

Is the S_2N_2 Ring a Singlet Diradical? Critical Analysis of Alternative Valence Bond Descriptions

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Rival valence bond (VB) descriptions are investigated for the π -electron system of the S_2N_2 ring and of another sulfur-nitrogen ring, $S_4N_4^{2+}$, near equilibrium geometry. The lowest-energy compact spin-coupled generalized VB (SCGVB) description is provided in each case by the variational optimization of two configurations that are found to be symmetry related to one another. Optimization instead of symmetry-pure single-configuration SCGVB wavefunctions leads to the involvement of three-center SNS or NSN orbitals, which seems to be an unnecessary complication. In neither case is very much achieved from the mixing of competing solutions. Breathing orbital VB (BOVB) calculations for S_2N_2 confirm a structure with NN singlet diradical character to be more important than one with SS singlet diradical character, but the largest contribution (ca. 60%) turns out to be due to the symmetry-determined linear combination of four symmetry-equivalent structures that lack any obvious diradical character. Much the same pattern was consistently found when we used a simple but robust projection of our various SCGVB wavefunctions for S_2N_2 onto the basis of BOVB structures (plus an orthogonal complement).

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Introduction

Although various larger (SN)_x systems continue to attract considerable experimental and theoretical attention for a wide range of reasons, we should admit from the outset that our own particular interest in the planar S₂N₂ ring is associated with the somewhat unusual levels of attention that this small molecule has been receiving from theoretical chemists, particularly those specializing in valence bond (VB) theory. A spincoupled generalized VB (SCGVB, vide infra) description of S₂N₂, reported by Gerratt et al., [1] was interpreted at the time as being dominated by SS singlet diradical character but some other studies, [2],[3],[4],[5],[6] and especially the very careful breathing orbital valence bond (BOVB) work of Braïda et al., [2] have seriously questioned the validity of that interpretation. Indeed, Braïda et al.[2] were able to show by partly qualitative arguments that the SCGVB wavefunction, when stripped to some extent of the delocalization tails on the fully-optimized active orbitals, did in fact appear to be consistent with the BOVB viewpoint, in which a structure with NN singlet diradical character was the largest single contributor. We re-examine this issue in the present work, by means of a novel approach utilizing a robust projection of the SCGVB wavefunction onto the basis of the BOVB structures. While confirming the main gist of the conclusions of Braïda et al., [2] namely that Gerratt et al.[1] may have misinterpreted the results of their SCGVB calculations, we also show that the interpretation of the BOVB description could be interestingly more nuanced than had been supposed. Additionally, as was shown by Thorsteinsson et al.,[7] there can exist rival SCGVB-like solutions that are fairly close in energy to the one described by Gerratt et al..[1] We explore this issue further and then require different combinations of SCGVB solutions to fight it out without any prejudice in a variational boxing ring, thereby obtaining further insight into the bonding in the π -electron system of S_2N_2 . We then examine briefly certain analogous rival SCGVB descriptions of the corresponding π -electron system in the S₄N₄²⁺ ring.

Theoretical and computational details

The single-configuration SCGVB wavefunction for the valence $\pi\text{-space}$ of S_2N_2 is based on a single product of six singly-occupied nonorthogonal active orbitals π_μ that are expanded in the full basis set and it can be written in the following form: $^{[8]}$

$$\Psi_{\text{SCGVB}} = \mathcal{A} \left[\left(\prod_{i=1}^{18} \varphi_i \alpha \varphi_i \beta \right) \left(\prod_{\mu=1}^{6} \pi_{\mu} \right) \Theta_0^6 \right]$$
(1)

in which the φ_i are doubly-occupied inactive orbitals that accommodate the 'core' $S(2p_\pi)$ electrons and all of the σ system. The active-space total spin function Θ_0^6 is expanded in the full spin space of five linearly-independent modes of coupling together the spins of six electrons so as to achieve an overall singlet state, with the expansion coefficients known as spin-coupling coefficients.

Traditionally, wavefunctions of this type, as introduced by Gerratt, [9] have mostly been termed spin-coupled (SC or even SCVB), with an acknowledgement that the construction is in fact entirely equivalent to that of *full* generalized valence bond (full-GVB), as introduced by Goddard, [10] or they have been called (full-)GVB, with a mention of the equivalence to SC (or SCVB). It seems to the present authors to be undesirable to persist with different names for essentially identical calculations that might even have been carried out with the same codes. Accordingly, we use here instead the compound term spin-coupled generalized valence bond (SCGVB) that aims to encompass both sets of names, and we recommend that others consider doing the same.

Following fairly closely the construction used by Braïda et al., [2] our BOVB calculations for S₂N₂ were carried out using six VB structures (see Fig. 1). BOVB active orbitals can be described as 'strictly localized', in the sense that each of them is restricted so as to use only the basis functions that are associated with one atomic center. Accordingly, in this work, the symmetry-unique active π orbitals in each BOVB structure were optimized simultaneously as entirely separate linear combinations of the basis functions centered on the relevant nuclei. Where there are two electrons associated with the same center, they are accommodated with opposing spins in the same orbital. As can be seen from Fig. 1, BOVB structure 1 clearly corresponds to NN singlet diradical character and BOVB structure 2 to SS singlet diradical character. The four symmetry-equivalent BOVB structures **3** to **6** each feature instead one S–N π bond. (We subsequently also carried out S-BOVB calculations in which each of the doubly occupied active orbitals was allowed to 'split' into two singly occupied orbitals, but maintaining singlet coupling for the two orbitals.)



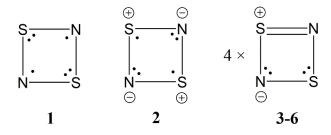


Figure 1. Structures used in BOVB calculations for S₂N₂.

All of our electronic structure calculations for S_2N_2 were carried out for the nuclear geometry and orientation shown in Fig. 2 and using the standard cc-pVQZ basis set. We have intentionally chosen the same idealized square geometry, close to experiment, that was used by Gerratt et al., [1] but we are confident that all of our key findings will be relatively insensitive to small changes to this geometry.

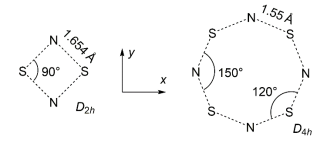


Figure 2. Geometries and orientations used for S_2N_2 and $S_4N_4^{2+}$.

Instead of optimizing each time the various inactive orbitals in our various VB descriptions of S_2N_2 , we have chosen to take those orbitals without any further reoptimization from an appropriate CASSCF description that should not introduce any significant bias for or against the various competing VB descriptions of the valence π systems. Following various numerical tests of different choices of CASSCF inactive spaces (see Table S1 in the Supporting Information) we selected a six electrons in eight orbitals expansion spanning $3B_{1u}+2B_{2g}+2B_{3g}+1A_u$, which we abbreviate to [3,2,2,1].

Note that we were unable in our BOVB and S-BOVB calculations to orthogonalize the active π orbitals to the two fixed 'core' π MOs (B_{1u}+B_{2g}) taken from the CASSCF, because of the strict localization constraints on the active orbitals. In order to avoid numerical problems, we identified in the S atomic basis set the 2p π contraction which contributes most to the inactive π orbitals and then constrained to zero its coefficients in

the expansions of the active orbitals. Such constraints were not necessary in any of the SCGVB calculations for S_2N_2 . We did, however, check that changes to SCGVB energies upon the application of such constraints are negligible.

Our subsequent SCGVB calculations for the valence π -space of the planar $S_4N_4^{2+}$ ring were carried out for the geometry and orientation shown in Fig. 2 and again using the standard cc-pVQZ basis set. The SN nuclear separation, the bond angles and the molecular point group (D_{4h}) were based on the crystallographic studies of Gillespie et al.^[11] but, as in the case of S_2N_2 , we do not expect any of our key findings to be sensitive to small changes to this geometry. Our single-configuration SCGVB wavefunction for the valence π -space of $S_4N_4^{2+}$ is based on a single product of ten singly-occupied nonorthogonal active orbitals π_μ that are expanded in the full basis set and it can be written in the following form: [8]

$$\Psi_{\text{SCGVB}} = \mathcal{A} \left[\left(\prod_{i=1}^{40} \varphi_i \alpha \varphi_i \beta \right) \left(\prod_{\mu=1}^{10} \pi_{\mu} \right) \Theta_0^{10} \right]$$
(2)

in which the active-space total spin function Θ_0^{10} is expanded in the full spin space, consisting of 42 linearly-independent modes of coupling together the spins of ten electrons so as to achieve an overall singlet state. For all of our frozen-core VB calculations on $S_4N_4^{2^+}$ we took the doubly-occupied inactive orbitals φ_i that accommodate the 'core' $S(2p_\pi)$ electrons and all of the σ system from a full-valence π -space CASSCF(10,8) wavefunction, without further optimization.

For a normalized wavefunction Ψ that is expressed as a linear combination of nonorthogonal VB structures or configurations Φ_k with expansion coefficients c_k , it is most usual to assess the relative importance of the various Φ_k according to their Chirgwin-Coulson weights, W_k , which may be defined according to: [12]

$$W_k = c_k \sum_l c_l \langle \Phi_k | \Phi_l \rangle \tag{3}$$

in which the W_k sum to unity, so as to satisfy the normalization condition for Ψ . This widely-used definition of weights has a number of useful properties except that, especially in the case of high values of the overlaps $S_{kl} = \langle \Phi_k | \Phi_l \rangle$, individual values of W_k can occasionally fall outside the physically-meaningful



range of 0 to 1. In addition to the Chirgwin-Coulson scheme, we also make some use in the present work of the inverse-overlap definition of Gallup and Norbeck:^[13]

$$w_k = |c_k|^2 / (S^{-1})_{kk}$$
(4)

where the values of w_k are usually renormalized so as to add to unity. Such values then necessarily lie in the physically-meaningful range (0 to 1).

The workhorse for all of the VB calculations reported here was the generalized multiconfiguration spincoupled (GMCSC) program developed bν Penotti^{[14],[15],[16],[17]} with CASSCF inactive orbitals and the required integrals over basis functions generated the GAMESS-US package.^{[18],[19]} depictions of SCGVB active orbitals were produced using Virtual Reality Markup Language (VRML) files generated with Molden. [20] Quantum theory of atoms in molecules (QTAIM) analysis^[21] was performed using AIMAII^[22] and with our own codes. Additional π -space CASSCF calculations were carried out in D_{2h} symmetry using MOLPRO.[23],[24]

Results and discussion

S_2N_2

It is useful to consider first a single-configuration SCGVB wavefunction for S2N2 that is based on active orbitals which span the space denoted a by Thorsteinsson et al.^[7] (vide infra) and which corresponds directly to the solution described by Gerratt et al.. [1] We found in the present work that certain symmetry relations amongst the active orbitals appeared spontaneously during the optimization of our frozen-core SCGVB(a) wavefunction for S_2N_2 , such that orbitals π_3 and π_4 can be generated from π_1 and π_2 , respectively, by reflection in the σ_{xz} mirror plane, whereas π_6 can be generated from π_5 by reflection in the σ_{vz} mirror plane. Spontaneity of this type is usually a good indication of wavefunction stability with respect to breaking spatial symmetry. Other symmetry properties of the active orbitals, such as the invariance of π_1 and π_2 to reflection in the σ_{vz} mirror plane and of π_5 to reflection in the σ_{χ_Z} mirror plane, also arose spontaneously. The symmetry-unique active orbitals π_1 , π_2 and π_5 , as depicted in the top row of Fig. 3, clearly include a three-center SNS function that potentially makes it relatively difficult to interpret this wavefunction directly and unambiguously in terms of the sorts of VB structures that are shown in Fig. 1. Nonetheless, Gerratt et al.^[1] used the forms of such SCGVB orbitals, together with the pattern of active-space spin coupling, to assert the dominance of SS singlet diradical character. As was mentioned in the Introduction, such an interpretation is clearly at odds with the BOVB work of Braïda et al..^[2] Indeed, those authors were able to use mostly qualitative arguments, stripping away 'delocalization tails', in order to suggest that the SCGVB wavefunction of Gerratt et al.^[1] could in fact be more consistent with the NN singlet diradical character that was observed in BOVB calculations than with the original claim of SS singlet diradical character.

We use here a robust numerical approach to establish the links between various SCGVB wavefunctions and BOVB descriptions. For this purpose, we turn now to our BOVB results, carried out with the same basis set and with the same choice of frozen core as our SCGVB calculations, so that we may compare like with like. Examining our frozen-core BOVB results, it can clearly be seen from Table 1 that the single BOVB structure with the lowest energy is structure 1 (see Fig. 1), whether we take the active orbitals directly from the BOVB wavefunction or perform further separate optimizations for each structure. On the other hand, the Chirgwin-Coulson weight of structure 1 in the BOVB wavefunction turns out to be just 30.1% (see top row of values in Table 2), so that it corresponds to a minority of the total wavefunction. A somewhat lower energy is given by the symmetry-determined linear combination of the four symmetry-equivalent structures 3-6, with a Chirgwin-Coulson weight of 59.0%. These various findings are on the whole rather similar to those reported by Braïda et al.^[2] and the general conclusions also turn out to be much the same from our corresponding frozen-core S-BOVB calculations (see Tables S4 and S6 in the Supporting Information). Additionally, the patterns of weights obtained with the Gallup-Norbeck scheme (see Tables S3 and S6 in the Supporting Information) are much the same as those from the Chirgwin-Coulson definition. The energy difference between structure 3 on its own and the symmetry-determined linear combination **3-6** clearly indicates significant resonance stabilization; this is, of course, strongly reminiscent of the energy stabilization in benzene that can be associated with the resonance of two dominant Kekulé structures.



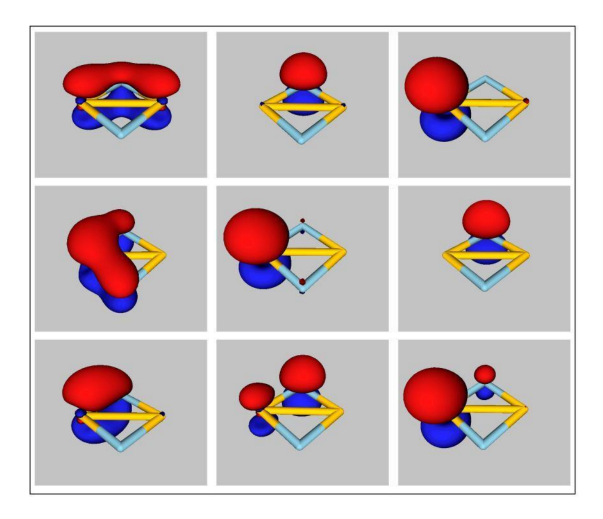


Figure 3. Symmetry-unique active orbitals π_1 , π_2 and π_5 (left to right) for frozen-core S_2N_2 wavefunctions: SCGVB(**a**) (top row); SCGVB(**b**) (middle row); SCGVB(**C**) (bottom row).

Table 1. Energies (in hartree) from frozen-core sixstructure BOVB calculations for S_2N_2 , where 'Fixed' signifies the use of active orbitals taken directly from the six-structure BOVB calculation and 'Relaxed' signifies further optimization.

Structures	Fixed	Relaxed
1	-903.79363	-903.81387
2	-903.65901	-903.69344
3	-903.70564	-903.71732
3-6	-903.88008	-903.88919
1-6	-903.94635	-903.94635

Although it is certainly true that structure ${\bf 1}$ is indeed the most important one in our BOVB or S-BOVB wavefunctions, the majority of each of those wavefunctions is instead associated with the symmetry-determined linear combination (Φ_{3-6}) of the four symmetry-equivalent structures ${\bf 3-6}$. Clearly, though, the interpretation put forward by Gerratt et al., $^{[1]}$ based on claims of dominant SS singlet diradical character in their SCGVB wavefunction, remains distinctly anomalous. With this in mind, it proves to be very

informative to use a fairly simple but robust numerical approach, that we now outline, to project the compact SCGVB(a) solution (based on just a single product of active orbitals) onto the corresponding BOVB representation.

Table 2. Chirgwin-Coudirectly from the froze S_2N_2 and where \boldsymbol{X} sign	n-core six-	structure	BOVB calc	ulations for
Wavefunction	1	2	3-6	Х
BOVB (1-6)	30.1%	10.9%	59.0%	_
SCGVB(a)	25.1%	13.2%	56.7%	5.0%
SCGVB(b)	27.1%	13.0%	54.7%	5.2%
SCGVB(C)	24.6%	13.3%	57.3%	4.7%
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	24.9%	13.3%	57.2%	4.6%

In general terms, we wish to consider the expansion of a normalized wavefunction Ψ in the following form:

24.6%

$$\Psi = d_1 \Phi_1 + d_2 \Phi_2 + d_{3-6} \Phi_{3-6} + d_X \Phi_X$$
(5)

12.0%

 $GMCSC(a \oplus b \oplus C)$

4.9%



in which Φ_X is envisaged as a Ψ -dependent normalized entity which is orthogonal to each of the normalized BOVB structures Φ_1 , Φ_2 and Φ_{3-6} . In other words, $d_X\Phi_X$ denotes any aspects of Ψ that cannot be expressed as a linear combination of the BOVB structures. It follows directly from Eq. (5) that:

$$\begin{pmatrix} \langle \Phi_{1} | \Psi \rangle \\ \langle \Phi_{2} | \Psi \rangle \\ \langle \Phi_{3-6} | \Psi \rangle \end{pmatrix} = \begin{pmatrix} 1 & \langle \Phi_{1} | \Phi_{2} \rangle & \langle \Phi_{1} | \Phi_{3-6} \rangle \\ \langle \Phi_{1} | \Phi_{2} \rangle & 1 & \langle \Phi_{2} | \Phi_{3-6} \rangle \\ \langle \Phi_{1} | \Phi_{3-6} \rangle & \langle \Phi_{2} | \Phi_{3-6} \rangle & 1 \end{pmatrix} \begin{pmatrix} d_{1} \\ d_{2} \\ d_{3-6} \end{pmatrix} \tag{6}$$

and so, given that we can calculate all of the overlap integrals that appear in Eq. (6) (see Tables S2 and S5 in the Supporting Information), it is very straightforward to solve for d_1 , d_2 and $d_{3\text{-}6}$, and then to compute the Chirgwin-Coulson weights W_k . The corresponding weight in Ψ of the normalized orthogonal complement Φ_X , i.e. $W_X = d_X^{\ 2}$, is most simply obtained from the requirement that the Chirgwin-Coulson weights must sum to unity. (This does of course correspond exactly to determining d_X using the normalization condition for Ψ .) Our overall scheme is entirely equivalent to the application of a projection operator \mathcal{P} , defined according to:

$$\mathcal{P} = \sum_{k,l} |\Phi_k\rangle (\mathbf{\Delta}^{-1})_{kl} \langle \Phi_l| \qquad (k,l = 1,2,3-6)$$
(7)

in which Δ is the 3×3 overlap matrix shown in Eq. (6). The Ψ -dependent orthogonal complement, which we have chosen here to denote as $d_X \Phi_X$, is then simply $(1-\mathcal{P})\Psi$.

Using the scheme that we have just described, the projection of the SCGVB(a) solution onto the basis of BOVB structures leads to the Chirgwin-Coulson weights that are reported in the second row of values in Table 2. The interpretation put forward by Gerratt et al. [1] does indeed turn out to be erroneous given that the single most important BOVB structure in SCGVB(a) is clearly 1, albeit with a weight that is slightly lower than in our total BOVB wavefunction, with BOVB structure 2 (corresponding to SS singlet diradical character) being far less important. Just as we observed for the total BOVB wavefunction, it is the symmetry-determined linear combination of the four symmetry-equivalent BOVB structures 3-6, with a Chirgwin-Coulson weight of

56.7%, which accounts for more than a half of the SCGVB(a) wavefunction. We find that the normalized orthogonal complement (Φ_X) , *i.e.* the part of the SCGVB(a) solution that is not described by this set of BOVB structures, has a weight of just 5.0%. (All of our key observations are much the same when using Gallup-Norbeck weights and/or if we project instead onto S-BOVB structures – see Tables S3, S4 and S6 in the Supporting Information.)

We now return to the observation of Thorsteinsson et al.^[7] that there can exist rival SCGVB-like solutions that are fairly close in energy to SCGVB(a). Whereas the SCGVB(a) active orbitals span $3B_{1u}+1B_{2q}+2B_{3q}+0A_u$, which we may abbreviate to **a**=[3,1,2,0], Thorsteinsson et al.^[7] suggested that there are various energetically nearby solutions which span various alternative distributions, including b=[3,2,1,0] and c=[2,1,2,1]. We find that the corresponding CASSCF(6,6) energies for active spaces a and c (see Table S1 in the Supporting Information) are particularly close to one another (differing by less than 0.35 millihartree), with the CASSCF(6,6) energy for active space **b** being inferior by 8.4 millihartree. In order to optimize the corresponding SCGVB(b) wavefunction, without it returning to the SCGVB(a) solution, we imposed the following constraints on the active orbitals: $\pi_3 = \hat{\sigma}_{vz}\pi_1$, $\pi_4 =$ $\hat{\sigma}_{yz}\pi_2$ and $\pi_6=\hat{\sigma}_{xz}\pi_5$. Other symmetry properties of the active orbitals, such as the invariance of π_1 and π_2 to reflection in the σ_{xz} mirror plane and of π_5 to reflection in the σ_{vz} mirror plane, arose spontaneously during the optimization. To a large extent, the resulting symmetry-unique active orbitals π_1 , π_2 and π_5 for the SCGVB(b) solution (shown in the middle row of Fig. 3) are somewhat reminiscent of those for SCGVB(a), except that the symmetry-unique three-center active orbital is now over NSN rather than SNS and it exhibits a larger contribution from the central atom of the triad. Just as we might have anticipated from the corresponding CASSCF(6,6) energies, the SCGVB(b) solution is indeed energetically inferior to SCGVB(a), with the difference being 8.8 millihartree (see Table 3).

It proves to be especially informative to consider the projection of this SCGVB(b) solution onto the BOVB basis because the resulting Chirgwin-Coulson weights, as reported in Table 2, turn out to be fairly similar to those that we have described above for SCGVB(a). In spite of the differences in the forms and, particularly, the locations of the three-center active orbitals, these two wavefunctions do in fact turn out to be rather similar in terms of their BOVB character. Furthermore



the overlap between the total SCGVB(a) and SCGVB(b) wavefunctions is 99.3% (see Table S8 in the Supporting Information) even though the overlap between the two normalized orthogonal complements (Φ_X) is a little lower (86.1%, see Table S8 in the Supporting Information). All of this apparent numerical similarity between the SCGVB(a) and SCGVB(b) solutions (except for their different energies) reinforces our suspicion that one can easily be misled about the degree of (say) SS or NN singlet diradical character when relying mostly on the visual inspection of SCGVB active orbitals that are not sufficiently well localized. It does now appear that Gerratt et al. [1] were misdirected in this way when (mis)interpreting their SCGVB(a) wavefunction in terms of dominant SS singlet diradical character. (As before, all of our key observations are much the same when using Gallup-Norbeck weights and/or if we project instead onto S-BOVB structures.)

Table 3. Energies (combinations of) froze	and Chirgv en-core SCGVI			
Wavefunction	Energy		Weights	
vvaverunction	(hartree)	а	b	С
SCGVB(a)	-903.98083	100%	_	1
SCGVB(b)	-903.97206	_	100%	_
SCGVB(C)	-903.99314	_	_	100%
VBCI(a ⊕ b ⊕ C)	-903.99326	-2.1%	7.9%	94.2%
$GMCSC(a \oplus b \oplus C)$	-904.00270	25.9%	19.0%	55.1%

When restricting SCGVB active orbitals to span particular active spaces, Thorsteinsson et al.^[7] found in some cases, such as c=[2,1,2,1], that the resulting SCGVB solution was symmetry broken. The proper full symmetry could be restored by using a twoconfiguration description in which the two sets of active orbitals were related by a particular D_{2h} symmetry operation, such as a reflection or a rotation. With this in mind, we chose here to carry out two-configuration SCGVB calculations using the GMCSC program. Even without specifying any symmetry constraints between the two orbital strings, we observed convergence to a symmetry-pure solution in which the two sets of active orbitals are related to one another by reflection in the σ_{vz} mirror plane. Additional symmetry relations emerged spontaneously within the two orbital strings, so that π_3 , π_4 and π_6 can be generated from π_1 , π_2 and π_5 , respectively, by a $\hat{C}_2(z)$ rotation. (Specific details of these orbitals are slightly different from some of those envisaged by Thorsteinsson et al.^[7] for their 'projected' c solution, prompting us to use a slightly different label.) The resulting energy for our variationally-optimized solution, which we label SCGVB($C_1 \oplus C_2$), or SCGVB(C) for short, is somewhat better than that of SCGVB(a) (see

Table 3), but the two sets of Chirgwin-Coulson weights (see Table 2) are fairly similar. Furthermore, the overlap between the total SCGVB(a) and SCGVB(C) wavefunctions is 99.4% (see Table S8 in the Supporting Information), with the overlap between the two normalized orthogonal complements (Φ_X) being a little lower (88.2%, see Table S8 in the Supporting Information). The symmetry-unique active orbitals π_1 , π_2 and π_5 for the SCGVB(C) solution, shown in the bottom row of Fig. 3, clearly all exhibit a significant degree of (at least) two-center character, rendering it somewhat difficult to interpret them directly and unambiguously in terms of relative contributions from the sorts of VB structures shown in Fig. 1. Relying instead on the projection onto BOVB structures, we can say that the largest net contributor (57.3%) is the symmetry-determined linear combination (Φ_{3-6}) of the four symmetry-equivalent BOVB structures **3-6**.

Just as there are CASSCF(6,6) solutions based on active spaces $\mathbf{e}=[3,1,1,1]$ and $\mathbf{f}=[2,2,2,0]$ which lie lower than that for $\mathbf{b}=[3,2,1,0]$ by ca. 3.0 and 1.6 millihartree, respectively (see Table S1 in the Supporting Information), it also proved possible to locate another single-configuration SCGVB solution which lies lower than SCGVB(\mathbf{b}) by ca. 2.3 millihartree. However, given the dominance of SCGVB(\mathbf{a}) and SCGVB(\mathbf{C}) we decided not to pursue this solution, or any of the higher lying ones, in any detail.

The close proximity in energy of different SCGVB descriptions, especially a and C, prompted us to wonder which of them would dominate variationally-optimized combinations of the individual solutions. Accordingly, we also considered a VBCI($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) description, in which we combined the SCGVB(a), SCGVB(b) and SCGVB(C) wavefunctions via a nonorthogonal CI calculation, without relaxing any of the active-space spin-coupling coefficients. As can be seen from Table 3, this multicomponent wavefunction gives only a very modest energy improvement over SCGVB(C), with the C component remaining overwhelmingly dominant (Chirgwin-Coulson weight 94.2%). The corresponding Gallup-Norbeck weights (See Table S7 in the Supporting Information) show the **C** component to be even more dominant. Projection of VBCI($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) onto the basis of BOVB structures gives weights that are very similar to those for SCGVB(C). Analogous outcomes are observed for the overlap between the VBCI($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) and SCGVB(C) wavefunctions, as well as for the overlap between the two orthogonal components, Φ_X (see Tables S8 and S9 in the Supporting Information).



Relaxing the various spin-coupling coefficients (without reoptimizing the active orbitals) produced only a very small energy lowering. All in all, other than making the resulting description more difficult to interpret directly, rather little is achieved by this mixing of the SCGVB(a), SCGVB(b) and SCGVB(C) wavefunctions.

The outcome is somewhat better, at least in terms of the total energy, if all of the active orbitals and activespace spin-coupling coefficients are simultaneously reoptimized (albeit with a limited number of suitable constraints so as to retain some distinction between the a, b and C components). We use the label $GMCSC(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$ for the resulting description. We observe from Table 3 that the reoptimized C component remains the largest contributor, but nearly 45% of the total is now due to the reoptimized **a** and **b** components. These changes to the weights of the different components of GMCSC($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) relative to those of VBCI($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$), as well as the lowering of the total energy, are accompanied by relatively small changes to the forms of the various symmetry-unique SCGVB active orbitals (see Fig. S1 in the Supporting Information). Variationally, this may well be the best π space wavefunction that has been applied so far to S₂N₂ but, due to the approach that we have adopted, it can still be interpreted in chemically-relevant terms.

Projection of GMCSC($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) onto the basis of BOVB structures produces weights that are not so different from those for our various other SCGVB descriptions (see Table 2). As before, BOVB structure 1 (corresponding to NN singlet diradical character) is found to be more important than BOVB structure 2 (SS singlet diradical character), but the largest contribution (58.5%) comes from the symmetry-determined linear combination of the four symmetry-equivalent BOVB structures 3-6, with no obvious diradical character. A further 4.9% is due to the normalized orthogonal complement (Φ_X) . (We find again that all of our key observations are much the same when using Gallup-Norbeck weights instead of those from the Chirgwin-Coulson scheme and/or if we project instead onto S-BOVB structures.)

There are substantial differences between existing estimates of the degree of diradical character in the singlet ground state of S_2N_2 , but it is important to note in this context, as was emphasized by Braïda et al., that significant singlet diradical character can co-exist with aromaticity in this molecule. At one extreme, Jung et al. argued that S_2N_2 should be regarded as a 2π -

electron aromatic system, without any significant diradical character. Tuononen et al. [4],[6] used a simple scheme that is based on the ratio of two CI coefficients in their CASSCF(22,16) description to estimate just 6% diradical character for this molecule. On the other hand, for the same CASSCF(22,16) wavefunction, they also considered a different form of analysis, based on idealized p_{π} orbitals, which assigned a weight of 34% for 1 and of 14% for 2. [4],[6] The BOVB calculations of Braïda et al.,[2] as well as our own BOVB and S-BOVB calculations, also suggest significant weights for these two diradical structures, with 1 being somewhat more important than 2, just as was found by Harcourt [3] when using a somewhat different VB approach.

Whereas Gerratt et al. [1] (mis)interpreted their S₂N₂ wavefunction in terms of dominant SS singlet diradical character, our projection of the SCGVB(a) description onto BOVB structures reveals a higher weight for 1 than for 2. Indeed, all of our projections of SCGVB-like wavefunctions for S2N2 onto the basis of BOVB or S-BOVB structures (and an orthogonal complement) show 1 to be more important than 2, but they also indicate that nearly 60% of the wavefunction is instead associated with the symmetry-determined linear combination of the four symmetry-equivalent BOVB structures 3-6. This relatively high combined weight for structures 3-6 is of course consistent with the wellestablished pattern for the atomic charges in which nitrogen is negative and sulfur is positive. [5] We find that the net QTAIM charges for the SCGVB(a) total electron density are numerically much the same as those just for the valence π space. This suggests that inferences about the bonding in the valence π system from considerations of the overall charge distribution, such as those discussed by Jung et al., [5] are not skewed by any significant charge separation in the σ -bonded framework. (Our QTAIM charges for SCGVB(a) do of course have the correct NS polarity. We also find that the QTAIM charges for the GMCSC($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) total electron density are very similar to those for SCGVB(a).)

$S_4N_4^{2+}$

We find that free optimization of a single-configuration SCGVB wavefunction for $S_4N_4^{2+}$ results in a solution that corresponds to one of the D_{2h} subgroups of the full molecular point group (D_{4h}) . Amongst many other deviations from full D_{4h} symmetry, such that the desired $^1A_{1g}$ wavefunction has a small but not negligible $^1B_{2g}$ contaminant, we observed that active orbitals π_1 and π_2 were each relatively close to being invariant under



reflection in the σ_{yz} mirror plane, but they were not exactly so. The various findings described above for S_2N_2 are suggestive that a suitable way forward would be to optimize a two-configuration SCGVB description in which the two sets of active orbitals are related to one another by an appropriate reflection or rotation. We label the resulting wavefunction as SCGVB($A_1 \oplus A_2$), or SCGVB(A_1) for short. Nonetheless we also consider a single-configuration SCGVB description in which the active orbitals are suitably constrained so as to ensure that the resulting wavefunction, which we denote SCGVB(A_1), still respects the full A_2 symmetry.

We found at convergence of our symmetry-pure SCGVB(A) solution that the two sets of active orbitals (i.e. those for A_1 and A_2) are related to one another by reflection in the σ_{xz} mirror plane. Additionally, we observed symmetry relations within each orbital string, such that reflection of π_1 and π_2 in the plane x=yyields π_3 and π_4 , respectively, and $\hat{\mathcal{C}}_2(z)$ rotation of π_1 and π_2 gives π_5 and π_6 , respectively. Similarly, reflection of π_1 , π_2 and π_9 in the plane x=-y yields π_7 , π_8 and π_{10} , respectively. For the optimization of the SCGVB(**B**) solution, we constrained active orbitals π_1 and π_2 to be exactly $\hat{\sigma}_{vz}$ invariant and it also proved necessary to add as constraints symmetry relations for π_3 - π_8 which emerged spontaneously for SCGVB(A). Additionally, after some experimentation in which we sought the lowest possible energy, we found that we had to constrain π_9 for SCGVB(**B**) to be invariant under reflection in the plane x = y as well as under $\hat{C}_2(z)$ rotation, with π_{10} generated from π_9 by a $\hat{C}_4(z)$ rotation.

The resulting symmetry-unique active orbitals π_1 , π_2 and π_9 from the SCGVB(**A**) and SCGVB(**B**) calculations are displayed in the top and bottom rows, respectively, of Fig. 4. Except for being more localized, the SCGVB(**A**) active orbitals are fairly reminiscent of those for the S_2N_2 SCGVB(**C**) solution (bottom row of Fig. 3). On the other hand, the symmetry-unique SCGVB(**B**) active orbitals include a three-center SNS function, just as was the case for the S_2N_2 SCGVB(**a**) solution (top row of Fig. 3). As can be seen from Table 4, solution SCGVB(**A**) lies somewhat lower than SCGVB(**B**), as we should have expected. Computing the energy of the A_1 (or A_2) component using the same active orbitals and spincoupling coefficients as in the SCGVB(**A**) solution, we

find that the energy lowering associated with mixing together these A_1 and A_2 components is $160\,\text{kJ/mol}$ (38 kcal/mol). The corresponding energy change from C_1 to SCGVB(C) in the case of S_2N_2 is $124\,\text{kJ/mol}$ (30 kcal/mol). In both cases, the optimization of the active orbitals allows the two configurations to become more different from one another, at the expense of the energy of each of them separately, with the consequence that the mixing of the two components corresponds to a significant energy lowering. (Additional data are available in Table S10 in the Supporting Information.)

Table 4. Energies and (combinations of) frozen $S_4N_4^{2+}$.	-		
Wavefunction	Energy	Wei	ghts
Waveranotion	(hartree)	Α	В
SCGVB(A)	-1807.29841	100%	_
SCGVB(B)	-1807.24746	_	100%
VBCI(A ⊕ B)	-1807.29856	95.1%	4.9%
GMCSC(A ⊕ B)	-1807.30558	73.0%	27.0%

In keeping with our experience for S₂N₂, combining the SCGVB(**A**) and SCGVB(**B**) descriptions of $S_4N_4^{2+}$ via a nonorthogonal CI calculation, thereby generating the VBCI(**A** ⊕ **B**) wavefunction (without relaxing any of the active-space spin-coupling coefficients), results in relatively little energy improvement, with SCGVB(A) remaining dominant (see Table 4). Just as was the case for the VBCI($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) description of S_2N_2 , relaxation of the various spin-coupling coefficients, without reoptimizing the active orbitals, produced only a very small energy lowering. Larger changes are achieved for $GMCSC(A \oplus B)$, in which all of the active orbitals and active-space spin-coupling coefficients simultaneously reoptimized (subject to a limited number of suitable constraints so as to retain some distinction between the A and B components). The reoptimized A component remains the largest contributor and there are only relatively small changes to the forms of the various symmetry-unique SCGVB active orbitals (see Fig. S2 in the Supporting Information). Amongst other changes from $VBCI(A \oplus B)$ to GMCSC($\mathbf{A} \oplus \mathbf{B}$), the $\langle \mathbf{A_1} | \mathbf{A_2} \rangle$ overlap is reduced from 0.675 to 0.571 whereas $\langle \mathbf{A} | \mathbf{B} \rangle$ goes down from 0.951 to 0.847 (see Table S11 in the Supporting Information).



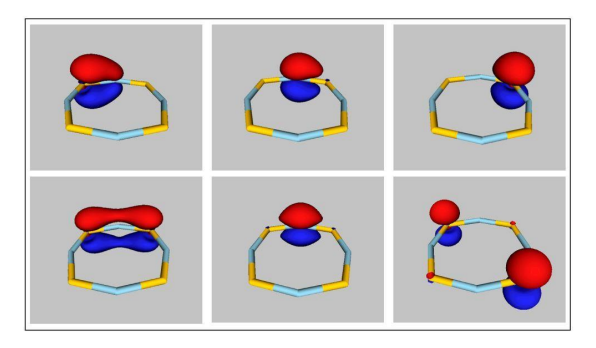


Figure 4. Symmetry-unique active orbitals π_1 , π_2 and π_9 (left to right) for frozen-core S₄N₄²⁺ wavefunctions: SCGVB(**A**) (top row); SCGVB(**B**) (bottom row).

Summary and conclusions

We have carried out a detailed comparative analysis of a number of alternative VB descriptions of the π -electron system in S₂N₂. As has been shown before,^[7] S₂N₂ at its idealized square geometry, close to experiment, offers multiple energetically close π -space '6 electrons in 6 orbitals' CASSCF solutions and thus also various competing SCGVB descriptions. Although the active space $2B_{1u}+1B_{2q}+2B_{3q}+1A_u$, or c=[2,1,2,1] for short, gives the lowest CASSCF(6,6) energy for the calculations carried out here, the corresponding a=[3,1,2,0] solution lies less than 0.35 millihartree higher. (The ordering of these two solutions was reversed in the calculations of Thorsteinsson et al., [7] who used a somewhat smaller basis set) We find that the energetically preferred single-configuration SCGVB wavefunction is related to active space a. This SCGVB(a) solution, which corresponds to the one described by Gerratt at al., [1] appears at first sight of the orbitals and active-space total spin function to correspond to dominant SS singlet diradical character. Somewhat higher in energy is a single-configuration SCGVB(b) solution, which appears at first sight to be dominated by NN singlet diradical character. As was pointed out by Thorsteinsson et al., [7] the occurrence of negative overlaps for the SS and NN singlet diradical orbital pairs, with nodal planes between the two participating

centers, means that there can be no question of any cross-ring bonding in either of the SCGVB(a) or SCGVB(b) descriptions. Given the occurrence of threecenter orbitals in both descriptions, as well as delocalization tails, it is of course easy to be misdirected by an examination of the shapes and locations of such active orbitals. In reality, in spite of featuring active orbitals that are visually somewhat different, these two SCGVB solutions turn out to be rather similar to one another (except in terms of their total energies).

The corresponding single-configuration **SCGVB** wavefunction for S_2N_2 active space c is symmetry broken and so we optimized instead a twoconfiguration SCGVB wavefunction, which we label SCGVB(C). (Specific details of the two orbital strings are slightly different from some of those envisaged by Thorsteinsson et al.^[7] for their 'projected' **c** solution, prompting us to use a slightly different label.) Although our variationally-optimized SCGVB(C) solution turns out to be energetically preferred over SCGVB(a) by more than 12 millihartree, it is still reasonable to wonder whether C would dominate a variationally-optimized combination. A nonorthogonal CI with fixed active orbitals and fixed spin-coupling coefficients, VBCI(a \oplus b \oplus C), produced rather little energy improvement over SCGVB(C), with C being by far the dominant component. Subsequent relaxation of the spin-coupling coefficients achieved relatively little for



the energy whereas simultaneous reoptimization also of the active orbitals, in the GMCSC($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) description, yielded a modest energy improvement of \mathbf{ca} . 9.5 millihartree. Nonetheless, although the reoptimized \mathbf{a} and \mathbf{b} components collectively account for nearly 45% of the total GMCSC($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) wavefunction, the largest contribution is due to the reoptimized \mathbf{C} component.

Subsequent calculations for the 10-electron π space of the D_{4h} S₄N₄²⁺ ring produced a symmetry-broken singleconfiguration SCGVB solution, unless suitable constraints were placed on the orbitals, as was done for SCGVB(B) description. Our optimal twoconfiguration solution, which we label SCGVB(A), turns out to be energetically preferred over SCGVB(B) by nearly 51 millihartree. Whereas the symmetry-unique SCGVB(B) active orbitals are found to include a threecenter SNS function, just as was the case for the S₂N₂ SCGVB(a) solution, the SCGVB(A) active orbitals are more reminiscent of those for the S_2N_2 SCGVB(C) solution, except for being more localized. We found that the mixing of SCGVB(A) and SCGVB(B) generates hardly any energy improvement unless the active orbitals and the spin-coupling are simultaneously reoptimized. The resulting GMCSC($\mathbf{A} \oplus \mathbf{B}$) solutions lies lower than SCGVB(A) by ca. 7 millihartree, with the reoptimized A component remaining the largest contributor.

For both of the S_2N_2 and $S_4N_4^{2+}$ rings at their idealized geometries, close to experiment, our preferred compact SCGVB description of the π -electron system is provided by the variational optimization of two configurations which turn out to be symmetry related to one another. The optimization instead of symmetry-pure single-configuration SCGVB wavefunctions leads to the involvement of three-center SNS or NSN orbitals, which now seems to be an unnecessary complication. In neither ring system is very much achieved from the mixing of such competing solutions.

As is to be expected, our BOVB and S-BOVB calculations for S_2N_2 confirm that structure ${\bf 1}$ is more important than structure ${\bf 2}$, $^{[2],[3],[4],[6]}$ but the largest contribution (ca. 60%) turns out to be due to the symmetry-determined linear combination of the four symmetry-equivalent structures ${\bf 3-6}$. Much the same pattern was consistently found when we used a simple but robust projection of our various SCGVB, VBCI and GMCSC wavefunctions for S_2N_2 , including the very general GMCSC(${\bf a} \oplus {\bf b} \oplus {\bf C}$) construction, onto the basis of BOVB or S-BOVB

structures (plus an orthogonal complement). In particular, it does indeed now appear that Gerratt et al.^[1] were misdirected by active orbitals that are not sufficiently localized when (mis)interpreting their SCGVB(a) wavefunction in terms of dominant SS singlet diradical character.

These results reconcile one of the long-standing arguments in the applications of modern VB theory and help to establish a consistent VB description of the electronic structure of S_2N_2 . This description incorporates significant levels of resonance, dominated by the combined contribution of the four symmetry-equivalent structures **3-6**, each of which features a sulfur-nitrogen double bond but lacks any obvious diradical character, followed by the structure with NN singlet diradical character, **1**, and with the structure with SS singlet diradical character, **2**, coming only last.

Keywords: S_2N_2 and $S_4N_4^{2+}$; spin-coupled generalized valence bond (SCGVB); π -electron rings; breathing orbital valence bond; generalized multiconfiguration spin-coupled.

Additional Supporting Information may be found in the online version of this article.

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Supporting Information

Is the S₂N₂ Ring a Singlet Diradical? Critical Analysis of Alternative Valence Bond Descriptions

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Table S1. Tests of different sets of CASSCF inactive orbitals for S₂N₂.

(a) Spaces spanned

	B _{1u}	B_{2g}	B_{3g}	Au	
π full-valence (fv)	2	1	1	0	[2,1,1,0]
a	3	1	2	0	[3,1,2,0]
b	3	2	1	0	[3,2,1,0]
С	2	1	2	1	[2,1,2,1]
d	2	2	1	1	[2,2,1,1]
е	3	1	1	1	[3,1,1,1]
f	2	2	2	0	[2,2,2,0]
max(a,b,c,d,e,f)	3	2	2	1	[3,2,2,1]

(b) CASSCF energies (in hartree) and selected differences (in millihartree) for different choices of inactive orbitals. (E_{RHF} = -903.91753 hartree)

Inactive orbitals			Energy				
mactive orbitals	fv =[2,1,1,0]	a =[3,1,2,0]	b =[3,2,1,0]	c =[2,1,2,1]	max=[3,2,2,1]	а – с	b – c
Variational	-903.96018	-903.98159	-903.97320	-903.98194	-903.99550	0.35	8.73
fv =[2,1,1,0]	-903.96018	-903.98140	-903.97315	-903.98190	-903.99536	0.51	8.76
a =[3,1,2,0]	-903.95998	-903.98159	-903.97314	-903.98183	-903.99548	0.24	8.69
b =[3,2,1,0]	-903.96013	-903.98152	-903.97320	-903.98192	-903.99547	0.40	8.71
c =[2,1,2,1]	-903.96015	-903.98148	-903.97319	-903.98194	-903.99545	0.46	8.75
<i>max</i> =[3,2,2,1]	-903.96005	-903.98157	-903.97318	-903.98189	-903.99550	0.31	8.71

(c) Energies (in hartree) for additional CASSCF descriptions.

Inactive orbitals	Energy d =[2,2,1,1] e =[3,1,1,1] f =[2,2,2,0]			
mactive orbitals	d =[2,2,1,1]	e =[3,1,1,1]	f =[2,2,2,0]	
variational	-903.97044	-903.97614	-903.97487	
<i>max</i> =[3,2,2,1]	-903.97032	-903.97613	-903.97483	



Figure S1. Symmetry-unique active orbitals in the frozen-core GMCSC($\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C}$) wavefunction for S_2N_2 . Orbitals shown in the same order as in Fig. 3.

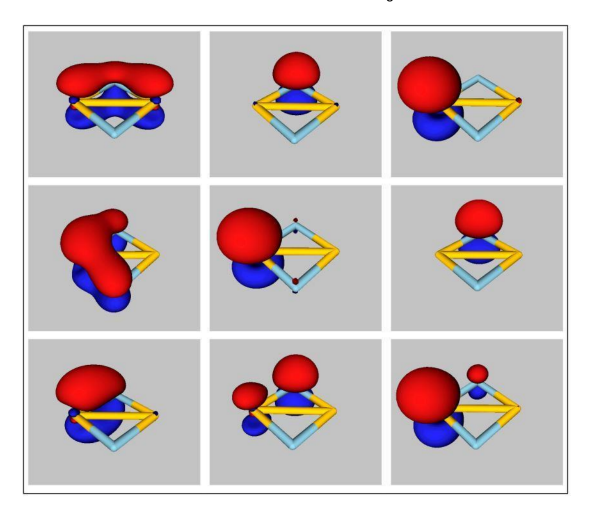


Table S2. Overlap integrals for frozen-core six-structure BOVB wavefunctions for S_2N_2 (where **X** signifies a normalized orthogonal complement).

1	2	3-6
1.000	0.130	0.472
0.130	1.000	0.475
0.472	0.475	1.000
	0.130	1 2 1.000 0.130 0.130 1.000 0.472 0.475

	1	2	3-6	1-6	X
SCGVB(a)	0.689	0.575	0.906	0.973	0.225
SCGVB(b)	0.703	0.568	0.899	0.973	0.227
SCGVB(C)	0.686	0.577	0.909	0.975	0.217
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.688	0.577	0.909	0.975	0.215
$GMCSC(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.688	0.564	0.912	0.974	0.222



Table S3. Inverse-overlap (Gallup-Norbeck) weights, with **1** to **6** taken directly from the frozencore six-structure BOVB calculations for S_2N_2 and where **X** signifies a normalized orthogonal complement with weight d_X^2 .

Wavefunction	1	2	3-6	Х
BOVB (1-6)	31.8%	7.4%	60.8%	-
SCGVB(a)	25.5%	10.2%	59.3%	5.0%
SCGVB(b)	28.7%	10.1%	56.0%	5.2%
SCGVB(C)	24.8%	10.2%	60.2%	4.7%
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	25.1%	10.3%	60.0%	4.6%
$GMCSC(\mathbf{a}\oplus\mathbf{b}\oplus\mathbf{C})$	24.5%	8.6%	62.0%	4.9%

Table S4. Energies (in hartree) from frozen-core six-structure S-BOVB calculations for S_2N_2 , where 'Fixed' signifies the use of active orbitals taken directly from the six-structure BOVB calculation and 'Relaxed' signifies further optimization.

Structures	Fixed	Relaxed
1	-903.79978	-903.85801
2	-903.66658	-903.71685
3	-903.71852	-903.73523
3-6	-903.89354	-903.90623
1-6	-903.96054	-903.96054

Table S5. Overlap integrals for frozen-core six-structure S-BOVB wavefunctions for S₂N₂ (where **X** signifies a normalized orthogonal complement).

	1	2	3-6
1	1.000	0.136	0.471
2	0.136	1.000	0.469
3-6	0.471	0.469	1.000

	1	2	3-6	1-6	X
SCGVB(a)	0.685	0.575	0.905	0.972	0.232
SCGVB(b)	0.701	0.566	0.895	0.970	0.240
SCGVB(C)	0.683	0.577	0.910	0.975	0.218
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.685	0.577	0.909	0.975	0.217
$GMCSC(\mathbf{a}\oplus\mathbf{b}\oplus\mathbf{C})$	0.689	0.563	0.914	0.977	0.210



Table S6. Chirgwin-Coulson and inverse-overlap weights, with **1** to **6** taken directly from the frozen-core six-structure S-BOVB calculations for S_2N_2 and where **X** signifies a normalized orthogonal complement with weight d_X^2 .

(a) Chirgwin-Coulson weights

Wavefunction	1	2	3-6	Χ
S-BOVB (1-6)	28.9%	11.3%	59.8%	_
SCGVB(a)	24.5%	13.4%	56.8%	5.4%
SCGVB(b)	26.9%	13.0%	54.4%	5.7%
SCGVB(C)	24.1%	13.3%	57.8%	4.8%
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	24.4%	13.3%	57.6%	4.7%
$GMCSC(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	24.5%	11.9%	59.2%	4.4%

(b) Inverse-overlap (Gallup-Norbeck) weights

Wavefunction	1	2	3-6	Х
S-BOVB (1-6)	29.7%	7.8%	62.5%	-
SCGVB(a)	24.5%	10.3%	59.8%	5.4%
SCGVB(b)	28.2%	10.0%	56.1%	5.7%
SCGVB(C)	23.8%	10.2%	61.3%	4.8%
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	24.1%	10.2%	61.0%	4.7%
$GMCSC(\mathbf{a}\oplus\mathbf{b}\oplus\mathbf{C})$	23.9%	8.5%	63.2%	4.4%

Table S7. Overlap integrals and Gallup-Norbeck weights for combinations of frozen-core SCGVB wavefunctions for S_2N_2 .

Wavefunction		Ove	rlaps	Weights			
wavefunction	⟨a b⟩	⟨a C⟩	$\langle \mathbf{b} \mathbf{C} \rangle$	$\langle C_1 C_2\rangle$	а	b	С
VBCI(a ⊕ b ⊕ C)	0.993	0.994	0.989	0.810	0.0%	0.9%	99.1%
$GMCSC(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.861	0.911	0.886	0.744	19.0%	13.6%	67.3%



Table S8. Overlaps between various (combinations of) frozen-core SCGVB wavefunctions for S_2N_2 .

-	SCGVB(a)	SCGVB(b)	SCGVB(C)	$VBCI(\mathbf{a}\oplus\mathbf{b}\oplus\mathbf{C})$	$\overline{GMCSC(a\oplusb\oplusC)}$
SCGVB(a)	1	0.99271	0.99422	0.99467	0.99205
SCGVB(b)	0.99271	1	0.98897	0.99047	0.98797
SCGVB(C)	0.99422	0.98897	1	0.99994	0.99670
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.99467	0.99047	0.99994	1	0.99680
$GMCSC(\mathbf{a}\oplus\mathbf{b}\oplus\mathbf{C})$	0.99205	0.98797	0.99670	0.99680	1

Table S9. Overlaps between the normalized orthogonal complements arising from projections of various (combinations of) frozen-core SCGVB wavefunctions for S_2N_2 .

(a) BOVB

_	SCGVB(a)	SCGVB(b)	SCGVB(C)	$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	$\overline{GMCSC(a\oplusb\oplusC)}$
SCGVB(a)	1	0.86145	0.88248	0.89080	0.84333
SCGVB(b)	0.86145	1	0.78478	0.81253	0.77044
SCGVB(C)	0.88248	0.78478	1	0.99890	0.93459
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.89080	0.81253	0.99890	1	0.93626
$GMCSC(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.84333	0.77044	0.93459	0.93626	1

(b) S-BOVB

	SCGVB(a)	SCGVB(b)	SCGVB(C)	$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	$GMCSC(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$
SCGVB(a)	1	0.87484	0.88768	0.89617	0.84577
SCGVB(b)	0.87484	1	0.80272	0.82952	0.78086
SCGVB(C)	0.88768	0.80272	1	0.99888	0.93255
$VBCI(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.89617	0.82952	0.99888	1	0.93395
$GMCSC(\mathbf{a} \oplus \mathbf{b} \oplus \mathbf{C})$	0.84577	0.78086	0.93255	0.93395	1



Figure S2. Symmetry-unique active orbitals in the frozen-core GMCSC($\mathbf{A} \oplus \mathbf{B}$) wavefunction for $S_4N_4^{2+}$. Orbitals shown in the same order as in Fig. 4.

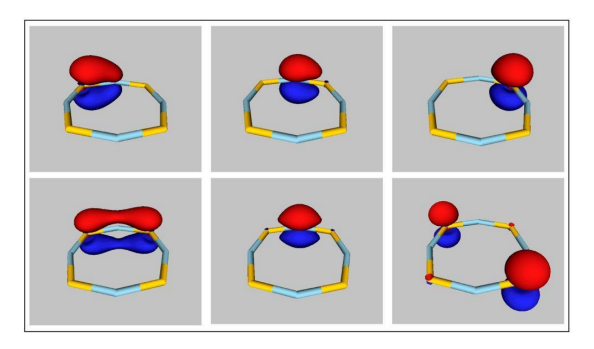


Table S10. Component energies (in hartree) and inter-component overlaps for two-configuration wavefunctions.

System	Source of orbitals and spin-coupling coefficients	Energy of single component	Inter-component overlap
S_2N_2	SCGVB(C)	-903.94607	0.810
$S_4N_4^{2+}$	SCGVB(A)	-1807.23764	0.675

Table S11. Overlap integrals and Gallup-Norbeck weights for combinations of frozen-core SCGVB wavefunctions for $S_4N_4^{2+}$.

Wavefunction		Overlap	Weights		
vvaverunction	$\langle A B\rangle$	$\langle A_1 A_2\rangle$	$\langle A_1 B\rangle$	Α	В
VBCI(A ⊕ B)	0.951	0.675	0.870	99.7%	0.3%
$GMCSC(\mathbf{A} \oplus \mathbf{B})$	0.847	0.571	0.751	86.4%	13.6%