Chloroplast evolution, structure and functions
Poul Erik Jensen¹ and Dario Leister¹,²*  

Addresses: ¹Copenhagen Plant Science Center (CPSC), Department of Plant and Environmental Sciences, University of Copenhagen, Thorvaldensvej 40, DK-1871 Frederiksberg C, Denmark; ²Plant Molecular Biology, Department of Biology I, Ludwig-Maximilians-University Munich, Großhaderner Str. 2, D-82152 Planegg-Martinsried, Germany  
* Corresponding author: Dario Leister (leister@lmu.de)

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Abstract
In this review, we consider a selection of recent advances in chloroplast biology. These include new findings concerning chloroplast evolution, such as the identification of Chlamydiae as a third partner in primary endosymbiosis, a second instance of primary endosymbiosis represented by the chromatophores found in amoebae of the genus Paulinella, and a new explanation for the longevity of captured chloroplasts (kleptoplasts) in sacoglossan sea slugs. The controversy surrounding the three-dimensional structure of grana, its recent resolution by tomographic analyses, and the role of the CURVATURE THYLAKOID1 (CURT1) proteins in supporting grana formation are also discussed. We also present an updated inventory of photosynthetic proteins and the factors involved in the assembly of thylakoid multiprotein complexes, and evaluate findings that reveal that cyclic electron flow involves NADPH dehydrogenase (NDH)- and PGRL1/PGR5-dependent pathways, both of which receive electrons from ferredoxin. Other topics covered in this review include new protein components of nucleoids, an updated inventory of the chloroplast proteome, new enzymes in chlorophyll biosynthesis and new candidate messengers in retrograde signaling. Finally, we discuss the first successful synthetic biology approaches that resulted in chloroplasts in which electrons from the photosynthetic light reactions are fed to enzymes derived from secondary metabolism.

General characteristics of chloroplasts
The first photosynthetic eukaryotes originated more than 1000 million years ago through the primary acquisition of a cyanobacterial endosymbiont by a eukaryotic host, which subsequently gave rise to glaucophytes (whose photosynthetic organelles are called “cyanelles”), red algae (containing “rhodoplasts”) and green algae and plants (with “chloroplasts”). Other major photosynthetic eukaryotic lineages arose when eukaryotic hosts engulfed a free-living photosynthetic eukaryote (e.g. red or green alga), initiating secondary and tertiary endosymbioses [1]. Therefore, chloroplasts are organelles that are characteristic of plant and green algal cells, but still exhibit many prokaryotic features.

During evolution, the cyanobacterium-derived genome has undergone a dramatic reduction in size, mainly as a result of outright gene loss and the large-scale transfer of genes to the nuclear genome. Thus, the genomes of modern chloroplasts (plastomes) contain only 120-130 genes, most of which encode components of the organelle’s gene expression machinery and its photosynthetic apparatus, and are organized in nucleoids that show both prokaryotic and eukaryotic features. However, chloroplasts contain many more protein species than their plastomes can code for. Hence, the majority of chloroplast proteins are now encoded by the nuclear genome and must be imported post-translationally into the organelle [2].

Apart from photosynthesis, chloroplasts are capable of performing many other specialized functions that are essential for plant growth and development — nitrate and sulphate assimilation, and the synthesis of amino
acids and fatty acids, chlorophyll and carotenoids. To carry out these tasks, their membrane systems are equipped with specialized transport functions. The outer and inner envelope membranes mediate the import and sorting of proteins and the exchange of metabolites, while protein complexes in the thylakoid membranes implement the proton and electron transport processes that are an essential part of the photosynthetic light reactions. The thylakoids of land plants, where photosynthesis takes place, display an intricate architecture, with regions of stacked and appressed thylakoid membranes forming so-called grana. Moreover, plastids communicate with the nucleus by retrograde signaling to adjust the expression of nuclear genes according to the metabolic and developmental state of the organelle.

**Chloroplast evolution**

**Primary endosymbiosis: a ménage à trois?**

New evidence implies that primary endosymbiosis might have been more complex than has been envisaged hitherto, possibly involving a “ménage à trois”. Members of the genus *Chlamydia* are obligate intracellular bacteria, which include important pathogens of humans and other animals, and are found as endosymbionts in amoebae and insects. Although Chlamydiae are not found in plants, an unexpected number of chlamydial genes share significant similarities with plant genes [3,4], and these often contain a plastid-targeting signal [5]. In several studies, between 21 and 55 genes were shown to be transferred between Chlamydiae and primary photosynthetic eukaryotes [6-8]. This suggests that a protist lineage that could enter into a symbiosis with a particular cyanobacterium was routinely infected by an ancestor of extant *Chlamydia* that facilitated the establishment of the cyanobacterial endosymbiont by Chlamydia-to-protist lateral gene transfer [7]. Recent studies suggest that the chlamydial symbiont compartment was probably not the site of any essential biochemical pathway and was maintained only until all possible chlamydial genes had been transferred to the host [6]. Thus, the two critical steps in primary plastid endosymbiosis might have been the secretion of effector proteins into the host cytosol by intracellular chlamydial pathogens, together with the maintenance of the afflicted host by the cyanobiont, which supplied photosynthetic carbon to a chlamydia-controlled assimilation pathway [6]. If such complex interactions were indeed necessary for the establishment of the primary endosymbiotic relationship between plastid and host cytoplasts, this could explain why endosymbiotic relationships between heterotrophs and photoautotrophs were so rarely successful in the long term [7].

**A second primary endosymbiosis**

Evidence has also emerged for an independent instance of the primary endosymbiotic acquisition of a cyanobacterium — by the rhizarian amoeba *Paulinella chromatophora* about 60 million years ago [9,10]. This organism contains stably transmitted cyanobacterium-like photosynthetic organelles termed “chromatophores”, the genome of which encodes about a quarter of the protein-coding genes that can be found in its free-living relative *Synechococcus* WH5701 [10]. Eleven putative pseudogenes were identified, indicating that reductive genome evolution is ongoing. More than 30 expressed genes have been transferred from the chromatophore to the nuclear genome of the host. In the case of three photosynthetic genes that now reside in the nucleus, biochemical evidence indicates that their products are synthesized in the amoeba cytoplasm and delivered to the chromatophores, where they form complexes with chromatophore-encoded subunits [11]. This highlights *P. chromatophora* as an exceptional model for the study of early events in the generation of an organelle, and suggests that protein import into bacterial endosymbionts might be more widespread than is currently assumed [11].

**What is the basis for the longevity of kleptoplasts?**

Kleptoplasts are a special case of transient internal photosynthetic symbionts in otherwise non-photosynthetic eukaryotes. In contrast to some lineages, in which the cells of photosynthetic symbionts are retained in their entirety (“photosymbionts”), other eukaryotes collect and retain only the chloroplasts of photosynthetic species, generating structures termed “kleptoplasts” (reviewed in [12]). The most dramatic kleptoplast association known to date occurs in the sacoglossan sea slug *Elysia chlorotica*, which can maintain photosynthetically active kleptoplasts derived from ingested xanthophyte algae for up to 10 months [13]. This gives these animals their distinctive green colour, which is why they are also called “leaves that crawl”, “solar-powered slugs” or “photosynthetic slugs”. It is widely assumed that the slugs survive starvation by means of kleptoplast photosynthesis, yet direct evidence for this is lacking. Moreover, the inference that kleptoplasts require many proteins in order to support a photosynthetic lifestyle implies that essential genes for photosynthesis have been transferred by lateral gene transfer (LGT) from the alga to the slug, and in fact one instance of a tentative transfer has been reported so far [14]. However, no evidence for massive LGT has been obtained, and genome- and transcriptome-wide approaches actually argue against it [15,16]. Recently, doubts have been raised as to whether these molluscs are actually dependent upon photosynthesis, and the role of light in the survival of the sea slugs was reinvestigated [17]. Surprisingly, photosynthesis was
found not to be essential for the slugs to survive months of starvation, which explains the lack of LGT from alga to animal in these species. A possible explanation for the longevity of the sacoglossan kleptoplast was suggested previously: plastids that remain photosynthetically active within slugs for periods of months share the property of encoding FtsH, a D1 quality-control protease that is essential for photosystem II repair [18]. A replenishable supply of chloroplast-encoded FtsH could, in principle, rescue kleptoplasts from D1 photodamage, thereby influencing plastid longevity in sacoglossan slugs.

Chloroplast structure

Nucleoids

A single mesophyll chloroplast can contain up to 300 chromosomes, which are organized into complex structures called “nucleoids”, each consisting of 10-20 copies of the plastid genome, together with RNA and various proteins [for a recent review see [19]]. Owing to their endosymbiotic origin and the fact that photosynthetic metabolism goes on all around them, nucleoids have a unique composition and organization, and display features typical of prokaryotic nucleoids, as well as attributes of eukaryotic chromatin. Nucleoids contain all the enzymes necessary for transcription, replication and segregation of the plastid genome (reviewed in [20]). In addition, mRNA processing and editing, as well as ribosome assembly, take place in association with the nucleoid, suggesting that these processes occur cotranscriptionally. However, few nucleoid proteins have been characterized in detail [19,21].

Proteomic analysis of nucleoid preparations has identified new DNA-binding proteins, some of which were not inherited from the prokaryotic ancestors [22,23]. One group of proteins in particular have been described, which contain a so-called SWIB domain that has previously been shown to be part of chromatin remodelling complexes in yeast. This domain is present in 20 proteins in Arabidopsis, and at least four of these are located in the chloroplast [23]. The SWIB-domain proteins in chloroplasts are small proteins with a high isoelectric point and a high lysine content and might serve as functional replacements for the bacterial histone-like, DNA-binding HU proteins known from Escherichia coli [23]. Thus one of them, SWIB-4, has a histone H1-like motif and binds to DNA, and recombinant SWIB-4 has been shown to induce compaction and condensation of nucleoids, and functionally complements an E. coli mutant that lacks the histone-like nucleoid structuring protein H-NS [23].

The two suppressor of variegation 4 (SVR4) proteins (SVR4 and SVR4-like), originally identified in Arabidopsis [24], both have orthologues in all dicot and monocot plants sequenced so far, whereas the moss Physcomitrella patens and spike moss Selaginella moellendorffii contain only one gene copy, indicating that a gene duplication took place in the progenitor of vascular plants [25]. Inactivation of either SVR4 or SVR4-like in Arabidopsis results in seedling lethality [24,26]. Both proteins are localized in the chloroplast, expressed during early stages of chloroplast development, and contain 20% negatively charged amino acid residues [26]. Given the inherent risk of random aggregation of the negatively charged nucleic acids and basic proteins, such as histones and ribosomal subunits [27-29], SVR4 and SVR4-like could function as negatively charged molecular chaperones that mimic nucleic acids or serve as decoys [28,30] to allow for the establishment of productive DNA/RNA-protein interaction in developing chloroplasts, where dramatic rearrangements in nucleoid organization take place [26].

Thylakoid architecture

A structural hallmark of thylakoid membranes in plants are the so-called “grana” (reviewed in [31]). Grana cylinders are made up of stacks of flat grana membrane disks with a diameter of about 300-600 nm, which are enwrapped in (and interconnected by) the unstacked stroma lamellae. Tightly curved margins form the periphery of each discoid sac. For a typical granum from Arabidopsis thaliana the membrane bilayers are on average 4.0 nm thick, lumen thickness is 4.7 nm and disks are separated by a 3.6 nm gap [32].

The exact three-dimensional architecture of grana is still under debate, and two quite different types of models have been proposed: the “helical” model and several “fork/bifurcation” models. In the helical model, thylakoids comprise a network of stroma lamellae, which wind around grana stacks as a right-handed helix, connecting individual grana disks via narrow membrane protrusions (Figure 1). The grana are connected to each other solely by the stroma lamella helices, which are tilted at an angle ranging from 10 to 25°, with respect to the grana stacks [33-35], and make multiple contacts with successive layers in the grana through slits located in the rims of the stacked disks. The fork/bifurcation models, on the other hand, postulate that the grana themselves are formed by bifurcations of stroma lamellae. Thus, Arvidsson and Sundby (1999) suggested that a granum contains piles of repeat units, each containing three grana discs, which are formed by symmetrical invaginations of a thylakoid pair caused by the bifurcation of the thylakoid membrane [36] (Figure 1). Shimoni et al. (2005) presented another model in which grana discs are paired units formed by
simple bifurcation of stroma thylakoids [37] (Figure 1). Here, the granum-stroma assembly is formed by bifurcations of the stroma lamellar membranes into multiple parallel discs. The stromal membranes form wide lamellar sheets that intersect the granum body roughly perpendicular to the long axis of the granum cylinder [37,38]. In this model, adjacent granum layers are joined not only through the stroma lamellae, but via the bifurcations and through direct membrane bridges. This model can also be used to explain the rearrangements seen in thylakoids during state transitions [39]. The mutual incompatibility of the helical and bifurcation models has led to a great deal of debate [33,35,38,40,41], but recent tomographic data clearly support the helical model [33,40].

**Lateral heterogeneity of thylakoids**

The term “lateral heterogeneity” refers to the observation that grana and stroma lamellae differ in their protein composition. Photosystem II and light-harvesting complex (LHC)II are concentrated in the grana, while photosystem I with its LHCI and the chloroplast ATP synthase are localized in the unstacked thylakoid regions, that is the stroma lamellae and grana end membranes. The cytochrome $b_{6}f$ complex (Cyt $b_{6}f$) can be found in both appressed and non-appressed regions of thylakoids (reviewed in [41,42]). The NDH complex and the PGR1-PGR5 heterodimer – the two thylakoid complexes specifically involved in cyclic electron flow – are less abundant than the aforementioned four major thylakoid complexes and are located in the stroma lamellae [43–45], where they can functionally interact with photosystem I as an electron donor. While the bulkiness of the NDH complex precludes its location in grana, PGR1 homodimers have been detected in grana [43].

Detection of several of the major thylakoid multiprotein complexes in margin-enriched fractions of thylakoids by biochemical methods has been reported in some experiments. However, the marked curvature of thylakoid membranes at the grana margins is essentially incompatible with the presence of the larger multiprotein complexes at these sites. Therefore, grana margins have been thought to be essentially protein-free (reviewed in [42]). However, following the recent demonstration, by immunogold labelling, that CURT1 proteins — small polypeptides with two transmembrane regions and a putative N-terminal amphipathic helix — are localized to grana margins [46], this view must be revised. Interestingly, the CURT1 proteins appear to

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**Figure 1. The helical model of thylakoid architecture**

A fretwork of stroma lamellae, which wind around the ascending grana stacks as a right-handed helix connects to individual grana discs via narrow membrane protrusions (indicated by dotted circles in the side view). Adapted with permission [31] 2014. Journal of Experimental Botany doi:10.1093/jxb/eru090.
control the level of grana stacking, which points to an unsuspected role of grana margins in regulating the fraction of thylakoid membranes incorporated into the appressed regions that make up grana.

**Chloroplast functions**

**Chloroplast proteins: mutants and proteomes**

Estimates for the size of the chloroplast proteome in Arabidopsis range from 2000 [47] to 4400 (http://www.plastid.msu.edu/) different proteins. In the course of the Chloroplast 2010 Project (http://www.plastid.msu.edu/), homozygous mutants for several thousand nuclear genes with chloroplast functions were identified and phenotypically characterized. Despite extensive screening, for several hundred genes no homozygous mutant alleles were discovered, suggesting that these might represent genes with essential functions. More recently, lines that had failed to yield any homoygotes when grown in soil were tested for homozygous lethality owing to defects either in seed or seedling development [48,49]. Mutants arrested at various stages of seed development (and with defects in seedling development that responded to supplementation with sucrose, amino acids or to CO2 enrichment) were indeed uncovered. This resulted in an annotation of more than 200 pubically available Arabidopsis mutants, including 36 and 33 genes with one and two, respectively, independent seed- or seedling-development-defective mutant alleles. The study also resulted in the submission of 521 homozygous mutants and 128 seed stocks segregating for lethal alleles to the Arabidopsis Biological Resource Center (ABRC; http://www.arabidopsis.org).

Proteomics usefully complement the reverse genetics approach to chloroplast function outlined above. Driven by recent advances in bioanalytical and computational technologies, the strategy allows for identification, and reasonably accurate quantification, of thousands of proteins in complex mixtures, as well as the ability to characterize post-translational modifications, such as acetylation, glycosylation and phosphorylation (reviewed in [50]). An attempt to obtain a high-quality inventory of the plastid proteome has led to the identification of 1564 and 1559 proteins for maize and Arabidopsis, respectively [51]. These estimates were based on both manual curation of published experimental information, including more than 150 proteomics studies devoted to different subcellular fractions, and new quantitative proteomics experiments on plastid subfractions. These figures correspond to an estimated 40% and 50% of all plastid proteins in maize and Arabidopsis, respectively — the most comprehensive inventory assembled so far. Recently, members of the Arabidopsis proteomics community decided to develop a summary aggregation portal that is capable of retrieving proteomics data from a series of online resources on the fly [52]. The web portal is known as the MASCt Gator and can be accessed at the following address: http://gator.masc-proteomics.org.

**Chlorophyll biosynthesis**

Biosynthesis of chlorophyll takes place in the plastid, and the initial steps in the pathway leading to protoporphyrin IX are common to the biosynthesis of other tetrapyrroles, such as heme. Important discoveries in chlorophyll biosynthesis include the demonstration that plastid glutamyl-transfer RNA is involved in the formation of glutamate-1-semialdehyde [53], which is subsequently converted into 5-aminoethylgulonic acid — the universal precursor of tetrapyrrole biosynthesis in all organisms (reviewed in [54]), and the finding that the enzyme Mg-chelatase (which catalyzes the insertion of the Mg2+ ion into protoporphyrin IX) contains three different protein subunits: ChlH, ChlI, and ChlD [55,56]. Later, the GENOME UNCOUPLER4 (GUN4) was found to bind both the substrate and the product of the Mg-chelatase, thereby dramatically enhancing the activity of the enzyme [57]. GUN4 also reduces the threshold Mg2+ concentration required for activity at low porphyrin concentrations [58], and it was proposed to have a protective function in tetrapyrrole trafficking [59] and to control Mg-chelatase activity at physiologically significant Mg2+ concentrations [58]. More recently, it was shown that GUN4 interacts with the ChlH subunit of the enzyme [60,61].

One of the least understood steps in chlorophyll biosynthesis is the formation of the isocyclic “fifth” ring (ring-E), which is catalyzed in plants by the aerobic cyclase system (ACS). The overall cyclase reaction is a six-electron oxidation proposed to occur in three sequential steps: (a) hydroxylation of the methyl-esterified ring-C propionate by incorporation of atmospheric oxygen; (b) oxidation of the resulting alcohol to the corresponding ketone; (c) reaction of the activated methylene group with the γ-mesocarbon of the porphyrin nucleus in an oxidative reaction involving the removal of two protons to yield the “fifth” ring [62]. At the biochemical level, the ACS requires both soluble and membrane-bound chloroplast fractions and, in barley, at least two mutants exist (*xantha-l* and *viridis-k*) which are defective in the membrane components [63]. Thus, the ACS may be composed of three gene products: a soluble protein and two membrane-bound chloroplast fractions and, in barley, at least two mutants exist (*xantha-l* and *viridis-k*) which are defective in the membrane components [63]. Thus, the ACS may be composed of three gene products: a soluble protein and two membrane-bound components — one encoded in barley by *Xantha-l* and the other by *Viridis-k*. So far, only AcsF (which corresponds to Xantha-l in barley, CRD1 in Chlamydomonas or CHL27 in Arabidopsis) has been unambiguously identified [63–65]. As diiron enzymes
are known to perform hydroxylation and cyclization of keto intermediates, AcsF could be involved in one or more of the proposed cyclase steps. Recent progress has come from pull-down experiments using FLAG-tagged versions of the two AcsF-like gene products in Synechocystis in combination with protein mass spectrometry, which have identified the soluble YCF54 protein as a new putative subunit of ACS [66]. Inactivation of the Synechocystis\textit{ycf54} gene resulted in significantly reduced chlorophyll levels, marked accumulation of the substrate of the cyclase, Mg-protoporphyrin IX methyl ester, and only traces of its product, protochlorophyllide, indicating that YCF54 is essential for the activity and/or stability of the oxidative cyclase. Future experiments must clarify whether YCF54 is the long-sought soluble component of the cyclase system, or whether it functions in AcsF synthesis/maturation or in cyclase assembly. Low chlorophyll accumulation \textit{A} (LCAA), the tobacco homologue of YCF54, might have an additional role in the feedback-control of 5-aminolevulinic acid biosynthesis [67]. The second pathway mediating CEF involves the so-called “NAD(P)H dehydrogenase complex” or “NDH complex”. Although the plant NDH complex is related to the NADH dehydrogenase complexes of bacteria and mitochondria, its function and composition are enigmatic. Recently, the Shikanai group has identified three novel subunits of plant NDH (CRR-31, -J and –L) and their functional characterization clearly indicated that CRR-31 supplies a docking site for ferredoxin [71,72]. Therefore it can be concluded that the plant NDH complex accepts electrons from ferredoxin rather than NAD(P)H. Consequently, the authors of the first study proposed that the term “NDH” be retained, but used to mean “NADH dehydrogenase-like complex” rather than “NAD(P)H dehydrogenase complex” [71]. In a strict sense, the NDH complex is also an

### Photosynthesis: new proteins and new functions

It comes as a surprise to learn that some proteins that are directly involved in the light reactions of photosynthesis have remained unidentified until very recently. Thus, although antimycin A-sensitive cyclic electron flow (AA-sensitive CEF), which serves to recycle electrons from ferredoxin to plastocyanin, was discovered by Arnon and co-workers more than 50 years ago, it is only a few years since the proteins responsible were identified. A role in AA-sensitive CEF has been attributed to the two thylakoid proteins PGR5 [68] and PGRL1 [69] ever since their identification, but this assignment has remained controversial. Indeed, current technical limitations still preclude unequivocal clarification of their precise function in CEF \textit{in vivo}, but recent biochemical experiments have shown that PGRL1-PGR5 complexes possess ferredoxin-plastoquinone reductase (FQR) activity \textit{in vitro} [43]. Consequently, PGRL1-PGR5 complexes in flowering plants appear to shuttle between photosystem I and the cytochrome (Cyt) \textit{b}$_{6}$ complex, whereas in the green alga Chlamydomonas PGRL1 (but not PGR5) has been detected in a photosystem I cytochrome \textit{b}$_{6}$ supercomplex that has intrinsic CEF activity [70].

### Table 1. Accessory factors involved in the assembly of thylakoid multiprotein complexes in plants and cyanobacteria

| PSII | PSI | Cyt \textit{b}$_{6}$ | cpATPase | NDH |
|------|-----|------------------|-----------|------|
| HCF136 [103]/YCF48 [104] | YCF3 [85,86] | CCS1 [105] | ALB4 [106] | AtCYP2-2 [107] |
| ALB3 [108,109]/Sr147 [110] | YCF4 [87,88] | CCB1 [111,112] | AtCGL160 [113] | CRR1 [114] |
| YCF39 [115] | YCF39 [116] | CCB2 [111,112] | CRR6 [118] | Synechocystis YCF3 [119] |
| LPA1 [120]/REP27 [121] | PPD1 [122] | CCB2 [111,112] | CRR7 [123] | |
| LPA2 [124] | Y3P1 [125] | DAC [126] | CRR41 [127] | |
| LPA3 [128] | PBF1 [129] | | CRR42 [127] | |
| Sr2013 [130] | HCF101 [131–133] | | NDF5 [134] | |
| Psb27 [135–138] | RubA [140,141] | | PAM68L [75] | |
| LPA19 [139] | Psb28 [142] | | | |
| Psb29/THF1 [143–145] | | | | |
| PratA [146] | | | | |
| Ptc [147] | AtCYP38 [148] | | | |
| PAM68 [149] | | | | |

**Organism in which the assembly factor was functionally characterized:** A, Arabidopsis, C, Chlamydomonas, S, Synechocystis or Synechococcus, T, tobacco. **Abbreviations:** cpATPase, chloroplast ATP synthase; Cyt, cytochrome; PSI, photosystem I; PSII, photosystem II.

Table 1. Accessory factors involved in the assembly of thylakoid multiprotein complexes in plants and cyanobacteria
inventory of assembly factors identified in Arabidopsis, Chlamydomonas, tobacco or cyanobacteria (Synechocystis or Synechococcus) (Table 1). Interestingly, two photosystem I assembly factors are encoded by chloroplast genes: ycf3 and ycf4 [85-88]. An important function in the chloroplast protein import machinery has been recently assigned to another chloroplast open reading frame, ycf1 [89]. However, such a tentative assignment of an important function of the encoded Tic214 protein is somehow at variance with the observation that chloroplast genomes of Poaceae species lack the ycf1 gene.

**Novel retrograde signals**

The term “retrograde signalling” refers to the idea that signals emanating from chloroplasts or mitochondria can modulate nuclear gene expression. Proposed almost 30 years ago, the initial notion that a single plastid signal might regulate the expression of nuclear genes involved in plastid biogenesis has since expanded to accommodate the insight that multiple signals are produced by plastids. While the ultimate effects of retrograde signalling on nuclear gene expression have now been clearly defined, many aspects of the initiation and transmission of the signals, and their mode of action, remain unresolved, speculative or controversial [90,91]. Relevant signals are thought to be derived from various sources, including (a) the pool of reactive oxygen species (ROS), (b) the reduction/oxidation (redox) state of the organelle, (c) organellar gene expression, and (d) the tetrapyrrole pathway. More recently, “brand-new” retrograde signaling pathways have been described that involve (e) metabolites — particularly 3'-phosphoadenosine 5'-phosphate (PAP) [92] and methylenylthritol cyclodiphosphate (MeCp) [93] — and (f) a carotenoid derivative (β-cyclotetral [β-CC]) [94].

**Synthetic biology**

Synthetic biology can broadly be defined as “the deliberate (re)design and construction of novel biological and biologically based systems to perform new functions for useful purposes, that draws on principles elucidated from biology and engineering” (http://www.erasynbio.eu/index.php?index=32).

The plastome, at least in some species, such as tobacco, tomato and Chlamydomonas, can be manipulated by genetic transformation with large constructs made up of foreign or synthetic DNA segments [95]. In fact, due to its prokaryotic origin, the chloroplast genome offers many advantages for genetic engineering because its genes are organized in operons and many are co-expressed from a single promoter as a polycistronic transcript that may subsequently be processed further into monocistronic mRNAs. Moreover, no position effects or epigenetic
gene-silencing mechanisms, like those observed with nuclear transgenes, have been reported in chloroplasts [96]. These features make the chloroplast compartment especially amenable to the application of synthetic biology to goals such as the sustainable synthesis of chemicals and high-value products. Lu et al. [97] successfully demonstrated this by expressing the tocochromanol pathway (which produces tocopherols and tocotrienols, collectively called “vitamin E”) in the chloroplasts of tobacco and tomato and achieving up to a tenfold increase in total tocochromanol accumulation. This represents a prime example of how overexpression of enzymes in the chloroplast can redirect photosynthetically generated carbon skeletons from the endogenous isoprenoid biosynthetic pathway into the production of higher levels of tocopherols and tocotrienols.

It is highly desirable that novel pathways introduced into the chloroplast should be able to tap directly the chemical energy derived from sunlight in the form of ATP, NADPH or even photo-reduced ferredoxin. One group of enzymes which could potentially be used for this purpose are the cytochrome P450 mono-oxygenases (P450s), which are represented in all biological kingdoms and constitute one of the largest superfamilies of enzymes known [98]. Most P450s are located in the endoplasmatic reticulum, where they act as key enzymes in the biosynthesis of a large number of high-value bioactive natural compounds. Many of these compounds are normally made in very small quantities and are difficult to produce by chemical synthesis due to their often complex structures [99]. P450s generally obtain the electrons needed for their catalytic reactions from NADPH or NADH, but bacterial and mitochondrial P450s are also known to accept electrons from ferredoxin. Therefore, a direct link between photoreduced ferredoxin and P450s is possible if the evolutionary compartmentalization of the photosystems in the chloroplasts and of the majority of the P450 pathways in the endoplasmatic reticulum can be broken down.

The potential value of combining P450-mediated monooxygenation reactions with photosynthesis was first demonstrated in vitro when spinach chloroplasts were brought together with microsomes from yeast expressing a fusion between a P450 from rat (CYP1A1) and a reductase. This mixture supported the light-driven conversion of the P450 substrate 7-ethoxycoumarin into 7-hydroxycoumarin [100]. More recently, it was shown in vitro that electrons supplied by photosystem I purified from barley could be transferred with high efficiency to a P450 (CYP79A1) from Sorghum bicolor via ferredoxin, thus eliminating the need for an NADPH recycling system and a reductase [101]. Subsequently, it was shown that the P450-catalysed pathway for the biosynthesis of dhurrin (a cyanogenic glycoside) can be transferred from the cytosolic endoplasmatic reticulum of S. bicolor into the tobacco chloroplast [102]. To this end, fusion proteins between a chloroplast transit peptide and the coding regions of two P450 enzymes and a uridine 5’-diphosphate (UDP) glucosyltransferase, which together constitute the route to dhurrin biosynthesis, were successfully expressed in the chloroplasts of transiently transformed tobacco leaves. Interestingly, the chloroplast was able to provide the heme cofactor for the proper assembly of the P450s, the tyrosine and UDP-glucose substrates. The electron-demanding P450-catalysed synthesis of dhurrin was driven by directly tapping into light-driven reduction of ferredoxin by photosystem I (Figure 3). Thus, this example demonstrates that P450s that normally reside in the endoplasmic reticulum membranes can be targeted to the chloroplast and inserted into the thylakoids and can act as receptors for electrons from the light reactions of photosynthesis for use in the biosynthesis of dhurrin.

**Abbreviations**

AA-sensitive CEF, antimycin A-sensitive cyclic electron; ACS, aerobic cyclase system; Cyt, cytochrome; FNR, ferredoxin NADP⁺ oxidoreductase; GUN4, GENOME UNCOUPLER4; FQR, ferredoxin plastoquinone reductase;
LHC, light-harvesting complex; MECPP, methylethylthiol cyclodiphosphate; NDH, NAD(P)H dehydrogenase; PAP, 3'-phosphoadenosine 5'-phosphate; SVR4, suppressor of variegation 4; UDP, uridine 5'-diphosphate.

Disclosures
The authors declare that they have no disclosures.

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