MECHANISM OF PROTON TRANSFER IN BACTERIORHODOPSIN

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Photoinduced proton pumping in bacteriorhodopsin (bR) was extensively studied using multiple experimental methods and utilizing various theoretical modelling approaches. These studies usually refer to the well-resolved structural data of bacteriorhodopsin. However, despite the obtained results, the origin of the proton pumping force initiated by the electronic excitation of retinal remains questionable. Using quantum chemical calculations, we have revealed that the retinal molecule after its excitation is fixed in the ground state of 13-cis,15-syn configuration, as a result of interaction with specific protein residuals. Reaching this fixed configuration, the proton is first transferred to the aspartic acid No. 85 (Asp-85) residue from the water molecule, which is subsequently restored by the proton initially located in the Schiff base. We discuss the challenges and approaches to modelling the proton transfer in bR and demonstrate that the process, which starts from the electronic excitation of the retinal molecule, is mainly due to the detailed arrangement of the protein environment.

Keywords: photoinduced proton transfer, membrane proteins, bacteriorhodopsin, retinal, isomerization

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1. Introduction

Bacteriorhodopsin (bR) is the light-transducing protein in the purple membrane of Halobacterium salinarum [1, 2]. This hydrophobic protein contains seven transmembrane helices (A through G, see Fig. 1), which span the membrane at small angles to the perpendicular and surround the centrally located retinal linked to G-helix by a protonated Schiff base (SB) with Lys-216 [3–6]. As a result of the light absorption, the retinal yields the transition into the 13-cis,15-syn form, which triggers a cyclic reaction that comprises the series of spectral intermediates usually named J, K, L, M, N and O, reflecting protein stages of the proton pumping across the membrane [7–13]. Proton transport starts from the middle of the protein structure, with the proton transfer from the SB to aspartic acid 85 residue labelled as Asp-85 or D85 [14]. Subsequently, the proton is transferred to the extracellular side of the membrane and the SB is reprotonated from the cytoplasmic side completing the cycle. In addition to the ability of the proton pumping of bR, it was also experimentally demonstrated that a negative chlorine (Cl−) or positive proton (H+) exchanger could be converted into a passive channel for Cl− by substitution of only two key residues [15, 16]. Site-directed mutagenesis combined with various spectroscopic methods, crystallographic data, in conjunction with quantum chemical and molecular dynamics simulations, revealed many details of the structural changes of the protein backbone in the course of the bR photocycle [8]. However, the main questions concerning the exact pathway of energy conversion from the electronic excitation of retinal into the proton pumping force in this photoactive protein, as well as the unidirectionality of proton transfer processes, are still unanswered.
In the present paper, we discuss the challenges for the modelling of bR related to its intermediate size, as it is too large for the fully quantum-chemistry-based description, but nevertheless, the system of helices and residues bound to them cannot be described statistically, in terms of the uniform dielectric environment. Moreover, the energy exchange between various parts of the whole system, such as the excited electronic states, transferred protons, twisted retinal and bent helices, is not unidirectional, as none of them serves as the energy sink. Correspondingly, the calculations can be fulfilled by keeping in mind the energetic states of the entire system, even if it is impossible to take into account all the specific interactions. As an example, we address the proton transfer event associated with the deprotonation of the SB and the protonation of the Asp-85 residue. We show that the irreversibility of the proton transfer can be achieved in terms of quantum chemical calculations if the 13-cis,15-syn configuration of the retinal defines the initial conditions for the proton pumping. This isomerization does not correspond to the minimum of energy for the retinal separated from the rest of the complex. However, we argue that it might be defined as the state corresponding to the proper energy redistribution in the full system at this configuration when all interactions with the residues at the bent helices are taken into account. We would also like to stress one of the active hypothesis that according to our analysis, proton transport starts from the middle of the protein, with one of the water molecules donating its proton to Asp-85 with a subsequent reprotonation from the SB.

The rest of the paper is structured as follows. In Section 2, we address the challenges of the modelling of the proton transfer. A review of the molecular dynamics and quantum chemical calculations of the photoisomerization is also presented in Section 2. The methods used for quantum chemical calculations are briefly described in Section 3. The results obtained by modelling the retinal isomerization and its link to the initial proton transfer steps are presented in Section 4. Discussion of the obtained results is presented in Section 5, and Section 6 contains the conclusions of our work.

2. Challenges in modelling of bacteriorhodopsin

2.1. Charge transfer

Modelling of charge transport is usually based on the assumption that the system can be effectively separated into the dynamical part and the environment. This assumption is at the heart of the Marcus theory of electron transfer with the transfer rate given by

$$k_{et} = \frac{2\pi}{\hbar} |H_{12}|^2 \frac{1}{\sqrt{4\pi\lambda k_B T}} \exp \left( \frac{(\lambda + \Delta G^0)^2}{4\lambda k_B T} \right),$$

where $|H_{12}|$ is the coupling between the initial and final states, $\lambda$ is the total reorganization energy of the environment, $k_B$ is the Boltzmann constant, $T$ is the absolute temperature, and $\Delta G^0$ is the total Gibbs free energy change for the transfer reaction. It was shown that such an approach can be successfully used to describe proton transport as well [17, 18]. However, there are two caveats for a specific bR case, both of them related to the smallness of the system and the corresponding impossibility of the separation of the dynamical subsystem and its environment. First, it is very difficult to define the reorganization energy, with all the bR residues reacting to the proton transfer events in a rather dynamical, not statistical manner. Second, in the calculation of the differences of free energies between the initial and final states, the direct electrostatic interaction between the transferring...
proton and charges on the residues provides an essential impact on the result. In previous works based on the energy conceptions [19, 20], these both aspects are not taken into account properly. Free energies corresponding to proton sites (based on the related pK_a factors) are usually calculated with the electrostatic interactions considered. The rest of the system is included by calculating the reorganization energy and non-vacuum dielectric permittivity, the electrostatic interaction of the protons with electrons being completely ignored, in particular. Water molecules, which are shown to be crucial for the proton transfer in bR [21–23], are considered electrically neutral, non-dissolved and dipole-less. While this approach can be appropriate for large protein systems [24], its applicability for relatively small systems, such as bR, is questionable.

A proper description of the full bR dynamics could be based on the inclusion of the influence of all helixes and residues into the Gibbs free energy and the consideration of various vibrational degrees of freedom as the environment. In this case, the reorganization energy would be small, and the functional form of Eq. (1) becomes close to a delta-function. With small dissipation, the transfer between the states would be almost resonant, but slightly downhill with respect to the energy of the full system. As mentioned above, the bR complex is too large for such quantitative treatment, but this approach might be used for qualitative discussion. In particular, if the exact dynamics of some fragment of the full complex is being calculated, the initial and final energies of this fragment alone do not have to be minimal for given configurations, as the requirement of the energy minimum for the relaxed system applies only to the whole complex. Moreover, when two final states are available, the system can have a higher probability to be transferred to the more energetic of the two, because it is impossible to unload a large amount of energy to the vibrational environment in order to proceed to the minimal energy state, i.e. if the system is in the inverted Marcus regime. In some sense, our approach is comparable to the ‘reaction coordinate models’ [23, 24], where strongly-coupled environmental degrees of freedom (often high frequency modes) were specifically included into the dynamical part.

2.2. Molecular dynamics and quantum chemical calculations

Mimicking photoinduced processes in bR is not a trivial task since the resulting outcome is sensitive to the protein environment, that has to be taken into account properly. For this purpose, molecular dynamics (MD) simulations have been intensively used [4, 27–37]. Indeed, the rotation of SB as a result of the retinal isomerization (see Fig. 3) can be obtained by modelling the retinal with its protein environment [34, 35]. However, in order to achieve it, multiple approximations should be used due to the limits of computational resources including algorithms themselves. One very crucial approximation suggests that the retinal structure can be subdivided into quantum mechanics (QM) and molecular mechanics (MM) segments, and the division between them is set across covalent bonds. For QM/MM simulations various approximations can be used for the description of MM and QM counterparts. The definition of the QM–MM boundary is the major problem by constructing covalent bonding and adaptive schemes [4, 27, 33, 37]. Hayashi and co-authors [34, 35] exploited only a part of the retinal structure with the attached SB using the QM approach, while the rest of the molecule was parameterized. QM simulations of full retinal do not demonstrate isomerization and/or rotations of some parts of the molecule as follows from experimental data [36, 37]. Moreover, the SB bound to the retinal does not rotate in the vacuum models but instead only the rotation of the β-ring is obtained in the excited state of retinal [58, 59]. Starting from the excited state, the structure optimization procedure restores the initially optimized ground state structure, which implies that the minima of the potential energy surfaces in the ground- and excited-states are not separated by any significant energy barrier. Furthermore, by analyzing the isomerization process of retinal up to 90 deg, it was concluded that the DFT methodology is not applicable due to the fact that the system is reaching the conical intersection region in the course of isomerization [40].

Molecular dynamics simulations with molecular mechanics force fields are performed in order to relate deformations of the bR helices to the proton transfer mechanism by assuming that
The L-intermediate follows from the retinal \textit{all-trans} to 13-\textit{cis} photoisomerization step corresponding to the K-intermediate \cite{41, 42}. If the water molecules (especially W402) are not taken into account, the retinal pumping of the proton is expected from the 13-\textit{cis},15-\textit{anti} structure and not from the 13-\textit{cis},15-\textit{syn}, in opposite to the crystallographic data \cite{42}. Other studies based on three transfer mechanisms in terms of QM/MM calculations \cite{43} also indicate the presence of the 13-\textit{cis},15-\textit{anti} configuration. More accurate approach for the simulation of the protonated retinal active centre is based on the force field methodology. The water molecular clusters with additional possible water molecules in accordance with several crystallographic data were taken into consideration \cite{44}. As follows from this type of studies, the proton transfer from retinal interacting with the water molecule (W402) is more complicated since the presence of the energy barrier appears for the proton pathway \cite{45}, while the NMR studies report opposite evidences \cite{46}.

The protonated retinal after its excitation is known as the J state of bR corresponding to a complication of the excitation dynamics of the retinal \cite{45–47}. In the vicinity of the local minimum of the first singlet excite state (S_1), the retinal coupled to the SB remains almost planar, but the single and double bonds turn to become inverted according to the MD simulations and \textit{ab initio} modelling \cite{29, 30, 32, 37, 48}. As follows from these data, the existence of a small barrier between \textit{trans} and \textit{cis} structures in the excited states could partially explain the fluorescence effects related to the structural evolution of retinal bound to the protonated SB in bR \cite{29}. Our studies of the retinal-type conjugated systems support this conclusion: the excited state S_1 demonstrates an inversion between single and double bonds, that should be caused by the presence of the second singlet excited state (S_2) properties (also known as 1^1B_u type state, corresponding to its symmetry according to polyene models), appropriate for

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\textbf{Fig. 2.} Schematic representation of the photoinduced initial processes: initial state of the retinal (RET) coupled with the protonated Schiff base (SBH\textsuperscript{+}); photoinduced isomerization of the retinal and subsequent proton release \cite{5, 40}. The Schiff base (SB) rotation angle presents the dihedral angle described by atoms 12-13-14-15.
polyene and carotenoid molecules [49]. It follows from the equation-of-motion coupled cluster singles and doubles model (EOM-CCSD) approach [50] that the lowest excited state of the retinal bound to the protonated SB is of a similar origin to that of the polyene type 1\(^1\)B\(^u\)\(^+\) state. In the case when retinal is bound to the non-protonated SB, the optically forbidden state of symmetry 1\(^1\)A\(^g\)^\(-\) turns to be the lowest excited state due to the electron correlations. The EOM-CCSD is computationally expensive and the excited state energies are largely overestimated. The lowest S\(^1\) state of the retinal bound with the deprotonated SB should be of the 1\(^1\)A\(^g\)^\(-\) symmetry while the lowest S\(^1\) state of the retinal bound to the protonated SB should be of the 1\(^1\)B\(^u\)\(^+\) symmetry. Thus, the retinal energy surfaces should be more complicated by considering the retinal isomerization together with the possible flexibility of helices. Thus, energy transfer channels, which might fix the retinal in the 13\(^\text{-cis}\),15\(^\text{-syn}\) and 13\(^\text{-cis}\),15\(^\text{-anti}\) configuration (Fig. 2), and finally pass the absorbed light energy from the retinal into the helices, are expected.

Deprotonation of the SB is usually attributed to the first proton transfer step in bR [10]. The protonation of Asp-85 upon the deprotonation of SB together with the photoisomerization of retinal from the initial all\(^\text{-trans}\) configuration to the 13\(^\text{-cis}\) configuration is followed by the release of a proton via a network of protein residues and bound water molecules [51–56] to the extracellular membrane surface [57, 58]. As follows from studies based on theoretical calculations, various possible pathways for retinal deprotonation in the ground state, which at moderate approximations agrees with the experimental data [48, 59], are expected.

3. Methods

To reveal the connection between the retinal photoisomerization and the initial steps of proton transfer, the studies of possible energy redistribution channels between the electronic excitation of the retinal and the protein surrounding are carried out. For these purposes, stationary density functional theory (DFT) calculation, two-layer own N-layered integrated molecular orbital and molecular mechanics (ONIOM), and quantum molecular dynamics (QMD) are used. Recently it was demonstrated that by considering simple polyene type molecules the excited states of retinal have a multireference character, which cannot be taken into consideration within the DFT approach [45, 50, 60]. However, the proton transfer is taking place from the ground state of the retinal as a result of the excitation relaxation, thus causing several bR stages [11, 18, 53, 62, 63]. Therefore, the DFT approaches should be applicable to provide quite accurate descriptions.

It is known that the bR in the L state reflects changes of the active centre environment in comparison to the initial state [62]. Thus, to simplify the description of the initial steps of proton pumping, these two states will be mainly considered. The initial structures are taken from the protein data bank (PDB). The PDB codes for the structures are 1C3W [5] and 2NTW [64], respectively. The structure 1C3W provides crystallographic data for bR in the initial state with the all\(^\text{-trans}\) configuration of the retinal and the 2NTW provides the crystallographic information for the bR in the intermediate state with the 13\(^\text{-cis}\),15\(^\text{-syn}\) configuration of the retinal (Fig. 2).

For QM calculations, DFT modelling is performed using the Gaussian software package [50] with HSEH1PBE, CAM-B3LYP and B3LYP functionals, and cc-pVDZ and 6-31G* basis sets. Similar approaches were used previously [27, 46, 49]. It should be noted that the HSEH1PBE and CAM-B3LYP functionals involve long-range corrections. According to our studies, the HSEH1PBE functional provides the same results as the B3LYP functional. Here we report the results of computations with the HSEH1PBE functional with the cc-pVDZ basis set. The CAM-B3LYP functional, which results in the β-ring rotation when the all\(^\text{-trans}\) retinal is optimized, is also applied for the optimization of the retinal structure in the excited state. The QM/MM or QM/QM in terms of ONIOM approach are carried out by adding relevant residuals of the protein and water molecules into the calculation scheme using molecular mechanics with the universal force field (UFF) or semiempirically using PM6 Hamiltonian methods. In all cases, the lower layer was always frozen while the higher layer was free of changes in ONIOM methodology. Quantum molecular dynamics (QMD) is performed with
Gaussian 9 version at the B3LYP/cc-pVDZ level by using the atom-centred density matrix propagation (ADMP) molecular dynamics model at different temperatures: 1, 70 and 300 K. ADMP was performed with 63 atoms taken into account, time steps were 0.1 fs, fictitious electronic mass was 0.1 amu, velocity scaling was performed all the way through the simulation, the dynamics were performed with converged SCF results at each point, and the temperature was set to determine the fixed nuclear kinetic energy according to the Gaussian overlay documentations [50].

4. Results

4.1. Modelling of retinal isomerization

The proton transfer correlates with the photoinduced transition of the retinal from its initial trans-configuration to the 13-cis,15-syn isomer on μs timescale [65]. Thus, first, we consider the photoisomerization process in terms of QM/MM by taking into account the structural details of the part of the protein from the vicinity of the photoactive centre (retinal) as follows from crystallographic data [5, 64]. The protonated SB, which is a proton donor, becomes connected to the proton acceptor Asp-85 via the water molecule (labelled W402) as a result of the retinal isomerization. Additional water molecules, labelled W401 and W406, as well as the protein two tryptophan residues labelled Trp-86, Trp-182 and tyrosine residue (labelled Tyr-185) are also present in the vicinity of the active centre and might play a decisive role in the photoinduced proton transfer. In order to repeat calculations with a higher basis set the threonine residue (labelled Thr-90) may also be required. The all-trans configuration of the retinal together with the protein residues and bound water molecules mentioned above is taken from the crystallographic data.

As follows from the calculations, the all-trans retinal (corresponding to the ground state) contains only one global minimum along the rotational pathway, while the optimization in the excited state results in the presence of two possible minima (Fig. 2(a)). One of these minima is at about −92 deg that correlates well with the structure defined from the crystallographic data 2NTW [64]. Moreover, the ground state in this configuration can also be found when the protein residues Trp-86, Trp-182, and Tyr-185 are taken into account, thus, suggesting a conical intersection between these two states (Fig. 3(b)). Similar results are obtained both in QM/MM or QM/QM (ONIOM) studies by adding Trp-86, Trp-182, and Tyr-185 residues into the calculation scheme and using molecular mechanics UFF.

Fig. 3. Schiff base rotation energy surface of the retinal bound to the protonated Schiff base along the dihedral angel indicated in Fig. 2: 0 deg corresponds to the all-trans retinal configuration in vacuum (a); −92.4 deg represents the 13-cis,15-syn retinal isomer together with the Trp-86, Trp-182 and Tyr-185 residues in vacuum (b). The dots represent the dihedral scan increment.
and in the semiempirical PM6 calculations. Thus, the minima corresponding to the excited states of $13$-$cis$, $15$-$syn$ isomer of the retinal appear within the supermolecular (several molecules in the same QM calculations) approach containing the retinal Trp-86, Trp-182, and Tyr-185 residues in agreement with the earlier suggestions for the ground state stability. However, this structure does not follow from the calculations of the ground state, so it can be obtained for the excited state minima only (Fig. 3). Accordingly, to obtain the minimum for the $13$-$cis$, $15$-$syn$ isomer of retinal in the ground state, additional interactions should be taken into account.

Thus, to examine the stability of the configuration of active centre corresponding to the retinal in the $13$-$cis$, $15$-$syn$ configuration, we initially fix this configuration and use the QMD simulations of the structure at different temperatures (Fig. 4).

The initial structures used for the calculations are chosen to be optimized either in the ground state or in the first excited state with the SB rotated by 90 deg. The results presented in Fig. 4 demonstrate that the ground state structure (E0) remains almost the same, while in the excited state (E1), this structure reaches other values of the rotation angle (retinal of cis isomer) on the time scale less than 0.5 ps at low temperatures. At higher temperatures (70 and 300 K), the isomerization process starts at around 30 fs and experiences transition into another structural arrangement in about 100 fs and, thus, the initial state evidently cannot survive for several µs, as expected from the experimental observations. Accordingly, we conclude that QMD simulations do not support the evidence of the appearance of $13$-$cis$, $15$-$anti$ retinal isomer without taking into account the essential interaction with the protein environment that has been ignored so far.

4.2. Initial step of proton transfer

To model the initial stage of the proton transfer, let us keep the key atoms taken from the crystallography data (with the retinal in the $13$-$cis$, $15$-$syn$ configuration and water molecules as it is in 2NTW [64]) fixed (see Fig. 5). The modelling data obtained by using QM calculations demonstrate the proton displacement from W402 to the Asp85 group (Fig. 6(a)) that opens the pathway of the proton transfer from the SB to W402 (Fig. 6(b)).

![Fig. 4. Possible evolution of the retinal demonstrating the isomerization as follows from the ADMP type quantum molecular dynamics in the ground state (E0) and in the first excited state (E1) at 1, 70 and 300 K temperatures, respectively. Initially, the optimized 13-cis,15-syn configuration is assumed (Fig. 3(b)). Here $\alpha$ denotes the rotation angle (see Fig. 2). The inset shows the simulation results for the structure in the E1 state at 1 K up to 1 ps.](image-url)
Fig. 6. The proton pathway from the protonated SB (in the 13-cis,15-syn configuration of retinal) to Asp-85 via mediating water molecule (W402): (a) energy surfaces between the water molecule (W402) and Asp-85 at the presence of the 13-cis,15-syn retinal and other water molecules (W401, W406); (b) energy surfaces between the SB and water molecule (W402) due to the presence of Asp-85 group. Here $R_{\text{N-H}}$ defines the distance between the nitrogen atom in SB and the hydrogen atom, as shown in Fig. 5. The computations were performed using the HSEH1PBE functional.

5. Discussion

5.1. Energy redistribution during the photocycle

Let us discuss now how the energy supplied by the photon is redistributed over various degrees of freedom in the transitions between the intermediates shown in Fig. 7, keeping in mind that these transitions are almost resonant with a small dissipation. Immediately after the photon absorption, the energy is entirely in the electronic degrees of freedom of the excited state of the retinal. The excited electronic state cannot relax back to the all-trans ground state because the internal degrees of freedom are not able to accept all the energy of the electronic excitation. Instead, the conformation to the 13-cis form (J to K transition)
allows the excitation energy to be transferred to the strain of the retinal. It should be noted that the part of the energy is still contained in the electronic subsystem, as the ground state energy of the 13-cis isomer is larger than that of all-trans. In the consequent transition between K and L intermediates, the retinal is more relaxed and the energy is redistributed to the protein resulting in bending of the C-helix and thus in modification of the energies of the Coulomb interaction between residues [63, 66].

These modified interactions compensate for the huge acid dissociation constant (pK_a) differences between the SB and Asp-85 (more than 11 and 2.5, respectively, for the bR states) and allow the initial proton transfer between these residues (L to M_1 transition). To make the transfer possible, the pK_a factors of both protonable sites are modified bringing them to the pK_a of about 6.5. Accordingly, the energy delivered to the proton is much less than half of the electron-volt as follows from the initial pK_a difference. The protonation of Asp-85 modifies the electrostatic interactions in the proton release group, so the corresponding proton energy becomes larger than the chemical potential of the extracellular space and the proton can be released (M_1 to M_2 transition). The deprotonation of SB leads to conformational changes (relaxation of the C-helix and bending of B- and F-helices), which, in turn, allow both the increase of the pK_a of SB to 8.3 and the decrease of the pK_a of Asp-96 to 7 [8]. Moreover, it leads to the penetration of water molecules in the space between the Asp-96 residue and SB. As a result, the reprotonation of the SB from Asp-96 via the water bridge becomes energetically favourable (M_2 to N_1 transition). Consequently, the proton energy level of Asp-96 again becomes smaller than the chemical potential of the cytoplasmic side and the Asp-96 residue is reprotonated (N_1 to N_2 transition). Accordingly, in the series of events the bR complex releases the high-energy proton to the extracellular side and takes the low-energy proton from the cytoplasmic side, i.e. the proton is effectively pumped. The energy gain can be estimated from the difference between the pK_a of Asp-96 residue at the initial state (about 11) and the potential of the hydrogen (pH) factor of extracellular space, which is about 0.3 eV for pH = 5. The proton translocations from and to Asp-96 give rise to a new sequence of conformational changes, such as the relaxation of B- and F-helices, and the modifications of electrostatic interactions, so the retinal

![Fig. 7. The photocycle of bR showing the main proton transfer events and the main mechanical transformations of the protein and its environment.](image-url)
Reisomerization to the *all-trans* configuration is now possible (N to O transition). The reprotonation of SB leads to the decrease of the pKₐ of Asp-85 and its proton is slowly transferred downhill to the proton release group (O to bR transition) completing the photocycle.

5.2. General description of energy redistribution processes

Let us discuss now how the obtained results of our calculations of the initial step of proton transfer are related to a general description of the energy redistribution processes. Proton transport correlates with the protein conformational changes at various bR stages during the photocycle, especially with the bending of C- and F-helices (Fig. 7). In particular, the proton transfer to the extracellular side is related to the C-helix, while the proton uptake from the cytoplasmic side is related to the F-helix.

As follows from our quantum chemistry studies, the rotation energy surface cannot explain the initial energy transfer without the account of interaction with the protein helices. Indeed, the protonated 13-cis,15-syn retinal configuration cannot occur without the support of the protein because it is absent in the ground state structure (Fig. 3(a)), while the relevant minima might be present in the first excited state if three structural parts from the bR are taken into account (Fig. 3(b)). Thus, the energy should be evidently translocated to the C-helix for the rearranged positions of three water molecules (Fig. 8(a)), which appear as a result of the isomerization in the vicinity of the protonated Schiff base. This complex dynamics of molecules and protein residuals in the vicinity of the excited retinal could result in irreversible proton transfer by (i) temporarily shifting the OH-group from W402 and subsequently protonating Asp-85 (Fig. 8(b)) and (ii) translocating the proton from the SB group to W402 (Fig. 8(a)). Thus, the initial energy should be transferred to the protein in the initial step.

As a result of such configuration changes, the energy of the electronic states of retinal should be modified as schematically shown in Fig. 8. According to our calculations, the proton should be transferred to Asp-85 (with the help of the water molecule) from the 13-cis,15-syn configuration, which is not of the lowest energy state of the retinal taken alone. To have the retinal fixed in this configuration, we have to postulate that the energy of this state is decreased as a result of the electrostatic

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Fig. 8. Energy model for the retinal electronic excited states and possible excitation energy uptake by the protein of bR. The red lines mark the energy surfaces of the retinal, the green lines correspond to the energy changes of the protein.
interaction with the bent helices. The ground state of the retinal at the 13-cis,15-syn conformation with the account of this interaction should be almost at the same energy level as it is for cis/trans minima. Thus, we should have two similar centres, which differ by the water translocation but with similar structural parts causing different irreversible proton transfer events (Fig. 8).

The energy translocation from the retinal to protein helices in the primary step allows us to qualitatively determine the energy swap in the protein, as shown in Fig. 8. Proton transport is initiated from the middle of the structure by step-by-step pushing and uptaking protons over all the bR proteins resulting in $pK_a$ changes.

6. Conclusions

In conclusion, we have discussed the challenges and approaches to modelling the proton transfer in bR and suggested that the retinal molecule responsible for the light absorption cannot be completely separated from the protein environment. In the spirit of the Marcus transport theory, all the energies of the initial and final states of the transfer event should be taken from the whole system, including the energies of the bent helices, stretched bonds and modified electrostatic couplings to the charged and electric-dipole-possessed residues and water molecules. If the part of the whole system is involved in specific exact or high-level numerical calculations, it does not have to be at its ground state (calculated for this isolated part), as one of the excited states can correspond to the energy minimum for the whole system.

For the illustration, we have discussed molecular dynamics and performed quantum chemistry calculations of the initial proton transfer event in bR, i.e. the deprotonation of SB and the protonation of Asp85 residue, with the retinal isomerization and the energy of the retinal electron state taken into account. We have shown that the proton transport becomes irreversible if the retinal is initially in the 13-cis,15-syn configuration, which is not the ground state. However, this state might correspond to the energy minimum of the whole system including the energy of the modified electrostatic interaction with the residues of the bent C-helix. We have revealed the actual mechanism of the proton transfer, as initially the Asp-85 residue is protonated by the water molecule which is subsequently restored by the SB proton. In some sense, it is comparable to the proton transfer through the whole complex, as it is not the situation when the single proton is transferred from the cytoplasmic side to the extracellular space, but instead, proton transport is initiated from the middle of the structure.

We believe that the outlined approach will offer additional prospects for future research because the restrictions on the initial and final states can be lifted. Correspondingly, molecular dynamics and quantum chemistry calculations can be performed in a much wider range of parameters and allow detailed descriptions of individual transfer events. We also believe that eventually all these individual events will be combined in the unified self-consistent picture of the bR machinery.

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Santrauka

Bakteriorodopsine (BR) vykstanti protono pernaša plačiai nagrinėjama daugeliu eksperimentiniais ir teoriniai metodais. Pastariejį tyrimai paremti ypač detaliais kristalografiniais BR duomenimis. Vis dėlto, nepaisant turimų duomenų, retinalyje vykstanti protono pernaša, iniciuota elektroninio sužadinimo, nera viškai aiški. Naudodami kvantinės chemijos skaičiavimus, nagrinėjome retinalio molekulę, kuri yra fiksuota 13-cis,15-syn konfigūracijos pagrindinėje būsenoje ir susidaro dėl baltyminės aplinkos įtakos. Naudodami šią fiksuotą būseną, nustatėme, kad protono pernaša į asporto rūgštį N r. 85 (Asp-85) vyksta tarpininkaujant vandens molekule, o analogiškoje aplinkoje taip pat dalyvaujama vel protonuojant Šifo bazę. Aptarėme BR vykstančio protonų perdavimo modeliavimo iššūkius ir metodus, analizuodami procesą, kai molekulės elektroninio sužadinimo savybės nulemia baltyminė aplinka.