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SPi	Journal Code	Article ID	Dispatch: 25-FEB-20	CE: Akila
	JCC	26185	No. of Pages: 6	ME:

Received: 13 December 2019

Revised: 14 February 2020 Accepted: 17 February 2020

DOI: 10.1002/jcc.26185

FULL PAPER

Resolving a puzzling anomaly in the spin-coupled generalized valence bond description of benzene

Lu T. Xu¹ | David L. Cooper² | Thom H. Dunning Jr¹

Washington, Seattle, Washington, USA 15 ²Department of Chemistry, University of 16 Liverpool, Liverpool, UK Correspondence David L. Cooper, University of Liverpool, Liverpool L69 7ZD, UK. Email: dlc@liverpool.ac.uk 20 Thom H. Dunning Jr, Department of 22 Chemistry, University of Washington, Seattle, WA 98195 Email: thdjr@uw.edu Funding information Chemical Sciences, Geosciences and

Biosciences Division, Office of Science,

U.S. Department of Energy

¹Department of Chemistry, University of

Abstract

In an earlier study of benzene, Small and Head-Gordon found that the spin-coupled generalized valence bond (SCGVB) wave function for the π system predicted a distorted (non-D_{6h}) geometry, one with alternating CC bond lengths. However, the variations in the energy were very small and the predictions were made using a very small basis set (STO-3G). We re-examined this prediction using a much larger basis set (aug-cc-pVTZ) to determine the dependence of the energy of benzene on the distortion angle, $\Delta\theta_{CXC}$ ($\Delta\theta_{CXC}$ = 0° corresponds to the D_{6h} structure). We also found a distorted geometry with the optimum $\Delta \theta_{\text{CXC}}$ being 0.31° with an energy 0.040 kcal mol⁻¹ lower than that for the D_{6h} structure. In the optimum geometry, adjacent CC bond lengths are 1.3861 Å and 1.4004 Å. Analysis of the SCGVB wave function led us to conclude that the cause of the unusual non-D_{6h} geometry predicted by the SCGVB calculations seems to be a result of the interaction between the Kekulé and Dewar components of the full SCGVB wave function. The addition of doubly ionic configurations to the SCGVB wave function leads to the prediction of a D_{6h} geometry for benzene and a dependence on $\Delta \theta_{CXC}$ essentially the same as that predicted by the complete active space self-consistent field wave function.

KEYWORDS

benzene, Dewar structures, Kekulé structures, non-D_{6h} structure, SCGVB

9 1 | INTRODUCTION

41 The concept of aromaticity is one of the key concepts in organic 42 chemistry and has profound implications for the structures and reac-43 tivities of organic molecules. The prototype aromatic molecule, ben-44 zene, is planar with all of the CC bonds of equal length-a D_{6h} 45 structure. Traditionally, the D_{6b} structure of benzene is attributed to 46 aromaticity, that is, resonance between the two Kekulé structures of 47 benzene. More recently, there has been a spirited debate in the litera-48 ture about the role played by the σ and π bonds in determining the 49 symmetric structure of benzene. Shaik and coworkers have argued 50 that the electrons in the π orbitals of benzene favor a geometry with 51 unequal bond lengths and that the hexagonal structure of benzene is a result of the σ bonds (see the summary of these discussions in Shaik 52 et al. ^[1]). Given the importance of benzene in organic chemistry, it is 53

not surprising that others addressed this question, reporting evidence92to the contrary; see, for example, Glendening et al.[2] However, defini-tive statements as to the role of the σ and π bonds in determining the94structure of benzene requires their contributions to the total energy95to be cleanly separated and, even for the simplest level of theory, that96is, Restricted Hartree-Fock (RHF) theory, there are terms in the97energy that couple the σ and π orbitals ($J_{\sigma\pi}, K_{\sigma\pi}$).98

Against this backdrop, Small and Head-Gordon^[3] reported that a 99 spin-coupled generalized valence bond (SCGVB) wave function that 100 kept the σ orbitals doubly occupied but included all possible spin func-101 tions for the six electrons in the six SCGVB π orbitals predicted a 102 structure for benzene that has alternating CC bond lengths. However, 103 the basis set used in that study was very small (STO-3G) as was the 104 predicted energy lowering (less than 0.05 kcal mol⁻¹). Nonetheless, 105 this is a surprising result given that the SCGVB wave function for the 106

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electrons in the π orbitals of benzene includes the two Kekulé spin 1 2 coupling modes plus the three Dewar (para-bonded) spin coupling 3 modes, which, taken together, treat the CC bonds symmetrically. 4 Thus, it is surprising that the SCGVB wave function predicts a non-5 D_{6h} structure for benzene. We note that the topic of symmetry break-6 ing in the benzene molecule has also been investigated for a variety of 7 SCGVB-related methodologies by Van Voorhis and Head-Gordon,^[4,5] Lawler, Beran, and Head-Gordon,^[6] and Parkhill and Head-Gordon.^[7] 8

9 The current article reports high level calculations on the benzene molecule using the RHF, SCGVB, complete active space self-10 consistent field (CASSCF), and CCSD(T) methods. For the latter 11 12 method, calculations are reported for correlating the electrons in the π system as well as the full valence ($\sigma + \pi$) system. A much larger basis 13 14 set was used in these studies than in Small and Head-Gordon^[3]: the aug-cc-pVTZ basis sets for the carbon and hydrogen atoms.^[8,9] The 15 aug-cc-pVTZ basis sets are expected to yield results close to the com-16 17 plete basis set limit for the RHF, SCGVB, and CASSCF methods.

2 THEORETICAL METHODS

The SCGVB wavefunction considered here for the six electrons in the π system of the benzene molecule is:

> $\Psi_{\text{SCGVB}} = \hat{a}\varphi_{C\pi_1}\varphi_{C\pi_2}\varphi_{C\pi_3}\varphi_{C\pi_4}\varphi_{C\pi_5}\varphi_{C\pi_6}\Theta_{0,0}^6$ (1)

27 The σ orbitals are taken to be doubly occupied in the wave func-28 tion and are not listed in Equation (1). The set of orbitals in Equation (1), $\{\varphi_{C_{\pi_i}}\}$, are the six singly occupied carbon π orbitals, one for 29 each electron. The spin function for the six electrons in the six carbon 30 π orbitals, Θ_0^0 , 0, is a linear combination of the five linearly independent 31 32 ways that the spins of these six electrons can be coupled to yield a state with S = 0 and $M_S = 0$.^[10] Using the Rumer spin functions, these 33 34 spin couplings correspond to the two Kekulé spin couplings and the 35 three Dewar (para-bonded) spin couplings. In the SCGVB calculation all of the orbitals, including the doubly occupied σ orbitals, as well as 36 the coefficients of the five spin functions are fully optimized.

38 We also considered three additional wave functions for the electrons in the π orbitals: (a) the RHF wave function,^[11] which restricts 39 the π orbitals to be doubly occupied; (b) the CASSCF wave 40 function,^[12] which includes all of the configurations generated by dis-41 tributing the six electrons in the six π orbitals (i.e., full configuration 42 interaction in that space); and (c) a CCSD(T) wave function^[13,14] corre-43 44 lating only the electrons in the π system, leaving the orbitals in the σ 45 system doubly occupied. We also report the results of CCSD(T) calcu-46 lations for all of the valence electrons to help establish the "ground 47 truth" for the variation of the energy of benzene with the distortion 48 angle.

Following the approach used by Small and Head-Gordon^[3] see 49 also Lawler et al., $^{[6]}$ we define a deformation angle, $\Delta \theta_{\text{CXC}}$, which is a 50 measure of the deviation from the D_{6h} structure of benzene. A value 51 52 of $\Delta \theta_{CXC} = 0^{\circ}$ corresponds to the D_{6h} structure, while a nonzero value of $\Delta \theta_{CXC}$ gives adjacent bond angles of $\Delta \theta_{CXC+} = (60^{\circ} + \Delta \theta_{CXC})$ and 53



FIGURE 1 Definition of geometrical parameters for benzene [Color figure can be viewed at wileyonlinelibrary.com]

 $\Delta \theta_{CXC-}$ = (60° – $\Delta \theta_{CXC}$); see Figure 1. The resulting adjacent CC bond lengths are given by $R_{CC+} = 2R_{CX}\sin(\frac{1}{2}\theta_{CXC+})$ and $R_{CC-} = 2R_{CX}\sin(\frac{1}{2}\theta_{CXC+})$ $(\frac{1}{2}\theta_{CXC})$. Note that for D_{6h} geometries, $R_{CC} = R_{CX}$.

Most of the calculations presented in this study were performed with the Molpro suite of quantum chemical programs (version 2010.1).^[15,16] In particular, the CASVB module in Molpro was used to perform most of the SCGVB calculations^[17,18] with additional results obtained using an older program.^[19]

RESULTS AND DISCUSSION 3

In Table 1, we list the CC and CH bond lengths, total energies, and energy differences, relative to the SCGVB energy, for the aug-cc-86 pVTZ basis sets at $\Delta \theta_{CXC} = 0^{\circ}$. The first four calculations in this table 87 refer to calculations on the π system of benzene; the last calculation 88 includes all of the electrons in the valence orbitals. For each of the 89 methods of interest we optimized R_{CX} and R_{CH} for $\Delta \theta_{CXC} = 0^{\circ}$, with 90 91 $R_{\rm CC} = R_{\rm CX}$.

The variation in R_{CH} is very modest for all of the methods that 92 treated only the π system: RHF (1.0733 Å), SCGVB (1.0732 Å), CAS-93 SCF (1.0733 Å), and CCSD(T) (1.0734 Å). As expected, the variation in 94 R_{CX} was somewhat larger, with the RHF method predicting 95 R_{CX} = 1.3829 Å and the SCGVB and CASSCF methods predicting 96 1.3931 Å and 1.3921 Å, respectively. The CCSD(T) calculations on the 97 π system predicted a value of R_{CX} in between these three values, 98 1.3888 Å, that is, a longer bond than predicted by the RHF calcula-99 tions and a shorter bond than predicted by the SCGVB and CASSCF 100 calculations. The CCSD(T) calculations for the full valence space, 101 $(\sigma + \pi)$, predicted larger values for both R_{CH} (1.0839 Å) and R_{CX} 102 (1.3980 Å). 103

The energy of the RHF wave function is 39.4 kcal mol⁻¹ higher 104 than that of the SCGVB wave function. This is a measure of the non-105 dynamical correlation energy in the π system of benzene. The CASSCF 106

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TABLE 1 Structure and energies for the RHF, SCGVB, CASSCF, and CCSD(T) wave functions of benzene (in Hartrees) at $\Delta\theta_{CXC} = 0^{\circ}$. Distances (R_{CC} , R_{CH}) are in Ångstroms, and total energies (E_b) are in Hartrees. Except for the last row, the relative energies are with respect to the SCGVB energy and are in kcal mol⁻¹. Basis set: aug-cc-pVTZ

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	Method	$R_{\rm CC} = R_{\rm CX}$	R _{CH}	E _h	ΔΕ	57
π -System Only ^a	RHF	1.3829	1.0733	-230.782351	39.40	58
	SCGVB	1.3931	1.0732	-230.845140	0.0	59
	CASSCF	1.3921	1.0733	-230.852789	-4.80	60
	CCSD(T)	1.3888	1.0734	-230.899220	-33.94	61
Full valence	CCSD(T)	1.3980	1.0839	-231.820465	-578.09 ^b	62

^aAll of the electrons were included in the π -System Only calculations, although the σ orbitals were kept doubly occupied. ^bRelative to the CCSD(T) calculations for the π -System Only.



FIGURE 2 Contour plots of one of the six SCGVB π orbitals for benzene: (a) top view of the orbital and (b) side view of the orbital. Contours are shown from 0.05 to 0.25 in increments of 0.05 [Color figure can be viewed at wileyonlinelibrary.com]

energy, on the other hand, is only 4.8 kcal mol^{-1} lower than the 30 SCGVB energy. Thus, the SCGVB wave function accounts for 89.1% 31 32 of the difference between the CASSCF and RHF energies, very similar 33 to the 89.6% reported by Karadakov and Cooper for the 6-311G(d,p) basis set.^[20] The CCSD(T) calculations for the π system of benzene 34 35 yielded an energy 33.9 kcal mol⁻¹ lower than the SCGVB energy. This is a measure of the dynamical correlation energy of the benzene π sys-36 tem. Thus, in this case the non-dynamical correlation energy is slightly 38 larger than the dynamical correlation energy. For comparison, the 39 CCSD(T) calculations on the full valence ($\sigma + \pi$) system gives an energy that is 578.1 kcal mol⁻¹ lower than the CCSD(T) calculation on the π 40 41 system alone.

One of the SCGVB π orbitals in benzene is plotted in Figure 2. The 422 43 other five orbitals can be obtained by rotating this orbital around the ring in 60° increments. This orbital is very similar to the SCGVB orbitals 44 reported earlier for benzene by the Gerratt group.^[21-23] As can be seen, 45 the SCGVB π orbital it is largely localized on one of the carbon atoms, but 46 47 is polarized toward the carbon atoms on each side of that atom. In this 48 way, the SCGVB π orbitals of benzene reflect the bonding motif found in 49 the Kekulé spin couplings, where each carbon atom is bonded to both 50 neighboring carbon atoms in the combined Kekulé spin couplings.

Figure 3 is a plot of the relative energies of the distorted benzene 5**1F3** 52 molecule, referenced to the energy at $\Delta \theta_{CXC} = 0^{\circ}$ (D_{6h} symmetry) for the RHF, SCGVB, CASSCF, and CCSD(T) calculations on the π system 53



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FIGURE 3 Variation of ΔE with the distortion angle, $\Delta \theta_{CXC}$, for the RHF, SCGVB, CASSCF, and CCSD(T) wave functions for the π system. (R_{CX} , R_{CH}) have been fixed at the optimum values for each method at $\Delta \theta_{CXC} = 0^{\circ}$ [Color figure can be viewed at wilevonlinelibrary.com]

with the values of R_{CX} and R_{CH} fixed at the optimum values for $\Delta \theta_{CXC} = 0^{\circ}$. Note that the curves from the RHF and CASSCF calculations are virtually on top of one another. We also carried out full valence CCSD(T) calculations as a function of $\Delta \theta_{CXC}$; the resulting curve is essentially indistinguishable from that for the CCSD(T) calculations on the π system only and, therefore, is not shown. The first feature to note is that the energies of the RHF, CASSCF, and CCSD(T) wave functions increase monotonically with increasing $\Delta \theta_{CXC}$, that is, the optimum geometries are for $\Delta\theta_{CXC}$ = 0°, yielding a D_{6h} geometry for benzene. The curves for the RHF and CASSCF calculations lie essentially on top of one another, while that for the CCSD(T) calculations is only slightly higher. The energy of the SCGVB wave function, on the other hand, initially decreases with increasing $\Delta \theta_{CXC}$, having a minimum of -0.040 kcal mol⁻¹ at $\Delta\theta_{CXC} = 0.31^{\circ}$. The resulting optimum bond lengths are: R_{CC+} = 1.4004 Å and R_{CC-} = 1.3861 Å, that is, a difference of $\Delta R_{\rm e}$ = 0.0143 Å. This is in basic agreement with the

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FIGURE 4 The difference in the CASSCF and SCGVB energies as a function of the distortion angle, $\Delta \theta_{CXC}$. (R_{CX} , R_{CH}) have been fixed at the optimum values for each method at $\Delta \theta_{CXC} = 0^{\circ}$

findings of Small and Head-Gordon,^[3] who reported a minimum of approximately -0.047 kcal mol⁻¹ at $\Delta\theta_{CXC} \approx 0.39^{\circ}$.

We also carried out SCGVB calculations that optimized R_{CX} and R_{CH} for each distortion angle, $\Delta \theta_{CXC}$. The changes in R_{CX} and R_{CH} were sufficiently small, for example, ~10⁻⁴ Å for R_{CX} and ~10⁻⁶ Å for R_{CH} at $\Delta \theta_{CXC} = 0.31^{\circ}$, that any changes from the results described above were negligible.

To gain insights into the underlying reason or reasons that the SCGVB calculations predict a non-D_{6h} structure for benzene, we first calculated the difference between the energies of the CASSCF and SCGVB wave functions as a function of $\Delta \theta_{CXC}$. This difference is plot-34=4 ted in Figure 4 and provides no obvious reason for the unusual depen-dence of the SCGVB energy on distortion angle: the energy difference increases steadily from $\Delta \theta_{CXC} = 0^{\circ}$ to $\Delta \theta_{CXC} = 1^{\circ}$, increas-ing nearly linearly in the vicinity of $\Delta \theta_{CXC} = 0.31^{\circ}$.

Next, we calculated separately the energies of the Kekulé and Dewar components of the SCGVB wave function using the optimum SCGVB orbitals for each distortion angle. The energies of the resulting SCGVB(Kekulé) and SCGVB(Dewar) wave functions increase with increasing $\Delta \theta_{CXC}$, predicting a D_{6b} geometry for benzene, although their **3-5** behavior near $\Delta \theta_{CXC} = 0^{\circ}$ is certainly unusual (see Figure 5). Given that, individually, the two components of the full SCGVB wave function pre-dict that benzene has a D_{6h} geometry, we are led to conclude that the cause of the unusual non-D_{6h} geometry predicted by the SCGVB calcu-lations seems to be a result of the interaction between the Kekulé and Dewar components of the full SCGVB wave function.

We then calculated the interaction between the Kekulé and Dewar components of the SCGVB wave function (H_{12}) as well as the overlap of these components (S_{12}) . These results, as a function of the distortion angle, are plotted in Figure 6. Again, there is little hint in these plots about the cause of the non-D_{6h} geometry of benzene 

FIGURE 5 Variation of ΔE with the distortion angle, $\Delta \theta_{CXC}$, for the SCGVB wave function and its Kekulé and Dewar components. The optimum orbitals from the full SCGVB calculations are used in all calculations



FIGURE 6 Dependence on distortion angle, $\Delta \theta_{CXC}$, of the variation in the interaction energy, H_{12} (solid line), and overlap, S_{12} (dashed line), for the Kekulé and Dewar components of the benzene SCGVB wave function

predicted by the SCGVB wave function, although the magnitude of 100 the overlap of the Kekulé and Dewar components (0.992–0.980) is 101 surprisingly large and may be implicated in the unusual prediction of 102 the non-symmetric structure for benzene. With this in mind, we also 103 examined the energy of the orthogonal complement to the Kekulé 104 space and the value of the corresponding off-diagonal Hamiltonian 105 matrix element. We found for both quantities that the variation with 106 1

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FIGURE 7 Variation of ΔE with the distortion angle, $\Delta \theta_{CXC}$, for 20 the SCGVB+ (ion = 2) wave function with SCGVB orbitals (blue. dashed) and SCGVB+ (ion = 2) wave function with optimized orbitals 22 (red, solid) [Color figure can be viewed at wileyonlinelibrary.com]

24 distortion angle, $\Delta \theta_{\text{CXC}}$ is smooth and monotonic with no unusual 25 features near either $\Delta \theta_{CXC} = 0^{\circ}$ or $\Delta \theta_{CXC} = 0.31^{\circ}$.

26 Finally, we investigated the addition of ionic terms to the SCGVB wave function, a topic explored earlier by Van Voorhis and Head-Gor-27 don.^[5] As first noted by Coulson and Fischer,^[24] the SCGVB wave func-28 tion largely accounts for singly ionic structures, so that the first ionic 29 30 configurations of importance are the doubly ionic ones. If we simply add 31 the doubly ionic configurations to the SCGVB wave function, we obtain the dashed curve in Figure 7, which has a remarkable resemblance to the 327 corresponding curve for the Kekulé wave function in Figure 5. The 33 34 energy of this wave function is 0.47 kcal mol⁻¹ higher than the CASSCF 35 energy. If we optimize the orbitals for the SCGVB+ (ion = 2) wave func-36 tion, the energy decreases to -230.852311 Eh, which is 0.30 kcal mol⁻¹ 37 higher than the CASSCF energy. Furthermore, the SCGVB+ (ion = 2) 38 wave function predicts that benzene has a D_{6h} geometry. In fact, the 39 dependence of the energy on distortion angle for the SCGVB+ (ion = 2) wave function, see Figure 7, is essentially identical to that for the CAS-40 SCF wave function, differing by just 0.009 kcal mol⁻¹ at $\Delta \theta_{CXC} = 1^{\circ}$. 41 42 Clearly, higher ionic configurations make only minor improvements to 43 the SCGVB+ (ion = 2) wave function. Further numerical experiments indi-44 cated that the ion = 2 structures in which the negative charges are para 45 to one another are of particular importance in reproducing the correct 46 dependence of the energy on $\Delta \theta_{CXC}$.

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CONCLUSIONS 4 49

Small and Head-Gordon^[3] found that the SCGVB wave function for 51 52 benzene predicted a non-D_{6h} geometry, a surprising result given that 53

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the combinations of the Kekulé and Dewar spin couplings in the 54 SCGVB wave function can describe all CC bonds equally. The distor-55 tion from a D_{6h} geometry was very small ($\Delta \theta_{CXC} = 0.39^{\circ}$) as was the 56 energy lowering, $\Delta E = -0.047$ kcal mol⁻¹. Since those calculations 57 used a very small basis set for the carbon and hydrogen atoms (STO-58 3G), we decided to repeat this study, obtaining similar results with 59 an aug-cc-pVTZ basis set, namely, $\Delta \theta_{\text{CXC}}$ = 0.31° and ΔE = 60 -0.040 kcal mol⁻¹. The aug-cc-pVTZ basis set should provide results 61 very close to the complete basis set limit for the SCGVB method, so 62 this puzzling anomaly is an inherent feature of the SCGVB description 63 of benzene. 64

In an attempt to understand the reason for this unusual prediction 65 by the SCGVB method, we separated the full SCGVB wave function 66 into its two components, the two Kekulé and three Dewar modes of 67 spin coupling, and determined the dependence of the energies of 68 these two components on the distortion angle, $\Delta \theta_{CXC}$, using the opti-69 mum SCGVB orbitals for that angle. We found that the Kekulé and 70 Dewar energies increased with increasing $\Delta \theta_{CXC}$, predicting that ben-71 zene has a D_{6h} geometry, although the behavior of the resulting cur-72 ves near $\Delta \theta_{CXC} = 0^{\circ}$ was unusual and unexpected. These results 73 indicate that the non-D_{6h} geometry predicted by the SCGVB calcula-74 tions could be a result of the interaction between the Kekulé and 75 Dewar components of the full SCGVB wave function. However, 76 examination of the dependence of this interaction, as well as the over-77 lap of the two components, did not provide any insights into the 78 underlying cause of this curious anomaly in the SCGVB description of 79 benzene. 80

In agreement with Van Voorhis and Head-Gordon,^[5] we found 81 that adding the doubly ionic configurations to the SCGVB wave func-82 tion (the singly ionic configurations are already largely subsumed in 83 the SCGVB wave function) led to the prediction of a D_{6h} geometry for 84 benzene. In fact, if the orbitals in the SCGVB+ (ion = 2) wave function 85 are optimized, the total energy as well as the energies as a function of 86 distortion angle, $\Delta \theta_{CXC}$, are essentially identical to those of the CAS-87 SCF wave function. The addition of the doubly excited configurations 88 to the SCGVB wave function clearly corrects the deficiencies in the 89 SCGVB description of benzene, although the orbitals must be 90 reoptimized. Analyzing the doubly ionic configurations contributing to 91 the SCGVB+ (ion = 2) wave function, we found that those configura-92 tions with the negative charges para to one another were of particular 93 importance in reproducing the correct dependence of the energy 94 95 on $\Delta \theta_{CXC}$.

ACKNOWLEDGMENTS

This research was supported in part by the Center for Scalable Pre-98 dictive Methods for Excitations and Correlated phenomena (SPEC), 99 which is funded by the U.S. Department of Energy, Office of Sci-100 ence, Basic Energy Sciences, Chemical Sciences, Geosciences and 101 Biosciences Division, as part of the Computational Chemical Sci-102 ences Program at Pacific Northwest National Laboratory. We 103 thank one of the reviewers for bringing the work of Van Voorhis 104 and Head-Gordon to our attention. 105

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ORCID

2 Thom H. Dunning Jr D https://orcid.org/0000-0002-3290-6507

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How to cite this article: Xu LT, Cooper DL, Dunning Jr TH. Resolving a puzzling anomaly in the spin-coupled generalized valence bond description of benzene. *J Comput Chem*. 2020;

1-6. https://doi.org/10.1002/jcc.26185

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