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Modeling the Fluorescence of Protein-Embedded Tryptophans with ab Initio Multiconfigurational Quantum Chemistry: The Limiting Cases of Parvalbumin and Monellin

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We show that a quantum-mechanics/molecular-mechanics strategy based on ab initio (i.e., first principle) multiconfigurational perturbation theory can reproduce the spectral properties of a tryptophan residue embedded in the contrasting hydrophobic and hydrophilic environments of parvalbumin and monellin, respectively. We show that the observed absorption and emission energies can be reproduced with a less than 3 kcal mol$^{-1}$ error. The analysis of the computed emission energies based on a protein disassembly scheme and protein electrostatic potential mapping allows for a detailed understanding of the factors modulating the tryptophan emission. It is shown that for monellin, where the tryptophan is exposed to the solvent, the fluorescence wavelength is controlled not only by the distribution of the point charges of the protein—solvent environment but also by specific hydrogen bonds and, most important, by the environment-induced change in chromophore structure. In contrast, in parvalbumin, where the chromophore is embedded in the protein core, the structure and emission maxima are the same as those of an isolated 3-methylindole fluorophore. Consistently, we find that in parvalbumin the solvation does not change significantly the computed emission energy.

Introduction

The computer-aided design of unnatural proteins with specific optical properties, such as color and luminescence, represents a complex problem. In these cases, the quantum chemical method employed must be capable to describe both ground and electronically excited states of the protein chromophore. In particular, the computational description of a luminescent (e.g., fluorescent) protein implies the use of methodologies capable of predicting the excited state equilibrium structure and reactivity of fluorophores characterized, even for singlet states, by mixtures of open-shell and charge transfer characters. The ab initio (i.e., first-principle) complete-active-space self-consistent-field (CASSCF) method$^1$ is a multiconfigurational method offering maximum flexibility for an unbiased (i.e., with no empirically derived parameters and avoiding single-determinant wave functions) description of the electronic and equilibrium structure of the ground and excited states of a molecule. Furthermore, the CASSCF wave function can be readily used for subsequent multiconfigurational second-order perturbation theory$^2$ computations (CASPT2) of the dynamic correlation energy of each state ultimately allowing for a quantitative evaluation of energy gap between different electronic states.$^{3,4}$

In recent studies$^5$ we have focused on the prediction of the spectra of proteins featuring cationic or anionic chromophores. In particular, we have shown that an ab initio CASPT2//CASSCF quantum chemical procedure (equilibrium geometries and electronic energies are determined at the CASSCF and CASPT2 levels, respectively) allows, when used as part of a quantum-mechanics/molecular-mechanics (QM/MM) method, for the evaluation of the excitation energy of the retinal protonated Schiff base chromophore (treated quantum mechanically) of rhodopsin (treated using molecular mechanics force fields), of the conjugated bases of the para-hydroxy-benzylidene imidazolone fluorophore$^7$ embedded in the β-barrel scaffold of the green fluorescent protein, and of the para-hydroxy-cinnamyl chromophore of the photoactive yellow protein.$^8$ On the other hand, the simulation of proteins featuring neutral chromophores/ fluorophores has, to our knowledge, never been attempted using the same technology.

The fluorescence of tryptophan, due to its 3-methylindole (3-MIfluor) moiety, is sensitive to the molecular environment.$^9$–$^{13}$ The 3-MIfluor emission maxima fall in the 308–355 nm$^{14}$ range, with the most blue-shifted emissions associated with apolar environments. For this reason, its fluorescence spectrum and intensity have been used to gather structural information and follow protein structural changes such as during folding or denaturation.$^{15}$ Due to its spectral properties and wide distribution, tryptophan appears as an ideal candidate for high-level (i.e., “brute force”) quantum chemical investigation of a protein-embedded neutral fluorophore. Accordingly, one specific goal of the present study is to gain an atomic-level understanding of the factors controlling the fluorescence wavelength of simple proteins containing a single tryptophan residue with respect to the isolated gas-phase 3-methylindole (3-MIgas). This information can be used to rationalize/predict quantitatively the fluorescence of different proteins or protein mutants where a tryptophan probe has been inserted in certain key positions.$^{16}$

In the following, we use a CASPT2/CASSCF/AMBER/6-31G* QM/MM protocol to compute the tryptophan residue absorption ($\lambda_{\text{abs}}$) and emission ($\lambda_{\text{em}}$) maxima values of two
very different proteins. These are the F102W mutant of carp parvalbumin\(^1\) (Parv), a calcium-binding protein, and monellin\(^18\) (Mone), a sweet protein extracted from the African serendipity berry Dioscoreophyllum cumminsii. While, due to their high cost, CASPT2/CASSCF/AMBER computations cannot presently be used for screening a large set of proteins, the chosen proteins represent different limiting cases. In fact, according to their crystallographic structures, in Parv the single tryptophan residue is located in a substantially hydrophobic cavity that does not contain solvent (water) molecules (see left side of Figure 1). In contrast, in Mone, the tryptophan residue is located on the external surface of the protein in direct contact with the solvent (see right side of Figure 1). The fact that the observed value of \(\lambda_{\text{max}}\) of the two proteins differs by <25 nm (≤6.5 kcal mol\(^{-1}\) difference in excitation energy) makes them challenging candidates for modeling and analysis.

Below we show that, for both Mone and Parv, the computed \(\lambda_{\text{max}}\) and \(\lambda_{\text{fmax}}\) values reproduce the observed quantities with a blue-shifted error of <8 nm (equivalent to a ≤2.5 kcal mol\(^{-1}\) error). These results compare well with those reported for charged chromophores such as, for instance, the \(\lambda_{\text{max}}\) computed for a set of retinal proteins with a <35 nm (<3.0 kcal mol\(^{-1}\)) blue-shifted error. Notice also that the relative \(\lambda_{\text{max}}\) changes were reproduced with a <2.0 kcal mol\(^{-1}\) error. Such limited errors allow for an atomic-level analysis of the strongly red-shifted fluorescence of Parv and Mone with respect to 3-MI\(_{\text{gas}}\).

Below we demonstrate that in Parv such \(\lambda_{\text{max}}\) change arises from the stabilization of the \(1_{\text{L}}\) emitting state (with respect to the ground state) due, exclusively, to the anisotropy of the positive electrostatic potential generated by the protein. On the other hand, the larger \(\lambda_{\text{fmax}}\) change seen in Mone is due to different effects including a change in the 3-MI\(_{\text{fluor}}\) geometry (accounting for ca. 50% of the observed red-shift with respect to 3-MI\(_{\text{gas}}\)) and a positive electrostatic potential imposed, cooperatively, by the protein residues and average solvent configuration.

### Methods and Models

A full description of our QM/MM protocol and protein model building is given in the Supporting Information. Briefly, our QM method is based on a hydrogen link-atom scheme\(^16\) with the frontier placed at the C\(_{\alpha}\)C\(_{\beta}\) bond of the Trp102 and Trp3 side chains of Parv and Mone, respectively. The selected CASSCF active space comprises the full \(\pi\)-system of 3-MI\(_{\text{fluor}}\) (ten electrons in nine orbitals). The MM (we use the AMBER force field) and QM segments interact in the following ways: (i) the QM electrons and the full set of MM point charges interact via the one-electron operator, (ii) stretching, bending, and torsional potentials involving at least one MM atom are described by the MM potential, and (iii) QM and MM atom pairs separated by more than two bonds interact via standard van der Waals potentials. CASSCF/6-31G**/AMBER geometry optimization is carried out with the GAUSSIAN03\(^20\) and TINKER\(^21\) programs.

As detailed below, we define two types of protein models featuring different levels of approximation. The first (Parv-MD and Mone-MD) are fully solvated models in which the average solvent configuration is determined via molecular dynamics (MD) equilibration (see left side of Scheme 1). The second is a cruder model that lacks the solvent in Parv (Parv in vacuo) and features a 3-MI\(_{\text{fluor}}\) shielding shell of solvent molecules equilibrated in the absence of the protein (i.e., featuring the configuration of the pure solvent) in Mone (Mone \textit{shell}) (see right side of Scheme 1). Parv in vacuo and Mone \textit{shell} are analytical tools used to help to determine the factors controlling fluorescence shifts.

The Parv-MD model is based on the F102W mutant and is derived from monomer A of the crystallographic structure deposited in the Protein Data Bank (PDB) archive as file 1B8R,\(^17\) while model Mone-MD is derived from monomer A deposited in the PDB archive as file 1IV7.\(^18\) To get globally neutral models, one sodium and two chloride ions have been added to Parv and Mone, respectively. The average configuration of the solvent surrounding the proteins has been produced according to the following protocol: the PDB structures are embedded in large rectangular boxes (60 × 63 × 53 and 48 × 60 × 53 Å\(^3\)) equivalent to 3569 and 2623 solvent molecules placed within 8 Å from any given Mone and Parv atom, respectively) of TIP3P waters that were energy minimized at the MM level for 2000 steps using the steepest descent method and the AMBER force field. Then, 500 ps MD simulations of the solvent were performed using the Sander module of the Amber 7.0 package\(^22\) with the standard parameters. Both the MM energy minimization and MD simulation were carried out using periodic boundary conditions to simulate the solvent bulk. Coordinates coming from the last frame were used to build the final QM/MM models. This required a CASSCF/6-31G*/AMBER geometry optimization relaxing the coordinates of the 3MI\(_{\text{fluor}}\) (the full QM subsystem) together with those of the TIP3P water molecules and side chains within 5 Å from any given QM atoms (due to such threshold, no solvent molecules are optimized for Parv).

The optimizations have been stopped when the maximum force is <0.003 u.a./bohr and the rms is <0.0005 bohr. The equilibrium structures for the ground state (S\(_0\)) and for the first and second singlet excited states (S\(_1\), S\(_2\)) were calculated. The S\(_2\) relaxation always produces a structure very close to the crystallographic one. Notice that during the S\(_1\) and S\(_2\) optimization the side chains and solvent molecules surrounding the fluorophore may change position/orientation to adapt to the corresponding excited state charge distribution of 3-MI\(_{\text{fluor}}\). In Mone-MD, the 39 relaxed solvent molecules interact with the equilibrated fixed outer solvent configuration. The outer solvent extends 16, 11, and 16 Å away from the closest 3-MI\(_{\text{fluor}}\) atom along the \(x\), \(y\), and \(z\) directions of the solvent box (corresponding to more than four layers of water surrounding the fluorophore). In Parv, the solvent box extends 24, 14, and 17 Å away from the fluorophore located at the center of the protein matrix. During the QM/MM calculations, long-range electrostatic effects are not included beyond the solvent box boundaries. Since the 3-MI\(_{\text{fluor}}\) fluorophores of Parv-MD and Mone-MD are located in different regions of the box we cannot exclude different accuracies in the evaluation of the corresponding excitation energies. Again, notice that the coordinates of the full protein backbone and side chains located >5 Å away from the QM atoms are kept fixed at their crystallographic values (these are considered average values). Due to the excessive computational cost, no second derivative computations could be performed to rigorously determine the nature of the stationary point.

The reduced Parv in vacuo and the Mone \textit{shell} models are derived using the crystallographic structures and QM/MM optimization protocol seen above but feature a different treatment of the solvent environment. Parv in vacuo does not include the solvent at all. In Mone \textit{shell} the solvent is represented by a shell of solvent molecules set up by placing the systems in a rectangular box of TIP3P water molecules positioned within 8 Å from any given atom of the protein using the xleap module of the Amber package.\(^22\) This module provides an initial configuration of the solvent corresponding to a snapshot of MD
equilibrated waters in the absence of the solute. To determine a relaxed solvent configuration, this is then minimized for 2000 steps (including counterions) using the steepest descent method (without MD equilibration) while keeping the solute fixed. The resulting configuration provides a solute cavity that, in contrast with the Mone-MD cavity, is not adapted to the solute point charges. The final QM/MM model is constructed by discarding all solvent molecules except those forming a solvent sphere of 8 Å radius centered on the 3MI fluor chromophore. The equilibrium structures for Mone shell is then determined by CASSCF/6-31G*/AMBER geometry optimization to relax the coordinates of the QM chromophore as well as the MM residues and water molecules within 5 Å from any given atom of the 3MI fluor moiety. The positions of the remaining solvent molecules (in this model, an ca. 3 Å shield of frozen solvent molecule provides a cage keeping the solvent shell in the correct position and density), protein backbone, and distant residues were kept frozen during QM/MM calculations. We use this model to test the effect of the correct orientation of the solvent molecules surrounding the fluorophore.

In all cases, the residue charges are described by the standard AMBER force field,23 and thus the residue polarizability or/and dispersion effects are not explicitly treated. As originally pointed out by Warshel,24,25 a correct QM/MM protein model should include the solvent and account for the solvent and protein polarizability. On the other hand, the effect of the residue polarizability and dispersion on the absolute excitation energy has been shown to be limited (for br, Warshel et al.26 estimated an effect of <1500 cm⁻¹; see also the work by Ren et al.,27 Matsuura et al., and Rajamani et al.28) and shall fall into the reported error (however, notice that cancellation effects cannot be excluded). Due to cancellation effects, the error on excitation energy changes will be even smaller.

In the context of the present work, the prepared Mone-MD and Parv-MD structures are assumed to provide acceptable representation of the average environment of the chromophore/fluorophore (i.e., given the high >40% content of solvent in the crystals—see the PDB files—we assume that the crystallographic structures provide a suitable protein average structure in solution, and possible large-amplitude low-frequency fluctuations affecting the average structure are assumed to have a small effect on the observed \( \lambda_{\text{max}} \) values). Furthermore, within our fixed charge model, the excitation energy should be mainly determined from the residues belonging to the chromophore cavity. This assumption has been assessed29 by evaluating the \( \lambda_{\text{max}} \) of rhodopsin with a full protein model and with a reduced protein model consisting of the 27 residues surrounding the retinal chromophore and two crystallographic waters. It is shown that the two \( \lambda_{\text{max}} \) values differ by <1 kcal mol⁻¹.

For all protein models, at the S₀ equilibrium geometries a CASPT2 computation is carried out, using the MOLCAS-6 program30,31 to evaluate the vertical excitation energy for the \((S_0 \rightarrow S_1 \text{ and } S_0 \rightarrow S_2)\) transitions (assumed to match the energy of the corresponding \( \lambda_{\text{max}} \) values) and the associated oscillator strength. The emission \( \lambda_{\text{max}} \) values from the first and second excited states (\(S_1 \rightarrow S_0 \text{ and } S_2 \rightarrow S_0\) transitions) are evaluated by computing the \(S_1 \rightarrow S_0\) and \(S_2 \rightarrow S_0\) vertical energy gaps via CASPT2 computations using a 0th-order three-root state average CASSCF wave function at the \(S_1\) and \(S_2\) equilibrium geometry, respectively.

Again, notice that while the AMBER charges account for \(S_0\) polarization effects in a mean-field way23 no polarizable residue is included in the protein model. The same charges are used for the excited state computations without introducing an ad hoc dielectric constant. Also, notice that, in this work, there is no scaling of the computed CASPT2/CASSCF/AMBER excitation energies.

Results and Discussion

The changes in the tryptophan fluorescence as a function of the protein structure are a manifestation of the diversity of protein environments.15 Since the stiff bicyclic framework of the 3-MI fluor structure does not allow for significant deconjugation of its π-system, the observed spectral change is expected to originate mainly from a change in the electrostatic potential acting on the fluorophore centers. In other words, as pointed out by Callis and co-workers,14 the \( \lambda_{\text{max}} \) value and intensity must depend on the “electrostatic potential landscape”. Most important, also the fluorescence lifetime and its decay dynamics are affected by such a landscape.13 In this context, the possibility to employ ab initio (i.e., unbiased) multiconfigurational quantum chemistry opens up new perspectives. Indeed, in the past, CASPT2/CASSCF photochemical reaction path computations32 have shown that (nonadiabatic) hydrogen and charge (electron) transfer mediated by conical intersections provide the main channel of lifetime modulation (e.g., through fluorescence quenching) of synthetic fluorophores such as azaalkanes and ketones.34–36 Thus, the development of CASPT2/CASSCF/AMBER protocols for the evaluation of excited state reaction paths6 and trajectories37 should allow for the extension of such studies to protein fluorophores.

As a preliminary step toward the investigation of the tryptophan fluorescence and its decay dynamics in proteins with
multiconfigurational quantum chemistry tools, here we focus on the origin of the $\lambda_{\text{max}}$ value. For other photoactive proteins, four different molecular factors have been proposed to play a role at the molecular level: (i) the increased or decreased oscillator strengths can be compared with the variation in the observed extinction coefficients (10 $\text{M}^{-1} \text{cm}^{-1}$) energies, computed at the corresponding CASSCF/6-31G* optimized geometries, are given in italics. Computed spectral parameter data for the parent gas-phase indole (not reported below) using the same protocol and the (C,N[3s2p1d]/H[2s]) ANO-S basis are reported in ref 38.

In Table 1, we report the excitation energies, evaluated at the ground state ($S_0$) equilibrium structure, and the emission energies, evaluated at the $1L_1$ equilibrium structure, for the Parv-MD and Mone-MD models together with those of the reference system 3-MI$_{\text{gas}}$. In their $S_0$ equilibrium geometry, the proteins display a slightly lower $1L_0$ state (see the Supporting Information). In contrast, in the $L_1$ excited state geometry, $S_1$ always corresponds to the $1L_1$ state, thus displaying charge transfer character (i.e., a larger oscillator strength and dipole moment change). Such $1L_0$/1$L_1$ inversion does not occur in 3-MI$_{\text{gas}}$ where the covalent $1L_0$ state is always lower (12 kcal mol$^{-1}$ at GS-3-MI$_{\text{gas}}$). The computed emission energies reproduce the observed values with systematically blue-shifted errors of $\pm 2.5$ kcal mol$^{-1}$ (see Figure 2A). The previous (semiempirical) INDO/S-CIS based study by Callis predicted, for Mone and Parv, red-shifted $\lambda_{\text{max}}$ values (with respect to the available experimental data). On the basis of these data, it was proposed that, in Mone, the $\lambda_{\text{max}}$ value is controlled by the polarization of the indole ring by the electric fields of both the solvent (water) and protein charges, while in Parv, the $\lambda_{\text{max}}$ is mainly controlled by the geometrical polarization of the external water molecules. However, a quantitative analysis of factors i–iv has never been reported.

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adapted to the solvent-exposed 3-MI$_\text{fluor}$ and surrounding Mone charge distribution. In conclusion, the effect of solvent is, as expected, limited in Parv where the fluorophore is embedded in a hydrophobic cavity but large (e.g., showing a 3-fold error increase with respect to the observed $\lambda_{\text{fmax}}$ value) in Mone where the fluorophore is placed at the surface of the protein and in direct contact with the solvent. In the next section, we will provide a rationalization for these effects on the basis of the electrostatic potential acting on the 3-MI$_\text{fluor}$ centers.

The error coming from the use of the 6-31G* basis with a correlated CASPT2/CASSCF wave function has been investigated by comparing the excitation energies computed for 3-MI$_\text{gas}$ with ANO-S data from the literature (see Table 1). We also recomputed the $\lambda_{\text{fmax}}$ of the full models Mone-MD and Parv-MD using the same ANO-S (C,N[4s3p1d]/H[2s]) basis set with respect to the 6-31G* basis set. This yields a smaller blue-shifted error (see Figure 2A and the Supporting Information) indicating that, as expected, a better basis increases the computational accuracy. However, since we focus on $\lambda_{\text{fmax}}$ changes and to avoid excessive computational costs, below we focus on the 6-31G* results.

**Effect of the Fluorophore Environment.** The limited, relative error in excitation energies computed for Mone-MD, Parv-MD, and 3-MI$_\text{gas}$ prompts for an analysis of the factors determining the $\lambda_{\text{fmax}}$ values. Accordingly, the emission energy of each model is compared with that of the isolated fluorophore taken with its protein-optimized geometry (Figure 2B) and with that of protein deprived of the solvent (Figure 2C).

Inspection of the data in Figure 2B shows that in Parv the change in 3-MI$_\text{fluor}$ geometry cannot be responsible for the observed red-shifted emission. In fact, the corresponding 3-MI$_\text{fluor}$ has an excitation energy that is only slightly higher than the reference quantity (i.e., 3-MI$_\text{gas}$). This is obviously due to a limited protein-induced change in fluorophore excited state geometry with respect to the gas phase. Such a conclusion is confirmed by the data in Figure 3A pointing to a $<0.015$ Å difference between the geometrical parameters of 3-MI$_\text{gas}$ and of the Parv-MD fluorophore in both the ground and excited states. Furthermore, the same quantity is close to the excitation energy computed for the fluorophore of Parv in vacuo (see Figure 2B) demonstrating that the solvent has a small effect on the 3-MI$_\text{fluor}$ structure (see Figure 3A). This conclusion is further supported by the data in Figure 2C where we show that removal of the solvent from Parv-MD does not significantly change the excitation energy (i.e., this value is very close to the excitation energy of Parv in vacuo seen in Figure 2A).

In contrast to Parv, the data of Figure 2B show that, in Mone, the isolated fluorophore must undergo a large structural change. In fact, the corresponding excitation energy is significantly lower with respect to 3-MI$_\text{gas}$. Again, this is consistent with the structure of Figure 3A that shows large deviations of the ground state and excited state 3-MI$_\text{fluor}$ structures from the 3-MI$_\text{gas}$ reference. Comparison of the Mone-MD excitation energies in Figure 2A and 2B reveals that a decrease $>50\%$ of the excitation energy with respect to 3-MI$_\text{gas}$ is due to a change in the fluorophore structure. This change is very sensitive to the structure of the solvent shell surrounding 3-MI$_\text{fluor}$ (or, in other words, to the solvent averaged configuration model). Indeed, the excitation energies of the Mone-MD and Mone shell fluorophores in Figure 2B are very different reflecting the limited change in the structure of the Mone shell fluorophore with respect to 3-MI$_\text{gas}$ (see Figure 3A). A correct solvent shell configuration also induces the structural change that leads to an enhanced dipole moment change and charge separation (see the Supporting Information). Comparison of the Mone data in Figure 2A and 2C indicates that the point charges both of the protein and of the solvent contribute, concurrently, to further decrease the excitation energy of the Mone fluorophore. In conclusion, the contributions of factor i and iii are not critical for Parv but are both important when the fluorophore is in contact with the solvent (as in Mone). The comparison between the Mone-MD and Mone shell data in Figure 2B indicates that the structure change of 3-MI$_\text{fluor}$ is due to factor iii.

A common basis for the discussion of the effects of the protein residues and solvent molecules is provided by the charge transfer nature of the spectroscopic state of tryptophan (see Figure 3B for the case of 3-MI$_\text{gas}$). In fact, in 3-MI$_\text{gas}$ $38\%$ of negative charge ($\pi$-electron density), originally located on the pyrrole moiety, is shifted toward the benzene ring upon the $S_0\rightarrow 1L_a$ transition. Thus, an electrostatic potential stabilizing the positive charge on the $1L_a$ pyrrole moiety or stabilizing the negative charge on the $1L_a$ benzene moiety will result in a decreased $1L_a\rightarrow S_0$ emission energy. Notice that while the $1L_a$ equilibrium structure of the chromophore remains planar an extensive bond length rearrangement occurs to accommodate the charge transfer (see Figure 3A). As already mentioned above, both the entity of the charge transfer and the bond readjustment pattern are different in the systems investigated here. The Parv fluorophore shows changes very close to those of 3-MI$_\text{gas}$, while as discussed above, in Mone these changes are different.

According to Figure 4A, the anisotropic electrostatic potential (generated from the residue and solvent point charges) acting on the chromophore centers of Parv-MD and Mone-MD is
positive and localized on the benzene ring. This potential must thus decrease the $S_1 - S_0$ energy gap relative to the isolated chromophore consistently with the computed $\lambda_{\text{max}}$ red-shift. In Parv, the electrostatic potential is due exclusively to the protein charges.

Comparison of the Mone-MD and Mone shell electrostatic potential in Figure 4A shows that this is very sensitive to the configuration of the solvent shell surrounding 3-MI fluoro. If this configuration is not adapted to the protein and fluorophore charges (i.e., does not correctly represent the average configuration of the solvent), the potential acting on the QM atoms is negative and not positive. Consistently, with the excitation configuration of the solvent shell surrounding 3-MI fluoro. If this configuration is not adapted to the protein and fluorophore charges and solvent molecules in the close vicinity of the fluorophore are responsible for the described solvent effect and not the bulk.

The electrostatic potential acting on the centers of the Mone fluorophores in the absence of the Lys44 charges (Mone-Lys44) is clearly less positive on the pyrrole moiety and becomes very negative on the pyrrole moiety (see Figure 4B). This unveils a counterbalancing effect of Lys44 residue charges and solvent shell. To explain this finding (i.e., a negative potential induced by the removal of the positive Lys44 charge) one must focus on the fact that in Mone the 3-MI fluoro moiety and solvent shell are adapted to the protein environments. Since the water molecules have a large dipole moment their average orientation will be determined by the interaction with the protein partial or fully charged groups. Thus, in Mone, the positively charged Lys44 residue, which, with respect to 3-MI fluoro, is located opposite to the solvent (see the right structure in Figure 5A), will presumably orient the water molecules of the first solvation shell (also through enhanced polarization of the fluorophore π-system) in such a way to point their negatively charged oxygens toward 3-MI fluoro. This specific orientation leads to a negative potential (with the largest intensity on the positively charged pyrrole moiety). The $\lambda_{\text{max}} - S_0$ gap of the Mone-MD-Shell model with respect to Mone-MD indicates that only the solvent molecules in the close vicinity of the fluorophore are responsible for the described solvent effect and not the bulk.

In Figure 5B we report the results of an analysis of the effect of selected residues and solvent molecules on the emission maximum of the 3-MI fluoro of Mone. It is apparent that the Lys44 residue has a red-shifting effect similar to that induced by the full protein. It is also apparent that both the Lys44 residue alone and a specific hydrogen bonded water molecule (linked to the N-H bond of the pyrrole moiety) alone have similar red-shifting effects on 3-MI fluoro. It is interesting to see that when both effects are present one still gets a similar red-shift pointing to a different
The idea that the residues (or solvent molecules) in direct contact or close to the chromophore/fluorophore control the optical properties has also been investigated for the visual photoreceptor rhodopsin.5 CASPT2//CASSCF/AMBER excitation energy computations demonstrate that only the residues in the chromophore cavity affect significantly the excitation energy and therefore the absorption wavelength.29 In our Parv model, the electrostatic potential acting on the fluorophore is again positive (see Figure 4B). Therefore, the effect of the hydrogen-bonded Val43 residue must be such to increase the positive potential at the level of the benzene moiety or decrease the positive potential at the level of the indole moiety. As shown in Figure 4B, removal of Val43 leads to a more negative potential on the N–H region of indole. Thus, it is the latter effect to occur in Parv.

Conclusions

The work of Callis and co-workers14 has established that a QM/MM protocol based on semiempirical quantum chemical methods and a suitable fluorophore charge scaling can successfully reproduce the fluorescence maxima of 19 tryptophan-containing proteins with a ≈15 nm error. In that study, all protein—solvent systems (with a fixed 1La reference geometry for the fluorophore moiety) were modeled via a 30 ps trajectory and the emission maxima determined by averaging the emission energies of many snapshots. While such a protocol cannot be used with state-of-the-art ab initio QM methods, in the past we have provided evidence that both absorption and emission can be simulated using the CASPT2//CASSCF/AMBER protocol. In particular, using a crystallographic structure and a suitably prepared solvent box to model the average protein—solvent environment,6,7 it has been shown that the absorption or fluorescence of anionic and cationic biological chromophores/fluorophores can be reproduced within <5 kcal mol−1.

Above, the CASPT2//CASSCF/AMBER protocol has been used to study the spectroscopy of two very different protein-embedded tryptophans as examples of neutral biological fluorophores. As shown in Figure 2, and in spite of the absence of empirical parameters, the ab initio CASPT2//CASSCF level allows us to reproduce the observed changes in λf max value (i.e., with respect to 3-MI) with errors smaller than those obtained using not scaled semiempirical methods. Most important, the error is of the same magnitude as that found in proteins containing charged chromophores. Therefore, our study extends the validity of the protocol and paves the way to first-principle simulations of the photophysics of protein-embedded fluorescent probes.

We believe that the results presented above will have an impact on future studies of the fluorescence lifetime and decay dynamics of tryptophan. Indeed, previous mechanistic studies of the fluorescence decay dynamics of synthetic fluorophores demonstrate that the decay may occur via aborted chemical or electron transfer reactions mediated by conical intersection channels. This fact calls for the use of multiconfigurational quantum chemical theories (e.g., CASSCF) where real (non-avoided) crossings between potential energy surfaces of the same spin multiplicity are properly represented.34–36

Consistently with previous work, we have confirmed that in Parv the environmental effects on the λf max value are mainly due to stabilization of the emitting charge-transfer state. On the other hand, it has been possible to unveil that, in contrast to semiempirical studies,14 the ab initio CASPT2//CASSCF treatment points to a minor effect of the external solvent on the λf max change when a fluorophore is embedded in a substantially
hydrophobic protein matrix. In contrast, we have shown that when the fluorophore is in contact with the solvent (e.g., in Mone) the solvent contribution to the $\lambda_{\text{max}}$ change is very important and also affects the fluorophore structure.

The red-shift values induced by the protein and solvent changes are summarized in Figure 6. For Parv, the computed CASPT2//CASSCF/AMBER and “scaled-charge” INDO/S-CIS/CHARMM emission shifts have very different magnitudes. In fact, the semiempirical data point to a large and dominating effect of the solvent surrounding the protein (i.e., not in contact with 3-MI$_{\text{fluo}}$). In contrast, the ab initio data point to a dominating effect of the protein (consistently with the red-shift value of the Parv in vacuo unsolvated model). On the other hand, for Mone, the ratio between the solvent and protein effects is more balanced. However, in this case the INDO/S-CIS/CHARMM emission points to a protein dominating effect, while the CASPT2//CASSCF/AMBER protocol yields a solvent effect that is larger than the protein effect. Of course, as shown in the figure (see full model bars), these effects are far from being additive as the electrostatic field imposed by the solvent is modified by that of the protein, and this depends on the protein and solvent spatial locations with respect to the fluorophore.

At least three of the specific $i$–$iv$ factors mentioned above have been found to affect the fluorescence color in Mone and Parv. The first mechanism corresponds to factor i. In fact, in Mone, the bond-length pattern of 3-MI$_{\text{fluo}}$ is very different from that of the reference gas-phase fluorophore. It seems that the change is induced by the solvent shell structure. A second important factor is factor ii. This is related to the effect of specific and strategically located protein cavity residues. In Mone, the Lys44 residue with its positive charge placed above the benzene moiety and in Parv the Val43 residue that forms hydrogen bonds with the pyrrole moiety of the fluorophore constitute clear examples. The impact of specific interactions is estimated of the order of 2 kcal mol$^{-1}$ (see Figure 5B). Of course, cooperative effects in the same category (i.e., contributions coming from many fractional changes) may have a considerable weight. The third mechanism is related to the direct electrostatic interaction with the solvent and corresponds to factor iii. It is found that this factor is coupled with factor i and ii. In fact, in Mone both the fluorophore $\pi$-system polarization and the geometrical reorganization of the first solvent shell seem to account for part of the red-shifting effect. On the other hand, interactions with specific solvent molecules (e.g., a single water molecule in Mone) account for the emission energy decrease.

In the near future, the development of accurate tools for the simulation of the changes in tryptophan fluorescence and fluorescence lifetimes as a function of the residue position in a peptide backbone may constitute one important tool for the design of unnatural proteins with wanted properties. These tools will have to be potent enough to be able to describe the mechanism of internal residue or solvent mediated quenching. In particular, they should be able to deal with large and rapid changes in the structure and electronic wave function that often characterize photoinduced abortive photochemical or electron transfer reactions. Presently, it is accepted that these processes imply the evolution toward a conical intersection funnel connecting the emitting excited state to the ground state or to a lower nonemitting excited state (e.g., conical intersections are, by definition, regions of rapid change in electronic structure). The investigation of these processes calls for unbiased quantum chemical methods featuring highly flexible wave functions and for the tools of the emerging field of computational photochemistry. Above we have provided evidence that, for two very different proteins, the use of QM/MM strategy based on multiconfigurational second-order perturbation theory can potentially give access to a quantitative mapping of the excited state reaction paths and, in turn, resolve the atomic-level mechanism controlling the fluorescence quenching/decay. Recent work has established that the evaluation of realistic time scales (including the biexponential character of such processes) via scaled-CASSCF/AMBER trajectories is becoming a reality. Given the steady increase in computer performances, the continuation of such exploratory research effort, in our and other laboratories, appears to open new perspectives for the future of de novo light-sensitive protein design.

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**Supporting Information Available:** The QM/MM scheme, models, solutions models, tables, and optimized structures. This material is available free of charge via the Internet at http://pubs.acs.org.

**References and Notes**
