Bachelor’s Thesis

Steering excited state dynamics of biological chromophores by external electric fields.

prepared by

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Thesis period: 1st May 2011 until 1st September 2011

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# Nomenclature

## Greek & Latin Letters

<table>
<thead>
<tr>
<th>Variable</th>
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<tbody>
<tr>
<td>$\alpha$</td>
<td>Single Bond Torsion Angle</td>
</tr>
<tr>
<td>$\beta$</td>
<td>Double Bond Torsion Angle</td>
</tr>
<tr>
<td>$S_0$</td>
<td>Ground State Potential Energy Surface</td>
</tr>
<tr>
<td>$S_1$</td>
<td>First Excited State Potential Energy Surface</td>
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</table>

## Abbreviation

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Meaning</th>
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<tbody>
<tr>
<td>TISE</td>
<td>Time-Independent Schrödinger Equation</td>
</tr>
<tr>
<td>PES</td>
<td>Potential Energy Surface</td>
</tr>
<tr>
<td>HF</td>
<td>Hartree-Fock</td>
</tr>
<tr>
<td>RHF</td>
<td>Restricted Hartree-Fock</td>
</tr>
<tr>
<td>CI</td>
<td>Configuration Interaction</td>
</tr>
<tr>
<td>CSF</td>
<td>Configuration state function</td>
</tr>
<tr>
<td>MO</td>
<td>Molecular Orbital</td>
</tr>
<tr>
<td>CASSCF</td>
<td>Complete Active Space Self-Consistent Field</td>
</tr>
<tr>
<td>pCA</td>
<td>Para-Couramic Acid</td>
</tr>
<tr>
<td>PYP</td>
<td>Photoactive Yellow Protein</td>
</tr>
<tr>
<td>pCK$^-$</td>
<td>Para-Couramic Ketone</td>
</tr>
<tr>
<td>SB</td>
<td>Single Bond</td>
</tr>
<tr>
<td>DB</td>
<td>Double Bond</td>
</tr>
<tr>
<td>SB$_{\text{min}}$</td>
<td>Single Bond Twisted Minimum Structure</td>
</tr>
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</table>
### Nomenclature

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>DB&lt;sub&gt;min&lt;/sub&gt;</td>
<td>Double Bond Twisted Minimum Structure</td>
</tr>
<tr>
<td>EOM-CCSD</td>
<td>Equation of Motion Coupled Cluster for Single and Double Excitations</td>
</tr>
<tr>
<td>QM/MM</td>
<td>Quantum Mechanics / Molecular Mechanics</td>
</tr>
<tr>
<td>FC</td>
<td>Franck-Condon</td>
</tr>
<tr>
<td>HOMO</td>
<td>Highest Occupied Molecular Orbital</td>
</tr>
<tr>
<td>LUMO</td>
<td>Lowest Unoccupied Molecular Orbital</td>
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Introduction

Photochemical processes play a key role in biological systems. In such processes, photons are absorbed by biological molecules and trigger chemical reactions. Photoisomerization, a type of photochemical process which produces an isomer of the original molecule, plays a crucial role in many biological systems. In photoisomerization, the energy landscape of the excited molecule changes when a photon is absorbed, causing the molecule to rearrange into an isomer. Several different applications of photoisomerization are observed in nature. In the retinal chromophore, which is part of the photoreceptors in most vertebrates, photoisomerization is responsible for the initiation of visual phototransduction [23]. In Halobacterium salinarium the same chromophore functions as a molecular engine which pumps protons through a cell membrane, [2]. In both of these cases, the initial reaction is triggered by ultra-fast double bond isomerization on time scales less than a picosecond. This type of reaction is not only ultra-fast but is also very efficient and therefore offers great potential for many applications. Ultra-fast photoisomerization is connected to a conical intersection where the potential energy surfaces of the ground and excited state intersect, leading to a fast and radiationless decay of the excited molecule back to the ground state. For a deeper insight into protein functions it is crucial to understand and even control the photoisomerization process. A possible control
1. Introduction

mechanism is already provided in the protein environment. Strong, local electric fields are established by hydrogen bonds or charged side chains. In the following thesis, a method for steering this photoisomerization processes via external electric fields is presented. Electric fields are relatively easy to apply on small scales, and the current feasible permanent field strengths are sufficient to affect the energy levels of molecules with more than a few electrons.

As a proof of concept, one of the simplest biological systems, an analogue of the photoactive yellow protein (PYP) chromophore, has been selected for numerically study of photoisomerization processes with high accuracy. Time-resolved simulations can be accomplished with reasonable effort, so that conical intersections can be observed. The investigated molecule is treated in vacuo, not only for computational efficiency but also to neglect further non-field-dependent influences. The reaction pathway is investigated to determine whether the pathway is sensitive to the presence of electric fields. Following a short theoretical introduction on quantum chemistry methods, the biological origin, structure and charge distribution of the selected chromophore will be discussed. Subsequently, the topologies of the ground and first excited state under the influence of electric fields are examined. Finally, time-resolved simulations are performed that qualitatively validate the topologies and allow prediction of reaction rates and absorption spectra which can be experimentally verified.
2.1. Methods of Quantum Chemistry

2.1.1. Born-Oppenheimer Approximation

The first step for modeling the quantum mechanical properties of molecules is to derive the time-independent Schrödinger equation (TISE) and its Hamiltonian $\mathbf{H}$, which is given as the sum over the kinetic $\mathbf{T}$ and potential energy $\mathbf{V}$ for each particle. The potential energy terms will only include the Coulomb interaction between all particles of the molecule if relativistic effects are neglected (which is reasonable as biomolecules are generally composed of first and second row elements). Thus, the TISE and its Hamiltonian can be written as

$$H \psi(r_e, R_N) = E \psi(r_e, R_N)$$ (2.1)

$$H = T_N + T_e + V_{NN} + V_{eN} + V_{ee}$$ (2.2)

where $E$ is the total energy of the molecule and $\psi$ an eigenfunction. The subscript $N$ belongs to the nuclei and $e$ to the electrons whereas the nucleus-nucleus dependency is represented by $NN$, the electron-nucleus one by $eN$ and the electron-electron
2. Theory

dependency by $ee$. The wave function $\psi$ depends on the nuclear $R_N$ and electronic positions $r_e$. In general, there does not exist an analytical solution for the TISE (Eq. 2.1) and even numerical calculations are computationally intensive. J. Robert Oppenheimer and M. Born introduced an approximation in 1927 to simplify the TISE which allows the TISE to be separated into an electronic and a nuclear equation and solved separately.

The electronic TISE

$$
(T_e + V_{eN} + V_{ee} + V_{NN})\psi_e(r_e; R_N) = U(R_N)\psi_e(r_e; R_N)
$$

needs to be solved where $U(R_N)$ represents the electronic eigenvalue. $\psi_e$ is a function of $r_e$ and relies only parametrically on the nuclear positions $R_N$. In other words, the nuclear positions are fixed. This assumption is physically because the mass of an electron is orders of magnitude smaller than the mass of the nuclei ($m_e << m_n$). Hence, the velocities of the electrons are much higher than that of the nuclei, and therefore the electronic movement can be considered to adjust instantly to the nuclear movement.

$U(R_N)$ forms an potential energy surface (PES) for the nuclei (see Eq. 2.3), and thus the nuclear TISE is

$$
(T_N + U)\psi(R_N) = E\psi(R_N).
$$

As a consequence of the Born-Oppenheimer approximation different electronic states create different but well-defined potential energy surfaces that cannot intersect. This consequence of the approximation does not always reflect reality and can lead to errors, see Section 2.2.

### 2.1.2. Time Independent Hartree-Fock Theory

Even after applying the Born-Oppenheimer Approximation the electronic TISE cannot be solved analytically, and therefore numerical approximation methods are required. The basis for many numerical approximation methods with different depth of theory is given by the Hartree-Fock method. As mentioned in Section 2.1.1, the electronic wave function and energy depend only parametrically on the nuclear
coordinates \( \mathbf{R}_N \). Hence, the Hamiltonian of Eq. 2.3 follows as:

\[
\mathbf{H}_e = -\frac{\hbar}{2m_e} \sum_{i=1}^{n} \nabla_i^2 + \sum_{\alpha=1}^{N} \sum_{i=1}^{n} \frac{Z_{\alpha}e^2}{r_{\alpha i}} + \sum_{j=i+1}^{n} \frac{e^2}{r_{ij}} + \sum_{\alpha=1}^{N-1} \sum_{\beta=\alpha+1}^{N} \frac{Z_{\alpha}Z_{\beta}e^2}{r_{\alpha\beta}}
\]

(2.5)

where \( i \) and \( j \) are the electron indices and \( \alpha \) and \( \beta \) the nuclear ones. Furthermore, \( e' = \frac{1}{4\pi\varepsilon_0} \) is the prefactor of the Coulomb potential. \( r_{ij} = |\mathbf{r}_i - \mathbf{r}_j| \), for instance, is the distance between two particles. Thus, Eq. 2.5 represents a n-body problem, which makes it impossible to solve analytically. The task is to find the eigenfunctions \( |\psi\rangle \) and the energy \( E \) corresponding to the Schrödinger equation (Eq. 2.3). In Hartree-Fock (HF) theory, the total wave function is expressed through a single Slater determinant which is an antisymmetrized product approach of single electron wave functions and therefore obeys the Pauli principle. Together with the mean field interaction of the electrons, the n-body problem splits up into n coupled 1-body problems. The total wave function follows as

\[
|\psi_{SL}\rangle = \frac{1}{\sqrt{N}} \left| \begin{array}{cccc}
\phi_1(x_1) & \phi_2(x_1) & \ldots & \phi_N(x_1) \\
\phi_1(x_2) & \phi_2(x_2) & \ldots & \phi_N(x_2) \\
\vdots & \vdots & \ddots & \vdots \\
\phi_1(x_N) & \phi_2(x_N) & \ldots & \phi_N(x_N)
\end{array} \right|
\]

(2.6)

where \( (x_N) \) represents the spin and spacial coordinates \( x = \{\mathbf{r}, \omega\} \) of the n-th electron and \( \phi_N \) the n-th spin-orbital wave function which is the product of the spacial and spin wave function.

After applying \( \mathbf{H}_e \) to \( |\psi_{SL}\rangle \) the eigenvalues \( E \) become:

\[
E = \sum_{i=1}^{n} h_i + \frac{1}{2} \sum_{i,j} (C_{ij} - A_{ij}) \quad \text{with,}
\]

(2.7)

\[
C_{ij} = \int \frac{\phi_i^*(\mathbf{x}_i)\phi_j^*(\mathbf{x}_j)e^2}{|\mathbf{r}_i - \mathbf{r}_j|} \phi_i(\mathbf{x}_i)\phi_j(\mathbf{x}_j)d\mathbf{x}_id\mathbf{x}_j
\]

(2.8)

\[
A_{ij} = \int \frac{\phi_i^*(\mathbf{x}_i)\phi_j^*(\mathbf{x}_j)e^2}{|\mathbf{r}_i - \mathbf{r}_j|} \phi_j(\mathbf{x}_i)\phi_i(\mathbf{x}_j)d\mathbf{x}_id\mathbf{x}_j
\]

(2.9)

here \( h_i \) represents the \( i \)-th one electron energy, \( C_{ij} \) the Coulomb integral and \( A_{ij} \) the exchange integral. For a detailed derivation see [17]. The classical analogue of \( C_{ij} \).
is the repulsion energy between one electron occupying spin-orbital $\phi_i$ and another electron occupying spin-orbital $\phi_j$. The exchange integral on the other hand has no classical analogue. To get as close as possible to the real electronic ground state energy, the derived energy has to be minimized. For this purpose, the variational principle is used with the orthonormalization constraint ($\langle \phi_i | \phi_j \rangle = \delta_{ij}$) of the wave function.

\[ \delta L = \delta E - \sum_{i,j} \lambda_{ij} (\langle \delta \phi_i | \phi_j \rangle - \langle \phi_i | \delta \phi_j \rangle) = 0 \]  

(2.10)

At this point, it is reasonable to define the Fock-operator $\mathbf{F}$ which takes $\delta E$ to the following form:

\[ \mathbf{F}_i = \mathbf{h}_i + \sum_j (\mathbf{C}_j - \mathbf{A}_j) \]  

⇒ $\delta E = \sum_i (\langle \delta \phi_i | \mathbf{F}_i | \phi_i \rangle + \langle \phi_i | \mathbf{F}_i | \delta \phi_i \rangle)$  

(2.12)

Inserting Eq. 2.12 into Eq. 2.10 the Hartree-Fock equation can be written as:

\[ \mathbf{F}_i \phi_i = \sum_j \lambda_{ij} \phi_j \]  

(2.13)

Subsequently, the last equation can be diagonalized, leading to an eigenvalue equation whose eigenvalues represent the canonical molecular orbital energies $\varepsilon_i$.

\[ \mathbf{F}_i \phi_i' = \varepsilon_i \phi_i' \]  

(2.14)

Hence, the ground state energy is given as follows:

\[ E = \sum_i \varepsilon_i - \frac{1}{2} \sum_{i,j} (\mathbf{C}_{ij} - \mathbf{A}_{ij}) \]  

(2.15)

The idea is to improve the $\phi_i'$ iteratively, by improving the electron densities in $\mathbf{C}$ and $\mathbf{A}$ until a convergence criterion is reached. Nevertheless, the HF energy is still an upper bound for the real ground state energy. Summing up HF theory, there are two important aspects to keep in mind:

- The wave function is expressed through a single Slater determinant which fulfills the Pauli principle.
- The electrons only notice the presence of each other via a mean field which is created by their charge density.
The latter is a drawback in HF theory because electron-electron correlation effects are neglected.

The question of the orbital initial conditions remains. Fortunately, every eigenfunction of a Hermitian operator can be expanded after a complete orthonormal basis where \( a_i \) are the molecular orbital (MO) coefficients and \(|\chi_i\rangle\) are the atomic orbitals.

\[
|\phi\rangle = \sum_{i=0}^{n} a_i |\chi_i\rangle 
\]  

(2.16)

For numerical calculations, only a small number of elements of this expansion can be taken into account which are summarized in a basis set. Thus, the selection of the basis set has a serious influence of the accuracy of the HF energies as well as the molecular orbitals.

### 2.1.3. Configuration Interaction Method

As previously pointed out, HF theory fails when electron-electron correlation cannot be neglected. Unfortunately, an accurate prediction of binding energies and energy barriers, which is very important for this work, requires a more advanced level of quantum theory.

Over the past 50 years many different approximations for improving HF theory have been developed. One often used advanced approximation method is the Configuration Interaction (CI) method.

Due to mathematical complexity, only an outline of the CI is presented. For a more detailed derivation see [17].

The starting point of a CI calculation is MOs that have been either arbitrary chosen or calculated with a lower level of theory, which are composed of occupied and virtual (unoccupied) orbitals. In practice, Hartree-Fock MOs \( (\Phi'_i) \) are often used as the starting point of a CI calculation. The next step is to form configuration state functions (CSF) \( \Phi_i \) by either a single Slater determinant or a linear combination of Slater determinants which include the \( \Phi'_i \). The trial wave function \( \psi \) can be expressed through a linear combination of the configuration state function (CSF).

\[
\psi = b_0 \Phi_0 + \sum_S b_S \Phi_S + \sum_D b_D \Phi_D + \sum_T b_T \Phi_T + \ldots = \sum_i b_i \Phi_i 
\]

(2.17)

Here 0 indicates no, \( S \) single, \( D \) double, and \( T \) triple excited Slater determinants relative to the starting configuration. The excited Slater determinants are created
2. Theory

by removing electrons of occupied orbitals, and placing them into virtual ones. The CI coefficients \( b_i \) are determined by the variational principle, similar to Section 2.1.2. Thus, not only the electronic correlation between already occupied orbitals but also between virtual ones are taken into account.

If the molecule has more than a few electrons, it will be impractical to consider all orbitals to evaluate \( \psi \) because the MO and CI coefficients have to be optimized independently. The computational effort required is infeasible, even for supercomputers. The number of CSFs (\(#_{\text{CSF}}\) ), which has to be optimized in the CI method, is given by

\[
#_{\text{CSF}} = \frac{c!(c + 1)!}{(\frac{1}{2}n)!(\frac{1}{2}n + 1)!(c - \frac{1}{2})!(c - \frac{1}{2}n + 1)!} \tag{2.18}
\]

where only singlet states are allowed (\(S = 0\)). \( c \) is the number of one electron basis functions and \( n \) the number of electrons. Table 2.1 shows the number of CSFs for the CH\(_3\)OH (\(n = 18\)) which demonstrates that even for comparatively small molecules an extensive basis set induces a large computational effort. In fact, with a small basis set the full CI is achievable at least for a few electrons but this does not guarantee an accurate solution. For high accuracy calculations a larger basis set is required, and therefore only a small number of orbitals can be included. The included orbitals are consequently chosen such that they span the active space.

Table 2.1.: Number (\#) of SCFs for CH\(_3\)OH for different basis sets [20]

<table>
<thead>
<tr>
<th>basis set</th>
<th>(c)</th>
<th>(#) CSFs</th>
</tr>
</thead>
<tbody>
<tr>
<td>STO-3G</td>
<td>14</td>
<td>(1 \cdot 10^6)</td>
</tr>
<tr>
<td>6-31G**</td>
<td>50</td>
<td>(7.6 \cdot 10^{17})</td>
</tr>
</tbody>
</table>

![Figure 2.1: Schematic overview of active and inactive space of the CASSCF method](image)

In the complete active space self consistent field method (CASSCF) the full CI is performed within the active space. The challenge is to find the right active space...
2.2. Photoisomerization

Photoisomerization plays a crucial role in many reactions in biochemistry and is the focus of this thesis. This chapter provides a qualitative description of two different kinds of photoisomerization processes.

![Diagram of photoisomerization](https://via.placeholder.com/150)

**Figure 2.2.** Schematic overview of photoisomerization. The reaction coordinate $q$ is plotted against the energy of states $S_0$ and $S_1$. (a) shows an adiabatic transition from $S_1$ to $S_0$ via emission of a photon. (b) shows a non-adiabatic transition from $S_1$ to $S_0$.

The system is situated in its ground state $S_0$ at a local minimum of a potential energy surface. To visualize this surface, at least one reaction coordinate $q$ is needed, which is plotted against the energy of the system. The reaction coordinate could be, for instance, a bond length, a dihedral angle or any other attribute that characterizes a reaction. In the ground state the system is forced to remain in the minimum because of a barrier $E_0$ that cannot be crossed. When the system is excited by a photon, it evolves from the ground state to the first excited state $S_1$. Due to the instant excitation of the system, the value of the reaction coordinate is fixed. In other words, the excitation must be represented vertically in Figure 2.2. However, in the $S_1$ state a barrier $E_1$ can also be located at the Franck-Condon region. In contrast to $E_0$, the barrier $E_1$ is low and consequently can be passed. The system evolves to a deeper local minimum that causes a change in $q$, for example, a single
2. Theory

bond could convert to double bond, or the variation of an angle can lead to a twisted structure. Two different cases are considered in this work:

- After spending a time $\tau$ in the minimum of $S_1$ the system falls back to the ground state via emitting a photon. This transition is referred to as a fluorescent decay with $\tau$ being the fluorescent lifetime. Once returned to $S_0$ two possible paths can be taken. Either the systems goes back to the starting minimum (cis) or it evolves to another minimum (trans) that causes a permanently change in $q$. The cycle is schematically shown in Figure 2.2a.

- In the second case, the energy surfaces $S_0$ and $S_1$ intersect or almost intersect when $S_0$ has its maximum and $S_1$ has its minimum. The system can evolve from $S_1$ to $S_0$ as a result of its vibrational modes. To be precise, the energy difference between the surfaces has to be on the order of or smaller than the separation between the vibrational energy levels, see [12]. At this point, the Born-Oppenheimer approximation (see Section 2.1.1) collapses because the different potential energy surfaces are no longer well-defined. The point where both surfaces intersect is called conical intersection and needs a different mathematical description. The conical intersection causes a radiationless decay (no photons are emitted) to the ground state. As mentioned before the system can evolve in two different ways. The radiationless cycle is illustrated in Figure 2.2b.

Figure 2.2 is a simplified model for photoisomerization. In reality the PES does not depend on just one reaction coordinate but on degree of freedom minus six coordinates. Therefore, the intersection is given by a multi-dimensional seam (see Figure 2.3 for three-dimensional PESs).

---

Figure 2.3: Three-dimensional potential energy surfaces [11]
3.1. Natural Origin and Structure of pCK−

One of the most frequently described systems for photochemistry, especially for cis-trans photoisomerization, is given by the photoactive yellow protein (PYP) and its chromophore para-coumaric acid (pCA). The PYP was originally found in the salt tolerant bacterium *Halorhodospira halophila* and has the role of a photoreceptor which is involved in the photomotility. To avoid damages, caused by ultra-violet radiation, the bacterium detects blue light and moves to the opposite direction of the gradient. Precisely, the chromophore absorbs blue light which triggers a fully reversible photo cycle with several intermediates on time scales between a few hundred femtoseconds and seconds [15, 24].

![Figure 3.1: Chromophore pCA in its natural environment as part of the photo active yellow protein (PYP)](image-url)
3. The Biological System

covalent thioester bond with a cysteine residue (Cys69). Furthermore, it experiences steric constraints imposed by the geometry of the active site as well as hydrogen bonds with several amino acids. In detail, it is Glu46 and Tyr42 on the phenolate side and Cys69 on the carbonyl side. Moreover, the chromophore is influenced by the electrostatic interaction of Arg52. In PYP, the chromophore is present in its anionic form which is suggested to play a crucial role in cis-trans photoisomerization and thus the biological function of the protein.

The chromophore which is investigated in the thesis is an analogue of the pCA. The acid group is substituted by a methyl group to maintain the anionic character where the net charge is -1. Figure 3.2 represents the structure of the p-coumaric ketone (pCK\(^-\)) chromophore. Moreover, in the following work, only the ultra-fast cis-trans photoisomerization process which has already been observed in pCK\(^-\) \([3]\) will be taken into account. In this work, three energy minima are important, two in the first excited state and one in the ground state, where each minimum corresponds to a different geometrical conformation.

![Figure 3.2: pCK\(^-\) and its minimum structures. (Bottom) planar minimum structure. All carbon atoms are aligned in the same plane; (Top left) single bond minimum structure SB\(_{\text{min}}\). The torsion angle \(\alpha = 90^\circ\) relative to the planar minimum; (Top right) double bond minimum structure DB\(_{\text{min}}\). The torsion angle \(\beta = 90^\circ\) relative to the planar minimum.](image)
In Figure 3.2 (bottom) both the phenolate and alkene moiety of the chromophore are located in the same plane and therefore this structure will be labeled as planar minimum. The second minimum (SB$_{\text{min}}$) can be reached by a single bond (SB) twist of 90° and the third minimum (DB$_{\text{min}}$) by a double bond (DB) twist of 90°. Figure 3.2 illustrates the possible structures. In contrast to the planar minimum, the SB$_{\text{min}}$ and DB$_{\text{min}}$ are only accessible via the first excited state $S_1$ (see Section 2.2) where the transition from the planar minimum to the respective excited state minimum is achieved on time scales less than one picosecond which is supported by several ab initio calculations ([3, 9]) as well as experimental data [6]. These time scales are not only increasing the reaction efficiency but are necessary to observe the isomerization process in simulations.

### 3.2. Charge Distribution and First Excited State

The pCK$^-$ chromophore contains 86 electrons which are occupying 43 orbitals in the ground state. However, the important fact is that the first excited state is energetically well separated from its two neighbors.

Table 3.1.: Energy difference for excited states of pCK$^-$, calculated using EOM-CCSD and Pople’s 6-31G* basis set.

<table>
<thead>
<tr>
<th>$\Delta S_0S_1$</th>
<th>$\Delta E$</th>
<th>wavelength</th>
<th>oscillator strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta S_0S_2$</td>
<td>2.7505 eV</td>
<td>450.76 nm</td>
<td>0.2565</td>
</tr>
<tr>
<td>$\Delta S_0S_3$</td>
<td>4.3450 eV</td>
<td>285.35 nm</td>
<td>0.0977</td>
</tr>
<tr>
<td>$\Delta S_0S_4$</td>
<td>4.5273 eV</td>
<td>273.86 nm</td>
<td>0.0216</td>
</tr>
</tbody>
</table>

Table 3.1 lists the energy differences from the ground to the first, second and third excited state as well as its corresponding excitation wavelength and oscillator strength. The separation of $S_1$ is very favorable as it ensures that only the first excited state gets occupied when the stimulating photons are within a certain energy interval. Otherwise, the photochemistry which will be investigated in the following chapters would be much more complicated. Another important fact for the following calculations is the symmetry of the ground and first excited state. According to Groenhof et al. [3, 9] as well as an EOM-CCSD calculation, both states belong to $\pi$ orbitals, making the $S_0/S_1$ transition a $\pi/\pi^*$ transition.

$^1$The oscillator strength refers to the transition from the ground state to the respective excited state.
3. The Biological System

Figure 3.3.: HOMO and LUMO for single excitation as well as the mulliken charge distribution for both states and each minimum structure respectively. Note the charge transfer between the HOMO and LUMO.

Figure 3.3 presents the highest occupied molecular orbitals (HOMO) and the lowest unoccupied molecular orbitals (LUMO) for all minimum structures after performing a CASSCF(12e,11o) calculation. The computational details are listed in the following Section 4.1. For visualizing the charge distribution of all minimum structures, I assume that most of ground/first excited state, especially its transition, is essentially modulated by the HOMO/LUMO. In Figure 3.3a the electron density in the HOMO is slightly shifted to the phenolate ring compared to the LUMO where it is more favored over the alkene moiety. The density shift involves a charge transfer when the chromophore gets excited. Hence, the dipole moments $\mu_0$ and $\mu_1$ are pointing in different directions, even though the difference between both dipole moments is small. On the contrary, if pCK$^-$ evolves to one of the other minimum structures, the shift becomes much more obvious which is demonstrated in Figure 3.3b and Figure 3.3c. For SB$_{\text{min}}$ (Fig. 3.3c) the electron density of the HOMO is exclusively placed on the phenolate ring, compared to the LUMO where it is completely transferred to the carbonyl tail. As a matter of fact, the charge separation is increased through the SB twist and therefore the dipole moment of the respective state is strengthened.
On the other hand, the DB twist (Fig. 3.3b) has the opposing effect. Here the LUMO occupies only the ring structure, whereas the HOMO is only located on the alkene moiety. However, the significant effect is given by the change of directions of the dipole moments and its absolute values, whether the chromophore is situated in SB\textsubscript{min} or DB\textsubscript{min}. For more quantitative evidence of the charge transfer please refer to the Mulliken charge distribution at the bottom of each structure in Figure 3.3. The charge transfer will play a key role for understanding the result in the subsequent chapters.
The aim of this chapter is the investigation of the topology of the $S_0$ and $S_1$ state of pCK$^-$ in vacuo with respect to external electric fields. The first section deals with the computational details, followed by the discussion of the topologies, for which both possible reaction coordinates ($\alpha$ and $\beta$), see Figure 3.3 on page 14, with their corresponding energies are evaluated. Subsequently, the field influence of the two excited state minimum structures is presented, followed by a short examination on the active space dependency of the topologies.

4.1. Computational Details

The calculations reported in this chapter have been achieved with the ab initio software package MOLPRO [26]. The first approach of evaluating the single point energies was obtained via the restricted Hartree-Fock (RHF) method. Afterwards, a geometry optimization has been performed. Subsequently, the canonical Hartree-Fock MOs were used as starting guesses for CASSCF calculations. Triplet states have not been considered, so the spin quantum number always stayed $S = 0$. The active space was chosen with respect to the expected $\pi/\pi^*$ transition (see Section
4. Static Calculations

and required 6 occupied (12 electrons) and 5 unoccupied MOs to include all π orbitals.

To be precise, Table 4.1 shows the necessary swapping of the orbitals to fulfill the required active space, including only π orbitals which inhibits n/π* transitions. Furthermore, the CASSCF calculations were performed using the state averaging method of $S_0$ and $S_1$ where both weights were taken equally as 0.5. Throughout all calculations Pople’s basis set 6-31G* was applied. The electric field was added as finite dipole field to the one electron Hamiltonian (see [14]).

Table 4.1.: Active space selection for CASSCF (12e,11o)

<table>
<thead>
<tr>
<th>replaced MO numbers</th>
</tr>
</thead>
<tbody>
<tr>
<td>41↔35</td>
</tr>
<tr>
<td>39↔34</td>
</tr>
<tr>
<td>47↔50</td>
</tr>
<tr>
<td>48↔54</td>
</tr>
</tbody>
</table>

Moreover, the field direction equals the direction of the ground state dipole moment $\mu_0$ which was calculated after the CASSCF(12e,11o) performance of the planar minimum structure. Thus, positive field values point parallel to $\mu_0$, whereas negative ones point anti-parallel. Figure 4.1 shows the orientation of the electric field, the ground and first excited state dipole moment. I believe the orientation of the electric field is reasonable because even in proteins the isomerization process is much faster than the adjustment of the excited molecule to the induced electric field lines, and therefore only at the Franck-Condon region the electric field points in the same direction as $\mu_0$. While simulating the single and double bond twisting processes, an adjustment to the field within 1 ps could not be observed. To obtain the PES the torsion angle, $\alpha$ or $\beta$, was successively increased from the PM to the SB$_{\text{min}}$ or DB$_{\text{min}}$ and beyond by $\Delta = 5^\circ$ with respect to a constant field strength. The twisting was achieved by holding the phenolate ring fix and rotating the carbonyl tail.

In the remaining thesis E represents the electric field pointing in the same direction as the ground state dipole moment of the planar minimum.

Figure 4.1.: Schematic overview of the electric field orientation
Moreover, geometry optimizations have been performed for selected angles and field strengths. Table 4.2 lists the default convergence criteria of MOLPRO.

Table 4.2: Convergence thresholds used by MOLPRO

<table>
<thead>
<tr>
<th>Threshold</th>
<th>Threshold [a.u.]</th>
<th>Threshold [a.u.]</th>
</tr>
</thead>
<tbody>
<tr>
<td>gradient</td>
<td>$1 \times 10^{-2}$</td>
<td>$3 \times 10^{-4}$</td>
</tr>
<tr>
<td>energy</td>
<td>$1.7 \times 10^{-6}$</td>
<td>$1 \times 10^{-6}$</td>
</tr>
<tr>
<td>step length</td>
<td>$1 \times 10^{-3}$</td>
<td>$3 \times 10^{-4}$</td>
</tr>
</tbody>
</table>

(a) CASSCF
(b) Geometry optimization
4. Static Calculations

4.2. Double Bond Twist

The DB twist corresponds to the change of the torsion angle $\beta$, see Figure 3.2 on page 12. The starting position of the rigid scan is the planar minimum, which is located at $\beta = 180^\circ$. As the second coordinate the electric field strength was taken into account. Therefore, the field strength was varied from $-50 \text{ MV/cm} \leq E \leq 50 \text{ MV/cm}$, with $\Delta E = 5.5 \text{ MV/cm}$.

![Figure 4.2:](image)

Figure 4.2.: The total energy of $S_0$ and $S_1$ is plotted against the electric field strength $E$ and the DB torsion angle $\beta$.

At $\beta = 90^\circ$ and $E = 50 \text{ MV/cm}$ Figure 4.2 shows a global maximum for $S_0$ and a minimum for $S_1$. The corresponding structure can be identified as the double bond twisted minimum DB$_{\text{min}}$ (see Fig. 3.2 on page 12). This PES indicates a photoisomerization including a conical intersection which leads to a radiationless decay. As a matter of fact, the general tendency can be observed that a stronger positive electric field lifts the energy of $S_0$. At the Franck-Condon (FC) region ($\beta \approx 180^\circ$) this is reasonable as the field direction equals the direction of the dipole moment $\mu_0$. On the other hand, the energy is lowered when the electric field points anti-parallel to the dipole moment. Both dipole moments $\mu_0$ and
4.2. Double Bond Twist

μ₁ show almost the same orientation at the FC region, see the charge distribution in Figure 3.3a on page 14, and therefore both surfaces elevate with increasing field strength. If β is twisting, the present dipole moment will change as well, thus, it does not point in the same direction as the external field anymore. Nevertheless, the energy gradient in field direction of S₀ (see Figure 4.2) is increased while increasing the field strength, leading to global maximum for β = 90°. The reason lies on the charge separation of DBₘᵢₙ, see Section 3.2. The absolute value of the dipole moment is increased, so that even the projection of μ₀ on the electric field vector, is larger than μ₀ of the planar minimum. To get a qualitative idea of the different dipole moments, see Figure 4.3. Unfortunately, quantitative data of the dipole moments cannot be presented as the chromophore has a non-zero net charge, and therefore the values of the dipole moments depend on the chosen origin.

For β < 90° the energy gradient in field direction decreases again as the charge separation is reversible. In contrast, the energy of S₁ becomes more negative for β = 90° and positive field strengths. Here the charge separation again increases the absolute value of μ₁ but it is even more important that the charge separation also partly inverts its direction in comparison to the field orientation. Hence, a component of μ₁ points anti-parallel to E, leading to a local minimum for E = 50 MV/cm and β = 90°.

Figure 4.4 on the next page shows the energy difference |S₁ – S₀| as function of β and E. Particularly, for E = 50 MV/cm and β = 90° an intersection point can be identified.
4. Static Calculations

Figure 4.4.: The energy difference between \( S_1 \) and \( S_0 \) is plotted against \( \mathbf{E} \) and \( \beta \).

The PESs were created without geometry optimization, otherwise the computational effort would be too high. Nevertheless, two energy two-dimensional section planes were calculated considering geometry optimizations, namely the planes for \( \mathbf{E} \in \{0 \text{ MV/cm}; 50 \text{ MV/cm}\} \). It was not possible to perform geometry optimizations while applying electric fields neither with the MOLPRO nor the GAUSSIAN \(^8\) software package. To obtain the optimized energies, geometry optimizations were performed without applied field and second energy calculations with applied field were started afterwards. I thus assume that the applied field strengths are not high enough to change the geometry of the molecule. The results are presented in Figure 4.5. The data reveal that the intersection of both states is also observable when applying the field and secondly, the reaction barrier disappears. Without field, the barrier is about \( 18 \text{ kJ/mol} \) high, but with applied field only a saddle point instead of a maximum is recognizable.
4.2. Double Bond Twist

The key results of the DB topology examination are: At first, positive field strengths favor the DB twist and lead to conical intersections and secondly, the energy barrier that needs to be crossed to evolve into DB\textsubscript{min} disappears while applying positive fields. Note, that the field orientations are relative to the ground state dipole moment of the planar minimum, see Section 4.1.
4. Static Calculations

4.3. Single Bond Twist

The SB twist corresponds to the rotation of $\alpha$ (see section 3.1). Similar to Section 4.2, the PESs were obtained without geometry optimization, yet the two interesting section planes were treated separately with more accuracy. The evaluated interval for $E$ and the step size $\Delta E$ are the same as in Section 4.2, the SB torsion angle $\alpha$ was varied from $180^\circ$ to $60^\circ$ with $\Delta \alpha = 5^\circ$. Figure 4.6 illustrates a global maximum of $S_0$ and a global minimum of $S_1$ for $E = -50 \text{ MV/cm}$ and $\alpha = 90^\circ$.

In contrast to the DB twist, this point is achieved by a strong electric field opposite to the dipole moment of the planar minimum. The corresponding structure can be identified as the single point minimum structure $\text{SB}_{\text{min}}$ (see Fig. 3.2 on page 12). The reason for the local maximum in $S_0$ and minimum in $S_1$ - similar to the double bond topologies - is given by the strong charge separation of $\text{SB}_{\text{min}}$, which shows exactly the opposite behavior of the $\text{DB}_{\text{min}}$ (see Section 3.2 and Figure 4.7). Here the gradient in field direction of $S_1$ is steeper than of $S_0$, because for $\alpha = 90^\circ$, $\mu_0$ is almost pointing perpendicular to the field direction. Instead of a conical
4.3. Single Bond Twist

intersection, Fig. 4.6 still shows an energy gap, even though the electric field strength is comparatively high. Due to the gap, surface hopping between both states is rather unlikely, forcing the system to stay at $S_1$. If the barrier near the minimum of $S_1$ is high and broad enough, the only possibility of falling back to $S_0$ will be via emission of a photon. A similar behavior of the surfaces is also observable at the FC region ($\alpha \approx 180^\circ$) where both surfaces are lifted with increasing field strength as the respective dipole moments point in the same direction. However, an apparent difference between the surfaces in Fig. 4.2 is the smoothness of the curves. $S_1$ is rather even, whereas $S_0$ exhibits a larger curvature. The opposite behavior is demonstrated in Figure 4.6 where $S_0$ is flat compared to $S_1$, leading to the assumption that the molecule evolves much quicker from the planar minimum to $SB_{\text{min}}$ than to $DB_{\text{min}}$ with respect to different field orientations. Further support comes from the observation of different heights of the energy barriers at the $S_1$ level. Figure 4.2 exhibits a much higher barrier than Figure 4.6 where no barrier is recognizable, leading to the conclusion that the SB twisted structure is favored, at least without applied field.

![Figure 4.7](image1.png)

Figure 4.7.: Schematic overview of the orientations of the dipole moments

![Figure 4.8](image2.png)

Figure 4.8.: The energy difference between $S_1$ and $S_0$ is plotted against $E$ and $\alpha$

Figure 4.8 shows the energy difference between both states where an energy gap is observed for even a negative field strength of $-50 \text{MV/cm}$. After performing geometry...
optimizations of the two section planes where $E \in \{-50 \text{ MV/cm}, 0.0 \text{ MV/cm} \}$, the observed energy barrier without electric field was negligibly small and completely disappeared when applying negative field strengths. The field dependency of the PES as described previously, e.g. the energy gap, is still present and thus, can be confirmed by the optimized curves, see Figure 4.9.

Figure 4.9.: Section planes of the DB topology. (Top left) with negative applied electric field $E = -50 \text{ MV/cm}$, (bottom left) without electric field. (Top right) zoom of the reaction barrier with applied field and without (bottom right).

The SB topology shows that negative field strengths favor the single bond twist. In detail, the very small energy barrier entirely vanishes and the energy gap between $S_0$ and $S_1$ becomes smaller, but is still present, at the examined field strengths.
4.4. Influence of the Field & Active Space

The energy gap between $S_1$ and $S_0$ of both, the single and double bond twisted structures, decreases linearly with the favoring field strength (see Figure 4.10). The absolute slope of the SB twisted structure is stronger than the slope of the DB twisted structure. Moreover, I suggest that even within $SB_{\text{min}}$ a conical intersection can be forced which would require, according to extrapolation, a field strength of approximately $-70 \text{ MV cm}^{-1}$.

![Figure 4.10](image)

Figure 4.10.: Energy gap between $S_0$ and $S_1$ plotted against the electric field strength of the single and double bond twisted structure. The fitted parameters of the linear regression are: $DB_{\text{min}}$ slope=$(-0.024 \pm 0.001) \text{ eV cm}^{-1} \text{ MV}^{-1}$, intercept=$(1.116 \pm 0.005)$ eV; $SB_{\text{min}}$ slope=$(0.033 \pm 0.001) \text{ eV cm}^{-1} \text{ MV}^{-1}$, intercept=$(2.230 \pm 0.002)$ eV.

Additionally, the topologies for the single and double bond twist were calculated, with a reduced active space, including six electrons and six orbitals. The topologies are presented in the appendix, see Fig. A.1 on page 43 and Fig. A.2 on page 44. In comparison to the larger active space, both SB topologies show the same curvature, except for their total energy values caused by the varied active space. However, the DB topology (Fig. A.2) demonstrates an evident difference to its larger active space analogue at first sight. The conical intersection is already present for a field strength of $25 \text{ MV cm}^{-1}$, followed by an increasing energy gap that cannot be reasonable as the angle did not twist and therefore both dipole moments still point in the same direction. Due to the smaller active space, the electric field is strong enough to induce a transition from the first excited state to the ground state, given by the swap of the CI coefficients. The transition is graphically supported by the gradients in field direction of $S_0$ and $S_1$ for $\beta = 90^\circ$ where the slope of $S_0$ before the conical intersection equals the slope of $S_1$ afterwards, and vice versa.
After performing the single point energy calculations, this chapter is about the time-resolved simulations, including new aspects as well as confirmation of earlier results. First, the computational details and the simulation setup are discussed, followed by the question if the predicted single/double bond twist and the radiationless decay of the latter can be observed in simulations. Finally, the field dependency of the absorption spectra and a proposition for a spontaneous emission rate is presented.

5.1. Simulation Setup and Computational Details

5.1.1. Simulation Setup

The simulation setup can be split into two parts: Firstly, the chromophore which was placed at the center of the simulation box and secondly, two charged plates which are embedding the chromophore. The plates are modulated with variable charged ions to imitate a parallel plate type capacitor.
5. Dynamics

To get a better idea of the setup, especially of the dimensions see Figure 5.1 and 5.2a. To understand the relative field orientations which will only be referred to as positive or negative in the following sections, see also Figure 5.1. If the plate next to the phenolate ring is positively charged and the other one negative, the field will be positively defined and vice versa. Additionally, the field lines\(^1\) are illustrated in Figure 5.2b representing the electric field distribution between both plates.

Figure 5.1.: Schematic illustration of the simulation setup

![Simulation setup](image1)

Figure 5.2.: (a) shows the simulation setup. The chromophore is placed in the center of the imitated plate type capacitor; (b) visualizes the field lines of both plates. Note that in the center of both plates the electric field lines are parallel.

The field lines are parallel to the line between the center of both plates thus, the field is homogeneous as expected from a parallel plate type capacitor. At the edges they are curving because of the limited plate size. Nevertheless, at the center, where

\(^1\)The field lines are calculated via the Poisson-Boltzmann equation plug-in of VMD [16]
the chromophore is located, the field strength can be calculated as for an infinite plate type capacitor. Overall, each plate contains 400 ions with a total area of $A = (6 \times 6) \text{nm}^2$. The total electric field strength can be expressed as

$$E = \frac{400 \cdot \tilde{q} \cdot e}{\epsilon_0 \cdot A}$$

(5.1)

where $\tilde{q}$ is the variable charge of an ion, $e$ the elementary charge and $\epsilon_0$ the dielectric constant. For $\tilde{q} = 0.01$ the field strength becomes $E = 20.3 \text{MV/cm}$.

5.1.2. Computational Details

To perform time-resolved simulations the QM/MM software GROMACS [1] was used. The chromophore was treated with quantum mechanics similar to the static calculations, see Section 4.1. The active space included six orbitals with six electrons, otherwise a meaningful simulation time would not have been affordable. On the contrary to the previous active space, instead of taking the canonical HF orbitals, the natural bond orbitals (NBO) were used as basis for the active space which was successively reduced where again only $\pi/\pi^*$ orbitals were taken into account. The minor selection criterion was obtained by the one electron density matrix. More precisely, orbitals with an eigenvalue of almost 2.0 or 0.0 were ruled out.

Table 5.1.: MO selection steps for reducing the active space

<table>
<thead>
<tr>
<th>active space</th>
<th>replaced MOs</th>
</tr>
</thead>
<tbody>
<tr>
<td>(10e;11o)</td>
<td>(39,38)$\leftrightarrow$(37,36)</td>
</tr>
<tr>
<td>(10e,10o)</td>
<td>-</td>
</tr>
<tr>
<td>(10e,9o)</td>
<td>-</td>
</tr>
<tr>
<td>(8e,8o)</td>
<td>-</td>
</tr>
<tr>
<td>(8e,7o)</td>
<td>47$\leftrightarrow$45</td>
</tr>
<tr>
<td>(6e,6o)</td>
<td>41$\leftrightarrow$40</td>
</tr>
</tbody>
</table>

1 Instead of 12 electrons the starting active space contains only 10 as the orbitals are not in correct order.

For the detailed orbital selection, see Table 5.1. Most of the simulations were performed via the GAUSSIAN plug-in for GROMACS. The MOLPRO plug-in was only used for calculating the transition dipole moments (see Section 5.5) and to double check selected results. Again the 6-31G* basis set and the state averaging method for the ground and first excited state were applied with both weights taken equally as 0.5. The time step was set to 0.5fs and the convergence energy and gradient threshold for GAUSSIAN was appointed to $10^{-6}$ a.u. which is consistent with a moderate accuracy but is a necessity for the required simulation times (1-2 ps).

For the molecular mechanics part, including the angle and dihedral dependency of the chromophore, the amber 99 force field [5] was
applied. Furthermore, all starting momenta were randomly generated but obey the Maxwell distribution for 300 K. All simulations have been performed using periodic boundaries with a box size of $20 \times 10 \times 10$ nm as well as a coulomb cutoff with a radius of 5 nm. To ensure that the ions do not interact with each other exclusions were applied. Moreover, to keep the chromophore always at the center of the box, a position constraint for the oxygen atom, located at the ring structure, was placed, otherwise the chromophore would move due to its net charge.

5.1.3. Ground State Ensemble

The first step before simulating excited state dynamics was the establishment of a ground state ensemble. After a classical MD simulation over 30 ns (0.1 ps step size), six trajectories with a total length of 2000 steps each were received using QM/MM. The applied electric field had a strength of $20 \text{ MV cm}^{-1}$, so the molecule could adjust to the field lines. Thus, the same situation as in the static calculations, where the electric field was applied parallel to $\mu_0$ of the planar minimum, was obtained. The positive strength was arbitrarily chosen with the only constraint that it is strong enough to adjust the chromophore. I might have chosen negative strengths as well, but the necessary simulation time for adjusting the chromophore to the field lines would be much higher as it has to rotate around 180° (see the starting position in Figure 5.1 on page 30). Finally, for getting the ensemble new starting structures were extracted from the equilibrium trajectories. More precisely, the first structure was obtained after the 800th step followed by three more every 400 steps, so that overall the ensemble contained 24 structures.
5.2. Control of the Single/Double Bond Twist

The focus of this section lies on the question whether or not the excited state dynamics of pCK$^-$ can be controlled by electric fields. Therefore, three different field strengths $E \in \{-20 \text{ MV/cm}, 0 \text{ MV/cm}, 20 \text{ MV/cm}\}$ were chosen according to the results of the static simulations in Chapter 4, although they did not match the maximal/minimal used field strengths.

![Figure 5.3: Relative frequencies of single and double bond twisted structures for different field strengths. For each strength 24 simulations were performed.](image)

For every field strength the generated ensemble had been simulated in the excited state as long as all trajectories were either in SB$_{\text{min}}$ or DB$_{\text{min}}$. The results are illustrated in Figure 5.3 where the relative frequencies are shown for different field strengths. The effect which is supported by the static investigations can be approved by the simulations. If the field strength is positive i.e. see Figure 5.1, almost all trajectories will go into the DB twist. On the contrary, the SB twist is favored by a negative field, although the distribution is not as clear as for the DB. According to the field influence of SB$_{\text{min}}$, see 4.10 on page 27, the distribution will even more favor the SB twist while increasing the negative field strength. Even though the results confirm the previously calculated PES, the predicted predominance of the SB structure - according to the topologies (see Figure 4.6 on page 24 and Figure 4.2 on page 20) - could not be found. The topologies were performed with respect to
5. Dynamics

one degree of freedom, so they serve as a oversimplified model but do not reflect reality. On the contrary, the time-resolved simulations depend on on all internal coordinates. As an example of the discussed trajectories, Figure 5.4 shows the time evolution of the energy gap between both states and its corresponding torsion angle, whereas the field strength was set to zero. Both excited state twisting processes behave related, when the single/double bond angle equals 90° the energy gap becomes minimal which implies that the $SB_{\text{min}}/DB_{\text{min}}$ is reached. Moreover, in this example the SB twist occurs faster than the DB twist which is in good agreement with the described curvature of the PESs and its missing barrier, see Section 4.3.

![Figure 5.4: Example of the single and double bond twist. The time evolution of the energy gaps $|S_{1} - S_{0}|$ and torsion angles without electric field is shown.](image-url)
5.3. Radiationless Decay

The next step is to focus on the DB twist and its expected radiationless decay. The goal is to determine the rate of the radiationless decay, assuming a simple exponential decay. The field strength $E = 20 \text{ MV cm}^{-1}$ was chosen for the simulation of the ensemble. According to Figure A.2 on page 44, the used field strength favors the conical intersection; for a smaller field the simulation time would increase exponentially. Unfortunately, within the time interval of 3.5 ps only 14 of 23 radiationless decays could be observed. To determine the rate the number of observed decay events $N(t)$ is assumed to be proportional to $e^{kt}$ where $k$ represents the decay rate. Figure 5.5 shows a half-logarithmic plot for $N(t)$ where $k$ is determined as the slope which is $k = (0.80 \pm 0.13) \text{ ps}^{-1}$.

![Half-logarithmic plot of observed number of radiationless decays to determine the decay rate $k$.](image)

Figure 5.5.: Half-logarithmic plot of the observed number of radiationless decays to determine the decay rate $k$. The obtained parameters are: slope=$0.80 \pm 0.13$ ps$^{-1}$ and intercept=0.65 ± 0.21.

The obtained rate is in close agreement with experimentally observed fluorescence decay rates in solution. Espagne et al. [7] measured the fluorescence decay rate for pCK$^-$ in aqueous solution using steady-state spectroscopy to be 0.77 ps$^{-1}$. Even though they did not directly apply electric fields, the strong dipole moments of the water molecules induce local electric fields. I suggest that these induced fields are in the order of the applied external electric fields and therefore accelerate the decay. In any case, according to [3], hydrogen bonds are stabilizing the single and double bond twisted minimum, with respect to $S_0$. Moreover, they lower the energy gap between $S_0$ and $S_1$ in the DB minimum, while the gap of the SB minimum is increased. Figure 5.6 shows the influence of hydrogen bonds for the

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2The 24th was falling into the SB$_{\text{min}}$. 

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5. Dynamics

Figure 5.6.: Hydrogen Bonds linked to the SB/DB twisted structure. Additionally, the mulliken charge distributions are shown.

According to simulations of Boggio-Pacqua et al., within the SB twist three water molecules are involved nearby the carbonyl tail, whereas one weak hydrogen bond is located at the phenolate group. In the case of the DB twist, at least two strong hydrogen bonds were found to influence the phenolate group but only one or two weak bonds were found influencing the carbonyl tail. According to the mulliken charge distribution, the hydrogen bonding patterns do not only reflect the charge distribution (see Figure 3.3 on page 14) but rather increase the charge transfer of the respective excited state minimum.

As an example for the surface hopping Figure 5.7 shows the time-resolved DB twist and the following intersection. At 0.2 ps the system evolves from the planar minimum into DB_{min} and at 0.6 ps the energy gap becomes zero, the chromophore falls back to the ground state and finally moves back to the planar minimum (trans-isomer).

Figure 5.7.: Example for a radiationless decay transition. The time evolution of the DB torsion angle and its corresponding energy gap is shown.
As previously pointed out, in addition to the trans-isomerization also the cis-isomerization exists. Here $\beta$ changes from 90° to 0° (see Figure 5.8), so that all carbon atoms are aligned in the same plane. The number of cis/trans transitions was counted for the observed radiationless decays (see Figure 5.5). 10 of the 14 trajectories were falling into the trans minimum but, at least, four trajectories showed the cis-isomer. The quantum yield of the cis-isomer is approximately 29%, which is in good agreement with the observed quantum yield for the cis-isomerization of pCA in PYP experimentally [15] as well as in former calculations [10].

![Figure 5.8.: Cis-isomer of the pCK− chromophore](image)
In this section the absorption spectra of pCK$^-$ are investigated. The focus lies on the question if and how the absorption spectra are influenced by electric fields. The initial point was again the ground state ensemble in the planar minimum where each structure got excited and the energy gap between $S_0$ and $S_1$ was calculated. Subsequently, this procedure was repeated for five different field strengths and on the basis of the data normalized histograms were created and normal distributions were fitted against them. Note that each absorption spectrum is based on the same ground state ensemble. An arbitrary example for such a histogram as well as its Gaussian fit is shown in Figure 5.9. Even though the ensemble is comparatively small, a normal distribution can be recognized.

Figure 5.9.: Single absorption spectrum for $E = -20 \text{ MV/cm}$.

Figure 5.10.: Gaussian fitted absorption curves for five different field strengths.
The influence of the electric field is illustrated in Figure 5.10 where an obvious shift of the field spectra is observable. For negative field strengths the absorption wavelength are larger than for positive ones which is in good agreement with the static calculations (see Section 4.3 and 4.2) where the energy gap increases with positive field strengths. Furthermore, the shift seems to be proportional to the field strength and also the shapes of the Gaussians, which are determined by the standard deviations, are very similar. However, this is reasonable as, according to the linear Stark Effect, the electric field couples with the dipole moment $\mu_i$ of the respective state. The electric field is treated as perturbation, leading to the first order energy correction

$$E^{(1)} = \langle \psi_i^0 | H_1 | \psi_i^0 \rangle = -E \cdot \langle \psi_i^0 | \mu | \psi_i^0 \rangle = -E \cdot \langle \mu_i \rangle.$$  

(5.2)

Hence, whether or not the system gains energy depends only on the direction of the electric field relative to $\mu_i$. The positive field strengths in Figure 5.10 matches the direction of $\mu_0$ but not of $\mu_1$ (see Section 3.2), and therefore the energy difference between $S_0$ and $S_1$, only caused by the electric field is described by:

$$\Delta E = \pm |E| \cdot (|\langle \mu_0 \rangle| - |\langle \mu_1 \rangle| \cos \vartheta)$$  

(5.3)

where $\vartheta$ is the angle between the electric field and $\mu_1$. The energy shift is linear with the applied field strength as the second term is constant, because only the planar minimum structure was excited.

## 5.5. Spontaneous Emission

The last section is based on several assumptions. If the chromophore evolves into the SB minimum, the energy gap will be too large for a radiationless decay. Assuming that the barriers next to the minimum are high enough, the chromophore is forced to stay at SB$_{\text{min}}$ until a radiative decay can occur. This section provides the rate for a spontaneous emission from the first excited state to the ground state at SB$_{\text{min}}$ and its dependency on the electric fields.

The rate $\Gamma$ for spontaneous emission is proportional to the Einstein coefficient (for
5. Dynamics

a derivation see [19]) and can be expressed as

\[ \Gamma_{1\rightarrow0} = \frac{64\pi^4\nu_{10}^3}{3hc^2} \cdot |\langle 1|\hat{d}|0 \rangle|^2 \] (5.4)

where \( \nu_{10} \) is the frequency of the photon, being proportional to the energy gap between \( S_1 \) and \( S_0 \) and \( \hat{d} \) the transition dipole operator. To obtain \( \nu_{10} \) and \( \langle 1|\hat{d}|0 \rangle \) the SB minimum was simulated about 0.5 ps, while applying different field strengths. Overall, the field was varied from \(-40 \text{ MV cm} \leq E \leq 0 \text{ MV cm} \) with a step size of \( \Delta E = 4 \text{ MV cm} \) where three trajectories for each strength had been performed. In contrast to the previous section, all trajectories started with the same structure, otherwise structural change would lead to strong differences within the same field strengths. To achieve the rate for one field strength, the mean value of both, the transition dipole moments and energy gaps including all 500 steps, belonging to one trajectory, had been taken. Then the mean value of all three trajectories was evaluated. Figure 5.11 shows the rate dependency of the field strength which is apparently linear at least for the investigated interval. Furthermore, a linear regression had been performed with exclusion of the strongly deviated value \( E = -33 \text{ MV cm} \). The obtained linear function is:

\[ \Gamma(E) = 1.1 \frac{\text{cm}}{\text{ns MV}} \cdot E + 72.5 \text{ ns}^{-1} \] (5.5)

and thus, can be evaluated as direct proportional to the applied field strength. The intercept depends on the starting structure and therefore has less significance. According to Eq. 5.4 and the linear dependency of Eq. 5.5, the transition dipole moment \( |\langle 1|\hat{d}|0 \rangle| \) is proportional to \( \frac{1}{E_{1} - E_{0}} \).

Figure 5.11.: Field dependency of the rate for spontaneous emission from \( S_1 \) to \( S_0 \) at the single bond minimum. The fitted parameters are: slope \( m = (1.1 \pm 0.2) \frac{\text{cm}}{\text{ns MV}} \); intercept \( b = (72.5 \pm 5.5) \text{ ns}^{-1} \).
Conclusion & Outlook

In this thesis, the topology of the ground and first excited state for both possible torsion angles ($\alpha$ and $\beta$) of the pCK$^-$ chromophore were presented. The electric fields show a significant effect on the topologies of the single and double bond twisted structure. In detail, not only the applied field strength but also the orientation of the field determines whether the single bond or double bond twist is favored. The single bond twist is supported by negative field strengths and the double bond twist is supported by positive field strengths where the orientation is given relative to the ground state dipole moment of the planar structure. Moreover, by increasing the positive field strength the surfaces can be forced to intersect. In the second part, the achieved topologies were validated through dynamic simulations. The predicted influence of the field could be observed. Even if the field strengths of the time-resolved simulations are ten times larger than experimentally feasible ones - at least by using a capacitor - I think that the field influence can still be experimentally validated because the relative frequency of the single/double bond twisted structure increases linearly with it corresponding field strength (see Figure 5.3 on page 33). Additionally, the rate of the radiationless decay was obtained; it is similar to experimentally measured rates in aqueous solution, but without external electric fields. I suggest that the water molecules induce local electric fields being of the same magnitude
as the applied external ones. Furthermore, the field dependency of the absorption spectra was examined as well as an attempt was made to predict the radiative decay rate.

The next consequent step would be the experimental validation of the theoretically obtained results. Nakabayashi et al. [21] examined the field-induced changes in absorption and fluorescence of the GFP chromophore and Espagne et al. [6] also did absorption and fluorescence experiments with pCK−, although without electric fields.

Besides the experimental validation, further numerical calculations could include the effect of solvents additionally to the field influence which could favor the double bond twist even more. Another interesting question would be, if the single bond twist could be completely inhibited by an additional ethylene group connected to the phenolate ring.

However, the steering of excited state dynamics by electric fields makes several applications imaginable, for instance, a pixel could be established where the electric field acts as a switch for either color (SB twist) or no color (DB twist). Another possible application could be a molecular engine as the double bond isomerization is fully reversible, thus light energy would be converted into kinetic energy. Nevertheless, I hope that the results of this work will trigger further investigations of the influence of electric fields in photoisomerization and maybe it will form the basis of future technology.
Figure A.1.: PESs of $S_0$ and $S_1$ after performing CASSCF(6e,6o) without geometry optimization. The total energy of $S_0$ and $S_1$ is plotted against the electric field strength $E$ and the SB torsion angle $\alpha$. 
Figure A.2.: PESs of \( S_0 \) and \( S_1 \) after performing CASSCF(6e,6o) without geometry optimization. The total energy of \( S_0 \) and \( S_1 \) is plotted against the electric field strength \( E \) and the DB torsion angle \( \beta \).
Bibliography


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Bibliography


Acknowledgement

Ich möchte zuallerst meinem Betreuter Dr. Gerrit Groenhof danken, ohne dessen Idee diese Arbeit gar nicht erst entstanden wäre. Besonders durch seine open-door policy, so stand er mir fast täglich bei kleinen und größeren Problemen mit Rat und Tat zur Seite, trug er erheblich zum Erfolg dieser Arbeit bei.


Zusätzlich möchte ich noch Prof. Dr. Helmut Grubmüller danken, dass er sich bereit erklärt hat diese Arbeit als Zweitkorrektor zu bewerten.

Als letztes danke ich meiner Freundin Imke für ihre emotionale und vor allem orthografische Unterstützung, sowie meiner Mutter für ihre finanzielle Hilfe, ohne die mein Studium und damit diese Arbeit nicht möglich wäre.
Erklärung

nach §13(8) der Prüfungsordnung für den Bachelor-Studiengang Physik und den Master-Studiengang Physik an der Universität Göttingen:

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Göttingen, den September 6, 2011

(Fabian Knoch)