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Quantifying aromaticity with electron delocalisation measures*

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Aromaticity cannot be measured directly by any physical or chemical experiment because it is not a well-defined magnitude. Its quantification is done indirectly from the measure of different properties that are usually found in aromatic compounds such as bond length equalisation, energetic stabilisation, and particular magnetic behaviour associated with induced ring currents. These properties have been used to set up the myriad of structural-, energetic-, and magnetic-based indices of aromaticity known to date. The cyclic delocalisation of mobile electrons in two or three dimensions is probably one of the key aspects that characterise aromatic compounds. However, it has not been until the last decade that electron delocalisation measures have been widely employed to quantify aromaticity. Some of these new indicators of aromaticity such as the PDI, FLU, $I_{\rm NG}$, and $I_{\rm NB}$ were defined in our group. In this paper, we review the different existing descriptors of aromaticity that are based on electron delocalisation properties, we compare their performance with indices based on other properties, and we summarise a number of applications of electronic-based indices for the analysis of aromaticity in interesting chemical problems.

1. Introduction

The study of aromatic species goes back to 1825 when Michael Faraday obtained benzene by distillation and named it "dicarburet of hydrogen". In this work, Faraday found that the empirical formula of benzene was CH. He already noted that "dicarburet of hydrogen" was much less reactive than "monocarburet of hydrogen" (trans-2-butene). Such a decreased reactivity was considered to be an experimental characteristic of aromatic compounds ever since. In 1834, Mitscherlich determined that the compound synthesised by Faraday had the molecular formula C₆H₆.² Since it was obtained from the distillation of benzoic acid (from gum benzoin) and lime, he named the compound as benzin, which became benzene when translated into English. In 1865, a century and a half ago, August Kekulé⁴ proposed the molecular structure of benzene consisting of a sixmembered ring (6-MR) of carbon atoms with alternating single and double bonds. Pyridine, the first heteroaromatic compound, was synthesised by Thomas Anderson⁵ in 1868 through studies on the distillation of bone oil and other animal matter. In 1911, Willstätter and Waser⁶ synthesised cyclooctatetraene, an

eight-membered carbon ring with alternating single and double bonds that had a reactivity very different from benzene and was classified as antiaromatic. The first synthesised inorganic heteroaromatic compound was borazine, B₃H₆N₃, obtained in 1926 by a reaction of diborane with ammonia. In 1954, Doering and Knox8 prepared the tropylium cation, C₇H₇⁺, which was considered the first verification of the Hückel rule.9 Four years later, Winstein introduced the homoaromaticity concept while studying the 3-bicyclo[3.1.0]hexyl cation. Three years later, the synthesis of the first organic derivatives of closo-dodecaborate and closodecaborate by the group of Muetterties¹¹ was the beginning of closo-borane chemistry and three-dimensional aromaticity, the type of aromaticity that characterises fullerenes.12 The identification 13,14 of the planar triplet ground states of C₅H₅ and C₅Cl₅ in the late 60's provided experimental support for the existence of triplet aromaticity as predicted by Baird. 15 In 1982 Roper et al. 16 synthesised the first metallabenzene, an osmabenzene, thus initiating a new group of aromatic species, the so-called metalloaromatic compounds.¹⁷ Interestingly, metallabenzene species were proposed first by Thorn and Hoffmann from theoretical calculations three years before. 18 More recently, in 2001 Boldyrev, Wang, and coworkers¹⁹ detected a series of bimetallic clusters containing Al₄²⁻, the first all-metal aromatic cluster known, face-capped by an M⁺ cation (M = Li, Na, Cu). The same group detected Ta₃O₃⁻, the first discovered metallic cluster showing δ-aromaticity.²⁰ In 2003, Herges et al. synthesised the first Möbius aromatic hydrocarbon.21 This kind of aromatic species were already predicted by Heilbronner forty years before on purely theoretical grounds.22

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 $[\]dagger$ This paper is dedicated to the memory of our colleague Prof. Dr Tom Ziegler who passed away on 24th March 2015

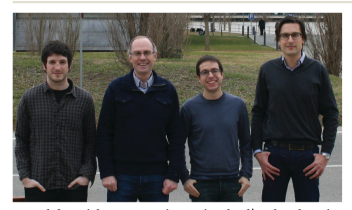
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Despite the nearly two centuries of intense study and continuous progress described in the previous paragraph, the interest in aromaticity has not decreased and still stimulates the creativity of a number of contemporary chemists. Considered like a chemical unicorn by Frenking and Krapp,²³ aromaticity is not a property directly observable and it lacks a well-founded physical basis. Therefore, its definition and quantification remain elusive. Chen and Schleyer²⁴ defined aromaticity as "a manifestation of electron delocalisation in closed circuits, either in two or three dimensions." Electron delocalisation is without doubt one of the key aspects of aromatic compounds. Since 1932 when Pauling^{25,26} introduced the concept of resonance, it is well established that the delocalisation of electrons in a molecule can stabilise it. Another relevant feature of aromatic compounds is their symmetry. Despite not all aromatic compounds are symmetric and not all symmetric cyclic compounds are aromatic, it is generally the case that the most archetypal aromatic compounds are highly symmetric and possess degenerate highest-occupied molecular orbitals that are fully occupied resulting in a closed-shell structure or have the samespin half-filled electronic structure. This is the case of benzene, but also of triplets $C_5H_5^+$, C_{60}^{10+} , Al_4^{2-} or *closo* borane clusters like B₆H₆²⁻. The closed-shell or the same-spin half-filled electronic structure is the origin of several rules of aromaticity such as the 4n+ 2 Hückel, 9 4n Baird, 15 2n + 2 Wade-Mingos, 27,28 2(n + 1) 2 $Hirsch^{29}$ or the $2n^2 + 2n + 1^{30}$ rules. This particular electronic

structure of aromatic species explains their substantial energetic stabilisation and, consequently, their low reactivity. Moreover, it results in a variety of unusual chemical and physical properties, including a tendency toward bond equalisation, unusual reactivity, and characteristic spectroscopic and magnetic features.

It is worth noting that the bond length equalisation observed in benzene and derivatives is enforced by the σ -electrons and not by the π -electron system. The latter is distortive and favours the D_{3h} structure of benzene over the D_{6h} one. This somewhat paradoxical observation was suggested first by Longuet-Higgins and Salem in 1959. 31 Berry 22 used this suggestion to account for the observed increased frequency of the b₂₁₁ Kekulé vibrational mode when going from the ground ${}^{1}A_{1g}$ to the first ${}^{1}B_{2u}$ excited state. The same idea was reinforced later on by the work of Haas and Zilberg, 33,34 and especially by that of Hiberty, Shaik, and co-workers^{35,36} and others^{37,38} Finally, more recently Pierrefixe and Bickelhaupt 39-41 showed that the regular geometry of benzene is a consequence of how σ and π overlaps depend on bond distances. In aromatic annulenes the authors confirmed that bond equalisation is due to the σ electrons whereas the π -electron system favours double-bond localisation.

The quantification of physicochemical properties that reflect some manifestation of the aromatic character of molecules is used to evaluate their global or local (e.g., individual



From left to right: Ferran Feixas, Miquel Solà, Eduard Matito and Jordi Poater

Ferran Feixas (1983) obtained his PhD from the Universitat de Girona (UdG) in 2011 for a thesis on the analysis of chemical bonding and aromaticity under the supervision of Prof. Solà, Dr Poater, and Dr Matito. He was awarded a Beatriu de Pinós fellowship for a two-year postdoctoral position at the group of Prof. McCammon at the University of California, San Diego (UCSD) to work in the field of protein dynamics and drug design. In 2014, he joined the group of Prof. Swart at the Institute of Computational Chemistry and Catalysis (IQCC, UdG) as a Beatriu de Pinós postdoctoral researcher. He has co-authored more than 30 publications.

Eduard Matito (1980) obtained his PhD (Univ. Girona, 2006) working on the role of electron correlation in chemical bonding descriptors and the design of new aromaticity indices with Profs. Solà and Duran. After graduation, he worked as a Marie-Curie

postdoctoral fellow in the group of Prof. Cioslowski (Univ. Szczecin) and as a postdoc in the groups of Profs. Christiansen (Univ. Aarhus, Denmark) and Ugalde (Euskal Kerriko Unibertsitatea, Basque Country), completing a postdoctoral training of five years. In 2012 he joined the UdG as a junior group leader at the Institute of Computational Chemistry and Catalysis. He has co-authored more than 60 publications.

Jordi Poater (1977) obtained his PhD in Chemistry in 2003 from the UdG with the analysis of molecular structure and aromaticity by means of electronic-based criteria. In 2005 he was awarded a Marie Curie postdoctoral fellowship at the group of Prof. Dr F. M. Bickelhaupt at the Vrije Universiteit Amsterdam (VUA). And in 2007 he was awarded a Ramón y Cajal position at the Institute of Computational Chemistry and Catalysis of the UdG. Right now Dr Poater holds a position as a senior associate researcher at the VUA. He has co-authored more than 90 publications.

Miquel Solà (1964) obtained his PhD from the Autonomous University of Barcelona in 1991. In 1993 he moved to the UdG as an assistant researcher and was appointed assistant professor in 1997. In 2001, he received the Catalan Distinction for the Promotion of University Research (young scientist category). Since 2003, he holds a full professor position in the UdG. He was awarded with the ICREA Academia Prize in the 2009 and 2014 editions. In 2013 he received the Physical Chemistry prize of the Spanish Royal Society of Chemistry. He serves in the Editorial Board of several journals and has co-authored more than 300 papers.

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rings in a polycyclic arene) aromatic character. ^{42–44} This leads to the countless existing measures of aromaticity based on the structural, ^{45,46} magnetic, ^{24,47} energetic, ⁴⁸ and electronic ⁴⁹ properties of molecules. These indicators provide indirect measures of aromaticity that, to some extent, are somewhat arbitrary. Moreover, for a series of compounds different descriptors of aromaticity do not provide the same aromaticity ordering and some descriptors fail to correctly quantify certain changes in aromaticity in particular examples. ⁵⁰ For this reason, it is widely accepted that the concept of aromaticity should be analysed by employing a set of aromaticity descriptors. ^{24,51,52}

The importance of electron delocalisation in aromatic species is universally recognised. It is thus reasonable to employ electron delocalisation as a tool to construct new aromaticity indicators. The problem is that electron delocalisation like aromaticity is not observable and, therefore, there is no unique way to measure it. Moreover, when devising new aromaticity descriptors three steps should be followed: development, assessment, and application. In the next sections, we describe firstly the most usual ways to quantify electron delocalisation; second, we discuss how these measures can be used to define indicators of aromaticity; third, we examine how electronic-based indices compare with other existing descriptors; and finally, we show some applications of these electronic descriptors of aromaticity carried out in our research group.

2. How can electronic delocalisation be measured?

Electron localisation/delocalisation is central in several fundamental chemical phenomena such as conjugation, hyperconjugation, and aromaticity that are important to explain the structure, stability, magnetic properties, and reactivity of many molecules. Several tools have been developed to quantify delocalisation in a molecule and to provide new insights into chemical bonding. These descriptors of delocalisation have been reviewed in several studies. ^{49,53–55} We just briefly mention here some of the most important and describe in more detail those utilised for the definition of aromaticity indicators.

Descriptors of electron localisation/delocalisation can be broadly grouped into three classes, namely, those computed from the wavefunction, those constructed directly from the electron density and its derivatives, and those that are derived from first- and higher-order density matrices. Among the first group of electron delocalisation indicators, we can mention the use of weights of resonance structures and energies in valence bond theory. 56,57 Also belonging to this group are the measures of delocalisation obtained with the block-localised wavefunction (BLW) method of Mo and coworkers. 58,59 In addition, in the framework of the molecular orbital (MO) theory, different techniques are available for the localisation of MOs to find the regions where electron pairs are located. To date the natural bond orbital (NBO) analysis of Weinhold et al. 60,61 has been one of the most used methods for localising bonds and lone pairs. The importance of electron delocalisation is assessed in NBO

analyses by approximated second-order perturbative expressions. Another similar tool used for obtaining patterns of chemical bonding is the adaptive natural density partitioning (AdNDP) method developed by Zubarev and Boldyrev. 62,63 AdNDP represents the electronic structure in terms of n-centre 2-electron (nc-2e) bonds. Starting from n=1, AdNDP recovers Lewis bonding elements (1c-2e and 2c-2e objects) and delocalised bonding elements (for n>2), which can be associated with the concepts of delocalisation and aromaticity. An even more recent method is the orbital localisation procedure based on the electron localisation function (ELF-LOC) of Alcoba, Tiznado, and coworkers. $^{64-66}$ This procedure localises the molecular orbitals in regions that have the highest probability for finding a pair of electrons, providing also a chemical bonding description in terms of nc-2e bonds.

In the second group of electron delocalisation descriptors, we can refer to the Laplacian of the electron density $(\nabla^2 \rho(\mathbf{r}))^{67,68}$ the ellipticity along the bond path $(\varepsilon = \lambda_1/\lambda_2 - 1)^{69}$ the noncovalent interaction index (NCI), 70 the inhomogeneity measures of the electron density,⁷¹ the source function,^{72,73} the single exponential decay detector, 74 and a recent electron delocalisation index obtained from electron population analysis.⁷⁵ The electron delocalisation indicators collected in the third group are probably the most abundant. The reason is that the location of electron pairs involves two spatial coordinates and, therefore, methods based on functions of two (or more) positions such as the first-order density matrix and the two-electron density or pair density are more suitable to analyse electron localisation and delocalisation. This group includes the localised orbital locator, 76 the electron delocalisation range function,⁷⁷ the parity function,⁷⁸ the analytical method by Proynov to calculate the population of effectively unpaired electrons, 79 the Fermi hole density maps, 80,81 the domain-averaged Fermi holes, 82,83 the Laplacian of the exchange-correlation density,84 the electron localisation function (ELF), 85-88 and the electron localisability indicator. 89,90 Also belonging to this group are the methods for computing the probability of finding a certain number of electrons in a given volume. 91-93 And finally, although not derived directly from density matrices, the analysis of the linear response kernel (LRK), 94-96 a function of two position variables, could also be included in this third group of electron localisation/delocalisation descriptors.

Because of its importance in the definition of aromaticity descriptors, we will refer with more detail to the electron sharing indices (ESIs). These ESIs are electron localisation/delocalisation descriptors that can also be classified into the third group. They are defined from the spinless two-electron density or pair density, $\gamma(\vec{r}_1, \vec{r}_2)$. This function can be interpreted as the probability density of having simultaneously two electrons at positions \vec{r}_1 and \vec{r}_2 , regardless of the position of the other N-2 electrons. It can be separated into an uncorrelated part and a part that collects all exchange and correlation effects,

$$\gamma(\vec{r}_1, \vec{r}_2) = \rho(\vec{r}_1)\rho(\vec{r}_2) + \gamma_{XC}(\vec{r}_1, \vec{r}_2) \tag{1}$$

The uncorrelated part of the pair density is given by the product $\rho(\vec{r}_1)\rho(\vec{r}_2)$ and provides the probability of finding simultaneously

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two independent electrons at positions \vec{r}_1 and \vec{r}_2 , irrespective of the position of the other N-1 electrons. The exchange–correlation density, $\gamma_{\rm XC}(\vec{r}_1,\vec{r}_2)$, which is the difference between $\gamma(\vec{r}_1,\vec{r}_2)$ and $\rho(\vec{r}_1)\rho(\vec{r}_2)$, gives a measure of the degree to which density is excluded at \vec{r}_2 because of the presence of an electron at \vec{r}_1 . The localisation and delocalisation indices (LIs and DIs) were defined by Bader and coworkers^{97,98} from the double integration of the exchange–correlation density over two atomic domains:

$$\lambda(A) = -\int_{A} \int_{A} \gamma_{\text{XC}}(\vec{r}_1, \vec{r}_2) d\vec{r}_1 d\vec{r}_2$$
 (2)

$$\delta(A, B) = -2 \int_{B} \int_{A} \gamma_{XC}(\vec{r}_{1}, \vec{r}_{2}) d\vec{r}_{1} d\vec{r}_{2}$$
 (3)

The DI or 2c-ESI, $\delta(A,B)$, gives a quantitative measure of the number of electron pairs delocalised or shared between atomic basins A and B, whereas the LI, $\lambda(A)$, is a measure of the average number of electrons localised on atom A. The double integration over the whole space of the exchange–correlation density gives -N electrons, and, therefore, the following sum rule is followed:

$$N(A) = \lambda(A) + \frac{1}{2} \sum_{B \neq A} \delta(A, B) \tag{4}$$

Eqn (2) and (3) can be applied at any level of theory, provided that the first and second-order density functions are known. At the Hartree–Fock (HF) level or with the density functional theory (DFT) approach (in this case we use the non-interacting wavefunction derived from Kohn–Sham orbitals)⁹⁹ the expressions for the LI and DI are,

$$\lambda(A) = \sum_{ij} S_{ij}^{2}(A) \tag{5}$$

$$\delta(A,B) = 2\sum_{ij} S_{ij}(A)S_{ij}(B) \tag{6}$$

where the summations run over all the pairs of occupied spin molecular orbitals (SMOs) of the molecule. The $S_{ij}(A)$ terms in eqn (5) and (6), needed to compute LIs and DIs, are the overlaps between SMOs integrated within the basin of atom A and require integration over atomic domains. In the Quantum Theory of Atoms-in-Molecules (QTAIM) these domains are the *atomic basins* defined as the regions in real space bound by zero-flux surfaces in $\rho(\vec{r})$ or by infinity. 67,68,100,101 However, other atomic partitions like the Mulliken-like partitioning in the Hilbert space spanned by the basis functions, the fuzzy-atom approach, 102 or the topological fuzzy Voronoi cell (TFVC) 103 can be used for the integration. The 2c-ESIs of organic covalent bonds give qualitatively the same results regardless of the atomic partition used. 104,105

The definition of the ESI can be generalised to analyse multicentre delocalisation or sharing of electrons. It is possible to define a multicentre DI or Mc-ESI¹⁰⁶ between the M centres A_1 and A_M :

$$I_{A_1,\ldots,A_M} = \sum_{i_1\ldots i_M} n_{i_1}\cdots n_{i_M} S_{i_1i_2}(A_1) S_{i_2i_3}(A_2)\cdots S_{i_Mi_1}(A_M) \quad (7)$$

where n_i are the occupancies of the natural orbitals. This formulation assumes that the electron sharing occurs only

between neighbouring atoms. Bultinck and coworkers¹⁰⁷ suggested another definition that takes into account all possible permutations of the M atoms, the multicentre index,

$$MCI(A_1,...,A_M) = \frac{1}{2M} \sum_{P(A_1,...,A_M)} I_{A_1,...,A_M}$$
 (8)

where P represents all possible permutations between centres A_1 and A_M . Indeed, in a HF calculation, the MCI represents the M-centre moment of the M-variate probability distribution restricting one electron in each atomic region; i.e., for a given molecule the MCI reflects the simultaneous electron sharing between all centres, regardless of the position of the atoms in the molecule. This interpretation is only valid in general if the M-order reduced density matrix (M-RDM)—which carries a large computational cost-is used for the calculation of the MCI. In practice, the calculation of MCI is performed using eqn (7) and (8), which is only exact for single-determinant wavefunctions, or using other approximations to the M-RDM. 108 The main drawback of MCI is that the computational cost grows very fast with the number of centres. Although this downside does not represent a problem for most systems, it prevents the calculation of expanded porphyrins and other Möbius systems.

One should also mention another important pitfall of multicentre indices: the atomic partition. Both I_{A_1,\dots,A_M} and MCI are very sensible to the atomic partition, QTAIM and TFVC partitions being the most reliable ones. ¹⁰⁹ In addition, the accuracy of the numerical integration over the atomic basins becomes an issue for large strings of atoms, as those occurring in molecules with large rings. In this sense, rings of more than ten members need very accurate calculations or the use of the Hilbert-space partition that provides analytical atomic overlaps, thus avoiding the integration hassle.

3. Electron delocalisation measures as indicators of aromaticity

In this section we will describe the electron delocalisation aromaticity indices and give the expression of the most important ones. In order to illustrate the performance of these indicators we will include the values for a small set of representative molecules: benzene, cyclohexane, borazine, pyridine, $\mathrm{Al_4}^{2-}$, and the transition state of the Diels-Alder cycloaddition reaction between butadiene and ethylene that we have calculated at the CCSD/cc-pVDZ level of theory. We will consider a ring structure of M atoms represented by the string $\mathscr{A} = [A_1, A_2, \ldots, A_M]$, where the elements are ordered according to the connectivity of the atoms in the ring.

The aromaticity indices calculated from the wavefunction include an energetic criterion using the adiabatic resonance energies calculated from the BLW method. From the group of indices based on the first- and higher-order density matrices, we can mention the ELF_{π} descriptor of Santos and coworkers, which has been used, among others, to account for excited state aromaticity and the para linear response (PLR) indicator. The PLR was defined by Sablon and coworkers as the LRK

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integrated over the atoms in the para position of a 6-MR. Several aromaticity indices defined in the last few years depend also on the topology of the electron density. Krygowski used the electron density calculated at the ring critical point (RCP), i.e. the lowest density point in a molecular ring plane 112 and other relevant quantities borrowed from the QTAIM. 113,114 On his side, Noorizadeh and Shakerzadeh used the Shannon entropy of the electron density calculated at the bond critical point to define an aromaticity index.115

Electron delocalisation has been tightly connected to aromaticity from the very beginning. The old concept of bond order-that later evolved to the electron sharing index, egn (3)—was brought up by Coulson in 1939 and it was first applied within the Hückel molecular orbital method to study the electronic structure of aromatic molecules. 116 July was among the first to suggest an index based on the uniformity of the interatomic distances, an indicator that a few years later he modified to take into account the charge gradient between bonded atom pairs in a ring. 117,118 In 1983, Jug suggested that aromaticity could be measured by the minimal bond order in a given ring structure¹¹⁹ and, two years afterwards, Bird defined a new measure of aromaticity based on the statistical deviation of the bond orders in the ring. More recently, Matta et al. 121 and Bultinck and coworkers 107 suggested the θ and BOIA aromaticity indices. The former is based on the electron delocalisation of the atoms and the latter uses the 2c-ESI in a ring. These indices compared the corresponding magnitudes to some reference values, usually taken from benzene. In this line, we suggested 122 the aromatic fluctuation index (FLU) that employs both quantities, the 2c-ESI and the atomic delocalisation of each ring member. FLU measures the uniformity of the electron delocalisation along the molecular ring and its bonding difference with respect to some aromatic reference, and it is given by:122

$$FLU(\mathscr{A}) = \frac{1}{M} \sum_{i=1}^{M} \left[\left(\frac{\delta(A_i)}{\delta(A_{i-1})} \right)^{\alpha} \left(\frac{\delta(A_i, A_{i-1}) - \delta_{ref}(A_i, A_{i-1})}{\delta_{ref}(A_i, A_{i-1})} \right) \right]^2,$$
(9)

where $A_0 \equiv A_M$, the atomic delocalisation is defined as:

$$\delta(A_i) = \sum_{A_i \neq A_i} \delta(A_i, A_j), \tag{10}$$

and α is a simple function to make sure that the first term in eqn (9) is always greater or equal to 1,

$$\alpha = \begin{cases} 1 & \delta(A_i) > \delta(A_{i-1}) \\ -1 & \delta(A_i) \le \delta(A_{i-1}) \end{cases}$$
 (11)

The CC and CN bond reference values are taken from benzene and pyridine in their ground state. FLU is close to 0 in aromatic species and increases as the molecule departs from the aromatic reference. Like any aromaticity index based on reference values it depends critically on the model aromatic molecules chosen and it cannot be used to study reactivity. 123 Moreover, it is difficult to compare molecules with different ring patterns. However, FLU gives a good account of the aromaticity for

Table 1 The CCSD/cc-pVDZ electronic-based aromaticity indices for a set of representative molecules and the transition state of the Diels-Alder cycloaddition (TS-DA). With the exception of FLU, for all indicators the larger the index, the more aromatic the species are. Values multiplied by 1000

	FLU	PDI	$I_{ m ring}$	MCI	$I_{ m NG}$	$I_{ m NB}$
Benzene	0	58.9	32.3	47.3	38.7	38.3
Pyridine	11.9	59.1	30.4	44.2	38.3	37.9
Borazine	_	10.4	0.9	1.0	21.1	20.2
Cyclohexane	339	6.3	0.2	0.2	16.7	15.9
TS-DA	315	51.0	21.5	31.0	36.1	35.7
$\mathrm{Al_4}^{2-}$	_	_	74.9	258.3	53.8	75.7

ground-state organic molecules. 124 These facts are illustrated by the numbers in Table 1, where we can see that FLU wrongfully assigns a very low aromatic character to the transition state of the Diels-Alder reaction, it does a good job in other organic molecules but it cannot assign a value to Al_4^{2-} or borazine due to the lack of reference systems.

The para-delocalisation index (PDI) measures the electron delocalisation across the ring by averaging the 2c-ESI of the three para-related positions in a 6-MR. 125 PDI is, obviously, limited to rings of six members and it suffers to describe molecular rings containing atoms with lone-pairs (it finds pyridine more aromatic than benzene) or to explain the small distortions around the equilibrium geometry (see Section 4).^{50,126} On the other hand, it does a good job in polycyclic aromatic hydrocarbons. 127 Ángyán 128,129 showed the connection between the 2c-ESI and the LRK for the QTAIM partition, which suggests that the PLR and the PDI could be measuring the same effect in a molecular ring.

 $I_{\rm ring}$ was defined by Giambiagi and coworkers¹³⁰ as their own multicentre index, eqn (7), applied to ring structures, assuming the obvious link between cyclic electron delocalisation and aromaticity. The MCI of Bultinck can also be used to quantify aromaticity. 107 Unlike I_{ring} , which only takes into account the Kekulé arrangement of the atoms in a ring, MCI considers all possible arrangements. The Kekulé structure contribution is usually the most important one and, therefore, I_{ring} and MCI rarely give disparate results. In such eventuality, the ring crosscontributions play a prominent role, as it happens in Al₄²⁻, where the delocalisation between non-bonded atoms is half as large as the delocalisation of neighbouring pairs. Unlike PDI and FLU, these multicentre indices do not have severe restrictions that limit their range of application, except for the numerical accuracy problems already mentioned. However, both I_{ring} and MCI multiply a number of overlaps that depend on the ring size, leading to a size-extensivity problem. Since overlap values are lower than one in absolute value, the MCI becomes artificially large for small rings, as we can see in the case of Al₄²⁻, which has an MCI value an order of magnitude larger than benzene. The latter issue was recently solved in our group by taking the Mth root of these two indices. 131 In addition, we also proved that taking into account the appropriate normalization factor these indices showed a good linear correlation with the topological resonance energy per π electron (TREPE).

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The normalised I_{ring} and MCI values were named I_{NG} and I_{NB} , ¹³¹ their expression reading:

$$I_{\rm NB}(\mathscr{A}) = \frac{\pi^2 G(N_{\pi})}{4 M N_{\pi}} I_{\rm ring}(\mathscr{A})^{1/M} \tag{12}$$

$$I_{\rm NB}(\mathscr{A}) = C \frac{G(N_{\pi})}{M N_{\pi}} {\rm MCI}(\mathscr{A})^{1/M} \tag{13}$$

where N_{π} is the number of π electrons of the atoms in the ring, $G(N_{\pi})$ equals 1 and -3 for aromatic and antiaromatic systems, respectively, and C = 1.5155 is a proportionality constant. These indices have the serious disadvantage of relying upon $G(N_{\pi})$, which might not be known a priori. However, one can give up the correlation with TREPE in favour of practicality and use the unnormalised MCI and I_{ring} , or their Mth root when comparing rings of different sizes. Finally, it is worth noting that-unlike $I_{\rm ring}$ and $I_{\rm NG}$ —both MCI and $I_{\rm NB}$ admit local and global counterparts depending on whether the index is calculated just for the atoms in the ring or all the atoms in the molecule. This feature opens up the possibility of interesting analyses of the effect of individual rings on the global aromaticity of a molecule, which to the best of our knowledge has not been explored thus far.

All electronic indices used in this section as well as the Harmonic Oscillator Model of Aromaticity (HOMA) and TREPE indices can be calculated using the software generated in our group, ESI-3D, 132 in conjunction with AIMPAC, 133 AIMAll or APOST-3D¹³⁵ packages to generate the atomic overlaps in the pertinent partition. These programmes read wavefunction or formatted checkpoint files from Gaussian, Gamess, Turbomole and NWChem, among other software programs. ESI-3D also reads the output of Gaussian, permitting the calculation of the electronic indices within a Hilbert-space partition.

A longstanding goal within the field of aromaticity has been the classification of molecules into different groups (aromatic, non-aromatic, antiaromatic) to rationalise structure, reactivity, and molecular properties. This classification was usually done on the grounds of chemical intuition or by examination of the occupied MOs. The discovery of new and more exotic molecules makes this classification more and more complicated, and for the past twenty years several aromaticity indices have been constructed to simplify this task. However, there has been such a proliferation of indices (often giving opposite answers) that is barely impossible to unambiguously characterise new molecules. To gain insight into the nature of aromatic molecules or to design and discover new molecules, aromaticity indices need to be validated.

4. Assessing the performance of aromaticity indices using a test set

A central tenet in the study of aromaticity is that it cannot be unambiguously defined and no single property exists that can be taken as a direct measure of aromaticity. Thus, directly comparing indicators of aromaticity for a given set of related compounds, e.g. polyaromatic hydrocarbons (PAH), is a daunting task as some of them measure distinct properties. In the literature several examples showing poor correlations between descriptors of different nature (i.e. magnetic-, energy-, structural-, or electronic-based indicators) have been reported, which prevents the use of this strategy to assess its performance. As illustrated by Bultinck, favourable correlations are observed, in general, among a set of electronic-based indices (MCI, PDI, BOIA, or θ), while comparing these descriptors with NICS gives poor correlations. 136 Similarly, work reported by Bickelhaupt et al. provided evidence that magnetic aromaticity is orthogonal to structural and energetic aromaticity for [5]paracyclophane. 137 In addition, aromaticity indices do not often give consistent results for long-established cases and these failures sometimes do not stand out in correlations of large data sets. The apparent contradictions obtained are occasionally justified by invoking the multidimensional character of aromaticity. 138 Therefore, it is not possible to discern between spurious results given by a descriptor and deviations due to the multidimensional character of aromaticity. These studies advocate for the use of more than one descriptor of aromaticity. While we agree on such an approach, one should first know the range of applicability of these indices, in order to choose the most appropriate ones.

In the last section we have seen how electronic aromaticity can be measured in a variety of flavours. The question now is how accurately electronic-based descriptors perform in a more diverse and complete set of systems and how they compare to other well-known descriptors of aromaticity. To gain insight into the accuracy and range of applicability of a particular descriptor it is crucial to assess the advantages and drawbacks of each measure in a variety of simple well-established situations before extending its application to more complex systems. If descriptors are not properly validated incorrect conclusions may arise when they are applied to characterise potential aromatic molecules. The aim of this section is to propose a simple protocol for assessing the performance of existing and prospective aromaticity descriptors and to provide a list of recommendations that may serve as a guide for future studies involving more complex systems.

In a series of papers we analysed whether a descriptor is able to properly account for the trends of aromaticity in a variety of well-established cases. For example, it is clear that benzene is more aromatic than toluene and a good descriptor of aromaticity should capture this fact. Since the development of the first descriptors, authors focused on identifying failures of descriptors in particular cases. However, there is a need for a more robust methodology to assess the performance of an index. The use of tests or training sets comprising a number of representative molecules is a common practice in other theoretical chemistry fields, such as density functional theory development, to judge the validity and the range of applicability of a new method. In 2008, we proposed to extend this idea to the field of aromaticity suggesting the use of a test set as a tool to assess the performance of aromaticity descriptors. 50,139 To this end, we proposed a series of tests with well-known trends that can be used to appraise the performance of existing and new indices of aromaticity (see Fig. 1). In this sense we are not

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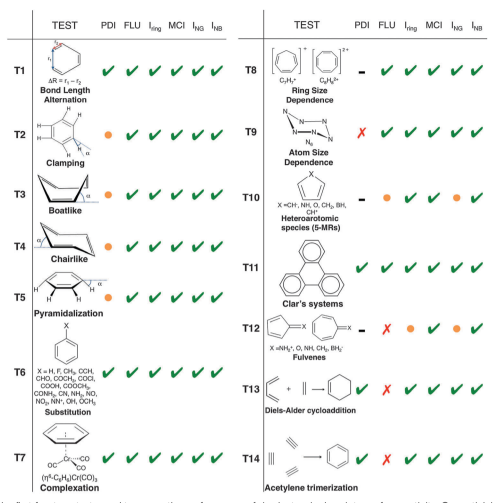


Fig. 1 Summary of the first fourteen tests used to assess the performance of six electronic descriptors of aromaticity. Green tick implies that the index follows the expected trend of aromaticity, red wrong mark indicates a major failure, while orange round represents a minor failure. See ref. 50 for a more detailed discussion.

strictly comparing indices between them but we are analysing their behaviour in a variety of cases that account for situations found along the spectra of aromatic molecules.

To set up a test of aromaticity we considered some guidelines. First, all tests must be based on well-established cases. Accumulated chemical experience provides several series of compounds with predictable trends of aromaticity. Second, controversial examples reported in the literature should be avoided e.g. whether the inner or the outer ring in anthracene is more aromatic. Third, the size of the systems should be relatively small so that the test can be readily performed. Initially, we proposed a total of fifteen tests including: five benzene distortions, one test that evaluates the substitution effects in the benzene ring, another that analyses the consequences of complexation, a couple that studies ring and atom size dependence, two that checks the trends of aromaticity in a series of heteroaromatic species of different sizes, one for reproducing the expected trends in a series of Clar's systems, another that studies the changes in aromaticity in a series of fulvenes, and two tests that assess the changes in aromaticity along the reaction path of two chemical reactions. Later on we added a

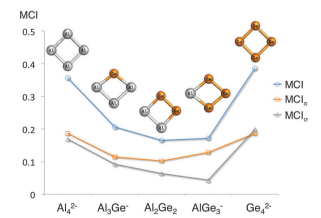


Fig. 2 Variation of multicentre indices MCI, MCI_{π} , and MCI_{σ} (in electrons) along the series $Al_4{}^2$, Al_3Ge^- , Al_2Ge_2 , $AlGe_3^+$, and $Ge_4{}^2$ (adapted from ref. 154).

test for performance appraisal in all-metal clusters. We propose now to exclude the test corresponding to the aromaticity trends in a series of heteroaromatic 6-MRs (Test 10 in ref. 50) due to its Review Article Chem Soc Rev

slightly controversial results. Therefore the final set comprises a total of fifteen tests (see Fig. 1 and 2).

We have used these tests to assess the performance of some of the electronic-based indices described in the previous section. If a descriptor is to be used in complex systems, first it should properly reproduce simple cases as the ones gathered in this test set. We anticipate that most of the indices evaluated fail in some of these well-established cases. Once the pitfalls and the range of applicability of a particular descriptor of aromaticity are validated, it can be applied to drive the discovery of new aromatic compounds. Otherwise, the inadequate use of an indicator can lead to the incorrect classification of a compound as aromatic (or antiaromatic) and the wrong assignment of its molecular properties. Evidently, these tests are not limited to electronic indices and we encourage and challenge researchers in the field to assess the performance of their indices using a test set instead of doing direct comparisons. This will provide valuable information about the limitations of the indices and help to avoid misinterpretations. For the sake of comparison, we also choose the simplest aromaticity index, HOMA, and the most popular one, NICS, to assess their performance in our test set. 50

4.1 Benzene distortions

Benzene is the archetypical organic aromatic molecule; it is planar and shows perfect bond length equalisation. However, when benzene is fused to another aromatic ring (e.g. to form naphthalene) the characteristic bond length equalisation is broken as is observed, for example, in C₆₀. In addition, pyramidalisation of CC bonds is commonly exhibited in large, strained systems, illustrating the certain ring flexibility of benzene. It is well known that in-plane or out-of-plane distortions alter the natural electron delocalisation patterns of benzene leading to a situation where the electrons are more localised. Thus, by altering planarity or bond lengths with respect to benzene's reference geometry a slight decrease in the aromatic character is expected. Common in-plane deformations found in benzene-containing molecules are bond-length alternation and clamping, while boat-like, chair-like, and pyramidalisation are found among the most frequent out-of-plane deformations. 126 A satisfactory descriptor should reproduce the decrease of aromaticity when the five above-mentioned distortions are applied. We proposed to determine the quality of existing electronic aromaticity indices by means of such deformations. As Fig. 1 shows, all multicentre indices evaluated in this study perfectly reproduce the decrease of aromaticity when a distortion is applied. The correct performance of referencebased indices like FLU and HOMA is certainly good in these particular tests because they actually measure aromaticity as the deviation from benzene's electronic distribution and geometry, respectively. Interestingly, delocalisation between carbons in the para position (PDI) is quite sensitive to the BLA distortion but not to the other four distortions (clamping, pyramidalisation, boat-like, and chair-like). We compared the trends obtained by electronic-based indices with structural (HOMA) and magnetic (a number of NICS descriptors

comprising NICS(0), NICS(1), NICS(1)_{zz}, and NICS(0)_{πzz}), which are, in general, in line with the expected trends. Only NICS(0) seems to be non-sensitive to clamping while HOMA tends to overestimate the loss of aromaticity for in-plane distortions. We recommend the use of multicentre indices and FLU for a quantitative analysis of distorted benzene-related compounds while PDI is only recommended for systems presenting bond length alternation. The concept of aromaticity can have a major impact on the design of benzene-based molecules like graphene, nanotubes, and fullerenes. For example, Martín-Martínez *et al.* employed multicentre indices to characterise the properties of several graphene nanoribbons and carbon nanotubes. 140

4.2 Substitution and complexation

Substitution and complexation to an aromatic ring are also common features found in a vast number of molecules spanning the whole chemical space. The nature of the substituent is key to explain the reactivity of an aromatic ring while complexation of an ion to aromatic rings plays a major role, for example, in molecular recognition. Therefore, these situations need to be considered to assess the proper performance of aromaticity descriptors. As shown by Krygowski et al., the effects of substituents in benzene, either electron-donating or electron-accepting groups, alter the π -delocalisation inducing partial localisation of π -electrons. Therefore, a decrease in aromatic character with respect to benzene is expected for substituted benzenes and this trend should be reproduced by aromaticity descriptors. We selected a representative set of sixteen substituted benzenes as a test set. As shown in Fig. 1, all electronic indices are in conformity with this statement indicating a loss of aromaticity with respect to benzene. In contrast, NICS(0) and NICS(1) assign a higher aromatic character to several substituted benzenes, although this trend is corrected by the more robust $NICS(1)_{zz}$ and $NICS(0)_{\pi zz}$ descriptors. Structural indices as HOMA also suffer minor deviations from the expected tendency. A similar behaviour is observed when a metal atom is complexed to benzene, as in the case of $(\eta^6-C_6H_6)Cr(CO)_3$. Surprisingly, it was found that the aromatic character of a benzene ring complexed to CrCO3 was 40% higher than benzene itself according to magnetic-based descriptors. Analysing the characteristics of the complexed ring in detail, we observe several deformations including pyramidalisation, loss of planarity, ring expansion, and bond length alternation. Charge transfer between highest occupied π -orbitals of benzene and LUMO orbitals of the CrCO₃ moiety is mainly responsible for such a ring distortion. We studied this particular case in detail by means of electronic-based descriptors and NICS-scan techniques concluding that a significant loss of aromaticity is observed in contrast to the original predictions. 143 NICS(0) and NICS(1) values are strongly affected by ring currents involved in the Cr-C₆H₆ bonding, unlike the out-of-plane zz counterparts $(NICS(0)_{\pi zz})$ and $NICS(1)_{zz}$ that provide the correct trend. In general, to study the effects of substitution and metal complexation all electronic indices perform remarkably well and their use is recommended. These observations pave the way

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towards the study of more complex systems involving substitution and complexation as the ones described in Section 5.

4.3 Ring-size dependence

Most aromatic molecules contain 6-MRs. However, there are a number of molecules with as few as three ring members and, in contrast, large rings such as porphyrins. Here, we explore the applicability of electronic-based indices to describe the aromatic character of rings with different sizes. To study ring-size dependence of aromaticity descriptors, we proposed the series C₆H₆, C₇H₇⁺, and C₈H₈²⁺ where a reduction of aromatic character is expected as the size of the ring increases while the number of π -electrons remains constant. Interestingly, FLU, MCI, I_{ring} , I_{NG} , and $I_{\rm NB}$ are all in agreement with the expected trend. PDI cannot be computed for rings other than 6-MRs. In comparison to electronic indices, magnetic-based NICS(0) places incorrectly the C₇H₇⁺ species. HOMA results are in line with the expected trend. A second relevant question is whether the values obtained for a set of rings with different size are comparable between them. $I_{\rm ring}$ and MCI are not normalised and thus may suffer ringsize dependence as some of us demonstrated before.131 Their normalised versions, I_{NG} and I_{NB} , are specifically designed to account for systems of different sizes and expected to show the correct behaviour when increasing the ring size. The substantial computational cost and the numerical accuracy problems associated with the calculation of multicentre indices for large rings leave FLU as one of the few electronic-based measures that can be applied beyond rings of ten members.

4.4 Atom size dependence

Since the very beginning, aromaticity rapidly broadened its realm to atoms other than carbon. Atoms of different types are commonly found to form rings that display typically aromatic properties. Substituting carbon for other atoms perturbs the π -delocalisation patterns of such molecules with respect to benzene. Therefore, by substituting carbon-hydrogen fragments by nitrogen atoms to form a chair-like N₆ ring one expects a reduction of the aromatic character. All multicentre indices and FLU account for the decrease of aromaticity when going from C₆H₆ to N₆. In contrast, PDI values increase significantly from 0.103 e in benzene to 0.131 e in N₆, showing that PDI is quite dependent on the atom nature. The same trend is observed for NICS(1) and NICS(1)zz, which assign a higher aromatic character to N6, revealing the atom-size dependence of these descriptors. The structure-based HOMA index gives the proper results. However, FLU and HOMA are limited by available reference values, which significantly restrict their applicability to rings containing elements other than carbon. In general, we recommend the use of multicentre indices to study rings with a variety of elements including metalloaromatic systems such as metallacycles and allmetal clusters. These indices shall aid the design and characterization of all-metal clusters with specific properties (see Section 5.2).

4.5 Heteroaromatic series, Clar's systems, and fulvene series

To complete the proposed test set with other relevant situations displayed by aromatic molecules, we included a total of three more tests. First, we introduced a test to predict the proper trend of aromaticity along the following well-established heteroaromatic series C_4H_4X (X = CH⁻, NH, O, CH₂, BH, CH⁺). Second, we proposed a test that analyses the effect of fusing aromatic rings represented by five Clar systems. Third, we included a test to assess the expected trend of aromaticity in 5-MR and 7-MR fulvenes with different substituents. In general, all electronic-based indices passed the three tests (see Fig. 1). The only exception to properly account for the predicted order of aromaticity in a series of fulvenes is FLU. Magnetic (NICS(0), NICS(1), NICS(1)_{zz}, NICS(0)_{πzz}) and structural indices (HOMA) are also in line with the expected trends for all three tests. Only NICS(0) and NICS(1) incorrectly predict C₄H₄NH to be more aromatic than C₅H₅⁻. Therefore, electronic based indices are a good choice for this kind of systems. In particular, fulvenes, termed aromatic chameleons by Ottosson and coworkers, 144 display a wide range of interesting properties. The application of electronic descriptors to describe the tuneable aromatic character of these species at their excited states will be highlighted in Section 5.3.

4.6 Chemical reactions

Reactivity is strongly linked to the concept of aromaticity. A considerable number of reactions present reactants, products or transition states (TS) with a clear aromatic or antiaromatic character. For example, aromaticity determines the regioselectivity of Diels-Alder reactions in fullerenes as discussed in Section 5.1.6. In a recent review, Schleyer, Wu, Cossío, and Fernández have extensively discussed the concept of transition state-aromaticity and its role in pericyclic, pseudopericyclic, and non-pericyclic reactions. 145 In this work the authors point out the importance of properly determining the aromaticity of transition states to understand the reaction mechanism. Are electronic indices capable of recovering the aromaticity changes along the reaction path? To answer this question, we suggested the study of Diels-Alder cycloaddition and the acetylene trimerisation. In the case of the Diels-Alder reaction it is well-known that the reaction takes place through a boat-like aromatic transition state. Along the reaction path, we expect to observe a peak of cyclic electron delocalisation in the ring at the vicinity of the TS. PDI, multicentre, and magnetic indices clearly reproduce this trend. In contrast, FLU and HOMA break down in this test, showing a steady increase of aromaticity from reactants to products. 123 The case of the [2+2+2] trimerisation of acetylene is slightly different. An increase of aromaticity is expected when going from reactants to TS, after this point it reduces somewhat the aromatic character until a final increase is observed to form benzene as a final product. Again, this trend is perfectly reproduced by all electronic indices besides FLU. HOMA shows exactly the same erroneous behaviour. These findings support the idea that reference-based aromaticity indices such as HOMA and FLU should never be used to study changes in aromaticity along a reaction path. 223 On the other hand, NICS indices correctly reproduce the shape of the curve but assign a more aromatic value to the TS than to benzene itself. Recently, Mandado and Ponec assessed the performance of MCI for a number of pericyclic reactions further proving the validity of this electronic measure for chemical reactivity. 146 In general, electronic-based multicentre indices perform better for

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this test set than other indices analysed either structural or magnetic.

4.7 All-metal clusters

The recent discovery of metallabenzenes, 147 heterometallabenzenes, 148,149 metallabenzynes, 150 metallacyclopentadienes, 151 all-metal 152,153 and semimetal clusters, etc. has prompted a revolution in the study of aromaticity.¹⁷ At variance with the classical aromatic organic molecules that only possess π -electron delocalisation, these compounds have σ -, π -, δ - and ϕ -electron delocalisation, which can be even combined to give double or triple aromaticity, the so-called multifold aromaticity. This latter issue makes most of the classic indicators of aromaticity not valid to discuss these complex systems, ^{49,50} and for such a reason more general and reliable indices of aromaticity are needed. These indices must fulfil two requirements: first, to be free from reference values (since it is difficult to choose a most aromatic reference molecule); and second, indices should be separable into σ -, π -, δ -, and ϕ -components of aromaticity. Multicentre electronic delocalisation indices and NICS are among the few indicators that fulfil these two requirements for rings of arbitrary size. 139,154

The new aromaticity test consists of the 4-MR series of valence isoelectronic species $[X_nY_{4-n}]^{q\pm}$ (X, Y = Al, Ga, Si, and Ge; n=0-4), which should exhibit particular aromaticity trends. 154 Let us focus on the series from $\mathrm{Al_4}^{2-}$ to $\mathrm{Ge_4}^{2+}$; similar trends are expected for the other series. We anticipate a steep decrease in aromaticity when going from Al₄²⁻ to Al₃Ge⁻ due to the reduction of symmetry and to the substitution of one Al atom by a more electronegative Ge atom. Although more arguable, we anticipate a smooth aromaticity reduction going from Al₃Ge⁻ to Al₂Ge₂. The same decrease should occur from $Ge_4^{\ 2+}$ to Al_2Ge_2 ; giving the following trend of aromaticity: $Al_4^{2-} > Al_3Ge^- \ge Al_2Ge_2 \le$ $AlGe_3^+ < Ge_4^{2+}$. From Fig. 2, it is observed how both total MCI and its π -component (MCI $_{\pi}$) successfully provide the expected order of aromaticity, showing a concave U shape. NICS indices were submitted to the same test but only NICS(0) $_{\pi}^{\text{RCP}}$ provided the expected trend. In these rings, with different atomic sizes, the calculation of NICS should be performed at the ring critical point (RCP), as suggested by Morao and coworkers. 155 The different analogous series analysed yield similar conclusions. Therefore, we concluded that electronic indices are also suitable to analyse aromaticity in all-metal and semimetal clusters.

As we have seen, none of the indices analysed is infallible and some of them fail in a number of simple cases, putting forward important limitations of these indicators. These results are summarised in Fig. 1 and 2. We hope that the results of this test set will provide a collection of guidelines that will drive the choice of aromaticity indicators in future studies and will lead to the construction of more robust and applicable descriptors.

5. Some applications

In this section, we present some of the most relevant applications carried out by our research group using the above defined electronic-based measures of aromaticity.

5.1 Electronic measures of aromaticity in classical aromatic compounds

5.1.1 Polycyclic aromatic hydrocarbons and derivatives. In our first application, a set of planar and bowl-shaped PAHs, together with C_{60} and C_{70} fullerenes, were evaluated by means of PDI and FLU indices, showing that these indices could identify regions of local aromaticity and antiaromaticity in PAHs and fullerenes. PDI and FLU values for C_{60} and C_{70} indicated the relatively weak local aromaticity of the 6-MRs and the non-aromatic or antiaromatic character of the 5-MRs in fullerenes. 125,127,156

Acenes, phenacenes, and non-planar helicenes are three different series of benzenoid compounds. Using FLU and PDI 157 we found that the most reactive inner rings are somewhat more aromatic than the outer ones in [n]acenes, whereas for [n]phenacenes the external rings are clearly the most aromatic. And for [n]helicenes, despite the departure from planarity, we observed almost the same aromaticity trends as in [n]phenacenes. 158

5.1.2 Substituent effects on aromaticity. The analysis of the substituent effects on benzene proved the high resistance of aromatic systems to disrupt the π -electron structure in electrophilic aromatic substitution reactions. In particular, changes in PDI when going from benzene to substituted benzene derivatives are small and correlated with Hammett substituent constants. 142 The same behaviour was observed for the complexation of a lithium cation to a series of PAHs. 159 Substituent effects were also studied in 4-substituted-1,2-benzoquinones. Results show that only the keto group in the meta position is affected by the electron-donating/ attracting power of the substituent, whereas the para-related C=O is not. Although MCI and FLU display small changes, these indices agree on assigning a more aromatic character to rings with electron-donating substituents. 160 On the other hand, the substituents do have a large effect on the aromaticity of pyrazoles and imidazoles with N-substituents.161 We found that the imidazole ring is more stable than the pyrazole one. The reason for the relative energy difference was attributed to the weakness of the NN bond in the latter, and not to a higher aromaticity of the former, as both rings present similar MCI and FLU values. 162 By comparing the corresponding substituted series of benzene, the latter appears not only to be more aromatic, but also more robust towards substitution effects as denoted by the slope of the FLU^{1/2} vs. $\sigma_{\rm R}$ correlation (see Fig. 3). A recent work showed that 1-indenones and their aza derivatives are more stable than 2-indenones because their 6-MR is more aromatic. 163 Interestingly, tetrafluorination of the 6-MR in such compounds hardly causes any change in the local aromaticity of this ring, thus confirming that the aromaticity of benzene rings is quite robust.

5.1.3 Discrepancy with magnetic measure of aromaticity. Pyracylene was the first case in which we showed that the calculation of NICS at the ring centre should be analysed with caution, as NICS(0) wrongly assigned an increase of the local aromaticity of 6-MRs upon distortion from planar to pyramidalised pyracylene. ¹⁶⁴ In contrast, NICS(1) calculated 1 Å above pyracylene reported the expected decrease of aromaticity upon bending. Both PDI and ring currents contradicted NICS, giving the expected reduction of aromaticity with bending. Another conflicting case of NICS' performance is found in [2,2]paracyclophane, in which

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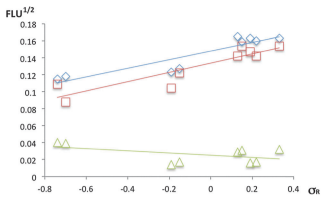


Fig. 3 Correlation between $FLU^{1/2}$ and the Hammett substituent constant σ_R . Blue diamonds, imidazole (Im); red squares, pyrazole (Pz); and green triangles, benzene (Bz) (adapted with permission from ref. 161).

the NICS predicted decrease of local aromaticity of the stacked rings was not real, but caused by the coupling of the magnetic fields generated by these two rings. The same problem was experienced when the local aromaticity of a series of polyfluorene compounds with an increasing number of π -stacked layers was analysed. NICS shows a spurious increase of aromaticity that is due to the coupling between the magnetic fields generated by the π -stacked rings, whereas PDI, FLU, and HOMA showed that aromaticity does not change due to π -stacking. Finally, NICS also fails when attributing an aromatic character to the (HF)₃ ring. 167

5.1.4 Aromaticity in DNA bases. First, we analysed how the aromaticity of the rings of the guanine-cytosine base pair was affected when a metal cation is coordinated to the N₇ position of guanine (see Scheme 1, M = Ca²⁺, Cu⁺, and Cu²⁺). Such an interaction causes a change in the strength of the hydrogen bonds $(N_1-H\cdots N_3 \text{ and } N_2-H\cdots O_2 \text{ become stronger, whereas})$ O₆···H-N₄ weakens). In turn, these alterations affect the aromaticity of the 5- and 6-MRs of the nucleobases. The observed increase of the aromaticity of the guanine and cytosine 6-MRs due to the interaction with Cu⁺ and Ca²⁺ was attributed to the strengthening of hydrogen bonding in the guanine-cytosine pair that stabilises the resonance structure with a π -sextet in the 6-MRs. On the other hand, the reduction of aromaticity in the 5- and 6-MRs rings of guanine due to the interaction with Cu²⁺ is caused by the oxidation process, which removes a π electron disrupting the π -electron distribution.

 N_7 N_7 N_1 N_1 N_2 N_3 N_4 N_4

Scheme 1 The guanine-cytosine base pair interacting with the metal cation ($M = Ca^{2+}$, Cu^+ , and Cu^{2+}). Labels of the atoms involved in the H-bond interactions are shown.

In a subsequent work, we studied the change of aromaticity in a series of size-expanded benzobases (xDNA). The insertion/addition of a benzene ring in the natural DNA bases reduces the local aromaticity of both 5- and 6-MRs of DNA bases according to PDI and FLU values. The degree of aromaticity of the added 6-MR of xDNA bases varies in the order xA > xG \sim xC > xT. Interestingly, an inverse correlation between the HOMO-LUMO gap of the size-expanded benzobases and the aromaticity of the inserted/added benzene ring was observed. ¹⁶⁹

5.1.5 Hückel's rule and beyond. Hückel's rule is essential for the comprehension of aromaticity in organic molecules. The fact that it is only valid for conjugated monocyclic systems prompted the appearance of new rules that could describe local aromaticity in PAHs. Clar's rule¹⁷⁰ is probably the most successful extension of the Hückel rule to PAHs. 171 According to Clar's rule, the Kekulé resonance structure with the largest number of disjoint aromatic π -sextets is most important to characterise PAHs. π -sextets are defined as six π -electrons localised in a single benzene-like ring separated from adjacent rings by formal CC single bonds. Clar's rule was validated by PDI values calculated in the rings of a set of PAHs. 172 For instance, phenanthrene has two π -sextets located in the outer rings, which are the most aromatic rings in the molecule according to PDI. The referred Hückel rule was also checked from π -electronic delocalisation measures. In particular, we studied a series of systems in which two electrons were either added or removed. In an N-electron aromatic system, the addition of two electrons leads to an antiaromatic N + 2 species in which we expect these extra two electrons to be essentially localised. Similarly, the extraction of two electrons to give an antiaromatic N-2 species should also reduce the delocalisation in the system. Our results show that there is an important increase of electronic delocalisation (of about 1 e) when going from antiaromatic $4n\pi$ systems to aromatic $(4n + 2)\pi$ systems. The change in π -electronic delocalisation when we move from a $(4n + 2)\pi$ -aromatic to a $4(n + 1)\pi$ -antiaromatic species by adding a pair of electrons is much smaller (see Fig. 4). FLU, PDI, and MCI criteria, as well as the cross terms of the total π -electron delocalisation, correctly assign an aromatic or antiaromatic character to each system, regardless of the number of electrons. Therefore, these indices provide aromaticity values in agreement with Hückel's rule. 173,174

4n Baird's rule represented the extension of the renowned 4n + 2 Hückel rule to open-shell species, ¹⁵ while Hirsch's $2(n + 1)^2$

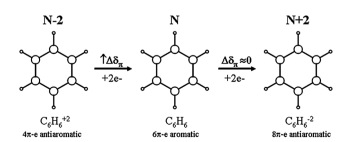


Fig. 4 Change in π -electronic delocalisation from aromatic to antiaromatic systems with the successive addition of 2 electrons (adapted from ref. 174).

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 $rule^{29}$ of aromaticity has considered the analogue of the 4n + 2rule for spherical systems. Our group recently extended the latter rule to open-shell spherical molecules, providing evidence that spherical species having the same-spin half-filled last energy level and the rest of the levels being fully filled $(2n^2 + 2n + 1)$ electrons and S = n + 1/2) are aromatic.³⁰ For instance, neutral C₆₀ appears to be non-aromatic according to MCI and NICS(1)zz; whereas singlet C_{60}^{10+} that obeys the $2(n+1)^2$ rule is found to be aromatic. On the other hand, C_{60}^{19+} with S = 9/2 and C_{60}^{1-} with S = 11/2following the $2n^2 + 2n + 1$ rule appear to be even more aromatic than C₆₀¹⁰⁺. This new rule may become a powerful tool to study the stability of high-spin spherical molecules. In Section 5.3 we comment on the aromaticity of excited states in connection with the Baird rule.

5.1.6 Aromaticity determines the regioselectivity of Diels-Alder reactions in fullerenes. The reactivity and regioselectivity of Diels-Alder (DA) reaction involving empty fullerenes is generally favoured for [6,6] bonds, whereas in endohedral metallofullerenes [5,6] bonds are commonly more reactive. 175,176 When a metal cluster is encapsulated inside a fullerene there is a charge transfer from the metal cluster to the fullerene. In the case of M_3N units (M = Sc, Y, Gd...) formally six electrons are transferred to the carbon cage. We decided to analyse the effect of adding electrons to the fullerene cage by calculating the reaction profile for the DA reaction of cyclopentadiene (Cp) to the [6,6] and [5,6] bonds of C_{60}^{n-} (n = 0-6) species. The C_{60}^{n-} (n > 0) was taken as a model for the cage of endohedral metallofullerenes. It was found that the reaction becomes more exothermic (and the barrier is reduced) for the [5,6] attack when n increases from 0 to 6 electrons (see Fig. 5). On the other hand, for the [6,6] addition the exothermicity is somewhat reduced (and the barrier increases) when n increases. For n = 4-5, there is a change in the regionelectivity of the process and [5,6] becomes the preferred attack.

To understand the change of regioselectivity in C₆₀ upon reduction, we calculated the MCI for the 5- and 6-MRs of C₆₀ⁿ⁻ showing that the aromaticity of the 5- and 6-MRs increase and decrease, respectively, with the successive addition of electrons to the C₆₀ molecule (see Fig. 5). Because the DA addition leads

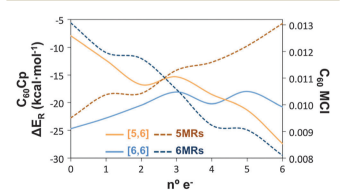


Fig. 5 Change in the reaction energy of the DA between Cp and $C_{60}^{\ \ n-}$ and in the aromaticity (dashed lines) of five- and six-membered rings of C_{60}^{n-} as a function of number of electrons added (n). Reproduced from ref. 176 with permission from the Royal Society of Chemistry.

to a change from planar sp² to tetrahedral sp³ for the C atoms of the attacked bond, when the addition occurs on a [6,6]-bond type, the π -conjugation of two 5-MRs and two 6-MRs is lost. On the other hand, when the addition is on a corannulenic [5,6]bond, the conjugation vanishes in three 6-MRs and one 5-MR. For neutral C₆₀, the preferred attack is the [6,6] addition because the aromaticity of only two of the most aromatic 6-MRs is lost (the [5,6] attack affects the aromaticity of three 6-MRs). On the other hand, for C_{60}^{6-} the most favourable addition is at the [5,6] bond because the aromaticity of only one of the most aromatic 5-MRs is lost. In this case, the changes in aromaticity between a 6-MR and a 5-MR upon reduction determine the regioselectivity of the DA additions to fullerenes. 176

5.1.7 Aromaticity in metalloporphyrins. Assessing the aromatic character of rings beyond ten members corresponds to one of the current challenges of electronic descriptors of aromaticity. Porphyrins and their derivatives are macrocycles with highly delocalised π -electrons. They play a key role in a number of biological processes and catalysis. Some of the molecular properties exhibited by porphyrins can be rationalised in terms of their aromatic character. To this end, structural and magnetic descriptors have been used to estimate the role of aromaticity in a variety of porphyrin-related systems. However, the large size and peculiarities of these molecules prevent the application of some well-known measures. Observations based on structural indices and ring currents reveal that the inner porphyrin ring is significantly more aromatic than the outer ring.

Porphyrins are often coordinated to a metal atom, which plays a major role in determining shape, reactivity, and molecular properties. The effects of metal coordination and the spin state on the electronic delocalisation patterns of metalloporphyrins were unknown. For this reason, we decided to explore the chemical bonding and aromaticity patterns in a series of metalloporphyrins that include first row transition metals (Sc-Zn), alkaline earth metals (Mg, Ca), and second-row transition metals (Ru, Pd, Ag, Cd) in a variety of spin states. 177 The quantification of aromaticity in metalloporphyrins is challenging because the metal placed in the ring centre makes the calculation of some well-known descriptors such as NICS quite troublesome. Therefore, magnetic indices were discarded for this study. The use of electronic indices in porphyrins is scarce because of the computational cost and the accuracy problem associated with multicentre indices. As we mentioned above, FLU is one of the few electronic indices that can be applied to measure aromaticity in large rings. Considering that FLU performs remarkably well for heteroaromatic 5-MR (see Section 4) it was our descriptor of choice to study aromaticity patterns in a number of metalloporphyrins. In comparison to the metal free porphyrin, FLU values indicated that the aromatic character of the inner ring (16π -electrons) is hardly affected by the metal complexation whereas the electron delocalisation of the outer ring $(20\pi\text{-electrons})$ is significantly enhanced by the presence of the metal (see Fig. 6). This was notably clear in the case of 5Ru where the value of FLU is roughly equal for inner and outer rings. Therefore, FLU is an attractive alternative to structural

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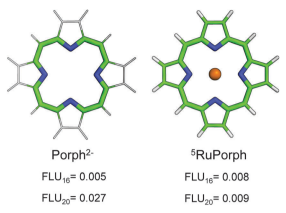


Fig. 6 FLU values obtained for the inner (16-MR) and outer (20-MR) rings of the free-based and ⁵Ru porphyrins. Electron delocalisation patterns are highlighted in green for both systems.

and magnetic indices for the analysis of aromaticity in porphyrins and metalloporphyrins. Otero et al. made also use of an approximation of MCI employing Aihara's circuit resonance energy to study the effect of aromaticity on the thermodynamic stability of hydroporphyrins. 178 Their study provides a good starting point for the application of multicentre indices towards larger systems. Recently, the synthesis and characterization of expanded porphyrins with Hückel and Möbius topologies showing aromatic and antiaromatic character has gained a lot of attraction. 179 Identifying methods to properly characterise the aromaticity of these peculiar systems may help on the design of expanded porphyrins.

5.2 Metalloaromaticity

One of the most important findings in metalloaromaticity was the discovery of the Al₄²⁻ cluster by Boldyrev, Wang, and coworkers. ¹⁹ This cluster was the first example of all-metal species with σ - and π -aromaticity. It is well known that π -electrons in benzene are distortive and it is the σ -skeleton which is responsible for the D_{6h} symmetry of benzene. We wondered whether the σ - and π -electrons in Al₄²⁻ favoured the D_{4h} or the D_{2h} structure. To solve this question, we performed an energy decomposition

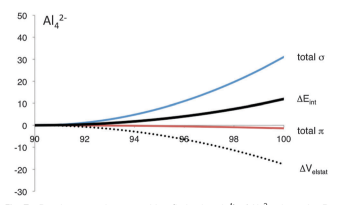


Fig. 7 Bond-energy decomposition (in kcal mol⁻¹) of Al_4^{2-} along the D_{4h} to D_{2h} distortion measured with the θ angle (in deg., defined in Fig. 8) from delocalised to localised structure (adapted from ref. 153).

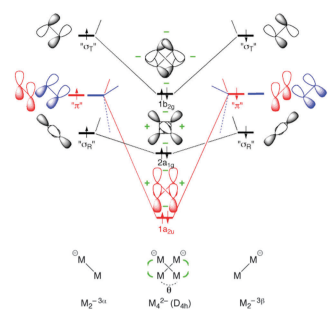


Fig. 8 Schematic molecular orbital interaction diagram of ${\rm Al_4}^{2-}$ constructed from two $[M_2]^{...}$ fragments at the BP86/TZ2P level (adapted from ref. 153). The positive and negative signs in green indicate whether there is a gain or loss of interaction when going from the D_{4h} to D_{2h} structure.

analysis (EDA, see Fig. 7) showing that the π -electrons in ${\rm Al_4}^{2-}$ prefer the D_{2h} structure but the σ -electrons force the double bond to delocalise, leading to the regular D_{4h} geometry.¹⁵³ This analogous behaviour to benzene was explained through the corresponding MO diagram (see Fig. 8), where it is observed how both the π and the radial- σ (σ_R) orbitals induce distortion, whereas the tangential occupied orbitals (σ_T) are the ones responsible for the D_{4h} structure. This different character of σ_R and σ_T orbitals is also supported by MCI indices that show a more important contribution to the total σ -aromaticity from the radial than from the tangential orbitals.

The geometrical conformers of $M_2N_2^{2-}$ (M and N = B, Al, and Ga) clusters that are valence isoelectronic with Al₄²⁻ were studied with a gradient embedded genetic algorithm. 180 For $Al_2B_2^{\ 2-}$ and $Ga_2B_2^{\ 2-}$, the C_{2v} structures with a short B-B bond are more stable than the D_{2h} alternate structure. On the other hand, the latter symmetry is the most stable for $Ga_2Al_2^{2-}$. Clusters with B atoms prefer the C_{2v} structure (with X-X, Y-Y, and two X–Y bonds) to form strong B–B bonds. In the $Ga_2Al_2^{2-}$ species this is not possible and the two structures are almost thermoneutral with D_{2h} (with four X-Y bonds) being somewhat more stable. Finally, for all cyclic $M_2N_2^{2-}$ isomers we found large MCI_{σ} and MCI_{π} values, similar to those obtained for Al_4^{2-} .

 $\mathrm{NaMg_3}^-$ and $\mathrm{Na_2Mg_3}$ clusters containing the *cyclo*- $\mathrm{[Mg_3]}^{2-}$ unit are two of the very few electronic species with π -bonding without the occurrence of a σ -framework. Interestingly, the aromaticity switches from σ to π when the Mg₃²⁻ unit coordinates Na atoms to give NaMg₃⁻ and Na₂Mg₃ clusters. The distance between the coordinated Na atom and Mg3 can be used to tune the aromaticity and prompt an unprecedented switch from σ - to π -aromaticity.

The aromaticity of all-metal clusters with transition metal complexes involving d and f orbitals implies more complicated analyses due to the large number of electrons involved and the inclusion of relativistic effects. The occurrence of highly delocalised valence electrons occupying the large angular momentum orbitals in transition metals gives rise to multifold aromaticity. To study this phenomenon, we analysed the aromaticity of the series Cu₃+, Y₃-, La₃-, Ta₃O₃-, Hf₃, ⁵Ta₃-, and ³Hf₃ with the MCI index. ¹⁸³ Cu₃ was confirmed to present exclusive σ-aromatic character; whereas Y₃⁻ and La₃⁻ present σ - and π-aromaticity, supported by MCI_{σ} and MCI_{π} values. On the other hand, Ta₃O₃⁻ is the first cluster presenting both π - and δ-aromaticity, as confirmed by the small value of MCI_{σ}, and the twice as large MCI_{π} and MCI_{δ} values. The aromaticity patterns of Hf3 were even more complex, showing prominent σ -, but also significant π - and δ -orbital contributions (the so-called threefold aromaticity). According to MCI, the open-shell molecules $^5\mathrm{Ta_3}^-$ and $^3\mathrm{Hf_3}$ show σ + π + δ and σ + π aromatic character, respectively.

In a recent work¹⁸⁴ we analysed the metalloaromaticity of the 5-MR of the $M(XC_3H_3)(PH_3)_2$ species (M = OsH₃, OsCl₃, OsCl₂, RuCl₂, RhCl₂, and IrCl₂ and X = NH, O, S, CH⁻, and CH⁺). Our results showed that the heterometallacycle 5-MR has a low aromatic character (MCI values of about 0.01 e) except for X = CH⁺ compounds in which this 5-MR is antiaromatic as denoted by negative MCI values. Finally, it is worth mentioning that Möbius metalloaromaticity was found in the Cu(OC₃H₃NH)₂ and related complexes and confirmed through I_{NB} values. 149

From these analyses, it becomes clear that the characterization of metalloaromaticity is much more complex than in classic organic compounds and that multicentre electronic indices are well suited for analysing aromaticity in metalloaromatic species. In general, one should make a judicious choice of the indicators used to evaluate aromaticity. Aromatic indices should not be limited to classify existing molecular systems. The main purpose of aromaticity indices is actually to improve our understanding in order to design new compounds with compelling properties.²⁴

5.3 Aromaticity of excited states

In a recent review, Ottosson and co-workers underlined the importance of applying the concept of aromaticity to rationalise excited state properties and reactions. 144 Aromaticity and antiaromaticity effects observed in the excited states may play a similar role in understanding reactivity and molecular properties as in the ground state. To gain insight into the nature of these effects we need to know which ground-state descriptors of aromaticity are transferable to excited states. One of the main difficulties that one encounters is that few descriptors of aromaticity can be easily employed to assess the aromatic character of excited states. Karadakov made the first calculations at the CASSCF level of magnetic aromaticity using NICS and other measures to describe the aromatic character of low-lying singlet and triplet excited states of benzene and related compounds. The results obtained were shown to be in line with Baird's rule. Among electronic-based descriptors only

Table 2 Values of FLU, Iring, and MCI for low-lying singlet, triplet, quintet, and septet states of C_6H_6 , C_4H_4 , and C_8H_8 at the B3LYP/6-311++G(d,p) level of theory. All units are in a.u. Reproduced from ref. 185 with permission of the Royal Society of Chemistry

	State	FLU	$I_{ m ring}$	MCI
C_6H_6	So	0.000	0.0478	0.0721
$D_{6\mathrm{h}}$	T_1	0.025	0.0028	-0.0015
011	Q_1	0.029	0.0011	0.0451
C_4H_4	S_0	0.104	0.0054	0.0101
$D_{ m 2h}$	T_1	0.012	0.0385	0.1271
C_8H_8	S_0	0.051	0.0244	-0.0005
$D_{ m 4h}$	T_1	0.001	0.0071	0.0271
	Q_1	0.029	0.0001	0.0013
	Septet ₁	0.033	0.0000	0.0178

 ELF_{π} has played a major role in studying the aromatic character of lowest-lying triplet states in fulvenes but no attempts have been done to quantify the electronic delocalisation in electronic states of higher energy.144 To bridge this gap, we proposed to generalise the use of electronic indices, PDI, FLU, I_{ring} , and MCI to study the aromaticity of a set of simple molecules in a number of excited states. 185 To this end, by means of DFT and CASSCF calculations, we studied the aromaticity patterns of the low-lying singlet, triplet, quintet, and septet excited states of benzene, cyclobutadiene, and D_{4h} cyclooctatetraene. The results of electronic indices based on DFT calculations are in agreement with the generalization of Baird's rule proposed by Soncini and Fowler to account for high-order multiplicities such as quintet and septet states. 186 That is, compounds with $(4n + 2)\pi$ -electrons that are aromatic in their lowest-lying singlet state should also be aromatic in their lowest-lying quintet state and antiaromatic in their lowest-lying triplet and septet states. On the contrary, molecules with $4n\pi$ -electrons are antiaromatic in their lowest-lying singlet and quintet states and aromatic in their triplet and septet states. These trends are perfectly followed by the MCI values summarised in Table 2, while FLU and $I_{\rm ring}$ show some discrepancies. However, CASSCF-based calculations do not assign an aromatic character to the lowestlying quintet state of benzene and the lowest-lying septet state of cyclooctatetraene. On the other hand, our calculations clearly showed the aromatic character of the vertical S2 and T1 states of cyclobutadiene and cyclooctatetraene in agreement with Baird's rule. This new methodology paves the way towards the possibility of studying the role of aromaticity in reactions occurring in the excited state.

6. Conclusions

Over the last few decades there has been a remarkable expansion in the number of different types of aromatic systems and in our understanding of aromaticity. The field of aromaticity is in constant evolution and the variety of molecules that present properties related to aromaticity is growing exponentially. It is our opinion that the field of aromaticity has been enriched

(and not cheapened as pointed out by Hoffmann 187 recently) by all these fascinating new aromatic compounds. One of the main difficulties faced by researchers interested in aromaticity characterization is the lack of a physical basis for this property, which makes its quantification difficult. Despite the great progress in the field over the last few years, the range of applicability of many descriptors is still limited to simple organic compounds. In this review, we have shown that indices based on electron delocalisation measures are all-round indicators of aromaticity that outperform most of the classical structural- and magnetic-based indices. Despite the success of these indicators, there is still room for improvement. Although the definition of the I_{NB} and I_{NG} indicators represented a big step forward in the correct direction, more efforts need to be carried out in order to extend the applicability of these indices. The major difficulty is describing aromaticity in large rings or in tridimensional circuits. Developers of electronic based indices should also work to enhance the scope of applicability towards larger and more complex systems. For example, systems like expanded porphyrins are still out of the scope of multicentre descriptors. Future challenges include the generalization of some of these tools to study the aromaticity of such intricate molecules and the possibility of obtaining both local and global measures in porphyrins, borane clusters, fullerenes or nanotubes.

Acknowledgements

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