Towards elucidation of dynamic structural changes of plant thylakoid architecture

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Long-term acclimation of shade versus sun plants modulates the composition, function and structural organization of the architecture of the thylakoid membrane network. Significantly, these changes in the macroscopic structural organization of shade and sun plant chloroplasts during long-term acclimation are also mimicked following rapid transitions in irradiance: reversible ultrastructural changes in the entire thylakoid membrane network increase the number of grana per chloroplast, but decrease the number of stacked thylakoids per granum in seconds to minutes in leaves. It is proposed that these dynamic changes depend on reversible macro-reorganization of some light-harvesting complex IIb and photosystem II supracomplexes within the plant thylakoid network owing to differential phosphorylation cycles and other biochemical changes known to ensure flexibility in photosynthetic function in vivo. Some lingering grana enigmas remain: elucidation of the mechanisms involved in the dynamic architecture of the thylakoid membrane network under fluctuating irradiance and its implications for function merit extensive further studies.

Keywords: dynamic structure; fluctuating irradiance; grana; lateral heterogeneity; plant thylakoid architecture; thylakoid protein complexes

1. INTRODUCTION

The static view of the structural organization of the unique plant thylakoid membrane network inferred from transmission electron micrographs, being but snapshots in time, cannot capture their highly organized and dynamically regulated ultrastructure. To survive and thrive under ever-fluctuating light, plants have evolved long-term acclimation strategies to optimize photosynthetic efficiency and resource utilization [1–3] that are intertwined with vital short-term structural and functional flexibility under fluctuating irradiance [4–7]. Granal stacks in higher plants have been selected during evolution for the integrated, multifaceted advantages and optimization of photosynthesis they confer in diverse and ever-fluctuating light environments [8–11]. In this article, we emphasize that the elaborate dynamic structural changes of more but shorter grana versus fewer but taller grana and vice versa observed in hours, days to seasons in chloroplasts of shade and sun plants are mimicked in seconds to minutes in leaves in response to increasing fluctuating light. Although there have been few examples of snapshots that capture dynamics from experiments with leaves under controlled fluctuating conditions, we suggest that they are caused by the reversible phosphorylation of thylakoid proteins and the various changes associated with non-photochemical quenching (NPQ), the D1 protein repair cycle and the photoprotection of non-functional photosystem II (PSII) under very high irradiance. This highly dynamic regulation of ultrastructure and function of plant thylakoids is intriguing yet still puzzling, especially in the remarkably dynamic three-dimensional architecture of plant thylakoids in vivo. Hence, some lingering grana enigmas concerning the dynamic structural changes in the architecture of the thylakoid network in vivo are also discussed.

2. SUPRAMOLECULAR ORGANIZATION AND THYLAKOID ARCHITECTURE UNDER ACCLIMATION AND FLUCTUATING IRRADIANCE

The highly dynamic structural organization of the continuous thylakoid membrane network of higher plant chloroplasts is intriguing, especially in its elaborate three-dimensional architecture. The continuous thylakoid network that encloses one internal aqueous lumenal space is structurally differentiated into two distinct morphological regions: cylindrical, tightly appressed granal thylakoids (grana stacks) are interconnected by single stromal thylakoids whose outer
surfaces face the stroma. This elaborate structural membrane architecture is accompanied by compositional and functional differentiation with respect to the location of the thylakoid pigment–protein complexes termed lateral heterogeneity. PSIII/light-harvesting complex II (LHCCI) supercomplexes and extra LHCCI are mainly segregated in dynamically regulated grana, whereas photosystem I (PSI) and ATP synthase are confined to stromal thylakoids and end granal membranes, while cytochrome (cyt) b$_{6}$/f complexes are reversibly located between stacked and unstacked thylakoid regions ([12–15] and references therein).

Long-term acclimation of plants grown in nature or controlled conditions related to light quantity and quality is well understood [1,2]. With long-term acclimation, the ratio of granal to stromal membrane domains in higher plant chloroplasts is highly variable: the chloroplasts of sun and high-light grown plants have more grana with fewer (5–16) stacked thylakoids per granum, while shade and low-light acclimated plants have fewer grana per chloroplast with more stacked thylakoids, with Alocasia macrorhiza having giant grana with up to 160 stacked thylakoids [3]. Relative to PSI cyanophyl shade and light plants (lower chlorophyll (Chl) a/Chl b ratios) have fewer core PSI II complexes served by larger light-harvesting antennae compared with sun and high-light plants (higher Chl a/Chl b ratios) with more core PSI II complexes served by smaller light-harvesting antennae; these striking adjustments of the PSII/PSI reaction centre stoichiometry modulate grana stacking [2]. Shade and low-light plants have more chlorophyll for maximal light capture at the expense of electron transport, photophosphorylation and carbon fixation, resulting in lower maximal rates of photosynthesis which saturate at low irradiance. Conversely, sun and high-light plants, being limited in electron transport rather than in light energy capture and conversion, have greater amounts of cyt b$_{6}$/f complexes, ATP synthase and mobile plastoquinone, plastocyanin and ferredoxin, to support greater maximal rates of photosynthesis which saturate at higher irradiance [2].

Many of these modulations of composition are fully reversible [16], and in turn are accompanied by changes in photosynthetic function and thylakoid membrane network organization [2]. These long-term modulations of composition are so beautifully orchestrated that even Chl a/Chl b ratios can serve as simple indices of light intensity acclimation in plants. The Chl a/Chl b ratios of pea leaves are linearly correlated with the content and activity of cyt b$_{6}$/f complex, ATP synthase and Rubisco, but inversely related to LHCCI and LHCI content [1,2], and the extent of thylakoid stacking [17].

Acclimation to high irradiance is also linked to the phenomenon of photo-inhibition. Although a detailed discussion of photo-inhibition is beyond the scope of this article, it should be emphasized that from a functional viewpoint, exposure to excess irradiance brings about a stable, long-term regulation of PSII that may be ‘locked in under sustained high light’, particularly in shade and low-light plants [18,19]. Thus, photo-inhibition should be regarded not only as a damaging event, but also as a photoprotective strategy of ecological relevance. Here too, grana may be of critical importance, perhaps preventing premature degradation of D1 and D2 proteins by harbouring non-functional PSIIs deep in appressed thylakoid domains under high irradiance [17].

Besides long-term acclimation (hours, days, weeks or seasons), plants also need reversible, dynamic regulation of photosynthesis from seconds to minutes to allow structure to intertwine with function under fluctuating light quality and quantity. Plants undergo rapid, reversible phosphorylation cycles involving some LHCCI and PSI II core proteins depending on the interplay between the two main kinases, STN7 and STN8, and their respective phosphatases according to irradiance ([7] and references therein). Low irradiance mainly stimulates phosphorylation of LHCCI proteins and, to a lesser extent, some PSI core proteins that cause the STN7 kinase-mediated phosphorylated LHCCI to laterally migrate from stacked grana to nearby stroma thylakoids, thereby decreasing thylakoid stacking [20,21]. Increasing irradiance induces mainly STN8 kinase-dependent phosphorylation of some PSI core proteins and also the STN7-mediated phosphorylation of the minor antenna complex CP29 [22,23]. Also in high light, the rapid build-up of ΔpH stimulates the activation of violaxanthin de-epoxidase (resulting in accumulation of zeaxanthin) and the protonation of various PSI II antenna proteins [4–6]. Together, these cause NPQ, the transformation of LHCCI/PSII into a state that dissipates excess light energy. NPQ is accompanied by changes in the structure and organization of the thylakoid membrane [24–26].

3. DYNAMIC STRUCTURAL CHANGES IN GRANA IN LEAVES UNDER FLUCTUATING IRRADIANCE: A GRANA CONUNDRUM

(a) In darkness, antisense Arabidopsis leaves lacking native LHCCI trimer have a different thylakoid architecture compared with wild-type leaves

Antisense plants lacking one or more of the six specific PSII antenna Chl a/b proteins usually maintain PSII function but never possess the full orchestral suite of dynamic thylakoid molecular strategies observed in vivo [5]. Arabidopsis antisense plants, asLhcb2, lacking the main LHCCIb trimers involved in grana stacking, increase the content of Lhcb5, which assembles as compensatory Lhcb5 trimers in vivo [27]. These novel trimers function as an alternative PSII antenna enabling somewhat normal macro-organization of PSII supercomplexes to be attained, although the extra LHCCIb-only domains of wild-type chloroplasts are absent. Nevertheless, the asLhcb2 chloroplasts retain well-preserved grana stacking, apparently similar to wild-type chloroplasts [28], although isolated antisense thylakoids are extremely unstable [27,29].

We investigated the architecture of the thylakoid membrane network of chloroplasts with wild-type and asLhcb2 Arabidopsis leaves that were dark-adapted for 15 h, particularly the number of grana per chloroplast and the number of appressed thylakoids per granum. The size and number of grana in two-dimensional thin chloroplast sections from these leaves were assessed by image analysis, according to the method of Rozak et al. [28]. The asLhcb2 plants had higher numbers of non-appressed grana per chloroplast and also much smaller grana, with a different internal structure as well as a more dynamic arrangement of the thylakoid membrane. The whole chloroplast was more filamentous and the grana were much smaller than in wild-type leaves [28]. These results are consistent with the observation that the asLhcb2 leaves show lower rates of photosynthesis and carbon fixation than wild-type leaves [28].

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Table 1. Ultrastructural analysis of chloroplasts of dark-adapted leaves from wild-type and asLhcb2 Arabidopsis, and after 1 h growth light (150 μmol photons m⁻² s⁻¹). Fifteen chloroplasts were used for the measurement of height and width in wild-type and asLhcb2, respectively. Values are given as means ± s.e.

<table>
<thead>
<tr>
<th></th>
<th>wild-type m</th>
<th>asLhcb2 n</th>
<th>100 × (n – m)/m</th>
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</thead>
<tbody>
<tr>
<td>grana per chloroplast in dark</td>
<td>37 ± 8</td>
<td>63 ± 12</td>
<td>70%</td>
</tr>
<tr>
<td>thylakoids per granum in darka</td>
<td>5.5 ± 0.4</td>
<td>3.8 ± 0.2</td>
<td>-31%</td>
</tr>
<tr>
<td>diameter (μm) of grana in dark</td>
<td>0.47 ± 0.04</td>
<td>0.48 ± 0.03</td>
<td>2%</td>
</tr>
<tr>
<td>chloroplast section profile area (μm²) in dark</td>
<td>15.4 ± 1.5</td>
<td>12.2 ± 1.1</td>
<td>-21%</td>
</tr>
<tr>
<td>grana per chloroplast in 1 h growth light</td>
<td>67</td>
<td>53</td>
<td></td>
</tr>
<tr>
<td>thylakoids per granum in 1 h growth lightb</td>
<td>3.9</td>
<td>3.9</td>
<td></td>
</tr>
</tbody>
</table>

The number of thylakoids per granum was calculated by dividing the height by the average thickness of one thylakoid in grana. Granal height, width and cross-sectional area were measured with ANALYSIS program (Olympus Soft Imaging System).

Values are mean granum sizes calculated from a frequency distribution of the number of thylakoids per granum. Adapted from E-H Kim, PhD thesis, Australian National University, 2006.

The dark-adapted antisense chloroplasts had an increase of 70% per cent in the number of grana per chloroplast: 63 in antisense chloroplasts compared with 37 in the wild-type (table 1). Concomitantly, in dark-adapted asLhcb2 chloroplasts, the number of appressed thylakoids per granum was diminished by 31% per cent compared with wild-type chloroplasts, resulting in somewhat comparable extents of grana stacking in antisense and wild-type chloroplasts (table 1). Surprisingly, asLhcb2 leaves have smaller chloroplasts (table 1), with each chloroplast possessing more grana with fewer stacked thylakoids per granum than found in wild-type leaves. This unexpected thylakoid profile was not observed visually by Andersson et al. [28] or us until the image analysis was undertaken.

Thus, the overall grana membrane architecture of Arabidopsis leaves was profoundly changed in the dark-adapted antisense chloroplasts compared with wild-type chloroplasts. Notably, a gain in grana per chloroplast with a loss of stacked thylakoids per granum, with no difference in the diameter of grana in the dark, means a significant increase in the total granal end membrane area in asLhcb2 chloroplasts compared with that of wild-type chloroplasts.

(b) Dynamic structural changes in Arabidopsis in wild-type and asLhcb2 leaves in response to irradiance

When dark-adapted wild-type Arabidopsis leaves were illuminated under growth light for 1 h to reach steady-state photosynthesis, the number of grana per chloroplast section increased from 37 to 67, while the number of stacked thylakoids per granum decreased from 5.5 to 3.9 (table 1). By contrast, antisense leaves exhibited little or no increase in the number of grana per chloroplast or in the number of stacked thylakoids per granum (table 1). The distribution of granal size (thylakoids per granum) between dark- and light-adapted leaves is shown in figure 1a (for wild-type) and figure 1b (for asLhcb5 leaves). The change in the distribution of granal size in asLhcb2 leaves was hardly shifted between dark and 1 h growth light (figure 1d), in marked contrast to the wild-type (figure 1c). It is remarkable that the grana characteristics of light-adapted plants are now virtually the same in the wild-type and antisense plants. Thus, the antisense plants not only lose the ability to show dynamic changes in grana structure, but also even in the dark-adapted state they resemble the light-adapted state of the wild-type.

The explanation of these differences between wild-type and antisense plants resides in the differences in Lhcb and Lhca protein composition. Although Lhcb5-containing S and M trimers accumulate in response to the absence of Lhcb1 and Lhcb2, the additional trimers are absent [27]. Could the loss of these trimers in the antisense plants lead to the constitutive adaption of the light-adapted state found in the wild-type? In the wild-type, the change in grana structure from dark to light almost certainly occurs because of LHCCIb phosphorylation. Therefore, because Lhcb5 lacks a phosphorylation site, it would be predicted that the antisense plants would show no light-induced change in thylakoid stacking. Indeed, the state 1–2 transitions, the expression of changes in antenna organization caused by the reversible LHCCIb phosphorylation, are absent in antisense plants [28]; the plants were found to be locked in state 2, which in wild-type plants is the phosphorylated state. Thus, the depletion of LHCCI from the grana, either genetically in the antisense plants or by phosphorylation in the wild-type, underlies the increase in frequency and reduction in size of grana stacks. In addition, the antisense thylakoids also possess an increased content of Lhca1–4 proteins per PSI reaction centre, contributing to a larger PSI antenna, thereby compensating for the absence of phosphorylated LHCCI [29]. However, the antisense plants are extremely vulnerable to both rapid ever-changing environmental conditions and very low growth irradiance [31], demonstrating the importance of phosphorylation of LHCCI under fluctuating low irradiance and consistent with the requirement for STN7 kinase for optimal growth [7].

(c) Rapid dynamic structural and functional changes in the architecture of the entire thylakoid network in spinach leaves following transitions in irradiance

The remodelling of the grana in Arabidopsis wild-type leaves upon transition from darkness to light is very similar to that observed by Rozak et al. [30], who compared spinach plants before and after a transition from growth light (300 μmol photons m⁻² s⁻¹) with low
light or shade (10 μmol photons m⁻² s⁻¹) by neutral density shade or self-shading by another leaf). Changes in the overall grana number and size in the attached leaves were quantified by measuring the two-dimensional areas of grana in cross section as seen in transmission electron microscopy thin sections [30]. There is a striking decrease in grana number per chloroplast, but a gain of larger grana (table 2). In particular, transfer to very low irradiance from growth light causes a substantial decrease in the number of grana per chloroplast, but a concomitant increase in the number of appressed thylakoids per grana stack, with a net 9 per cent increase in thylakoid appression (table 2). Consequently, there is a significant decrease in the number of end granal membranes and possibly also the interconnecting stromal thylakoid area. As

Table 2. Comparative measurements of the two-dimensional ultrastructural characteristics of chloroplasts from spinach leaves that were light-adapted (2 h at 300 μmol photons⁻² s⁻¹) or switched from the light-adapted state to low irradiance/self-shade or higher irradiances. Values are given as means ± s.e.

<table>
<thead>
<tr>
<th>light treatment</th>
<th>grana per chloroplast profile</th>
<th>10³ × grnum size (μm²)</th>
<th>calculated thylakoid appression (μm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>light-adapted (2 h at 300)a</td>
<td>44.5 ± 2.1</td>
<td>36 ± 1</td>
<td>23.4</td>
</tr>
<tr>
<td>10 min of low irradiance (10)</td>
<td>30.3 ± 1.6</td>
<td>49 ± 2</td>
<td>25.6 (+9%)b</td>
</tr>
<tr>
<td>10 min self-shade (approx. 10)</td>
<td>32.6 ± 1.4</td>
<td>49 ± 2</td>
<td>27.5 (+17%)</td>
</tr>
<tr>
<td>20 min self-shade (approx. 10)</td>
<td>30.3 ± 1.6</td>
<td>46 ± 2</td>
<td>24.0 (+6%)</td>
</tr>
<tr>
<td>10 min of 800 μmol m⁻² s⁻¹</td>
<td>49.2 ± 2.8</td>
<td>33 ± 1</td>
<td>23.0 (−2%)</td>
</tr>
<tr>
<td>10 min of 1500 μmol m⁻² s⁻¹</td>
<td>46.0 ± 2.0</td>
<td>51 ± 2</td>
<td>42.0 (+79%)</td>
</tr>
</tbody>
</table>

aNumbers in brackets in the first column indicate irradiance in μmol photons m⁻² s⁻¹.
bValues in parentheses in the last column represent the percentage change from the light-adapted value. From Rozak et al. [30] with permission.
above, these changes are interpreted in terms of de-phosphorylation of LHCII upon transfer to shade, as evidenced by the alteration of the 77 K fluorescence emission spectra [30].

Rozak et al. [30] extended their study to observing plants 10 min after transfer to a moderate irradiance (800 μmol photons m⁻² s⁻¹) and also to a very high irradiance (1500 μmol photons m⁻² s⁻¹). A shift to moderate irradiance induces a small (statistically not significant) increase in grana per chloroplast and a smaller granal size (table 2), slightly enhancing the character of the growth light state (in comparison with the shade state). In contrast, the shift from growth irradiance to very high irradiance greatly increases granal size with no change in granal frequency. This results in a vast increase in thylakoid appression by 79 per cent relative to that found in the growth irradiance (table 2).

This rapid response to exposure to very high light is intriguing, and deserves further study. Its molecular basis cannot be deduced at present, although we suggest that it is caused by one or more of the biochemical changes that have been recorded upon longer exposure to high irradiance. At high irradiance, the STN7 kinase-mediated phosphorylation of LHCIIb trimers is inhibited [32] and dephosphorylated LHCII would return to the appressed membranes. At the same time, the STN8 kinase induces particularly phosphorylation of certain PSII core proteins, D1, D2, CP43, the PsbH proteins of PSII and the Ca²⁺-sensing receptor protein [7,23]. As the rate of damage to PSII increases, there is lateral movement of non-functional PSII monomers to nearby stroma-exposed regions for D1 protein repair [33]. A further event occurring in very high light is the phosphorylation of the minor antenna complex CP29 by the STN7 kinase, an event thought to de-stabilize the LHCII/PSII supercomplex [23]. These changes in the balance of phosphoproteins under very high irradiance may well cause the tremendous increase in membrane appression and the accompanying increase in grana size (table 2). However, other features of the thylakoid also change in high light, principally the formation of NPQ. Under the influence of the increased ΔpH, the organization of the PSII/LHCII supercomplexes within stacked granal thylakoids is remodelled; this involves the structural re-organization of some LHCII, which dissociates from PSII supercomplexes, and then is aggregated [5,6,24–26].

The interplay between these responses to high light and how they might influence the granal structure is poorly understood. The return of dephosphorylated LHCIIb proteins to stacked granal domains may help to protect non-functional PSII from further photodamage (via enhancement of NPQ), and elicit some as yet unidentified PSII/LHCII remodelling within the stacked grana domain that is reminiscent of the structural organization found in very low irradiance or darkness. This may be aided by the dissociation of some PSII supercomplexes. Concomitantly, the dominance of phosphorylated PSII core proteins may widen the stromal gap between appressed grana thylakoids to allow photoinactivated PSII supercomplexes in the grana to generate dimeric and then monomeric non-functional PSII units that laterally migrate to nearby stroma-exposed thylakoid domains for D1 protein cycle repair [21,22,33].

(d) Integrated responses of thylakoid membranes to fluctuating irradiance

Thus, the number of grana per chloroplast and the number of thylakoids per granum can be reversibly altered within the whole thylakoid membrane network over the entire range of fluctuating irradiances (figure 2). That is, grana provide integrated and multifaceted functional advantages by facilitating mechanisms in a matter of seconds to minutes to cope with rapid fluctuating irradiances that fine-tune the dynamic flexibility of structure with function of the plant thylakoid network in vivo [1,4–8,14,15,22–26,30,34,35]. This grand design depends in part on the interplay of the two major kinases that are influenced by light intensity and quality, the STN7-dependent phosphorylation of some LHCII trimers and CP29 versus STN8-dependent phosphorylation of some PSII core proteins, including CP43. These alterations are governed by intricate changes in phosphorylation profiles, according to the redox state of the PQ pool coupled to the redox state of cyt b₆f, and stromal redox components beyond PSI [7]. In addition, the ΔpH induces major changes in thylakoid structure that result in NPQ decreasing membrane thickness and inducing LHCIIb aggregation. The extent of each of these events depends upon the initial irradiance, the magnitude of the transition in irradiance and the resulting degree of the rapid perturbation of the redox and energy states of the chloroplast. It should also be remembered that what is low, high or very high irradiance for such dynamic structural changes is different for diverse plant species, and even for the same plant species that have been acclimated mainly to low or high growth irradiance. In all cases, the thylakoid membrane network responds to rapidly maximize the efficiency of photosynthesis in limiting light, while avoiding photodamage in excess light.

With a dynamic shift from darkness to growth irradiance, fewer grana per chloroplast with more appressed thylakoids per granal stack in dark-adapted attached leaves rapidly reorganize to provide more grana with fewer appressed thylakoids per granal stack to ensure efficient light-harvesting and increased abundance of end granal membranes. Conversely, with a shift from growth irradiance to low light or self-shade, there were fewer grana per chloroplast, but much larger grana. Hence this important structural strategy helps all plants with very different compositions due to acclimation in various light environments to have the same maximum constant quantum yields in limiting light [36,37]. It is of particular relevance to note that these rapid short-term, reversible transitions of ultrastructure mimic the long-term, acclimation ultrastructural profiles of shade and low-light plant versus sun and high-light plant chloroplasts. Significantly, alteration in the amount and organization of LHCII in the grana is one of the common biochemical features of both these types of change. Similarly, STN7 kinase is so far the only enzyme known that is common to short-term transitions and long-term acclimation [38].
Clearly, the dynamic macroscopic structural changes of the thylakoid network under fluctuating irradiance need to be further explored. Important questions demand a better understanding of the regulation of flexible three-dimensional molecular architecture of the thylakoid membrane network in vivo, including the following:

— How are more grana per chloroplast with fewer appressed thylakoids per granum converted so rapidly to fewer grana per chloroplast with more appressed thylakoids and vice versa? How does the size of the grana increase so rapidly upon transfer from growth light to very high light? How do the transitions between these very different low light and high light responses occur? These are mind-boggling questions and some insights into the complexity involved have come from structural analysis of chloroplasts in the phosphorylated and unphosphorylated states [39]. Atomic force microscopy, scanning and transmission electron microscopy and confocal imaging reveal marked structural reorganization of the membranes at the interface between the stacked grana and unstacked stromal thylakoids. The reorganization of the membrane architecture is suggested to involve both fission and fusion events similar to those observed in the membranes of mitochondria, Golgi apparatus and endoplasmic reticulum [39].

— How do these rapid transitions interface with the acclimation changes that become embedded when the transition in irradiance is sustained?

— Do the chloroplasts of shade-acclimated species differ from those of sun plants in their dynamic ultrastructural responses to fluctuating irradiance? Are the amounts of ‘extra’ Lhcb2 trimers not attached to the PSII/LHCII supercomplexes increased in shade plants?

— In the light, the Mg²⁺ efflux from the lumen and the cytosol to the stroma is counter-balanced by the Ca²⁺/H⁺ antiporter which transports Ca²⁺ from the stroma to the granal lumen in exchange for a proton efflux [40]. How does this ionic redistribution elicit varying widths of grana lumen and maybe stroma lumen in leaves in darkness or diverse fluctuating irradiances?

— What happens in the LHCII/PSII remodelling in appressed grana thylakoids as dephosphorylated LHCII returns from nearby stroma thylakoid domains upon transition to increasingly high irradiance? Do they reorganization within stacked granal thylakoids enhance photoprotection, particularly of the remaining non-functional PSII centres? Antagonism between LHCIb phosphorylation and ΔpH-dependent NPQ has been observed, suggesting that dephosphorylation is a prerequisite for maximum quenching [41,42].

‘Given the complexity and pleomorphic nature of the thylakoid lamellar system, deducing its three-dimensional structure from two-dimensional data is precarious’ [15, p. 159]. Further three-dimensional structural studies with leaves under dynamic transient changes in diverse, controlled light conditions are needed.
4. TOWARDS UNDERSTANDING DYNAMIC STRUCTURAL CHANGES IN THE MACROSCOPIC THYLAKOID STRUCTURE IN VIVO: LINGERING GRANA ENIGMAS

The exquisite complexity of the three-dimensional architecture of the static thylakoid membrane network is being explored by electron microscope topography [15,43–46]. Recent three-dimensional cryo-tomography work [43–46] confirms the helical model of Paolillo [47] by showing that the stromal thylakoids indeed wind around granal stacks in the form of multiple right-handed helices at an angle of 20–25° around granal stacks in the form of multiple right-handed helices [48]. Through bridges located at the bifurcation points at the marginal rim of the grana [15,39,48], the granal lumen appears to be narrower (despite its high concentration of PSII oxygen-evolving complex proteins) than the stromal lumen. Kirchhoff et al. [50] examined vitreous sections of unfixed leaf samples of dark- and light-adapted Arabidopsis leaves by cryo transmission electron microscopy: they demonstrated that the granal thylakoid lumen dramatically expands by 96 per cent from 4.7 nm in the dark to 9.2 nm in the light in vivo. This light-induced expansion of the granal lumen greatly alleviates the restrictions imposed on protein diffusion by increasing protein mobility, especially plastocyanin mobility, in the light; it also aids in the manifold processes associated with the D1 protein repair cycle [50]. This observed expansion of the granal lumen in the light contradicts earlier suggestions following observations on isolated thylakoids [11,51] that the lumen may contract in the light. Conceivably, lumenal contraction or expansion in the light depends on the duration and intensity of illumination. Initially, as Mg$^{2+}$ exits the lumen driven by a proton influx, the lumen volume could contract to simultaneously satisfy the conditions of acid–base equilibrium, Donnan equilibrium, osmotic equilibrium and electron neutrality [52]. On the other hand, with prolonged illumination in the presence of Ca$^{2+}$, the Ca$^{2+}$/H$^+$ antiporter [40] could accumulate a substantial concentration of Ca$^{2+}$ accompanied by Cl$^-$ in the lumen; such an accumulation of the two ionic species will be accompanied by an influx of water, thereby expanding the lumenal volume.

(b) Granal margins

Recent three-dimensional tomographic data of vitreous sections of intact chloroplasts and plunge-frozen suspensions of isolated thylakoids demonstrated that monomeric ATP synthase is confined to the minimally curved, almost flat stromal regions of end grana membranes and stroma thylakoids [46]. Further three-dimensional tomographic analysis of an isolated granal stack revealed that its granal margins are only 3–4 nm wide [49]. The results from these studies are consistent with the early hypothesis of Murphy [53] that the granal margins are protein-free. Clearly, the granal margins are both too narrow and much too curved to contain membrane-spanning protein complexes. This means that the structural assignment of one of the submembrane fractions isolated by Yeda Press treatment followed by aqueous polymer two-phase partition [54], the so-called granal margins, needs to be reconsidered. Although the composition and functional characterization of ‘granal margin’ fractions have been well demonstrated in many biochemical studies over five decades, their structural assignment as ‘granal margins’ per se is no longer valid.

(c) Stromal thylakoids

Stromal thylakoids are defined as thylakoids whose outer membrane surface is exposed to the stroma, but recent advances in three-dimensional structure suggest that there are instead different regions of stroma-exposed membranes throughout the continuous membrane network: those that are directly linked to granal discs, the junctional connections or frets and the end granal membranes at the top and bottom of each granal stack, which differ from the large sheets of intervening stromal thylakoids that are not so directly linked to granal stacks.

(d) Junctional slits

Recently, the neglected junctional slits that directly link stacked granal thylakoids to stromal lamellae at the granal margins (sometimes glimpsed in two-dimensional electron micrographs as staggered lamellar membrane protrusions from granal thylakoids) are being revealed by three-dimensional cryo-tomography [43,44]. Some stromal thylakoids merge with successive granal discs with approximately 35 nm junctional slits, while others...
are much wider (approx. 400 nm) so that one stromal thylakoid forms a planar sheet with only one nearby granal disc [43,44]. Are these junctional slits actually the stroma-exposed fraction that has previously been termed ‘granal margins’ [54] and thus the submembrane domain where much of the phosphorylation/deshphosphorylation-dependent regulation of excitation energy distribution between PSII and PSI as well as the D1 protein repair cycle occurs?

(c) End granal membranes

The flat stromal-exposed end granal thylakoid sacs are unique; only the outer membrane is exposed to the stroma, while the inner, opposing membrane surface needs to be appressed to the next adjacent grana thylakoid, resulting in different membrane compositions in the opposing membranes facing each other across the lumen of the end granum thylakoid sac. This is an unusual situation. The outer stromal-facing end granal membranes possess monomeric ATP synthase, PSI, cyt b/f and PSIIβ, while the inner opposing membrane facing the lumen contains the oxygen-evolving complexes of the PSII/LHCII arrays.

The transition from darkness to increasing irradiance leads to an increase in end granal membranes seen in the few dynamic structural changes reported here. What functional significance could an increase in end granal membranes have in the rapid restructuring of the entire thylakoid membrane architecture? The more abundant end granal membranes can accommodate more ATP synthase, which would be strategically placed to optimize the proton circuit that drives ATP synthesis. Protons only need to diffuse a short distance to the ATP synthase located in the end granal membranes, when compared with diffusion to an ATP synthase in the distant stromal lamellae. Therefore, an increased abundance of end granal membranes may speed up or increase the efficiency of the proton circuit and photophosphorylation. As for the regulation of linear versus cyclic photophosphorylation under fluctuating irradiance, further experimentation is still needed, not least an accurate method for quantifying cyclic electron flow.

5. CONCLUDING REMARKS

Although it is not fully understood how the hierarchical assembly of PSII/LHCII supercomplexes in the dark and very low irradiance compares to the hierarchical disassembly of PSII/LHCII supercomplexes under high irradiances, it seems that although functional and structural stability will be enhanced by fewer, but taller grana per chloroplast in the shade and low light, they nevertheless have the capacity for surprisingly highly dynamic structural flexibility in response to transient bursts of high light. This remarkable robustness of plant LHCII/PSII depends on the complementary logic inherent in this marvellous dynamic molecular machine, which is exquisitely regulated both under rapid fluctuating irradiance and long-term acclimation. The structural differentiation of plant thylakoids into stacked granal and unstacked stromal domains is an example of the ‘specialized compartmentation’ that has occurred during evolution as a strategy to regulate increasingly complicated pathways and achieve both dynamic and long-term control over cellular functions and responses [1]. Thus, grana formation in higher plant chloroplasts fine-tunes photosynthesis by eliciting many dynamic strategies, including the ability to ensure that all plants have constant, high quantum yields at limiting light and balance photochemical utilization with photoprotection by NPQ to regulate maximal linear electron transport, to perform the ingenious D1 protein repair cycle at higher irradiance and to protect non-functional PSIIs ‘parked’ in the stacked granal domains at very high irradiance.

Epilogue

‘We shall not cease from exploration
And the end of all our exploring
Will be to arrive where we started
And know the place for the first time’

(T. S. Elliot, The Waste Land 1922)

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REFERENCES

7 Tikkanen, M. & Aró, E.-M. 2012 Thylakoid protein phosphorylation in dynamic regulation of photosystem...


