Tailoring CIPSI expansions for QMC calculations of electronic excitations: the case study of thiophene

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The perturbatively selected configuration interaction scheme (CIPSI) is particularly effective in constructing determinantal expansions for quantum Monte Carlo (QMC) simulations with Jastrow-Slater wave functions: fast and smooth convergence of ground-state properties, as well as balanced descriptions of ground- and excited-states of different symmetries have been reported. In particular, accurate excitation energies have been obtained by the pivotal requirement of using CIPSI expansions with similar second-order perturbation corrections for each state, that is, similar estimated errors with respect to the full configuration interaction limit. Here we elaborate on the CIPSI selection criterion for excited states of the same symmetry as the ground state, generating expansions from a common orbital set. Using these expansions in QMC as determinantal components of Jastrow-Slater wave functions, we compute the lowest, bright excited state of thiophene, which is challenging due to its significant multireference character. The resulting vertical excitation energies are within 0.05 eV of the best theoretical estimates, already with expansions of only a few thousand determinants. Furthermore, we relax the ground- and excited-state structures following the corresponding root in variational Monte Carlo and obtain bond lengths which are accurate to better than 0.01 Å. Therefore, while the full treatment at the CIPSI level of this system would be quite demanding, in QMC we can compute high-quality excitation energies and excited-state structural parameters building on affordable CIPSI expansions with relatively few, well chosen determinants.

1. INTRODUCTION

The accurate description of photoinduced phenomena relies on the balanced treatment of the multiple electronic states involved in the excitation process. Recently, we have demonstrated the ability of quantum Monte Carlo (QMC) methods to yield chemically accurate ground- and excited-state structures as well as vertical and adiabatic excitation energies for small, prototypical molecules [1, 2]. We used wave functions of the Jastrow-Slater form where the determinantal component was generated in an automatic manner with the configuration interaction using a perturbative selection made iteratively (CIPSI) approach. If the expansion was then fully optimized together with the Jastrow factor in QMC [3, 4], a handful of CIPSI determinants was found to be sufficient to provide well converged geometries and excitation energies.

Importantly, we showed [2, 5] that a balanced QMC description of the ground and excited states also at different geometries could be achieved by generating CIPSI expansions characterized by the same second-order perturbation (PT2) energy correction, that is, the same “error” with respect to the full CI limit. Furthermore, these “iso-PT2” expansions were found to have similar values of the CI variance which is another useful measure of the error of a CIPSI wave function.

If the excited states investigated belong to a different symmetry class than the ground state, one can perform the expansions either separately, stopping when the same target PT2 energy correction or CI variance is reached [5], or concurrently with a common set of orbitals. In the latter case, the difference in symmetry and the relatively dominant single-reference character of the states investigated aided the CIPSI selection, and we heuristically found that a rather straightforward common selection criterion closely approaches the iso-PT2 condition [2].

If the states are of the same symmetry, separate expansions do not guarantee orthogonality, and the preferred route is the use of concurrent CIPSI expansions on a common set of orbitals. In this case, however, rendering their balanced description is more difficult and the simple selection scheme employed in Ref. [2] proves inadequate, especially if some of the relevant states are strongly multi-configurational.

Here, we propose a simple and effective modification of the selection criterion to enable the construction of nearly iso-PT2 CIPSI expansions for multiple states of the same symmetry, and illustrate the scheme on the challenging case of thiophene. This molecule forms the backbone of a class of π-conjugated donor polymers in organic solar-cell devices [6–11] and the computation of its electronic excited states has been the subject of several theoretical investigations [12–21]. In particular, the accurate prediction of the lowest, bright excited state of thiophene is difficult for traditional ab-initio methods, with different levels of theory spanning a range of about 0.5 eV as illustrated below. The multireference character of this state calls in fact for the use of highly-correlated electronic structure methods and its inherent complexity renders this a perfect test case for our modified CIPSI selection approach.

By simply assigning a higher weight in the CIPSI selection criterion to the state showing a slower convergence, we succeed in generating CIPSI expansions for the ground and excited states of thiophene fulfilling the basic condition of similar PT2 energy corrections and CI variances. The re-
sulting Jastrow-Slater wave functions yield variational Monte Carlo (VMC) vertical excitation energies which are lower and in closer agreement with the diffusion Monte Carlo (DMC) counterparts than those provided by the default selection scheme. Furthermore, we obtain converged estimates of the QMC vertical excitation energy already with compact expansions containing a few thousand determinants, and our best estimates are within 0.05 eV of the reference coupled cluster (CC) value. Finally, we compute the optimal ground- and excited-state geometries in VMC, following the relevant root and generating CIPSI expansions with similar PT2 corrections for both states along the optimization path. The optimal VMC structural parameters are in excellent agreement with the CASPT2 or CC estimates, namely, within 0.01 Å for the bond lengths and, in the excited state, 1° for the bond angles.

This article is organized as follows: We describe the modified CIPSI selection procedure for multiple states of the same symmetry in Section II, and present the computational details in Section III. The construction procedure of the wave functions with the modified selection scheme is detailed in Section IV A. We present the QMC vertical excitation energies in Section IV B and the optimal ground- and excited-state VMC structures in Section IV D. We conclude in Section V.

II. METHODS

In this work, we consider excited states that are not the lowest in their symmetry class. In the QMC calculations, we describe them together with the lower-energy states of the same symmetry via a set of Jastrow-Slater wave functions with different CI coefficients but the same Jastrow and orbital parameters:

$$\Psi_n = J \sum_{k=1}^{N_{\text{det}}} c_k D_k$$

where $N_{\text{det}}$ is the total number of determinants and the index $n$ denotes an electronic state. We use a Jastrow factor which describes electron-nucleus and electron-electron correlations ($J_{\text{2-body}}$), and guarantees that Kato’s cusp conditions are satisfied at the inter-particle coalescence points. To ensure that the common non-linear (Jastrow and orbital) parameters offer a reasonable description of all states of interest, we optimize them by minimizing the state-average (SA) energy \cite{22} defined as

$$E_{\text{SA}} = \sum_{n=1}^{N_{\text{states}}} w_{n}^{\text{QMC}} \frac{\langle \hat{H} | \Psi_n \rangle}{\langle \Psi_n | \Psi_n \rangle},$$

where the weights, $w_{n}^{\text{QMC}}$, sum up to one and are kept fixed during the optimization. Orthogonality between the states is maintained through the (linear) CI coefficients whose optimal values are obtained by solving a generalized eigenvalue problem in the basis of the determinants multiplied by the Jastrow factor.

As described in Ref. \cite{5}, we alternate a number of optimization steps of the non-linear parameters with a step of optimization of the linear coefficients. For the former, we follow the down-hill gradient of the SA energy in a scheme inspired by the stochastic reconfiguration approach for a single state \cite{23}, and solve the relevant equations in a low-memory conjugate-gradient implementation \cite{24}. For the latter, we use a memory-efficient Davidson diagonalization method that allows the computation of the lowest energy eigenvalues without explicit construction of the entire Hamiltonian and overlap matrices \cite{24,25}. Combining this optimization scheme with the fast generation of the quantities needed in the QMC estimators \cite{3,4}, we can optimize QMC wave functions for ground and excited states containing large determinantal expansions and several thousand parameters.

To construct the determinantal component of these wave functions, we employ an improved CIPSI approach that allows us to iteratively select the most important determinants required for the balanced description of multiple electronic states. Starting from an initial reference subspace, $\mathcal{S}$, given by the union of determinants describing the states of interest, an external determinant $|\alpha\rangle$ is selected based on its weighted second-order perturbation (PT2) energy contribution obtained via the Epstein-Nesbet partitioning of the Hamiltonian \cite{26,27}.

$$e_{\alpha} = \sum_{n=1}^{N_{\text{states}}} w_{n} \delta E_{\alpha,n}^{(2)},$$

where

$$\delta E_{\alpha,n}^{(2)} = \frac{|\langle \alpha| \hat{H} | \Psi_n^{\text{CIPSI}} \rangle|^2}{\langle \Psi_n^{\text{CIPSI}} | \hat{H} | \Psi_n^{\text{CIPSI}} \rangle - \langle \alpha | \hat{H} | \alpha \rangle},$$

and $\Psi_n^{\text{CIPSI}}$ is the current normalized CIPSI wave function for state $n$. In this partitioning scheme, the first-order energy correction is zero by definition. The determinant $|\alpha\rangle$ is added to $\mathcal{S}$ if its energy contribution $e_{\alpha}$ is higher than a threshold, and the threshold is automatically adjusted so that the number of determinants in $\mathcal{S}$ is increased by a certain percentage at every iteration.

We have recently shown that the use of iso-PT2 CIPSI expansions results in a balanced description of the relevant states when complemented by a Jastrow factor and fully optimized in QMC, these expansions were found to yield accurate QMC estimates of the excitation energies for relatively small numbers of determinants \cite{2}.

When interested in the lowest-energy states of different symmetries, one can in principle perform the expansion separately for each state until the corresponding perturbation energy contribution is equal to a target value. Alternatively, one can generate the expansions using one set of orbitals for all states and enlarging the union space, $\mathcal{S}$, through a single threshold. Following this last scheme, we were able to obtain matched PT2 corrections \cite{2} by simply choosing the weights in the selection step (Eq. 3) as:

$$w_{n} = w_{n}^{\text{max}} = \frac{1}{\max(c_{k,n}^2)},$$

where the index $k$ runs over all determinants in the current $\Psi_n^{\text{CIPSI}}$. Such a choice follows Ref. \cite{28}, with modifications...
due to the fact that we perform the CIPSI selection in the basis of determinants and not of configuration state functions (CSFs). The resulting expansions for states of different symmetry have different sizes since they do not share any common determinant (determinants of a given symmetry have zero coefficients in the expansion for a state belonging to a different symmetry class).

Here, we are interested in states of the same symmetry expanded on the same set of determinants, where a CSF may have non-zero and non-negligible overlap with all states of interest. In this case, we find that the use of the simple weights (Eq. 5) does not yield iso-PT2 expansions. To improve this balance, we explore a simple modification to the selection scheme where we multiply the weights of Eq. (5) by user-given “state-average” weights, $w_n^{SA}$, i.e.

$$w_n = w_n^{cmax} \times w_n^{SA}.$$  \hspace{1cm} (6)

In addition to the PT2 correction, we investigate the behavior of the CI variance, $\sigma_{CI}^2$, which is defined as the variance of the full CI (FCI) Hamiltonian:

$$\sigma_{CI}^2(\Psi_n^{CIPSI}) = \sum_{\alpha \in \text{FCI}} \langle \Psi_n^{CIPSI} | \hat{H} | \alpha \rangle \langle \alpha | \hat{H} | \Psi_n^{CIPSI} \rangle - \langle \Psi_n^{CIPSI} | \hat{H} | \Psi_n^{CIPSI} \rangle^2,$$ \hspace{1cm} (7)

and goes to zero as the CIPSI wave function approaches the FCI limit. Since the CI variance is also an indicator of the quality of the CIPSI wave function, one can match the CI variances of the states of interest together with the PT2 energy correction or as an alternative to the PT2 criterion. We note that the CI variance should not be confused with the QMC variance which is defined in terms of the exact Hamiltonian.

### III. COMPUTATIONAL DETAILS

All QMC calculations are performed with the program package CHAMP [29]. We employ the Burkatzki-Filippi-Dolg (BFD) scalar-relativistic energy-consistent Hartree-Fock (HF) pseudopotentials and correlation consistent Gaussian basis sets that have been specifically constructed for these pseudopotentials [30, 31]. When unclear, we append the “(BFD)” suffix to these basis sets to avoid confusion with the corresponding all-electron basis sets. For most test QMC calculations, we use a minimally augmented double-\(\zeta\) (maug-cc-pVDZ) basis set, where the basis on the heavy atoms are augmented with s and p diffuse functions. Final calculations are computed with the fully augmented double (aug-cc-pVTZ) and triple (aug-cc-pVQZ) basis sets. All diffuse functions are obtained from the corresponding all-electron Dunning’s correlation-consistent basis sets [32]. We employ a two-body Jastrow factor including electron-electron and electron-nucleus correlation terms [33].

We optimize all parameters (Jastrow, orbital, and CI coefficients) of the Jastrow-Slater wave function in VMC as described above. We employ equal weights $w_n^{QMC}$ in the state-average energy of Eq. (4) and a guiding wave function $\Psi_2 = \sum_n |\Psi_n|^2$ in the sampling to ensure a reasonable overlap with all states of interest [22]. In the VMC geometry optimization, we relax the structure without symmetry constraints and simply follow the path of steepest descent for the root of interest by appropriately rescaling the interatomic forces and using an approximate constant diagonal Hessian. After convergence, we perform 40 additional optimization steps to estimate the optimal average structural parameters. In the DMC calculations, we treat the pseudopotentials beyond the locality approximation using the T-move algorithm [34] and employ an imaginary time step of 0.02 a.u. with single-electron moves. This time step yields DMC excitation energies converged to better than 0.01 eV also for the simplest wave function employed here (see S2).

We carry out the CIPSI calculations with Quantum Package [35] using orbitals obtained from complete active space self-consistent field (CASSCF) calculations performed with the program GAMESS-US [36, 37], correlating six electrons in five \(\pi\) orbitals in a minimal CAS(6,5). The CIPSI expansions are constructed to be eigenstates of \(\hat{S}^2\) and are mapped into the basis of CSFs, effectively reducing the number of optimization parameters in VMC. The PSI4 package [38] is used to compute the reference ground state geometry and the Dalton package [39, 40] is used to compute the vertical excitation energies within the iterative approximate coupled cluster singles, doubles, and triples (CC3) approach. We perform the \(n\)-electron valence perturbation theory (NEVPT2) calculations using the Molpro 2019.2 code [41, 42].

All vertical excitation energies are computed on the fixed ground-state structure optimized at the CC3 level with an all-electron aug-cc-pVTZ basis and the frozen-core (FC) approximation. Unless explicitly stated, the calculations presented below are computed with the BFD pseudopotentials and the corresponding basis sets, irrespective of the level of theory.

### IV. RESULTS AND DISCUSSION

We focus here on the lowest-lying, bright \(\pi \rightarrow \pi^*\) singlet excited state of the thiophene molecule (C\(\text{H}_2\text{S}\)). The ground-state structure of thiophene has C\(\text{2v}\) symmetry and the ground and the targeted excited state belong to the A\(_1\) irreducible representation. The accurate computation of this excited state is challenging because of its multi-reference character: in a CASSCF calculation with the minimal active space correlating six electrons in five \(\pi\) orbitals, one finds that the two dominant transitions, HOMO−1 \(\rightarrow\) LUMO and HOMO \(\rightarrow\) LUMO+1, account for almost 60% and 20% of the wave function, respectively, while higher-order double excitations with four unpaired electrons make up for another 7%. The ground state, on the other hand, is single reference, dominated by the HF determinant which alone accounts for more than 90% of the wave function in the same calculation.

The nature of this excited state leads to difficulties in estimating the corresponding vertical excitation energy, which we find to span a range of about 0.5 eV, between 5.60 and 6.07 eV, across different levels of theory as illustrated in Fig. 1. Time-dependent density functional theory (TDDFT)
FIG. 1. Vertical excitation energy of thiophene for the $\pi \rightarrow \pi^*$ singlet transition to the lowest, bright excited state, computed with different approaches. All calculations are performed with the BFD pseudopotentials and the corresponding aug-cc-pVTZ basis set with the exception of the all-electron CC3/aug-cc-pVTZ (FC) calculation. We also include schematic representation of thiophene, where yellow, blue, and white denote sulfur, carbon, and hydrogen, respectively.

in combination with different exchange-correlation functionals incorrectly places a $\pi \rightarrow \pi^*$ state of $B_2$ symmetry lower than the $A_1$ state by about 0.1 eV or less [19]. All tested wave function methods (CASPT2, NEVPT2, QMC, and CC3), instead, identify our state of interest as the energetically lowest singlet excited state, in qualitative agreement with experimental observations [17, 43–45]; however, they yield very different excitation energies.

Estimating the FCI excitation energy of thiophene at the CIPSI level is challenging because of the size of the FCI space. Consequently, CIPSI calculations are limited to relatively small basis sets and yield extrapolated energy differences characterized by uncertainties as large as 0.08 eV [46]. Since, within a small basis set, CC3 was found to yield an excitation energy of thiophene compatible with the FCI estimated by extrapolating very large CIPSI calculations [46], one can use this level of theory to compute the reference excitation energy for this state of thiophene. Furthermore, the CC3/aug-cc-pVTZ value was shown to differ from the complete basis set limit at the same level of theory by only 0.02 eV [47]. Therefore, since we use a slightly different geometry than the one of Refs. [46] and [47] and the QMC calculations are done with pseudopotentials, we compare our QMC results to the CC3/aug-cc-pVTZ (BFD) excitation energy computed with our geometry, which we believe to be an accurate estimate of the exact vertical excitation energy for our study.

A. Modified selection criterion for CIPSI

For the ground and excited states of thiophene, we construct CIPSI expansions of increasing length from subsequent CIPSI iterations. To test the different selection criteria, we use the

FIG. 2. CI results for the ground- (GS) and excited-state (ES) of thiophene with CIPSI expansions of increasing size: total energy (a); difference of the ground- and excited-state PT2 energy corrections $\Delta E_{PT2}$ (b) and CI variances $\Delta \sigma^2_{CI}$ (c); vertical excitation energy $\Delta E_{CI}$ (d). We employ two reweighting schemes in the selection criterion, namely, $w_{cmax}^n$ and $w_{cmax}^n \times w_{SA}^n$, and different choices of state-average weights, $w_{SA}^n$. The aug-cc-pVDZ basis set is used.

maug-cc-pVDZ basis set and the common set of CAS(6,5) orbitals.

We first adopt the simple reweighting scheme (Eq. 5) that we successfully used for states of different symmetry [2] and plot in panels (b) and (c) of Fig. 2 the resulting differences $\Delta E_{PT2}$ between the PT2 energy corrections and $\Delta \sigma^2_{CI}$ between the CI variances of the ground and excited states (black symbols labeled “cmax”). We find that the convergence of the ground state is faster than that of the excited state: $\Delta E_{PT2}$ decreases and $\Delta \sigma^2_{CI}$ increases over the whole range of expansion sizes considered (see also Table S3). The faster convergence of the ground state is further reflected in the excitation energy $\Delta E_{CI}$, shows in panel (d), which also grows
with the expansions’ size. This is consistent with the observation that partitioning the determinants in each expansion based on the relative contribution to each state gives an insufficient number of “excited-state” determinants, namely, about 1.3-1.6 times more determinants contributing to the excited than to the ground state (see Table S4) compared to a significantly higher ratio of about 2.4-2.9 in our previous calculations with matched PT2 energy corrections for formaldehyde and thioformaldehyde [2]. Therefore, all indications are that the description of the states is increasingly unbalanced in favor of the ground state as the number of determinants gets larger, at least within the range explored.

To achieve a more even description of the two states, we therefore modify the selection criterion by introducing the state-average weights, $w_{n}^{SA}$ (Eq. 6), and place a larger weight on the excited state. We show the resulting difference of the ground- and excited-state PT2 energy corrections and CI variances for sets of expansions generated with different choices of the state-average weights (“$cmax \times SA$”) in Fig. 2. The modified scheme clearly represents an improvement on the bare “$cmax$” criterion and the use of 0.425 and 0.575 as $w_{n}^{SA}$ weights for the ground and the excited state, respectively, leads to nearly optimal matching of PT2 corrections and CI variances for all expansion sizes.

Partitioning the determinants based on their dominant relative contributions to the two states reveals that the ratio between “excited-state” and “ground-state” determinants is now increased to about 1.9-2.1 as shown in Table S4. Furthermore, an inspection of the CI energies $E_{CI}$ obtained with both schemes, plotted in panel (a) of Fig. 3, shows that, for comparable sizes of the expansion, the modified selection criterion slows down the convergence of the ground state and, to a much lesser extent, speeds up the excited state. The CI excitation energy, compared in panel (d) with the CC3 value obtained in the same basis, is also significantly reduced from about 6.6 eV with the “$cmax$” criterion to 5.9 eV for the larger expansions considered here. For comparison, the CC3/aug-cc-pVTZ excitation energy is 5.65 eV. Importantly, we find that the ideal weights depend rather weakly on the orbital basis employed in the determinant selection (see Fig. S1) and also on the basis set (see Figs. S2 and S3), being always very close to 0.4 and 0.6. Finally, we note that, while we set here the state-average weights manually, one can in principle devise a scheme where the values of $w_{n}^{SA}$ are dynamically adjusted during the CIPSI iterations to enforce the iso-PT2 condition as closely as possible.

B. Impact of selection on QMC-CIPSI excitation energies

In Fig. 3 we illustrate the impact of the CIPSI selection on the VMC and DMC vertical excitation energies of thiophene computed with the corresponding Jastrow-CIPSI wave functions fully optimized at the VMC level. To this aim, we consider the CIPSI expansions generated with the old “$cmax$” and the new “$cmax \times SA$” scheme and the ideal weights.

For all but the smallest expansion sizes, the VMC excitation energies corresponding to the “$cmax$” determinantal components settle around about 5.9 eV. Performing DMC calculations for these Jastrow-CIPSI wave functions decreases the excitation energy, which remains however more than 0.1 eV higher than the CC3 reference. Therefore, while both VMC and DMC substantially improve on the starting CIPSI excitation energy, which is as high as 6.6 eV for the largest expansions considered here (see Fig. 2), the bias of the CIPSI selection towards the ground state is reflected in the QMC overestimation of the excitation energy.

When we use the expansions obtained with the “$cmax \times SA$” reweighting scheme, we observe that the convergence of the ground state is somewhat slowed down with respect to “$cmax$” expansions of comparable size, while the VMC excited-state energies are largely unaffected (see Table S1). This leads to reduced VMC excitation energies, which quickly
converge to about 5.8 eV. The VMC correction on the CIPSI excitation energy is now smaller and the same holds for the improvement of DMC upon VMC.

For comparison, in Fig. 3 we also plot the QMC excitation energies computed with Jastrow-Slater wave functions built with three different CAS expansions: the minimal CAS(6,5), a CAS(10,9) correlating two additional occupied \( \sigma \) and two unoccupied \( \sigma^* \) orbitals on the C−S bonds, and a CAS(10,14) that further includes the five 3d orbitals of the S atom. For the largest active space, the expansions are truncated with a threshold of \( 5 \times 10^{-4} \) on the CSF coefficients and the union of the CSFs of the ground and excited states is retained. While increasing the active space lowers the QMC total energies (see Table S1), in all three cases, the VMC excitation energies are compatible within statistical errors and rather comparable to their DMC counterparts, which remain higher the DMC values obtained with the CIPSI expansions and either criterion.

C. Best-quality QMC excitation energy

The use of the \( \text{"cmax } \times \text{SA"} \) criterion to generate iso-PT2 CIPSI expansions has the desired effect of yielding a more balanced description of the two states also at the QMC level. Furthermore, the calibration of the state-average weights we have carried out for the small maug-cc-pVDZ basis helps us in selecting an appropriate range of values also when constructing wave functions with other basis sets.

In particular, since the use of a maug-cc-pVDZ basis set is not sufficient for an accurate treatment of the vertical excitation energy of thiophene, we generate new CIPSI expansions with both the aug-cc-pVDZ and aug-cc-pVTZ basis sets, using the \( \text{"cmax } \times \text{SA"} \) scheme, and find that the choice of weights, 0.4 and 0.6, leads to well matched PT2 contributions and CI variances for both basis sets. As shown in Fig. 4 the resulting VMC and DMC excitation energies are red-shifted relative to comparable expansions in the smaller maug-cc-pVDZ basis, and lie within less than 0.05 eV of our best CC3 reference value for both basis sets.

Finally, we also estimate the vertical excitation energies by matching the VMC variances for the wave functions \[48, 49\]. To this aim, we linearly fit the VMC ground- and excited-state variances separately against their corresponding VMC variances \( \sigma^2 \) and then compute the excitation energy as \( E_{\text{ES}}^\text{fit}(\sigma^2) - E_{\text{GS}}^\text{fit}(\sigma^2) \). As shown in the inset of Fig. 4, we find that the estimate of the excitation energy falls below the CC3 reference and decreases with increasing number of determinants, deviating for the largest expansions by about 0.1 eV. Therefore, while the adoption of the iso-PT2 \( \text{"cmax } \times \text{SA"} \) reweighting scheme yields a more consistent estimate of the excitation energies, we observe a somewhat less predictable behavior when matching the VMC variances.

D. Optimal ground- and excited-state structures

To optimize the structure of thiophene in the ground and excited states in VMC, the maug-cc-pVDZ basis set is found to be sufficiently accurate as verified with the use of the aug-cc-pVDZ basis set in Table S7, and we will therefore proceed with this cheaper basis set to determine the structural parameters of the minima.

For the ground state, we start from the geometry and wave functions previously employed to compute the vertical excitation energies of Table II for selected values of the number of determinants. For all chosen expansion sizes, we obtain very accurate geometries with differences of about 5 mÅ in the bond lengths with respect to the reference values as shown in Table II.

The excited-state global minimum is the S-puckered structure of symmetry \( C_s \) shown in Fig. 5, as obtained in previous studies employing the ADC(2) \[19\], DFT/MRCI \[15, 50\], and MS-CASPT2 \[16\] approaches. It is interesting to note that

**FIG. 4.** VMC (filled) and DMC (empty circles) vertical excitation energies \( \Delta E_{\text{QMC}} \) of thiophene versus the number of determinants for CIPSI expansions generated with the aug-cc-pVDZ (aug-D) and aug-cc-pVTZ (aug-T) basis sets and the \( \text{"cmax } \times \text{SA"} \) scheme. The VMC excitation energy with the aug-T basis is also estimated as the difference \( \Delta E_{\text{QMC}}^\text{fit} = E_{\text{ES}}^\text{fit}(\sigma^2) - E_{\text{GS}}^\text{fit}(\sigma^2) \) of the fits of the energies against the VMC variance of the two states (inset; the line of the fit difference is solid over the range of variances covered by the wave function used, with lower values of \( \sigma^2 \) corresponding to larger numbers of determinants).
TABLE I. VMC and DMC ground- and excited-state energies (a.u.), and vertical excitation energies (eV) of thiophene for increasing CIPSI expansions in fully optimized Jastrow-Slater wave functions and different basis sets. The “cmax × SA” selection criterion is used.

| No. det | No. param | VMC E(GS) | VMC E(ES) | DMC E(GS) | DMC E(ES) | ∆E |
|---------|-----------|-----------|-----------|-----------|-----------|-----|
|         |           | E(GS)     | E(ES)     | E(GS)     | E(ES)     |     |
| aug-cc-pVDZ | 1023 | 3084 | -35.28511(30) | -35.07512(31) | 5.714(12) | -35.35234(29) | 6.690(11) |
|         | 11122 | 9412 | -35.30335(28) | -35.09350(29) | 5.710(11) | -35.35884(26) | 6.698(10) |
|         | 30615 | 16370 | -35.31022(28) | -35.09988(28) | 5.724(11) | -35.36062(28) | 6.699(11) |
| aug-cc-pVTZ | 1019 | 7442 | -35.29270(28) | -35.08212(29) | 5.730(11) | -35.35744(28) | 6.710(11) |
|         | 5051 | 14656 | -35.30375(27) | -35.09326(28) | 5.728(11) | -35.36164(26) | 6.679(10) |
|         | 10755 | 19674 | -35.30913(27) | -35.09948(27) | 5.705(10) | -35.36438(22) | 6.707(09) |
|         | 31581 | 33363 | -35.31600(26) | -35.10596(27) | 5.715(10) | -35.36534(46) | 5.678(17) |
| CC3/aug-cc-pVDZ (BFD) | -0.67 | 1037 | 2002 | 1.4281(2) | 1.3662(1) | 0.06(5) |
|         | -0.64 | 2614 | 2823 | 1.4282(2) | 1.3681(2) | 0.09(3) |
|         | -0.59 | 5605 | 4106 | 1.4290(2) | 1.3669(1) | -0.08(3) |
|         | -0.54 | 11003 | 6326 | 1.4279(4) | 1.3676(4) | 0.05(4) |
| CASPT2 | 1.430 | 1.372 | 1.720 | 0.00 |
| CCSD(T) (BFD) | 1.425 | 1.368 | 1.717 | 0.00 |
| CCSD(T) (FC) | 1.430 | 1.372 | 1.728 | 0.00 |
| CC3 (FC) | 1.430 | 1.372 | 1.729 | 0.00 |
| ES | -0.67 | 1663 | 3630 | 1.4396(7) | 1.4161(6) | 1.7626(5) | 24.84(2) |
|         | -0.63 | 3752 | 4957 | 1.4388(7) | 1.4151(3) | 1.7655(4) | 25.58(3) |
|         | -0.59 | 8304 | 7271 | 1.4383(1) | 1.4144(2) | 1.7709(1) | 25.75(2) |
|         | -0.546 | 15815 | 10278 | 1.4422(7) | 1.4112(4) | 1.7725(6) | 26.01(7) |
| CASPT2 | 1.448 | 1.423/1.416 | 1.782/1.778 | 26.7 |
| ADC(2) | 1.422 | 1.419 | 1.796 | 28.2 |
| DFT/MRCI | 1.436 | 1.394 | 1.799 | 27.4 |

**TABLE II.** Optimal VMC ground- and excited-state bond lengths (Å) and angles (deg) of thiophene using “cmax × SA” CIPSI expansions and the maug-cc-pVDZ basis.

| State | δE<sub>PT2</sub> | No. det | No. param | C−C | C=C | C−S | δ<sub>CCCS</sub> |
|-------|-----------------|---------|-----------|-----|-----|-----|----------------|
| GS    | -0.67           | 1037    | 2002      | 1.4281(2) | 1.3662(1) | 1.7201(2) | 0.06(5) |
|       | -0.64           | 2614    | 2823      | 1.4282(2) | 1.3681(2) | 1.7202(1) | 0.09(3) |
|       | -0.59           | 5605    | 4106      | 1.4290(2) | 1.3669(1) | 1.7218(1) | -0.08(3) |
|       | -0.54           | 11003   | 6326      | 1.4279(4) | 1.3676(4) | 1.7223(4) | 0.05(4) |
| CASPT2 |                 |         |           | 1.430 | 1.372 | 1.720 | 0.00 |
| CCSD(T) (BFD) |         |         |           | 1.425 | 1.368 | 1.717 | 0.00 |
| CCSD(T) (FC) |         |         |           | 1.430 | 1.372 | 1.728 | 0.00 |
| CC3 (FC) |         |         |           | 1.430 | 1.372 | 1.729 | 0.00 |
| ES    | -0.67           | 1663    | 3630      | 1.4396(7) | 1.4161(6) | 1.7626(5) | 24.84(2) |
|       | -0.63           | 3752    | 4957      | 1.4388(7) | 1.4151(3) | 1.7655(4) | 25.58(3) |
|       | -0.59           | 8304    | 7271      | 1.4383(1) | 1.4144(2) | 1.7709(1) | 25.75(2) |
|       | -0.546          | 15815   | 10278     | 1.4422(7) | 1.4112(4) | 1.7725(6) | 26.01(7) |
| CASPT2 |                 |         |           | 1.448 | 1.423/1.416 | 1.782/1.778 | 26.7 |
| ADC(2) |                 |         |           | 1.422 | 1.419 | 1.796 | 28.2 |
| DFT/MRCI |                |         |           | 1.436 | 1.394 | 1.799 | 27.4 |

**FIG. 5.** Optimal ground- and excited-state structures of thiophene.

TDDFT predicts instead a C-puckered minimum with the S-puckered structure actually being a transition state [18]. Since the excited state has the same symmetry as the ground state also when the molecular symmetry is lowered from C<sub>2v</sub> to C<sub>s</sub>, we need to follow the path of steepest descent for the second root in optimizing the excited-state structure. We do not impose any symmetry constraints and start from a slightly distorted geometry, where we re-optimize the same wave functions of Table I selected for the ground-state structural optimization. In the subsequent steps, we generate iso-PT2 CIPSI.
wave functions along the whole minimization path for both the ground and the excited state.

The final converged structural parameters are shown in Table II; the optimized geometry is S-puckered in agreement with previous correlated calculations, with an elongation of all bonds, in particular of the former C=C and the C–S bonds. The bond lengths and angles obtained with Jastrow-CIPSI wave functions are within 10 mA and 1° of the corresponding CASPT2 values, respectively.

V. CONCLUSIONS

In this work, we present a systematic investigation on how to obtain balanced CIPSI expansions for multiple states of the same symmetry and corresponding high-quality QMC excitation energies and optimal excited-state geometries. We focus on a case, the lowest-energy bright state of thiophene, which is characterized by a significant multi-reference character.

To this aim, we adapt here the CIPSI selection scheme to treat multiple states of the same symmetry with wave functions expressed on a common set of determinants, by introducing additional weights in the energy threshold for the selection step. This enables us to obtain expansions for the two states with the same PT2 energy corrections and CI variances that is, similar estimated errors with respect to the full configuration interaction limit. Importantly, we find that the modification introduced in the selection scheme are largely independent of the basis set size and orbital choice as the rate of convergence of the energy appears to be governed by the intrinsic multi-reference character of the excited state.

In practice, for thiophene, the new criterion slows down the convergence of the ground state at every CIPSI iteration and, to a lesser extent, accelerates the one of the excited state, making the quality of the two CI wave functions more similar. Using these expansions as determinantal components in Jastrow-Slater wave functions leads to DMC excitation energies within 0.05(2) eV of the theoretical best estimate available, already when the expansions comprise only about 5000 determinants. With these Jastrow-CIPSI wave functions, structural relaxation in the ground state yields VMC bond lengths which are compatible with the reference values to better than 0.01 Å.

Following the second root in the geometry optimization while maintaining an iso-PT2 description of the ground and excited states, we obtain optimal VMC excited-state bond lengths and angles, which are within less than 0.01 Å and 1° of the best available estimates, respectively.

In summary, also in a case like thiophene where a reliable treatment fully at the CIPSI level is quite demanding, we are able to generate balanced Jastrow-Slater wave functions for multiple states and determine accurate excited-state properties in QMC, using relatively short CIPSI expansions. Therefore, when increasing the number of electrons or the size of the basis set, we expect that QMC in combination with compact and balanced CIPSI expansions will remain a viable route to deliver high-quality excited-state potential energy surfaces in the domain of applications where selected configuration interaction becomes instead intractable.

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SUPPINFO

CIPSI energies, PT2 energy corrections, and CI variances; VMC and DMC ground- and excited-state energies and vertical excitation energies computed with the maug-cc-pVDZ and aug-cc-pVTZ basis sets; dependence of the DMC energies on the time step; dependence of the $w^n$ weights on the choice of orbitals and basis set; partitioning of the CSFs between the two states; comparison with other levels of theory; ground- and excited-state VMC geometries obtained with the aug-cc-pVDZ basis set; ground-state CC3 geometry.

[1] M. Dash, S. Moroni, A. Scemama, and C. Filippi, J. Chem. Theory Comput. 14, 4176 (2018).
[2] M. Dash, J. Feldt, S. Moroni, A. Scemama, and C. Filippi, J. Chem. Theory Comput. 15, 4896 (2019) [https://doi.org/10.1021/acs.jctc.9b00476].
[3] C. Filippi, R. Assaraf, and S. Moroni, J. Chem. Phys. 144, 194105 (2016).
[4] R. Assaraf, S. Moroni, and C. Filippi, J. Chem. Theory Comput. 13, 5273 (2017).
[5] A. Cuzzocrea, A. Scemama, W. J. Briels, S. Moroni, and C. Filippi, J. Chem. Theory Comput. (2020).
[6] J. Roncali, Chem. Rev. 92, 711 (1992).
[7] F. Zhang, D. Wu, Y. Xu, and X. Feng, J. Mat. Chem. 21, 17590 (2011).
[8] L. Bian, E. Zhu, J. Tang, W. Tang, and F. Zhang, Prog. Polym. Sci. 37, 1292 (2012).
[9] Y.-X. Xu, C.-C. Chueh, H.-L. Yip, C.-Y. Chang, P.-W. Liang, J. J. Intemann, W.-C. Chen, and A. K.-Y. Jen, Poly. Chem. 4, 5220 (2013).
[10] R. Meerheim, C. Körner, and K. Leo, Appl. Phys. Lett. 105, 125_1 (2014).
[11] C. Duan, K. Gao, F. J. Colberts, F. Liu, S. C. Meskers, M. M. Wienk, and R. A. Janssen, Adv. Energy Mat. 7, 1700519 (2017).
As Jastrow factor, we use the exponential of the sum of two fifth-order polynomials of the electron-nuclear and the electron-electron distances, respectively, and rescale the inter-particle distances as \( R = (1 - \exp(-\kappa r)) / \kappa \) with \( \kappa \) set to 0.6 a.u. We employ different electron-nucleus Jastrow factors to describe the correlation of an electron with C, S, and H. The total number of free parameters to be optimized in the 2-body Jastrow factor is set to 0.6 a.u. We employ different electron-nucleus Jastrow factors to describe the correlation of an electron with C, S, and H. The total number of free parameters to be optimized in the 2-body Jastrow factor is 17 for thiophene.