Communication: GAIMS—Generalized Ab Initio Multiple Spawning for both internal conversion and intersystem crossing processes

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PERSPECTIVES

[Communication: GAIMS—Generalized](http://dx.doi.org/10.1063/1.4943571) *Ab Initio* **Multiple Spawning [for both internal conversion and intersystem crossing processes](http://dx.doi.org/10.1063/1.4943571)**

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Full multiple spawning is a formally exact method to describe the excited-state dynamics of molecular systems beyond the Born-Oppenheimer approximation. However, it has been limited until now to the description of radiationless transitions taking place between electronic states with the same spin multiplicity. This Communication presents a generalization of the full and *ab initio* multiple spawning methods to both internal conversion (mediated by nonadiabatic coupling terms) and intersystem crossing events (triggered by spin-orbit coupling matrix elements) based on a spin-diabatic representation. The results of two numerical applications, a model system and the deactivation of thioformaldehyde, validate the presented formalism and its implementation. ^C *2016 AIP Publishing LLC.* [\[http:](http://dx.doi.org/10.1063/1.4943571)//[dx.doi.org](http://dx.doi.org/10.1063/1.4943571)/[10.1063](http://dx.doi.org/10.1063/1.4943571)/[1.4943571\]](http://dx.doi.org/10.1063/1.4943571)

I. INTRODUCTION

Nonadiabatic molecular dynamics—coupled electronnuclear dynamics beyond the Born-Oppenheimer approximation—is a key technique for investigating photochemical and photophysical processes of molecules. However, most of the nonadiabatic methods commonly employed focus only on internal conversion (IC), i.e., electronic state transitions that conserve spin multiplicity. They often neglect *intersystem crossing* (ISC) events which couple electronic states with differing spin multiplicity due to relativistic spin-orbit coupling (SOC) .^{[1](#page-6-0)} Far from being a curiosity, ISC plays an important role in the deactivation process of organic molecules^{[2](#page-6-1)[–4](#page-6-2)} and metal complexes^{[5](#page-6-3)} that are used in energy-related devices.^{[6](#page-6-4)[,7](#page-6-5)}

Apart from grid-based exact quantum dynamics for low dimensional problems,^{[8](#page-6-6)} few nonadiabatic dynamics methods have been extended to include SOC effects and, therefore, to describe $ISC⁹$ $ISC⁹$ $ISC⁹$. As one example, the Multiconfiguration Time-Dependent Hartree (MCTDH) technique has been used to understand the role of intersystem crossing in benzene photophysics.^{[10](#page-6-8)} Trajectory surface hopping (TSH), which approximates the dynamics of a nuclear wavepacket by a swarm of independent classical trajectories, has also been modified to incorporate SOC effects.[11](#page-6-9)[–15](#page-6-10) Freedom in the choice of different spin representations^{[11](#page-6-9)[–14](#page-6-11)[,16](#page-6-12)} or hopping schemes^{[17](#page-6-13)} has resulted in a plethora of TSH algorithms, which are, however, prone to shortcomings of the independent trajectory approximation.^{[18](#page-6-14)[,19](#page-6-15)} In particular, the spin-diabatic representation for including SOC in TSH has been questioned due to a lack of rotational invariance for electronic population dynamics.[13](#page-6-16) It can further lead to problematic results whenever sublevels are grouped in single multiplet states. 13

An ideal method for describing both IC and ISC processes would be (i) derivable from *first principles*, (ii) able to treat medium to large molecular systems, (iii) compatible with onthe-fly calculations of electronic quantities, and (iv) able to adequately describe coherence and decoherence effects during nonadiabatic events. Full Multiple Spawning $(FMS)^{20,21}$ $(FMS)^{20,21}$ $(FMS)^{20,21}$ $(FMS)^{20,21}$ or its *ab initio* version $\text{AIMS},^{22,23}$ $\text{AIMS},^{22,23}$ $\text{AIMS},^{22,23}$ $\text{AIMS},^{22,23}$ fulfills all of the previously listed requirements, combing the computational efficiency of trajectory-based methods with quantum dynamics that is *in principle* exact (in the limit of a large enough basis set).

In short, FMS represents a nuclear wavefunction by a swarm of coupled frozen Gaussian functions following classical trajectories. $22-24$ $22-24$ The number of trajectory basis functions used to describe the nuclear wavepacket is adapted during the dynamics—through so-called "spawning" events to accurately describe wavepacket bifurcation in nonadiabatic regions. In the limit of a complete basis set and exact evaluation of all the requisite matrix elements, FMS constitutes a formally exact solution of the time-dependent Schrödinger equation (TDSE). In the following, we show how FMS and AIMS can be extended to the description of both IC and ISC processes and present the resulting Generalized FMS and AIMS (GFMS and GAIMS) methods and their implementation.

II. THEORY

 $\hat{H}_{mol}(\mathbf{r},\mathbf{R}) = \hat{T}_N + \hat{H}_{el}(\mathbf{r},\mathbf{R})$

The original derivation of the FMS method begins with the standard molecular Hamiltonian, 22 22 22 defined as the sum of the nuclear kinetic energy operator and the electronic Hamiltonian,

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where \hat{T}_e is the electronic kinetic energy operator and $\hat{V}_{ee}(\mathbf{r})$, $\hat{V}_{eN}(\mathbf{r}, \mathbf{R})$, and $\hat{V}_{NN}(\mathbf{R})$ are the electron-electron, electron-
pucleus, and pucleus-pucleus potential energy operators nucleus, and nucleus-nucleus potential energy operators, respectively. The collective variables for electronic and nuclear positions are denoted by \bf{r} and \bf{R} , respectively. The Hamiltonian in Eq. [\(1\)](#page-1-1) is non-relativistic and therefore neglects all terms related to both nuclear and electronic spin.

Including SOC effects in the framework of the timedependent Schrödinger equation necessitates the derivation and approximation of new extended electronic Hamiltonians from the Dirac-Coulomb-Breit equation.^{[25](#page-6-22)} The Breit-Pauli Hamiltonian is commonly employed and its most important parts^{[26](#page-6-23)} comprise the non-relativistic Hamiltonian of Eq. (1) , a scalar relativistic part containing the mass-velocity and Darwin terms, and a spin-orbit coupling Hamiltonian.^{[25](#page-6-22)[,27](#page-6-24)} We note that scalar relativistic effects, when important, are often accounted for in quantum chemistry by using effective core potentials.[28](#page-6-25) Incorporating the SOC Hamiltonian in the electronic Hamiltonian, $27,29-31$ $27,29-31$ $27,29-31$ we obtain a molecular Hamiltonian accounting for the electronic spin s,

$$
\hat{H}(\mathbf{x}, \mathbf{R}) = \hat{T}_N + \hat{H}_{el}(\mathbf{r}, \mathbf{R}) + \hat{H}_{SOC}(\mathbf{x}, \mathbf{R})
$$
 (2)

with $\mathbf{x} = (\mathbf{r}, \mathbf{s})$. Inserting Eq. [\(2\)](#page-2-0) into the time-dependent Schrödinger equation leads to the starting equation for our extension of FMS including SOC,

$$
i\frac{\partial \Psi(\mathbf{x}, \mathbf{R}, t)}{\partial t} = \hat{H}(\mathbf{x}, \mathbf{R}) \Psi(\mathbf{x}, \mathbf{R}, t).
$$
 (3)

 $\frac{\partial t}{\partial t}$ The following derivation will use electronic states obtained from a spin-free electronic Hamiltonian as a basis, i.e., each electronic state is an eigenstate of both the total spin operator \hat{S}^2 and the spin projection operator \hat{S}_z .^{[13](#page-6-16)} The time-dependent molecular wavefunction is represented by a Born-Huang expansion 32 in the aforementioned spin-diabatic electronic basis,

$$
\Psi(\mathbf{x}, \mathbf{R}, t) = \sum_{J} \sum_{M_{S_J} = -S_J}^{S_J} \Omega_J^{M_{S_J}}(\mathbf{R}, t) \Phi_J^{M_{S_J}}(\mathbf{x}; \mathbf{R}), \quad (4)
$$

where $\Phi_J^{M_{S_J}}(\mathbf{x};\mathbf{R}) = |$ J, M_{S_J} j. is the electronic wavefunction for the *J*th state with spin multiplicity $(2S_J + 1)$ and spinprojection eigenvalue M_{S_J} . While the formalism is completely general, we focus our attention on singlet $(S = 0)$ and triplet $(S = 1)$ electronic states, the latter having M_S values of -1 , 0, or +1. The FMS ansatz^{[20,](#page-6-17)[21](#page-6-18)} for the nuclear wavefunction becomes

$$
\Omega_{J}^{M_{S}J}(\mathbf{R},t) = \sum_{k'=1}^{N_{J,M_{S}J}(t)} C_{k'}^{J,M_{S}J}(t) \chi_{k'}^{J,M_{S}J} \left(\mathbf{R}; \overline{\mathbf{R}}_{k'}^{J,M_{S}J}(t),\right)
$$

$$
\overline{\mathbf{P}}_{k'}^{J,M_{S}J}(t), \overline{\gamma}_{k'}^{J,M_{S}J}(t), \alpha_{k'}^{J,M_{S}J}\right), \tag{5}
$$

which expresses the nuclear wavefunction for the electronic which expresses the nuclear wavefunction for the electronic
state $|J, M_{S_I}\rangle$ as a linear combination of multidimensional J, M_{S_J} frozen Gaussians $\chi_{k'}^{J,M_S}$ and corresponding complex coefficients $C_{k'}^{J,M_S}$ (*t*). Each term of the linear combination is called a trajectory basis function (TBF). The timedependent position $\overline{\mathbf{R}}_{k'}^{J,M_S}$ *(t)* and momentum $\overline{\mathbf{P}}_{k'}^{J,M_S}$ *(t)* centers for each frozen Gaussian are propagated using classical Hamilton's equations, while the nuclear phase $\overline{\gamma}_{k'}^{J, M_S}$ (*t*) is time-evolved semiclassically.^{[22](#page-6-19)} We note here that the time-dependent parameters in the Gaussian functions could be time-evolved in different ways, leading to techniques for nonadiabatic dynamics like the direct dynamics variational multi-configurational Gaussian^{[33](#page-6-29)} (DD-vMCG) or the multiconfiguration-Ehrenfest (MCE) approach, 34 for example.

Inserting Eqs. (5) and (4) in Eq. (3) leads to a set of equations of motion for the complex amplitudes in Eq. [\(5\).](#page-2-1) After left-projection by $\left\langle \chi \right\rangle$ $\int_k^{I, M_{S_I}} \Phi_I^{M_{S_I}}$ μ , we obtain

$$
\frac{d\mathbf{C}^{I,M_{S_{I}}}(t)}{dt} = -i(\mathbf{S}^{-1})^{II,M_{S_{I}}M_{S_{I}}} \left\{ \left[\mathbf{H}^{II,M_{S_{I}}M_{S_{I}}} - i\dot{\mathbf{S}}^{II,M_{S_{I}}M_{S_{I}}} \right] \mathbf{C}^{I,M_{S_{I}}}(t) + \sum_{J} \sum_{M_{S_{J}}=S_{J}}^{S_{J}} \mathbf{H}^{IJ,M_{S_{J}}M_{S_{J}}}\mathbf{C}^{J,M_{S_{J}}}(t) \right\},
$$
(6)

where bold symbols indicate vectors or matrices in the basis of Gaussian functions. The overlap matrices are defined by $S_{k k'}^{IJ, M_{S_I} M_{S_J}}$ $\binom{IJ, M_{S_I} M_{S_J}}{K, k'} = \left\langle \right\rangle$ $\ddot{}$ I, M_{S_I} \overline{x} $\left\langle \begin{array}{c} J,M_{SJ}\ k' \end{array} \right\rangle$ $\delta_{IJ}\delta_{M_{S_I}M_{S_J}}$ and $\dot{S}^{IJ, M_{S_I}M_{S_J}}_{k \ \ k'}$ $\binom{I J, M_{S_I} M_{S_J}}{k, k'} = \left\langle$ $\ddot{}$ I, M_{S_I} $\frac{\partial}{\partial t}$ $\bigg|$ χ $\left\langle \begin{array}{c} J,M_{SJ}\ k' \end{array} \right\rangle$ $\delta_{IJ}\delta_{M_{S_I}M_{S_J}}$. A general Hamiltonian matrix element in Eq. (6) has the form

$$
H_{kk'}^{IJ,M_S}{}_{I}^{M_S} = \left\langle \chi_{k}^{I,M_S}{}_{I} \right| \hat{T}_{N} \left| \chi_{k'}^{J,M_S}{}_{J} \right\rangle_{R} \delta_{IJ} \delta_{M_S}{}_{I}^{M_S}{}_{J} + \left\langle \chi_{k}^{I,M_S}{}_{I} \right| E_{I}^{el} \left| \chi_{k'}^{J,M_S}{}_{J} \right\rangle_{R} \delta_{IJ} \delta_{M_S}{}_{I}^{M_S}{}_{J} - \sum_{\rho=1}^{3N} \left\langle \chi_{k}^{I,M_S}{}_{I} \right| \left(\mathbf{d}_{IJ}^{M_S}{}_{I}^{M_S}{}_{J} \right) \frac{1}{\rho m_{\rho}} \frac{\partial}{\partial R_{\rho}} \left| \chi_{k'}^{J,M_S}{}_{J} \right\rangle_{R} \delta_{M_S}{}_{I}^{M_S}{}_{J} - \sum_{\rho=1}^{3N} \frac{1}{2m_{\rho}} \left\langle \chi_{k}^{I,M_S}{}_{I} \right| \left(D_{IJ}^{M_S}{}_{I}^{M_S}{}_{J} \right) \right\rangle_{R} \delta_{M_S}{}_{I}^{M_S}{}_{J} + \left\langle \chi_{k}^{I,M_S}{}_{I} \right| \left\langle \Phi_{I}^{M_S}{}_{I} \right| \hat{H}_{SOC} \left| \Phi_{J}^{M_S}{}_{J} \right\rangle_{X} \left| \chi_{k'}^{J,M_S}{}_{J} \right\rangle_{R}.
$$
\n(7)

The first two terms, as well as the electronic diagonal contributions from the fourth term, couple TBFs evolving on the same electronic state with the same sublevel (Fig. $1(a)$, dotted arrows). E_I^{el} is the electronic energy for state *I*, and $m_ρ$ is the mass corresponding to nuclear degree of freedom ρ .

The third and fourth terms in Eq. [\(7\)](#page-2-5) couple TBFs evolving on different electronic states but having the same S and M_S values (Fig. [1\(a\),](#page-3-0) dashed arrows). These terms depend on the firstorder nonadiabatic coupling vectors $\mathbf{d}_{IJ}^{M_{S_I}M_{S_J}}$ and second-

FIG. 1. (a) General representation of the coupling pattern between TBFs in GFMS for a case with two singlet and two triplet states. The Gaussian shape symbolizes the different TBFs (black = singlet, blue = triplet), where a continuous line is used for $M_S = 0$ states, while dashed and dotted lines represent $M_S = -1$ and $M_S = 1$ for triplet states. These TBFs evolve along a classical trajectory represented by the filled circle. Intrastate (dotted arrow) and nonadiabatic (dashed arrows) couplings are present between TBFs evolving in electronic states with the same spin multiplicity (as in the standard FMS method), while GFMS introduces an important number of additional couplings due to SOC (plain arrows). Note that a more detailed representation of the couplings requires the use of separate arrows for each possible coupling between sublevels (not shown). (b) Schematic representation of the GFMS method. A TBF is initiated in S_m at time $t = 0$ (gray) and will at a later time spawn TBFs both in T_n (blue) and in S₀ (black).

order nonadiabatic coupling $D_{IJ}^{M_{S_I}M_{S_J}}$, where the latter is usually neglected in nonadiabatic molecular dynamics.^{[22](#page-6-19)} The novelty of GFMS resides in the last term of Eq. [\(7\)](#page-2-5) which allows for amplitude transfer between electronic states with different spin multiplicity, according to the rules of SOC (Fig. $1(a)$, continuous arrows), and fully preserves rotational invariance. In addition, this last term can also couple TBFs evolving on states that have the same spin multiplicity *S*, but only if the conditions $S_I = S_J > 0$ and $\Delta M_S = 0, \pm 1$ are fulfilled. It is important to note that any definition of the operator \hat{H}_{SOC} can be used in Eq. [\(7\).](#page-2-5)

A key feature of the FMS method is that it uses an adaptive basis set to ensure an accurate description of nonadiabatic processes. The number of TBFs describing the nonadiabatic processes. The number of TBFs describing the nuclear wavefunction in state $|I, M_{S_I}\rangle$, $N_{I,M_{S_I}}(t)$, will indeed

change in time as a result of spawning events. In short, a TBF entering a region of strong nonadiabaticity—detected using an effective coupling—can under certain conditions spawn a new TBF on the coupled electronic state (Fig. $1(b)$). Upon spawning, the size of the matrices in Eq. [\(6\)](#page-2-4) is extended and the resulting coupled propagation of the expanded set of TBFs allows for an exchange of nuclear amplitude between electronic states. For detailed discussions about the spawning algorithm between same-spin states, the reader is referred to previous works. $22,24,35$ $22,24,35$ $22,24,35$ In GFMS, the spawning algorithm needs to be extended to allow for spawning between spinorbit coupled states. Based on an already proposed effective coupling between diabatic states, 22 22 22 we suggest to measure the effective SOC strength between state *I* and state *J* at the position of TBF *k* as

$$
\Lambda_{IJ}^{eff}(\mathbf{R}_k) = \frac{\left(\sum_{M_{S_I}=-S_I}^{S_I} \sum_{M_{S_J}=-S_J}^{S_J} \left| \left\langle \Phi_I^{M_{S_I}}(\mathbf{R}_k) \left| \hat{H}_{SOC} \right| \Phi_J^{M_{S_J}}(\mathbf{R}_k) \right\rangle_{\mathbf{x}} \right|^2 \right)^{1/2}}{\left| E_J^{el}(\mathbf{R}_k) - E_I^{el}(\mathbf{R}_k) \right|}.
$$
(8)

This rotationally invariant spawning measure indicates the overall coupling between the sublevels of state *I* and those of state *J*. If $\hat{\Lambda}_{IJ}^{eff}(\mathbf{R}_k)$ is higher than a certain threshold value, the spawning mode is triggered and a new TBF will be created in each sublevel of the electronic state *J* (see Fig. [1\(b\)\)](#page-3-0).^{[36](#page-6-32)}

Ab initio multiple spawning^{22-[24](#page-6-21)} uses FMS nuclear dynamics combined with *ab initio* electronic structure calculations, allowing for an on-the-fly solution of the molecular time-dependent Schrödinger equation. Two approximations simplify the application of FMS to molecules, namely, the saddle-point and the independent first generation (IFG) approximations. The (zeroth-order) saddle-point approximation is used to compute the integrals over nuclear degrees of freedom that appear in the Hamiltonian matrix.^{[22](#page-6-19)} Extending the saddle-point approximation to the calculation of SOC matrix elements in Eq. [\(7\)](#page-2-5) is straightforward,

$$
\left\langle \chi_{k}^{I, M_{S_I}} \middle| \left\langle \Phi_{I}^{M_{S_I}} \middle| \hat{H}_{SOC} \middle| \Phi_{J}^{M_{S_J}} \right\rangle_{\mathbf{R}} \right\vert \mathcal{H}_{k'}^{J, M_{S_J}} \right\rangle_{\mathbf{R}} \n= \left\langle \chi_{k}^{I, M_{S_I}} \middle| H_{SOC, IJ}^{M_{S_I} M_{S_J}} \middle| \chi_{k'}^{J, M_{S_J}} \right\rangle_{\mathbf{R}} \n\approx H_{SOC, IJ}^{M_{S_I} M_{S_J}} \left(\mathbf{R}_{k'}^{(c)} \right) \left\langle \chi_{k}^{I, M_{S_I}} \middle| \chi_{k'}^{J, M_{S_J}} \right\rangle_{\mathbf{R}}, \tag{9}
$$

where $\mathbf{R}_{kk}^{(c)}$ $\frac{k}{k}$ is the centroid position between TBFs *k* and k' . It is important to realize that the quality of the saddlepoint approximation is expected to be especially good for SOC matrix elements, as they are usually slowly varying with respect to the nuclear position. The IFG approximation proposes that the initial TBFs, whose initial positions and momenta are usually sampled from a Wigner distribution, are run independently, i.e., the interactions between initial TBFs are neglected. Each initial TBF however remains fully coupled with any child TBFs it will generate in the course of the nonadiabatic dynamics. For more information on these approximations, the interested reader is referred to different reviews on $\text{AIMS.}^{9,22,37,38}$ $\text{AIMS.}^{9,22,37,38}$ $\text{AIMS.}^{9,22,37,38}$ $\text{AIMS.}^{9,22,37,38}$ $\text{AIMS.}^{9,22,37,38}$ $\text{AIMS.}^{9,22,37,38}$

To summarize, GFMS is a generalization to the *in principle* exact method FMS for the description of both IC and ISC processes. The GAIMS technique, which is amenable to molecules, is obtained by applying the IFG and saddle-point approximations to GFMS.

III. TEST APPLICATIONS

The proposed AIMS extension to SOC is first tested on a model system recently proposed by Persico and co-workers.^{[13](#page-6-16)} The model comprises a singlet (S_1) and a triplet state (T_1) , which cross at $x = 10$ bohrs and both have a dissociative character (continuous curves in Figs. $2(a)-2(c)$). All SOC matrix elements between the singlet and the triplet sublevels change sign at a given position, *x*s, that can therefore be used as a parameter to tune the strength of intersystem crossing processes. For example, $x_s = 10$ bohrs leads to weak coupling between the electronic states, as the SOC is small around the point of crossing between the singlet and the triplet state (Fig. $2(a)$). In contrast, when the sign change of the SOC takes place away from the states crossing point, e.g., $x_s = 8$ bohrs, the intersystem crossing is strong as $H_{SOC, S_1T_1}^{00}$
 $H_{IOC, S_1T_2}^{00}$   and $H_{SOC, S_1T_1}^{01}$ $=$ $H_{SOC, S_1T_1}^{0-1}$ are equal to 219.5 and 155.2 cm⁻¹, respectively, at the point of crossing (Fig. [2\(c\)\)](#page-4-0). Therefore,

varying *x*^s allows testing GAIMS for different SOC strength conditions.

The GAIMS dynamics is based on 200 initial conditions, sampled from the Wigner distribution of the initial Gaussian wavefunction, as defined in Ref. [13](#page-6-16) (its corresponding probability density is represented in Fig. $2(a)$). As mentioned in Sec. [II,](#page-1-2) GAIMS uses both the IFG and the saddle-point approximation, meaning that perfect agreement with an exact solution is not expected. It is however only within these two approximations that GAIMS can be routinely applied to molecular systems, and the goal in the following is to validate the general accuracy of GAIMS with respect to exact calculations. GAIMS reproduces the exact results, obtained by solving the TDSE, 13 13 13 both qualitatively and quantitatively, within a maximum deviation of 7%, for three different cases of SOC coupling strength (right panels of Fig. [2\)](#page-4-0). Furthermore, Fig. [2](#page-4-0) shows the results for uncorrected spin-diabatic TSH $dynamics, ¹³$ $dynamics, ¹³$ $dynamics, ¹³$ with the three sublevels of the triplet state grouped into a single TSH amplitude. In this approximated formalism, TSH does not capture sign changes for the SOC and is unable to qualitatively describe ISC events, whenever the sign change occurs as the two states come close in energy. To fix this problem, a phase factor that forces a sign change in the effective SOC at the crossing point x_s can be added, leading to an excellent agreement with the exact result.^{[13](#page-6-16)} However, this simple fix is limited, since it requires *a priori* knowledge of x_s . Moreover, the contracted spin-diabatic approach to TSH is difficult to generalize for a larger number of electronic states. $\frac{13}{12}$ $\frac{13}{12}$ $\frac{13}{12}$ In contrast, such problems do not exist in the GAIMS method, and its results for this one-dimensional system highlight the accuracy of the IFG and the saddle-point approximations. Importantly, this model system also validates the naive spawning criterion $(Eq. (8))$ $(Eq. (8))$ used to spawn TBFs in

FIG. 2. GAIMS applied to a one-dimensional model system with one singlet and one triplet state. ((a)-(c)) Description of the model system—potential energy curves (black and gray lines), SOC matrix elements (dashed lines), for the case (a) $x_s = 10$ bohrs, (b) $x_s = 9.5$ bohrs, and (c) $x_s = 8$ bohrs. ((d)-(f)) Population in the singlet state for the three corresponding x_s values. GAIMS (blue) is compared with an exact solution of the time-dependent Schrödinger equation (red) and TSH in a spin-diabatic representation with no phase factor (gray), taken from Ref. [13.](#page-6-16)

a state with different spin multiplicity. All the runs leading to the results presented in Fig. [2](#page-4-0) indeed underwent only one spawning event. This behavior contrasts with that of the spin-diabatic TSH dynamics, where a large number of *hops* between states are observed. 13 As discussed in detail in the AIMS literature, $22,35$ $22,35$ improving the spawning criterion would surely result in an even better match with the exact result. We intend to investigate potential improvements in the spawning criteria for ISC in future research.

Having shown that GAIMS provides an accurate description of ISCs for different SOC strengths, we now move to an example of its real*raison d'être*, namely, the study of ISC processes in molecular systems. We choose for this purpose the nonadiabatic dynamics of thioformaldehyde, H_2CS , in its first singlet excited state. The first excited electronic state (S_1) of H_2CS has $n\pi^*$ character in the Franck-Condon region and its 0-0 transition energy has been experimentally determined at 2.033 eV.^{[39](#page-6-35)} Two triplet states lie close to S₁: T₁(n π ^{*}) and T₂($\pi\pi^*$). When the excited-state dynamics of H₂CS is initiated in S_1 , we therefore expect to observe a direct application of the El-Sayed rules^{[40](#page-6-36)}—S₁(n π^*) should be more strongly coupled to $T_2(\pi \pi^*)$ via SOC than to $T_1(n\pi^*)$ as a result of the change in orbital type. We performed GAIMS dynamics considering the first four electronic states of H_2CS (S₀, S₁, T₁, and T₂), using SA(4)-CASSCF(4/3)/6-31G* in Molpro^{[41](#page-6-37)} for 20 Wigner-sampled initial conditions. This level of theory places S_1 at 2.26 eV above the ground state in the Franck-Condon region. We present here the first 200 fs of dynamics in S_1 and will mostly comment on the GAIMS algorithm. This application is intended to provide a molecular test system for GAIMS dynamics and does not seek to obtain a complete and quantitatively accurate physical picture of the nonradiative relaxation of H2CS (which might require larger basis sets and dynamic electron correlation).

The small yet sizable population of T_2 shortly after the beginning of the excited-state dynamics confirms the El-Sayed rule (Fig. [3\)](#page-5-0), while T_1 appears to be only weakly populated in the same time window. This immediate, yet weak, population of triplet states upon photoexcitation in S_1 has also been observed for the parent molecule acetone, 42 although the population transfer is even weaker in this latter case. The total number of TBFs in the different electronic states grows quickly during the dynamics (orange line in Fig. [3\)](#page-5-0), reaching a total value of 326 TBFs, among which 306 evolve in triplet states. From an initial TBF evolving in S_1 , GAIMS rapidly starts to spawn TBFs in both T_2 and T_1 , even though a sizable amount of population is eventually only transferred to T_2 . To further analyze the dynamics, we present in Fig. [3](#page-5-0) (upper panel) a depiction of the $C=_S$ bond length and electronic population for each of the TBFs. This representation highlights the growing number of TBFs in T_1 (red) and T_2 (blue) over the course of the simulation and depicts the different dynamics they experience, evolving in electronic states of differing electronic character. Hence, the dynamics of TBFs in T_1 is closer to those in S₁—both exhibiting an $n\pi^*$ character—while the $\pi\pi^*$ character of TBFs evolving in T_2 is consistent with the longer average C=S bond length. As noted before, population transfer between singlet and triplet TBFs due to SOC is limited in the time scale of

FIG. 3. GAIMS dynamics of thioformaldehyde after photoexcitation to S_1 . Upper panel: $C = S$ bond length for all TBFs produced during GAIMS dynamics. The width of each line is proportional to the population carried by the TBF. TBFs are associated with the S_1 (light gray), T_1 (red), or T_2 (blue) electronic state. Lower panel: population of the two triplet states during GAIMS dynamics averaged over 20 initial conditions (light area indicates the standard error). The total number of TBFs is given in orange.

this simulation and the projection is mostly dominated by the TBFs evolving in S_1 .

IV. CONCLUSION

In this Communication, we presented a generalization of the full and *ab initio* multiple spawning methods to the description of internal conversion and intersystem crossing processes, both treated on an equal footing. The derivation of GFMS and GAIMS uses a spin-diabatic formalism and the implementation of GAIMS has been validated both with a model system and with a molecular application, the nonadiabatic dynamics of thioformaldehyde. This work will be followed by an extensive study of the interplay between the TBFs and the development of improved rules for ISC-triggered spawning that will minimize the number of unpopulated TBFs on triplet states. GAIMS opens the door for complete simulation of deactivation pathways in molecules and, when combined with GPU-accelerated electronic structure codes,[43–](#page-6-39)[46](#page-6-40) will be used to study the competition between internal conversion and intersystem crossing in both organic molecules and organometallic complexes.

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