Blue Light-excited Light-Oxygen-Voltage-sensing Domain 2 (LOV2) Triggers a Rearrangement of the Kinase Domain to Induce Phosphorylation Activity in Arabidopsis Phototropin1*

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Phototropin1 is one of the blue light (BL) receptor proteins in plants and shows BL-dependent kinase activation. The BL-excited light-oxygen-voltage-sensing domain 2 (LOV2) is primarily responsible for the activation of the kinase domain; however, the molecular mechanism by which conformational changes in LOV2 are transmitted to the kinase domain remains unclear. Here, we investigated BL-induced structural changes of a minimum functional fragment of Arabidopsis phototropin1 composed of LOV2, the kinase domain, and a linker connecting the two domains using small-angle x-ray scattering (SAXS). The fragment existed as a dimer and displayed photoreversible SAXS changes reflected in the radii of gyration of 42.9 Å in the dark and 48.8 Å under BL irradiation. In the dark, the molecular shape reconstructed from the SAXS profiles appeared as two bean-shaped lobes in a twisted arrangement that was 170 Å long, 80 Å wide, and 50 Å thick. The molecular shape under BL became slightly elongated from that in the dark. By fitting the crystal structure of the LOV2 dimer and a homology model of the kinase domain to their inferred shapes, the BL-dependent change could be interpreted as the positional shift in the kinase domain relative to that of the LOV2 dimer. In addition, we found that lysine 475, a functionally important residue, in the N-terminal region of LOV2 plays a critical role in transmitting the structural changes to the kinase domain, which is critical for signaling, suitably changing the structure to activate the kinase in response to conformational changes in the adjoining LOV2.

Phototropin (phot)2 is one of the blue light (BL) receptor proteins in plants (1, 2). It plays several important roles to maximize the efficiency of photosynthesis: namely phototropism (3), chloroplast movement (4, 5), stomata opening (6), and leaf expansion (7). Most plants possess two isoforms of phot, designated as phot1 and phot2 (1). Both isoforms redundantly regulate stomata opening (6) and chloroplast accumulation depending on fluence rate of BL (3). However, only phot2 is responsible for the photoavoidance response in chloroplast relocation (4). Regarding the phototropic response, phot2 works under high irradiance condition, whereas phot1 predominantly initiates the response under low irradiance condition (3).

The structure of plant phot, which consists of ~950–1000 amino acid residues and two flavin mononucleotide (FMN) molecules (1), folds into three functional domains (Fig. 1A). The N-terminal half contains two light-oxygen-voltage-sensing domains (LOVs) (designated LOV1 and LOV2) (8, 9), which belong to the PAS (Per-Arnt-Sim) superfamily found in heterodimeric interaction in cellular signaling. Each LOV domain receives BL via the FMN molecule bound to the pocket formed by the characteristic α-helix/β-strand-mixed structure known as the α/β-scaffold (10, 11). The C-terminal half of phot is a serine-threonine kinase (STK) domain classified into group VIII of the AGC family (12). The linker region, which has α-helix lying on the α/β-scaffold in the N-terminal quarter, connects LOV2 and STK (13). The LOV1 domain acts as a site for dimerization of Arabidopsis thaliana (At) phot1 and phot2 (14–17), whereas BL-activated LOV2 is primarily responsible for inducing autophosphorylation of phot (18) and phosphorylation of substrate proteins such as ABCB19 (19), PKS4 (20), and BLUS1 (21) in the downstream signaling pathways.

Structural and spectroscopic studies on phot and its functional domains have focused on the molecular mechanism for how BL-excited phot acquires the kinase activity. Upon BL irradiation, phot undergoes a characteristic photoreaction cycle (22–24). A transient adduct is formed between FMN and a conserved Cys residue (S390 intermediate state with an absorption maximum at 390 nm) (22, 23), which then is thermally broken to the dark state (D450 with an absorption maximum at 450 nm). In the S390 of LOV2, a couple of small structural changes in the side chains of the amino acid residues near FMN are thought to...
Structure and Kinase Activation of Arabidopsis Phototropin 1

be the primary step necessary for the activation of STK (18). Under BL irradiation, the Jα-helix unfolds, and then its interaction with the α/β-scaffold would change (13, 25–27).

To date, we have been investigating the minimum functional unit of phot, LOV2-linker-STK fragment (L2LK) (28, 29), to understand how the conformational changes in LOV2 propagate to STK. For example, the BL-induced rearrangement of LOV2 and STK was found in the L2LKS of At phot2 (30) and Chlamydomonas reinhardtii (Cr) phot (31, 32) through structural studies by using small-angle x-ray scattering (SAXS). Mutation studies demonstrated that amino acid residues critical to the phosphorylation activity of STK are localized around the N-terminal regions of LOV2 (33, 34). From these studies, we envision a scenario whereby the linker and its adjoining region mediate the propagation of small conformational changes in BL-excited LOV2 to STK (Fig. 1B).

In the case of At phot1, mutagenesis studies of phot1 L2LK (P1L2LK) (residues 449–996; Fig. 1A) are in progress to identify the residues necessary for the BL-dependent kinase activity (34, 35). Cys-512 residue in LOV2 is critical for the adduct formation with FMN in the photoreaction cycle. Asp-806 residue in STK is essential for the kinase activity (18). Lys-475 residue in the Aα/Aβ gap of the N terminus of LOV2 and Lys-636 residue in the linker region would have important roles in the BL-dependent kinase activity. Although the arrangement of LOV2 and STK in phot1 remains unknown, the dimer structures of LOV1 and LOV2 are now known (Fig. 1C) (14–17). Therefore, the structural roles of the lysine residues are difficult to ascertain. Here, we study the molecular structures of wild-type and mutated P1L2LK by SAXS to better understand the molecular mechanism responsible for BL-induced kinase activity of phot1.

Results

Photoreaction Kinetics of the Wild-type P1L2LK—The purity of the wild-type P1L2LK was greater than 98% as indicated by the sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) pattern. Size exclusion chromatography (SEC) indicated that P1L2LK probably forms a dimer in solution in the concentration range used in the spectroscopic and SAXS measurements (0.5–2.3 mg ml⁻¹) (Fig. 2A). The absorption spectra of D₄₅₀ and S₃₉₀ states were almost identical to those of the recombinant phot1 LOV2 (24).

Under successive BL irradiation, the amount of accumulated S₃₉₀ depended on the fluence of BL because of the dark (thermal) reversion from S₃₉₀ to D₄₅₀. For SAXS measurements, almost all of the P1L2LK in solution sample must be converted to the S₃₉₀ state under BL irradiation. To estimate the fluence rate necessary to accumulate almost completely S₃₉₀, we measured the amount of S₃₉₀ under BL irradiation by using our custom-made cell holder mounted on the UV-visible spectrometer. This device is limited to measuring the UV-visible spectra under a fluence rate of less than 200 μmol m⁻² s⁻¹ because of the stray light under high fluence rate. Therefore, we measured the amount of S₃₉₀ under a fluence rate of less than 200 μmol m⁻² s⁻¹. Then the fluence rate necessary for the almost complete accumulation of S₃₉₀ was estimated to be 400 μmol m⁻² s⁻¹ by an extrapolation (see Fig. 2B, inset).

Within 120 s after starting BL irradiation of 200 μmol m⁻² s⁻¹, the amount of L2LK in S₃₉₀ reached saturation. The thermal dark reversion took more than 400 s after turning off BL at 293 K (Fig. 2C). After BL irradiation of 200–400 μmol m⁻² s⁻¹, P1L2LK undergoes photoreaction. This result indicated little molecular damage in P1L2LK under the BL irradiation of the fluence rate of 400 μmol m⁻² s⁻¹.

BL-induced Conformational Changes of the Wild-type P1L2LK—Little x-ray radiation damage to the samples was found as verified by very small differences between the absorption spectra and SDS-PAGE patterns of specimens before and after the exposure to x-rays within one round of DK1-BL1-DK2 exposures. To avoid x-ray radiation damage during several rounds of x-ray exposure, SAXS profiles of the three states were recorded through only one round of exposure as done in our previous SAXS studies on the LOV2-linker-kinase fragment of At phot2 (30) and full-length phot of Chlamydomonas (31).
Structure and Kinase Activation of Arabidopsis Phototropin1

For each P1L2LK solution, we sequentially recorded the SAXS profiles of P1L2LK in the order of DK1, BL1, and DK2 states (Fig. 3A). The scattering intensities of BL1 increased in $S < 0.003 \text{ Å}^{-1}$ but decreased in $0.004 < S < 0.010 \text{ Å}^{-1}$ for the DK1 state (Fig. 3A). The SAXS profiles of the DK2 state resembled those of the DK1 state rather than the BL1 state. The profile difference, calculated by subtracting the profile of the BL1 state from that of the DK1 state, is negative in $S < 0.003 \text{ Å}^{-1}$ with a maximum at approximately $S = 0.005 \text{ Å}^{-1}$. The profile difference obtained by subtracting the profile of the BL1 state from that of the DK2 state is positive in $S < 0.003 \text{ Å}^{-1}$ with a minimum at approximately $S = 0.005 \text{ Å}^{-1}$. Because these two cases are almost opposite in sign, the SAXS changes reflecting BL-induced conformational alternation of P1L2LK were probably photo reversible.

When taking into consideration the reciprocity between the length of the scattering vector and the distance between any pairs of large electron density clusters, such as functional domains, the changes in $0.004 < S < 0.010 \text{ Å}^{-1}$ suggest rearrangement of domains separated by $\sim 150 \text{ Å}$. In this regard, differences in the $P(r)$ functions between the DK1 and BL1 states were prominent at $r > 100 \text{ Å}$ (Fig. 3C).

The Guinier plots for SAXS profiles of P1L2LK in the DK1, BL1, and DK2 states could be approximated by straight lines in $S^2 < 40 \times 10^{-6} \text{ Å}^{-2}$ (Fig. 3D). The calculated $C/(S = 0, C)$ and $R^2(C)$ values displayed almost linear concentration dependences (Fig. 3E), indicating their monodispersive properties. The apparent molecular weight estimated from $I(S = 0, C = 0)$ indicated that P1L2LK forms a dimer both in the dark and under BL. This result is consistent with the crystal structure of LOV2-J (17) and the SEC prior to the SAXS measurements. Because the $I(S = 0, C = 0)$ value of the DK1 state is almost the same as that of the BL1 state, the dimer structure is kept under BL irradiation.

The P1L2LK in the BL1 state had an $R_g(C = 0)$ of $48.8 \pm 0.5 \text{ Å}$ that was significantly larger than that of either the D1K date ($42.9 \pm 0.5 \text{ Å}$) or the DK2 state ($43.5 \pm 0.5 \text{ Å}$). This increase in $R_g(C = 0)$ suggests an expansion and/or hydration changes in P1L2LK under BL. The slightly greater $R_g(C = 0)$ of the DK2 state than that of the DK1 state indicates that the thermal reversion of the overall structure proceeded more slowly than did the structural relaxation around the FMN monitored in the absorption spectra (Fig. 2B).

**Molecular Shape of the Wild-type L2LK—**Each molecular shape, restored by applying the GASBOR program to the observed SAXS profile, gave a theoretical scattering curve that closely matched those observed with $\chi^2$ values of 2% (Fig. 4A). Following the principal component analysis (PCA) and K-means clustering analysis, the molecular shapes of DK1, BL1, and DK2 were calculated from 472, 380, and 307 restored models, respectively.

The averaged molecular shape of P1L2LK in the DK1 state appeared as two bean-shaped lobes in a twisted arrangement along the long molecular axes (Fig. 4B). Its dimensions are 170 Å long, 80 Å wide, and 50 Å thick. The molecular shape of the BL1 state was slightly different from that of the DK1 state. The two lobes were elongated and simultaneously tilted by $\sim 30^\circ$ against the 2-fold symmetry axis of the dimer. Then the dimen-
sion along the molecular axis was enlarged by ~20 Å from that of the DK1 state. The molecular shape and dimensions of P1L2LK in the DK2 state are more similar to those of the DK1 state than of the BL1 state.

Structure Changes of the Mutated P1L2LK under BL—Structural roles of Lys-475 and Lys-636 residues located at the interface between LOV2 and STK were studied through SAXS for K475A/D806N and K636A/D806N double mutants. K475A and K636A mutations cause reduction of the kinase activity of STK (34, 35). D806N mutation results in the loss of the phosphorylation activity of STK because the side chain of the Asp-806 residue is expected to stabilize the position of a magnesium ion necessary for the ATPase activity as well as various kinases (36). Therefore, the D806N mutation was introduced to avoid heterogeneity of P1L2LK in regard to phosphorylation in the purification (29). As a reference, the C512A/D806N mutant was used to examine whether BL irradiation is essential for structural changes because C512A mutation results in the loss of adduct formation capability under BL irradiation. Prior to SAXS measurements, the dark reversion kinetics of D806N, K475A/D806N, and K636A/D806N mutants were determined through UV-visible absorption measurements.
Structure and Kinase Activation of Arabidopsis Phototropin1

Prior to the x-ray measurements, the relaxation times of the $S_{390}$ states of D806N, K475A/D806N, and K636A/D806N were determined by measuring their UV-visible absorption. The relaxation time in D806N was 93 s, which is nearly the same as that of the wild-type (91 s). This result indicates that the mutation in STK has little influence on the photoreaction of LOV2 as well as in the case of LOV2-linker-kinase of At phot2 (30). In contrast, the K475A and K636A mutations yielded relaxation times of 69 (34) and 78 s (35), respectively.

We measured SAXS of four mutants in which amino acid residues important for phot activity were substituted to alanine (Fig. 5A). We qualitatively examined whether BL-induced changes in structure of the mutated P1L2LK were similar to that of the wild-type P1L2LK by comparing the difference in SAXS profiles between DK1 (DK2) and BL1 (Fig. 5B) with those of the wild type (Fig. 2B).

The SAXS changes in D806N mutant were similar to those of the wild type (Fig. 5B), suggesting this mutation had little influence on the BL-induced conformational changes. The D806N mutation is only critical for ATPase activity of STK as well as D720N mutation in the phot2 LOV2-linker STK fragment (30). Therefore, SAXS measurements for double mutants with D806N and the other single mutations reflect the influence of the single mutation on the structure and function of P1L2LK.

Regarding the C512A/D806N mutant, which lacks the capability to form $S_{390}$ under BL, little profile change around the single mutation on $S_{390}$ decreased with a half-life time of 239 s (relaxation time of 345 s) under the approximation of single exponential decay. Regarding the absorption change, the decay of $S_{390}$ proceeded significantly faster than the deoligomerization of K475A/D806 double mutant. The decrease in the $S_{390}$ half-life time was 89 s (decay constant of 128 s), which is similar to the decay speed in a previous report (34). Therefore, the decay of $S_{390}$ proceeded significantly faster than the deoligomerization of K475A/D806 double mutant. The decrease in the scattering intensity depends on the dissociation of oligomers, probably caused by the relaxation of the BL-induced conformational changes. It should be noted that this series of 14 SAXS measurements was conducted with attenuated x-ray beam to avoid radiation damage. In fact, the absorption and the SDS-PAGE patterns after the x-ray exposures displayed little differences from those before the measurement.

The K636A mutation in the middle of the linker region resulted in the reduction of the BL-dependent kinase activity to 30% of the wild type (35). The similarity of profile differences among the wild-type P1L2LK and K636A/D806N and D806N
mutants suggests that the BL-induced rearrangement of functional domains is necessary for kinase activity (Fig. 5, A and B).

**Discussion**

The present study on P1L2LK has revealed that BL-excited LOV2 causes global conformational changes detectable by SAXS. Based on the present results, we now discuss the arrangement of functional domains and the structural roles of lysine residues composing the A’α/Aβ and linker regions for the BL-induced kinase activity of P1L2LK.

Arrangement of the Functional Domains in P1L2LK—The present SAXS study and the SEC measurement revealed the dimeric association of P1L2LK. Here, we discuss the arrangement of LOV2 and STK domains in the restored molecular shapes of P1L2LK dimer (Fig. 6A). For the molecular shape of DK1, the crystal structure model of LOV2-Jα dimer (17) fit to the center part of the molecular shape well. In addition, two homology models of STK can be placed in the bean-shaped lobes such that the N-terminal domain of STK faces both the Jα-helix and the A’α/Aβ gap. For BL1, the LOV2-Jα subunits...
Structure and Kinase Activation of Arabidopsis Phototropin 1

Lys-475 of the A'α/αβ gap region. In contrast, both lysine residues are difficult to digest in the dark (34). Taken together, the BL-induced conformational changes in the α-helix and A'α/αβ gap likely drive the positional shifts of STK (Fig. 6B). In the proposed arrangement of the functional domains, it is difficult for each STK to phosphorylate the other STK in the dimer.

The rearrangement of LOV2 and STK under BL is observed in monomeric At phot2-LOV2-linker-STK (30) and in monomeric Cr phot (31). In the case of Cr phot, the residues composing the α-helix and adjoining region are known to be critical for the BL-dependent kinase activity (33). The rearrangements of LOV2 relative to STK observed in P1L2LK are similar to those observed in At phot2-LOV2-linker-STK and Cr phot under BL. Although the quaternary structures and amino acid sequences are different among the three phots, molecular mechanisms in the BL-induced rearrangements of LOV2 and STK would be similar among them. To discuss further these molecular mechanisms, we must study the roles of amino acid residues composing the interface between LOV2 and STK critical for transmitting BL-induced conformational changes in LOV2 to STK.

The Roles of Lys-475 and Lys-636 in BL-induced Kinase Activation—The wild type and D806N mutants displayed photoreversible changes in their SAXS profiles (Fig. 5B), whereas the C512A/D806N mutant did not. Thus, the photoreversible change around S = 0.005 Å⁻¹ of the SAXS profiles could be useful to examine whether BL-induced conformational changes occur in mutated P1L2LK.

Here we speculate the mechanism in oligomer formation/deformation of K475A/D806N mutant. In our structural model proposed (Fig. 6A), K475A residue located at the interface between LOV2 and STK would contribute to the drastic changes of surface properties under BL irradiation. The conformation of K475A is thought to depend on the small conformational changes in the αC-β scaffold of LOV2. Then the surface structures change to increase intermolecular interactions among P1L2LK in specimens. The loop structures in the linker region may contribute to the association of P1L2LK under BL, but the structure of this region is unclear due to the lack of the crystal structure of P1L2LK.

The different SAXS profiles and the lack of BL-induced kinase activity in the K475A mutant suggest that the K475A mutation probably causes conformational changes to prevent the propagation of a structural signal from LOV2 to STK. The photoreversible association/dissociation implies that the Lys-475 mutation induces conformational changes different from that of the wild type. Therefore, Lys-475 and the adjoining amino acid residues may function as a switch that regulates the propagation of structural changes from LOV2 at the S390 state to STK.

The profile difference of the K636A/D806N mutant was similar to those observed for the wild type and D806N mutant. In addition, the Lys-636 mutation only reduced the BL-dependent kinase activity. Therefore, the capability to rearrange functional domains under BL is kept in the K636A mutant. However, the reduced kinase activity implies that lysine is more functionally suitable than alanine at position 636 to correctly transmit the structural changes from LOV2 to STK.

In the transmission of conformational changes from LOV2 to STK, Lys-475 is critical, whereas Lys-636 may play a role in

![FIGURE 6. A, arrangement of the LOV2-Jα dimer (Protein Data Bank code 4HHB) and the two STKs in the molecular shapes of DK1, BL1, and DK2 (white mesh). The homology model of STK is built from the crystal structure of cAMP-dependent protein kinase in the unliganded state (Protein Data Bank code 1J3H) as a template. The dashed circles indicate the interface between LOV2 and STK. The dashed lines in the perpendicular view of the BL1 model indicate the molecular axes of STKs. In this arrangement, it is difficult for one subunit to make contact with the other subunit for intramolecular phosphorylation. B, schematic illustration of the BL-induced conformational changes of L2LK based on the experimental results presented here.](image-url)
Structure and Kinase Activation of Arabidopsis Phototropin 1

Experimental Procedures

Sample Preparation—We prepared wild-type P1L2LK, one single mutant (D806N), and three double mutants (C512A/D806N, K475A/D806N, and K636/D806N), all of which have histidine tags attached to their N-terminal end with the expression vectors reported previously (28, 34, 35). The D806N mutation for the double mutants was introduced through PCR-based site-directed mutagenesis with the following two primers: 5′-GGGATACCTTATGTCGAAATTCTGTTCTATGCTG-3′ and 5′-GGGATACCTTATGTCGAAATTCTGTTCTATGCTG-3′. The vectors for the three double mutants were prepared from the vector used for D806N mutation following a procedure already reported (28).

The P1L2LK peptides were overexpressed in Escherichia coli BL21 (DE3) and purified by means of nickel affinity chromatography and SEC as described previously (28). Because SAXS measurements require sample solutions with a high purity of 95% or greater, each peptide was purified further through anion exchange chromatography using a Mono Q column (GE Healthcare) (30). The purity of each specimen was examined by SDS-PAGE using Coomassie Brilliant Blue staining. To estimate the molecular weights of the purified fragments, SEC was used with a Sephacryl S-200 column (GE Healthcare). Purified P1L2LK was stored in a buffer containing 20 mM Tris-HCl, pH 7.8, 100 mM NaCl, 10% (w/v) glycerol, and 1 mM EGTA.

Measurement of UV-Visible Absorption Spectra—Absorption spectra and the time course of absorption changes during thermal decay of S390 to D450 were measured at 293 K using a U-3310 spectrophotometer (Hitachi-High Technologies, Japan). The sample chamber of the spectrophotometer was modified to measure the spectrum of sample solution under continuous BL irradiation at a fluence rate of less than 200 μmol m⁻² s⁻¹. For a high fluence rate of greater than 200 μmol m⁻² s⁻¹, it is difficult to measure the spectrum under continuous BL irradiation because stray light significantly increasing inside the sample chamber prevents the measurement. When carrying out the scan measurement just after shutting out BL, the scanned spectrum contains the spectrum from P1L2LK fragment in the thermal recovery process. Thus, we estimated the spectrum of S390 by extrapolating those recorded under a fluence rate of 200 μmol m⁻² s⁻¹.

The P1L2LK solutions, at a concentration of 37 μM in 20 mM Tris-HCl, 500 mM NaCl, 1 mM EDTA, 10% (w/v) glycerol, pH 7.8, were measured after a centrifugation at 100,000 g for 30 min at 277 K. A blue photodiode illuminator (λ_max = 460 nm) was used for the light activation of fragments. The fluence rate of the photodiode was changed in the range of 10–200 μmol m⁻² s⁻¹ to estimate the amount of S390 intermediate in sample solutions under continuous BL irradiation. The time course of absorption changes under both BL illumination at a fluence rate of 190 μmol m⁻² s⁻¹ and in darkness was monitored at 447 nm (28).

SAXS Measurement—The SAXS measurements were performed at BL45XU of SPring-8. The wavelength of the x-ray was 1.000 Å, and the camera distance was ~2500 mm. The temperature of a specimen cell was maintained at 293 K. The SAXS patterns of wild-type and mutant P1L2LK were optimally transmitting these conformational changes to STK. Through SAXS measurements of the other mutated P1L2LKs, the structural roles of residues should be investigated further based on the kinase activity and SAXS changes under BL.

In the beamtime at the synchrotron facility, the longest time to measure the thermal reversion is 15 min for each sample solution. To measure how long the thermal reversion takes, management of the measurement schedule should be taken into consideration in the near future.

Dimerization of Phot1—The present SAXS evidence as well as the quaternary structures of LOV1 (15, 16) and LOV2-1α (17) implies that full-length Phot1 probably exists as a dimer. However, the arrangement of the LOV1 dimer relative to the LOV2 dimer is still unknown. To date, the molecular shape of monomeric full-length Chlamydomonas phot is known (31). In Chlamydomonas phot, LOV1 and LOV2 likely have direct contacts, probably because the number of amino acid residues linking LOV1 and LOV2 is ~80. In contrast, the corresponding linker in phot1 is composed of approximately twice the number of amino acid residues as that of linker in phot1 is composed of approximately twice the number of amino acid residues linking LOV1 and LOV2 in Chlamydomonas phot. The arrangement of the LOV1 dimer relative to the LOV2 dimer and two STKs is difficult to speculate (Figs. 1A and 7). However, the N-terminal region composed of 200 amino acid residues may contact the active site cleft of STK within a subunit (Fig. 7) because the phosphorylation sites of Phot1 known so far are predominantly located in the N-terminal region (37).

To characterize the arrangement of LOV1 further, a SAXS study or crystal structure analysis of full-length Phot1 is necessary, although we have not yet succeeded in the preparation of full-length Phot1. In this regard, it is of interest to study whether monomeric Phot1 can activate the signaling system in vivo through modifying the residues necessary for dimerization for LOV1 (15) and LOV2 (17).
recorded by a PILATUS 300K-W. Each successive SAXS pattern of the wild type was accumulated as 12 time-resolved frames at a 5-s exposure, whereas those of the mutants were accumulated as 20 frames at a 1-s exposure. Intensities of the incident x-ray beam were kept constant during the experiments by the top-up operation of the SPring-8 storage ring.

For each specimen solution, SAXS patterns were measured sequentially in the dark (the DK1 state), under BL of 450 μmol m−2 s−1 after preirradiation for 5 min (the BL1 state), and then in the dark after a dark adaptation for 15 min (the DK2 state). By comparing SAXS profiles of the DK1 state and the BL1 state, we could examine whether BL-induced conformational changes occur in P1L2LK under the BL1 state. In addition, by comparing SAXS profiles from the BL1 state and the DK2 state, structural changes during the thermal reversion can be monitored.

When recording the SAXS profiles in the thermal reversion of the K475A/D806N mutant, the x-ray beam was attenuated to 6% of the original flux intensity by using aluminum foil with a thickness of 800 μm. This attenuation is necessary to reduce radiation damage during successive exposures to x-rays. The specimen solution was irradiated by BL for 40 s. Then the SAXS profile was recorded for 30 s as time-resolved frames of a 10-s exposure by using a PILATUS 3X2 M detector. The schedule of x-ray exposure is given under “Results.”

SAXS Analysis—Each two-dimensionally recorded SAXS pattern was reduced to a one-dimensional profile through circular averaging from which the background scattering of buffer solution was subtracted. The Guinier approximation (38) was applied to the SAXS profiles. The scattering intensity I(S, C) at a scattering vector S and a protein concentration C is expressed using the forward scattering intensity, I(0, C), and the radius of gyration of the solute, R_g(C), as follows.

\[ I(S, C) = I(0, C) \exp \left[ -\frac{4\pi^2}{3} R_g(C)^2 S^2 \right] \]  
(Eq. 1)

where λ and 2θ are the x-ray wavelength and the scattering angle, respectively. Under dilute conditions, the concentration dependences of I(S = 0, C) and R_g(C)^2 were approximated as follows.

\[ \frac{I(S = 0, C)}{I(S = 0, C)} = \frac{K C}{M_w} + 2 A \cdot C \]  
(Eq. 3)

\[ R_g(C)^2 = R_g(C = 0)^2 - B \cdot C \]  
(Eq. 4)

where K is a geometrical constant in the SAXS measurement, M_w is the apparent molecular weight of protein, A is the second virial coefficient, and B depends on the interprotein interactions (36). Assuming a partial specific volume of 0.74 cm^3 g−1 for soluble proteins, the M_w of solute is determined by reference to I(S = 0, C = 0) of hen egg white lysozyme serving as a reference. The pair correlation functions P(r) were calculated using the program GNOM (39).

Molecular Shape—The molecular shapes of P1L2LK were predicted by applying the ab initio structure determination program GASBOR (40) to the x-ray scattering profiles in S < 0.04 Å−1. This program minimizes the discrepancy between the experimental (I_exp(S)) and the calculated (I_model(S)) scattering profiles in retaining compact packing of the dummy residues representing the molecular structure. The discrepancy between the experimental and calculated profiles is monitored using the following equation.

\[ \chi^2 = \frac{N}{N - 1} \sum_{j=1}^{N} \left[ \frac{(I_{exp}(S_j) - K I_{model}(S_j))}{\sigma(S_j)} \right]^2 \]  
(Eq. 5)

where N is the number of data points, K is a scale factor, and \sigma(S_j) is the statistical error of I_exp(S_j) at the scattering vector S_j.

The optimum number of dummy residues is that which gives the smallest \chi^2 values as an average. This was searched for by varying the number between 450 and 650 per subunit with a step of 25. For each number, the GASBOR calculations were conducted independently 20 times, each time starting from different models. For the optimum number of dummy residues, GASBOR calculations began from 576 different models for each state. Calculations were performed on a supercomputer system with 576 cores of Intel Xeon CPU X5690 (3.7 GHz per core).

Each model shape is expressed as an assembly of voxels of 4 × 4 × 4 Å^3 for subsequent analyses. After superimposition, all restored molecular shapes are classified into several groups by applying PCA followed by K-means clustering. In PCA, molecular shapes in a multidimensional space were projected onto the plane spanned by the first and second principal components. After applying the K-means clustering, we obtained the best molecular shape as the average for models composing the largest class. The details of the analysis will be published elsewhere. Finally, the molecular shapes are conveyed as density maps of the dummy residues in 4 × 4 × 4 Å^3 cubes. It should be noted here that the mirror images of the molecular models could also explain the measured scattering profiles. The molecular models are illustrated using PyMOL (41).

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