

Understanding Radicals via Orbital Parities

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We introduce analysis of orbital parities as a concept and a tool for understanding radicals. Based on fundamental reduced one- and two-electron density matrices, our approach allows us to evaluate a total measure of radical character and provides spin-like orbitals to visualize real excess spin or odd electron distribution of singlet polyyradicals. Finding spin-like orbitals automatically results in their localization in the case of disjoint (zwitterionic) radicals and so enables radical classification based on spin-site separability. We demonstrate capabilities of the parity analysis by applying it to a number of polyyradicals and to prototypical covalent bond breaking.

INTRODUCTION

Importance of radicals for chemistry cannot be overestimated due to their role as reactive intermediates¹, magnetic² and optical properties³, and biological function⁴. Theoretical chemistry has achieved impressive results in understanding their structure and reactivity as well as in accurate calculation of their properties^{5,6}. At the same time, the general picture is still surprisingly non-uniform, leading to several widely used classifications^{7,8}.

The conventional definition of a radical as a system with unpaired electron(s) is intuitive, but barely applicable in cases where the electronic structure is not described by a simple valence-bond picture. On the other hand, a physical view based on quantum numbers of the electronic wave function, the total spin S^2 and its projection m_s , is not insightful for some systems. An iconic example of those are singlet diradicals^{5,7-9}. For them both quantum numbers are zero, suggesting a simple singlet molecule. Neither is spin density insightful, as it is zero everywhere¹⁰. However, spectroscopic and chemical properties of these molecules reveal radical behavior.

An ideal theoretical framework for understanding radicals should provide a numerical measure for radical (polyyradical) character of the molecule and a function for the unpaired electron distribution suitable for visualization. In addition, the insights into the ability of multiple radical sites to react independently would be a bonus as it helps classifying polyyradicals into disjoint and non-disjoint^{7,9}, also known as zwitterionic and covalent⁸, types.

There exist numerous valuable theoretical approaches for understanding radicals based on various properties: natural occupation numbers¹¹, configuration interaction coefficients^{12,13}, collectivity number¹⁴, hole-particle density¹⁵, distribution of unpaired electrons^{10,16,17}, natural¹⁸ or partial orbital occupations¹⁹. Many of these theories are conceptually involved and often based on specific or complicated wave-function types.

Here we propose an alternative framework for understanding radicals based on orbital parity, inspired by the theory of Mott insulators ("mottness")²⁰. Our approach is conceptually simple, not restricted to particular types or systems and wave functions, and offers a qualitative measure of the radical character. Moreover, our method provides a transformation to a spin-like orbital basis suitable for visualizing the unpaired

electron distribution. This basis may or may not be localized, thus, making it possible to distinguish between disjoint and non-disjoint polyyradicals. In addition, spin-like orbitals can be used to simplify post-processing quantum chemical calculations. There already exist successful theoretical approaches to radical chemistry based on orbital transformations²¹⁻²³. However, we argue that using the transformation based on orbital parity optimization demonstrates a number of lucrative features.

RESULTS AND DISCUSSION

The energy of the molecule in the normalized electronic state $|0\rangle$ is given as²⁴

$$E = \sum_{p,q} D_{pq} h_{pq} + \frac{1}{2} \sum_{pqrs} d_{pqrs} g_{pqrs} + h_{nuc}, \quad (1)$$

where indices p, q, r, s run over spatial orbitals; D_{pq} and d_{pqrs} are one-electron and two electron reduced density matrix (1-RDM and 2-RDM) elements in the molecular orbital (MO) basis, respectively; h_{pq} and g_{pqrs} are one-electron integrals and two electron integrals, respectively, and h_{nuc} stands for nuclei Coulomb repulsion. RDMs are defined as follows:

$$D_{pq} = \sum_{\sigma} \langle 0 | \hat{a}_{p\sigma}^{\dagger} \hat{a}_{q\sigma} | 0 \rangle, \quad (2)$$

$$d_{pqrs} = \sum_{\sigma\sigma'} \langle 0 | \hat{a}_{p\sigma}^{\dagger} \hat{a}_{r\sigma'}^{\dagger} \hat{a}_{s\sigma'} \hat{a}_{q\sigma} | 0 \rangle \quad (3)$$

In these equations σ, σ' correspond to the spin functions and run over spin-up, \uparrow , and spin-down, \downarrow , states, whereas $a_{p\sigma}^{\dagger}$ and $a_{p\sigma}$ are standard second-quantization creation and annihilation operators.

We now define the parity operator for orbital p using the number operators $\hat{n}_{p\uparrow} = \hat{a}_{p\uparrow}^{\dagger} \hat{a}_{p\uparrow}$ and $\hat{n}_{p\downarrow} = \hat{a}_{p\downarrow}^{\dagger} \hat{a}_{p\downarrow}$ (expectation values of these, $n_{p\uparrow}$ and $n_{p\downarrow}$, being spin-orbital occupations):

$$\hat{P}_p = (-1)^{\hat{n}_{p\uparrow} + \hat{n}_{p\downarrow}}. \quad (4)$$

The parity expectation value is calculated from RDMs:

$$P_p = \langle 0 | \hat{P}_p | 0 \rangle = 1 - 2D_{pp} + 2d_{pppp}. \quad (5)$$

$n_p \uparrow$ and $n_p \downarrow$ take values between 0 and 1. For single determinant methods, the spin-orbital occupations are exactly

TABLE I: Relation between spin-orbital occupation numbers and orbital parities for a single Slater determinant.

$n_{p\uparrow}$	$n_{p\downarrow}$	P_p
0	0	1
1	0	-1
0	1	-1
1	1	1

0 (empty) or 1 (occupied), whereas for more general, multi-configurational, wave functions they are non-integer. Orbital occupation is the sum of $n_p \uparrow$ and $n_p \downarrow$. From equation (4) we see that P_p takes values from -1 to 1. For the single-determinant wave function singly-occupied orbitals have $P_p = -1$, whereas for vacant or doubly-occupied ones $P_p = 1$ as listed in Table I. Thus, there is a clear mapping between orbital occupation number and its parity: half-filled orbitals are spin-like.

In the case of multiconfigurational wave functions, the simple mapping between occupations and parities does not generally hold. Singly occupied molecular orbitals (canonical, natural *etc.*) may not possess spin-like character. However, it may be possible to minimize the parity, bringing it close to -1 with an orbital transformation. This can be achieved by pairwise orbital rotations by an angle ϕ and satisfying the minimum conditions:

$$\frac{dP_p(\phi)}{d\phi} = 0, \quad (6)$$

and selecting the solutions corresponding to the parity minimum, where $P_p(\phi)$ is the orbital parity in the transformed molecular orbital basis. Therefore, within our framework **a radical is a system where one or more orbitals with $P_p = -1$ can be found** by orbital rotations. We will refer to such orbitals as **spin-like**. Diradicals have two spin-like orbitals, triradicals have three *etc.* In trivial cases such as those in Table I, the transformation from the canonical basis is a unit matrix.

In case of polyyradicals, parity optimization may result in orbital localization. **Spatial distribution of the spin-like orbitals provides a tool to classify radicals into the disjoint and non-disjoint classes:** in the former, the spin-like orbitals are localized on different sites.

For the general multiconfigurational wave functions required to capture electron correlation, **parities are not integer (see equation (5)) and can naturally serve as measures of radical character.** As opposed to some criteria introduced for diradicals $P_p(\phi)$ are a general criterion, *i. e.* applicable to all radical types.

RDMs, available from standard quantum chemistry software, are needed to perform the spin-mapper orbital transformation that searches for spin-like orbitals. The latter has been derived and implemented by us and is described in more detail in the Appendix.

As a first example, we apply the orbital parity formalism to singlet diradicals⁷⁻⁹. We consider benzene isomers a pyridine-based analogue of p-benzene²⁵ shown in Figure 1.

Orbital parity analysis of the CASSCF(8,8)²⁶ wave functions of o- and m-benzynes reveals a weak diradical character with minimum parities of -0.5 and -0.3, respectively (see Figure 8). This is to be expected due to close proximity of the supposed unpaired electrons, which are able to couple with each other contributing to a triple bond²⁷. In contrast to this, p-benzene and its analogue are proper diradicals with two spin-like orbitals each ($P \approx -0.9$).

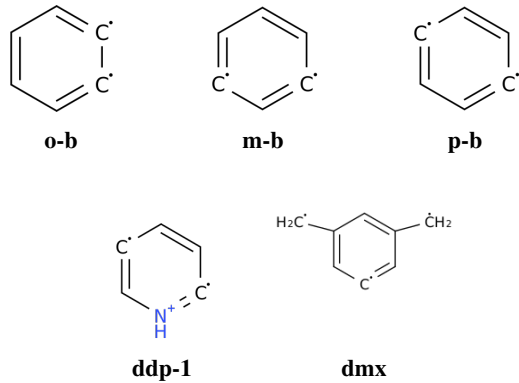


FIG. 1: Chemical structures of the considered radicals: **o-b** - ortho-benzene; **m-b** - meta-benzene; **p-b** - para-benzene; **ddp-1** - 2,5-didehydropyridinium cation; **dmx** - 5-dehydro-m-xylylene.

For p-benzene, parity optimization results in the spin-like orbitals being a symmetric and antisymmetric combination of the canonical HOMO and LUMO. This is to be expected for the multi-configurational wave function dominated by the two configurations: doubly-occupied HOMO with empty LUMO and *vice versa*. Whereas canonical orbitals are delocalized, spin-like ones are localized at different carbon atom as shown in Figure 3a indicating that p-benzene is a disjoint (zwitterionic) diradical. This transformation (45° rotation) is a well-known result of generalized valence-bond (GVB) theory^{7,28,29}. Although the canonical HOMO and LUMO in p-benzene have occupation numbers close to one, their parities are not spin-like (see Figure 3a).

Pyridine-based diradical is less symmetric than p-benzene, which results in a more general spin-mapper transformation: rotation angle different from 45°. GVB-transformed orbitals are similar to the spin-like ones, but exhibit non-optimal parities (see Figure 3b).

As shown in Figure 8, parity analysis generally agrees with the odd electron distribution^{10,16,17} results for overall diradical character, whereas Yamaguchi's function¹¹ underestimates it. Importantly, the odd electron distribution function $u(\mathbf{r})$ (see the Appendix) shows similar trends as the two spin-like orbitals, but does not provide information on the separability of the radical sites.

We go on by applying the parity-based approach to a triradical, 5-dehydro-m-xylylene, or **dmx** (see Figure 1)³⁰. This remarkable molecule was the first organic compound to known to violate Hund's rule as it has an open-shell doublet rather than quartet ground state³¹. Our state-averaged CASSCF(9,9) calculation confirms this, revealing three almost perfectly

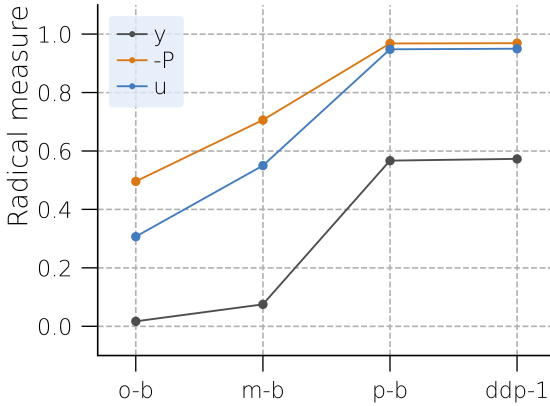


FIG. 2: Radical measures (parities, odd electron distribution functions, and diradical characters) for benzenes and analogues: $-P$ – negative parity; u – odd electron density; y – Yamaguchi’s diradical character. $-P$ is the average over the two spin-like orbitals (are identical due to symmetry). u is averaged over the two canonical frontier orbitals.

spin-like orbitals in the ground state shown in Figure 4b. The sp^2 -hybrid one localized on the benzene ring (rightmost) has the unpaired electron on it, and is identical to the canonical orbital (4a) with perfect parity of -1. The other two spin-like orbitals localize (although not completely) on the different methylene groups. These two orbitals are to a large extent (weight of ca. 0.8) linear combinations of the two symmetric delocalized canonical orbitals (see Figure 4a). The latter already exhibit almost spin-like parities of ca. -0.9. If those parities were optimal, the ground state of **dmx** would be a non-disjoint doublet diradical. However, parity optimization results in localization on the different sites, providing evidence that the ground state is rather a disjoint doublet diradical. The lowest triplet state exhibits very similar localized spin-like orbitals as the one with the same perfect parities, revealing its disjoint character.

An archetypal case of radical emergence is homolytic dissociation of a covalent bonds²⁴. Whereas around the equilibrium bond distance a singlet molecule has little if any radical character, the two separated fragments constitute a perfect disjoint diradical with singly-occupied orbitals localized on each side. This is illustrated with our parity-based approach for dilithium (Li_2) dissociation computed with a simple CASSCF(2,2) wave function (see Figure 5). Modestly negative parities of the two active orbitals at around the equilibrium separation (slight diradical character) abruptly fall down in the dissociation region, reaching the limiting value of -2 at bond distance larger than 6 Å. The energy converges to its plateau at approximately the same distance. Interestingly, the minimum parity is achieved for localized 2s-orbitals (obtained from the canonical ones by GVB transformation) at all interatomic distances.

Next, we consider the evolution of polyradical character in linear oligoacenes with the increase in the number of benzene

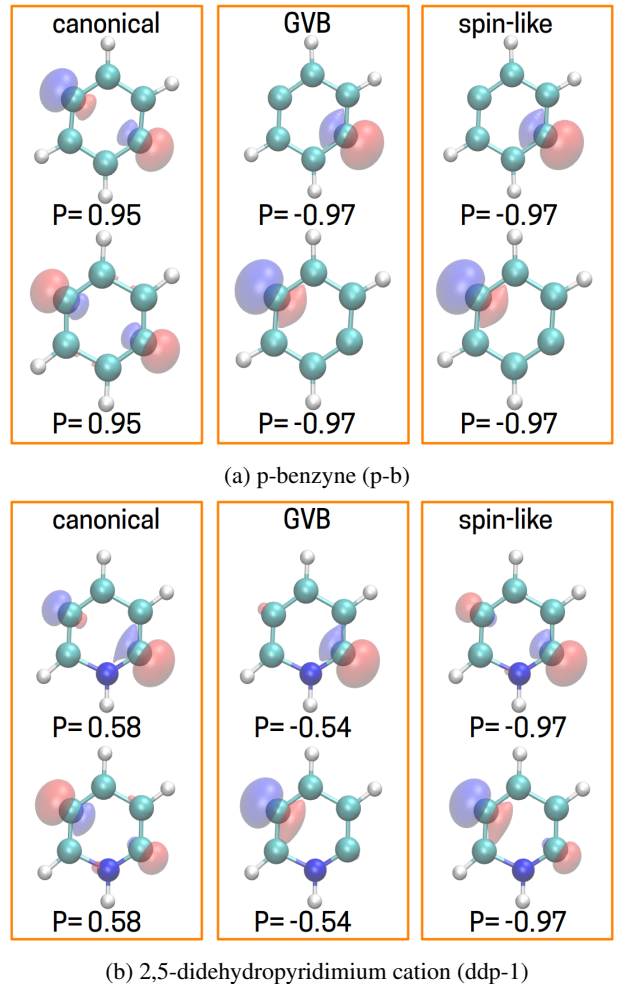
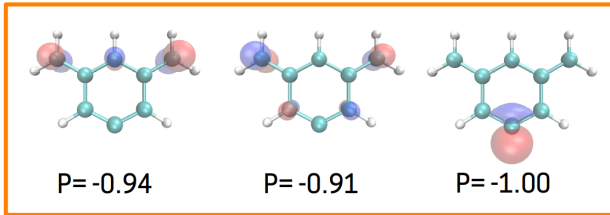


FIG. 3: Molecular orbitals of diradicals and their parities: canonical, generalized-valence-bond (GVB) transformed, and spin-like.

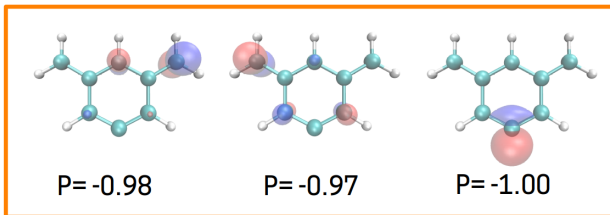
rings³². It is instructive to analyze qualitative simple CASCI wave functions in the minimum active space (2,2) including only the HOMO (ϕ_1) and the LUMO (ϕ_2) for the two lowest singlet states. These calculations reveal a gradual increase in the measures of radical character for the ground state (see Figure 6) along the series.

For decacene, almost ideal diradical orbital parities of -1 are reached, its wave function being $\Psi_0 = 1/\sqrt{2}(|\phi_1\bar{\phi}_1\rangle - |\bar{\phi}_2\phi_2\rangle)$, where the bar denotes the opposite spin. The weight of the excited configuration (second term) in the smaller molecules is less than 0.5. The excited state of all homologues is an open-shell singlet and thus a perfect diradical: $\Psi_1 = 1/\sqrt{2}(|\phi_1\bar{\phi}_2\rangle - |\bar{\phi}_1\phi_2\rangle)$.

Whereas in the excited state delocalized HOMO and LUMO are spin-like, in the ground state minimum parity is achieved for the GVB transformed-orbitals. These orbitals are localized on the opposite fringes of the oligoacene along the short axis (see Figure 6, top). Therefore, the ground state of decacene is a disjoint (zwitterionic) singlet diradical, whereas its excited state (and that of all homologues) is a non-disjoint



(a) Canonical orbitals



(b) Spin-like orbitals

FIG. 4: Molecular orbitals of the 5-dehydro-m-xylylene (dmx) triradical and their parities: canonical and spin-like.

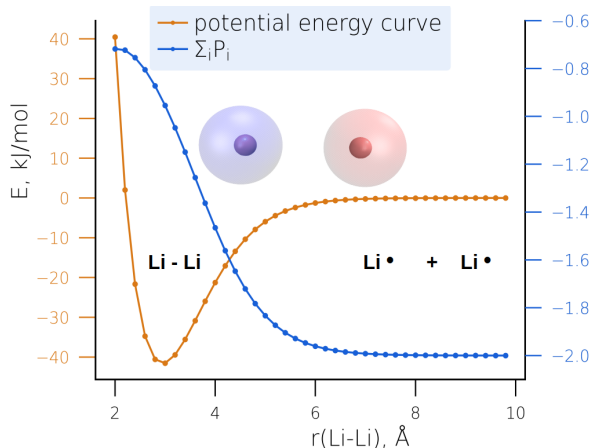


FIG. 5: Dissociation of dilithium, Li_2 . Potential energy curve and sum of the two active orbitals' parities for the singlet ground state. Optimal-parity orbitals (centre) at atomic separation of 5 Å are localized on lithium atoms: red - positive; blue - negative.

(covalent) singlet diradical.

Interestingly, for decacene the 1-RDM, \mathbf{D} for the two states are almost identical, yielding identical odd electron distributions \mathbf{u} (computed as $\mathbf{u} = 2\mathbf{D} - \mathbf{D}^2$)^{17,33} shown in the Appendix. This approach reveals a proper spatial odd electron distribution $u(\mathbf{r})$, but does not provide insights into zwitterionic/covalent characters of diradicals.

A more realistic state-averaged CASSCF(4,4) calculation of decacene with HOMO-1 and LUMO+1 included in the active space reveals a more complex picture. As expected, the radical character of the frontier-orbital subspace is reduced: the corresponding parities are -0.85 for the ground state and -0.81 for the previously considered excited state Ψ_1 . At

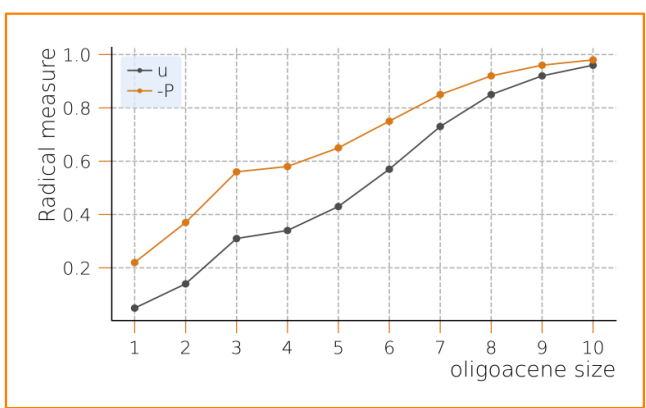
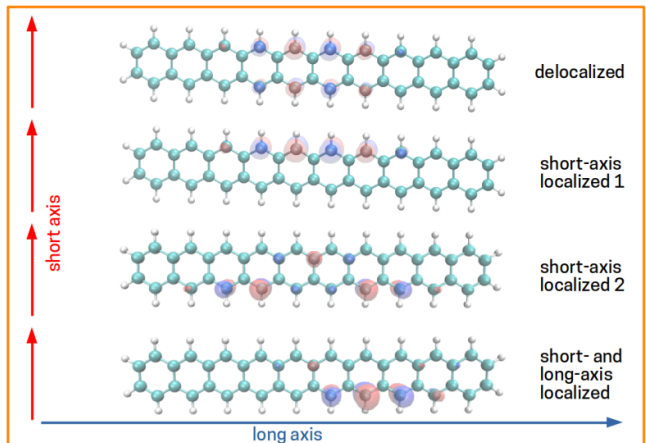


FIG. 6: Linear oligoacenes. Top: orbitals of decacene with considerable spin-like character in several electronic states from the CASSCF(4,4) wave function. Bottom: radical measures (parities and odd electron distributions functions) for the ground state from the CASCI(2,2) wave function, shown for each of the two most spin-like orbitals.

the same time, near frontier orbitals (HOMO-1 and LUMO-1) contributes to the total radical character: spin-like orbitals in this subspace reach parities of -0.35 and -0.15 for the ground and excited states, respectively. Remarkably, for the two states the most spin-like orbitals in the near-frontier space are delocalized (short-axis localized) if their counterparts in the frontier space are short-axis localized (delocalized).

The state-averaged CASSCF(4,4) wave function reveals another excited state with the four most spin-like orbitals ($P = -0.65$) localized along both short and long axis (see Figure 6), *i.e.* in the "corners" of the molecule. Those are obtained by mixing all four active orbitals.

Similar localization patterns and evolving polyyradical character in various states of oligoacenes have been shown by Yang *et al.*³⁴ by manual wave function analysis, whereas spin mapper approach arrives at these results automatically. Despite conceptual simplicity the actual orbital transformation matrices are non-trivial (see the SI) due to the complicated nature of state-averaged CASSCF wave function and can barely

be reproduced by hand.

CONCLUSION

We have introduced a theoretical approach to understanding radicals based on orbital parities, P . For single-determinant wave functions $P = -1$ for the singly-occupied and $P = 1$ for the doubly-occupied orbitals. Thus, there exist a one-to-one correspondence between occupation numbers and parity. For general multiconfigurational wave functions this is not the case. Therefore, the spin-like orbitals are obtained via the orbital transformation minimizing their parities (spin-mapper transformation) and can be different from canonical and natural orbital. Values of P and the quantity of spin-like orbitals provide qualitative measures of radical character. Visualizing spin-like orbitals proves useful to understand spatial distribution of the unpaired spins, whereas their localization properties reveal the radical type (disjoint/non-disjoint).

We have demonstrated the power of the methodology quantifying radical character of a number of complex di- and polyyradicals in several electronic states of different multiplicity, revealing their localization type and visualizing spin-like orbitals. We believe that the orbital-parity approach to radical chemistry, is simple, general, and versatile, being based on fundamental quantities (reduced density matrices) and versatile. It can contribute to the understanding of fundamental radical chemistry by means of theory and provides a useful tool for quantum chemistry practitioners.

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CONFLICT OF INTEREST

The authors declare no conflicts of interest.

Appendix A: Parity optimization

We apply an iterative procedure to determine the spin-like orbital basis in which the orbital parities are extremal. For this we perform a sequence of unitary pairwise rotations by an angle θ of the fermionic operators:

$$\begin{aligned}\hat{a}_{q\sigma} &= \cos \theta \hat{a}_{i\sigma} + \sin \theta \hat{a}_{j\sigma}, \\ \hat{a}_{p\sigma} &= -\sin \theta \hat{a}_{i\sigma} + \cos \theta \hat{a}_{j\sigma},\end{aligned}\quad (\text{A1})$$

with the same rotation being performed for the hermitian conjugates of the operators. From the reduced density matrices we can compute the parity of an orbital ϕ_p , which results from the linear combination of orbitals ϕ_i and ϕ_j , as $\langle P_p \rangle_0(\theta)$. The orbital parity $\langle P_p \rangle_0(\theta)$ is an analytic, 2π -periodic function of the rotation angle θ . We find extremal points θ_n of the function $\langle P_p \rangle_0(\theta)$ in the domain $\theta \in [0, 2\pi]$ from

$$\frac{d\langle P_p \rangle_0}{d\theta} \Big|_{\theta_n} = 0, \quad (\text{A2})$$

and select solutions θ_n that satisfy

$$\frac{d^2\langle P_p \rangle_0}{d\theta^2} \Big|_{\theta_n} \neq 0. \quad (\text{A3})$$

The analytic expression for the derivatives of the function derived and implemented.

Appendix B: Computational details

CASSCF³⁵ calculations were carried out with the **PySCF** package version 2^{36–38}. All calculations were done using triple- ζ Def2-TZVP basis set (with the exception of oligoacenes, for which a smaller def2-SVP basis was used)³⁹ with the default auxillary density-fitting basis. The active space of the CASSCF calculations was comprised of all π -orbitals and the non-bonding orbitals. The initial orbitals used for CASSCF calculations were UHF natural orbitals²². The **Geometric** software package⁴⁰ has been used for geometry optimization. All the structures except oligoacenes were optimized for the ground-state using the corresponding CASSCF method.

Appendix C: Odd electron distributions for selected examples

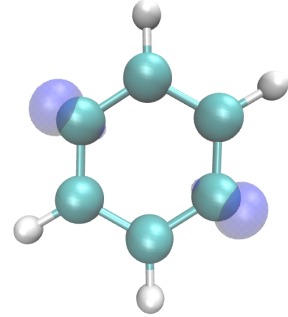


FIG. 7: Odd electron distribution for p-benzene from the CASSCF(8,8) wave function.

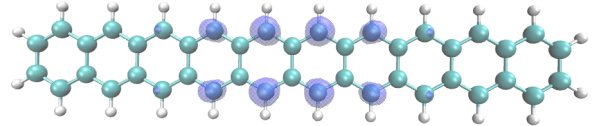


FIG. 8: Odd electron distribution decacene from the CASCI(2,2) wave function: the function is identical for the ground and excited singlet states.

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