically performed to ensure the absence of imaginary frequencies for local minima.

Bond orders

The Mulliken bond population analysis,²⁶ the Wiberg²⁷ and the Mayer^{28,29} bond indices were calculated using the MultiWFN package.³⁰ The delocalization index (DI)^{31,32} was obtained using AIMall.³³ We have also carried out an electron localization function (ELF) topological and population analysis,^{3,4} which is frequently adopted as a descriptor of atomic shells and covalent bonds using the TopMoD package.³⁴ For simplicity, the ELF approach description is reported in Supporting Information.

Stretching force constant

The bond stretching force constant k was also calculated as an estimator of the bond strength in order to be compared with the IBSI. However, it makes no sense to use the standard vibrational normal-mode analysis to derive a stretching force constant of a particular bond in a polyatomic molecule. Actually, a normal mode of vibration commonly couples to others, it is delocalized over more than the two atoms of interest. To derive a force constant k associated with a local vibrational mode between two given atoms X and Y, a numerical procedure was used in which the second derivative of the potential energy is numerically assessed (see ESI). A purely localized bond force constant k is however difficult to determine because the geometry and the ED distribution in the XY surrounding must be kept frozen as much as possible during the numerical procedure so that k is only inherent in the XY bond. Two different procedures were considered. In the first one, the molecule is divided into two rigid fragments translating along the XY direction, leading to a force constant hereafter called k_1 . It cannot be computed for bonds belonging to a ring. In this first procedure, although non-bonding interaction initially present between separating fragments will contribute to a small extent to k_1 , k_1 is more intrinsic to the XY bond strength than the second procedure k_2 . In the second version, only X and Y are displaced in the molecular framework, leading to a force constant called k_2 . It is highly important to note that, in this second version, the fixed substituents attached to XY indirectly significantly contribute to the resulting k_2

via steric and electronic effects upon XY displacement. In the following, the k_1 and k_2 local force constants have been obtained and compared using the set1 of 184 molecules not containing rings (438 bonds). k_2 has also been determined for 51 ring-containing molecules (239 bonds, see supporting information). Anharmonic computations were performed for a set of diatomics using the program Gaussian.²⁴ For each diatomic molecule, the X matrix representing the anharmonicity is made up of only one term given in cm^{-1} and involved in the correction of the potential. This term has been employed to compare the anharmonicity contribution across some diatomics.

Results and discussion

A systematic study has been performed on 235 molecules and 677 different bonds covering a broad range of chemical bondings (non-covalent bonding, metal coordination and covalent bonds, see ESI). When we say bond property, we naturally think of bond order. That is the reason why we have first addressed the possible link between the IBSI and bond orders. This preliminary study reported in Figure 1 in ESI demonstrates that the IBSI is clearly not correlated with the examined bond orders: Mayer ($R^2 = 0.41$), Wiberg ($R^2 = 0.37$), Mulliken ($R^2 = 0.27$), Delocalization Index ($R^2 = 0.46$), ELF ($R^2 = 0.007$).

From a chemical perspective, the inspection of the IBSI computed for H-X bonds across the periodic table undoubtedly shows a certain degree of consistency: for instance, it gradually increases from left to right along the second period of elements, from a low value for H-Li (IBSI=0.06) to the largest value for H-F (IBSI=1.63) (see Figure 2 in ESI). Hence, we subsequently looked into a correlation between the IBSI and the stretching force constant concept.

IBSI versus stretching force constant

Actually, we may suppose that the more the electron sharing between the two atoms at equilibrium geometry is pronounced, the bigger the force to stretch the bond. In a preliminary

examination, we have compared the IBSI with the stretching force constant k_1 calculated for a series of 5 homonuclear diatomic molecules involving the diatomic oxygen.

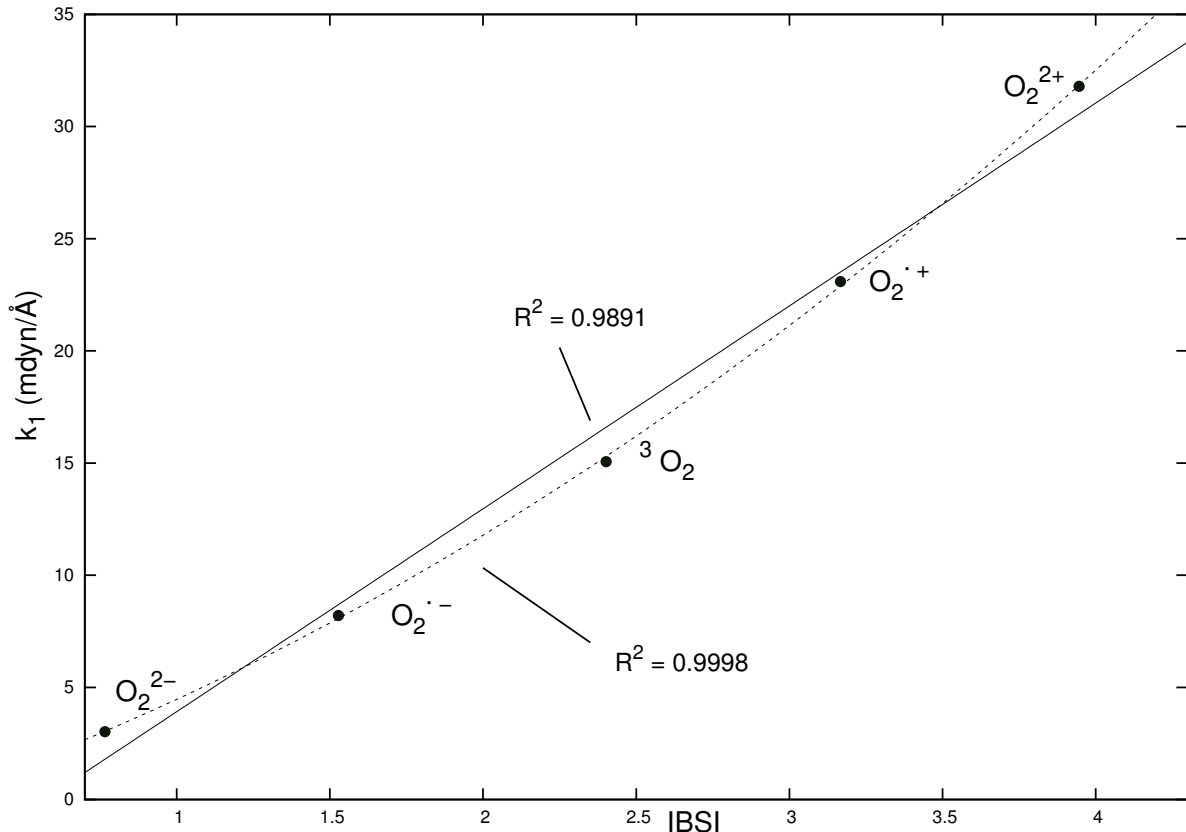


Figure 3: Relationship between the stretching force constant (scheme k_1) and the IBSI for 5 diatomic oxygen species; fitted linear (full line) and quadratic (dashed line) regression curves are reported.

As can be seen on Figure 3, the linear correlation between these two properties is very strong ($R^2 = 0.99$). We know that placing electrons on antibonding orbitals will decrease the stability of a molecule. This is what happens across the series from O_2^{2+} to O_2^{2-} . Accordingly, the conventional bond order calculated as $\frac{1}{2}[(\text{number of bonding electrons}) - (\text{number of antibonding electrons})]$ decreases. The lower this bond order, the weaker the bonding. The IBSI fully reflects this trend over a wide range of chemical bonds in this series, taking values between 3.95 for O_2^{2+} down to 0.77 for O_2^{2-} . Although this set is limited to 5 molecules, the observed trend is noteworthy.

This preparatory work has been extended to 438 bonds (set1). The result is reported on Figure 4 (k_1 versus IBSI). The obtained linear correlation is surprisingly high with 93% of the force constant variance being explained by the IBSI. A close inspection of the plot on Figure 4 (and also on Figure 3) reveals that the [straight](#) line is not the best shape to represent the data. An attempt to fit a quadratic function to the data significantly improves the

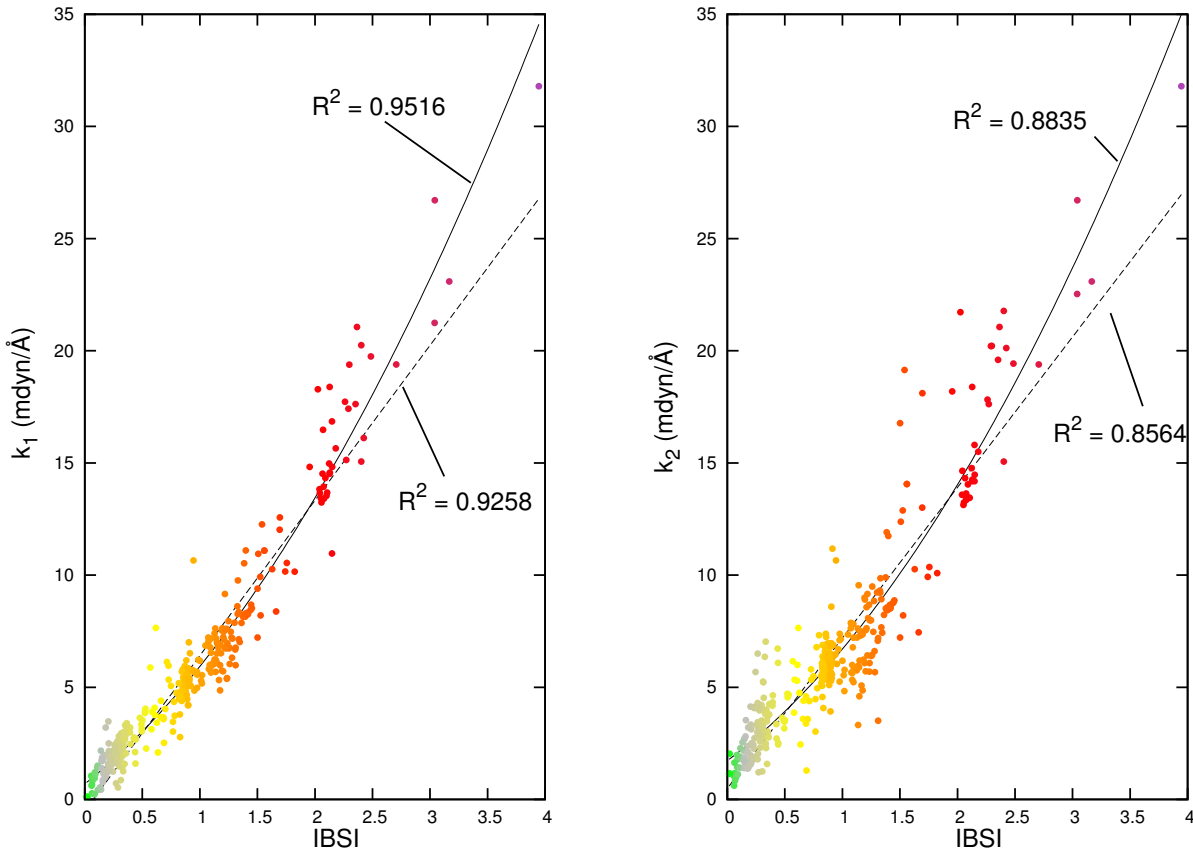


Figure 4: Relationship between the stretching force constant and the IBSI for the set1 of 438 bonds in 184 molecules (with no rings since k_1 would not be calculable in that case) for (a) scheme k_1 and (b) scheme k_2 ; fitted linear and quadratic regression curves are reported. The points are colored using the color scale given in Figure 5.

correlation, with $R^2 = 0.95$ on Figure 4 (left panel). It seems clear from the data presented here that the IBSI is deeply connected with the stretching bond force constant concept. The spread of the points about the fitted line is relatively narrow in spite of the very different chemical bonds explored here: from weak non-covalent interactions (hydrogen and halogen

bonding) to metal coordination and strong covalent bonds. Based on our work, one observes that the IBSI values range from 0.02 to 3.95 and that three domains can be distinguished as illustrated on Figure 5. In this work, covalent bonds are characterized by an IBSI value larger than 0.15. Transition metal coordination occurs in the IBSI range 0.15-0.6. Ionic pair interaction as well as hydrogen- and halogen-bonding involve IBSI value lower than 0.15. This scale and limits are only indicative for the set of 677 examined bonds and has been proposed to make life simpler for IBSI users.

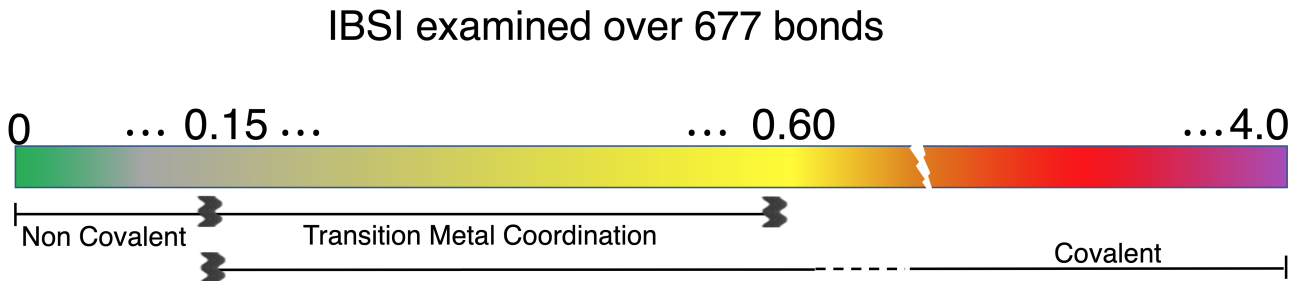


Figure 5: IBSI scale obtained on the set of 677 studied bonds. Limits are only indicative for the set of 677 bonds examined here.

In this work, 10 simple ionic compounds with large ionic character (like NaCl, KCl, LiH, LiF, LiCl, ...) have also been investigated. Such species are stable mainly because of the electrostatic attraction between atoms and the energy required to break the bond may be as large as for covalent bonds. However, simple ionic compounds exhibits low stretching force constant values compared to covalent bonds.³⁵ Here, the corresponding IBSI values are predicted to be very small (< 0.15), below the IBSI covalent domain ($0.15 < \text{IBSI} < 4.00$). This fits very well with the low value of the force constant k_1 ($< 3 \text{ mdyn}/\text{\AA}$) calculated for these ionic species. But it does not reflect the strong bond dissociation energy expected for such very polar molecules with respect to dissociation either into the neutral atoms or into ions. Clearly, the IBSI is not related to the bond dissociation energy but rather to the local stretching force constant.

For the sake of convenience and to save CPU time, the 6-31G** basis set has been used

throughout this work for molecules made up of elements in rows 1-3. But the influence of adding diffuse functions has been addressed. Adding a diffuse function to this basis set (namely, 6-31+G**) does not change the k_1 /IBSI correlation. However, in absolute term, the most important changes are observed for those systems being weakly bonded, like for instant for Li-F, Mg-Cl in CH_3MgCl , and certain non-covalent bondings (see ESI). In that case however, it is not attributable to the change in geometry. In other respects, it turns out that adding polarization functions to the LANL2DZ basis set (using either LANL2DZdp for the main group elements and the hydrogen atom, or LANL2TZf for the metal) does not change the force constant k_1 /IBSI correlation. However, it has to be noticed that, actually, both IBSI and k_1 can experience significant variations, in absolute terms. This is notably observed for metal-ligand bonding (IBSI(Pt-Cl)=0.216-0.278, IBSI(Pt-N)=0.312-0.360). This is partly because of the geometry change, but not only. As a consequence, the LANL2DZ or LANL2TZ basis sets augmented with polarization functions, although significantly more CPU-expensive, are advised.

Differences between IBSI and the force constant k

Undeniably, the IBSI is strongly connected to the physically grounded bond strength concept. The force constant k being the second derivative of the energy with respect to the internuclear distance at the equilibrium state, \AA^{-2} appears in the unit of k . In comparison, although the IBSI is dimensionless (see Equation 1), its value is, by construction, inversely proportional to the square of the bond length d^2 , which is consistent with the unit of k . But, despite the good agreement between the IBSI and the force constant k , the fraction of the variance unexplained is 5% (quadratic fit). We can try to find reasons for that.

It is highly important to notice that the static IBSI calculation is fully inherent to the considered atom pair while the force constant k_1 accounts for side effects. Actually, the local force constant may have a significant dependency on the chemical environment surrounding the given bond XY. For instance, non-bonded interactions between adjacent substituents on the two neighboring carbon atoms of 1,2-dibromoethane impact to some degree the C-C

stretching local force constant value. The effect of the bonding environment on the force constant k can be gauged by comparing the correlations k_1 /IBSI and k_2 /IBSI for the same set of bonds. Indeed, the alternative numerical differentiation scheme k_2 is less intrinsic to the studied bond than force constant k_1 (see Computational details section). As expected (see Figure 4, right panel), this second scheme k_2 yields a poorer quadratic correlation with the IBSI using the same set of molecules without rings ($R^2 = 0.88 < 0.95$ for k_1). This is not surprising since k_2 (atoms X and Y moving in the rigid molecular framework) is more sensitive to the substituents attached to the studied bond than k_1 (rigid fragmentation along the XY bond). This is further evidenced by investigating a second set of bonds XY exclusively located in rings (see ESI, Figure 4, $k_2=f(\text{IBSI})$). Stretching such bonds substantially disrupts the ED in the bond surrounding. The resulting determination coefficient dramatically drops to 0.74. This clearly demonstrates that the presence of ring constraints in the vicinity of the considered bond significantly affects the deduced force constant k_2 , but not the IBSI, hence deteriorating the IBSI- k_2 quadratic correlation. Conversely, for diatomic molecules XY where no environment effect can be invoked, the k_1 and k_2 schemes are identical and lead to a good quadratic correlation with the IBSI ($R^2 = 0.95$, See Figure 6).

In fact, the IBSI and force constant k are structurally different. The second derivative of the potential energy with respect to the geometric displacement corresponds to the curvature of this function. Hence, k indirectly depends on the total structure ED reorganization upon stretching the bond embedded in its molecular framework. In comparison, the static IGM- δg^{pair} approach exclusively focuses on ED between atoms X and Y. More precisely, it captures the ED gradient reorganization between two states: the real system and a non-interacting reference (the IGM), which are only different in the middle of the two atoms where the phenomenon of contragradience takes place (see Figure 2). Therefore, the IGM approach does not account for any ED reorganization outside the bond region. Moreover, this ED gradient reorganization (δg^{pair}) is estimated solely based on a single geometry ED, without considering any ED "breathing" between the two states, in contrast with the calculation of k .

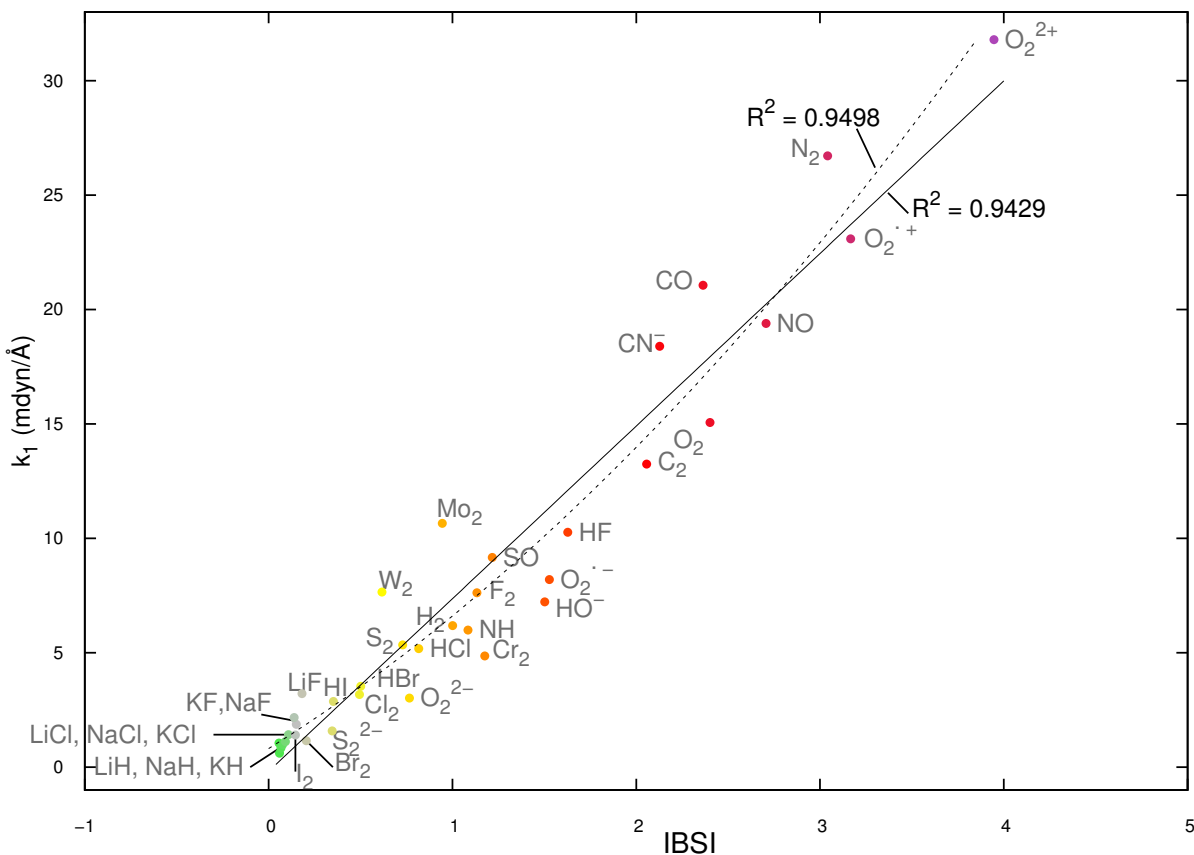


Figure 6: Relationship between the stretching force constant and the IBSI for 36 diatomics; fitted linear and quadratic regression curves are reported. The points are colored using the color scale given in Figure 5.

This might partly explain why diatomics featuring strong multiple bonding (N_2 , CO , CN^- , MoO_2 , W_2) lie all above the [two fitted lines](#) on Figure 6 showing the correlation between k_1 and IBSI. Actually, such multiple bonds undergo strong ED relaxation upon infinitesimal stretching used to numerically assess the force constant k .

Another way of calculating a stretching force constant associated with just one bond length is to apply the Konkoli-Cremer adiabatic internal modes (KC-AIM).^{36,37} This approach is particularly well suited to obtain a stretching force constant k^a not contaminated by coupling with other modes (in contrast with normal modes). As with the local k_1 and k_2 numerical schemes used in our work, the KC-AIM approach probes the bond strength in a non-destructive manner (it leaves the considered bond undissociated). However, the

KC-AIM approach has the additional benefit of allowing the rest of the molecule to relax upon considering a perturbation to the considered bond length. Based on KC-AIM mode force constants k^a taken from reference³⁸ for 39 molecules and 145 bonds, an additional comparison has been reported between IBSI and k^a (see ESI Figure 6). It shows a significant correlation between these two properties ($R^2 = 0.92 - 0.95$), very similar to the one obtained between IBSI and local force constants k_1 and k_2 (see Figure 4) previously approximated numerically by a three-point central difference formula. But again, a small fraction of the variance remains unexplained. k^a , k_1 and k_2 are dynamic descriptors: a local force constant reflects the potential energy curvature associated with the vibration within its chemical environment. For its part, the IBSI provides a means for assessing the intrinsic bond strength without being affected by adjacent atoms. It measures the strength of the attachment between two considered atoms through their degree of ED contragradience. It does not rely on estimating any potential curvature, and then does not depend on how neighboring atoms would affect the vibration. Rather, it quantifies the ED clash between two given atoms as compared to the contragradience-free system (virtual) having exactly the ED of the true system. That is the reason why the IBSI and force constant are conceptually different. The static bond descriptor IBSI can be utilized in any molecular situation, equilibrium geometry or even along a reaction path where the force constant is no longer connected to the restoring force concept.

Another reason can be given to explain differences between IBSI and the force constant k . The level of theory used could rise differences in the treatment of both k and the IBSI. Actually, the IBSI depends on the ED and the geometry (d^2) and then, to a certain degree, on the level of theory chosen. Similarly, the second-order three-point central difference formula employed to approximate the second derivative k also depends, to a certain extent, on the level of theory employed. Based on case studies, the IBSI has been proven to be only slightly dependent on the level of theory (see ESI), and the DFT(M06-2X/6-31G**) level of theory has been chosen here. This popular functional is recommended for applications involving

main-group elements and non-covalent interactions.²⁵ Of course, small changes in ED and geometry might be expected upon using more sophisticated methods.

In other respects, avoided crossing, as observed in C_2 between states $X^1\Sigma_g^+$ and $B'^1\Sigma_g^+$ ³⁹ may affect the stretching force constant derived from the ground-state adiabatic potential energy curve. In the presence of such avoided crossing, the stretching force constant obtained using a single determinant wave function cannot serve as an accurate bond strength descriptor, in absolute terms. Neither does the IBSI, since such avoided crossing affects in the same way the electron density used to obtain the IBSI at the equilibrium geometry. Similarly, a DFT level of theory based on a single Slater determinant description cannot describe appropriately diatomics having multireference character. However, in our study, Figures 5 and 8 (diatomic systems) aim at proving that IBSI and force constant, given a same level of theory (DFT here), are connected. An in-depth description of bonding of diatomics (like C_2 ,³⁹ Mo_2 or W_2 ⁴⁰ or F_2 ⁴¹) would require highly accurate and multi-reference methods, very CPU-expensive approaches, beyond the scope of this work.

For the set of diatomic molecules, another factor has been investigated to explain the remaining difference between the IBSI and k_1 obtained within the harmonic approximation. Anharmonic corrections to vibrational frequencies have been computed using the Gaussian program.²⁴ However, diatomics exhibiting significant anharmonicity could not be associated to the most important deviations observed on the k_1 /IBSI graph.

In summary, the force constant k and IBSI, although both strongly correlated to the bond strength concept, do not rigourously express the same physical content. The IBSI is intrinsically linked to the atom pair considered and provides a very efficient way of measuring a local bond strength in molecular situation, without any bias due to the bonding environment, even in cyclic structures. In practice, the IBSI has the advantage of requiring a single point energy calculation.

IBSI: scope of application and limitations

The IGM- δg descriptor allows for the proper identification of ED contragradience domains in the molecule. The IBSI quantifies this contragradience in the region between two atoms and only considers the orbital contributions of these two atoms. Compared to the local stretching force constant k^a obtained by the Konkoli-Cremer procedure, the IGM-IBSI scheme additionally provides information through the local distribution of ED contragradience not necessarily uniform nor located at the center of the atom pair. In most cases, in the space region in the middle of the considered pair, the ED contragradience dominantly arises from two atomic sources. Then most often, the ED contragradience domains are well separated in space, as illustrated in Figure 7 of the manuscript showing a TS structure. But, what if a third atom brings some ED gradient in the considered atom pair region ? When does this occur ? To illustrate this point, the interactions in the B_2H_6 molecule have been identified and characterized thanks to the full IGM approach with $\delta g = |\nabla \rho^{IGM}| - |\nabla \rho|$. As can be seen in the Figure 8 of ESI, two contragradient domains very slightly overlap around each H_{bridge} atom. Then, one may wonder whether the B_1-H_{bridge} bond strength can be assessed based solely on atom pair contragradience. To which extent B_2 is really involved in the B_1-H_{bridge} contragradience domain (if it is) cannot be assessed within the current IGM approach. Even though this could be achieved, it would make little sense to keep a single geometrical parameter (B_1-H_{bridge} distance) in the IBSI definition in that case. The user must be aware that the IBSI approach relies on atom pairwise interactions. Despite this, the IBSI faithfully reflects the relative weakness of the B_1-H_{bridge} bond (IBSI=0,47) compared to the $B_1-H_{terminal}$ bond (IBSI=0.55). To our knowledge, this kind of situation should be limited to very small ring systems with atoms having electrons more easily spread. Examining the cyclopropane molecule (see Figure 9 in ESI), in addition to the well separated 2-center contragradience domains (C-C), a 3-center domain appears in the center of the ring. Despite this, the IBSI correctly describes the C-C bond strength. Actually, the relative C-C bond strengths estimated with the local KC-AIM force constant in propane ($k^a = 4.066$ mdyn/ \AA) and cyclopropane (4.137 mdyn/ \AA) fully support the IBSI rating (0.839 and 0.892,

respectively; see ESI in section Local KC-AIM force constants versus IBSI).

The IBSI approach is not restricted to well behaved 2-center bonding systems. Delocalized aromatic systems are suitably described. For instance, the C-C intrinsic bond strength in C_6H_6 (IBSI = 1.197) is found in between that of a single bond (IBSI = 0.833 in C_6H_{12}) and a double bond (IBSI = 1.396 in C_2H_4), as supported by the local KC-AIM C-C force constants k^a : 6.600 mdyne/Å in C_6H_6 , in between 3.923 mdyne/Å in cyclohexane and 9.911 mdyne/Å in ethene (see ESI). The ability of the IBSI approach to properly describe the bond strength in a 3-center 2 π -electron system is also observed with the allyl cation (IBSI C-C = 1.169). In other respects, bonding having a 3-center 4-electron character as in certain hypervalent molecules⁴² or in systems exhibiting hydrogen or halogen bonds⁴³ is also well described since the contragradience domains are well separated in such cases. For instance, the F-Cl...NH₃ system exhibits IBSI values of 0.691 (F-Cl) and 0.101 (Cl...N), as supported by the relative local KC-AIM force constants k^a : 2.687 mdyne/Å (F-Cl) and 0.311 mdyne/Å (Cl...N).⁴³

Another kind of bonding deserves attention: when a ligand is bound to a metal via a conjugated π -system like in metallocenes. For illustration purpose, the interaction between the cyclopentadienyl anion (Cp) and the dication Zn^{2+} has been reported in ESI (Figure 10). The IGM approach unambiguously reveals the region of interaction. The Cp... Zn^{2+} contragradience domain takes the form of a continuous ring for this η^5 hapticity mode. The IBSI value obtained for each C... Zn^{2+} pair is 0.194, which clearly reveals a bonding weaker than pure covalent bonds like C-C in Cp (IBSI=1.081). However, in such case, although the AIM analysis predicts a bond critical point between the carbon atom and Zn^{2+} , the use of the pairwise IBSI scheme is tricky since there are more than two atoms simultaneously involved in the coordination to the transition metal.

The agostic interaction is another type of challenging interaction, occurring between an electron deficient metal and a σ -bond located close to the metal.⁴⁴ For illustrative purpose, the complex $[Ti(Cl_2)C_2H_5]^+$ has been reported in ESI (Figure 11). The IGM approach

reveals this agostic interaction with the contragradience domain being located between C-H and titanium. The relative weakness of this interaction is correctly predicted by the IBSI calculation (IBSI=0.105), compared to the Cl-Ti coordination (IBSI=0.353). However, this agostic interaction involves three atoms while the IBSI scheme is intended for pairwise interactions.

To summarize, we are fully confident that the IBSI approach correctly describes the intrinsic bond strength between atom pairs in the vast majority of molecules, including transition state structures, non-covalent hydrogen bonds and halogen bonds, as long as contragradience domains are well separated. However, there are limitations in interpreting IBSI for interactions with complex bonding scenarios where more than two atoms are involved in a bonding situation. Metallocenes and even non-covalent stacking interaction, like in the benzene dimer, belong to this category. It is beyond the scope of this work to explore more in details such cases. However, whether the IBSI approach can be used is usually obvious by inspection of the molecular geometry and the shape of the contragradience isosurfaces.

Bridge between local ED descriptor and bond strength

To our knowledge, only one previous study reported such a bridge between a local ED-based descriptor and a bond stretching force constant. Very recently, considering a set of 53 molecules, a strong linear correlation ($R^2 = 0.97$) was presented between the localized stretching force constant of a given bond and the potential energy density distribution integrated over the surface between the two Bader’s atomic basins.¹⁵ In a similar way as in our study, the authors divided the resulting integral by the internuclear separation. However, this approach needs to characterize the ED topology (by determining critical points and basins, ...) prior to any calculation. In contrast, a compelling feature of the IGM model is to provide an automatic workflow that delivers the IBSI without any ED topology analysis. We believe that the IBSI is a powerful and robust tool for interpretation accessible to a wide community of chemists (organic, inorganic chemistry, including transition metal complexes and reaction mechanisms).

A typical application of IBSI

A review of possible applications of the IBSI is beyond the scope of this study. However, in order to show the possibilities offered by the IBSI compared to the Mayer bond index we have reported on Figure 7 a transition state structure (TS) taken from a recent study on the reactivity of thiolates with cisplatin.⁴⁵ The reactant is the complex $[\text{Pt}(\text{NH}_3)_2(\text{H}_2\text{O})(\text{cystein})]$. The mechanism proceeds with the intramolecular substitution of the water molecule by the $-\text{CO}_2^-$ group of the cystein, which finally acts as a (S,O)-bidentate chelating agent in the final product.

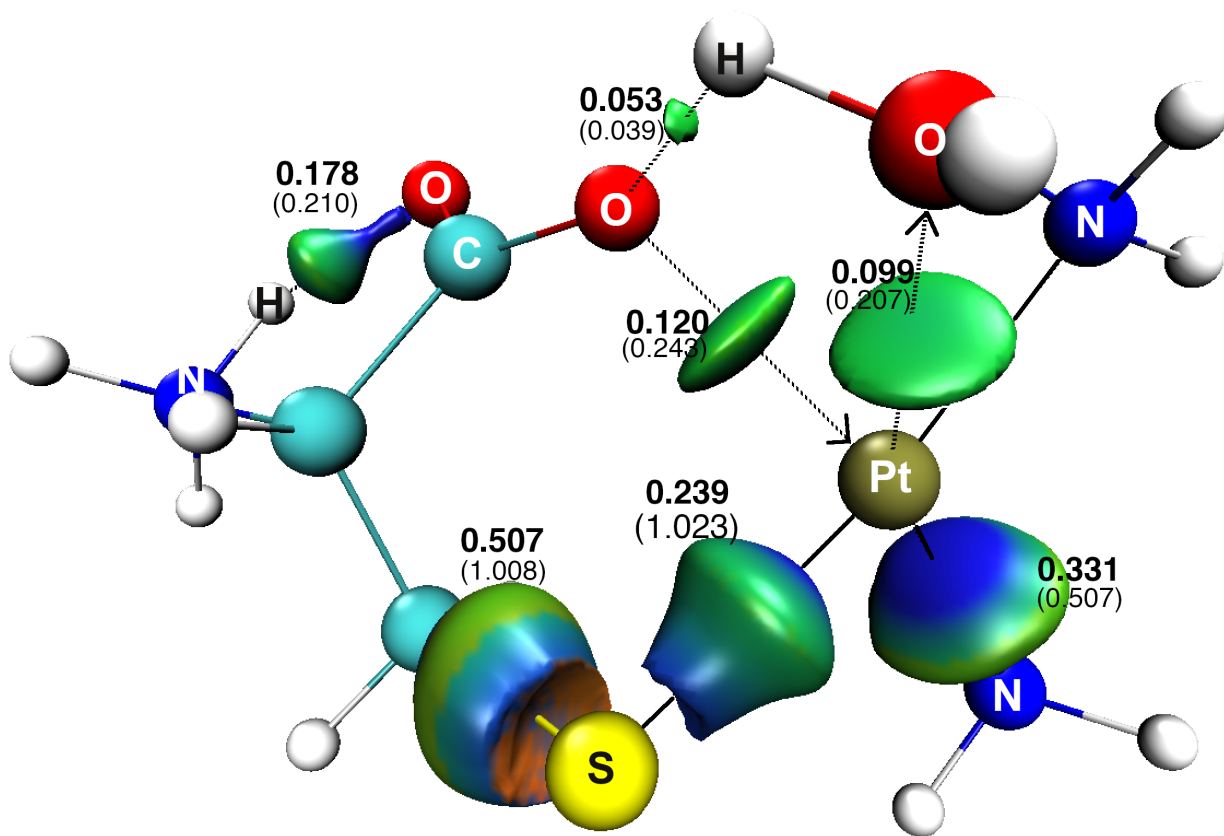


Figure 7: Bond Strength Index (IBSI) and $0.045 \delta g$ isosurfaces for selected bonds in a transition state involving a platinum complex (M06-2X LANL2DZ level of theory); isosurfaces colored according to the BGR scheme over the range $-0.2 < \text{sign}(\lambda_2)\rho < 0.2$ a.u.; Mayer bond order in parentheses.

According to the Mayer analysis (MBO), the C-S and S-Pt bonds exhibit an identical bond multiplicity of about 1.0. The IBSI provides an additional information: the covalent bond C-S is stronger (IBSI=0.507) than the S-Pt coordinate bond (IBSI=0.239). This is consistent with the strength of metal ligand bonds known to be intermediate between that of covalent bonds and non-bonded interactions. In other respects, the MBO predicts a formal half-bond between the nitrogen and platinum atoms (0.507 pair) and a single bond between the sulfur and platinum atoms (1.023 pair). It may be then naively expected that the corresponding metal - ligand bond strengths should correlate with these bond orders at a ratio of 1(N-Pt):2(S-Pt). However, the IBSI predicts an inverse strength ratio with the N-Pt bond being significantly stronger (IBSI=0.331) than the S-Pt one (0.239).

This TS has one imaginary vibrational frequency with a negative force constant no longer connected to the restoring force concept. Nevertheless, the IBSI stays attached to this notion, allowing for internally probing bond strength in TS structures and along reaction pathways. In the course of this reaction, the breaking Pt...OH₂ and forming -CO₂⁻...Pt chemical bonds undergo electronic structure changes. As expected, the MBO analysis at the TS shows that very few electron is shared between these reacting atoms (around 0.2 pair), in accordance with the small degree of orbital overlap achieved between the corresponding atoms at the TS. This decreased orbital overlap also weakens the two corresponding bonds, in the IBSI range 0.10-0.12: at the boundary between non-covalent bonding and coordinate covalent bonds (see Figure 5).

Usually, in hydrogen bonding, little or no electron pair is shared by the valence shells of the two atoms, making the interpretation of the Mayer bond order generally quite difficult. Here however, the MBO detects an uncommonly large electron sharing (0.210) for NH...O compared with OH...O (0.039) in this TS. In a complementary way, the IBSI analysis reveals that the picture of the hydrogen bond is deceptively simple for NH...O hydrogen. Actually, its bond strength (0.178) is outside the known H-bonding range (generally ≤ 0.15 on the IBSI scale, see Figure 5) and more than three times larger than the OH...O one (0.053).

This is because the former exhibits a short distance (1.6 Å) and a strong acid/base character between the two charged entities -NH_3^+ and -CO_2^- . In light of this short example, we see that the IBSI and MBO analyses provide a complementary information to explore chemical bonding and electronic effects.

Conclusions

In this work, we present for the first time the Intrinsic Bond Strength Index (IBSI). It is a post-processing tool very efficient to internally probe the strength of a given chemical bond in molecular situation. This quantitative tool complements the conventional visual usage of the IGM approach which provides colored isosurfaces highlighting the spatial localization of interactions in molecules. The theoretical advantage of the IGM method rests on the definition of a non-interacting reference system (the so-called Independent Gradient Model). This feature allows the quantitative determination of bond strength. In this regard, to the best of our knowledge, the IBSI is distinct from other ED-based approaches like SEDD, DORI or NCI. Based on the large range of compounds explored in this work (235 species and 677 atom pairs), we suggest to approximately distinguish between weak interactions: $\text{IBSI} \lesssim 0.15$, coordinate covalent bond: $0.15 \lesssim \text{IBSI} \lesssim 0.6$, and covalent bonding: $\text{IBSI} \gtrsim 0.15$ (these limits being only indicative).

The IBSI is defined in a parameter-free manner and its implementation is very easy: it only requires a wavefunction file as input. It has been shown to be little dependent on the level of theory and basis set. From the practical point of view, an attractive feature of the IGM-IBSI approach is to provide an automatic workflow that delivers the bond strength index without any prior characterization of the ED topology (unlike within the AIM theory). In other words, it is not an ED basin-based method like ELF or AIM.

From a theoretical perspective, the current work provides a quantitative bridge between a local electron density-based descriptor and the physically grounded bond strength concept. Actually, the IBSI emerges from the IGM formulation of electron sharing between two atoms. This index results from the integration of the local descriptor IGM- δg^{pair} bond signature. Through a quadratic relationship, a tight link ($R^2 = 0.95$) has been established between the IBSI and the localized stretching force constant of a chemical bond. This result underpins the IGM formalism and its two key-components: the non-interacting reference definition and the gradient based partition. [Future work could also consider to seek a correlation between the IBSI and ED characteristics at bond critical points.](#) It has however to be noticed that by construction the δg descriptor is limited to theoretical ED models, which rules out the use of experimental data.

Differences exist between the IBSI and a local force constant. It is shown that the IBSI relates to the intrinsic bond strength. It is obtained by focusing only on the atom pair considered. No expensive calculations of energy second derivatives are needed to obtain the IBSI. In contrast, a local force constant calculation may depend on the bonding environment and the electron density reorganization occurring during bond stretching. Furthermore, along reaction paths, even in regions with imaginary vibrational frequencies where the force constant is no longer connected to the restoring force concept, the IBSI stays attached to this notion, extending its scope beyond equilibrium geometries.

Clearly, this index covers a broad range of bonding situations and is not limited to closely related molecules. This makes it suitable for following bond formation and breaking in chemical reactions. In particular, the IBSI turns out to be a suitable indicator for predicting a realistic intrinsic strength of a given chemical bond in transition states. Changes in stretching force constants upon bond polarizing of a reaction partner is a key element of the unified reaction valley approach^{37,46,47} used for the detailed mechanistic analysis of

elementary chemical reactions. Hence, changes in IBSI occurring in the course of a chemical reaction might contribute to gain insights into reaction phases each of which leads to a change in chemical bonding. More generally, this index has the advantage of being applicable to diverse fields in chemistry and future work will be devoted to applications for systems in organic and inorganic chemistry, transition metal complexes and reaction mechanisms. The next release of IGMPlot including the calculation of the IBSI will be made available at <http://igmpplot.univ-reims.fr>.

Acknowledgement

This work was supported by a Grant (J. Klein, Master’s thesis) from the Université de Reims Champagne-Ardenne and the ICMR laboratory (Institut de Chimie Moléculaire de Reims). We thank the MaSCA (Maison de la Simulation de Champagne-Ardenne, France) for various supports including computing facilities (<http://romeo.univ-reims.fr>) and the CRIANN computational center (<http://www.criann.fr>) for additional support. We thank Prof. F. Bohr for helpful discussions.

Supporting Information Available

Detailed molecule sets used throughout this study and computed properties (IBSI, k^a , k_1 , k_2 , bond orders). Investigation of the basis set and method dependence on the IBSI.

This material is available free of charge via the Internet at <http://pubs.acs.org/>.

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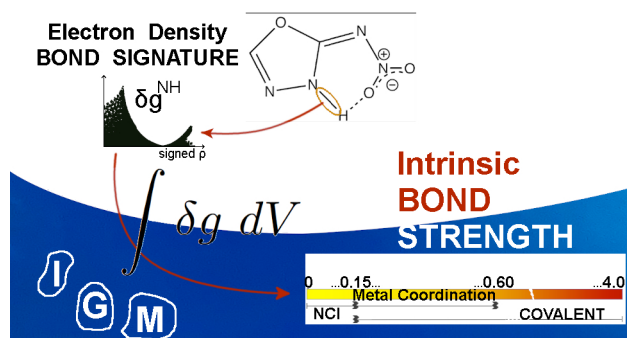


Figure 8: For Table of Contents Only