Anaerobiosis Induced State Transition: A Non Photochemical Reduction of PQ Pool Mediated by NDH in Arabidopsis thaliana

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Abstract

Background: Non photochemical reduction of PQ pool and mobilization of LHCII between PSII and PSI are found to be linked under abiotic stress conditions. The interaction of non photochemical reduction of PQ pool and state transitions associated physiological changes are critically important under anaerobic condition in higher plants.

Methodology/Findings: The present study focused on the effect of anaerobiosis on non-photochemical reduction of PQ pool which trigger state II transition in Arabidopsis thaliana. Upon exposure to dark-anaerobic condition the shape of the OJIP transient rise is completely altered where as in aerobic treated leaves the rise is unaltered. Rise in Fₙ and F₉ was due to the loss of oxidized PQ pool as the PQ pool becomes more reduced. The increase in Fₐ,⁰ was due to the non photochemical reduction of PQ pool which activated STN7 kinase and induced LHCII phosphorylation under anaerobic condition. Further, it was observed that the phosphorylated LHCII is migrated and associated with PSI supercomplex increasing its absorption cross-section. Furthermore, evidences from crr2-2 (NDH mutant) and pgr5 mutants (deficient in non NDH pathway of cyclic electron transport) have indicated that NDH is responsible for non photochemical reduction of the PQ pool. We propose that dark anaerobic condition accelerates production of reducing equivalents (such as NADPH by various metabolic pathways) which reduce PQ pool and is mediated by NDH leading to state II transition.

Conclusions/Significance: Anaerobic condition triggers non photochemical reduction of PQ pool mediated by NDH complex. The reduced PQ pool activates STN7 kinase leading to state II transition in A. thaliana.

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NADP reductase (FNR), leads to ΔpH and ATP production [22,23,24]. It has been reported that heat stress enhances the dark reduction of PQ pool also indicating the stimulation of cyclic electron transport around PSI [16,25]. This could be due to higher ATP demand under heat stress resulting in a higher NADPH/ATP ratio favouring non photochemical reduction of PQ pool which in turn provide ATP.

Apart from light, heat and nitrogen deficient conditions anaerobiosis was reported to reduce PQ pool of the photosynthetic electron transport chain [4,15,26,27]. The effect of anaerobiosis on reduction of PQ pool was well studied by OJIP transients [28]. Anaerobic condition results in oxygen-depletion inhibiting the terminal oxidase that generally keeps the PQ-pool in oxidized state ultimately resulting in the reduction of the PQ pool [29]. Similarly, transition to state II was also observed in A. thaliana when oxidative phosphorylation is inhibited by inhibitors of respiratory electron transport [30]. This phenomenon was reported to be due to a rapid drop in the ATP content, stimulation of glycolysis and an increase in the NADPH level, which in turn results in non photochemical reduction of the PQ pool [30,31].

The mechanism of state transitions under anaerobic condition is well documented in C. reinhardtii. However, the composition of the photosystems and its light harvesting components are different ranging from cyanobacteria to C. reinhardtii and higher plants. This is evident from cyanobacteria where the light harvesting complexes are phycobilisomes [32]. Phosphorylation of LHCII is well studied in C. reinhardtii and higher plants under low light conditions. However, the phosphorylation and migration of LHCII differs among these organisms. In higher plants only 15–20% of the LHCII is transferred to PSI whereas in C. reinhardtii ~80% of the antenna is migrated to PSI upon state II [33,34,35,36]. It has been studied that state I to state II transitions induces a switch from linear to cyclic electron transport. Such changes in electron transport have not been well studied in higher plants. Further, in C. reinhardtii, NDH-2 participates in chlororespiration whereas NDH-1 is involved in higher plants chlororespiration [37]. However, the detailed mechanism of PQ reduction and the involvement of NDH and cyclic electron transport under anaerobic condition are still unclear in higher plants.

The present study deals with the effect of anaerobiosis on non photochemical reduction of PQ pool and on the mechanism of state transitions in wt A. thaliana and mutants crr2-2 (defective in NdhB subunit of NDH complex) and pgr5 (defective in PSI CET). Post illumination studies were carried out to monitor changes in non photochemical reduction of PQ pool during anaerobic conditions when compared to aerobic conditions. We investigated the changes in electron transport, phosphorylation pattern of LHCII, and changes in absorption cross-section of PSI and PSII in wt and stn7. The importance of NDH in non photochemical

Figure 1. Effect of anaerobiosis on Chl a fluorescence transients in A. thaliana leaves. A. Raw Chl a fluorescence transients of dark anaerobic treated leaves for different time periods (0 to 20 min). B. Fv/Fm ratio for 0 and 20 min of anaerobic treatment. C. Normalized Chl a fluorescence transients of anaerobic treated leaves for different time periods (0 to 20 min). D. Normalized Chl a fluorescence transients of recovered leaves to aerobic condition from 20 min of anaerobic condition.
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Anaerobic treated leaves (Far red transition). E. Post illumination and stn7 Chl a that the Chl were harvested from 6–7 week old plants.

The measurements were carried out in N2-atmosphere. We note into the bag for 1–20 min; however, some outflow was allowed.

Anaerobic Treatment

Figure 2. Modulated Chl a fluorescence transients. A. Chl a fluorescence transient recorded during light to dark transition. B. Post illumination transients of Chl a fluorescence in aerobic treated leaves (dark transition) of wt and stn7. C. Post illumination transients of Chl a fluorescence in anaerobic treated leaves (dark transition) of wt and stn7. D. Post illumination transients of Chl a fluorescence in aerobic treated leaves (Far red transition). E. Post illumination transients of Chl a fluorescence in recovered leaves to aerobic condition from anaerobic condition (dark transition).

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reduction of the PQ pool leading to LHCII phosphorylation is also studied under dark anaerobic condition.

Materials and Methods

Growth Conditions of Arabidopsis thaliana

A. thaliana wild type (wt), stn7, cry2-2 and pgr5 mutant plants were grown in controlled environment chambers at 100–120 μmol m⁻² s⁻¹, with 8 h light/16 h dark periods. Leaves were harvested from 6–7 week old plants.

Aerobic condition within 20 min following a 20-min of N2-gas treatment. All the experiments were repeated thrice and obtained similar results.

Immunoblotting

For immunoblotting analysis, proteins separated by SDS-PAGE were transferred onto a polyvinylidene difluoride (PVDF) membrane (Bio Rad) using transblot apparatus (Bio Rad), according to manufacturer’s instructions. Blots were probed with anti-phosphothreonine polyclonal antibodies (1/2500 dilution) from New England Biolabs (Cell Signaling Technologies, UK) to detect protein phosphorylation at a threonine residue. Chl concentrations were determined by using Porra et al [38]. Equal Chl concentration (2 μg) was loaded on each lane.

77 K Fluorescence Emission Measurements

Isolated thylakoid membranes from different treatments were diluted to10 μg Chl ml⁻¹ and excited at 436 nm and emission spectra at 77 K were recorded in the range of 600–780 nm. Low temperature fluorescence emission spectra were measured at 77 K using a (Perkin Elmer, LS-55) fluorescence spectrophotometer. Emission spectra were recorded at a speed of 1 nm s⁻¹. Band width was 5 nm for both excitation and emission. Raw spectra were normalized at 685 nm for comparing fluorescence emission bands emanating from PSI.

The Fast OJIP Fluorescence Transient Measurements

Chl fluorescence fast induction curves were measured using Chl fluorimeter (PEA, plant efficiency analyzer, Hansatech, King’s Lynn, Norfolk, UK). Dark adapted leaves were excited by an array of three light-emitting diodes peaking at 630 nm at a photon flux density of 3000 μmol photons m⁻² s⁻¹. The fast fluorescence transients (OJIP) were measured [39] from aerobic and anaerobic treated leaves.

Post Illumination Transients for Monitoring Non Photochemical Reduction of PQ Pool

Chl fluorimeter (dual PAM100, Heinz Walz) was used to monitor Chl a fluorescence. Actinic light (AL) illumination was provided by an array of 635-nm LEDs illuminating the surface of the leaf, and applied 1 ms after turning on the 460 nm measuring beam. Post illumination measurements were carried out by applying AL of 35 μmol photons m⁻² s⁻¹ for 8 min and turned off later for 100 s [40]. The maximal fluorescence in the dark-adapted state (Fm) was determined by a 0.8-s saturating (4000 μmol photons m⁻² s⁻¹) light pulse applied after the onset of AL light. The rise in Chl fluorescence in dark is termed as F."
transient reflects the kinetics and heterogeneity involved in filling up of the redox carriers involved in electron transport from PSII to PSI, consequently, it can be used as a sensitive tool to investigate the photosynthetic apparatus in vivo under different environmental conditions [39,49].

Fig. 1A demonstrates O-J-I-P Chl a raw fluorescence transients of anaerobic treated wt leaves from 0 to 20 min. As the duration of anaerobic treatment increased, the rise in the fluorescence transient was significantly altered, increasing minimal fluorescence ($F_{20\text{m}}$) and decreasing fluorescence yield at P phase ($F_{p}$). At 0 min of anaerobic treatment (aerobic condition) there was lower $F_{20\text{m}}$ and showed $\sim$1, $\sim$3, and $\sim$30 ms for J, I, and P phase, respectively. At the end of anaerobic treatment (20 min), the Chl a fluorescence transient drastically varied when compared with 0 min of treatment (aerobic condition). The $F_{o}$ level increased with the time of exposure to anaerobic treatment while the $F_{p}$ level decreased, suggesting a significant decrease in the photochemical yield of $F_{r}/F_{p}$ (in this context) from 0.8 to 0.5 (Fig. 1B). This result was expected because of partial reduction of PQ pool. Also, there has been drastic increase in fluorescence at J phase ($F_{j}$) when the exposure time to anaerobic condition is increased. The rapid increase in J phase might be due to larger reduction of PSII acceptor side and the J step might be the starting point for the over reduction of the PQ pool. This increase in fluorescence yield at 2 ms is usually interpreted as an evidence for accumulation of the reduced QA, possibly due to decreased electron transport beyond QA$^{-}$ [39]. However, the increase in $F_{o}$ level by the moderate heat treatment was ascribed to back transfer of electron from the secondary Qb$^{\cdot}$ to the stable primary acceptor QA of PSII [50].

The $S_{m}$ normalized total complementary area above the O-J-I-P transient reflecting multiple turnover of QA reduction events and $N_{r}$, turn over number: number of QA reduction events between time 0 and $t_{fr}$ [42] were drastically reduced as the exposure time increased to anaerobic condition stating that number of multiple turnover QA reduction events were almost abolished during anaerobic condition (Data not shown). The above fluorescence parameters ($S_{m}$ and $N$) further indicate that the electron transport has been drastically decreased in anaerobic treated leaves. The area above the OJIP fluorescence transient is generally assumed to be a measure for the number of electrons that have to flow through the electron transport chain in order to reduce the redox carriers of PSII and PSI [46,51,52]. In the present study, the reduced area above the fluorescence transient indicates that the electron flow from PSII to PQ pool and PSI is decreased (Fig. 1A,C,D).

Fig. 1C,D are the O-J-I-P transients normalized at the O ($F_{o}$) and P ($F_{p}$) step in order to see the changes in the transients more clearly. The drastic increase in J phase and drop from J to P phase indicates that reoxidation of PQ was still monitored in anaerobic treated leaves for 20 min. In treatments like DCMU, there was no drop in J to I phase and attains maximal fluorescence at J phase and follows same yield of fluorescence at I and P phase [53]. When the leaves were relaxed to aerobic condition the reappearance of typical OJIP transient was monitored and this suggests that the changes in redox status of the PQ pool are reversible (Fig. 1D). Earlier reports suggested that under anaerobic conditions PTOX...
was found to be inactivated and the PQ pool was kept in highly reduced state [19,20,21]. In this study, chlorophyll fluorimeters are used to measure the redox state of PQ pool (by PEA) and NDH activity (by PAM, post illuminations studies).

Chl a Fluorescence Analysis for Non Photochemical Reduction of PQ Pool

Changes in redox state of PQ pool are further analyzed by post illumination transients (after illuminating with weak actinic light) during aerobic and anaerobic treatment (Fig. 2A–E). Fig. 2A describes Chl a fluorescence during light to dark transition. The apparent rise in \( F_o \) (Chl fluorescence increase) after a light to dark transition was used as a measure of non photochemical reduction of the PQ pool and referred as \( F_o' \) [54,55]. Rise in \( F_o' \) level over a period of 100 s after turning off the actinic light was not detectable during dark aerobic treatment indicating the absence of non photochemical reduction of PQ pool in wt and \( stn7 \) (Fig. 2B). Further, upon dark-anaerobic treatment, a rise in \( F_o' \) after switching off the actinic light indicated that non photochemical reduction of the PQ pool was operative in both wt and \( stn7 \) (Fig. 2C). This change in Chl fluorescence level is generally ascribed to the reduction of PQ pool via NDH activity [56,57] and this activity was not observed under aerobic condition (Fig. 2B,E). Earlier, it has been reported that increase in \( F_o' \) can be ascribed to increased electron transfer from stromal reductants to PQ pool and Cyt b6/f mediated by NDH [25]. In higher plants, plastidial NDH complex mediates chlororespiration and cyclic electron transport in thylakoid membranes [19,26,56].

When the actinic light was turned off and further illuminating with far-red background which preferentially excites PSI drives the oxidation of PQ pool. When anaerobic treated leaves were illuminated with far red light (electron flow from PSI is accelerated), there is no rise in \( F_o' \) indicating that PSI light oxidized the PQ pool (oxidation of PQ pool by PSI and re-reduction of PQ pool resumed back when switched off the far red light) (Fig. 2D). Anaerobic treated leaves when recovered back to aerobic conditions tend to resume their original state with the PQ pool under oxidized state where non photochemical reduction of the PQ pool was not observed (Fig. 2E). Taken together these observations, it can be suggested that the PQ pool was kept in oxidized state during dark- aerobic conditions.
while the PQ pool was under reduced state during dark-anaerobic conditions which is indicative of operation of alternative electron transport mechanisms resulting in non photochemical reduction of PQ pool. Earlier reports suggested that reduction of PQ pool was due to the inability of mitochondrial respiration to dispose metabolically-generated electrons under O₂ free environment leading to the accumulation of reducing equivalents throughout the cell [58].

Reduction of PQ Pool Under Anaerobic Condition Triggered State Transition

*stn7* mutant (the expression of the gene responsible for STN7 kinase was blocked) was used to study the role STN7 kinase in phosphorylation of LHCII under anaerobic condition [59]. Since the fluorescence transient rise in \( F_{v} \) (Fig. 1) and \( F_{v}/F_{m} \) (Fig. 2C) is indicative of the redox state of the PQ pool and the redox status of the PQ pool is linked to the LHCII phosphorylation [60], we further aimed to study the changes in LHCII phosphorylation by immunoblotting analysis. Immuno detection with anti phospho-threonine antibodies enabled us to determine changes in phosphorylation levels of PSII proteins especially LHCII. Wt leaves treated under dark aerobic conditions did not exhibit LHCII phosphorylation (Fig. 3A, lane 1). When leaves were exposed to anaerobic condition for 20 min, phosphorylation levels of LHCII were increased drastically (Fig. 3A, lane 2). Dark-anaerobiosis for 20 min would lead to reduction of PQ pool and subsequently activates STN7 kinase which phosphorylates LHCII. STN7 kinase responded to oscillations of the PQ pool redox state in a similar manner as in low light or state II adapted leaves [61]. However, in *stn7* mutant though non photochemical reduction of PQ pool was monitored, LHCII phosphorylation was not observed in both dark aerobic and anaerobic treated leaves (Fig. 3A lane 3 and 4). LHCII phosphorylation was also earlier reported during temperature treatment (in dark), due to non photochemical reduction of PQ pool by stromal reductants [15,16,25]. Similarly, Umate et al., (2008) have shown LHCII phosphorylation in anaerobic treated Tobacco leaves [62].

In order to show whether the increase in LHCII phosphorylation is contributing to increase in the PSI absorbance cross-section (phosphorylated LHCII is migrated from PSII to PSI), we further carried out 77 K fluorescence emission analysis. After excitation at 436 nm, PSI gives a significant fluorescence signal peaking at 726 nm, while PSII fluorescence peaks appear.
mostly at 685 and 695 nm. Fig. 3B shows, the 77 K fluorescence emission spectra of thylakoid membranes isolated from dark aerobic, dark anaerobic adapted wt leaves. For monitoring the changes in PSI (F726), the spectra were normalized at 685 nm which corresponds to PSII emission. In dark, there was no increase in the F726 showing that absorbance cross section of PSI was less due to reduced levels of phosphorylated LHCII (Fig. 3B). In anaerobic condition, the increase in PSI fluorescence emission was due to increased absorption cross-section by binding of LHCII migrated from PSII supercomplex (Fig. 3B). Fluorescence emission at F726 was found to be increased in LL adapted leaves as reported previously [59,63]. In \textit{stn7}, there is no significant difference between aerobic and anaerobic treated leaves (Fig. 3C) indicating that the phosphorylation of LHCII by STN7 kinase is operational under anaerobic condition in \textit{wt}. The above results indicate that under anaerobic condition the absorption cross-section of PSI was increased in \textit{wt}. These results are consistent with the previous reports on anaerobic treatment in \textit{C. reinhardtii} where similar kind of observations were made (LHCII is migrated to PSI under anaerobic conditions which leads to state II) [64].

Effect of Anaerobiosis on Fluorescence Induction Curves Under Actinic Light Illumination in Mutants Deficient in NDH and PGR5 Mediated Cyclic Electron Transport

The increase in NDH activity was mainly estimated by slow fluorescence rise after actinic light illumination. To investigate the role of chlororespiration and cyclic electron transport under
anaerobic conditions, chlororespiratory mutant crr2-2 (defective in processing of the mRNA of ndhB) and cyclic electron transport mutant pgr5 (deficient is defective in the non-NDH pathway of PSI CET) were used for the present study. These mutants were analyzed for Chl a fluorescence (Fig. 4) and phosphorylation studies (Fig. 5). Exposure of dark-aerobically treated leaves to actinic light induced initial sharp rise in Chl fluorescence, followed by a small rise in the transient which slowly quenched to a steady-state fluorescence after few min in wt and crr2-2 (in the presence of light wt could operate NDH dependent cyclic electron transport, whereas crr2-2 could not operate under anaerobic condition) (Fig. 4A,C). The initial quenching in fluorescence yield in pgr5 was found to be very slow when compared to wt and crr2-2 (Fig. 5B). Upon anaerobic treatment wt and crr2-2 mutant exhibited almost identical transients implying that mutation of NDH dependent cyclic electron transfer did not affect photochemistry under low actinic light.

In pgr5 mutant, steady state fluorescence (fluorescence quenching was drastically reduced) was observed after few sec of actinic light illumination and this could be due to PSI acceptor side limitation under anaerobic condition (Fig. 4F). The high fluorescence yield in wt and mutants during anaerobic condition could be attributed to the reduction of PQ pool due to inactive PTOX. The increased reduction state of QA, under anaerobic condition likely reflects a high electron pressure on the thylakoid redox system involving PTOX and NDH [19,56].

**Changes in LHCII Phosphorylation in Mutants crr2-2 and pgr5 Upon Anaerobic Treatment**

To monitor the changes in LHCII phosphorylation, wt and mutants were subjected to dark aerobic and dark anaerobic treatment. LHCII phosphorylation levels were found to be increased in wt during anaerobiosis confirming that non-photochemical reduction of the PQ pool is contributing to the LHCII phosphorylation and leading to a transition to state II. Similarly, when crr2-2 was subjected to identical conditions as that of wt, LHCII phosphorylation was not observed indicating that NDH is important in dark reduction of PQ pool under anaerobic condition (Fig. 5A). When pgr5 was subjected to anaerobiosis LHCII phosphorylation was observed. This reduction could be due to the NDH activity of the pgr5 mutant (Fig. 5A). The increased F726 in pgr5 is similar to that of wt (Fig. 3B) (state II transition) in anaerobically treated leaves, indicating that mutation in PGR5 mediated cyclic electron transport did not affect state II transition (Fig. 5B). However, in crr2-2 mutant abolition of increase in F726 was monitored showing that NDH is important in dark reduction of PQ pool driving to state transition (Fig. 5C).

To further confirm that NDH is responsible for the non-photochemical reduction of the PQ pool during anaerobic conditions, mutants deficient in chlororespiration and cyclic electron transport were analyzed by post illumination transients (Fig. 6). Post illumination transient analysis showed that the wt, crr2-2 and pgr5 mutants did not exhibit an increase in $F_{v}'$, indicating the absence of NDH activity during dark aerobic condition leading to an oxidized state of PQ pool (state I) (Fig. 6A–F). However, upon anaerobic treatment an increase in $F_{v}'$ was monitored for wt indicating enhanced NDH activity (Fig. 6B and Fig. 2B,C). Post illumination transients in crr2-2 mutant after dark anaerobiosis exhibited a significant decrease in the $F_{v}'$ indicating the importance of NDH in non-photochemical reduction of the PQ pool (Fig. 6D). Similarly, when pgr5 mutant was used for the study, the $F_{v}'$ rise was completely absent in anaerobic treatment, however state II transition was effectively monitored (Fig. 6F, 5B). This shows the peculiar behavior of pgr5 under dark anaerobic condition.

In pgr5, alternative mechanisms like NDH activity may contribute to the ATP deficiency to a certain extent, but under such abiotic stress conditions promoting higher ATP demand, the impairment of ATP supply might have resulted in significant inhibition of photosynthesis [65,66]. The contribution of the PGR5-mediated alternative pathway is thus crucial in ATP generation and draining of electrons from PSI under acceptor limiting conditions in O2 free air.

**Proposed Mechanism of State Transition Under Anaerobic Condition**

Based on our studies we propose that under anaerobic conditions non-photochemical reduction of PQ pool mediated by NDH leads to activation of STN7 kinase to phosphorylate LHCII. The phosphorylated LHCII undocks from PSII and migrates to PSI thereby increasing its absorption cross-section and thus leading to state II transition (Fig. 7). The contribution of the components CRR2-2 (NdhB) and PGR5 during anaerobic conditions were well documented in the present study where depletion of crr2-2 affected the non-photochemical reduction of PQ pool as well as phosphorylation of LHCII. Under anaerobic conditions reducing equivalents will accumulate in the cell leading to a non-photochemical reduction of the PQ pool mediated by NDH activity (Fig. 7). The breakdown of starch and glycolysis and other metabolic processes have been reported to be increased under dark anaerobic condition leading to accumulation of reducing equivalents (NADPH) and further resulted in non-photochemical reduction of PQ pool by NDH activity [67].

**Conclusion**

The above findings indicate that anaerobiosis in A. thaliana would lead to an electron flow which would keep the PQ pool in reduced state. Results with OJIP transients have shown that the drastic increase in J phase under anaerobic condition was due to reduced state of PQ pool. Post illumination transients indicated that the non-photochemical reduction of the PQ pool is contributing to the mechanism of state transition where a transition to state II is observed during anaerobiosis in wt and stn7. The changes after a transition from dark aerobic to dark anaerobic condition is evident from the LHCII phosphorylation and increase in the PSI absorption cross section. The PQ pool reduction mediated by NDH under anaerobic conditions is confirmed by experiments with crr2-2 mutant. However, the contribution of the two major pathways (antimycin sensitive as well as antimycin insensitive pathways of cyclic electron transport) which are also involved in the non-photochemical reduction of the PQ pool mediated by NDH activity can be ignored under dark anaerobic condition. Though the present study focused on the importance of state transition under anaerobic conditions, the components which are associated with the reduction of PQ pool in presence of light anaerobic condition involving cyclic electron transport are not clear and needs further study.

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Author Contributions
Conceived and designed the experiments: RS SN. Performed the experiments: SN SK MT. Analyzed the data: RS SN SK. Contributed reagents/materials/analysis tools: RS. Wrote the paper: SN RS.

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