




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A natural scheme for the quantitative analysis of the magnetically induced molecular current density using an oriented flux-weighted stagnation graph. I. A minimal example for LiH

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A new natural scheme is introduced to analyze quantitatively the magnetically induced molecular current density vector field, \mathbf{J} . The set of zero points of \mathbf{J} , which is called its *stagnation graph* (SG), has been previously used to study the topological features of the current density of various molecules. Here, the line integrals of the induced magnetic field along edges of the connected subset of the SG are calculated. The edges are oriented such that all weights, *i.e.*, flux values become non-negative, thereby, an oriented flux-weighted (current density) stagnation graph (OFW-SG) is obtained. Since in the exact theoretical limit, \mathbf{J} is divergence-free and due to the topological characteristics of such vector fields, the flux of all separate vortices (current density domains) and neighbouring connected vortices can be determined exactly by adding the weights of cyclic subsets of edges (*i.e.*, closed loops) of the OFW-SG. The procedure is exemplified by the minimal example of LiH for a weak homogeneous external magnetic field, \mathbf{B} , perpendicular to the chemical bond. The OFW-SG exhibits one closed loop (formally decomposed into two edges), and an open line extending to infinity on both of its ends. The method provides the means of accurately determining the strength of the current density even in molecules with a complicated set of distinct vortices.

studies of the molecular magnetic response are highly relevant not only from a theoretical perspective.

In molecular magnetic response theory, the induced electronic current density, \mathbf{J} , is the key quantity from which all other magnetic response properties can be calculated in a quasi-classical fashion by evaluating the expectation value integrals. Hirschfelder has coined the term *subobservable* for quantities for which it is possible to define a formal quantum mechanical operator.¹ A series of reviews on the current state of research on magnetically induced molecular current densities is available.^{2–5} One branch of this research field is concerned with both the topological and the quantitative characterization of \mathbf{J} , an undertaking that is analogous to the topology analysis of the electron density based on the quantum theory of atoms in molecules (QTAIM).^{6,7} The \mathbf{J} field typically consists of multiple vortices, some embedded in each other. However, their boundary surfaces may never cross due to charge conservation. These surfaces are known as separatrix surfaces. Topological characterizations of the induced molecular current density field are fairly abundant in the literature^{6,8–26} and analogies between the QTAIM analysis and the SG graph analysis have been discussed.³ Unlike for bond paths in QTAIM, there are no established empirical rules or simple models of interpretation for the topology of the magnetically induced current density, yet. Large unsymmetric molecules in general do not seem to possess a connected stagnation graph, but rather a collection of isolated stagnation points (see below).

We have recently reported some progress on the quantitative characterization of the current-density field²⁷ from the secondary magnetic field that it induces, \mathbf{B}_{ind} , better known as the nucleus-independent chemical shift (NICS), which can be derived from the chemical shielding tensor, σ , by contracting it with the external magnetic field.²⁸ The obtained data are used, for example, to rationalize the aromatic or antiaromatic nature of a compound. For the quantification of the molecular current density, one is typically interested in its flux, Φ , through a particular surface, \mathcal{S} , chosen according to chemical intuition or other demands or model ideas, such as the “ring-current” model for molecular systems² or the integration based on the

1 Introduction

Practically all relevant physical changes in molecules are ultimately triggered by the electromagnetic force since the other three fundamental interactions, the weak, the strong and the gravitational interaction are effectively irrelevant to chemistry. Molecules are restricted to interacting with their environment *via* electronic or magnetic perturbations (that are not necessarily weak). Therefore,

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zero-flux electron density surfaces used in QTAIM.^{29,30} The idea underlying our recent work²⁷ is to apply the integral form of the Ampère–Maxwell law by evaluating a line integral of \mathbf{B}_{ind} instead of evaluating a surface integral of \mathbf{J} . The flux is then obtained by line-integration of $\mu_0^{-1}\mathbf{B}_{\text{ind}}$ over the boundary line $\partial\mathcal{S}$ of surface \mathcal{S} ,

$$\Phi_{\partial\mathcal{S}} = \iint_{\mathcal{S}} \mathbf{J} \cdot d\mathbf{s} = \frac{1}{\mu_0} \oint_{\partial\mathcal{S}} \mathbf{B}_{\text{ind}} \cdot d\boldsymbol{\ell}. \quad (1)$$

In this work, we employ the simple example of the lithium hydride molecule to illustrate in simple terms the essence of our method and show how this method can be naturally extended by applying it to the stagnation graph (*i.e.*, the set of zero points) of \mathbf{J} , such that the quantification of separate current vortices becomes exact, simple and possible to automate.

Analysis of the magnetically induced current density is done routinely for various molecules, including compounds of potential practical application such as porphyrinoids. Their current density field consists of multiple pathways, the strength of which is difficult to determine using the standard approach of placing an integration plane through the molecule and calculating the strength of the flux across the plane. In our scheme, one first needs to obtain the stagnation graph (SG) of the molecule and then perform the integration along the points of the SG as described in Section 2. Hence, it becomes possible to precisely characterize the magnetically induced current density in molecules where the vortices are hard to identify and discern.

This work is, furthermore, intended as the first of a series, where we outline the new method by increasingly complex examples. The upcoming works in the series are planned to contain studies on II. homo- and heterodiatomic molecules, III. Benzene, IV. Neutral and charged $[\text{C}_n\text{H}_n]^m$ type annulenes, V. Condensed ring systems and porphyrins, VI. Selected organo-metallic molecules. Work on them is already under way.

2 Results and discussion

We exemplify our proposed method with the lithium hydride molecule since this gives, to the best of our knowledge, the simplest case of a stagnation graph containing a closed loop and an open line. Thus, it is the simplest case of a molecule that shows two physically distinct (separable) vortices. The topology of the magnetically induced current density field, \mathbf{J} , of LiH has been discussed previously in great detail by Stevens and Lipscomb (1964),³¹ Keith and Bader (1993),⁷ and later by Pelloni, Lazzarotti and Zanasi (2009),¹⁸ so we will only give a summary here. General procedures to compute stagnation points and stagnation lines have been published, as well, and are freely available.^{9,10} Placing \mathbf{B} parallel to the z axis and perpendicular to the Li–H bond (lying along the y direction) results in a \mathbf{J} field composed of exactly two separate current vortices, each possessing a central stagnation line (see Fig. 1). Both vortices are separated from each other by a single closed surface \mathcal{X} of the exact topology and the approximate geometry of a sphere. The spherical vortex domain (inner/toroidal vortex) is completely enclosed in the global domain, extending

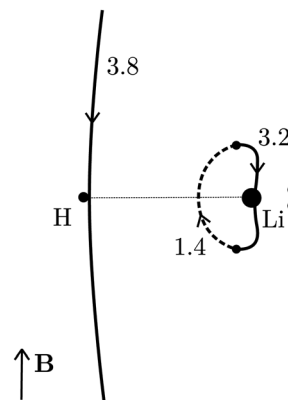


Fig. 1 The oriented flux-weighted current density stagnation graph (OFW-SG) of LiH. Adding weight values of line segments which form loops yield exactly all physically uniquely defined vortex fluxes (ring currents) in a molecule. For example, $1.4 + 3.2 = 4.6 \text{ nA T}^{-1}$ which is the total toroidal current flux in LiH. The total current of the hydride vortex is 3.8 nA T^{-1} . There are no other uniquely defined current fluxes in this orientation of the magnetic field. All possible results of a full quantitative current density analysis are, thus, combined in this flux-weighted stagnation graph.

over the entire molecular space (outer/hydride vortex). The domains are illustrated as streamlines in Fig. 2.

The outer vortex has an open stagnation line, ℓ_1 , extending from $z = -\infty$ to $z = \infty$ and lying in the (y, z) plane. Line ℓ_1

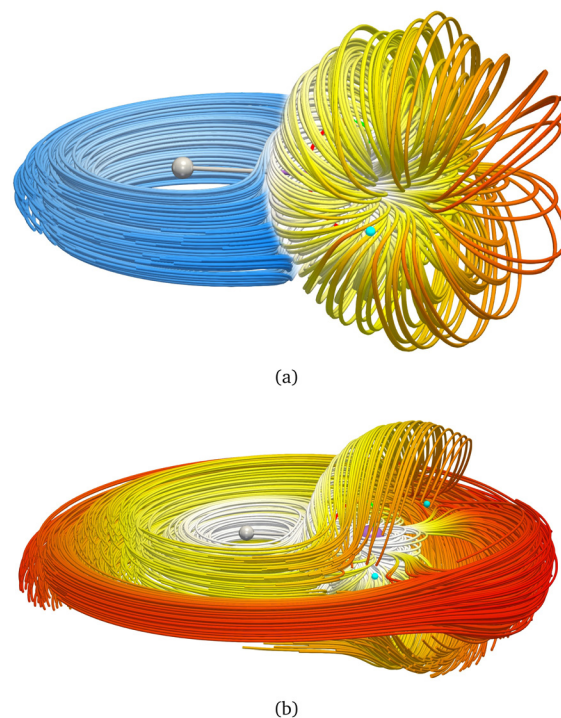


Fig. 2 A streamline representation of the magnetically induced current density field, \mathbf{J} , of LiH. (a) There are two domains, an inner vortex belonging to the lithium atom (illustrated in yellow) that is surrounded by a spherical surface to isolate it from the outer vortex surrounding the hydrogen atom and extending around the whole molecule (shown in blue). (b) The \mathbf{J} field of LiH. A logarithmic colour scale has been employed in both figures such that white is the highest current density.



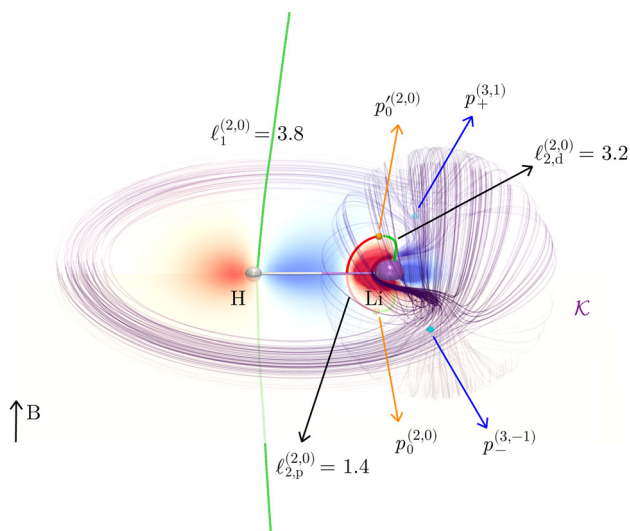


Fig. 3 The magnetically induced current density field \mathbf{J} of LiH in an external magnetic field, \mathbf{B} , perpendicular to the bond, contains a set of stagnation points (= zero points): the vortex critical lines ℓ_1 , $\ell_{2,p}$, $\ell_{2,d}$ (in green or red) of Collard–Hall index (2,0). The stagnation points p_0 and p'_0 (in orange) are arbitrarily selected points where the two parts (shown in green and red) of the closed stagnation line join. They are not topological (0,0) branch critical points. The two isolated torus or source- and sink-critical points p_+ and p_- with Collard–Hall index (3,1) and (3,-1) (*vide infra*), respectively, are shown in cyan. Streamlines that are closely passing the torus critical points are embedded in a topological and geometrically approximate spherical surface \mathcal{K} , which separates the main molecular diatropic vortex around ℓ_1 from the toroidal vortex around $\ell_2 = \ell_{2,p} \cup \ell_{2,d} \cup \{p_0, p'_0\}$. The flux Φ_{ℓ_1} in the main vortex amounts to 3.8 nA T⁻¹ while the toroidal vortex flux $\Phi_{\ell_2} = 4.3 (= 3.2 + 1.4)$ nA T⁻¹. The current density flux flowing away from the viewer is illustrated in red, while the opposite flow is shown in blue.

passes very close to the H atom, only a fraction of an atomic unit, and bends towards it. Above and below the (x, y) plane, ℓ_1 is bending slightly towards the center of the LiH molecule but straightens out at larger $\pm z$ heights. The current vortex around this stagnation line ℓ_1 is diatropic, thus, according to our convention, the vortical flow is clockwise if the external field \mathbf{B} is pointing in the z direction and the reader looks towards negative z . The approximately spherical inner vortex domain can be seen as inserted in-between the streamlines of the outer vortex which bypass this domain similarly to how a laminar-flowing fluid would pass around a ball fully submerged into it. This results in two isolated “toroidal stagnation points”[†] $p_+^{(3,1)}$ and $p_-^{(3,-1)}$ on the separatrix sphere, \mathcal{K} , where the outer flow diverges in (and converges from) all directions on \mathcal{K} , respectively.

The inner domain encloses the Li atom, and its radius extends roughly to the midpoint between the two atoms. The geometrical center of \mathcal{K} does not lie on the Li atom, but instead, it is significantly shifted away from the H atom. The stagnation line, ℓ_2 , of the current-density vortex is closed, which gives rise to a toroidal flow with a (metaphorically speaking) “reflux” region in the middle. On the side towards the H atom, the flux resembles that of a paratropic vortex, while on the remote side of the Li atom, the flux acts as if it were

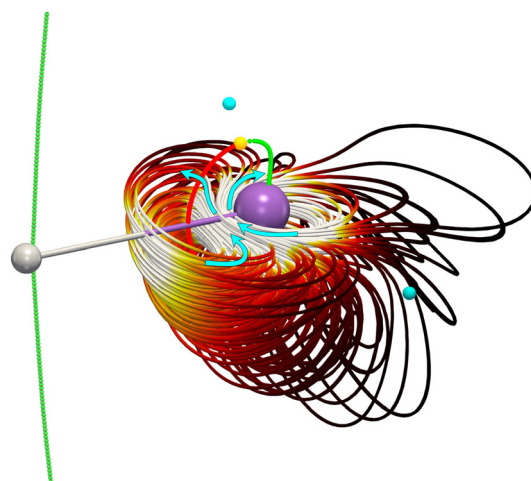


Fig. 4 The reflux region in the toroidal vortex of the magnetically induced current density field \mathbf{J} of LiH. The current flux of this vortex integrates to 4.6 nA T⁻¹, as can be easily derived from Fig. 3 by adding the weights of branches $\ell_{2,p}$ and $\ell_{2,d}$.

diatropic. The respective parts of the stagnation line are illustrated in red and green in Fig. 3 and 4. We define the two somewhat arbitrary stagnation points p_0 and p'_0 (illustrated in orange) where the apparent diatropic and paratropic parts of the critical stagnation line meet as the “branching points” of the closed loop. However, they are not (0,0) branch critical points in the topological sense. The streamline representations show that the current-density flow in the middle region is similar to the water circulating in a waterspout fountain with an inner tube (illustrated using cyan arrows in Fig. 4). The inner stream passes close to the Li atom, roughly perpendicular to both \mathbf{B} and the Li–H bond, lying in the (x, y) plane. It has the shape of a double S and contains a heteroclinic streamline line which connects the source critical point $p_+^{(3,1)}$ with the sink critical point $p_-^{(3,-1)}$ also inside \mathcal{K} . Critical points can be classified using the Collard–Hall indices (r, s) .[‡] Points p_- and p_+ are illustrated in cyan in Fig. 3 and 4.

The stagnation line of the inner vortex, ℓ_2 , is a topological circle (a closed loop) that we split into two branches, $\ell_{2,d}$ (with diatropic current vortices in the vicinity) and $\ell_{2,p}$ (with paratropic vortices in the vicinity), connected by the “branching points” p_0 and p'_0 illustrated in orange. Strictly speaking, this division is arbitrary, and ℓ_2 is actually one unbranched topological domain. Nevertheless, this scheme provides a simple example of the general method for determining the current flux that we are outlining here. We also note that this type of toroidal current vortex is, obviously, neither diatropic nor paratropic over its whole domain. Due to charge conservation, no current can pass from one vortex into or out of another one. In LiH, this means that the inner and the outer domains are perfectly separated by surface \mathcal{K} . Hence, a total current flux can be defined for each individual vortex. As a chemical interpretation, one would assign the diamagnetic, spatially unconfined current vortex around ℓ_1 to the hydride anion, which dominates the response and encloses a spatially confined toroidal vortex arising from the Li⁺ cation.



As we have shown previously,²⁷ and since \mathbf{B}_{ind} is vanishing at infinity in the z and y directions, the total current flux, Φ_{ℓ_1} , of the outer vortex can be obtained from the line integral

$$\Phi_{\ell_1} = -\frac{1}{\mu_0} \int_{\ell_1} \mathbf{B}_{\text{ind}} \cdot d\ell. \quad (2)$$

Here we have chosen the direction of stagnation line ℓ_1 such that Φ_{ℓ_1} becomes positive. In this way, every current–density stagnation graph can be uniquely oriented to yield a directed graph. Furthermore, to each “edge” (line segment) of a stagnation graph, a flux integral can be assigned, such that an oriented edge-weighted graph with strictly positive weights is obtained.

To obtain the current flux of the toroidal current inside \mathcal{K} , one can make use of the notion that the total flux is passing through the D-shaped closed stagnation line ℓ_2 (composed of $\ell_{2,d}$ and $\ell_{2,p}$). Thus,

$$\Phi_{\ell_2} = -\frac{1}{\mu_0} \oint_{\ell_2} \mathbf{B}_{\text{ind}} \cdot d\ell \quad (3)$$

$$= -\frac{1}{\mu_0} \left(\int_{\ell_{2,d}} \mathbf{B}_{\text{ind}} \cdot d\ell + \int_{\ell_{2,p}} \mathbf{B}_{\text{ind}} \cdot d\ell \right) \quad (4)$$

$$= \Phi_{\ell_{2,d}} + \Phi_{\ell_{2,p}}, \quad (5)$$

where again an orientation is obtained for each integral associated with $\ell_{2,d}$ and $\ell_{2,p}$. A simple numerical integration scheme has been applied (see Appendix B for details) through which we have obtained $\Phi_{\ell_1} = 3.8$, $\Phi_{\ell_{2,d}} = 3.2$ and $\Phi_{\ell_{2,p}} = 1.4$ nA T⁻¹. The corresponding surface integrals of \mathbf{J} over the (y, z) plane in the region left of ℓ_1 and over the surface enclosed by ℓ_2 yielded 4.0, and 4.6 nA T⁻¹, respectively, both in very good agreement with the corresponding values of the line integrals (3.8 nA T⁻¹ and 3.2 + 1.4 = 4.6 nA T⁻¹). Details about these computations are given in the appendix.

Any field \mathbf{J} partitioned into vortices separated by asymptotic surfaces (separatrices), *i.e.*, the field is decomposable into pairwise disjoint sets, has cyclic stagnation subgraphs defining the vortex, such as the pair of stagnation lines ℓ_1 and ℓ_2 in the example of the toroidal vortex in LiH. Setting this cyclic subgraph to $\partial\mathcal{S}$ in the line integral in eqn (1) gives the current flux, Φ , in this vortex. Then, in order to obtain the flux of a separate vortex, one only has to add the weights of the corresponding cyclic subgraph in the correct orientation (*i.e.*, a plus or minus sign).

Regarding the efficiency and feasibility of the proposed method, there are the following advantages:

- Instead of surface integrals, only line integrals have to be calculated, which roughly leads to scaling of order $\mathcal{O}(N)$ versus $\mathcal{O}(N^2)$.

- For integrals of the current flux for which the stagnation lines are (approximately) known, *e.g.*, the central axis in planar (approximately) symmetric rings, and which yield the total “ring” current,²⁷ no current density has to be calculated

directly but instead only the typically simpler chemical shielding tensors have to be computed.

- With this method, there is no ambiguity concerning the precise definition and localization of the vortices under consideration, which is in stark contrast to the commonly used schemes which rely on explicit or implicit topological knowledge that very often was present only in rudimentary form. So it seems highly likely that many of the previous studies where surface integrals of the current density have been calculated based on a casual inspection of streamline graphs or various *ad hoc* integration schemes are flawed due to ill-defined current domains.

On the contrary one has to face two methodological disadvantages using this method:

- For all but the simplest cases (total current in symmetric ring systems), the SG has to be determined. As outlined above, we suggest that this is only an ostensible disadvantage since integration without knowledge of the topology is at risk of being erroneous, particularly if not at least investigated visually.

- For vortices with open stagnation lines (such as the hydride vortex in our example), one has to integrate relatively far out (*i.e.*, approximately 100 or even 200 atomic units) into the shielding cone of the molecule in order to achieve convergence (see appendix C for one numerical and one analytical example). This is a consequence of the non-local nature of the shielding contributions with respect to the underlying \mathbf{J} field.

- As shown previously,¹⁷ only relatively small or highly symmetric molecules possess non-trivially connected stagnation subgraphs. Hence, only in such cases, the current–density field can be partitioned into physically well-defined separate (pairwise disjoint) current–density vortices. Thus, only in these cases, it is possible to unambiguously define flux integrals (other than the total molecular flux). This leads to the conclusion that concepts like “bond-currents”, “local ring currents”, or “semi-local ring currents” can actually not be well-defined for molecules without non-trivially connected stagnation subgraphs. Their definition would either require—to the best of our knowledge—a yet unknown unambiguous physical definition, or it should be completely abandoned until we arrive at a deeper understanding of the general topology of molecular current–density fields (a subject on which we are continuously working).

3 Conclusion and outlook

The presented approach effectively gives a complete analysis of all magnetically induced current–density vortices in a molecule. This can be achieved by evaluating the line integral from eqn (1) for each “edge” of the stagnation graph taken with the appropriate sign, *i.e.*, orienting it accordingly. The resulting OFW-SG can be condensed into a schematic diagram, as shown in Fig. 1. In principle, the strategy can be generalized to non-planar and non-cyclic molecules. Hence, the analysis can be



carried out for any molecule with a non-trivially connected stagnation graph.

Further research is needed to address cases which do not possess a non-trivially connected stagnation graph. This probably applies to the vast majority of molecules. However, one way to deal with this problem is to restrict the analysis to the pseudo-SG,¹³ where the z component of the induced magnetic field is set to zero, or related methods. Thereby, the topology of the current density field tends to become significantly simpler and usually gets partitioned into smaller separate vortices (current density domains). This comes at the cost that in such fields, the continuity condition is violated, which, strictly speaking, implies that the usual topological tools (such as the Poincaré–Hopf theorem⁸) are inapplicable. Apart from the usage of pseudo-SGs, we need to arrive at a deeper understanding of the topology of molecular current–density fields for the cases where the SGs are just a collection of isolated points. Ultimately, in this way we could bridge the gap between the intuitive ring-current model and the actual physical current–density field of molecules in general. From our point of view, this is a necessary requirement to give a physically meaningful and mathematically sound definition to any of the traditionally applied schemes for integration of the current density flux. Otherwise, the integration surfaces commonly used in current density analysis studies remain ambiguous and arbitrary to some extent.

Data availability statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Author contributions

RJFB: conceptualization, investigation, supervision and writing—original draft; MD: funding acquisition, data curation, formal analysis, investigation, software, visualization and writing—review & editing (according to the <https://casrai.org/credit/CRediTscheme>).

Conflicts of interest

There are no conflicts to declare.

Appendix A: quantum chemistry calculations

The quantum chemistry calculations were performed using Gaussian 16 Rev. C01.³⁴ The geometry was optimized at the density functional theory level, employing the B3LYP functional^{35,36} and the aug-cc-pVTZ basis set.³⁷ The nuclear magnetic shielding tensors were calculated using the continuous set of gauge transformation (GSGT) approach.^{7,38} The current density and the nuclear magnetic shielding tensors in specific points were calculated with the SYSMOIC program.¹⁰ It also provided the stagnation graph of the molecule. Two more basis sets from the Dunning family were

employed, the quadruple- ζ and the quintuple- ζ versions, to evaluate basis-set convergence of the integrated strength of the current density.

The current–density field can be inspected visually on a slice or using streamlines. In this work, we have visualized the results by obtaining the current density on a three-dimensional (3D) grid using the gauge-including magnetically induced current (GIMIC) method^{3,4,39–41} which is interfaced to quantum chemistry program packages such as Gaussian,⁴² and employing the Paraview⁴³ program to prepare the graphics. As the name suggests, the method employs gauge-including atomic orbitals (GIAOs)⁴⁴ rather than GSGT.

Appendix B: convergence between Ampère line integrals and current–surface integrals

A rectangular basis grid was set up in the (y, z) plane consisting of 50×100 points with a spacing of $0.01a_0$ in both dimensions and positioned such that the Li nucleus is centered on the right z edge between two grid points at a distance $0.005a_0$. The $2 \times (100 + 50) - 4 = 296$ boundary points $\{\mathbf{r}_i^b\}_{i \in \{1, \dots, 296\}}$ were used for the numerical line integration,

$$k_1 \sum_{i=1}^{295} \frac{\mathbf{B}_{\text{ind}}(\mathbf{r}_{i+1}^b) - \mathbf{B}_{\text{ind}}(\mathbf{r}_i^b)}{2} \cdot (\mathbf{r}_{i+1}^b - \mathbf{r}_i^b) + \frac{\mathbf{B}_{\text{ind}}(\mathbf{r}_1^b) - \mathbf{B}_{\text{ind}}(\mathbf{r}_{296}^b)}{2} \cdot (\mathbf{r}_1^b - \mathbf{r}_{296}^b) \quad (6)$$

$$\approx \frac{1}{\mu_0} \oint_{\partial \mathcal{S}} \mathbf{B}_{\text{ind}} \cdot d\ell,$$

with $k_1 = e^2 a_0^2 c / \hbar \times 10^9 \approx 28.179409$.

To approximate the surface integral over \mathbf{J} , we have used the dual of the base grid which yields $49 \times 99 = 4851$ points $\{\mathbf{r}_i^d\}_{i \in \{1, \dots, 4851\}}$ centered in the squares of the basis grid such that

$$k_2 \sum_i^{4851} \mathbf{J}_x(\mathbf{r}_i) \cdot 0.01^2 \approx \int_{\mathcal{S}} \mathbf{J} \cdot d\mathbf{s},$$

with $k_2 = a_0 / (100\mu_0) \approx 0.042111$ (Table 1).

Table 1 A numerical test for basis-set convergence between Ampère line integrals and surface integrals of \mathbf{J} across a plane. All values are given in $nA \text{ T}^{-1}$

Basis set	$\mu_0^{-1} \mathbf{B}_{\text{ind}} \cdot d\ell$	$\mathbf{J} \cdot d\mathbf{s}$	Difference
aug-cc-pVTZ	3.83	4.00	0.17
aug-cc-pVQZ	4.21	4.34	0.13
aug-cc-pV5Z	4.41	4.47	0.05



Appendix C: convergence of open-line integrals for different line lengths

C.1 Total current flux in benzene

The total current flux in benzene has been numerically integrated by using the r.h.s. of eqn (1) with a set of points at a fixed distance of 0.05 a.u. The points are placed either symmetrically along the z axis (forming a line) or on a semicircle in the x, z plane. In each of the cases, they extend from $-height$ to $+height$. The resulting line integral values are shown in Table 2. The value of the integral on the semicircle can serve as an indicator of the incompleteness of the finite line integral. Remarkably, an extension of ± 100 a.u. is required to achieve

Table 2 Numerical convergence of total current integral in benzene depending on the integration path height (given in atomic units) on the z axis. Current flux (susceptibility) values are given in nA T^{-1} . The support points of the numerical path integrals for height of 50 a.u. is shown in Fig. 5

Height	Path integral for path		
	Semi circle	Line	Semicircle \cup line
50	0.01821	11.20840	11.22661
100	0.00457	11.23570	11.24027
200	0.00107	11.24250	11.24357
400	-0.00000	11.24420	11.24420

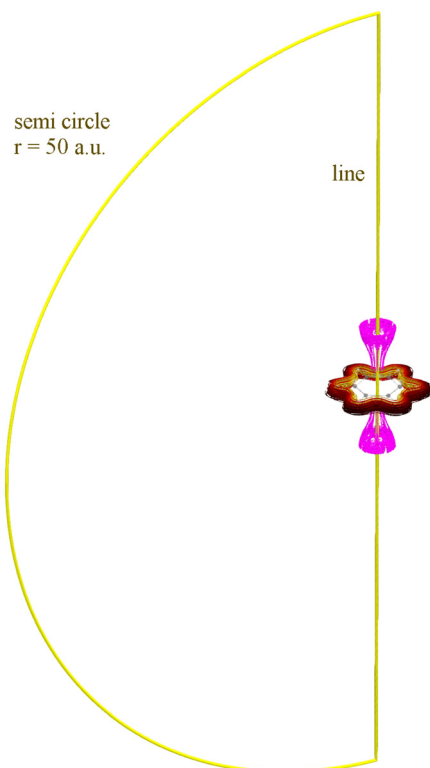


Fig. 5 A benzene molecule in a magnetic field perpendicular to the molecular plane with current density streamlines (yellow–red–brown), regions of numerically zero ($<10^{-30}$ nA T^{-1}) currents (magenta) and exemplary integration lines (yellow) as outlined in Section C.1 are shown.

numerical convergence to two digits accuracy of the current strength (which is recommended for intermediate results).

C.2 The flux in an infinitely thin wire loop obtained using the r.h.s. of eqn (1)

According to the Biot-Savart law, an infinitely thin wire loop of radius R placed onto the x, y plane, centered in the origin and sustaining current I yields a magnetic flux of

$$B(z) = \frac{\mu_0}{2} \frac{R^2 I}{(R^2 + z^2)^{3/2}} \quad (7)$$

along the z axis. Applying the r.h.s. of eqn (1) and integrating from $-z_0$ to z_0 gives a flux of

$$I(z_0) = \frac{I z_0}{R \sqrt{R^2 + z_0^2}} \quad (8)$$

For $R = 1$ a.u., we obtain $I(z_0) = I z_0 / \sqrt{1 + z_0^2}$. Thus, if we want to recover 99.9% of the total current (correct to one digit), z_0 needs to extend already to more than 22 a.u.

For molecules with a more diffuse current density distribution and larger ring-current radii, it is plausible that in order to arrive at two digits accuracy, one has to integrate out to a hundred atomic units or more.

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Notes and references

† According to Reyn,³² these two stagnation points can be classified as “three-branched saddle-node with stable two branched plane node” and they are connected by a heteroclinic streamline.

‡ The Collard–Hall index gives rank r and signature s of the Jacobian as an ordered pair (r, s) where s is the sum of the signs of the eigenvalues.³³

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