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State-averaged Monte Carlo configuration interaction applied to electronically excited states

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We introduce state-averaging into the method of Monte Carlo configuration interaction (SA-MCCI) to allow the stable and efficient calculation of excited states. We show that excited potential curves for H₂, including a crossing with the ground state, can be accurately reproduced using a small fraction of the full configuration interaction (FCI) space. A recently introduced error measure for potential curves [J. P. Coe and M. J. Paterson, J. Chem. Phys. 137, 204108 (2012)] is also shown to be a fair approach when considering potential curves for multiple states. We demonstrate that potential curves for LiF using SA-MCCI agree well with the FCI results and the avoided crossing occurs correctly. The seam of conical intersections for CH₂ found by Yarkony [J. Chem. Phys. 104, 2932 (1996)] is used as a test for SA-MCCI and we compare potential curves from SA-MCCI with FCI results for this system for the first three triplet states. We then demonstrate the improvement from using SA-MCCI on the dipole of the 2 ¹A₁ state of carbon monoxide. We then look at vertical excitations for small organic molecules up to the size of butadiene where the SA-MCCI energies and oscillator strengths are compared with CASPT2 values [M. Schreiber, M. R. Silva-Junior, S. P. A. Sauer, and W. Thiel, J. Chem. Phys. 128, 134110 (2008)]. We finally see if the SA-MCCI results for these excitation energies can be improved by using MCCIPT2 with approximate natural orbitals when the PT2 space is not onerously large. © 2013 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4824888]

I. INTRODUCTION

Monte Carlo configuration interaction (MCCI)⁴⁻² randomly augments a configuration space in an iterative scheme where those configurations whose absolute coefficient in the resulting solution of the time-independent Schrödinger equation in this space is less than a certain amount will eventually be deleted. This procedure, without prior knowledge of the important configurations or orbitals, has been demonstrated to produce a compact wavefunction which recreates much of the full configuration interaction (FCI) solution for systems comprising a few atoms. MCCI has recently been used for the calculation of vertical excitations of atoms and small molecules,⁴ potential curves of ground states,⁴ electron affinities, ionization energies, and multipole moments.⁵

The main goal of this paper is to investigate whether MCCI with a type of state-averaging (SA-MCCI) can be applied to the calculation of excited states particularly potential curves—including conical intersections and avoided crossings—using a small fraction of the FCI configurations and without prior knowledge of the important orbitals.

MCCI represents another approach for the calculation of excited states which may be expected to be particularly suitable for potential curves of small systems. For systems where there is a clearly dominant configuration in the ground-state then single-reference methods based on coupled-cluster such as EOM-CCSD (see, e.g., Ref. 6) may be used to efficiently find excited states, which may be multireference, as long as their main configurations have a substitution level from the ground-state appropriate to the method used. However along a potential curve, away from the equilibrium geometry, the ground-state wavefunction may have a number of important configurations and here multireference methods such as complete active space self-consistent field (CASSCF),⁷ can be used to capture much of the static correlation perhaps followed by second-order perturbation CASPT²⁸ to recover more of the dynamic correlation. However to appropriately choose the orbitals for inclusion in the active space requires insight and the calculation can become intractable if the active space is too large. Furthermore, the important orbitals may change along the potential curve. MCCI offers an alternative approach as it depends only on the cut-off value for configurations to be included so in principle, with sufficiently small cut-off and sufficiently long calculation time, it can find compact wavefunctions to describe excited states accurately at all points along a potential curve. Other recent stochastic approaches to excited states include those using projector or diffusion Monte Carlo in Slater determinant space to approach the FCI energy and wavefunction. For example, Ohtsuka and Nagase⁹ calculate an excited state by using the projector Monte Carlo results from all the states below to eliminate their component from the initial wavefunction. Booth and Chan¹⁰ use (Ĥ − S)² in the projection operator where S is chosen to be very close to the energy of the state of interest. Another interesting approach is model space quantum Monte Carlo (MSQMC)¹¹ which partitions configuration space and uses projector Monte Carlo to construct an effective Hamiltonian in the smaller space where Slater determinants

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can move from one space to the other if their importance passes a threshold. We note that MCCI is not used in this work to find the FCI energy, but rather a very compact wavefunction which captures much of the FCI result yet can be used for further calculations due to its tractable size.

The calculation of excited state potential curves in MCCI is more challenging than those of the ground state as during an MCCI calculation of, e.g., the first excited state of a given symmetry the method may remove configurations unimportant for the excited state to produce a set of configuration state functions (CSFs) that cannot describe the ground state. In this case the ground state within this set may become similar to the first excited state and the first excited state may be an approximation to a higher excited state. In the best case this will cause oscillations in the property of interest (e.g., the excited state dipole moment of CO in Ref. 5), but it can cause the diagonalization procedure to fail. We introduce a type of state-averaging to MCCI (SA-MCCI) to efficiently overcome this problem. We demonstrate that excited potential curves with conical intersections for H$_3$ and CH$_2$, and avoided crossings for LiF can be calculated using SA-MCCI. We then show that this method can prevent the oscillations seen in the MCCI dipole results for the $2^1A_1$ state of carbon monoxide.\(^5\) We finally look at the vertical excitations of organic molecules up to the size of butadiene using SA-MCCI and also consider if the approach of using approximate natural orbitals to construct small SA-MCCI wavefunctions followed by MCCIPT2\(^12\) can more accurately describe these excitation energies.

II. METHOD

We first present a brief recap of the standard MCCI algorithm\(^1,2\) which usually begins with the CSF constructed from occupied Hartree-Fock (HF) molecular orbitals:

1. A set of CSFs is randomly enlarged using symmetry preserving single and double substitutions.
2. The Hamiltonian matrix and overlap matrix are constructed and a diagonalization gives the wavefunction coefficients.
3. CSFs that have just been added are deleted if the absolute value of their coefficient is less than $c_{\text{min}}$.
4. The process continues and after every ten iterations all CSFs are considered for deletion (full pruning).

The integrals of the molecular orbitals are calculated using Molpro\(^13\) for the potential curves in this work, while for oscillator strength calculations of the vertical excitations we use Columbus\(^14\) due to the ease availability of the dipole integrals. We freeze core orbitals for the calculations of the integrals rather than in the MCCI calculation. This means that fewer calculations are needed in MCCI when constructing matrix elements and fewer integrals need to be stored.

We use a convergence check when calculating potential curves using MCCI as described in Ref. 12. Here the set of CSFs is not enlarged on the step following a full prune. The maximum difference in the energies after the last three full pruning steps is required to be less than $10^{-3}$ hartree for the calculation to cease. We employ sixty warmup iterations before the convergence check is implemented and run MCCI on twelve processors.

One approach when working with excited states of a certain symmetry would be to calculate the ground state then ensure that these configurations are always included in an excited-state calculation. For the vertical excitations of Ref. 3, important configurations from the lower states were permanently included\(^15\) to prevent the collapse of the excited state MCCI calculation. However, for a calculation of the first excited state with a fixed number of iterations this will require twice as many diagonalizations and the size of the excited-state Hamiltonian matrix will always be at least that of the ground state. This problem will clearly become worse as the number of states of interest increases. We, therefore, investigate a type of state averaging: in addition to the coefficients for the excited state $s$ of interest $c_{i,s}$ the coefficients $c_{i,1}$ of the ground state and all other states below $s$ in this symmetry and spin are calculated at each step and when coefficients are considered for removal they are constructed as

$$c_i = \sum_{j=1}^{s} |c_{i,j}|. \quad (1)$$

This aims to ensure that the set of CSFs found during the MCCI run offers a good description of both the ground and excited states. The size of the wavefunction will be slightly larger as some states that would be removed in a single-state calculation may have a large enough state-averaged value to be included. However, we can start from the HF reference so the states are built up together and the size of the MCCI wavefunction in the early iterations at least is smaller than when using a fixed ground state. Furthermore, we reduce the number of diagonalizations by using the Davidson-Liu algorithm\(^16\) to calculate the $s$ lowest eigenvalues and eigenvectors simultaneously. The initial guess of expansion vectors $b_k$ for the smaller Hamiltonian matrix that is solved in the Davidson algorithm appears to be problematic for excited states but rarely for the ground states as the HF configuration will tend to still have a large coefficient after the first addition of configurations in the latter case. Hence, to improve stability of the method, on the first iteration the ground state only is calculated, but new $b_k$ vectors from the ground and excited states of the reduced Hamiltonian matrix are created for use in the next iteration.

III. RESULTS

A. H$_3$ equilateral triangle

We test SA-MCCI on three hydrogens in an equilateral triangle using a cc-pVTZ basis and a bond length of 2 Å. At this geometry the ground and first excited state are degenerate and of different symmetry. To test the ability of SA-MCCI to describe excited states at a degeneracy we do not use symmetry. In this case standard MCCI is not able to calculate the excited state but SA-MCCI, as depicted in Fig. 1, works well. Three hundred fifty two CSFs were used to describe both states compared with 36162 SDs for the FCI when symmetry is not considered. We note that we needed to use symmetry to achieve degeneracy in the Molpro\(^13\) FCI calculations.
Given the following positions in angstrom H(1) (1, 0, 0), H(2) (−1, 0, 0) H(3) (0, \sqrt{3}, 0) we now vary the y position of H(3) to see how SA-MCCI behaves in the vicinity of the point of degeneracy. Again we do not use symmetry in the SA-MCCI calculation but depict the FCI results of \( A_1 \) and \( B_1 \) symmetry within \( C_{2v} \). Starting from the HF reference appears challenging for SA-MCCI here and often the excited state is much too high in energy. We attribute this to the configurations comprising the ground-state wavefunction as having zero matrix elements with the excited-state due to symmetry and it seems that SA-MCCI can sometimes fail to include sufficient states of the excited symmetry when we move away from the point of degeneracy. To overcome this we use the SA-MCCI states found for the point of degeneracy as the starting point for each calculation. As we now have sufficient states of both symmetry to begin with then this approach works well: Fig. 2 shows that with a cut-off of \( c_{\text{min}} = 10^{-3} \) the shape of the curves near the degeneracy are reproduced.

We may quantify the match of this curve using the measure introduced in Ref. 12. There it was shown that the standard deviation of the difference in energies (\( \sigma_{\Delta E} \)) can be used as a measure of the error between two potential curves. This takes into account all the points and that the curves may be shifted by a constant. We find \( \sigma_{\Delta E} = 7.7 \times 10^{-2} \) kcal/mol for the ground and \( \sigma_{\Delta E} = 8.6 \times 10^{-2} \) kcal/mol for the excited curve when the cut-off is \( 10^{-3} \). While at \( c_{\text{min}} = 10^{-4} \) the curve is almost indistinguishable from the FCI here \( \sigma_{\Delta E} = 9.7 \times 10^{-4} \) kcal/mol for the ground and \( \sigma_{\Delta E} = 1.4 \times 10^{-3} \) kcal/mol for the excited state. This measure is useful if one wants to quantify how well an approximate method gives the shape of a single curve. However, when we want to quantify how well an approximate method gives the shape of several curves such an approach applied to each curve may not be fair if it results in shifting an excited state curve by a different constant to the ground state. One way to account for this is to use the vertical excitation energies from the lowest state at each point of the curve, as such a quantity is independent of the overall constant. However, this would not detect if the ground curve was poor but the curves of the second and third states are good. An average of vertical excitation energies between all states could be used to prevent this. However, to facilitate easy comparison with results only for the ground-state, we extend our method of Ref. 12 to multiple states. Here we average our measure over the states of interest when all the curves from an approximate method may be shifted by a constant \( c \),

\[
A = \frac{1}{s} \sum_{j=1}^{s} \frac{1}{M} \sum_{i=1}^{M} (\Delta E_{i,j} - c)^2. \tag{2}
\]

Here \( \Delta E_{i,j} = E_{i,j}^{\text{FCI}} - E_{i,j}^{\text{approx}} \) and \( E_{i,j} \) corresponds to the energy of state \( j \) at geometry \( i \) when we consider \( s \) states and \( M \) geometries. We find the minimum of this error with respect to \( c \) by setting \( \frac{\partial A}{\partial c} = 0 \). This results in

\[
c = \frac{1}{s} \sum_{j=1}^{s} \frac{1}{M} \sum_{i=1}^{M} \Delta E_{i,j}. \tag{3}
\]

Putting this into Eq. (2) leads to a similar result to Ref. 12:

\[
\min_{c} A = \sigma_{\Delta E}^2, \tag{4}
\]

but now all states of interest, in addition to geometries, are considered for the calculation of the variance. Hence, we can use \( \sigma_{\Delta E} \) to quantify the match between a number of states and geometries and it will suffice to declare how many states are under consideration. For the two states near the point of degeneracy depicted in Fig. 2 we find \( \sigma_{\Delta E} = 8.1 \times 10^{-2} \) kcal/mol for \( c_{\text{min}} = 10^{-3} \) and \( 1.3 \times 10^{-3} \) kcal/mol for \( c_{\text{min}} = 10^{-4} \).

1. Excited potential curves and phase change

For another test of SA-MCCI we consider moving the third hydrogen on a circle of radius 0.4 Å centred on the previous point of degeneracy: (0, \( \sqrt{3} \), 0) Å. We also investigate if SA-MCCI can detect a phase change\(^{17}\) in this case. We now work with Slater determinants (SDs) rather than CSFs to allow more straightforward calculation of the overlap between SA-MCCI wavefunctions at different geometries. In addition...
we do not use a convergence check here. Due to the use of SDs we also use a smaller cut-off. We use the following co-ordinates in angstrom: H(1) (1, 0, 0), H(2) (-1, 0, 0), H(3) (0.4 cos(θ), √3/2 - 0.4 sin(θ), 0). Again we do not use symmetry. The energy curve is displayed in Fig. 3 where we see that no crossings occur. We note that the points at 90° and 270° were not smooth unless the previous point was used as the starting guess. This again suggests that SA-MCCI has difficulty when including states of a different symmetry when symmetry is not being exploited in the calculation. We find that the SA-MCCI curves with this small cut-off give a result that is almost the same as that of FCI (σΔE = 4.6 × 10^{-3}) and without symmetry MCCI used 2442 SDs on average compared with the full CI space of 36 162 SDs.

To calculate the overlap between SA-MCCI wavefunctions at different geometries we first use Molpro\textsuperscript{13} to calculate the overlap of the AOs between different geometries. The MO coefficients are then used to construct the MO overlaps between the two geometries. The overlap between SDs D1 and D2 at different geometries is\textsuperscript{18}

\[
\langle D_1|D_2 \rangle = \det(S),
\]

where S\textsubscript{ij} = ∫ φ\textsuperscript{*}i(x)φ\textsuperscript{j}(x)d\textit{x} and i labels the spin-MOs in D1 while j labels those in D2. As all coefficients are taken to be real, we find that with a step size of 10\textdegree the absolute overlap between consecutive SA-MCCI wavefunctions is always greater than 0.988 and by requiring the overlap to be positive we see that the wavefunction changes sign when we traverse the circle. The overlap seems to be large enough to suggest that SA-MCCI with SDs can display the expected phase change when traversing around the conical intersection in this case.

We next test SA-MCCI on systems that have a degeneracy or avoided crossing that is not a consequence of the symmetries of the molecule.

B. LiF avoided crossing

The adiabatic potential curves for the first two 1Σ\textsuperscript{+} states of LiF are known to exhibit an avoided crossing, this was seen, for example, by Bauschlicher, Jr., and Langhoff using FCI and SA-MRCl.\textsuperscript{19} There a change from ionic to covalent character was demonstrated for the ground state using the dipole moment while the opposite behaviour was seen for the excited state. It was later shown\textsuperscript{20} that CASPT2 could have difficulties with the curve of LiF: a double crossing was observed in this case rather than an avoided crossing. Later, multi-state CASPT2\textsuperscript{21} which uses multiple SA-CASSCF states as the reference, was demonstrated to be able to describe the curve.

FCI curves for LiF were also considered in Ref.\textsuperscript{22} including those for the 6-31++G* basis. Using this basis and three frozen cores we also calculate the FCI curves for a range of geometries with Molpro.\textsuperscript{13} The FCI space contains around 7 × 10\textsuperscript{6} SDs when considering symmetry. With FCI the gap at the avoided crossing at 13.5 bohrs is 0.036 eV. We return to using SA-MCCI with CSFs and a convergence check. We note that with c\textsubscript{min} = 5 × 10\textsuperscript{-4} the curve at large R was not as smooth as we would have liked as small undulations occurred after R = 15 bohrs. However, the avoided crossing gap occurred at 13.5 bohrs, in agreement with the FCI result, and was 0.053 eV. The results are displayed for c\textsubscript{min} = 3 × 10\textsuperscript{-4} which used 4186 CSFs on average and resulted in a gap of 0.035 eV at 13.5 bohrs (Fig. 4). For c\textsubscript{min} = 3 × 10\textsuperscript{-4} we find that for the two states considered σΔE = 0.325 kcal/mol while this error was almost doubled, although still not too large, at σΔE = 0.602 kcal/mol for the larger cut-off of c\textsubscript{min} = 5 × 10\textsuperscript{-4}.

Reference\textsuperscript{9} compared projector Monte Carlo FCI with SDs (PMC-SD) for excited states with standard FCI using the basis of Ref.\textsuperscript{19} (Li (9s 5p)/(4s 2p); F (9s 6p 1d)/(4s 3p 1d)). They found that PMC-SD almost gave the same energies as FCI when using around 5 × 10\textsuperscript{6} walkers. The FCI results are reported to 3 decimal places only in the paper so we recreate their FCI results using Molpro.\textsuperscript{13} Two frozen cores are employed which means there are around 7.7 × 10\textsuperscript{7} SDs in the FCI when considering symmetry while the mean number of CSFs at c\textsubscript{min} = 5 × 10\textsuperscript{-4} to describe both the ground and excited state in SA-MCCI was 2734.

We see in Fig. 5 that, although the SA-MCCI results are noticeably higher in energy than the FCI, the avoided crossing...
is reproduced and the shapes of the curves appear close to that of the FCI. The avoided crossing gap at 11.5 bohrs is 0.113 eV when using FCI while SA-MCCI gives 0.112 eV. We find for the potential curves from the two SA-MCCI states that $\sigma_{\Delta E} = 0.368$ kcal/mol. When using both the PMC-SD and FCI results to 3 decimal places as reported in Ref. 9 we see that $\sigma_{\Delta E} = 0.312$ kcal/mol. If we also restrict the SA-MCCI comparison to three decimal places then we find that the error in the SA-MCCI curves increases slightly to $\sigma_{\Delta E} = 0.372$ kcal/mol. Although the energies are higher in SA-MCCI the shape of the curves are very good with only a slightly larger error than PMC-SD despite using a wavefunction consisting only of a few thousand CSFs. However, given the paucity of points and that the $\sigma_{\Delta E}$ value for PMC-SD is based on results rounded to three decimal places caution should be used with the comparison.

1. Computational efficiency

In Ref. 23 the scaling of MCCI with Slater determinants was considered. The cost of iterative diagonalization was taken as $O(N^2)$ where $N$ is the number of configurations so a possible reduction of time when using MCCI is $O(N_{MCCI}^2/N_{FCI})$. However the number of iterations needed for MCCI may affect the scaling and it is noted that FCI programs can improve efficiency by exploiting the sparsity of the Hamiltonian matrix.

Comparing SA-MCCI directly to methods other than FCI is problematic because SA-MCCI can be applied to single-reference or multi-reference wavefunctions and ground states or excited states with or without degeneracies. Furthermore SA-MCCI produces a compact wavefunction that can be used in the subsequent calculation of properties. Efficient methods for specific problems such as single-reference wavefunctions would be expected to surpass MCCI in their domain of applicability but may be qualitatively wrong for other problems. CASSCF perhaps followed by perturbation or MRCI can be applied generally but its efficiency is dependent on the choice of active space and the time required to find an appropriate active space, which may be difficult for excited states, and it is not clear how to take these aspects into account. The suggestion of an appropriate active space for excited states could perhaps be a future application of MCCI with large cut-offs.

A comparison with FCI is still not straightforward as FCI memory requirements increase quickly with the size of the basis and so can prevent FCI calculations when MCCI can still be run. If FCI is possible then the MOLPRO13 FCI program used here scales as $O(N_{FCI}m^4)$ where $m$ is the number of orbitals. Hence if the fraction of FCI space needed by MCCI is sufficiently small then MCCI will be more efficient than FCI, but we cannot know beforehand if this is the case. Furthermore, the cut-off value and convergence criteria for the MCCI calculation will affect the balance between time and accuracy. With these caveats we briefly look at the computational cost of the results for LiF as they capture the behaviour of the FCI results, demonstrated by the small errors ($\sigma_{\Delta E}$), and the calculations are of reasonable duration. For the 6-31++G* basis the FCI calculation required 0.43 h per point on average while the SA-MCCI calculations used 0.09 h per point and 0.21 hours per point for cut-offs of $5 \times 10^{-4}$ and $3 \times 10^{-4}$, respectively. However the SA-MCCI calculation was run on 12 processors while the FCI program is not parallelized. SA-MCCI therefore requires 1.12 processor hours per point for the large cut-off and 2.52, for the smaller cut-off. For the results when using the basis of Ref. 19 the number of FCI configurations increases by an order of magnitude and now the FCI calculation uses 5.5 h per point and the SA-MCCI calculation for $c_{min} = 5 \times 10^{-4}$ required 0.09 h per point or 1.11 processor hours per point. Although when considering processor hours this implementation of SA-MCCI was less efficient than FCI for this system when the FCI space consisted of $\sim 7 \times 10^6$ SDs, it appeared more efficient when the FCI space was $7.7 \times 10^7$ however we acknowledge that the larger calculation was over a smaller range of geometries. These values approximately conform to the expected scalings in that the FCI space has increased by an order of magnitude and so has the FCI calculation time while the size of the SA-MCCI space is similar resulting in similar calculation time for the SA-MCCI calculation. We note that if the size of the FCI space were to continue to increase in magnitude then the FCI calculation would quickly become intractable and the small number of configurations used for the SA-MCCI wavefunctions means that there can be further efficiencies, compared with the FCI wavefunction, if they are later used for the calculation of properties.

C. CH2 conical intersections

Conical intersections of the $2^1A''$ and $3^1A''$ states of CH2 when using $C_s$ symmetry (1 $^3A_2$ and $2^3B_1$ when the molecule has $C_{2v}$ symmetry) were found directly in Ref. 25. When the angle of the carbon atom from the centre of mass of the hydrogens was at $\theta = 90^\circ$ a seam of conical intersection was reported. We test if by using the basis of Ref. 25 SA-MCCI can give degenerate energies when following the seam found in Ref. 25. Here the carbon to hydrogen distance R(CH) and hydrogen to hydrogen distance R(HH) are both varied. We see in Fig. 6 that the shape of the curves are as expected and that...
the $1^3A_2$ and $2^3B_1$ states from SA-MCCI are essentially degenerate for much of the curve although there is a small gap noticeable around $R_{HH} = 4$ bohrs. However, this only used in the region of $10^6$ CSFs (12 718 for the $2^3B_1$ state on average) compared to around $5 \times 10^5$ SDs that would be needed for a FCI in $C_{2v}$ symmetry. We note that the second-order CSF expansions using SA-CASSCF orbitals used to calculate the curve in Ref. 25 required around $6 \times 10^5$ configurations. It is not clear whether this small gap is due to SA-MCCI not being accurate enough here or that the FCI would also display a gap due to the FCI conical intersection seam for $\theta = 90^\circ$ occurring at slightly different geometries to the results of Ref. 25.

We now look at potential curves when $\theta \neq 90^\circ$, i.e., $C_s$ symmetry with a smaller basis so that we may compare with FCI results. Using a cc-pVDZ basis with one frozen core and $c_{\text{min}} = 5 \times 10^{-4}$ we fix the separation of the hydrogens at 2 Å and vary the angle and distance of the carbon atom from the centre of mass of the hydrogens.

The assignment of the curves in Fig. 7 is not straightforward when $\theta = 30^\circ$ as in the FCI triplet calculation the fourth state crosses the third around $R = 0.45$ and this state is not found in the triplet SA-MCCI calculation. Due to the use of CSFs in MCCI we can calculate the ground state quintet and find that this corresponds to most of the FCI curve which was not detected in the SA-MCCI triplet calculation. The occurrence of this state is attributed to the use of Slater determinants in a FCI calculation resulting in the possible inclusion of higher spin states. We see that there is not a crossing of the $2^1A''$ and $3^1A''$ states for the range considered. Despite the complexity of the potential energy surface we find that SA-MCCI performs well here: for the first three triplet states we find that $\sigma_{\Delta E} = 0.34$ kcal/mol.

At $\theta = 60^\circ$ there is a conical intersection between the second and third state at $R \approx 0.45$ Å and this is described well by SA-MCCI (Fig. 8). There is again a crossing between the third and fourth FCI states which is not detected as a triplet using SA-MCCI and as before we find that it is due to a ground-state quintet. When using the first three triplet states we find a good match between SA-MCCI and FCI with $\sigma_{\Delta E} = 0.28$ kcal/mol.

The FCI calculation has an SD space of around $1.1 \times 10^6$ when symmetry is taken into account while SA-MCCI with $c_{\text{min}} = 5 \times 10^{-4}$ required, on average, 7247 CSFs for $\theta = 60^\circ$ and 6425 CSFs for $\theta = 30^\circ$. With this very small fraction of the FCI space SA-MCCI was still able to produce accurate excited potential curves and demonstrate conical intersections between the second and third triplet states.

D. Dipole of the $2^1A_1$ state of CO

As alluded to in the Introduction, in earlier work it was found that the dipole of the first excited state of $A_1$ symmetry for carbon monoxide could be calculated with MCCI and was close to the FCI result, but there were large oscillations in the MCCI dipole during the calculation. These oscillations caused the dipole to fluctuate from around 0.2 e bohrs to almost 0.8 e bohrs. We attribute this to the MCCI set of CSFs at times being unable to describe the ground state so the first excited state is now a higher excited state. We now check if SA-MCCI can remedy this. The SA-MCCI dipole of the
first excited singlet state of CO is displayed in Fig. 9. On iteration 100 the state averaging type calculation used 18,765 CSFs while the standard calculation had 8,988 CSFs. Here the standard MCCI result was 0.61 e bohrs while the SA-MCCI result was 0.58 e bohrs which is slightly closer to the FCI result of 0.56 e bohrs. The erratic behaviour when not using state-averaging has been eliminated in the SA-MCCI case. The larger space means that more time is needed for the SA-MCCI calculation here but the lack of oscillations means that convergence of the calculation could be checked to finish the computation in much fewer iterations.

E. Vertical excitations of organic molecules

Electronic excitation energies of medium-sized organic molecules were calculated using CASPT2 and a range of coupled-cluster approaches in Ref. 26 when using a def1-TZVP basis. Vertical excitation energies using a geometry optimized using MP2/6-31G* were reported. FCI results are not available due to the very large size of the FCI space for these molecules when using this basis. We therefore compare MCCI results with these benchmarks for some of the smaller organic molecules. As the molecules in the ground state are in, or close to, their def1-TZVP equilibrium structure then one would expect methods based on a single reference to be appropriate except when the excited state has important configurations involving substitutions beyond the method employed. For example, EOM-CCSD performs well for many of the systems in Ref. 26 but for the 2 1A’ transition of formamide it gives 4.52 eV compared with 7.44 eV for CASPT2. These systems, therefore, represent a useful benchmark of MCCI even if MCCI would not be the first choice for the calculation of some of these states. Particularly of interest here are those excitations within a symmetry class to allow the efficacy of SA-MCCI to be observed.

For fairness, excitations energies to states of the same symmetry use the energy from the SA-MCCI calculation for the ground and excited states. While excitations between different symmetries use the energies from single state calculations unless otherwise stated. Core orbitals were frozen for each molecule: two for ethene; two for formaldehyde, three for cyclopentene, three for formamide and four for E-butadiene. Calculations were carried out at \( c_{\text{min}} = 10^{-4} \) then the resulting converged wavefunction was used as the starting point for a \( c_{\text{min}} = 5 \times 10^{-4} \) calculation.

In preliminary calculations we saw that the excitation energy to the 1 1B1 state of formamide appeared anomalously high given that the size of the FCI space is much less than cyclopentene and comparable with ethene. We started this calculation with a B1 configuration formed by exchanging the HOMO of B1 symmetry with the LUMO of A1 symmetry in the HF reference. We now label this configuration CSF1. MCCI should be insensitive to the choice of reference configuration as long as the symmetry is correct and the cut-off is low enough. It seems that the latter condition was not satisfied here as a CIS calculation suggests that the configuration formed by swapping the highest occupied A1 orbital with the lowest unoccupied B1 orbital in the HF reference (CSF2) is more important based on its coefficient.

To reduce the chance of the reference affecting the result and reduce the input needed, we modified MCCI so that on the first iteration all single substitutions of the HF reference that give a required symmetry are used to create the initial MCCI space.

In this case we get a lower excitation energy, and the same dominant configuration (CSF2) as suggested by CIS, at the end of the MCCI run. We find that when starting with CSF1 this initial configuration remains as the most important although CSF2 appears in the final wavefunction with a small coefficient (\( \sim 10^{-2} \)) in a \( c_{\text{min}} = 0.001 \) calculation. An EOM-CCSD calculation gives the second excited state of B1 symmetry with a dominant excitation which is the HOMO B1 to the LUMO A1. This suggests that MCCI has become confined in the second excited state when starting with CSF1 and that the other CSFs, at this cut-off, are such that they appear to support CSF1 so that it remains dominant even when CSF2 is included.

We use the approach of including all single-substitutions resulting in a required symmetry from the HF reference as the starting point for all subsequent calculations except for multiple states of the completely symmetric representation when this approach caused problems in the diagonalization.

In Table I we display the singlet excitation energies for organic molecules in order of increasing FCI space (Table II). The formaldehyde results of different symmetry are close to the CASPT2 results at a cut-off of \( 10^{-3} \) and become closer as \( c_{\text{min}} \) is lowered to \( 5 \times 10^{-3} \). The SA-MCCI result for the 2 1A1 excitation is fairly large compared with the CASPT2 result, but is still within 0.9 eV of it. For ethene the results agree very well with CASPT2 with the smallest cut-off MCCI excitation energy a little bit lower. The difference between the methods is more noticeable for cyclopentene where the \( c_{\text{min}} = 10^{-3} \) results are larger by more than 1 eV although the ordering of the states agrees. Reducing the cut-off reduces the difference to about 0.7 eV. The 0.001 cut-off result for the 1 1A’ state of formamide agrees surprisingly well with the CASPT2 result but this may be attributed to the ground state energy still being somewhat high when taken from the SA-MCCI 3 A’ calculation. The 2 1A’ state excitation is around 1.4 eV higher than...
The CASPT2 result at the lower cut-off but, interestingly, the 3 1 A' state excitation is lower at both cut-offs. E-butadiene had a similar problem with the larger cut-off giving essentially the HF wavefunction for the 1 1 A state and the use of the ground state energy of the 2 1 A state in its place leading to a better match for the 1 2 B excitation at this cut-off with CASPT2 than at the smaller cut-off. However, they are both more than 1 eV higher. The agreement with CASPT2 is poorest for 2 1 A here with a 3 eV difference even at \( c_{\text{min}} = 5 \times 10^{-4} \) suggesting that the fraction of the FCI space considered by MCCI is too small here. The 2 1 A state is considered a doubly excited state and the use of the ground state from the SA-MCCI 2 1 A calculation as the 1 1 A calculation gave essentially the HF energy.

### 1. Oscillator strengths

When using atomic units, the electric dipole oscillator strength may be written as

\[
f_{ab} = \frac{2}{3} \Delta E |D_{ab}|^2,
\]

where

\[
D_{ab} = \langle \Psi_a | \hat{r} | \Psi_b \rangle.
\]

As we freeze orbitals before producing the integrals then this needs to be taken into account in the transition dipole moment calculation by using

\[
D_{ab} = r_{\text{core}} \langle \Psi_a | \hat{r} | \Psi_b \rangle + \langle \Psi_a | \hat{r} | \Psi_b \rangle.
\]

The oscillator strength would perhaps be expected to be more sensitive to the details of the wavefunction than the excitation energy. The results are displayed in Table III and the absolute differences between MCCI and CASPT2 oscillator strengths are generally reasonably small. Although the percentage errors would be much higher than those of the excitation energies in many cases. The MCCI formaldehyde f values are similar in order of magnitude to those of CASPT2. The 1 2 B value for MCCI is around ten times less than that of CASPT2 but both results are very small. Given the agreement with the excitation energies the MCCI f value at around 1 being more than double that of CASPT2 for the 2 1 A transition is unexpected. The methods, however, agree well for ethene and also for cyclopropane although the very small 1 2 B values would have a large percentage error in the latter case. The 1 1 A values for formamide are small but agree well, however the result for 2 1 A is very different, when using MCCI at the smaller cut-off, to CASPT2. There is also a discrepancy for the 3 1 A' result but this is less marked. The e-butadiene SA-MCCI results are also fairly in good agreement with CASPT2.

### 2. MCCIPT2 results with approximate natural orbitals

Second-order perturbation theory on a MCCI wavefunction (MCCIPT2)\textsuperscript{12} was shown to improve the accuracy of the N\textsubscript{2} potential curve in Ref. 12 and approximate natural orbitals were demonstrated to be able to increase the efficiency of a MCCI calculation. We now investigate how MCCIPT2 with approximate natural orbitals fared in the calculation of the excited states. MCCIPT2 adapts a CI perturbation scheme from

| Molecule | Excitation | 0.001 | 0.0005 | CASPT2\textsuperscript{26} |
|----------|------------|-------|--------|-----------------|
| Formaldehyde | 1 1 A\textsubscript{2} | 4.45 | 4.17 | 3.98 |
| | 1 2 B\textsubscript{1} | 9.69 | 9.40 | 9.14 |
| | 2 1 A\textsubscript{1} | 10.54 | 10.14 | 9.31 |
| Ethene | 1 2 B\textsubscript{1u} | 8.73 | 8.57 | 8.62 |
| Cyclopropane | 1 2 B\textsubscript{2} | 8.02 | 7.44 | 6.76 |
| Formamide\textsuperscript{a} | 1 1 A' | 5.39 | 6.70 | 5.63 |
| | 2 1 A' | 9.11 | 8.85 | 7.44 |
| | 3 1 A' | 10.18 | 9.93 | 10.54 |
| E-butadiene\textsuperscript{b} | 1 2 B\textsubscript{u} | 7.58 | 7.79 | 6.47 |
| | 2 1 A\textsubscript{g} | 10.33 | 9.74 | 6.83 |

\textsuperscript{a}0.001 ground state from the SA-MCCI 3 1 A' calculation as the 1 1 A' calculation gave essentially HF energy.

\textsuperscript{b}0.001 ground state from the SA-MCCI 2 1 A\textsubscript{g} calculation as the 1 1 A\textsubscript{g} calculation gave essentially the HF energy.

### TABLE III. MCCI and CASPT2 oscillator strengths using the def1-TZVP basis.

| Molecule | Excitation | 0.001 | 0.0005 | CASPT2\textsuperscript{26} |
|----------|------------|-------|--------|-----------------|
| Formaldehyde | 1 1 A\textsubscript{2} | 0 | 0 | ... |
| | 1 2 B\textsubscript{1} | 1.60 \times 10^{-3} | 1.78 \times 10^{-3} | 1.3 \times 10^{-2} |
| | 2 1 A\textsubscript{1} | 1.12 | 1.07 | 0.451 |
| Ethene | 1 2 B\textsubscript{1u} | 0.494 | 0.443 | 0.513 |
| Cyclopropane | 1 2 B\textsubscript{2} | 3.11 \times 10^{-3} | 3.03 \times 10^{-3} | 10^{-2} |
| Formamide\textsuperscript{a} | 1 1 A'' | 0.242 | 0.224 | 0.234 |
| | 2 1 A' | 0.429 | 9.34 \times 10^{-2} | 0.479 |
| | 3 1 A' | 0.632 | 0.441 | 0.163 |
| E-butadiene\textsuperscript{b} | 1 2 B\textsubscript{u} | 1.07 | 1.19 | 0.783 |
| | 2 1 A\textsubscript{g} | 0 | 0 | ... |

\textsuperscript{a}0.001 ground state from the SA-MCCI 3 1 A' calculation as the 1 1 A' calculation gave essentially HF energy.

\textsuperscript{b}0.001 ground state from the SA-MCCI 2 1 A\textsubscript{g} calculation as the 1 1 A\textsubscript{g} calculation gave essentially the HF energy.
Ref. 29 to work with MCCI and employs an efficient approach for the checking and removal of duplicate CSFs in the PT2 space. Here any configurations that contribute more than a certain value to the perturbative energy estimate are added to the MCCI CSF space and a new MCCI wavefunction is found. This should mean that intruder states pose less of a problem when calculating the PT2 correction to excited state energies in this method. The PT2 part of the calculation is currently run in serial for CSFs.

We require a consistent cut-off and also the MCCIPT2 calculation to contain around $10^6$ CSFs or fewer so that the calculation could be completed in a reasonable time with the current version of the code. To this end we used a reasonably large cut-off of $c_{min} = 5 \times 10^{-3}$ for the MCCI calculations. We hypothesise that sufficient important configurations should be found, but that the MCCI wavefunction is compact enough so that the PT2 calculation is not excessively challenging when the FCI space becomes extremely large. To try to improve the efficiency of the calculation we used approximate natural orbitals (NOs) where QCISD NOs are employed for the ground-state while MCCI NOs are used for other calculations. The MCCI NOs are calculated using Slater determinants (SDs). When using a starting configuration consisting of all single substitutions resulting in a required symmetry from the HF wavefunction the Davidson-Liu algorithm had problems diagonalizing the Hamiltonian matrix on the first step. In this case we carried out a complete diagonalization of the Hamiltonian matrix and the eigenvectors of interest were then used to create the initial $b_k$ vectors for the Davidson-Liu routine.

Where excited states of the same symmetry were used SA-MCCI was utilised and the state average of the first-order reduced density matrix $\rho = \frac{1}{i} \sum_i \rho_i$, was used to attempt to fairly construct MCCI NOs. The procedure for these calculations is summarised below:

1. Calculation of SA-MCCI NOs using SDs with 50 iterations and $c_{min} = 5 \times 10^{-4}$ with a starting configuration comprising all symmetry consistent single substitutions for all but ground state calculations where the QCISD NOs were calculated.

2. SA-MCCI with NOs, a starting configuration comprising all symmetry consistent single substitutions and $c_{min} = 5 \times 10^{-3}$ until convergence in the energy of 0.001 hartree.

3. MCCIPT2 is then carried out on the converged wavefunction. Any CSF in the PT2 space that contributes more than the cutoff is incorporated and the process repeated until this does not occur.

We see in Table IV that the MCCIPT2 results are closer to the CASPT2 results than the MCCI results were. The PT2 calculations included from one million to a hundred million configurations which tended to increase as the FCI space did. Now the discrepancy is always within 1.1 eV, with formamide giving the largest difference. The formamide $3^1A'$ excitation is noticeably lower than the CASPT2 result using MCCIPT2 and using SA-MCCI which suggests that the FCI result may be lower than the CASPT2 estimate. The order of the states is the same except for cyclopropene where the close $1^1B_1$ and $1^1B_2$ excitations have swapped round. We repeated the cyclopropene calculations with $c_{min} = 3 \times 10^{-3}$ which resulted in a larger MCCI wavefunction and PT2 space: around $10^8$ configurations were needed for the ground state and $2 \times 10^8$ for the excited states in the PT2 calculation. The $1^1B_1$ transition was again higher in energy at 7.46 eV but was closer to the $1^1B_2$ transition occurring at 7.42 eV than before.

We also look at the excitation energies with MCCIPT2 when using the improved and larger def1-TZVP basis sets. The results are displayed in Table V and we see that the values are similar, particularly the different order of the cyclopropene excitations is again observed. The excitation energy for the $1^1B_n$ state of e-butadiene is now much higher than with the def2-TZVP basis but we note that both MCCIPT2 results are within 1 eV either side of the CASPT2 result in the def1-TZVP basis. However, this is the most challenging calculation due to the large space: before the removal of duplicate CSFs in the MCCIPT2 calculation there were more than half a billion CSFs for the $2^1A_g$ state of e-butadiene. Hence, a smaller cut-off calculation with the current

### Table IV. Excitation energies (eV) from MCCIPT2 with $c_{min} = 5 \times 10^{-3}$ compared with CASPT2 when using a def1-TZVP basis.

| Molecule | Excitation | MCCIPT2 | CASPT2$^{26}$ | PT2 configurations |
|----------|-----------|--------|--------------|-------------------|
| Formaldehyde | $1^1A_2$ | 3.98 | 3.98 | $4.2 \times 10^6$ |
| | $1^1B_1$ | 9.08 | 9.14 | $4.7 \times 10^6$ |
| | $2^1A_1$ | 10.22 | 9.31 | $5.8 \times 10^6$ |
| Ethene | $1^1B_{1u}$ | 8.20 | 8.62 | $5.8 \times 10^6$ |
| Cyclopropene | $1^1B_1$ | 7.35 | 6.76 | $4.3 \times 10^7$ |
| | $1^1B_2$ | 6.94 | 7.06 | $3.5 \times 10^7$ |
| Formamide | $1^1A''$ | 6.75 | 5.63 | $6.6 \times 10^7$ |
| | $2^1A'$ | 8.53 | 7.44 | $9.5 \times 10^7$ |
| | $3^1A'$ | 9.69 | 10.54 | $9.5 \times 10^7$ |
| E-butyadiene | $1^1B_u$ | 5.89 | 6.47 | $3.4 \times 10^7$ |
| | $2^1A_g$ | 7.18 | 6.83 | $1.6 \times 10^8$ |

### Table V. Excitation energies (eV) from MCCIPT2 with $c_{min} = 5 \times 10^{-3}$ when using the def2-TZVP basis.

| Molecule | Excitation | MCCIPT2 | PT2 configurations |
|----------|-----------|--------|-------------------|
| Formaldehyde | $1^1A_2$ | 3.43 | $1.2 \times 10^7$ |
| | $1^1B_1$ | 8.87 | $1.3 \times 10^7$ |
| | $2^1A_1$ | 10.04 | $1.5 \times 10^7$ |
| Ethene | $1^1B_{1u}$ | 8.03 | $1.1 \times 10^7$ |
| Cyclopropene | $1^1B_1$ | 7.53 | $1.1 \times 10^8$ |
| | $1^1B_2$ | 6.20 | $1.4 \times 10^7$ |
| Formamide | $1^1A''$ | 6.49 | $1.1 \times 10^7$ |
| | $2^1A'$ | 8.56 | $2.2 \times 10^8$ |
| | $3^1A'$ | 9.33 | $2.2 \times 10^8$ |
| E-butyadiene | $1^1B_u$ | 7.38 | $6.7 \times 10^7$ |
| | $2^1A_g$ | 7.67 | $3.1 \times 10^8$ |
version of the code would be unreasonably time consuming. We note that the ordering of the e-butadiene states agrees with Refs. 26 and 28 for both basis sets and approaches considered.

IV. SUMMARY

We have demonstrated that SA-MCCI can be used to overcome problems arising from the removal of configurations important for the ground state in an MCCI calculation. SA-MCCI was seen to be able to calculate conical intersections, avoided crossings, a dipole moment, and vertical excitations. The conical intersection due to symmetry of $H_3$ in the shape of an equilateral triangle was successfully calculated with SA-MCCI without using symmetry. Potential curves near the degeneracy and when considering moving one hydrogen around a circle centred on a vertex of the triangle were also accurately calculated. We showed that the measure of the error between two potential curves ($\sigma_{AE}$) that we introduced in Ref. 12 could also be used as a fair way to measure the error when comparing approximate potential curves for multiple states with the FCI results. The avoided crossing in the potential curve of the first two $1\alpha_1$ states of LiF was recovered when using SA-MCCI in two bases and the potential curves had low errors when compared with the FCI results despite only using a very small fraction of the FCI space: a few thousand CSFs were needed compared with FCI spaces of around $10^7$ SDs. The seam of conical intersections for the first three triplet states of CH$_2$ was fairly well reproduced by SA-MCCI although there was a slight deviation from degeneracy for some values. When using geometries corresponding to $C_1$ symmetry and a smaller basis then potential curves for the first three triplet states of CH$_2$ agreed well with the FCI results and the use of CSFs in MCCI allowed a FCI state not present in the SA-MCCI triplet calculation to be revealed as a quintet. We saw that SA-MCCI can also be useful in the calculation of properties of excited states: the oscillations in the MCCI dipole of the $2\alpha_1$ state of carbon monoxide were eliminated by using SA-MCCI.

We then considered the excitation energies of a collection of organic molecules up to the size of butadiene and compared SA-MCCI results with those of CASPT2 from Ref. 26 when using the def1-TZVP basis. We found that in general the excitation energies from SA-MCCI were larger than the CASPT2 results of Ref. 26 and the difference tends to be greater when the FCI space is much larger, while decreasing the cut-off lowers the MCCI excitation energy. The ordering of the states was the same. The results agreed reasonably well for formaldehyde, ethene, and cyclopropene in that SA-MCCI and CASPT2 were within 1 eV of each other. For formamide the difference was more than 1 eV and for e-butadiene it was as large as 3 eV suggesting that the FCI space is too large here for MCCI to describe sufficiently well at a cut-off of $c_{min} = 5 \times 10^{-4}$. The agreement in oscillator strength values was reasonable except for the $2\alpha_1$ state of formamide. We then looked at using MCXCIPT2 and approximate natural orbitals albeit with a larger cut-off of $5 \times 10^{-3}$ for the SA-MCCI calculation so that the MCXCIPT2 calculation would not have to consider more than a billion CSFs. The results were now in better agreement with those of CASPT2 with a difference of always less than 1.1 eV with formamide having the largest discrepancy. Interestingly, the order of the two fairly similar cyclopropene excitations in MCXCIPT2 was different from the CASPT2 results. This remained the same in MCXCIPT2 when the cutoff was lowered to $c_{min} = 3 \times 10^{-3}$ although the energies were now closer. We also used MCXCIPT2 with the def2-TZVP basis for the excitation energies of these systems. The only notable difference was that the $1\,^1B_u$ excitation of e-butadiene that was about 1.5 eV higher than in the def1-TZVP basis suggesting that more configurations needed to be considered from the very large FCI space ($10^8$) of this system to get a more stable value. This could be achieved by improving the efficiency of MCXCIPT2, perhaps through parallelisation of the PT2 duplicate removal, to allow smaller cut-offs to be used for these very large FCI spaces.

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