Water-stress induced downsizing of light-harvesting antenna complex protects developing rice seedlings from photo-oxidative damage

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The impact of water-stress on chloroplast development was studied by applying polyethylene glycol 6000 to the roots of 5-day-old etiolated rice (Oryza sativa) seedlings that were subsequently illuminated up to 72 h. Chloroplast development in drought environment led to down-regulation of light-harvesting Chl-proteins. Photosynthetic proteins of Photosystem II (PSII) and oxygen evolving complex i.e., Cytb559, OEC16, OEC23 and OEC33 as well as those of PSI such as PSI-III, PSI-V, and PSI-VI, decreased in abundance. Consequently, due to reduced light absorption by antennae, the electron transport rates of PSII and PSI decreased by 55% and 25% respectively. Further, seedling development in stress condition led to a decline in the ratio of variable (Fv) to maximum (Fm) Chl a fluorescence, as well in the quantum yield of PSII photochemistry. Addition of Mg²⁺ to the thylakoid membranes suggested that Mg²⁺-induced grana stacking was not affected by water deficit. Proteomic analysis revealed the down-regulation of proteins involved in electron transport and in carbon reduction reactions, and up-regulation of antioxidative enzymes. Our results demonstrate that developing seedlings under water deficit could downsize their light-harvesting capacity and components of photosynthetic apparatus to prevent photo-oxidative stress, excess ROS generation and membrane lipid peroxidation.
Water-stress is known to affect electron transport from water to NADP. Water-stress has been shown to damage the oxygen-evolving complex of PSII, as well as both the PSI and PSII reaction centers. Further, 30% PEG 6000 treatment of mature rice plants results in a reduction of photosynthetic electron transport. Also, water-stress-induced stomatal closure and decreased stomatal conductance limit CO₂ availability that results in diverting electrons from the photosynthetic electron transport chain to molecular oxygen generating ROS (O₂⁻) at the end of PS. At the same time, energy transfer from triplet state of PSII reaction center chlorophyll a, in its excited state and even from antenna chlorophylls (Chl), in their triplet state, to oxygen generates singlet oxygen (O₂) that often damages thylakoid membranes.

Rice seeds germinate beneath the soil and the seedlings grow in near-darkness till they emerge to the surface (skotomorphogenesis). During skotomorphogenesis, seedlings do not synthesize Chl, since the Chl biosynthesis pathway enzyme protochlorophyllide oxidoreductase (POR) (EC 1.3.3.1) requires light to photo-transform Phlide to Chlide. Therefore, in rice, the differentiation of etioplast to chloroplast does not take place in darkness. Upon exposure to light, as the seedlings grow, Chl biosynthesis and the associated green-up process transform the etioplasts into chloroplasts. These developing seedlings are very often exposed to drought, due to scarcity of rainfall. These developing seedlings are able to withstand water-stress for a long time and rebound upon rain fall. Therefore, it is important to understand its underlying mechanism of tolerance to water-stress.

Photosynthetic responses of mature plants and developing greening seedlings to water-stress are fundamentally different. In mature leaves, functional photosynthetic complexes are already formed and water-stress induces generation of ROS due to excess light absorption, which affect the photosynthetic apparatus. However, in water-stressed developing seedlings, there is the possibility to down-regulate Chl biosynthesis and downsize the synthesis and assembly of light-harvesting complexes of PSI and PSII, and to adapt plants not to absorb excess light, which is harmful. In our previous study, we have shown that PEG-induced water-stress during early rice seedling development minimizes Chl biosynthesis. Active photosynthesis complexes are formed by the assembly of apo-proteins and incorporation of chlorophyll in them. Therefore, reduced Chl synthesis in water-stressed developing seedlings is likely to affect chloroplast biogenesis. PEG-6000 has long been utilized as a reliable osmoticum for the simulation and constitutive maintenance of water-stress in plants under laboratory conditions, due to its property to act as a non-penetrating osmotic agent that decreases water potential of the cells. To understand the mechanism of relative tolerance of developing seedling to drought, we have studied the impact of water-stress on the photosynthetic apparatus during early seedling growth, as they emerge from skotomorphogenesis beneath the soil to photomorphogenesis stage above the soil.

Results
To understand the impact of water-stress on chloroplast biogenesis during the early photomorphogenesis, 5-d-old etiolated rice seedlings were treated with polyethylene glycol (PEG 6000) for 16 hours prior to their transfer to continuous cool-white fluorescent (plus incandescent) light (100 µmol photons m⁻² s⁻¹). In this paper, we have analyzed the structure and function of chloroplasts, ROS production, and associated processes to assess the role of water-stress on early seedling development.

Impact of Water-Stress on Chloroplast Development. Chl a fluorescence measurements. Fo, Fm and Fv/Fm ratio were monitored after water-stress treatment of etiolated rice seedlings for 72 h with 40 mM and 50 mM PEG, in the presence of light (100 µmol photons m⁻² s⁻¹). Seedlings were kept for 20 min in darkness (see Materials and Methods) before the initial (Fo), and the maximum (Fm), fluorescence was measured. The minimum fluorescence Fo decreased by 11% and 21% in the seedlings treated with 40 and 50 mM PEG compared to the controls, while Fm decreased by 32% and 63% respectively (see Fig. 1A).

The ratio Fv/Fm = (Fm – Fo)/Fm, a “proxy” of the maximum quantum yield of PSII photochemistry, declined in water-stressed samples; Fv/Fm calculated after 72 h of greening decreased by 10% and 22% in 40 and 50 mM PEG-treated seedlings (Fig. 1A).

Quantum yield of PSII (ΦPSII) and non-photochemical quenching (NPQ) during 300 s illumination with actinic light. As expected, ΦPSII declined at higher light intensity. Compared to the controls, ΦPSII, at the highest light intensity (~275 µmol photons m⁻² s⁻¹), was reduced by 33% and 58% in samples treated with 40 mM and 50 mM PEG (Fig. 1B). On the other hand, the NPQ ((Fm – Fm’)/Fm’) increased in response to increase in light intensity. It was higher in 40 mM and 50 mM PEG-treated seedlings than in controls by 21% and 29% under the highest (~275 µmol photons m⁻² s⁻¹) light intensity used (Fig. 1C).

Room temperature (298K) Chl a fluorescence spectra. Chl a fluorescence emission spectrum of thylakoid membranes isolated (after 72 h greening) had a peak at 684 nm; this is mostly, from PSI antenna. Upon the addition of Mg²⁺ in the suspension medium, this peak increased in the control as well as in the water-stressed samples (Fig. 2A). In water-stressed samples, the fluorescence intensity at 684 nm was substantially lower both in the presence and the absence of Mg²⁺, compared to the controls.

Low temperature Chl a fluorescence spectra. Low temperature (77 K) fluorescence emission spectra monitored after 72 h of light exposure are shown in Fig. 2B. Thylakoid membranes of control samples, suspended in a low salt medium (0 mM Mg²⁺), had a peak at 686 nm due to PSI and at 740 nm due to PSII; usually, PSII shows two emission bands at ~684 nm (from light-harvesting complex) and at 695 nm (from CP47), although a single peak has been observed in some cases (e.g. in Gonyaulax polyedra). Fluorescence emission spectra, shown here, were normalized at 686 nm. In water-stressed samples, the emission peak of PSI shifted from 740 nm to 738 nm. Upon addition of Mg²⁺ (4 mM), the PSI fluorescence emission at 740 nm (normalized at 686 nm; PSII) decreased in both the control and the water-stressed samples.
Figure 1. Chlorophyll a fluorescence measurements of control and water-stressed rice seedlings after 72 h of greening. (A) F₀, Fₘ, Fₚ/Fₘ = (Fₘ − F₀)/Fₘ, (B) “Operating” quantum yield (Φ) of PSII and (C) Non-photochemical quenching (NPQ). Five-day old etiolated seedlings were treated with 40 mM (or 50 mM) PEG 6000 dissolved in half strength MS nutrient soln., 16 h prior to their illumination with cool white fluorescent + incandescent light (100 μmol photons m⁻² s⁻¹), at 28 °C. Chl a fluorescence based parameters were measured with a PAM 2100 fluorometer. “Operating” PSII quantum yield (Φ) and NPQ were calculated as (Fₘ' − Fₜ)/Fₘ' and (Fₘ − Fₘ')/Fₘ' respectively. Each data point is the average of three replicates. The error bars represent standard deviations (SD). ANOVA: *p < 0.05; **p < 0.01; and ***p < 0.001.

Figure 2. Fluorescence emission spectra of thylakoid membranes isolated from leaves of control and water-stressed rice seedlings. (A) Room temperature and (B) Low temperature (77K) fluorescence spectra (E₄₄₀) of thylakoid membranes (3 μg Chl) suspended in 5 mM Hepes-NaOH buffer containing 0 or 4 mM MgCl₂ (pH 7.5), after 72 h of stress treatment. Seedlings were treated as in Fig. 1. Fluorescence emission spectra were recorded in ratio mode in a photon counting SLM-AMINCO 8000 spectrofluorometer. For 77 K spectral measurements, excitation and emission slit widths were set at 4 nm. For room temperature spectra, the excitation and emission slit widths were set at 8 nm and 4 nm, respectively. The room temperature spectra were corrected for photomultiplier tube response. Rhodamine B was used in the reference channel as a quantum counter. A tetraphenylibutadiene (TPD) block was used to adjust the voltage to 20000 counts per second in the sample as well as in the reference channels at excitation and emission wavelengths of 348 nm and 422 nm, respectively. Fluorescence spectra were measured three times and identical results were obtained.
PSII, PSI and the whole chain electron transport in isolated thylakoid membranes. To probe further results obtained from Chl a fluorescence measurements, we monitored partial PSII and PSI electron transport reactions, as well as the whole photosynthetic electron transport chain in thylakoid membranes isolated from control and water-stressed seedlings.

PSII activity. The partial PSII electron transport, which was measured polarographically as light-driven electron transport from H₂O to p-phenylenediamine (PD), increased as the greening process of etiolated control seedlings progressed (Fig. 3A). Further, in 50 mM PEG-treated seedlings, PSII activity was reduced by 35% and 55% after 48 h and 72 h of greening compared to that in the controls.

PSI activity. The partial PSI electron transport, which was measured polarographically as light-driven electron transport from ascorbate/DCIP to methylviologen (MV), increased in response to chloroplast development during light exposure. Similar to PSII, PSI electron transport decreased in water-stressed seedlings, compared to controls, but to a lower extent than that of PSII; i.e., by 20% and 25% after 48 h and 72 h of greening compared to that in the controls.

The whole (photosynthetic) electron transport chain. The whole chain electron transport through PSII and PSI, which was measured polarographically as light-driven electron transport from H₂O to methylviologen, was reduced by 35% after 48 h and 65% after 72 h of greening in stressed seedlings compared to control (Fig. 3C).

Light saturation curves of PSII and PSI electron transport reactions. PSII. To further ascertain if the inhibition of PSII reaction was due to reduction in the quantum yield of PSII photochemistry measured in limiting light intensities or it was in light saturated electron transport, we measured the rate of PSII reaction as a function of different light intensities, using thylakoid membranes isolated from the control as well as from water-stressed seedlings, after 72 h of greening. The dependence of PSII activity on light intensity showed typical saturation kinetics (Fig. 3D). Both the initial slope at limiting light intensities as well as light-saturated electron transport, were affected in PEG-treated seedlings. As compared to the control thylakoids, the percent inhibition of PSII reaction in water-stressed thylakoids was almost constant (nearly 50%) at all the light intensities used.

Figure 3. Photosynthetic polarographic measurements of thylakoid membranes isolated from control and water-stressed rice seedlings. (A) PSII, (B) PSI and (C) whole chain activities after 48 h and 72 h of greening, and light saturation curve of (D) PSII and (E) PSI reactions after 72 h of greening in thylakoid membranes isolated from control and water-stressed seedlings. Seedlings were treated essentially as in Fig. 1. Oxygen evolution/uptake by thylakoid membranes was measured by Oxy Lab, Hansatech. Thylakoids, equivalent to 20 µg of Chl, were used for each reaction. PSII, PSI and whole chain activities were measured as electron transport from H₂O to p-phenylenediamine (PD), Ascorbate/DCIP couple to methylviologen (MV) and H₂O to MV, respectively. For measurement of light saturation curve, different light intensities were obtained from a tungsten lamp using neutral density filters. Each data point is the average of three replicates. The error bars represent SD. ANOVA: *p < 0.05; **p < 0.01; and ***p < 0.001.
(Fig. 3D). The Eadie plot (i.e., the rate of oxygen evolution vs rate of oxygen evolution/light intensity in terms of % saturation) showed a straight line characterized by the equation $y = -28.73x + 159.1$ ($R^2 = 0.948$) for the control seedlings and $y = -33.05x + 80.65$ ($R^2 = 0.917$) for the water-stressed seedlings (insets in Fig. 3D). Both the intercepts on the abscissa and on the ordinate were reduced by nearly 50%.

**PSI.** In the PSI case, both the initial slope of electron transport rate at limiting light intensity as well as high light intensity i.e., for saturated electron transport rate, were reduced almost equally by ~30% (Fig. 3E). The Eadie plot showed a straight line with an equation $y = -27.62x + 1028$ ($R^2 = 0.983$) for the control seedlings and $y = -31.03x + 739.8$ ($R^2 = 0.979$) for the water-stressed seedlings (inset Fig. 3E). Intercepts on the abscissa and on the ordinates were reduced by nearly 30%.

**Immunoblot analysis of photosynthetic proteins.** To understand the mechanism of water-stress induced down-regulation of photosynthesis during the development of the photosynthetic apparatus in the rice seedlings, we performed immunoblot analysis of proteins involved in PSII, PSI and the inter-system electron transfer.

**PSII.** Most of proteins/pigment-protein complexes of PSII increased upon illumination of the seedlings. The abundance of cyt b559, an intrinsic membrane protein intimately associated with PSII reaction center, declined by 58% and 52% from that in the controls, in the developing rice seedlings after 24 and 72 h of water-stress treatment (Fig. 4). Further, light-harvesting pigment-protein complexes associated with PSII i.e., Lhcb2 and Lhcb1, were reduced by 30% and 63.4% after 24 h of water-stress treatment, and by 30% and 64% after 72 h treatment (Fig. 4).

**Oxygen Evolving Complex, OEC.** Most of the oxygen evolving complex proteins were severely reduced in the rice seedlings subjected to water-stress during chloroplast development. The OEC16 and OEC23 were reduced by ~75% and 66% respectively in water-stressed seedlings (Fig. 4) after 24 h of stress. Similarly, at this time point, other major OEC protein i.e., OEC33 was reduced by ~50% (Fig. 4).
Cytochrome b6/f. The protein abundance of cyt f, as well as the subunit IV of cyt b6/f complex declined by 4% and 15%, and by 40% and 7% after 24 h and 72 h of water-stress respectively during chloroplast biogenesis (Fig. 4).

PSI. In water-stressed seedlings, the protein abundance of the PSI subunit III (PsaF, 22 kD) was reduced by ~5%, while protein expression of the subunit V (PsaK, 17 kD) was severely reduced by ~80% after 24 h/72 h treatment (Fig. 4). Further, the PSI subunit VI (PsaH, 11 kD) was reduced by 88% and 67%, due to 24 h and 72 h water-stress. In contrast, PSI subunit IV (PsaE, 11 kD) had increased by 4 and 5 fold in water-stressed seedlings (Fig. 4). The PSI antenna proteins Lhca1 and Lhca4 were reduced by 75% and 46% after 24 h of water-stress treatment, and by 42% and 55% in seedlings water-stressed for 72 h.

Ultrastructure of chloroplast thylakoids. After 24–72 h of greening, both thylakoids and grana were well developed and contained starch granules in control samples. However, chloroplasts in the water-stressed seedlings had swollen thylakoids, disintegrated granal organization (mostly after 72 h) and fewer starch granules (Fig. 5).

H2O2 content. H2O2 content remained approximately the same in both water-stressed and the control rice seedlings, after 24 h of greening (Fig. 6A); however, after 72 h of greening, H2O2 content increased by 23% as compared to control samples.

Malondialdehyde (MDA) content. Thiobarbituric acid reactive substances (TBARS)/MDA equivalent production is an index of membrane lipid peroxidation. Since there was similar amount of H2O2 production in both the control and water-stressed seedlings (after 24 h of greening), we did not observe any significant increase in MDA content either. After 72 h of greening under water-stress, the MDA content increased by 50% above the control value, as expected (Fig. 6B).
**Figure 6.** Antioxidative response of seedlings. (A) H\(_2\)O\(_2\) content and (B) MDA content in control (Con) and stressed (50 mM PEG) rice seedlings after 24 h and 72 h of greening. Seedlings were treated as in Fig. 1. H\(_2\)O\(_2\) was measured by absorbance at 240 nm, against a standard curve and MDA was measured as TBARS equivalents. Leaf tissue (200 mg) was homogenized in 20% TCA solution with and without 0.25% TBA, boiled and centrifuged. Absorptions at 440, 532 and 600 nm were measured from supernatant. Unspecific absorption contribution from without-TBA solution was deducted appropriately (see M&M) from sample values. The error bar represents SD of three replicates. ANOVA: *p < 0.05; **p < 0.01; and ***p < 0.001.

**Figure 7.** Proteomics of PEG fractionated soluble and peripheral thylakoid proteins of rice leaves after 72 h of water-stress treatment in light. Five-day old etiolated seedlings were subject to water-stress as in Fig. 1. The 2D gel of control and treated samples were done several times till a consistent pattern was obtained. (A) Representative 2-D gel of PEG fractionated soluble proteins of control rice seedlings, (B) functional category distribution of 34 down-regulated and (C) 25 up-regulated differentially expressed proteins based on actual number of identified proteins. Soluble proteome was isolated and PEG fractionated in two steps; first upto 10% and later upto 20% to remove abundant proteins e.g. Rubisco. Protein (800 µg) from supernatant was treated with G-Biosciences Perfect Focus kit to remove impurities; the obtained pellet was air dried and dissolved in rehydration buffer. After rehydration, isoelectric focusing of proteins was done on a 17 cm IP strip overnight; strip was equilibrated and second dimension was run on a 12.5% SDS-PAGE. 2-D gel was stained with Coomassie Brilliant Blue (CBB), Imaged with UMAX PowerLook 2100XL Image Scanner. Spots were detected, and matched with the Image Master-2D Platinum 6.0 software to get differentially expressed proteins. Numbered spots showing down-regulated proteins in the representative gel from control seedlings are mentioned in text and tables starting with CBC3.
Differentially expressed proteins. Soluble and peripheral membrane proteome was analyzed by 2-D gel electrophoresis and MALDI-TOF/TOF or ESI-MS/MS analysis of spots, after 72 h of greening. Silver-staining as well as colloidal Coomassie Brilliant Blue (CBB)-staining was used to visualize the spots (Fig. 7A). In silver-stained gel, we identified 31 differentially expressed proteins; 10 were up-regulated and 21 were down-regulated. In CBB-stained gels, we further identified 15 up-regulated and 13 down-regulated proteins. In total, 34 down-regulated proteins and 25 up-regulated proteins were identified (Table 1): Silver-stained proteins are named starting with ‘SS’, and CBB-stained spots are named starring with ‘CB’; C3 and W3 denote spots from the ‘control’ and the ‘water-stressed’ samples, respectively, after 72 h of greening. Gene Annotations, Gene Index (GI) number, locus names, gene names and localization (wherever known or predicted) for identified proteins are provided in the Supplementary Table S1. Out of 59 differentially expressed identified proteins, for 18 Uniprot IDs, no gene models were retrieved from the rice database (http://ricedb.plantenergy.uwa.edu.au/). For the remaining 41 (59–18) Uniprot IDs, 56 corresponding gene-model/loci were retrieved as shown in the Supplementary Table S2. Enriched GO-terms (Biological Process) (p < 0.05) were glycolysis, tricarboxylic acid cycle, cellular carbohydrate metabolic process, oxidation-reduction, cysteine biosynthetic process, photosystem II stabilization, carbon fixation, hydrogen peroxide catalytic process, cysteine biosynthetic process from serine, phosphatidyl inositol metabolic process and phosphate metabolic process.

Category-wise distribution of down-regulated and up-regulated identified proteins is presented in Fig. 7A and B respectively. Differentially-regulated proteins were functionally distributed into 12 groups (Table 1).

Down-regulated proteins (Table 1, and Supplementary Table S1) were categorized into 9 groups (Fig. 7A), namely: (i) photosynthesis (‘light reaction’; OEC23, OEC33 and carbon fixation; Rubisco LSU, FBA, PGK, TPI); (ii) protein folding and degradation (PPIase and Lon protease); (iii) subunit; inorganic pyrophosphatase and mitochondrial formate dehydrogenase precursor); (iv) transcription/RNA stabilization (mTERF and CSD protein); (v) cell homeostasis (similar to WD40 domain containing and kinesin like); (vi) cell homeostasis (Cyt P450, TRX domain containing protein, cysteine synthase 1, glyceraldehyde 3-phosphate dehydrogenase (GPDH), malate dehydrogenase (MDH), aminotransferase like and anion transporting protein); (vii) mRNA processing (UPF0614 protein C14orf102); (viii) signaling; and (ix) unknown/uncharacterized proteins.

The proteins that were up-regulated (Table 1 and Supplementary Table S1) were distributed into 8 categories (Fig. 7B), which are: (i) signaling (receptor like kinase, putative receptor kinase, phosphatidyl inositol 4-phosphate 5-kinase 4 (PIP5K4), calcineurin-like phosphoesterase like, Ca2+ distributed into 8 categories (Fig. 7B), which are: (i) uncharacterized proteins. (ii) unknown/MRNA processing (UPF0614 protein C14orf102); (iii) signaling; and (iv) cytoskeleton organization (similar to WD40 domain containing polyprotein-like retrotransposon); (v) antioxidant defense mechanism (ascorbate peroxidase (APx1), and dehydro ascorbate reductase (DHAR)); (vi) cell homeostasis (Cyt P450), (vii) specific functions (Saat; IAA amino acid hydrolase ILR1-like 3); and (viii) uncharacterized proteins.

Discussion
During photomorphogenesis, Chl, proteins, and lipids are synthesized and assembled to form the functional photosynthetic apparatus. Earlier, Dalal and Tripathy39 showed that water-stress during transition from skotomorphogenesis to photomorphogenesis led to a 42% reduction in Chl content, due to down-regulation of expression of genes and proteins involved in Chl biosynthesis39. This must be, at least partly, responsible for the lower Fo and Fm (Fig. 1A), as well as the lower Chl a fluorescence in water-stressed rice seedlings (Fig. 4A). Shibata shift (blue shift of absorption maximum of chlorophyllide from 684 nm (aggregated form) to 672 nm (disaggregated form), discovered by K. Shibata42 that takes place within a few minutes of light exposure of etiolated seedlings was also impaired in stressed seedlings33. In higher plants, both Chl a and Chl b are bound to light-harvesting pigment-protein complexes, LHCs. The availability of Chl b is essential for the assembly and functioning of most LHC proteins43. Binding of Chls to the LHC proteins stabilizes the latter in the thylakoid membranes44. Further, ROS produced in chloroplasts of water-stressed plants could down-regulate the expression of genes involved in Chl biosynthesis and photosynthesis via retrograde signaling45–47. In water-stressed rice seedlings, there was not only a downregulation of Chl biosynthesis, but also a reduction in pigment-protein complexes; this, obviously, led to lower Chl a fluorescence (Fig. 2). Moreover, biosynthesis of PSI1 reaction center protein D1 and its repair were also inhibited by H2O2, and other photosynthetic inhibitors and uncouplers48–49. In water-stressed developing rice seedlings, water-stress imposed on four-week-old well developed green wheat plants had no effect on the leaf chlorophyll content, and on the abundance of ATP synthase, PSII, PSI, and light-harvesting complexes32. However, in these plants, thylakoid membranes and other proteins were oxidized, due to the generation of ROS as a consequence of excess light absorption by the LHCs31. Decreased LHCII is an essential protection mechanism that allows plants to survive under unfavourable conditions52. Our results demonstrate that germinating seedlings protect themselves from water-stress-induced oxidative stress and photo-damage by downsizing their light-harvesting antenna and photosynthetic reaction centers (Figs 4 and 6).

The decreased Fo and Fm (Fig. 1A) in water-stressed rice seedlings is most likely due to a reduced Chl content53, and/or stress-induced alteration in xanthophyll-cycle dependent non-radiative energy dissipation54. Alternatively, low fluorescence may be related to a state 2 induced during the water-stress treatment. A significant
| Spot No | Protein | Mowse Score | Mr [kDa]/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|-------------|------------|-----------------------------------|-------------------|-------------------|
| SSC3-2  | Ribulose bisphosphate carboxylase large chain precursor, putative [Oryza sativa (japonica cultivar−)] | 70.5 | 56.5/9.7 | gi|108862318 | 19.3%/10 | Down |
| SSC3-8  | Fructose-bisphosphate aldolase | 113 | 41.8/6.0 | gi|108864048 | 34.4%/13 | Down |
| SSC3-12 | Fructose-bisphosphate aldolase class-I [Oryza sativa Japonica Group] | 77.9 | 39.6/7.6 | gi|62732954 | 23.7%/8 | Down |
| SSC3-20 | hypothetical protein OsJ_30136 [Oryza sativa Japonica Group] | 137 | 26.3/9.7 | gi|125606445 | 60.0%/15 | Triose phosphate isomerase, chloroplastic Down |
| SSC3-23 | hypothetical protein OsJ_20474 [Oryza sativa Indica Group] | 109 | 30.5/7.7 | gi|125552851 | 43.7%/10 | Phosphoglycerate kinase, chloroplastic Down |
| SSC3-11 | Os01g0501800 [Oryza sativa (japonica cultivar-group)] | 143 | 35.1/6.0 | gi|115436780 | 53.8%/15 | OEC 33 Down |
| CBC3-49 | PSII Oxygen evolving complex protein 2 precursor, rice | 332 | 26.93/8.66 | T02B73 | 44%/8 | Down |

**1. Carbon fixation:**

**2. Unknown/Uncharacterized proteins:**

| Spot No | Protein | Mowse Score | Mr [kDa]/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|-------------|------------|-----------------------------------|-------------------|-------------------|
| SSC3-1  | Os04g0602020 [Oryza sativa (japonica cultivar-group)] | 45.8 | 31.6/9.1 | gi|115460610 | 24.3%/5 | Remorin C-terminal domain protein+DNA binding Down |
| SSC3-6  | hypothetical protein LOC_ Os11g39500 [Oryza sativa (japonica cultivar-group)] | 33.1 | 13.7/9.8 | gi|77551932 | 18.6%/3 | Hypothetical protein Down |
| SSC3-7  | hypothetical protein OsJ_32755 [Oryza sativa Indica Group] | 34.9 | 11.5/11.1 | gi|218184148 | 49.0%/3 | IGR family prot (unknown function) Down |
| SSC3-18 | Os11g0652000 [Oryza sativa (japonica cultivar-group)] | 56.4 | 11.3/10.7 | gi|115438911 | 60.8%/5 | Uncharacterized Down |
| SSC3-19 | unknown protein [Oryza sativa Japonica Group] | 44.3 | 12.7/10.4 | gi|47900462 | 28.6%/3 | Unknown protein Down |
| CBC3-39 | hypothetical protein [Oryza sativa Japonica Group] | 28.5 | 12.8/6.7 | gi|53791471 | 51.7%/2 | No match Down |
| CBC3-40 | hypothetical protein [Oryza sativa Japonica Group] | 59 | 9.9/7.8 | gi|57899958 | 46.4% | No match Down |
| CBW3-5  | hypothetical protein [Oryza sativa] | 45.2 | 15.2/11.5 | gi|10440615 | 59.4%/6 | No match Up |
| CBW3-26 | hypothetical protein [Oryza sativa Japonica Group] | 38.2 | 16.01/11.6 | gi|51534990 | 30.8%/4 | No match Up |
| CBW3-27 | hypothetical protein OsJ_26020 [Oryza sativa Indica Group] | 43.1 | 260.7/10.8 | gi|218199617 | 22.8%/8 | No match Up |
| CBW3-29 | Os12g0623600 [Oryza sativa (japonica cultivar-group)] | 34.0 | 32.6/7.8 | gi|115489646 | 18.8%/3 | Unknown function RmIC_Cupin domain, DUF1637 family Up |
| CBW3-31 | hypothetical protein OsJ_02300 [Oryza sativa Indica Group] | 46.4 | 12.5/9.9 | gi|125526034 | 37.9%/4 | No match Up |
| CBW3-32 | hypothetical protein [Oryza sativa Japonica Group] | 17.5/31.4 | 12.4 | gi|50251877 | 28.2%/32 | No match Up |

**3. Cell Homeostasis:**

Continued
| Spot No | Protein | Mowse Score | Mr (kD)/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|------------|------------|-----------------------------------|-------------------|-------------------|
| CBC3-45 | Putative malate dehydrogenase Oryza sativa (japonica cultivar-group) | 103 | 35.414/8.22 | Q6F361_ORYSA | 12%/4 | | Down |
| CBC3-46 | Malate dehydrogenase cytoplasmic Oryza sativa (japonica cultivar) | 221 | 35.415/5.75 | MDHC_ORYSA | 34%/17 | | Down |
| CBC3-47 | Putative malate dehydrogenase mitochondrial Oryza sativa (japonica cultivar-group) | 124 | 35.439/8.74 | Q4JA2_ORYSA | 12%/5 | | Down |
| CBW3-24 | Os04g0174100 Oryza sativa (japonica cultivar-group) | 36.9 | 27.7/9.0 | gi|115457106 | 21.7%/4 | (CYP family, fragment) or tyrosine N-monoxygenase | Up |

4. Retrotransposons:

| Spot No | Protein | Mowse Score | Mr (kD)/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|------------|------------|-----------------------------------|-------------------|-------------------|
| SS W3-7 | putative gag-pol polyprotein Oryza sativa (japonica cultivar-group) | 53.9 | 133.1/6.9 | gi|31126757 | 9.5%/8 | | Up |
| SS W3-8 | OSIGRa014621.3 Oryza sativa (indica cultivar) | 54 | 79.35/9.8 | gi|116309588 | 9.1%/7 | retrotransposon protein, putative, Ty1-copia subcla | Up |
| SS W3-11 | Hypothetical protein P0701E03.31 Oryza sativa (japonica cultivar-group) | 18.3 | 6.77/4.7 | Q69YD6_ORYSA | 10.6%/1 | plant disease resistance polyprotein-like | up |

5. Transcription factors and DNA/RNA binding proteins:

| Spot No | Protein | Mowse Score | Mr (kD)/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|------------|------------|-----------------------------------|-------------------|-------------------|
| CBC3-35 | Putative glycine rich protein 2 Oryza sativa-japonica; have four CCHC Zn fingers | 49 | 22.7/9.664 | Q6YUR8_ORYSA; BAD07599 | 5%/1 | | Contain S1-like CSD; cold shock domain protein 1 | down |
| CBC3-41 | Os04g0518200 Oryza sativa (japonica cultivar-group) | 66.7 | 10.2/72.15 | gi|115477308 | 25.3%/12 | Mito transcription termination factor family-related protein | down |
| SS W3-14 | Zinc finger CCCH domain-containing protein 31 Oryza sativa sub japonica GN = Os04g6665700 PE = 2 SV = 1 | 51.3 | 32.338/10.4 | C3H31_ORYSJ | 29.4%/5 | | | up |
| CBW3-4 | hypothetical protein Os_09486 Oryza sativa (japonica cultivar-group) | 45.7 | 20.5/4.9 | gi|125541662 | 13.8%/5 | Low Similarity (44%) to Setaria Zn-finger prot 7-like | up |
| CBW3-25 | hypothetical protein [Oryza sativa (japonica cultivar-group)] | 32.6 | 11.8/12.6 | gi|53793156 | 46.7%/4 | RNA binding glycine-rich protein like | up |

6. Signaling:

| Spot No | Protein | Mowse Score | Mr (kD)/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|------------|------------|-----------------------------------|-------------------|-------------------|
| SSC3-22 | Osb0g0146100 Oryza sativa (japonica cultivar-group) | 47.3 | 118.9/9.2 | gi|115466384 | 9.5%/9 | NBS-LRR disease resistant protein | | |
| SS W3-12 | Hypothetical protein OSIGRa014621.3 Oryza sativa (japonica cultivar-group) | 59.8 | 51.87/11.2 | Q6EQ30_ORYSA | 25.9%/6 | calcineurin-like phosphoesterase-like | up |
| SS W3-13 | hypothetical protein Os_14142 Oryza sativa (japonica cultivar-group) | 65.5 | 49.62/9.5 | gi|218194019 | 30.0%/12 | Putative receptor kinase (tyrosine) | up |
| SS W3-15 | Osb0g0483100 Oryza sativa (japonica cultivar-group) | 59.3 | 15.076/4.2 | gi|115479825 | 65.2%/5 | Ca²⁺ binding EF-hand 5 motif prot | up |
| CBW3-31 | hypothetical protein - OJNBA0016I09.17 Oryza sativa (japonica cultivar-group) | 49.5 | 30.4/10.7 | Q5QNA8_ORYSA | 26.4%/5 | Similar to receptor like kinase | up |
| CBW3-10 | Phosphatidylinositol-4-phosphate 5-kinase 4, putative, expressed Os04g0665700 PE = 2 SV = 1 | 53.5 | 84.93/9.7 | gi|108708236 | 13.5%/8 | | |

7. Energy balance:

| Spot No | Protein | Mowse Score | Mr (kD)/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|------------|------------|-----------------------------------|-------------------|-------------------|
| SSC3-15 | Osb0g0750100 Oryza sativa (japonica cultivar-group) | 57.2 | 26.2/4.8 | gi|115448701 | 23.5%/6 | H⁺ transporting ATP synthase, δ chain | down |
| SSC3-21 | putative inorganic pyrophosphatase Oryza sativa (japonica cultivar-group) | 155 | 31.8/5.7 | gi|46805452 | 42.3%/14 | | down |
| CBC3-34 | AP003518 NID: Oryza sativa (japonica cultivar-group) | 58 | 41.3/56.8 | BAD37348 | 6%/3 | Formate dehydrogenase, mitochondrial precursor | down |

8. Protein folding/de-folding and degradation:

| Spot No | Protein | Mowse Score | Mr (kD)/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|------------|------------|-----------------------------------|-------------------|-------------------|
| SSC3-3 | putative Lon protease Oryza sativa (japonica cultivar-group) | 33.1 | 96.9/7.0 | gi|50725794 | 6.9%/4 | | |
| CBC3-42 | Putative peptidylprolyl isomerase Oryza sativa (japonica cultivar-group) | 140 | 26.5/9.37 | Q75M32_ORYSA | 24%/5 | | | |

9. Antioxidant proteins:

| Spot No | Protein | Mowse Score | Mr (kD)/pI | Access key | Sequence Coverage/peptides matched | Protein orthologue | Up/down regulated |
|---------|---------|-------------|------------|------------|-----------------------------------|-------------------|-------------------|
| CBW3-28 | Osb0g0285700 Oryza sativa (japonica cultivar-group) | 66.4 | 27.02/5.3 | gi|115452337 | 54.8%/10 | APx1 | up |

Continued
10. Cell shape/cytoskeleton protein:

SSW3-6  Os04g0538800 [Oryza sativa (japonica cultivar-group)]  71.5  106.4/6.6  protein] (Protein mannose-6-phosphate transporter-like)  12.3%/8  Similar to Kinesin heavy chain (fragment)  up

SCC3-5  hypothetical protein Os2_28608 [Oryza sativa Indica Group]  39.9  57.6/0.0  protein-like  0.0%/3  Similar to Transducin/WD-40 repeat protein-like  down

CBW3-3  Hypothetical protein Os12g0465000 [Oryza sativa (japonica cultivar-group)]  28  14.1/10.2  similar (68% positives) to Saccharum hybrid R570, A0A059Q350 Acetylglucosaminyl transferase family protein 22.0%/2  up

11. mRNA processing:

CBC3-33  Hypothetical Protein OX1611_108.15.0-Oryza sativa (japonica cultivar-group)  39  132.188/5.57  protein] (Protein mannose-binding lectin)- Oryza sativa (japonica cultivar-group)  1%/1  UPF0614 C14orf102-like protein; have HAT and NRDE-2 domains  down

Table 1. Differentially expressed proteins in water-stressed rice seedlings categorized into different subgroups according to their function. Spot No. generated by UMAX PowerLook 2100XL Image Scanner (v 6.0, GE Healthcare); Numbered spots correspond to the identified proteins, refer the spots shown on the representative image in Fig. 7. Accession number from NCBI database. SC, sequence coverage; Mr, theoretical Molecular weight; pI, theoretical Isoelectric point; Fold change refers to increased (Up) or decreased levels (down) of protein content from control to water-stressed seedlings. Fold change ± 2 was used as threshold for protein identification. Orthologues were obtained by using Uniprot protein sequence and BLAST.

12. Proteins with specific functions:

SSW3-9  IAA-amino acid hydrodase ILR1-like 3 O.sativa sub japonica (group)  47.8  44.07/5.4  protein] (Protein mannose-binding lectin)- Oryza sativa (japonica cultivar-group)  23.0%/5  up

SSW3-10  IAA-amino acid hydrodase ILR1-like 3 O.sativa subsp. japonica  47.4  44.07/5.4  protein] (Protein mannose-binding lectin)- Oryza sativa (japonica cultivar-group)  17.0%/6  up

CBW3-2  Salt stress-induced protein (Salt protein) (Protein mannose-binding lectin)- Oryza sativa (japonica cultivar-group)  72  15.03/4.9  protein] (Protein mannose-binding lectin)- Oryza sativa (japonica cultivar-group)  45.8%/5  up

decrease in Fv/Fm, which is a proxy of PSII photochemistry efficiency37, in water-stressed developing seedlings (Fig. 1A), might have resulted from the down-regulation of synthesis and assembly of PSII reaction centers. This would have reduced not only the concentration of PSII reaction centers, but also of the oxygen evolving complex proteins, such as OEC33 (PsbO), OEC23 (PsbP) and OEC 16 (PsbQ) (Fig. 4). The observed changes in Chl a fluorescence spectra (Fig. 2) are in agreement with this observation. Furthermore, the “operating” quantum yield of PSII (ΦPSII) during actinic illumination also sharply declined in water-stressed developing seedlings (Fig. 1B), OEC33 is important for stabilization of Mn and Cl co-factors in water oxidizing complexes54; its reduction via RNAi results in a decrease in quantum yield of photosynthesis55. Beside providing appropriate amounts of calcium and chloride ions for water splitting reactions, OEC23 and OEC16 play an additional role in the assembly of PSII supercomplexes, and their decreased abundance was clearly correlated with reduced PSII-dependent O2 evolution54; therefore, down-regulation of OEC proteins also explains the impaired PSII-dependent water oxidation and oxygen evolution that we found in thylakoid membranes isolated from water-stressed developing seedlings (Fig. 3).

Higher NPQ in water-stressed seedlings compared to the controls at all measured PAR values, especially at low actinic light intensities (Fig. 1C) suggest an adaptive response of developing seedlings for increased dissipation of absorbed energy as heat to protect their thylakoid membranes from photodamage26. Down-regulation of δ-subunit of ATP synthase was observed in our proteome profile of water-stressed seedlings (Table 1). Impaired ATP synthase would have resulted in reduced transport of protons across the thylakoid membrane; the lowered dissipation of ΔpH and increased acidification of the lumen25 could lead to higher NPQ in (water) stressed seedlings even at low light intensities.

In water-stressed samples, reduced amount of Chl, LHCs and reaction center proteins result in reduced light absorption, and decreased amount of oxygen evolving complex proteins result in reduced utilization of the absorbed energy. Consequently, reduction of PSII activity was almost similar (nearly 50%) under limiting as well as under saturating light intensities. In the same vein, the down-sizing of the light-harvesting Chl-binding proteins of PSI i.e., LHCl (Lhca1 and Lhca4) and PSI core complex protein subunits III (PsaF), V (PsaK) and VI (PsaH) in water-stressed seedlings, resulted in diminished light absorption as well as light utilization by PSI, as was evident from the reduction of PSI reaction by 30% at limiting and higher light intensities (Fig. 3E). Relatively lower reduction of PSI activity than that of PSII (Fig. 3) suggests that development of PSI of developing rice seedlings is more tolerant to water-stress. The Eadie plot21,46 for control and treated seedlings revealed a marked drop in the quantum efficiency of PSI and PSII reactions and a substantial reduction of Vmax in stressed developing seedlings even at low light intensities.
seedlings (Fig. 3); these suggest a coordinated developmental down-regulation of light-harvesting and reaction center proteins of PSI as well as PSII in response to osmotic stress\(^{31,40}\).

Room temperature fluorescence emission spectrum of thylakoid membranes had a peak at 684 nm due to PSII, when they were excited at 440 nm (Fig. 2A). Water-stressed samples had a reduced fluorescence emission, probably due to a block in electron donation from water side of PSII; decreased abundance of oxygen evolving complex proteins (33 kDa, 23 kDa and 18 kDa) supports this hypothesis. Gross perturbation of structural organization of thylakoid membranes usually induces differences in fluorescence spectra at low temperature (77 K); these spectra usually have emission peaks at 685 nm (F\(_{685}\)), at 695 nm (F\(_{695}\)), which mostly originates from PSII CP43 and CP47 respectively\(^{38}\), and a F\(_{735-740}\) peak that originates mostly from PSI\(^{37}\). If a part of LHCI antenna is removed from PSI by detergent treatment, the inner antenna of the PSI reaction center fluoresces at 722 nm\(^{58}\), while isolated LHCI complexes fluoresce around 735–740 nm; this is consistent with the assignment by Briantais et al.\(^{59}\), the inner PSI antenna for F\(_{722}\), and for LHCI for F\(_{735-740}\). A shift to F\(_{730}\) from F\(_{740}\) in our water-stressed samples are due to a partial loss of components of LHCI.

In the absence of Mg\(^{2+}\), the F\(_{685}/F_{740}\) ratio in thylakoid membranes isolated from control seedlings was ~1.54 (Fig. 2B). In experiments by others, LHCIi has been shown to migrate closer to PSI\(^{35}\) in presence of Mg\(^{2+}\), leading to stacking of thylakoids into grana\(^{42}\) (see a review on role of ions by Kaňa and Govindjee)\(^{42}\). This results in efficient energy transfer from LHCIi to the reaction center of PSI, and consequently, an increase in PSI fluorescence at 686 nm and, thus, higher F\(_{686}/F_{740}\) [see Results] ratio (an increase of F\(_{686}/F_{740}\) ratio to 1.73 in the control samples with Mg\(^{2+}\), compared to 1.54 without Mg\(^{2+}\) (Fig. 2B)). In water-stressed seedlings, Mg\(^{2+}\)-induced increase in F\(_{686}/F_{740}\) was almost similar to control. Although a reduced grana stacking was observed in electron micrographs of the plastids of water-stressed seedlings, they retained the ability of Mg\(^{2+}\)-induced migration of LHCIi to PSI core complex.

As revealed from our proteomics studies, water-stressed rice seedlings downsized their carbon reduction cycle enzymes i.e., Rubisco LSU, fructose bisphosphate aldolase (FBA), triosephosphate isomerase (TPI) and phosphoglycerate kinase (3PGK) (Table 1). In contrast, in water-stressed well-developed plants, Rubisco content was often not affected, although its initial and total activity is known to decrease due to blocking of catalytic sites by 2-carboxyarabinitol-1-phosphate and other sugar phosphates\(^{93}\). Sugar phosphates can be removed from Rubisco by ATP-dependent enzyme Rubisco activase\(^{27}\), leading to its reactivation, hence undermining the role of Rubisco activase and ATP production in regulating carbon assimilation under drought. Drought-induced stomatal closure, and consequent decreased carboxylation, also result in down-regulation of protein abundance of Calvin-Benson cycle enzymes\(^{13–16}\).

Down-regulation of peptidyl prolyl isomerase (PPlase), in our water-stressed rice seedlings, suggest that there was a decrease in the ability to repair photosynthetic proteins, which, in turn, results in reduced PSI and PSII activities.

In mature plants, only minimal amounts of Chl and proteins are synthesized, mostly to replace those photodestroyed in PSI and PSII and LHCS, due to light. Therefore, it appears that mature plants do not downsize the components of the photosynthetic apparatus in response to drought. Several studies have demonstrated that abiotic stresses, including water-stress, cause ROS production due to an over-reduction of photosynthetic electron transport chain\(^{25,29,64,65}\). Abiotic stresses also impair reaction centers of well-developed seedlings or mature plants, leading to reduced transfer of absorbed light energy from light harvesting complexes to photodamaged reaction centers and consequently generation of higher amounts of singlet oxygen via photosensitization reactions of Chl\(^{27,28,31}\). Well developed rice plants treated for 48 h with 30% PEG 6000, a concentration similar to that used in our current experimental protocol, accumulated 2.5–4 fold higher H\(_2O_2\), which impaired Rubisco activity and promoted stomatal closure\(^{25}\). However, we showed that germinating seedlings protect themselves by downsizing their light-harvesting antenna and photosynthetic reaction centers to reduce ROS production (Figs 4 and 6). In addition, we found that antioxidative enzymes i.e., ascorbate peroxidase 1 (APx1) and dehydroascorbate reductase (DHAR), were up-regulated in developing stressed seedlings (Table 1) to neutralize comparatively smaller amount of ROS generated. Therefore, we did not observe any increase of H\(_2O_2\) in developing seedlings subjected to water-stress for 24 h, and there was only a small increase (23%) after 72 h. Consequently, membrane lipid peroxidation, monitored as MDA production did not increase after 24 h of stress treatment, but it increased by 50% only after 72 h.

In conclusion, we can clearly state that unlike mature plants, seedlings exposed to water-stress during early photomorphogenesis, protect themselves from photo-oxidative stress by downsizing their photosynthetic apparatus. Therefore, we recommend that attempts should be made to down-regulate the light absorption during stress condition by reducing Chl biosynthesis. This can be achieved by either using molecular marker assisted plant breeding methods or by using transgenic RNAi approaches, silencing the expression of gene encoding for glutamyl-tRNA reductase or other early enzymes responsible for the synthesis of 5-aminolevulinic acid, a precursor of Chl biosynthesis. The RNAi expression could be modulated by promoters that sense increased ROS (e.g., H\(_2O_2\)) produced during stress conditions.

**Materials and Methods**

**Plants growth conditions and stress treatment.** Seeds of drought-sensitive rice (Oryza sativa) cultivar Pusa Basmati-1 (PB-1) were obtained from the Indian Agricultural Research Institute, New Delhi, India. Five-day old etiolated seedlings, grown at 25 °C, were treated with half-strength MS\(^{36}\) solution, or half-strength MS solution +40 mM PEG (−0.73 MPa) or 50 mM PEG 6000 (−1.06 MPa; Merck, Kenilworth, NJ)\(^{35}\), 16 h prior to their transfer to continuous cool-white fluorescent (plus incandescent) light of 100 μmol photons m\(^{-2}\) s\(^{-1}\); low light intensity was used to avoid photodamage of the seedlings during greening. The seedlings were maintained for 3 days at 28 °C and at 75 % relative humidity in a Conviron plant growth chamber.
**Polarographic measurement of photosynthetic electron transport.** Approximately 5 g of leaves were homogenized in 40 ml of isolation buffer containing 0.4 M sorbitol, 0.05 M Hepes/KOH (pH 7.3), 1mM MgCl₂, and 1mM EDTA at 4°C, under green safe light (28 Chakraborty and Tripathy, 1992). The homogenate was passed through 8 layers of cheese cloth and 1 layer of Mira cloth, and centrifuged at 5,000 rpm for 7 min. The pellet was suspended in a buffer solution containing 0.4 M sorbitol, 0.05 M Tris (pH 7.6), 1 mM MgCl₂ and 1 mM EDTA; Chl concentration was estimated according to Porra et al.⁶⁷. Photosynthetic electron transport was measured at 25 ± 1°C with a Clark-type oxygen electrode (Hansatech, Kings Lynn, UK), by using 1 ml suspension of thylakoid membranes containing 20 μg Chl ml⁻¹; illumination was with white-light from a tungsten source of 1,500 μmol photons m⁻² s⁻¹. The whole electron transport chain, from H₂O to methylviologen (MV) (1 mM), was monitored as O₂ uptake⁶⁸,⁶⁹. PSII activity was monitored as O₂ evolution resulting from electron transport from H₂O to p-phenylenediamine (PD) (0.5 mM). The partial electron transport through PSI was measured as oxygen consumption. Ascorbate (1 mM)/DCIP (0.2 mM) couple was used as electron donor to PSI, and MV (1 mM) as electron acceptor⁶⁸,⁶⁹; the electron flow from PSII was blocked by 3-(3, 4-dichlorophenyl) 1,1-dimethyl urea (DCMU) (20 μM), and 1 mM sodiumazide was used as an inhibitor of catalase.

Light saturation curves were measured at several light intensities obtained by using neutral density filters (Balzers, Neugrüt, Lichtenstein).

**Transmission electron microscopy.** Leaves (in triplicate) were fixed with 2.5% glutaraldehyde⁷⁰ and 1% OsO₄ for 2–4 h, and then washed, dehydrated with acetone, cleared with epoxy propane/xylene, and infiltrated with resin containing araldite and toluene (up to 75% araldite). After embedding, araldite was polymerized at 50°C, and then at 60°C. Ultra-thin sections were cut, stained in saturated uranyl acetate in 50% ethanol, washed in 50% ethanol and water, and viewed in a Transmission Electron Microscope (TEM) (Phillips-CM-10, Eindhoven, Netherlands).

**Chlorophyll a fluorescence measurements.** Chl a fluorescence signal is a highly sensitive signature of diverse aspects of photosynthesis, and thus, it is widely used in photosynthesis research³⁸,⁷¹ (see chapters in Papageorgiou and Govindjee)⁷².

**Fluorescence emission spectra of isolated thylakoids.** Room temperature (RT; 298 K) and low temperature (77 K) fluorescence emission spectra of isolated thylakoids were recorded with a SLM-AMINCO-8000 spectroradiometer; for 77 K-spectra, the excitation and emission slit widths were 4 nm, while for RT-spectra, the excitation and emission slit widths were 8 nm and 4 nm respectively. Rhodamine-B was used in the reference channel as quantum counter. A tetraphenylbutadiene block was used to adjust the voltage in sample, as well as in reference channels to 20000 counts s⁻¹ at excitation and emission wavelengths of 348 nm and 422 nm, respectively. RT-spectra were corrected for the instrument response.

**Thylakoid isolation.** Leaves from control and water-stressed seedlings were grounded in Hepes-NaOH buffer, pH 7.6 (20 mM Hepes, 10 mM NaCl, 0.4 M sucrose, 0.5% BSA) and then centrifuged at 5,000 rpm for 7 min; pellet was suspended in 5 mM Hepes-NaOH buffer (pH 7.5) containing 0 or 4 mM MgCl₂. Thylakoids equivalent to 3–4 g Chl ml⁻¹ were used for RT- and 77K-spectra (25% glycerol) at excitation wavelength of 440 nm and emission wavelength range of 620–750 nm and of 620–780 nm, respectively⁷¹. Each experiment was performed three times.

**Chl a fluorescence induction curves.** Chl a fluorescence induction curves at 25°C were recorded on attached leaves, with a portable PAM-2100 fluorimeter (Walz, Effeltrich, Germany); all measurements were repeated 3 times. The initial (Fo) and maximum (Fm) fluorescence, was measured on seedlings that were dark-adapted for 20 min; the measuring light was red (650 nm), and of very low intensity (0.1 μmol photons m⁻² s⁻¹); its frequency was 0.6 KHz. Then, a 0.8 s saturation light pulse of approximately 8,000 μmol photons m⁻² s⁻¹ was applied to measure the maximum fluorescence, Fm. The Fo and the Fm values were used to calculate the ratio Fo/Fm = (Fm – Fo)/Fm, where Fm is the maximal variable fluorescence; this ratio represents the maximum quantum yield of PSII photochemistry. Light response curves were obtained by measuring fluorescence as a function of increasing actinic light intensity (4 to 270 μmol photons m⁻² s⁻¹; wavelength: 665 nm). The quantum yield of PSII (φPSII) was calculated as (Fm’ – F)/Fm’, where Fm’ is the maximum fluorescence at steady state in pre-illuminated samples, and F is fluorescence immediately before the application of the saturation pulse⁷². Non-photocatalytic quenching (NPQ) was calculated as (Fm – Fm’)/Fm’.

**Thylakoid isolation and western blotting.** 20 μg of plastid protein³⁸, estimated as described by Bradford⁴¹, was separated on SDS-PAGE three times for each photosynthetic target protein, transferred to nitrocellulose (NC) membranes and blocked with BSA solution. Subsequently, blots were incubated with polyclonal and heterologous primary antibodies (Supplementary Table S4) and alkaline phosphatase-conjugated antibodies and developed for color. Image J (NIH, USA) was used for quantification.

**Estimation of malondialdehyde (MDA).** 200 mg fresh tissue was homogenized in 20 % (w/v) trichloroacetic acid (TCA) alone (– TBA (Thiobarbituric acid)) or 20 % TCA (w/v) plus 0.25 % TBA (+TBA)⁴¹. Samples were vortexed, heated at 95°C in a water bath for 25 min, cooled, and centrifuged at 12,000 rpm for 10 min. Absorbance was read at 440, 532, and 600 nm. MDA equivalent was calculated as follows: A = (Abs 532 (TBA) – (Abs 600 (TBA)) / (Abs 532 (TBA) – (Abs 600 (TBA))); B = (Abs 440 (TBA) – Abs 600 (TBA) 0.057; and finally as: A = B/1,57,000) 10⁶ (in nmol ml⁻¹).
Determination of hydrogen peroxide. Leaf tissue (200 mg) was homogenized in 2 ml of 0.1 % (w/v) TCA solution on ice and centrifuged at 12,000g for 15 min. 0.4 ml of the supernatant was added to 0.4 ml of 10 mM potassium phosphate buffer (pH 7.0) and 0.8 ml of 1 M KI; the absorbance was measured at 390 nm. The leaf H₂O₂ content was measured using standard calibration curve of H₂O₂ (Thermo Fisher Scientific, MA).

Statistical analysis. All measurements were subjected to analysis of variance (ANOVA) to check for significance level of differences (defined as p < 0.05). Data is shown as mean ± standard deviation (SD).

2-Dimensional gel electrophoresis. Isolation of soluble and loosely bound thylakoid proteins. After 72 h of greening, two grams of tissue was homogenized in 10 ml of ice-cold Mg/NP-40 extraction buffer containing 0.5 M Tris-HCl, pH 8.3, 2% v/v NP-40, 20 mM MgCl₂, 2% v/v 3-mercaptoethanol, 1 mM phenylmethylsulfonyl fluoride and 1% w/v polyvinylpolypyrrolidone; this was repeated several times until a consistent pattern was obtained. After centrifugation at 12,000 g for 15 min at 4 °C, 50% (w/v) PEG–4000 stock solution was added to the supernatant to a final concentration of 10% PEG. The solution was incubated in ice and centrifuged at 1,500 g for 15 min each. The supernatant was adjusted to 15% (Silver-stained) or 20% (CBB-stained) PEG by adding 50% (w/v) PEG–4000 stock, incubated and centrifuged as described above. For silver- and CBB-stained gels, 150 µg and 800 µg of protein from resulting supernatant was treated with PERFECT-FOCUS™ kit from Geno Biosciences (St. Louis, MO); the pellet, thus obtained, was washed and air dried.

Isoelectric focusing and 2-D gel-electrophoresis. Air dried pellet was dissolved in rehydration buffer containing 8 M urea, 3% CHAPS, 20 mM DTT, 1 mM PMSF, and 2% ampholyte (GE–Healthcare, Buckinghamshire, UK). Protein extract was loaded on a 17 cm IEF strip of pH 3–10 (4–7 in case of silver-stained gels) (BioRad, CA). Rehydration was followed by electrophoresis at 250 V for 0.5 h, 10,000 V for 3.5 h, and 40,000 VH at 10,000 V.

After equilibration of the strips, proteins were separated on 12.5% SDS-PAGE. Gels were scanned with UMAX PowerLook-2100XL Image Scanner (GE–Healthcare). Spots were detected, matched, and edited with Image Master 2D Platinum 6.0 software (GE–Healthcare).

Mass spectrometry and database search. Protein spots showing two fold or higher difference (% vol) were excised, destained, digested and extracted[6]. For one set of spots, matrix-assisted laser desorption ionization (MALDI) with Bruker Autoflex II TOF/TOF mass spectrometer (Bruker Daltonics, Billerica, MA) was used[6]. The peptide mass data were searched against NCBI nr (non-redundant) Oryza sativa green plant database with the Mascot search engine (Matrix Science Ltd., UK). Two missed cleavage sites were allowed, cysteine was carbamidomethylated and methionine was allowed to be partially oxidized.

Other sets of proteins were detected with LC-MS/MS[12]. MS analysis was performed on Applied Biosystems 4000 Q-TRAP–system (Foster City, CA), equipped with Agilent 1200-NanoLC system (Santa Clara, CA). Proteins were excised manually, destained, digested and extracted. LC separation of peptides was performed on ZORBAX 300SB-C18 silica column, increasing the gradient from 3 to 97 % acetonitrile in 0.1% formic acid. The first MS detected ions were at scan range of 400–1600 m/z. Fragmentation spectra were submitted by Sequest algorithm to BioWorks (v3.3.1; Thermo Scientific) for searching a composite database of proteins using mascot (http://www.matrixscience.com/). Search settings applied were: trypsin as cleaving enzyme, maximally two missed cleavages, peptide mass tolerance 15 ppm, fragment ion tolerance 0.5 Da, and no differential post-translational modifications were allowed per peptide.

UNIPROT and University of Western Australia rice database (http://ricedb.plantenergy.uwa.edu.au)[7] were used for gene annotations of identified proteins. Gene ontology (GO) enrichment analysis was performed with ricearray (http://www.ricearray.org/analysis/go_enrichment.php) using default parameters.

Data availability. Most of the data generated or analysed during this study are included in this published article (and its supplementary information files). Any other remaining necessary data is available from the corresponding author upon reasonable request.

References
1. Pareek, A., Sopory, S. K. & Bohnert, H. J. Govindjee (eds) Abiotic Stress Adaptation in Plants, Physiological, Molecular and Genomic Foundation (Springer, Dordrecht, 2010).
2. Zhao, M. S. & Running, S. W. Drought-Induced reduction in global terrestrial net primary production from 2000 through 2009. Science 329, 940–943 (2010).
3. Lawlor, D. W. & Tezara, W. Causes of decreased photosynthetic rate and metabolic capacity in water-deficient leaf cells: a critical evaluation of mechanisms and integration of processes. Annal. Bot. 103, 561–579 (2009).
4. Pinheiro, C. & Chaves, M. M. Photosynthesis and drought: can we make metabolic connections from available data? J. Exp. Bot. 62, 869–882 (2011).
5. Chaves, M. M., Flexas, J. & Pinheiro, C. Photosynthesis under drought and salt stress: Regulation mechanisms from whole plant to cell. Annal. Bot. 103, 551–560 (2009).
6. Evans, J. R., Kaldenhoff, R., Genty, B. & Terashima, I. Resistances along the CO₂ diffusion pathway inside leaves. J. Exp. Bot. 60, 2235–2248 (2009).
7. Tezara, W., Mitchell, V., Driscoll, S. D. & Lawlor, D. W. Water stress inhibits plant photosynthesis by decreasing coupling factor and ATP. Nature 401, 914–917 (1999).
8. Miyazawa, S. I., Yoshimura, S., Shinzaki, Y., Maeshima, M. & Miyake, C. Deactivation of aquaporins decreases internal conductance to CO₂ diffusion in tobacco leaves grown under long term drought. Funct. Plant Biol. 35, 553–564 (2008).
9. Maroco, J. P., Rodrigues, M. L., Lopes, C. & Chaves, M. M. Limitations to leaf photosynthesis in field-grown grapevine under drought –metabolic and modelling approaches. Funct. Plant Biol. 29, 451–459 (2002).
10. Salekdeh, G. H., Siopongco, J., Wade, L. J., Ghareyazie, B. & Bennett, J. Proteomic analysis of rice leaves during drought stress and recovery. Proteomics 2, 1131–1145 (2002).
11. Liu, W. J. et al. Effect of water stress on photosystem 2 in two wheat cultivars. Biol. Plant. 50, 597–602 (2006).
37. Genty, B., Briantais, J. M. & Baker, N. R. The relationship between the quantum yield of photosynthetic electron transport and
40. Sayre, R. T. & Cheniae, G. M. Studies on the reconstitution of O₂-evolution in chloroplasts.
39. Govindjee, W . & Prezelin, D. B.B. & Sweeney, B.M. Chlorophyll a fluorescence of
35. Michel, B. E. & Kaufmann, M. R. The osmotic potential of polyethylene glycol 6000.
36. Chezen, O., Hartwig, W . & Newman, P . M. The different effects of PEG-6000 and NaCl on leaf development are associated with
32. Santel, H. J. & Apel, K. The protochlorophyllide holochrome of barley (Hordeum vulgare L.): the effect of light on the NADPH:protochlorophyllide oxidoreductase. FEBS J.
31. Tripathy, B. C., Mohapatra, A. & Gupta, I. Impairment of the photosynthetic apparatus by oxidative stress induced by in vivo: re-evaluation of the roles of catalase, α-tocopherol, non-photothermal quenching, and electron transport. Biochim. Biophys. Acta 1817, 1127–1133 (2012).
30. Schmitt, F. J. et al. Reactive oxygen species: re-evaluation of generation, monitoring and role in stress-signaling in photrophic organisms. Biochim Biophy Acta 1837, 835–844 (2014).
29. Tripathy, B. C., Mohapatra, A. & Gupta, I. Impairment of the photosynthetic apparatus by oxidative stress induced by in vivo: re-evaluation of the roles of catalase, α-tocopherol, non-photothermal quenching, and electron transport. Biochim. Biophys. Acta 1817, 1127–1133 (2012).
28. Schmitt, F. J. et al. Reactive oxygen species: re-evaluation of generation, monitoring and role in stress-signaling in photrophic organisms. Biochim Biophy Acta 1837, 835–844 (2014).
27. Miller, G., Suzuki, N., Ciftci-Yilmaz, S. & Mittler, R. Reactive oxygen species homeostasis and signaling during drought and salinity stresses. Plant Cell Environ. 33, 453–467 (2010).
26. Chakraborty, N. & Tripathy, B. C. Involvement of singlet oxygen in 5-aminolevulinic acid induced photodynamic damage of cucumber (Cucumis sativus L.) chloroplasts. Plant Physiol. 98, 7–11 (1992).
25. Zhou, Y ., Lam, H. M. & Zhang, J. Inhibition of photosynthetic oxygen evolution in rice stressed induced structural and functional reorganization of photosystem-II in wheat leaves. Physiol. Plant. 93, 771–777 (1995).
24. Giardi, M. T. et al. Long-term drought stress induces structural and functional reorganization of photosystem II. Planta 199, 118–125 (1996).
23. He, J. X., Wang, J. & Liang, H. G. Effects of water-stress on photochemical function and protein-metabolism of photosystem-II in greening wheat leaves. Physiol. Plant. 93, 771–777 (1995).
22. Alberte, R. S., Fiscus, E. L. & Naylor, A. W. The effects of water stress on the development of the photosynthetic apparatus in greening leaves. Plant Physiol. 55, 317–321 (1975).
21. Balch, J. L. & Gross, R. W. The protochlorophyllide oxidoreductase. FEBS J. 207, 405–411 (1997).
20. Sayre, R. T. & Chemiae, G. M. Studies on the reconstitution of O₂-evolution in chloroplasts. Plant Physiol. 69, 1084–1095 (1982).
19. Hodges Delong, J. M., Forney, C. F . & Prange, R. K. Improving the thiobarbituric acid-reactive substances assay for estimating lipid peroxidation in plant tissues containing anthocyanin and other interfering compounds. Plant Physiol. 207, 604–611 (1999).
18. Sharkey, T. D. & Seemann, J. R. Mild water stress effects on carbon-reduction-cycle intermediates, ribulose bis-phosphate oxygenase in intact isolated chloroplasts. J. Exp. Bot. 58, 1207–1217 (2007).
17. Mehlcr, A. H. Studies on reactions of illuminated chloroplasts: I. Mechanism of the reduction of oxygen and other Hill reagents. Arch. Biochim. Biophys. 33, 65–77 (1951).
16. Miller, G., Suzuki, N., Ciftci-Yilmaz, S. & Mittler, R. Reactive oxygen species homeostasis and signaling during drought and salinity stresses. Plant Cell Environ. 33, 453–467 (2010).
15. Chakraborty, N. & Tripathy, B. C. Involvement of singlet oxygen in 5-aminolevulinic acid induced photodynamic damage of cucumber (Cucumis sativus L.) chloroplasts. Plant Physiol. 98, 7–11 (1992).
14. Ford, K. L., Cassin, C. & Bacic, A. Quantitative proteomic analysis of wheat cultivars with differing drought tolerance. J. Exp. Bot. 68, 744–748 (1988).
13. Kottapalli, K. R. The protochlorophyllide holochrome of barley (Hordeum vulgare L.): the effect of light on the NADPH:protochlorophyllide oxidoreductase. FEBS J.
12. Gazanchian, A., Hajheidari, M., Sima, N. K. & Salekdeh, G. H. Proteome response of Elymus elongatum to severe water stress and recovery. J. Exp. Bot. 58, 291–300 (2007).
11. Kottapalli, K. R. et al. Physiology and proteomics of the water-deficit stress response in three contrasting peanut genotypes. Plant Cell Environ. 32, 380–407 (2009).
10. Ford, K. L., Cassin, C. & Baci, A. Quantitative proteomic analysis of wheat cultivars with differing drought tolerance. Front. Plant Sci. 2, 44 (2011).
9. Mirzaei, M. et al. Manipulating root water supply elicits major shifts in the shoot proteome. J. Prot. Res. 7, 517–26 (2014).
8. Faghihi, E. et al. Comparative physiology and proteomic analysis of two wheat genotypes contrasting in drought tolerance. J. Prot. 114, 1–15 (2015).
7. Robinson, S. P. & Portis, A. R. Involvement of stomatal ATP in the light activation of ribulose-1, 5-bisphosphate carboxylase/oxygenase in intact isolated chloroplasts. Plant Physiol. 86, 293–298 (1988).
6. Sharkey, T. D. & Seematz, I. R. Mild water stress effects on carbon-reduction-cycle intermediates, ribulose bis-phosphate carboxylase activity & spatial homogeneity of photosynthesis in intact leaves. Plant Physiol. 89, 1060–1065 (1989).
5. Graan, T. & Boyer, J. S. Very high CO₂ partially restores photosynthesis in sunflower at low water potentials. Planta 181, 378–384 (1990).
4. Bota, J., Medrano, H. & Flexas, J. Is photosynthesis limited by decreased Rubisco activity and RuBP content under progressive water stress? New Physiol. 162, 671–681 (2004).
3. Toivonen, P. & Vidaver, W. Variable chlorophyll a fluorescence and CO₂ uptake in water stressed white spruce seedlings. Plant Physiol. 86, 744–748 (1988).
2. Alberte, R. S., Fiscus, E. L. & Naylor, A. W. The effects of water stress on the development of the photosynthetic apparatus in greening leaves. Plant Physiol. 55, 317–321 (1975).
1. He, J. X., Wang, J. & Liang, H. G. Effects of water-stress on photochemical function and protein-metabolism of photosystem-II in wheat leaves. Physiol. Plant. 93, 771–777 (1995).
55. Dwyer, S. A. et al. Antisenes reductions in the PsbO protein of photosystem II leads to decreased quantum yield but similar maximal photosynthetic rates. *J. Exp. Bot.* 63, 4781–4795 (2012).

56. Kanazawa, A. & Kramer, D. M. In vivo modulation of nonphotochemical quenching (NPQ) by regulation of chloroplast ATP synthase. *Proc. Natl. Acad. Sci. USA* 99, 12789–12794 (2002).

57. Mullet, J. E., Burke, J. J. & Arnezen, C. J. Chlorophyll proteins of photosystem I. *Plant Physiol.* 65, 814–822 (1980).

58. Kuang, T. Y., Argyroudi-Akoyunoglou, J. H., Nakatani, H. Y., Watson, J. & Arntzen, C. J. The origin of the long-wavelength fluorescence emission band (77 degrees K) from photosystem I. *Arch. Biochem. Biophys.* 235, 618–627 (1984).

59. Briantais, J. M., Krause, G. H., Vernotte, C. & Weis, E. Chlorophyll a fluorescence of higher plants, chloroplasts and leaves in Light Emission by Plants and Bacteria (eds Govindjee, Ames, J. & Fork, D. C.) 539–583 (Academic Press, 1986).

60. Williams, R. S., Allen, J. F., Brain, A. P. R. & Ellis, R. J. Effect of Mg2+ on excitation energy transfer between LHClI and LHCl in a chlorophyll–protein complex. *FEBS Lett.* 225, 59–66 (1987).

61. Rubin, B. T., Chow, W. S. & Barber, J. Experimental and theoretical considerations of mechanisms controlling cation effects on thylakoid membrane stacking and chlorophyll fluorescence. *Biochim. Biophys. Acta* 634, 174–190 (1981).

62. Kaňa, R. & Govindjee. Role of ions in the regulation of light harvesting. *Front. Plant Sci.* 7, 1849 (2016).

63. Medrano, H., Parry, M. A. J., Socías, X. & Lawlor, D. W. Long term water stress inactivates Rubisco in subterranean clover. *Ann. Appl. Biol.* 131, 491–501 (1997).

64. Tripathy, B. C. & Oelmüller, R. Reactive oxygen species generation and signaling in plants. *Plant Sig. Behav.* 7, 1621–1633 (2012).

65. Zivcak, M. et al. Photosynthetic electron transport and specific photoprotective responses in wheat leaves under drought stress. *Photosynth. Res.* 117, 529–546 (2013).

66. Murashige, T. & Skoog, F. A revised medium for rapid growth and bio-assays with tobacco tissue cultures. *Plant Physiol.* 15, 473–497 (1962).

67. Tripathy, B. C. & Chakraborty, N. 5-aminolevulinic acid induced photodynamic damage of the photosynthetic electron transport chain of cucumber (*Cucumis sativus* L.) cotyledons. *Plant Physiol.* 96, 761–767 (1991).

68. Tripathy, B. C. & Mohanty, P. Zinc-inhibited electron transport of photosynthesis in isolated barley chloroplasts. *Plant Physiol.* 66, 1174–1178 (1980).

69. Graham, R. C. & Karnovsky, M. J. The early stages of absorption of injected horseradish peroxidase in the proximal tubules of mouse kidney: Ultrastructural cytochemistry by a new technique. *J. Histochem. Cytochem.* 14, 291–302 (1966).

70. Stibet, A. & Govindjee. On the relation between the Kautsky effect (chlorophyll a fluorescence induction) and photosystem II: Basis and applications of the OJIP fluorescence transient. *J. Photochem. Photobiol. B Biol.* 104, 236–257 (2011).

71. Papageorgiou, G. C. & Govindjee (eds) Chlorophyll a Fluorescence: A Signature of Photosynthesis. Advances in Photosynthesis and Respiration, 19. (Springer, Dordrecht, 2004).

72. Bradford, M. M. A rapid and sensitive method for the quantification of microgram quantities of protein utilizing the principle of protein-dye binding. *Annal. Biochem.* 72, 248–254 (1976).

73. Papageorgiou, G. C. & Govindjee (eds) Chlorophyll a Fluorescence: A Signature of Photosynthesis. Advances in Photosynthesis and Respiration, 19. (Springer, Dordrecht, 2004).

74. Velikova, V., Yordanov, I. & Edreva, A. Oxidative stress and some antioxidant systems in acid rain-treated bean plants: protective role of glutathione and ascorbate. *J. Exp. Bot.* 52, 1537–1546 (2001).

75. Kanazawa, A. & Kramer, D. M. Enzymatic determination of the chlorophyll and xanthophyll content of leaves. *Plant Physiol.* 76, 1057–1073 (2013).

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**Author Contributions**

B.C.T. designed the experiments and V.K.D. conducted all the experiments. Manuscript was written by V.K.D. and B.C.T.

**Additional Information**

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