The Physiological Role of Ascorbate as Photosystem II Electron Donor: Protection against Photoinactivation in Heat-Stressed Leaves

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Previously, we showed that ascorbate (Asc), by donating electrons to photosystem II (PSII), supports a sustained electron transport activity in leaves in which the oxygen-evolving complexes were inactivated with a heat pulse (49°C, 40 s). Here, by using wild-type, Asc-overproducing, and -deficient Arabidopsis (Arabidopsis thaliana) mutants (miox4 and vtc2-3, respectively), we investigated the physiological role of Asc as PSII electron donor in heat-stressed leaves (40°C, 15 min), lacking active oxygen-evolving complexes. Chlorophyll-a fluorescence transients show that in leaves excited with trains of saturating single-turnover flashes spaced 200 ms apart, allowing continual electron donation from Asc to PSII, the reaction centers remained functional even after thousands of turnovers. Higher flash frequencies or continuous illumination (300 μmol photons m⁻² s⁻¹) gradually inactivated them, a process that appeared to be initiated by a dramatic deceleration of the electron transfer from Tyr Z to P680⁺, followed by the complete loss of charge separation activity. These processes occurred with half-times of 1.2 and 10 min, 2.8 and 23 min, and 4.1 and 51 min in vtc2-3, the wild type, and miox4, respectively, indicating that the rate of inactivation strongly depended on the Asc content of the leaves. The recovery of PSII activity, following the degradation of PSII proteins (D1, CP43, and PsbO), in moderate light (100 μmol photons m⁻² s⁻¹, comparable to growth light), was also retarded in the Asc-deficient mutant. These data show that high Asc content of leaves contributes significantly to the ability of plants to withstand heat-stress conditions.

Heat stress is a serious threat to plants, which can lead to a drastic reduction in crop yield (e.g. Wahid et al., 2007; Allakhverdiev et al., 2008). Exposure of plants to elevated temperatures results in the inactivation of Rubisco activase (Barta et al., 2010) and the oxygen-evolving complex (OEC) of PSII, including the removal of the extrinsic proteins as well as the release of calcium and manganese ions from their binding sites (Nash et al., 1985; Enami et al., 1994; Yamane et al., 1998; Barra et al., 2005). Heat stress can also damage the D1 and D2 proteins (De Las Rivas and Barber, 1997; Yoshioka et al., 2006).

Under natural conditions heat stress mostly occurs together with light stress. Though the effects of both stresses have been studied extensively, there are only a few studies where the mechanism of damage caused by combination of these two stress factors was investigated. It has been shown that in the absence of heat-adaptation processes, with yet an unknown mechanism (s), the two stress factors act in strong synergy to inactivate PSII in plants (Havaux, 1994), and in corals they lead to photobleaching (Abrego et al., 2008).

Inactivation of PSII by excess light and damage to the D1 protein are most probably induced by singlet oxygen, which is produced via interaction with the triplet reaction center chlorophyll (chl; ³P680) arising from the recombination of the charge-separated state between P680 and the pheophytin electron acceptor (⁵P680’Phe); Aro et al., 1993; Vass and Cser, 2009). This acceptor-side damage, as it has recently been shown by point mutation at the quinone-binding pocket of PSII in Synechocystis sp. PCC 6803 (Larom et al., 2010), can be alleviated by efficiently draining the electrons from the primary quinone acceptor QA to an exogenous electron acceptor, cytochrome c. In other terms it appears that photoinhibition can be prevented by warranting a rapid electron transfer from the acceptor side of PSII. A priori, it cannot be ruled out that photoinhibition under heat-stress conditions occurs with the same mechanism since heat stress can lead to...
a deceleration of the electron transfer rate between QA and Q*B, the primary and secondary quinone acceptors (Ducruet and Lemoine, 1985). However, elevated temperatures affect mainly the donor side of PSII, in particular the OEC. Experiments on PSII preparations and on leaves with chemically inactivated OECs indicate that the very high sensitivity of PSII to light appears to be caused by the impaired electron donation from the OEC. This, in turn, results in strong accumulation of highly oxidizing radicals, P680+, TyrZ+, and superoxide (Chen et al., 1995) or hydroxyl radicals (Spetea et al., 1997), and leads to a rapid inactivation and degradation of PSII reaction centers (Callahan et al., 1986; Blubaugh and Cheniae, 1990; Jegerschöld and Styring, 1996). This type of photodamage is called weak light or donor-side-induced photoinhibition. In analogy with the acceptor-side photoinhibition, it can be assumed, in fact in vitro data strongly suggest, that a continuous electron flow to PSII from alternative donors, such as diphenylcarbazide (DPC) or ascorbate (Asc), can alleviate the photoinactivation of the reaction centers (Mano et al., 1997).

In this article we provide experimental evidence that Asc plays a protective role in photoinhibition in heat-stressed leaves. Asc is present in the lumen probably at millimolar concentration (Foyer and Lelandais, 1996), and in the absence of active OEC serves as a relatively rapid ($t_{1/2}$ approximately 25 ms) electron donor to PSII (Tóth et al., 2009), and thus might be capable of protecting PSII by supplying electrons to the reaction center. To test this hypothesis, we subjected intact leaves of wild-type, Asc-overproducing ($miox4$; Lorence et al., 2004), and -deficient mutant ($vtc2-3$; Conklin et al., 2000) Arabidopsis (Arabidopsis thaliana) plants to heat stress (40°C, 15 min) and investigated the time course and mechanism of photoinactivation of PSII.

**RESULTS**

**Electron Donation Rates from Asc to PSII in Arabidopsis Genotypes with Different Asc Contents**

To study the physiological role of Asc as a PSII donor, we compared wild-type Arabidopsis, Asc-overproducing ($miox4$; Lorence et al., 2004), and Asc-deficient mutants ($vtc2-3$; Conklin et al., 2000) grown under identical conditions, at approximately 150 μmol photons m$^{-2}$ s$^{-1}$, 8-h light/16-h dark, the temperature was kept between 20°C and 24°C, and the plants were used when they were 2 to 3 months old. The Asc-deficient mutants were about 30% smaller than the wild-type plants (as reported by Müller-Moulé et al., 2004 as well) and the plants were somewhat larger (Fig. 1A). The Asc content of the $vtc2-3$ mutant was about 25% of the wild type and the Asc content of the $miox4$ mutant was about 70% higher than that of the wild type determined on chl basis (Fig. 1B; there were only minor differences between the chl contents of the three genotypes; data not shown). We also determined the nonphotochemical quenching (NPQ) and its energy-dependent quenching (qE) component, and...
the rate of electron transport (ETR) in the leaves. NPQ and qE were highest in the Asc-overproducing mutant (miox4) and lowest in the Asc-deficient mutant (vtc2-3) and there were no significant differences between the three genotypes in terms of ETR values (Fig. 1B). The results obtained on the vtc2-3 mutant are in agreement with the results of Müller-Moule et al. (2002, 2003), who showed that the lower NPQ values are caused by the lower violaxanthin deepoxidase activity due to limitation in the available Asc in the lumen. The higher NPQ and qE values obtained on the miox4 mutant indicate that there is more Asc available in the lumen of the Asc-overproducing miox4 mutant than in the wild type. Determination of the luminal Asc content does not seem to be feasible since upon chloroplast isolation Asc is mostly lost (Ivanov and Edwards, 2000).

Detached leaves of wild-type, Asc-overproducing (miox4), and Asc-deficient (vtc2-3) Arabidopsis plants were subjected to heat stress at 40°C for 15 min in a water bath in the dark. This treatment was chosen to obtain a homogenous material with completely inactivated OECs and to separate the effects of heat and light stresses. The B thermoluminescence (TL) band (Vass, 2003) arising from recombination reactions between the S2 state of the OEC and QB+ induced by a saturating single turnover flash (STTF) was totally abolished by the heat treatment in all three genotypes, showing a total loss of oxygen-evolving activity (Fig. 1C). A 40°C heat treatment is physiologically relevant since leaf temperatures can easily reach 40°C in the field, especially under drought-stress conditions (Burghardt et al., 2008; Shahenshah and Isoda, 2010). This treatment leads to a moderate, less than 2-fold acceleration of the decay kinetics of the flash-induced electrochromic absorbance transient (Δ515), an indicator of the dissipation of the transmembrane electrical field (Junge, 1977): The t1/2 of the decay decreased from 55 to 34 ms, with similar behavior in the three genotypes. These data show that the permeability of the thylakoid membranes did not increase drastically as a result of the 40°C, 15-min heat treatment (in contrast to a 50°C heat treatment applied earlier that caused dissipation of the electric field within a few ms; Tóth et al., 2005); these data agree well with the results obtained on heat-treated Arabidopsis leaves (Krumova et al., 2010). Decrease in the initial amplitude of the flash-induced electrochromic absorbance transient was also observed, suggesting a partial inactivation of PSII reaction centers (data not shown; see below).

The fast chl a fluorescence (OJIP) transients (Govindjee, 2004; Lazár and Schansker, 2009) of untreated Asc-overproducing, Asc-deficient, and wild-type Arabidopsis plants were essentially the same, with identical Fv/Fm values (data not shown). The OJIP transient reflects the reduction of the electron transport chain including PSI (Schansker et al., 2005). In leaves containing PSII reaction centers with inactive OEC the K peak appears at around 300 μs with a concomitant disappearance of the J and I steps (Fig. 2). The K peak represents approximately one stable charge separation, with TyrZ as the electron donor (Srivastava et al., 1997; Tóth et al., 2007). After the K peak fluorescence intensity decreases to a level approaching Fm in a few ms due to the reoxidation of QA+ by QB. In leaves a second peak appears at around 1 s (Fig. 2). We have shown previously that this latter phase is due to electron donation by Asc to PSII, leading to a partial reduction of the electron transport chain as shown by P700 measurements as well (Tóth et al., 2009). In heat-stressed (40°C, 15 min) Asc-deficient plants this second rise phase was considerably smaller than in the wild type and it was somewhat retarded, whereas in the Asc-overproducing mutant its intensity was somewhat higher than in the wild type (Fig. 2). This is in agreement with our previous finding that the intensity of this peak depends on the Asc content of the leaves (Tóth et al., 2009).

The t1/2 of electron donation from Asc to TyrZ+ can be determined in samples with fully inactivated oxygen evolution by using two short (5-ms) light pulses and by varying the dark interval between them (Tóth et al., 2007, 2009). During the 5-ms light pulse one charge separation and the reoxidation of QA+ by QB takes place. The inset of Figure 2 shows that after a 2.3-ms dark interval following the first light pulse
there is no variable fluorescence, which is due to the full inactivation of oxygen evolution. However, with longer dark intervals, the K peak recovers following single exponential kinetics and the $t_{1/2}$ of the regeneration of the K peak can be used as the half-time of the rereduction of Tyr$\gamma$ by Asc (Tóth et al., 2009). The $t_{1/2}$ was approximately 30 ms in wild-type and miox4 leaves, whereas in the Asc-deficient mutant the electron donation was much slower, the $t_{1/2}$ was approximately 50 ms (Table I). These results on heat-stressed leaves (40°C, 15 min) are in a good agreement with our previous findings where plants were subjected to a relatively harsh heat pulse of 49°C, 40 s (Tóth et al., 2009).

**Asc-Dependent Multiple Turnover of PSII**

To ascertain the notion that in heat-stressed leaves Asc can continually support the electron transport through PSII, i.e. it sustains multiple turnovers of the reaction centers, without damage, we carried out experiments with trains of SSTFs. Figure 3A shows that when wild-type leaves were subjected to SSTFs that were spaced 200 ms apart, the K peak remained essentially unaffected even after 18,000 turnovers (flashing the leaves for 1 h). Similar data were obtained in vtc2-3 with 500-ms spacing between the flashes (data not shown). This shows that if the dark interval between the flashes is large enough to allow a continual electron donation from Asc to PSII ($t_{1/2}$ approximately 30 and 50 ms in the wild type and the Asc-deficient mutant, respectively), there is no apparent limitation for the turnover of the reaction centers. In other terms, the pool of Asc available to PSII reduction is evidently very large; also, under these conditions, no significant damage appears to occur to PSII. A closer spacing of the SSTFs (e.g. $\Delta t$ = 100 ms), which did not allow full regeneration of the K peak between flashes, led to the diminishment of the amplitude of the K peak (Fig. 3B). The decrease was strongest in vtc2-3 (the amplitude decreased by about 70% after 6,000 SSTFs) and weakest (30%) in miox4 (Fig. 3C). The decrease of the K peak progressed gradually. In the first phase, after about five to 10 flashes, the amplitude of the K peak decreased due to the approximately 9% increase in the $F_{20us}$ value. This was followed by a much slower decrease in the amplitude of the K peak that strongly depended on the Asc content. The recovery of the K peak in the dark was also biphasic (Fig. 3D). The fast recovery of the K peak, occurring with $t_{1/2}$ of about 4 s, can be accounted for by the recovery of non-Q$\beta$-reducing centers (Chylla et al., 1987). The large, hardly recovering phase indicates that SSTFs in long train cause irreversible damage to PSII.

**Inactivation of PSII Reaction Centers in Heat-Stressed Leaves Exposed to Continuous Illumination**

To investigate in more detail the photoinactivation of PSII and the influence of the Asc content, heat-stressed leaves (40°C, 15 min) were subjected to relatively strong continuous illumination (300 μmol photons m$^{-2}$ s$^{-1}$). This treatment resulted in relatively fast and gradual diminishment of the K peak as well as of the second

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**Table I.** Half-times of electron donation to PSII (in heat-stressed wild-type and vtc2-3 Arabidopsis plants) and from Asc + DPC (in DPC-treated and heat-stressed vtc2-3 Arabidopsis mutant) and the half-time of the decrease of the amplitude of the K peak (as determined in Fig. 3B) in heat- + light-treated wild-type and vtc2-3 Arabidopsis plants

| Parameters | Wild Type | vtc2-3 | vtc2-3 + 1 mM DPC |
|------------|-----------|--------|------------------|
| Half-time of electron donation to PSII | 30.5 ± 1.3 ms | 49.7 ± 3.6 ms | 30.4 ± 2.5 ms |
| Half-time of the decrease of the K peak in the light | 2.8 ± 0.5 min | 1.4 ± 0.1 min | 2.5 ± 0.3 min |
peak at around 1 s (Fig. 4A). The rate of the diminishment of the K peak followed exponential kinetics and depended strongly on the Asc content of leaves: In the wild type it exhibited a $t_{1/2}$ of 2.8 min, whereas the $vtc2$-$3$ and $miox4$ $t_{1/2}$ values were 1.2 and 4.1 min, respectively (Fig. 4B).

To examine the relative amount of active PSII reaction centers, leaves were incubated in the dark for 2 h in 3-(3',4'-dichlorophenyl)-1,1-dimethylurea (DCMU) solution after the heat + light treatment (Fig. 4C) to ensure that all PSII reaction centers are closed during the fluorescence measurement. It can be seen that the heat stress per se induced about 15% decrease in the $F_m$ value. Upon light treatment of heat-stressed samples, the fluorescence intensity gradually decreased and after 4 h the $F_m$ value almost equaled $F_0$. Again, the decrease in the amplitude was significantly faster in the Asc-deficient mutant than in the wild type (apparent $t_{1/2}$ 10 and 23 min, respectively). This difference was even more pronounced when comparing it with the Asc-overproducing mutant ($t_{1/2}$ approximately 51 min; Fig. 4D). Very similar rates were obtained in the presence of the protein synthesis inhibitor lincomycin, showing that in 4 h, no significant recovery occurred (data not shown). It is also interesting to note that in the light-treated leaves the fluorescence rise was strongly decelerated: In the wild type, the $t_{1/2}$ increased from 0.3 to 2.3 ms in 1 h; deceleration was even stronger in $vtc2$-$3$ (from 0.3–3 ms).

The fast decrease in the amplitude of the K peak and the slowdown of the fluorescence rise determined in the presence of DCMU suggest that the first step of photoinactivation is the slowdown of the TyrZ-P680$^+$ (as suggested by Blubaugh et al., 1991) and this is followed by a complete inactivation of the charge separation activity of PSII (see “Discussion” for more details). Our data also show that the rate of PSII inactivation depends strongly on the Asc content of the leaves. Asc is not only an alternative electron donor to PSII but also a direct scavenger of reactive oxygen species (ROS) produced during donor-side-induced photoinhibition (e.g. Chen et al., 1995; Spetea et al., 1997). To ascertain that the Asc dependence correlates with the ability of Asc to act as a PSII electron donor, we incubated intact $vtc2$-$3$ leaves in DPC solution (1 mM DPC, an artificial electron donor of PSII, with no ROS-scavenging properties) before the heat treatment. In DPC-treated $vtc2$-$3$ leaves the $t_{1/2}$ of electron donation to PSII became similar to that in the wild type (about 30 ms; Table I) and the fluorescence intensity at around 1 s also increased. In DPC-incubated leaves exposed to light, the K peak diminished more slowly than in untreated leaves ($t_{1/2}$s of 2.5 and 1.4 min, respectively), at a rate similar to that found in wild-type leaves (2.8 min; see also Fig. 4B). This is a strong indication that Asc slows down the inactivation of the PSII reaction centers by acting as PSII electron donor and not only as a scavenger of ROS.

Figure 4. Chl a fluorescence transients (A and C) and the time course of the decrease of the K peak (B) and of the $F_m$ value in the presence of DCMU (D) on heat-stressed wild-type (WT), Asc-overproducing ($miox4$), and -deficient ($vtc2$-$3$) Arabidopsis leaves exposed to continuous white light of 300 μmol photons m$^{-2}$ s$^{-1}$. The amplitude of the K peak on heat-stressed samples was calculated as $F_{300}$ - $F_{20}$, from traces shown in A, and data points were fitted with single exponentials, yielding $t_{1/2}$ values as indicated in the figure. DCMU treatments were performed after the light treatments of heat-stressed leaves (C and D).
It is also known that Asc is required for the formation of NPQ (more specifically, qE), because it is a substrate for violaxanthin deepoxidase (Hager, 1969). qE plays a photoprotective role and therefore we have to consider if the faster rate of PSII inactivation observed in the vtc2-3 mutant might be caused by smaller qE. Determination of NPQ and qE in heat-treated samples is not possible because $F_m$ can only be reached when DCMU is present (Fig. 4; see also Tóth et al., 2007). However, the data show that the $t_{1/2}$ of electron donation from Asc to PSII in heat-treated samples is slow compared to electron donation by active OECs (30–50 ms versus 0.1–1 ms; Babcock et al., 1976) and therefore, significant and persistent reduction of the electron transport chain, a prerequisite for the formation of qE, is very unlikely: Illumination of samples with completely inactivated OECs at 300 μmol photons m$^{-2}$ s$^{-1}$ would result in electron transport comparable to that occurring at around 10 μmol photons m$^{-2}$ s$^{-1}$ in the case of non-heat-treated samples. Light intensities below 40 μmol photons m$^{-2}$ s$^{-1}$ are insufficient for the formation of qE in any of the three genotypes studied (data not presented; see Müller-Moulé et al., 2002), therefore photoprotection by qE formation during our photoinactivation experiments is unlikely to occur.

As shown in Figure 5, in untreated wild-type leaves, fast oxidation of P700 was followed by rereduction in about 200 ms due to electrons arriving from PSI. In heat-treated leaves, in which the electrons were donated by Asc, the rereduction was slower, in good accordance with our earlier data obtained on leaves exposed to heat pulses (49°C, 40 s; Tóth et al., 2009). Light treatment of heat-stressed leaves caused further, gradual deceleration of the rereduction and after about 1 h only a very limited rereduction could be observed. These data show that upon light treatment of heat-stressed leaves, electron transport to PSI was strongly decelerated or even stopped and this might lead to some loss in the amount of P700, as indicated by the slight decrease in the amplitude after 1 and 4 h of illumination, compared to DCMU-treated leaves.

Photoinactivation was studied in leaves with partially inhibited OEC activity as well. To this end, wild-type leaves were subjected to a 15-min heat stress at 38°C, inactivating about 40% of oxygen evolution, as estimated on the basis of TL measurements (data not shown). Accordingly, the fluorescence transient (Fig. 6) carries the characteristics of both the untreated leaves and those that contain no active OECs (compare with Fig. 4A): The variable fluorescence decreased, albeit less than after the 40°C treatment, and $F_m$ peaked at around 1 s, similarly to the sample with fully inactivated OEC, but also exhibited a shoulder at 200 ms, characteristic of the untreated leaves. Illumination (300 μmol photons m$^{-2}$ s$^{-1}$) caused a further decrease of the variable fluorescence that was mostly due to the disappearance of the peak around 1 s, showing that the contribution of reaction centers with inactive OECs gradually decreased during the light treatment, i.e., they became inactivated by light as in leaves with fully inhibited oxygen evolution (Fig. 4A). Indeed, after 1 h illumination the OJIP transient exhibited the typical steps, but with lower amplitudes. After 4 h, a significant part of this loss was recovered in the light.

**Light-Induced Degradation of PSII Proteins in Heat-Stressed Leaves**

As shown above, after 4 h of illumination of heat-stressed leaves (40°C, 15 min; Fig. 4C), there was a complete loss of charge separation activity of PSII reaction centers. During acceptor-side photoinhibition, the inactivation of the reaction centers is accompanied by protein degradation, which is confined to the D1 protein (Schuster et al., 1988; Aro et al., 1993; van Wijk et al., 1994). To investigate if PSII reaction centers in heat-stressed leaves suffer the same or different light-induced damage, western-blot analyses were performed, using wild-type leaves.

Data in Figure 7 show that upon the heat stress about 20% of D1 protein was lost, which is in agreement with earlier results (Yoshioka et al., 2006; Yamashita et al., 2008). Light treatment on heat-stressed samples resulted in further, gradual degradation of D1: In 1 h D1 content decreased by about 40% relative to the untreated control and after 4 h approximately 65% of the D1 protein was lost. These data are in good agreement with data obtained with chl $a$ fluorescence measurements in the presence of DCMU (Fig. 4C). In contrast to acceptor-side photoinhibition, we also observed significant losses in the amount of CP43 and the PsbO protein; by the 4th h of the light treatment of
heat-treated leaves, 70% and 80% of CP43 and PsbO were lost, respectively. These data suggest that there was a complete disassembly of PSII reaction centers upon the light treatment of heat-stressed leaves.

**Time Course of PSII Inactivation and Recovery**

We investigated the effect of Asc also during the light-dependent recovery of PSII activity following photoinactivation in heat-stressed leaves. The experiments were carried out at moderate light intensity, 100 μmol photons m⁻² s⁻¹ that is comparable to the growth light. In the dark neither significant damage nor recovery occurs (Tóth et al., 2005). On the other hand, at 300 μmol photons m⁻² s⁻¹ recovery was very slow and some photobleaching occurred (data not shown). It can be seen that at 100 μmol photons m⁻² s⁻¹ the K peak substantially diminished in a few hours (Fig. 8A), indicating a significant perturbation of the charge separation activity of PSII reaction centers, similar to the case at the higher light intensity i.e. at 300 μmol photons m⁻² s⁻¹. This was followed by a gradual recovery of the OJIP transient, which was completed in about 24 h. To monitor the recovery of PSII activity in the vtc2-3, miox4, and wild-type plants, we used the \( \frac{F_{t=100\text{ms}} - F_{t=20\mu s}}{F_{t=100\text{ms}}} \) parameter introduced earlier (Tóth et al., 2005), rather than the \( \frac{F_v}{F_m} \) ratio, which in the presence of inactive OEC also depends on the Asc content of the leaves (compare with Fig. 2). Our data show that the rate of recovery was significantly faster in the wild type and in miox4 than in vtc2-3, which appeared to suffer more substantial damage (Fig. 8B).

**DISCUSSION**

**Asc as a PSII Donor**

In this work, by using Asc-overproducing (miox4), -deficient (vtc2-3), and wild-type plants, we show that Asc retards the photoinactivation of PSII in heat-stressed leaves with inactive OECs by acting as a PSII donor. We also provide evidence that photoinhibition in heat-stressed leaves occurs with a similar mechanism and time course as described for samples with chemically inactivated OECs (Blubaugh et al., 1991; Jegerschöld and Styring, 1996).

Our experiments on heat-stressed leaves subjected to trains of SSTFs show that the reaction centers are capable of tens of thousands of turnovers, without noticeable damage—provided that the electron donation rate by Asc warrants full rereduction of Tyr-Z⁺ between flashes. This depends on the flash frequency and the Asc content of the leaves (Fig. 3). These data show that the Asc pool was capable to support tens of thousands of stable charge separations in PSII. Indeed, it has been estimated that the Asc content of chloroplasts is about 25 to 50 mM (Eskling and Åkerlund, 1998; Smirnoff, 2000) and at least 4 mM in the thylakoid lumen (Foyer and Lelandais, 1996). The transport through the chloroplast envelope membranes is relatively fast and is rapidly equilibrated (Foyer and...
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Lelandais, 1996; it is probably equally fast through the thylakoid membrane (Mano et al., 2004). Oxidized Asc regenerates in the Halliwell-Asada cycle and probably by other means as well (Potters et al., 2002).

**Donor-Side-Induced Photoinhibition and Dependence on the Asc Content**

When the dark intervals between SSTFs were not sufficiently long (in wild type and vtc2-3, shorter than 100 and 200 ms) and appeared not to allow full regeneration of the K peak between flashes the amplitude of the K peak gradually diminished (Fig. 3). The extent of this diminishment, which was largely irreversible, depended strongly on the Asc content of the leaves and occurred in several minutes (Fig. 3C). In continuous light of 300 μmol photons m\(^{-2}\) s\(^{-1}\), the amplitude of the K peak decreased with 1.2, 2.8, and 4.1 min \(t_{1/2}\) in vtc2-3, the wild type, and mixo4, respectively. This diminishment of the K peak was irreversible, indicating damage to the reaction center. The fact that Asc in vtc2-3 could be replaced with DPC (Table I) points to the role of this endogenous alternative electron donor in photoprotection, and shows that the disappearance of the K peak is indeed correlated with limitation in electron donation. In the presence of DCMU variable fluorescence was significantly higher than in its absence (up to about 1 h of illumination), indicating that charge separation activity of PSII is yet retained, albeit the fluorescence rise gradually slowed down during the illumination period. Similar phenomenon was observed by Callahan and Cheniae (1985) in hydroxylamine-treated leaves. The deceleration of the fluorescence rise originates from slowing down of the electron transfer between Tyr\(_{x}\) and P680\(^*\), as suggested earlier by Blubaugh et al. (1991) based on electron paramagnetic resonance measurements. The disappearance of the K peak, together with the relatively high \(F_m\) value detected in the presence of DCMU, shows that in the heat- + light-treated samples the rate of Tyr\(_{x}\)-P680’ electron transfer becomes comparable to the rate of QA to Q\(_B\) electron transfer, i.e. there is a deceleration of several orders of magnitude, from about 100 ns to at least hundreds of μs.

It is known that heat stress might lead to a slowdown of electron transport between QA and Q\(_B\) (Ducruet and Lemoine, 1985). However, OEC is more sensitive to high temperature than the acceptor side of PSII and slowdown of electron transfer from QA to Q\(_B\) can be observed only above 42°C (Pospšíšil and Tyystjärvi, 1999). Thus, the occurrence of acceptor-side photoinhibition is quite unlikely. Also, our data showing that in the presence of DCMU much higher \(F_m\) values could be obtained than in its absence points to a much stronger limitation at the donor side of PSII as well. Further, in acceptor-side photoinhibition ROS are produced (e.g. Hideg et al., 1998), which would not be possible to slow down by DPC, an artificial PSII donor with no ROS-scavenging properties.

Gradual inactivation of charge separation activity of PSII in continuous light occurred on a time scale of tens of minutes, as reflected by the decrease of the \(F_m\) value in the presence of DCMU as well as by the redox changes of P700 (Figs. 4C and 5). Based on our data showing the parallel loss of CP43 and PsbO protein and on earlier experiments on isolated PSII and thylakoid membranes (Mori and Yamamoto, 1992; Shing and Shinghal, 1999), we suggest that donor-side-induced photoinhibition is accompanied by extensive protein degradation, during which probably the whole PSII reaction center disassembled. This is in contrast to acceptor-side photoinhibition where protein degradation is mostly confined to the D1 protein (Schuster et al., 1988; van Wijk et al., 1994). Again, the fluorescence transients in presence of DCMU clearly show that the rate of full inactivation of PSII reaction centers strongly depends on the Asc content of the leaves (Fig. 4D).

The rates of inactivation of PSII were very similar in the presence and absence of the protein synthesis inhibitor lincomycin (data not shown). This indicates that the damage and degradation of the D1 protein is...
Physiological Significance

In nature heat stress is usually accompanied by high light, and therefore, photoinactivation of PSII is very likely to occur at high temperatures. As concerns the physiological significance of the Asc-dependent retardation of photoinactivation of PSII, we must take into account that full protection could only be achieved at low frequencies of the exciting flashes, corresponding to very low light intensities, in the range of several μmol photons m⁻² s⁻¹. Higher flash frequencies or continuous illumination (100 and 300 μmol photons m⁻² s⁻¹) lead to losses in PSII activity (Figs. 4 and 8). Photoinactivation occurred in samples with partial OEC inhibition as well (38°C; Fig. 6) and even in the case of a very mild heat treatment (36°C, 1 min) PSII reaction centers did not recover on the time scale of tens of minutes (J. Frolec, personal communication), suggesting that the same mechanism is taking place as in the case of heat stresses abolishing all OEC activity. Our data showing complete inactivation of PSII (Figs. 4 and 6) and protein degradation (Fig. 7) suggest that recovery is preceded by a complete inactivation of PSII reaction centers and does not appear to occur directly from the OEC-inactivated state. When investigating the rate of recovery from the photoinactivated state, large differences were found between the Asc-deficient and the wild-type and Asc-overproducing plants (Fig. 8). This might indicate that since the rate of photoinactivation is faster in the Asc-deficient plants, during photoinactivation more ROS (hydroxyl radical or superoxide) can be produced, which in turn cause more severe damage, not only to the reaction centers but also other compounds of the thylakoid membranes; in addition, the repair processes might also be affected, which might also depend on the rate of ROS production (Nishiyama et al., 2006), and it is also conceivable that Asc acts as an electron donor during the synthesis of PSII units, before the extrinsic proteins are attached to the reaction center. In the light of these data and considerations, we propose that in heat-stressed leaves under natural conditions the role of Asc as an alternative PSII electron donor is to slow down the photoinactivation processes and by this means minimize the generation of ROS in the thylakoid membranes, and thus alleviate the damage to the entire photosynthetic machinery.

MATERIALS AND METHODS

Plant Material and Growth Conditions

Arabidopsis (Arabidopsis thaliana Columbia-0 [wild-type Arabidopsis]), Asc-deficient (vtc2-3; Conklin et al., 2000), and Asc-overproducing mutant (miox4; Lorence et al., 2004) plants were grown in a growth chamber under short-day conditions (8-h light, 16-h dark), at approximately 150 μmol photons m⁻² s⁻¹ in the light period. The temperature was kept between 20°C and 24°C.

The vtc2 mutants are affected in the Gal pathway of Asc biosynthesis, more specifically in GDP-ı-Gal phosphorylase, an enzyme that is at an branch point of between Asc synthesis and incorporation of ı-Gal into cell wall polysaccharides (Asc biosynthesis was reviewed by Linstner and Clarke, 2008). The miox4 mutant was created by constitutive expression of miox4, a gene encoding myoinositol oxygenase in Arabidopsis plants (Lorence et al., 2004).

Heat Treatments and Chemical Treatments

Heat treatment: Whole leaves were submerged in a water bath in darkness for 15 min. Complete inactivation of oxygen evolution was achieved by a treatment at 40°C or sometimes 41°C, depending on the age of the plants; inactivation of OEC was tested with the aid of TL measurements. To obtain partial inactivation of OEC, the heat treatment was performed at 38°C. Light treatments: Trains of SSTFs from a xenon lamp were given with dark intervals of 100, 200, and 500 ms for different time periods (10 min–1 h). Continuous light treatments of heat-treated leaves were carried out at 300 μmol photons m⁻² s⁻¹ and the treatment lasted for up to 4 h. For recovery experiments, heat-treated leaves were illuminated for 24 h at 100 μmol photons m⁻² s⁻¹, at 24°C. DPC treatment: vtc2-3 leaves were incubated in 1 mM DPC. Leaves were placed in petri dishes and covered with one layer of filter paper for 2 h; by this means, the leaves were exposed to white light of approximately 30 μmol photons m⁻² s⁻¹.

DCMU treatment: Whole leaves were incubated in 0.2 mM DCMU solution for 2 h in complete darkness after the heat + light treatment. The solution contained 0.2% dimethyl sulfoxide to dissolve the DCMU.

Determination of Asc Content

The Asc contents of wild-type Arabidopsis, vtc2-3, and miox4 mutants were determined by a spectroscopic method using the absorption at 265 nm of Asc (Takahama and Oniki, 1992).

TL Measurements

TL was measured using a custom-made TL apparatus described by Wiessler and Demeter (1988). Leaf discs were placed on a copper sample holder, connected to a cold finger immersed in liquid nitrogen. A heater coil, placed under the sample holder, ensured the desired temperature of the sample during the measurements. Thoroughly dark-adapted samples were illuminated at 1°C by a SSTF and TL was measured while heating the sample to 70°C in darkness with a heating rate of 20°C min⁻¹. The emitted TL was measured with a Hamamatsu end-window photomultiplier.

Measurement of Flash-Induced Electrochromic Absorbance Transients (ΔA515)

Electrochromic absorbance changes, induced by single turnover flashes, were measured at 515 nm, the maximum of the electrochromic transients, in a set up described by Büchel and Garab (1995). The time constant was set to 100 ms. All kinetic traces were collected with a repetition rate of 2 s⁻¹ and averaged. The transients were recorded at room temperature on detached Arabidopsis leaves.

Determination of ETR, NPQ, and qE; Measurement of the Oxidation-Reduction Kinetics of P700

The ETR, NPQ, and qE were determined by a Dual-PAM-100 instrument (Heinz Walz GmbH), on overnight dark-adapted leaves, with saturating pulses of 5,000 μmol photons m⁻² s⁻¹ during illumination at 430 μmol photons m⁻² s⁻¹ for 8 min (ETR, NPQ) and a subsequent dark adaptation for 2 min (qE). Redox changes of P700 were measured with the same instrument. The absorbance changes in continuous red light (5,000 μmol photons m⁻² s⁻¹) were measured at 830/870 nm with a time resolution of 60 μs.

OJIP Measurements

Fluorescence measurements were carried out at room temperature with a special version of the Handy-PEA instrument (Hansatech Instruments Ltd.)
that allows reducing the length of the measurement to 300 µs. Leaf samples were illuminated with continuous red light emitted by three LEDs (3,500 µmol photons m⁻² s⁻¹, 650 nm peak wavelength; the spectral half-width was 22 nm; the light is cut off at 700 nm by a near-infrared short-pass filter). The first reliably measured point of the fluorescence transient is at 20 µs, which was taken as F₀. The length of the measurements was 5 s or 5 ms. In the case of the double 5-ms pulses, the dark intervals between the light pulses were 2.3, 9.6, 16.9, 31.5, 38.8, 53.4, 75.3, 100, 200, or 500 ms.

Western-Blot Analysis

Leaf discs equivalent to a total area of 3.1 cm² cut from Arabidopsis leaves were frozen in liquid nitrogen and ground to a fine powder and then homogenized in 500 µL Laemmli buffer. The homogenates were incubated at 90°C for 5 min followed by a 20-min incubation at 37°C, and then proteins were separated by 15% denaturing SDS-PAGE. The proteins were blotted on nitrocellulose membranes using a semidry blotting system with methanol-containing buffer. The nitrocellulose membranes were blocked using 5% skim milk powder in Tris-buffered saline plus Tween 20 (TBST buffer) for 2 h and incubated with primary antibodies raised against PsbA (D1), PsbC (CP43), and PsbO (33 kD OEC protein; Agrisera AB) for 2 h in TBST buffer with 5% milk powder. The membranes were washed three times for 5 min in TBST buffer and incubated with goat anti-rabbit IgG horseradish peroxidase conjugate (Millipore) at a 1:1,000 dilution in TBST buffer with 5% milk powder for 2 h. Immunoblotted membranes were incubated for 5 min in ECL plus horseradish peroxidase substrate (GE Healthcare Bio-Sciences) and chemiluminescence was detected on Hyperfilm ECL photographic film (GE Healthcare Bio-Sciences). The developed film was digitalized and analyzed by 1D Scan software package.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers NM_118819.2 (ctc2) and NM_118759.4 (mixd4).

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The Physiological Role of Ascorbate as Photosystem II Donor

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