Benchmarking Density Functional Approximations for Diamagnetic and Paramagnetic Molecules in Nonuniform Magnetic Fields

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ABSTRACT: In this article, correlated studies on a test set of 36 small molecules are carried out with both wavefunction (HF, MP2, CCSD) and density functional (LDA, KT3, cTPSS, cM06-L) methods. The effect of correlation on exotic response properties such as molecular electronic anapole susceptibilities is studied and the performance of the various density functional approximations are benchmarked against CCSD and/or Møller–Plesset (MP2). Atoms and molecules are traditionally classified into “diamagnetic” and “paramagnetic” based on their isotropic response to uniform magnetic fields. However, in this article, we propose a more fine-grained classification of molecular systems on the basis of their response to generally nonuniform magnetic fields. The relation of orientation to different qualitative responses is also considered.

1. INTRODUCTION

Magnetic field effects pose unique challenges for quantum chemistry. Although the calculation of particular properties, most notably magnetic dipoles, nuclear shielding constants, and current densities induced by uniform magnetic fields, is nowadays routine, many other aspects have been subject to relatively few systematic studies, if any at all. Higher-order magnetic response and response to nonuniform magnetic fields are examples of this.1–4 Moreover, nonperturbative effects of strong magnetic fields alter the normal chemistry of small molecules, giving rise to an exotic and largely unexplored strong field chemistry.5–7 These challenges appear at all levels of theory, as for example, time-reversal and spin symmetry and other built in adaptations to zero-field settings need to be reconsidered. In density functional theory, in particular, an additional aspect is that magnetic field effects are formally beyond the scope of the standard mathematical formulation, necessitating extensions.8–11 Yet, the practically available density functional methods have almost exclusively been developed as heuristic approximations with the standard formulation, and approximations that properly incorporate magnetic fields are not yet mature enough to be practically useful.12–15 Insofar as available density functional methods produce useful estimates of magnetic field effects, there is a high risk due to error cancellations specific to the most common magnetic properties. For example, some functionals have been fitted to magnetizabilities or nuclear shielding constants.13 In the present work, we explore properties related to magnetic field gradients using several available density functional methods, including meta-GGAs that have recently emerged as particularly promising candidates for magnetic field effects, and compare to results at the second-order Møller–Plesset (MP2) and coupled-cluster singles and doubles levels (CCSD).

Quantum chemical computations of magnetic properties almost always rely on the assumption of weak-fields enabling formulations based on perturbation theory. However, for a higher-order magnetic response, in particular when London atomic orbitals (LAOs) are employed to enforce gauge-origin invariance and accelerate basis set convergence,14–17 the perturbative approach becomes increasingly more difficult. Ordinary Gaussian basis sets require very large basis sets for gauge-origin invariance.1,3,4,18,19 In this study, we have used LAOs in combination with a nonperturbative (finite field) approach. Integral evaluation for the LAOs, which are plane-wave/Gaussian hybrid functions,20,21 has been implemented in the London program20,22 and has been employed in the finite field computation of magnetic properties.23,24 earlier. Since the introduction of the magnetic field in the Hamiltonian only requires a modification of the one-electron part, no additional implementation is necessary for extension to post-Hartree–Fock methods. A nonperturbative approach also opens up the possibility of studying strong magnetic fields competing with the Coulomb forces and has led to the discovery of nonperturbative transition from closed-shell para- to diamag.
closed-shell molecules subject to transverse magnetic moments arising from induced orbital currents for a set of 36 MP2. An earlier study to assess the eect of its isotropic value decides the classi- when this tensor only has negative (or positive) eigenvalues. A clear-cut example of diamagnetism (or paramagnetism) occurs in this case, the isotropic magnetizability turns out to be a three-dimensional vector, the second-order magnetic susceptibility is a 3 \times 3 tensor. Conventionally, the sign of its isotropic value decides the classification of the molecule as dia- or paramagnetic. For example, the BH molecule is found to have a diamagnetic response to fields parallel to the chemical bond and a paramagnetic response to fields perpendicular to it. In this case, the isotropic magnetizability turns out to be paramagnetic as well leading to the classification of BH as a closed-shell paramagnetic molecule. However, a molecule such as square C_4H_4 shows a weak paramagnetic response to fields perpendicular to the molecular plane but an overall diamagnetic response.

The magnetic susceptibilities related to inhomogeneities in the magnetic field such as the anapole susceptibilities are independent of the magnetizabilities and may in some cases oppose the effects of the uniform component of the magnetic field. In our opinion, a classification should also encompass molecular response to nonuniform magnetic fields in general, as far as possible. In what follows, we propose a simple classification of magnetic response to nonuniform fields. The response of the electrons to inhomogeneities in the external magnetic field arises from both orbital effects and spin effects. Among the few studies of these effects is the work by Lazzeretti and co-workers on a perturbative formalism for the orbital response due to field gradients and some other studies at the Hückel-level, Hartree–Fock level, and correlated levels. While spin effects are certainly important and in most cases the dominant effect, this article is only concerned with orbital effects in closed-shell molecules. Further exploratory studies are planned but beyond the present scope.

The response to (transverse) magnetic field gradients may be quantified by the anapole moments, which couple linearly to the curl of the magnetic field. They may be considered to arise from the meridional currents in a toroidal charge distribution. They are antisymmetric under both spatial inversion and time-reversal. Nuclear anapole moments are studied by physicists in connection with parity violation with the first experimental evidence coming from measurements on the Cs atom. Experiments for measuring permanent and induced electronic anapole moments have been suggested. However, only special structures such as molecular nanorotors and ferroelectric nanostructures, and Dy clusters (single-molecule magnets) are expected to have permanent anapole moments. Anapole moments in metamaterials have also been observed with potential application in sensors.

On the other hand, induced anapole moments easily arise in molecules placed in external nonuniform fields and we can compute the corresponding susceptibilities. Both toroidal spin and/or orbital currents can give rise to anapole moments. Induced anapolar current densities in conjugated cyclic acetylenes and some small molecules have been studied. Spin and orbital contributions to anapole moments have been analyzed in a simple analytical model of diatomics and also using nonperturbative General Hartree–Fock theory. The orbital contributions have been estimated by both perturbative approaches and nonperturbative approaches. Faglioni et al. have derived the perturbative expressions for induced anapole moments.

The outline of the article is as follows. In Section 2, we define the Hamiltonian and the properties relevant to our study. Section 3.2 discusses our proposed classification of molecules. Section 3 presents our results on the effect of correlation on the anapole susceptibilities and the relative performance of the various density functional approximations. Finally, we conclude with the summary in Section 4.

2. HAMILTONIAN AND PROPERTIES

In this study, the nonuniform magnetic field has the form

$$\mathbf{B}_{\text{tot}}(\mathbf{r}) = \mathbf{B} + \mathbf{r}_b \mathbf{b} - \frac{1}{3} r_0 \text{Tr}(\mathbf{b})$$

(1)
where $\mathbf{B}$ is a uniform (position independent) component, $\mathbf{b}$ is a $3 \times 3$ matrix defining the field gradients, and $\mathbf{r}_h = r - \mathbf{h}$ is the position relative to some reference point $\mathbf{h}$. This form may be viewed as arising from a truncation of a Taylor expansion of a general magnetic field around $r = \mathbf{h}$ of linear order. The corresponding vector potential can be written as

$$A_{\text{tot}}(r) = \frac{1}{2} \mathbf{B} \times \mathbf{r}_h - \frac{1}{3} \mathbf{r}_h \times (\mathbf{r}_h \times \mathbf{b})$$

(2)

where $\mathbf{r}_h = r - \mathbf{g}_h$, $\mathbf{g}_h$ being the gauge origin. One can show that $\mathbf{B}_{\text{tot}} = \nabla \times A_{\text{tot}}$ and that the magnetic field is divergence free, $\nabla \cdot \mathbf{B}_{\text{tot}} = 0$. The symmetric part, $\mathbf{b} = \mathbf{b}^T$, can be set to zero and we focus on the antisymmetric part $C_{ij} = \epsilon_{ijk} b_{kj}$ of the matrix $\mathbf{b}$. We can then write

$$A_{\text{tot}}(r) = \frac{1}{2} \mathbf{B} \times \mathbf{r}_h - \frac{1}{3} \mathbf{r}_h \times (\mathbf{C} \times \mathbf{r}_h)$$

(3)

$$\mathbf{B}_{\text{tot}}(r) = \mathbf{B} + \frac{1}{2} \mathbf{C} \times \mathbf{r}_h$$

(4)

Furthermore, the antisymmetric part of $\mathbf{b}$ equals the curl of the magnetic field, $\nabla \times \mathbf{B}_{\text{tot}} = \mathbf{C}$, which is taken to be position independent.

The nonrelativistic Schrödinger–Pauli Hamiltonian is given by

$$\hat{H} = \frac{1}{2} \sum_i \frac{\mathbf{p}_i^2}{m} - \frac{1}{2} \sum_i \phi(r_i) + \sum_{i<j} \frac{1}{r_{ij}} + \sum_i B_{\text{tot}}(r_i) \cdot \hat{S}_i$$

(5)

where $\hat{r}_i = -i \nabla_i + \mathbf{A}_\alpha(r_i)$ is the mechanical momentum operator. Properties can be alternatively viewed as expectation values $\langle \Psi | \mathbf{O} | \Psi \rangle$ or as derivatives of the energy $E = \langle \Psi | \hat{H} | \Psi \rangle$ related to terms in a Taylor expansion. In this study, we ignore the spin-Zeeman term and thereby the spin-breaking induced by the nonuniform part of the magnetic field. The first-order orbital angular momentum

$$\mathbf{L}_q = \sum_i \langle \Psi | \hat{L}_q | \Psi \rangle, \quad \hat{L}_q = \mathbf{r}_q \times \hat{\mathbf{r}}_q$$

(6)

is with respect to an arbitrary reference point, $q$. Given the form of the magnetic vector potential above, it is $L_q$ with the reference point at the gauge origin that is the relevant magnetic dipole moment. The orbital anapole moment is similarly given by

$$\mathbf{a} = -\sum_i \langle \Psi | r_i \times \mathbf{L}_q | \Psi \rangle$$

(7)

Recalling that the current density can be obtained as the functional derivative $j = \delta E / \delta A_{\text{tot}}$, we can also identify the magnetic orbital dipole moment and anapole moment with linear and quadratic moments of the current density

$$\mathbf{J}_q = \int r \times j(r) \, d\mathbf{r}$$

(8)

$$\mathbf{a} = -\frac{1}{3} \int \mathbf{r}_h \times \left( \mathbf{r}_h \times j(r) \right) \, d\mathbf{r}$$

(9)

We note that the energy $E$ as well as expectation value properties like $\mathbf{J}_q$ and $\mathbf{a}$ can be obtained directly as functions of $\mathbf{B}$ and $\mathbf{C}$. We can thus define second-order properties from a Taylor expansion of the energy

$$E(\mathbf{B}, \mathbf{C}) \approx E_0 + \frac{1}{2} \mathbf{B} \cdot \mathbf{J}_q - \frac{1}{2} \mathbf{C} \cdot \mathbf{a} - \frac{1}{2} \mathbf{B}^T \chi \mathbf{B} - \mathbf{B} \cdot \mathbf{M}$$

(10)

where $\mathbf{J}_q$ and $\mathbf{a}$ are evaluated at $\mathbf{B}_{\text{tot}} = 0$. We can identify $\chi$ as the magnetizability tensor, and call $\mathbf{M}$ as the mixed anapole susceptibility tensor, and $\mathbf{A}$ as the anapole susceptibility tensor.

When the Hellmann–Feynman theorem is applicable, the expectation value quantities can be equated with energy derivatives

$$\mathbf{J}_q = 2 \frac{\partial E(\mathbf{B}, \mathbf{C})}{\partial \mathbf{B}}$$

(11)

$$\mathbf{a} = -2 \frac{\partial E(\mathbf{B}, \mathbf{C})}{\partial \mathbf{C}}$$

(12)

However, when LAOs are used, the basis set depends on the parameters $\mathbf{B}$ and $\mathbf{C}$ leading to a discrepancy between the expectation values and the energy derivatives, in general, except in the complete basis set limit.

Second-order susceptibilities may be defined as follows

$$\chi_{\alpha\beta} = -\frac{\partial^2 E(\mathbf{B}, \mathbf{C})}{\partial B_\alpha \partial B_\beta} \bigg|_{\mathbf{B}=0,\mathbf{C}=0}$$

(13)

$$\mathbf{A}_{\alpha\beta} = -\frac{\partial^2 E(\mathbf{B}, \mathbf{C})}{\partial C_\alpha \partial C_\beta} \bigg|_{\mathbf{B}=0,\mathbf{C}=0}$$

(14)

$$\mathbf{M}_{\alpha\beta} = -\frac{\partial^2 E(\mathbf{B}, \mathbf{C})}{\partial B_\alpha \partial C_\beta} \bigg|_{\mathbf{B}=0,\mathbf{C}=0}$$

(15)

One can also introduce the closely related, but inequivalent, quantities

$$\mathbf{A}_{\alpha\beta}' = \frac{\partial \mathbf{a}_\alpha(\mathbf{B}, \mathbf{C})}{\partial C_\beta} \bigg|_{\mathbf{B}=0,\mathbf{C}=0}$$

(16)

$$\mathbf{M}_{\alpha\beta}' = -\frac{1}{2} \frac{\partial \mathbf{J}_q(\mathbf{B}, \mathbf{C})}{\partial C_\beta} \bigg|_{\mathbf{B}=0,\mathbf{C}=0}$$

(17)

$$\mathbf{M}_{\alpha\beta}'' = \frac{1}{2} \frac{\partial \mathbf{a}(\mathbf{B}, \mathbf{C})}{\partial B_\beta} \bigg|_{\mathbf{B}=0,\mathbf{C}=0}$$

(18)

Again, in the basis set limit, equivalence is restored, i.e., $\mathbf{A} = \mathbf{A}'$ and $\mathbf{M} = \mathbf{M}'$. Note that the multiplicative factors in eqs 10, 15, 17, and 18 have been corrected from those reported in earlier publications$^{2,5,36}$ to be self-consistent with the other definitions. This implies that the $\mathbf{M}$ values reported in these publications should be halved. However, this has no implication on the conclusions of the two papers.

3. RESULTS AND DISCUSSION

Our test set contains 36 molecules (HF, CO, N₂, H₂O, HCN, HOF, LiH, NH₃, H₂CO, CH₄, C₂H₄, AlF, CH₃F, C₃H₄, FCCH, FCN, H₂S, HCP, HFCO, H₂C₂O, LiF, N₂O, OCS, H₂C₂O, PN, SO₂, O₂F₂, H₂O₂, BH, CH⁺, AlH, BeH⁺, SiH⁺, C₄H₄, FNO) and subsumes the test set of diamagnetic molecules in Tellgren et al.$^{12,23,36}$ Geometries for the molecules are as reported in earlier publications$^{2,5,36}$ and are also provided in the Supporting Information.
All calculations were performed using the London program.\textsuperscript{20,22} The density functional calculations used the previously reported implementations.\textsuperscript{12,35} The coupled-cluster calculations were performed using the previously reported implementation\textsuperscript{19} and exploratory calculations (results not included) also used a new functionality.\textsuperscript{70} The symmetric finite difference formula for numerical second derivatives of the energy was employed to compute the anapole susceptibilities $\mathcal{A}$ and $\mathcal{M}$. Step sizes of $\epsilon = 0.01$ au for B and $\epsilon^* = 0.005$ au for C were used. $\epsilon^*$ was chosen to be smaller as the effect of C on the local magnetic field is scaled by the interatomic distances in the molecule. The reference point, $\mathbf{b}$, for C was placed at the center of charge of the molecules in all cases. The error in the energy is quadratic in the step size within the limits to which the energy is converged while the error in the analytically computed moments (first derivative of energy) is linear. All numerical results presented in this article are given in SI-based atomic units—see the earlier work for the conversion factors to SI units.\textsuperscript{23}

The uncontracted aug-cc-pVTZ basis set has been employed for all of the computations. The name of the basis set is prefixed with “L” to denote the use of London atomic orbitals and “u” to indicate that the basis sets are uncontracted—Luu-cc-pCVTZ.

3.1. Current-Dependence and Meta-GGAs. The meta-GGA functional form allows a dependence on the kinetic energy density. In the absence of a magnetic field, the everywhere positive, canonical kinetic energy $\tau_{\text{can}} = 1/2 \sum \lvert \mathbf{V} \phi_i \rvert^2$, with summation over occupied orbitals $\phi_i$, is the natural choice. In the presence of a magnetic field, $\tau_{\text{can}}$ is gauge dependent and cannot be used. An obvious solution is to use the gauge independent, physical kinetic energy density $\tau_{\text{phys}} = 1/2 \sum (-\mathbf{V} + \mathbf{A}_0) \phi_i \phi_i^T$ instead. This choice has been suggested by Maximo and Scuseria.\textsuperscript{71} An alternative, with some theoretical aspects and numerical results in its favor,\textsuperscript{36,72–74} is to instead use Dobson’s gauge-invariant kinetic energy density

$$\tau_D = \tau_{\text{phys}} - \frac{1}{2} \frac{\mathbf{j} \mathbf{j}^T}{2 \rho} = \tau_{\text{can}} - \frac{1}{2} \frac{\mathbf{j} \mathbf{j}^T}{2 \rho} \quad (19)$$

where $\mathbf{j} = \text{Im} \sum \phi_i \nabla \phi_i$ is the paramagnetic current density and $\mathbf{j} = \text{Re} \sum \phi_i \nabla \phi_i (-\mathbf{V} + \mathbf{A}_0) \phi_i$ is the physical current density.

A previous work used a prefix “a” to denote meta-GGA functionals using the physical $\mathbf{r}_{\text{phys}}$ (e.g., aTPSS) and a prefix “c” to denote functionals using Dobson’s $\tau_D$ (e.g., cTPSS). In the present work, we only consider the latter type, specifically cTPSS and cM06-L.

3.2. Classification. Different response tensors in general have different physical dimensions and units, although this fact is somewhat obscured when working with atomic units. To account for this, we fix a length $l = l_0$ and define an auxiliary quantity $C' = lC$, and auxiliary response tensors $\mathcal{A} = l^2\mathcal{A}$ and $\mathcal{M} = l\mathcal{M}$. In atomic units, the numerical values of the quantities $\mathcal{A}$ and $\mathcal{M}$ remain unchanged by this transformation of response tensors to shared units. Next, we construct a 6 x 6 matrix of the form

$$\xi = \left[ \begin{array}{cc} \chi & \mathcal{M} \\ \mathcal{M}^T & \frac{1}{2} \mathcal{A} \end{array} \right]$$

which allows us to re-express the second-order energy in eq 10 as

$$E(\mathbf{B}, \mathbf{C}) \approx E_0 + \frac{1}{2} \mathbf{B}^T \mathbf{C} \left[ \mathbf{J}_0 \right] \left[ -a/l \right] - \frac{1}{2} \mathbf{C}^T \mathbf{J}_0 \mathbf{B} \xi \mathbf{C}'$$

$\xi$ is symmetric and has real eigenvalues, which we denote by $\alpha = (\alpha_0, \ldots, \alpha_6)$. If one or more eigenvalues are positive, i.e., energy decreases with any component of $\mathbf{B}$ or $\mathbf{C}$, we classify the system broadly as paramagnetic. Otherwise, if all eigenvalues are negative, we classify the system as diamagnetic. Diagonalization of the submatrices gives us further details of this behavior such as separate response to only B or C. Additionally, the trace of $\xi$ gives us a single number for the overall response to a generally nonuniform field. We also obtain an average eigenvalue as

$$\bar{\alpha} = \frac{1}{6} \sum_{i=1}^{6} \alpha_i = \frac{1}{6} \text{Tr}(\xi) \quad (22)$$

which is also the orientational average over all possible molecular orientations (with $\mathbf{B}$, $\mathbf{C}$ fixed). This procedure may be extended to increasingly nonuniform magnetic fields, such as those with curvatures and beyond. In this article, the magnetic field has a constant gradient and is of the form shown in eq 1.

The different eigenvectors of $\xi$ are visualized for four molecules in Figure 1. CCSD level tensors were used for HOF and FNO and MP2 level tensors for C4H4 and CH2O. Each six-dimensional eigenvector ($\mathbf{B}$, $\mathbf{C}$) is represented as a pair of three-dimensional vectors, displaced along the vertices of a hexagon to make clear which arrows form pairs. The length of each arrow is constant varies between molecules, and the sign is indicated by red (negative) and blue (positive) colors. In the plot, the arrows are sorted so that the magnitude increases clockwise around the hexagon. Note also that the eigenvectors and the molecular geometry are expressed in the same coordinate system, such that the angle between eigenvectors and bond axes is independent of the choice coordinate system and orientation of the molecule. If the mixed anapole susceptibility tensor vanishes, the eigenvalue problem does not couple the $\mathbf{B}$ and $\mathbf{C}$. This happens for cyclobutadiene, for which each eigenvector is of the form ($\mathbf{B}$, 0) or (0, $\mathbf{C}$). On the other hand, for FNO and CH2O, there is...
substantial coupling and some eigenvectors have B and C' components of similar magnitude. Of the molecules shown, C₄H₄, HOF, and CH₂O each have a single paramagnetic eigenvector with a magnitude that is small compared to the
diamagnetic eigenvalues—in the plot the corresponding arrows are barely visible. By contrast, FNO has two paramagnetic eigenvalues of large magnitude.

In Figure 2, we have plotted the eigenvalues, $\alpha_i^\tau$ of the super-tensor, $\xi^\tau$ in decreasing order. We have presented the values obtained with the most accurate methods we have studied in this article, viz. CCSD and MP2. However, the classification is robust and all methods studied by us including Hartree–Fock (HF) and DFT show the same qualitative classification (except for FNO). The geometry, energy, and properties of FNO are all extremely sensitive to correlation and need at least CCSD(T) level computations for reasonable accuracy. 75 The values of the susceptibilities cannot be determined with any reasonable accuracy with our set of methods. Moreover, the computations do not converge with LDA and cM06-L functionals. We have thus left this molecule out of the error statistics presented in Section 3. All conventionally paramagnetic molecules show at least one positive eigenvalue. In addition, FNO and HOF show positive eigenvalues arising from a paramagnetic response to some component of C. Thus, according to our criterion, the following molecules from our test set are paramagnetic: AlH, BH, BeH$, C_2H_4$, CH$^+$, CH$_2$O, SiH$, HOF$, and FNO. The nature of this net paramagnetic behavior is summarized in Table 1.

| molecule | B | C | (B, C) |
|----------|---|---|--------|
| AlH      | + | + | +      |
| BH       | + | + | +      |
| BeH$^+$  | + | + | +      |
| C$_2$H$_4$ | + | + | +      |
| CH$^+$   | + | + | +      |
| CH$_2$O  | − | + | +      |
| FNO      | − | − | −      |
| HOF      | − | − | −      |
| SiH$^+$  | + | + | +      |

Table 1. Selected Molecules Which Are Proposed to Be Classified by Us as Paramagnetic$^a$

$^a$For response tensors related to B, C, and jointly to (B, C), we show the sign of the average and maximum eigenvalue. The molecule FNO is extremely challenging for correlated theories, leading to different conclusions from different methods with respect to the net response to (B, C): diamagnetic (−) from CCSD and MP2; paramagnetic (+) from HF, KT3, and cTPSS; and not converged for LDA and cM06-L.

We note that, due to the Cauchy interlace theorem, adding dimensions will increase the maximum eigenvalue and decrease the minimum eigenvalue. Hence, with $\alpha_{\text{max}} = \max_{1 \leq i \leq 6} \alpha_{\tau i}$, $\alpha_{\text{max}} = \max_{1 \leq i \leq 6} \alpha_{\tau i}$ and $\alpha_{\text{min}} = \max_{1 \leq i \leq 6} \alpha_{\tau i}$, we have

$$\alpha_{\text{max}} \geq \max \left\{ \alpha_{\text{max}}^i l^2 \alpha_{\text{max}}^i \right\}$$

With a similar notation for the minimum eigenvalues, we get an inequality in the reverse direction. When $M = 0$, usually by reason of symmetry, equality is achieved such as for C$_2$H$_4$, C$_2$H$_2$, CH$_2$H$_2$, and N$_2$. Equality is also achieved for the largest eigenvalue of $M$ in CH$_2$O, CO, and HCP. The smallest eigenvalue also almost saturates in these cases. This may indicate that there is a limit to how large the orbital paramagnetic and/or diamagnetic response can be within the limits of the basis set and molecular symmetry. While the non-zero components of $M$ are small ($\sim 10^{-2} \text{ au}$) for CO and HCP, this is not the case for CH$_2$O ($M_{\text{MP2}} = 0.956$, $M_{\text{MP2}} = 0.760 \text{ au}$).

Most molecules studied are highly diamagnetic with respect to C but we must remember that here we are only studying the orbital response. The spin symmetry breaking caused by C will activate the spin-Zeeman term in the Hamiltonian leading to an overwhelmingly paramagnetic response to C. 75 The interplay of spin and orbital effects of nonuniform magnetic fields is discussed in an earlier publication. 25 BeH$^+$ is particularly strongly diamagnetic to C. The DFT computation with the cM06-L functional does not converge for BeH$^+$. FNO, on the other hand, is strongly paramagnetic with respect to C according to computations with all of the methods in our study.

3.3. Performance of Density Functional Approximations for Magnetic Susceptibilities. In this section, we discuss the relative performance of various wavefunction and density functional methods in computing anapole magnetizabilities, $A$ and $M$. We compare this performance with their accuracy in describing the magnetizability, $\chi$, the most well-studied among the magnetic response quantities.

To quantify errors in the response tensors relative to a reference method we rely on the Frobenius norm

$$\epsilon^\tau_{\text{method}} = \| \psi_{\text{method}} - \psi_{\text{ref}} \|_F = \left( \sum_{ij} \alpha_{ij}^\tau \right)^{1/2}$$

and similarly for $\chi^\tau$, $A^\tau$, and $M$. For the isotropic average, we use a similar notation to mean $\epsilon_{\text{method}}^\tau = \| \psi_{\text{method}} - \psi_{\text{ref}} \|_{F_i}$.

We present the error bars for the various methods as box and whisker plots where the median error is indicated by the horizontal line in the middle of the box and the top and bottom ends of the box indicate the third quartile and first quartile, respectively. The length of the box is thus the interquartile range. The top and bottom ends of the whiskers indicate the maximum and minimum errors considered in the estimation of the quartiles. Points beyond the whiskers are not considered in the statistics and are regarded as outliers. We superimpose swarmplots on the box and whisker plots to display the underlying data.

Values of isotropic magnetizability computed with Hartree–Fock theory are often reasonable in the absence of low-lying excited states with correlation contributions of the order of 1–3%. 76 However, DFT approximations, which are reasonably good for correlation energy and electric properties, such as BLYP or B3LYP, are mostly inaccurate for magnetic properties, often being worse than Hartree–Fock theory. This has prompted the development of exchange–correlation functionals tailored to magnetic properties such as the KT3 functional of Keal and Tozer. 13cTPSS has also been seen to be remarkably accurate for the same. 77 Computational studies have indicated that the effect of electron correlation on the isotropic magnetizability $(\chi/3)$ is often an order of magnitude lower than that on the anisotropic magnetizability. 77 The isotropic magnetizability is also less sensitive to the basis set size. These conclusions are borne out by our results, as shown in Figure 3. We have not studied BLYP and B3LYP as they are known to perform poorly for conventional magnetic properties.

While the accuracy of various DFT functionals for magnetizability has been well-studied, it remains to be seen if the same conclusions can be reached for more exotic properties such as anapole susceptibilities. In particular, we wish to explore if the
Figure 3. Errors in the isotropic magnetizability computed by various methods in the aug-cc-pCVTZ basis set relative to CCSD (top panel) and MP2 (bottom panel).
Figure 4. Errors in magnetizability tensor, $\chi$, of the conventionally diamagnetic molecules (negative $\tilde{\alpha}$) in our test set computed by various methods in the Luaug-cc-pCVTZ basis set relative to CCSD (top panel) and MP2 (bottom panel).
Figure 5. Errors in the magnetizability tensor, $\chi$, of the conventionally paramagnetic molecules (positive $\bar{\alpha}$) in our test set computed by various methods in the Luaug-cc-pCVTZ basis set relative to CCSD (top panel) and MP2 (bottom panel).
Figure 6. Errors in anapole susceptibility, $\mathcal{A}$, computed by various methods in the Luaug-cc-pCVTZ basis set relative to CCSD (top panel) and MP2 (bottom panel).
Figure 7. Errors in mixed anapole susceptibility, $\mathcal{M}$, computed by various methods in the Luaug-cc-pCVTZ basis set relative to CCSD (top panel) and MP2 (bottom panel).
Figure 8. Errors in the average eigenvalue of the super-tensor, $\zeta$, computed by various methods in the Luau-cc-pCVTZ basis set relative to CCSD (top panel) and MP2 (bottom panel) for diamagnetic molecules as classified by us.
Figure 9. Errors in the average eigenvalue of the super-tensor, $\zeta$, computed by various methods in the Luaug-cc-pCVTZ basis set relative to CCSD (top panel) and MP2 (bottom panel) for paramagnetic molecules as classified by us.
KT3 and cTPSS functionals continue to perform well for $\mathcal{A}$ and $\mathcal{M}$.

In the top panel of Figure 4, we can see that except the cM06-L functional all of the methods considered here perform reasonably for the conventionally diamagnetic molecules. The cTPSS functional and MP2 show similar accuracy with HF and KT3 also doing quite well. The bottom panel in Figure 4 samples a larger test set using MP2 as the benchmark. Among the density functionals LDA, KT3, and cTPSS perform similarly. It is interesting to note that the outlier for many of the methods in the top panel is HOF, which would be classified as paramagnetic by our proposed scheme. The trends for the paramagnetic molecules plotted separately in Figure 5 are much more surprising. All of the errors are much higher than for the diamagnetic molecules indicating how much more difficult it is to describe the paramagnetic response. MP2 no longer performs as well and is easily surpassed in accuracy by both KT3 and cTPSS. A similar conclusion was reached by Reimann et al. where even CCSD performed worse than cTPSS relative to CCSD(T) for the paramagnetic molecules. The bottom panel in Figure 5 is thus much less reliable as a measure of errors in density functionals. Between KT3 and cTPSS, cTPSS is somewhat more accurate for the paramagnetic systems studied by us.

The top panel of Figure 6 shows a reasonable description of $\mathcal{A}$ by most methods (except cM06-L) against CCSD. MP2 shows the highest accuracy with cTPSS following close behind. The bottom panel of Figure 6 also follows the same trends with MP2 as the reference. Here too, the paramagnetic molecules show a larger error than the diamagnetic ones with HOF behaving as the conventionally paramagnetic molecules.

Due to the smaller numerical values in the mixed anapole susceptibility tensor, $\mathcal{M}$, the errors appear to be smaller in Figure 7. cM06-L is no longer as bad although it is still the worst among the methods studied.

Finally, we present the errors in the quantity, $\tilde{\alpha}_z$—the average eigenvalue of our super-tensor, $\zeta$ in Figures 8 and 9. The molecules have been classified according to the criterion in Section 3.2. This may be considered as a condensed representation of all of the errors presented in Figures 3–7 allowing for the possibility of some error cancellation. The top panel indicates a comparable performance of MP2, KT3, and cTPSS in comparison with CCSD. The larger test set in bottom panel also fits with a similar performance. Molecules classified as paramagnetic by us again show the largest errors.

Our findings are summarized in Table 2.

### Table 2. Methods Studied by Us Relative to CCSD in the Increasing Order of Median Errors

| property | class | error |
|----------|-------|-------|
| $\text{Tr}(\zeta)$ | dia | cTPSS < MP2 < HF ≈ KT3 < LDA ≪ cM06-L |
| $\text{Tr}(\zeta)$ | para | cTPSS ≈ KT3 < MP2 < HF ≈ KT3 < LDA ≪ cM06-L |
| $\chi$ | dia | cTPSS ≈ KT3 < MP2 < HF < KT3 < LDA ≪ cM06-L |
| $\chi$ | para | cTPSS ≈ KT3 < MP2 < HF ≈ KT3 < LDA < cM06-L |
| $\mathcal{A}$ | dia | MP2 ≈ cTPSS ≈ KT3 < MP2 < HF < KT3 < LDA < cM06-L |
| $\mathcal{A}$ | para | MP2 ≈ cTPSS ≈ KT3 < MP2 < HF ≈ KT3 < LDA < cM06-L |
| $\mathcal{M}$ | dia | MP2 < LDA < cTPSS < KT3 < MP2 < HF < KT3 < LDA < cM06-L |
| $\mathcal{M}$ | para | MP2 < LDA < cTPSS < KT3 < MP2 < HF ≈ KT3 < LDA < cM06-L |
| $\tilde{\alpha}_z$ | dia | cTPSS ≈ KT3 < MP2 < HF < KT3 < LDA < cM06-L |
| $\tilde{\alpha}_z$ | para | cTPSS ≈ KT3 < MP2 < KT3 < HF < LDA < cM06-L |

### 3.4. Performance of Density Functional Approximations for Nonperturbative Effects

As we have seen in the previous subsection, molecules which show paramagnetic behavior, with respect to a component of $\mathbf{B}$ or $\mathbf{C}$, are particularly challenging for all of the theories. In this subsection, we further explore two molecules—BH as an example of paramagnetic behavior with respect to a component of $\mathbf{B}$ and HOF as the newly discovered example of paramagnetic behavior with respect to a component of $\mathbf{C}$.

In the top panel of Figure 10, the paramagnetic behavior of BH with respect to a field ($B_z$) perpendicular to the bond axis ($z$) is evident up to a critical field strength of about $B_z = 0.22$ au after which the quadratic term takes over. An analogous behavior is observed in HOF (bottom panel of Figure 10) with increase in the component of $\mathbf{C}$ ($C_z$) perpendicular to the molecular plane ($xy$). The turning point can be read off as $C_z = 0.048$ au. The depth of the minimum is, however, only $10^{-5}$ $E_H$ for HOF against a depth of $10^{-2}$ $E_H$ for BH.

We have tried to assess the capacity of various theories to describe the changes in the electronic structure arising from the application of increasing $\mathbf{B}$ and $\mathbf{C}$. The CCSD method has been chosen as the reference and energies are computed with MP2 and a few selected density functional approximations. Since the
zero-field energies themselves differ considerably, we have subtracted the zero-field energy computed with each method from all other data points thereby shifting all plots to a common starting point of zero. The energy difference between these shifted data points of various methods and CCSD is then plotted in Figure 11. For BH (top panel), cTPSS and KT3 work best with nearly parallel error curves. MP2 is surprisingly worse and the cM06-L functional gives a highly non-parallel error plot. For HOF (bottom panel), the error values themselves are an order of magnitude smaller than for BH with MP2 showing the best performance. KT3 and cTPSS follow the same trend of errors increasing with increasing $C_z$ as MP2. No flattening is observed even when HOF starts showing diamagnetic behavior after the turning point of $C_z = 0.048$ au unlike the plots for BH. The cM06-L functional yields a very non-parallel error curve in this case too. Although not shown here, the corresponding plots for BH vs $C_i$ $i = x, y, z$, also show increasing errors with increasing $C_i$.

4. SUMMARY AND CONCLUSIONS

In this article, we have suggested a new classification of magnetic behavior of molecules based on their response to a generally nonuniform field. We have demonstrated that paramagnetic behavior can arise in a molecule due to inhomogeneities in the field even when its response to a uniform field is diamagnetic as is the case for FNO and HOF. We have concluded that the susceptibilities of molecules—$\chi$, $\mathcal{A}$ and $\mathcal{M}$—thus classified as paramagnetic are more difficult to describe. Assuming that CCSD gives accurate results, KT3 and cTPSS are found to show the best performance among the DFT approximations with cTPSS being marginally better than KT3. The interquartile range for both functionals are narrow across all of the properties studied by us. cTPSS and KT3 also perform quite well for the more challenging paramagnetic molecules, even better than MP2 relative to CCSD. cM06-L is particularly bad for magnetic properties performing even worse than LDA. These conclusions are also found to hold in the strong-field regime as verified for some typical challenging molecules. Hartree–Fock is surprisingly reliable for diamagnetic molecules with a more or less constant error across all magnetic properties. The paramagnetic molecules are far more sensitive to correlation. The eigenvalues of $\zeta$, or even its higher dimensional analogues with additional parameters beyond $\mathbf{B}$ and $\mathbf{C}$, can serve as a concise measure for comparing the accuracy of various theories in describing general magnetic properties.
ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jctc.0c01222.

Molecular geometries and data corresponding to the plots in this article have been tabulated (PDF).

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Notes
The authors declare no competing financial interest.

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