FTIR Spectroscopy Revealing Light-Dependent Refolding of the Conserved Tongue Region of Bacteriophytochrome

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ABSTRACT: Bacteriophytochromes (BphPs) constitute a class of photosensory proteins that toggle between Pr and Pfr functional states through absorption of red and far-red light. The photosensory core of BphPs is composed of PAS, GAF, and PHY domains. Here, we apply FTIR spectroscopy to investigate changes in the secondary structure of Rhodopseudomonas palustris BphP2 (RpbP2) upon Pr to Pfr photoconversion. Our results indicate conversion from a β-sheet to an α-helical element in the so-called tongue region of the PHY domain, consistent with recent X-ray structures of Deinococcus radiodurans DrBphP in dark and light states (Takala, H.; et al. Nature 2014, 509, 245−248). A conserved Asp in the GAF domain that noncovalently connects with the PHY domain and a conserved Pro in the tongue region of the PHY domain are essential for the β-sheet-to-α-helix conversion.

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Phytochromes are red-light-sensing proteins found in plants, bacteria, cyanobacteria, and fungi that act as photochromic switches activated by distinct wavelengths in the red and far-red regions. In the dark, most phytochromes adopt a red-absorbing state known as Pr and upon light absorption convert to a far-red-absorbing state known as Pfr. The light activation mechanism involves an isomerization process about the C15=C16 double bond of the linear tetrapyrrole, changing its configuration from 15Z to 15E. Their light-sensing module is composed of PAS, GAF, and PHY domains and covalently connects with the PHY domain and a conserved Pro in the tongue region of the PHY domain. A conserved Asp in the GAF domain that noncovalently connects with the PHY domain and a conserved Pro in the tongue region of the PHY domain are essential for the β-sheet-to-α-helix conversion.

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conserved tongue between the Pr and Pfr states. Specifically, in Pr, the tongue region in the PHY domain assumed a loop-two stranded β-sheet conformation, whereas in Pfr, it assumed a loop-α-helix conformation. Questions arise about the significance of this switch in the secondary structure. Does it occur in solution, how is it related to Pfr formation, and which residues are key for the switch to take place? Here, we address these questions through FTIR spectroscopy on a DrBphP homologue, BphP from Rhodopseudomonas palustris (RpBphP).

Sample preparation, site-directed mutagenesis, and FTIR spectroscopy on BphPs were carried out as described before. Figure 2 shows the light-minus-dark FTIR spectra of the RpBphP2 PAS-GAF-PHY photosensory core domain (black), the PAS-GAF chromophore-binding domain (red), the PAS-GAF-PHY D202A mutant (blue), and the PAS-GAF-PHY P465T mutant (green). D202 in RpBphP2 represents a well-characterized, conserved Asp from the PASDIP motif of the GAF domain, while P465 represents a conserved Pro from the PRXSF motif of the PHY domain. The FTIR difference spectrum shows a multitude of positive and negative bands, some of which were assigned before in this protein, RpBphP2. BV-specific bands were identified before on the basis of ultrafast IR experiments on RpBphP2, where optical excitation results in BV-specific signals on femtosecond and picosecond time scales before any signals from the apoprotein arise. Other BV-specific bands were observed and assigned before in Cph1 on the basis of isotope labeling of the chromophore and apoprotein and will be discussed below.

Figure 2. Light-minus-dark FTIR spectra of RpBphP2 PAS-GAF-PHY (black), PAS-GAF (red), PAS-GAF-PHY D202A mutant (blue), and PAS-GAF-PHY P465T mutant (green).

The interpretation of the remaining Pfr−Pr FTIR signals of RpBphP2 PAS-GAF-PHY is made on the basis of earlier results. The 1734 cm⁻¹ band results from BV ring A C1=O, while that at 1704 cm⁻¹ is due to ring D C19=O. The 1689 (−)/1682 (+) cm⁻¹ band is due to amide I (loop/turn) or a propionate COOH, and the 1664 cm⁻¹ band corresponds to loop amide I, as previously assigned in Cph1. The 1612 (+) cm⁻¹ band was previously tentatively assigned to amide I (β-sheet structure) in Cph1. The 1595 cm⁻¹ band is due to the BV C−D methine bridge. The 1563 cm⁻¹ band corresponds to amide II, as previously assigned for Cph1. The 1540 (−) cm⁻¹ band was shown to belong to the BV chromophore but was not interpreted.

To further explore the origin of the prominent 1630 cm⁻¹ (−)/1653 (+) cm⁻¹ signal, we performed FTIR spectroscopy on a shorter construct, RpBphP2 PAS-GAF (also referred to as the chromophore binding domain (CBD)), and on the
RpBphP2 PAS-GAF-PHY D202A mutant. Both proteins are deficient in Pfr formation. Upon photoconversion, the D202A mutant becomes arrested in a Meta-R-like state in which the absorption characteristic of the Pr state is bleached and only a small induced absorption is evident, less red-shifted than that in the Pfr state of wild-type1,13 (Figure 3). The PAS-GAF protein converts to a red-shifted state that absorbs at 741 nm11 and fails to form the fully red-shifted Pfr state of wild-type PAS-GAF-PHY, which absorbs at 753 nm (Figure 3). The FTIR difference spectra of the PAS-GAF (Figure 2, red line) and the D202A mutant (Figure 2, blue line) proteins are very similar. Strikingly, they both mostly lack the negative band at 1630 cm\(^{-1}\). These observations suggest that the signaling mechanism in RpBphP3 might be distinct from canonical BphPs.

The other bands of RpBphP2 PAS-GAF and the RpBphP2 D202A mutant are similar to those observed in wild-type PAS-GAF-PHY. The 1738 (\(-\)) and 1712 (\(-\)) cm\(^{-1}\) bands are assigned to BV C1\(=\)O and C19\(=\)O, respectively. Femtosecond IR experiments showed that in the PAS-GAF domain, the C19\(=\)O band is indeed upshifted with respect to PAS-GAF-PHY.\(^{17}\) The remaining bands are conserved with respect to PAS-GAF-PHY. The 1643 (\(-\))/1656 (\(+\)) cm\(^{-1}\) feature either corresponds to amide 1, where it would correspond to loosening of a helical element in the PAS or GAF domain upon photoconversion, or to C=C stretches of the BV chromophore.\(^{17,21}\) In PAS-GAF-PHY, this feature is present as well and appears additive to the prominent 1630 (\(-\))/1653 (\(+\)) cm\(^{-1}\) feature.

Further evidence on the nature of the prominent 1630 (\(-\))/1653 (\(+\)) cm\(^{-1}\) signal in wild-type PAS-GAF-PHY comes from the RpBphP2 P465T mutant. Pro-465 is a highly conserved residue located at a key position in the tongue region (Figure 1).\(^{5,14}\) The P465T mutant forms a red-shifted state at 746 nm, similar to the PAS-GAF protein, and is hence deficient in Pfr formation (Figure 3). Its FTIR difference spectrum of the P465T mutant (Figure 2, green line) is almost identical to that of the PAS-GAF protein and lacks the 1630 (\(-\))/1653 cm\(^{-1}\) (\(+\)) \(\beta\)-sheet-to-\(\alpha\)-helix switch feature.

We conclude that the \(\beta\)-sheet-to-\(\alpha\)-helix switch observed with FTIR spectroscopy is only present in wild-type RpBphP2 composed of PAS-GAF-PHY domains. It is absent in the protein lacking the PHY domain, the D202A mutant, and the P465T mutant. These observations strongly suggest that the \(\beta\)-sheet-to-\(\alpha\)-helix switch as observed in FTIR difference spectroscopy corresponds to the loop-sheet to loop-helix switch in the tongue region of DrBphP recently observed with X-ray crystallography\(^{7}\) and solidifies the argument that this conformational switch is an integral part of phytochrome signaling in vitro and in vivo. Furthermore, our results indicate that refolding of the tongue is required to stabilize the BV chromophore in the fully red-shifted Pfr conformation.

The conserved Asp-202 is instrumental for the sheet–helix switch. It is likely that the \(\beta\)-sheet conformation is stabilized through the salt-bridge interaction with the conserved Arg-466, which is located in the tongue region. This salt bridge is broken in the Pr state, and Arg-466 is flipped out to the solvent (Figure 1 B).\(^{1}\) Additionally, the conserved Pro-465, part of the tongue and located in the loop adjacent to the double-stranded \(\beta\)-sheet, is conserved in the Pr state and at the beginning of the newly formed \(\alpha\)-helix in the Pfr state, is essential for the sheet-to-helix switch. The proline residue is unique in that its side chain is covalently bonded to the nitrogen atom of the peptide backbone. Consequently, the backbone of Pro cannot form a hydrogen bond, and its N\(\rightarrow\)C\(_{\alpha}\) rotation is rigid. Thus, prolines are frequently found in the first N-terminus turn of an \(\alpha\)-helix where the loss of the H-bond to the imino nitrogen does not cause significant effects.\(^{29}\) Proline and glycine are known to destabilize \(\alpha\)-helices because they disrupt the regularity of the \(\alpha\)-helical backbone conformation. This may be taking place in the Pr conformation of the protein.

Finally, in the unusual bacteriophytochrome RpBphP3, which converts from Pr to a near-red-absorbing state Pnr,\(^{18}\) the amino acid equivalent to P465 in RpBphP2 is actually a threonine (Thr480) (Figure 1 C). To our knowledge, this is the only BphP that lacks a conserved Pro in the PRXSF motif of the PHY domain. In light of the present results, our observations suggest that the signaling mechanism in RpBphP3 might be distinct from canonical BphPs.

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