See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/24264709

# Steric Effect: Partitioning in Atomic and Functional Group Contributions

RTICLE in THE JOURNAL OF PHYSICAL CHEMISTRY A · MAY 2009							
Impact Factor: 2.69 · DOI: 10.1021/jp8096583 · Source: PubMed							
CITATIONS	READS						
40	60						

# **3 AUTHORS**, INCLUDING:



Shubin Liu

University of North Carolina at Chapel Hill

155 PUBLICATIONS 5,483 CITATIONS

SEE PROFILE



Subscriber access provided by UNIV OF NORTH CAROLINA

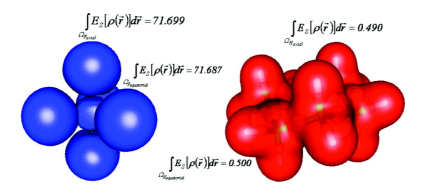
# **Article**

# Steric Effect: Partitioning in Atomic and Functional Group Contributions

Miguel Torrent-Sucarrat, Shubin Liu, and Frank De Proft

J. Phys. Chem. A, 2009, 113 (15), 3698-3702 DOI: 10.1021/jp8096583 Publication Date (Web): 12 March 2009

Downloaded from http://pubs.acs.org on April 13, 2009



# **More About This Article**

Additional resources and features associated with this article are available within the HTML version:

- Supporting Information
- Access to high resolution figures
- Links to articles and content related to this article
- Copyright permission to reproduce figures and/or text from this article

View the Full Text HTML



# Steric Effect: Partitioning in Atomic and Functional Group Contributions

# Miquel Torrent-Sucarrat, Shubin Liu, \*\* and Frank De Proft\*\*

Eenheid Algemene Chemie, Faculteit Wetenschappen, Vrije Universiteit Brussel, Pleinlaan 2, B-1050 Brussels, Belgium, and Research Computing Center, University of North Carolina, 211 Manning Drive, Chapel Hill, North Carolina 27599-3420

Received: November 01, 2008; Revised Manuscript Received: February 09, 2009

The recently proposed density-based quantification of the steric effect, based on an alternative energy partition scheme where the total electronic energy is decomposed into contributions from three independent effects, steric, electrostatic, and fermionic quantum, is investigated at the atomic and functional group levels in this work. Reasonable trends and linear relationships between theoretical and experimental scales of the steric effect at both group and entire molecular levels have been observed, providing further evidence that the newly defined quantity can serve as an intrinsic measurement of the steric effect for molecular systems.

### 1. Introduction

The steric effect is one of the most widely used concepts in chemistry and describes "the effect on a chemical or physical property (structure, rate, or equilibrium constant) upon introduction of substituents having different steric requirements." The effect is basically associated with the fact that atoms and larger parts of a molecule occupy a certain region of space. When atoms or groups are brought together, hindrance will be induced, resulting in changes in shape, energy, reactivity, etc. Steric effects arise from contributions ascribed to strain as the sum of nonbonded repulsions, bond angle strain, and bond stretches or compressions. For the purpose of correlation analysis or linear free-energy relation, various scales of steric parameters have been proposed in the literature. Taft<sup>2</sup> constructed a scale for the steric effect of different substituents, based on rate constants for the acid-catalyzed hydrolysis of esters in aqueous acetone. It was shown that  $log(k/k_0)$  was insensitive to polar effects, and thus, in the absence of resonance interactions, this value can be considered as being proportional to steric effects (and any others that are not field or resonance effects):

$$\log(k/k_0) = E_{\rm S}^{\rm Taft} \tag{1}$$

In this scale, hydrogen is taken to have a reference value of  $E_8^{\text{Taft}} = 0$ . Another scale of values was put forward by Charton.<sup>3</sup> These are independent of any kinetical data, as they were derived from van der Waals radii. Meyer<sup>4</sup> has used the volume of the portion of the substituent that is within 0.3 nm of the reaction center, derived from molecular mechanics calculations, to quantify the steric effect. Still other quantities from experimental measurements to probe the steric effect have been introduced.

In conceptual DFT or density functional reactivity theory,<sup>5,6</sup> chemical concepts are identified with response functions of the energy of the system with respect to either the number of electrons N or the external potential (for an isolated system, this is the potential due to the nuclei) or both. The response functions could be linked with concepts readily known by

chemists but, in most cases, vaguely or empirically defined. Also, theoretical justification could be provided for a number of principles, such as Sanderson's principle of electronegativity equalization,<sup>7,8</sup> Pearson's hard and soft acids and bases,<sup>9-11</sup> maximum hardness principles, <sup>10–13</sup> and, more recently, minimum electrophilicity principle. <sup>14–19</sup> To the best of our knowledge, parameters or concepts related to the steric effect have not previously been considered in this field until very recently. Weisskopf<sup>20</sup> has related the steric effect to what he has called the "kinetic energy pressure" in atoms and molecules. This concept has also been related to the quantum contribution from the Pauli Exclusion Principle (Fermi hole),<sup>21</sup> and different implementations have been proposed within the wave function theory framework.<sup>22–27</sup> Recently, one of us introduced, <sup>28</sup> based on considerations within the framework of density functional theory, the steric energy as the Weizsäcker kinetic energy.<sup>29</sup> This analysis was based on an alternative energy partition scheme, where the total energy was decomposed into independent contributions from steric, electrostatic, and fermionic quantum effects with a hypothetical bosonic reference state.<sup>30</sup> The Weizsäcker kinetic energy can be expressed as:

$$E_{\rm S}[\rho(\vec{r})] \equiv T_{\rm W}[\rho(\vec{r})] = \frac{1}{8} \int \frac{|\nabla \rho(\vec{r})|^2}{\rho(\vec{r})} \, \mathrm{d}\vec{r} \tag{2}$$

where  $\rho(\vec{r})$  stands for the electron density of the system, and  $\nabla \rho(\vec{r})$  denotes the density gradient. Appealing properties such as exclusiveness, repulsiveness, and extensiveness have been revealed, <sup>28</sup> and applications to information theory and internal rotation barriers of simple molecular systems have been carried out. <sup>31–34</sup>

In this Article, we will compute the steric energy for a number of different molecules and perform the integration of this quantity over different regions in space (atomic and fragmental regions), to determine their contribution to the overall steric energy. Next, we will investigate the possible correlation between computed values of the steric energies and the experimental scale of the steric effect,  $E_{\rm S}^{\rm Taft}$ , put forward by Taft. Note that the steric values of Taft that we have used in this work are from esterification and hydrolysis of aliphatic and ortho-substituted benzoated esters.  $^{35,36}$ 

 $<sup>\</sup>begin{tabular}{ll} * Corresponding & author. & E-mail: & shubin@email.unc.edu & (S.L.); \\ fdeprof@vub.ac.be & (F.D.P.). & \\ \end{tabular}$ 

Vrije Universiteit Brussel.

<sup>\*</sup> University of North Carolina.

 $E'_{S}[\rho(\vec{r})]^{b}$  $E'_{S,X}[\rho(\vec{r})]^b$ molecule  $E_{\rm S}[\rho(\vec{r})]$  $E_{\rm S.H}[\rho(\vec{r})]$  $E_{S,X}[\rho(\vec{r})]$  $E_{S,H}[\rho(\vec{r})]^b$  $CH_4$ 32.699 0.517 30.629 32.699 0.517 30.629 184.335 0.633181.803 SiH<sub>4</sub> 184.114 0.484182.178 1003.149 0.467 1001.280 1002.945 0.626 1000.442 GeH<sub>4</sub> NH<sub>3</sub> 44.011 0.548 42.367 44.011 0.548 42.367 211.794 0.470 210.382 211.931 209.917  $PH_3$ 0.671 AsH<sub>3</sub> 1068.220 0.459 1066.842 1067.976 0.689 1065.909 57.441 0.599 56.242 57.441 0.599  $H_2O$ 56.242  $H_2S$ 241.594 0.461 240.673 241.709 0.708 240.293 H<sub>2</sub>Se 1135.136 0.451 1134.234 1135.022 0.753 1133.516 HF 72.876 72.125 72.876 0.751 72.125 0.751 273.616 273.165 273.669 272.927 HCl 0.4510.7421203.957 0.444 1203.513 1203.932 1203.116 HBr 0.817

TABLE 1: Total Steric Energy,  $E_S[\rho(\vec{r})]$ , of a Series of First to Third Row Hydrides  $H_nX$ , Together with the Steric Energy Associated with the Hydrogen Atoms,  $E_{S,H}[\rho(\vec{r})]$ , and with the Heavy Atoms,  $E_{S,X}[\rho(\vec{r})]$ , in These Molecules<sup>a</sup>

<sup>a</sup> All values are in au. <sup>b</sup> The geometrical parameters have been restricted to the values of the first row systems (e.g., the HCl and HBr systems are evaluated using the equilibrium geometry of the HF). For more details, see the text.

## 2. Computational Details

In a recent contribution by Liu, <sup>28</sup> the steric energy,  $E_S[\rho(\vec{r})]$ , was identified with the Weizsäcker kinetic energy,  $T_{\rm W}[\rho(\vec{r})]$ . In his original paper, global values of this quantity were obtained, and integration of this property to atoms or functional groups was not considered. To obtain local values for this property for the purpose of, for example, probing the steric effect in a certain region of a molecular system, integration could be performed over a particular region of the space to obtain its contribution to the global value. Next, by taking appropriate atomic domains defined in one or another way, one can define the atomic contributions to the steric energy,  $E_{S,i}[\rho(\vec{r})]$ ,

$$E_{S,i}[\rho(\vec{r})] = \frac{1}{8} \int_{\Omega_i} \frac{|\nabla \rho(\vec{r})|^2}{\rho(\vec{r})} d\vec{r}$$
 (3)

in such a way so that

$$E_{S}[\rho(\vec{r})] = \sum_{i=1}^{N} E_{S,i}[\rho(\vec{r})]$$
 (4)

In this work, we have used the fuzzy Voronoi polyhedra to define the atomic domains. It consists of the decomposition of the integral of eq 2 over the 3D space into a sum of integrations over single-atom components using a weight function,  $\omega_i(\vec{r})$ , for nucleus i in the system at every point of space  $\vec{r}$  in such a way that:

$$\sum_{i=1}^{N} \omega_i(\vec{r}) = 1 \tag{5}$$

In this scheme, the numerical integration of  $E_S[\rho(\vec{r})]$  is determinated as a sum of contributions  $E_{S,i}[\rho(\vec{r})]$ :

$$E_{\mathbf{S}}[\rho(\vec{r})] = \sum_{i=1}^{N} E_{\mathbf{S},i}[\rho(\vec{r})] \cong \frac{1}{8} \sum_{i=1}^{N} \int_{\Omega_{i}} \omega_{i}(\vec{r}) \frac{|\nabla \rho(\vec{r})|^{2}}{\rho(\vec{r})} d\vec{r}$$
(6)

where  $\omega_i(\vec{r})$  has the value 1 in the vicinity of its own nucleus, but vanishes in a continuous and well-behaved manner near any other nucleus. The atomic weights  $\omega_i(\vec{r})$  used in this work are derived from the fuzzy Voronoi polyhedra proposed by Becke,<sup>37</sup> tuned by the Bragg-Slater set of atomic radii and following Becke's suggestion to increase the radius of hydrogen to 0.35 Å.38,39 Such atomic definition has already been successfully applied for the calculation of overlap populations, bond orders, valences, 40 or in several molecular energy decomposition schemes. 41,42 Any other disjoint (Atoms in Molecules 43) or fuzzy (Hirshfeld<sup>44</sup>) decomposition of the 3D space could also be used, although we expect that the differences will be marginal and the tendencies of the steric energies will be retained, especially because in this work only covalent bonds have been considered. Note that, in previous studies, it has been shown that larger differences of atomic charges and bonds orders between different partition schemes have been found in ionic systems, but not for the covalent ones. 40,45,46 It should be remarked, however, that it was shown previously that, different from other quantifications, the present approach to probe the steric effect is able to uniquely define the contributions at the atomic, functional group, and entire molecular levels by adopting Bader's zeroflux boundary condition of atoms in molecules (AIM).<sup>43</sup> All geometries were optimized at the B3LYP/6-311+G(2d,2p) level; the steric energy and its atomic contributions were evaluated at the same level of theory, using the wave functions and densities obtained form the Gaussian 03 program. 47 Each atom has been integrated using Chebyshev's integration for the radial part (40 points) and Levedev's quadrature (146 points) for the angular part.<sup>48</sup> This level of methodology allows achieving accuracies of the order of  $10^{-5}$  au.<sup>49-51</sup>

# 3. Results and Discussion

In the first part, we have considered the steric energy for a number of first, second, and third row hydrides  $H_nX$ . Table 1 lists the total steric energy  $E_S[\rho(\vec{r})]$  (i.e., the Weizsäcker kinetic energy) of these compounds, together with the steric energy associated with the hydrogen atoms in the molecule,  $E_{S,H}[\rho(\vec{r})]$ , and with the heavy atoms,  $E_{S,X}[\rho(\vec{r})]$ . In general, as can be expected, the total steric energy of these compounds increases with the size of the central heavy atom. However, the  $E_{S,H}[\rho(\vec{r})]$ decreases upon increasing size of the central atom in the hydride; for example, the values of the  $E_{S,H}[\rho(\vec{r})]$  for the ammonia, phosphine, and arsine are 0.548, 0.470, and 0.459 au, respectively. This fact indicates that apparently the hydrogen atoms are contributing less to the steric energy in the heavier hydrides than in the lighter ones, which seems counterintuitive, as one would expect the hydrogens to become more sterically hindered. Insight into this can be obtained by considering the different X-H bond distances in the different compounds. It indeed appears that the decreasing hydrogen atom contribution to the

**Figure 1.** Atomic and functional group steric energies for a few exemplary molecules. See text for discussion. All values are in au.

steric energy is dominated by the fact the bond distance increases with increasing size of the atom X in the hydride. Consequently, we have computed the steric energies for the different hydrides fixing the geometrical parameters to the values for the corresponding first row hydride (e.g., SiH<sub>4</sub> and GeH<sub>4</sub> are evaluated using the equilibrium geometry of CH<sub>4</sub>, whereas PH<sub>3</sub> and AsH<sub>3</sub> are evaluated using the equilibrium geometry of NH<sub>3</sub>). As can now be seen, the steric energies of the hydrogen atoms now increase considerably, and, as could be expected, the total steric energy remained approximately the same. For instance, the values of the  $E_{S,H}[\rho(\vec{r})]$  for the ammonia, phospine, and arsine now become 0.548, 0.671, and 0.689 au, respectively, reproducing the expected chemical tendency of the steric effect. This indicates that next to the intrinsic steric effect also the distance of the substituent to the relevant area on which a steric effect is measured plays an important role.

Next, we have considered the change in the steric effect on substituents in a number of typical compounds, such as the staggered and eclipsed conformations of ethane, the axial versus the equatorial positions in the chair form of cyclohexane, and the axial versus the equatorial positions in the trigonal pyramidal molecule PF<sub>5</sub>, whose results are given in Figure 1. The calculations confirm that the steric contribution of the hydrogen atoms and methyl groups in the staggered conformation of ethane is smaller than that in the eclipsed conformation, consistent with the picture from the global results reported earlier.32,33 It is also confirmed that the axial hydrogen atoms in the chair conformation of cyclohexane are contributing less to the steric energy than are the equatorial ones. Finally, it is also shown that the fluorine atoms in the equatorial position of PF<sub>5</sub> are contributing less to the steric energy than are the axial ones.

Then, one can conclude that the atomic and group steric energies can be used to discern the steric effect between atoms or groups of a same molecule. However, it is important to remark that they are only contributions to the overall steric energy. For instance, we cannot use the difference of the atomic steric energy between the axial and equatorial hydrogens of the chair form of cyclohexane (0.010 au, the result of 0.500 minus 0.490) and the difference of the atomic steric energy between the staggered and eclipsed conformations of ethane (0.001 au, the result of 0.511 minus 0.510) to affirm that the steric effect is more important for the cyclohexane than for the ethane. Thus,

TABLE 2: Total Steric Energy,  $E_S[\rho(\vec{r})]$ , of a Series of Compounds CH<sub>3</sub>-X, Together with the Steric Energy Associated with the Methyl Groups,  $E_{S,CH3}[\rho(\vec{r})]$ , for These Molecules<sup>a</sup>

X	$E_{\rm S}[\rho(\vec{r})]$	$E_{\mathrm{S,CH_3}}[\rho(\vec{r})]$	$d_{\mathrm{X-CH_3}}$	$E_{ m S}^{{ m Taft}\;b}$
Н	32.699	32.181	1.088	0.000
F	104.568	32.597	1.394	-0.460
CN	105.739	32.053	1.457	-0.510
OH	89.045	32.333	1.424	-0.550
OMe	120.577	32.282	1.413	-0.550
$NH_2$	75.544	32.202	1.466	-0.610
Me	64.213	32.107	1.528	-1.240
Et	95.690	32.084	1.529	-1.310
Pr	127.164	32.080	1.529	-1.600
<i>i</i> -Pr	127.134	32.060	1.532	-1.710
cyclohexyl	220.376	32.054	1.529	-2.030
<i>i</i> -Bu	158.595	32.056	1.533	-2.170
s-Bu	158.595	32.056	1.532	-2.370
t-Bu	158.555	32.038	1.537	-2.780
neopentyl	190.001	32.032	1.538	-2.980
Et <sub>3</sub> C	252.902	31.999	1.539	-5.040
Ph <sub>3</sub> C	619.626	31.980	1.557	-5.920

 $^a$  All values are in au, except for the distances, which are in angstroms. Also listed are Taft's steric effect values for the groups X,  $E_{\rm S}^{\rm Taft}$ .  $^b$  From refs 35 and 36.

the atomic and group functional steric energies can be used to contrast systems with similar geometries. In contrast, the total steric energy does not present these limitations, but it cannot be applied to predict the regioselectivity of the systems at difference to the atomic and group functional steric energies.

After checking that the partition of the Weizsäcker kinetic energy in atomic and functional group contributions can be used as a useful tool to evaluate the steric effect, we will analyze the possibility to establish a link between these theoretical values and the experimental ones. For that purpose, we will use the experimental constants obtained from linear free energy relationships mentioned in the Introduction, more specifically,  $E_{\rm S}^{\rm Taft}$ values.<sup>35,36</sup> Consequently, we have considered a number of first row and second row CH3-X type compounds, where we have investigated the relationship between the contribution of the CH<sub>3</sub> functional groups to the steric energy of the molecule and the  $E_{\rm S}^{\rm Taft}$  values of the groups X. Theoretical and experimental steric effect results are tabulated in Table 2, and the correlation of the two is plotted in Figure 2. From the figure, it is seen that there exists a reasonable linear correlation between theoretical and experimental scales of the steric effect for this set of first and second row compounds:

$$E_{\rm S}[\rho(\vec{r})] = -70.063E_{\rm S}^{\rm Taft} + 27.702$$
, with  $R^2 = 0.749(7)$ 

If the data point for the  $Ph_3C$  group is eliminated from the correlation analysis, the  $R^2$  becomes 0.754; eliminating both values for  $Ph_3C$  and  $Et_3C$  groups yields an  $R^2$  of 0.647.

The slope is negative, indicating that a larger value of  $E_S[\rho(\vec{r})]$  corresponds to a more negative experimental  $E_S^{Taft}$  value and thus a larger steric effect. We notice that the extensive property of the theoretical scale of eq 2 is dictated by its first-order density scaling homogeneity, but its genuine linear correlation with the experimental Taft's scaling is something theoretically unpredicted. Given that the experimental scale of the steric effect was based on rate constants for the acid-catalyzed hydrolysis of esters in aqueous acetone and the theoretical results are from gas-phase calculations at 0 K, the above correlation can be seen

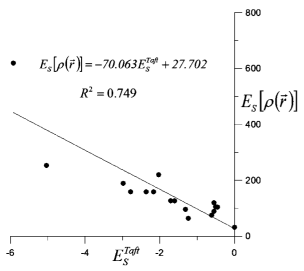


Figure 2. Linear relationship between the theoretical steric energy from eq 2 and Taft's experimental steric parameters based on eq 1 for 17 species listed in Table 2.

as the affirmative ratification that eq 2 is an intrinsic measure of the steric effect of a molecular system.

Moreover, in Table 2, we can also analyze the steric effect contribution of the CH<sub>3</sub> group,  $E_{S,CH_3}[\rho(\vec{r})]$ . For instance, the change of the hydrogen in the CH<sub>4</sub> for the F, OH, OMe, and NH<sub>2</sub> groups with more negative  $E_S^{Taft}$  values, that is, more steric effect, provokes an increase of  $E_{S,CH_3}[\rho(\vec{r})]$ . However, the remaining groups show smaller values of  $E_{S,CH_2}[\rho(\vec{r})]$  than the CH<sub>4</sub>, indicating that the tendencies between  $E_{S,CH_3}[\rho(\vec{r})]$  and  $E_S^{Taft}$ can be opposite. Taking into account the conclusions made in the first part of this work, we know about the relevance of the geometry of the molecules in the steric energy. So, to obtain a good description of the steric effect at least, it is necessary to consider two parameters,  $E_{S,CH_3}[\rho(\vec{r})]$  and  $d_{X-CH_3}$ . To prove this, a multilinear regression has been done using the  $E_{S,CH_3}[\rho(\vec{r})]$ ,  $d_{\rm X-CH_3}$ , and  $E_{\rm S}^{\rm Taft}$  values for 11 molecules of the Table 2. In the multilinear regression, we only consider the C-X sytems, where X is a carbon atom, to minimize the large geometry dependency of the group atomic steric energy, and we obtain the following expression:

$$E_{\rm S}^{\rm Taft} = 28.347 E_{\rm S, CH_3}[\rho(\vec{r})] - 56.740 d_{\rm H_3C-X} - 824.076$$
 (8)

To validate the utility of this equation, in Figure 3 we plot the linear regression between the experimental scale from Taft's steric parameters and the fitted theoretical steric energy from eq 8,  $E_S^{\text{theor}}[\rho(\vec{r}), d_{H_3C-X}]$ :

$$E_{\rm S}^{\rm theor}[\rho(\vec{r}), d_{\rm H_3C-X}] = 0.937 E_{\rm S}^{\rm Taft} - 0.168 \text{ with } R^2 = 0.937$$
(9)

When eliminating the data for the  $X = Ph_3C$  system from the correlation analysis, the  $R^2$  becomes 0.871; additionally eliminating the data for  $X = Et_3C$  systems yields  $R^2 = 0.908$ .

This analysis shows that the steric effect as given by  $T_{\rm W}[\rho(\vec{r})]$ and computed at both the global and the atomic or functional group level can be used as an estimate of the intrinsic steric effect, as, for example, quantified by the  $E_S^{Taft}$  values. It constitutes a first step to introduce the global steric effect and

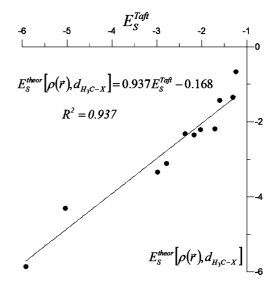


Figure 3. Comparison of the experimental scale from Taft's steric parameters and the fitted theoretical steric energy from eq 8 using  $CH_3$ group steric energy and the C-X bond distance (where X is a carbon atom). See the text for more details.

its atomic and functional group contributions in the applications of DFT-based reactivity indices to chemical problems.

### 4. Conclusions

In summary, based on a recent quantification of the steric effect in the framework of density functional theory where the total electronic energy was proposed to be decomposed into contributions from three independent effects, steric, electrostatic, and fermionic quantum, contributions of the effect at the atomic and functional groups levels are considered in this work. Reasonable linear relationships between theoretical and experimental scales at both group and entire molecular levels have been discovered, providing an additional piece of evidence affirming that the newly proposed definition serves as an intrinsic measurement of the steric effect for molecular systems. In addition, we have shown that the steric energy,  $E_S[\rho(\vec{r})]$ , can be a useful tool to measure the global steric effect between different molecules. However, if we want to compare the atomic and functional group contributions to the  $E_S[\rho(\vec{r})]$  between different molecules, it is essential to consider the contribution from the molecular geometry (distances and angles) as well.

Acknowledgment. We are grateful to Robert G. Parr and Lee G. Pedersen of the University of North Carolina and Paul W. Ayers of McMaster University, Canada, for their valuable comments and suggestions. M.T.-S. thanks the European Community for financial help through the postdoctoral grant MEIF-CT-2006-025362. F.D.P. acknowledges funding from the Free University of Brussels (VUB) and from the Fund for Scientific Research-Flanders (Belgium) FWO.

### References and Notes

- (1) IUPAC Compendium of Chemical Technology, 2nd ed.; compiled by McNaught, A. D., Wilkinson, A.; Blackwell Science: Cambridge, MA,
- (2) Taft, R.W., Jr. In Steric Effect in Organic Chemistry; Newman, M. S., Ed.; Wiley: New York, 1956; pp 556-675.
  - (3) Charton, M. J. Am. Chem. Soc. 1975, 97, 1552
  - (4) Meyer, A. Y. J. Chem. Soc., Perkin Trans. 2 1986, 1567.
- (5) Parr, R. G.; Yang, W. Density Functional Theory of Atoms and Molecules; Oxford University Press: Oxford, 1989.
- (6) Geerlings, P.; De Proft, F.; Langenaeker, W. Chem. Rev. 2003, 103, 1793.

- (7) Sanderson, R. T. Science 1951, 114, 670.
- (8) Sanderson, R. T. Chemical Bonds and Bond Energy; Academic Press: New York, 1976.
  - (9) Pearson, R. G. J. Am. Chem. Soc. 1963, 85, 3533.
  - (10) Pearson, R. G. J. Chem. Educ. 1987, 64, 561.
- (11) Pearson, R. G. Chemical Hardness: Applications from Molecules to Solids; Wiley-VCH: Oxford, 1997.
  - (12) Pearson, R. G. J. Chem. Educ. 1999, 76, 267.
  - (13) Parr, R. G.; Chattaraj, P. K. J. Am. Chem. Soc. 1991, 113, 1854.
- (14) Parr, R. G.; Von Szentpály, L.; Liu, S. B. J. Am. Chem. Soc. 1999, 121, 1922.
- (15) Chattaraj, P. K.; Sarkar, U.; Roy, D. R. Chem. Rev. 2006, 106, 2065
- (16) Noorizadeh, S.; Maihami, H. J. Mol. Struct. (THEOCHEM) 2006, 763, 133.
  - (17) Noorizadeh, S. Chin. J. Chem. 2007, 25, 1439.
  - (18) Noorizadeh, S. J. Phys. Org. Chem. 2007, 20, 514.
- (19) Xia, Y.; Yin, D.; Rong, C.; Xu, Q.; Yin, D.; Liu, S. B. J. Phys. Chem. A 2008, 112, 9970.
  - (20) Weisskopf, V. F. Science 1975, 187, 605.
  - (21) Luken, W. L.; Beratan, D. N. Theor. Chem. Acc. 1982, 61, 1432.
  - (22) Badenhoop, J. K.; Weinhold, F. J. Chem. Phys. 1997, 107, 5406.
  - (23) Schreiner, P. R. Angew. Chem., Int. Ed. 2002, 41, 3579.
- (24) Bickelhaupt, F. M.; Baerends, E. J. Angew. Chem., Int. Ed. 2003, 42, 4183.
  - (25) Weinhold, F. Angew. Chem., Int. Ed. 2003, 42, 4188.
- (26) Mo, Y.; Wu, W.; Song, L.; Lin, M.; Zhang, Q.; Gao, J. Angew. Chem., Int. Ed. 2004, 43, 1986.
  - (27) Mo, Y.; Gao, J. Acc. Chem. Res. 2007, 40, 113.
  - (28) Liu, S. B. J. Chem. Phys. 2007, 126, 244103.
    (29) von Weizsäcker, C. F. Z. Phys. 1935, 96, 431.

  - (30) Nagy, A. Chem. Phys. Lett. 2007, 449, 212.
  - (31) Liu, S. B. J. Chem. Phys. 2007, 126, 191107
  - (32) Liu, S. B.; Govind, N. J. Phys. Chem. A 2008, 112, 6690.
- (33) Liu, S. B.; Govind, N.; Pedersen, L. G. J. Chem. Phys. 2008, 129, 094104
- (34) Nagy, A.; Liu, S. B. Phys. Lett. A 2008, 372, 1654.
- (35) March, J. Advanced Organic Chemistry; John Wiley & Sons: New York, 1992; p 295.
  - (36) Taft, R. W., Jr. J. Am. Chem. Soc. 1952, 74, 2729.

- (37) Becke, A. D. J. Chem. Phys. 1988, 88, 2547.
- (38) Slater, J. C. J. Chem. Phys. 1964, 41, 3199.
- (39) Slater, J. C. Quantum Theory of Molecules and Solids; McGraw-Hill: New York, 1965; Vol. 2.
  - (40) Mayer, I.; Salvador, P. Chem. Phys. Lett. 2004, 383, 368.
- (41) Salvador, P.; Mayer, I. J. Chem. Phys. 2004, 120, 5046.
- (42) Salvador, P.; Mayer, I. J. Chem. Phys. 2007, 126, 234113.
- (43) Bader, R. F. W. Atoms in Molecules A Quantum Theory; Oxford University Press: Oxford, 1990.
  - (44) Hirshfeld, F. L. Theor. Chrm. Acc. 1977, 44, 129.
- (45) Matito, E.; Poater, J.; Solà, M.; Duran, M.; Salvador, P. J. Phys. Chem. A 2005, 109, 9904.
- (46) Matito, E.; Solà, M.; Salvador, P.; Duran, M. Faraday Discuss. 2007, 135, 325.
- (47) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Montgomery, J. A., Jr.; Vreven, T.; Kudin, K. N.; Burant, J. C.; Millam, J. M.; Iyengar, S. S.; Tomasi, J.; Barone, V.; Mennucci, B.; Cossi, M.; Scalmani, G.; Rega, N.; Petersson, G. A.; Nakatsuji, H.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Klene, M.; Li, X.; Knox, J. E.; Hratchian, H. P.; Cross, J. B.; Adamo, C.; Jaramillo, J.; Gomperts, R.; Stratmann, R. E.; Yazyev, O.; Austin, A. J.; Cammi, R.; Pomelli, C.; Ochterski, J. W.; Ayala, P. Y.; Morokuma, K.; Voth, G. A.; Salvador, P.; Dannenberg, J. J.; Žakrzewski, V. G.; Dapprich, S.; Daniels, A. D.; Strain, M. C.; Farkas, O.; Malick, D. K.; Rabuck, A. D.; Raghavachari, K.; Foresman, J. B.; Ortiz, J. V.; Cui, Q.; Baboul, A. G.; Clifford, S.; Cioslowski, J.; Stefanov, B. B.; Liu, G.; Liashenko, A.; Piskorz, P.; Komaromi, I.; Martin, R. L.; Fox, D. J.; Keith, T.; Al-Laham, M. A.; Peng, C. Y.; Nanayakkara, A.; Challacombe, M.; Gill, P. M. W.; Johnson, B.; Chen, W.; Wong, M. W.; Gonzalez, C.; Pople, J. A. Gaussian 03, revision B.04; Gaussian, Inc.: Pittsburgh, PA, 2003.
  - (48) Levedev, V. I.; Laikov, D. N. Dokl. Math. 1999, 59, 477.
- (49) Torrent-Sucarrat, M.; Salvador, P.; Geerlings, P.; Solà, M. J. Comput. Chem. 2007, 28, 574.
- (50) Torrent-Sucarrat, M.; Salvador, P.; Solà, M.; Geerlings, P. J. Comput. Chem. 2008, 29, 1064.
- (51) Torrent-Sucarrat, M.; De Proft, F.; Geerlings, P.; Ayers, P. W. Chem.-Eur. J. 2008, 14, 8652.

JP8096583