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1 **OHP2 is not required for *psbA* translation in *Chlamydomonas***

2

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22 **RUNNING TITLE**

23 D1 synthesis uncoupled from cofactor assembly

24

25 **ABSTRACT**

26 In land plants and cyanobacteria, co-translational association of chlorophyll (Chl) to  
27 the nascent D1 polypeptide, a reaction center protein of photosystem II (PSII), requires  
28 a Chl binding complex consisting of a short-chain dehydrogenase (HCF244/Ycf39) and  
29 One-Helix Proteins of the LHC superfamily (OHP1 and OHP2 in chloroplasts). Here,  
30 we show that an *ohp2* mutant of the green alga *Chlamydomonas reinhardtii* fails to  
31 accumulate core PSII subunits, in particular D1. Extragenic suppressors arise at high  
32 frequency, suggesting the existence of another route for Chl association to PSII. The

33 *ohp2* mutant can be complemented by the *Arabidopsis* ortholog. In contrast to land  
34 plants, where *psbA* translation is prevented in the absence of OHP2, ribosome profiling  
35 experiments show that the *Chlamydomonas* mutant translates the *psbA* transcript over  
36 its full length. Pulse labelling suggests that D1 is degraded during or immediately after  
37 translation. The translation of other PSII subunits is affected by assembly-controlled  
38 translational regulation (the CES process). Proteomics show that HCF244, a  
39 translation factor which associates with and is stabilized by OHP2 in land plants, still  
40 partly accumulates in the *Chlamydomonas ohp2* mutant, explaining the persistence of  
41 *psbA* translation. Several Chl biosynthesis enzymes overaccumulate in the mutant  
42 membranes. Partial inactivation of the D1-degrading FtsH protease restores a low level  
43 of PSII activity in an *ohp2* background, but not photoautotrophy. Taken together, our  
44 data suggest that OHP2 is not required for *psbAD1* translation in *Chlamydomonas*, but  
45 necessary for its stabilization.

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47

#### 48 **KEYWORDS**

49 One-Helix Protein, photosystem II, light harvesting-like family, D1 protein, *psbA* mRNA,  
50 ribosome profiling, targeted proteomics, control by epistasy of synthesis; cross-species  
51 complementation; FtsH protease

## 52 INTRODUCTION

53 Oxygenic photosynthesis, carried out by cyanobacteria and photosynthetic eukaryotes,  
54 is based on the light-driven electron transfer from water to NADPH, involving three  
55 complexes located in the thylakoid membrane and operating in series: photosystem II  
56 (PSII), cytochrome *b<sub>6</sub>f* (*Cytb<sub>6</sub>f*) and photosystem I (PSI). PSII is a multi-subunit  
57 complex, whose core complex is made up of 20-23 subunits: the reaction center  
58 proteins D1 (PsbA), D2 (PsbD), *Cytb<sub>559</sub>* (subunits PsbE and PsbF) and PsbI are  
59 encoded in the chloroplast genome by the *psbA*, *psbD*, *psbE/F* and *psbI* genes, as are  
60 the core antenna CP47 (PsbB) and CP43 (PsbC) and the additional subunits PsbH,  
61 PsbJ, PsbK, PsbL, PsbM, PsbT and PsbZ (reviewed in e.g. Gao et al., 2018). In  
62 eukaryotes, nucleus-encoded subunits complete the complex and build its peripheral  
63 chlorophyll (Chl) *a/b* containing light-harvesting antenna complexes (LHCs). Chl *a* and  
64 its derivative pheophytin (Phe) *a* are essential components of the intra-PSII electron  
65 transfer chain that allows the generation of a stable charge separated state after light-  
66 capture. The pigments and redox-active cofactors are scaffolded mainly by the  
67 homologous D1 and D2 polypeptides, which also provide Tyr residues involved in re-  
68 duction of P680<sup>+</sup> and residues liganding the manganese (Mn)-cluster where water-  
69 splitting occurs.

70 Given the tight folding of the enzyme and the high chemical reactivity of Chl cation  
71 radicals that can form from the light-excited states, it has long been hypothesized that  
72 specific mechanisms must coordinate the synthesis of the reaction center proteins,  
73 their intra-membrane insertion and their association with pigments. Over the years, the  
74 biogenesis of PSII has been dissected into a series of discrete assembly steps  
75 (reviewed in Nickelsen and Rengstl, 2013). Precursor D1 (pD1), co-translationally  
76 inserted into the membrane, binds to PsbI and specific assembly factors, forming a D1  
77 module that will associate with a D2 module formed between D2, *Cytb<sub>559</sub>* and other  
78 factors (Knoppová et al., 2022; Komenda et al., 2012). The reaction center (RC)  
79 subcomplex will first integrate CP47, then CP43, associated with other small subunits,  
80 forming the monomeric PSII core. Dimerization ensues, as well as association with  
81 LHCs integrated into the membrane by the CpSRP complex. On the luminal surface of  
82 the enzyme, cleavage of the C-terminal extension of pD1 and light-driven assembly of  
83 the Mn-containing cluster allow formation of the water-splitting PSII enzyme.

84 Over the years, numerous proteins have been found to catalyze various steps of  
85 this pathway, both during *de novo* assembly and during repair after photoinhibition

86 (Heinz et al., 2016; Lu, 2016). Some act by regulating gene expression, in particular  
87 translation of *psbA* which codes for the rapidly-turning over D1. Others act as a  
88 chaperone, binding an assembly intermediate until the next step can be completed. In  
89 this category, particular attention has been paid to proteins that could mediate the  
90 assembly of the cofactors. It is believed that Chl molecules are presented to the  
91 acceptor PSII subunits by specific carrier proteins that precisely mediate their insertion  
92 at the proper position, at the proper stage of assembly.

93 In *Arabidopsis*, a complex consisting of two conserved One-Helix Proteins, OHP1 and  
94 OHP2, and a protein related to short-chain dehydrogenase/reductases, HCF244, has  
95 been found to be essential for Chl integration into PSII, or for protection of the newly  
96 synthesized Chl-associated D1 during formation of the RC complex (Chotewutmontri  
97 and Barkan, 2020; Chotewutmontri et al., 2020; Hey and Grimm, 2018; Li et al., 2019;  
98 Myouga et al., 2018). In combination with OHP1, OHP2 is able to bind Car and Chl,  
99 the latter via specific residues of a Chl binding motif (Hey and Grimm, 2020), and the  
100 reconstituted heterodimer has unique photoprotective properties (Psencik et al., 2020).  
101 The **OHP1/OHP2/HCF244 Complex** (which we will hereafter call "OHC") is  
102 homologous to a complex of similar function described in cyanobacteria, where the  
103 relative stability of the assembly intermediates in core subunits mutants has allowed a  
104 fine dissection of the pathway (Komenda et al., 2012). OHP1 and OHP2 resemble  
105 cyanobacterial High Light-Inducible Proteins (HLIPs) encoded by the *hliA-D* genes  
106 (Komenda and Sobotka, 2016), in that they all present a single transmembrane domain  
107 showing the key residues for binding Chl (Engelken et al., 2010). While OHP1 clearly  
108 derives from HLIPs, the origin of OHP2 is less clear (Engelken et al., 2010). HliC and  
109 HliD have been found, together with the HCF244 homolog Ycf39, to associate with the  
110 D1 module and then to the RC subcomplex after binding of the D2 module (Chidgey  
111 et al., 2014; Knoppová et al., 2014). HliD can bind Chl and  $\beta$ -carotene (Staleva et al.,  
112 2015). The Chl synthase ChlG which carries an N-terminal HLIP domain co-  
113 immunoprecipitates with HliD/Ycf39 and it is believed that the complex can deliver  
114 newly synthesized Chl to D1 and/or D2 during or early after their translation (Chidgey  
115 et al., 2014; Staleva et al., 2015). For this process, Chl could also be scavenged from  
116 other Chl-binding proteins upon their degradation.

117 The phenotype of land plant mutants has revealed an additional function of the  
118 OHC, namely to regulate *psbA* translation. *Arabidopsis ohp1*, *ohp2* and *hcf244*  
119 mutants show strongly reduced PSII levels, while only weak or indirect effects are

120 observed on PSI accumulation (Beck et al., 2017; Li et al., 2019; Link et al., 2012;  
121 Myouga et al., 2018). In the three mutants, the other two partners of the OHC are  
122 undetectable or dramatically reduced, suggesting that formation of the complex is  
123 required for their stabilization (Beck et al., 2017; Hey and Grimm, 2020; Li et al., 2019).  
124 Interestingly, *ohp1* and *ohp2* mutants from *Arabidopsis* as well as an *hcf244* mutant  
125 from maize show a strong reduction (resp. 7-fold, 12-fold and 11-fold) of the ribosome  
126 footprint RF-Seq signal (RF-Seq; Chotewutmontri et al., 2020), indicating near  
127 complete inhibition of *psbA* translation (RF-Seq; Chotewutmontri et al., 2020). This has  
128 led to the hypothesis that the OHC regulates *psbA* translation by sensing the  
129 production of D1 properly loaded with Chl: as long as D1 remains bound to the OHC,  
130 HCF244 is unable to catalyze *psbA* translation initiation and D1 translation is halted  
131 (Chotewutmontri and Barkan, 2020). Such a tight coupling between assembly and  
132 translation is reminiscent of a process described as Control by Epistasy of Synthesis  
133 (CES) in *Chlamydomonas* (Minai et al., 2006; Wollman et al., 1999). In the PSII CES  
134 cascade, unassembled D1, as produced for example in the absence of D2, represses  
135 its own translation initiation. It was thus of particular interest to determine whether such  
136 an OHC-dependent translational regulation exists in *Chlamydomonas*.

137 Many other proteins of the PSII assembly pathway are conserved between  
138 cyanobacteria, land plants and *Chlamydomonas*, but their detailed working mechanism  
139 has only rarely been investigated in the alga (Spaniol et al., 2021). Among those acting  
140 at a site close to that of the OHC, another short-chain dehydrogenase/reductase,  
141 HCF173, has been found to be necessary for D1 synthesis and to bind the *psbA* 5'  
142 UTR, along with another protein of ill-defined function, SRRP1 (Link et al., 2012;  
143 McDermott et al., 2019; Schult et al., 2007; Watkins et al., 2020). ALB3, which belongs  
144 to a conserved family of protein integrases also found in bacteria and mitochondria,  
145 plays a major role in the insertion of integral thylakoid membrane proteins in general,  
146 and serves as a hub for many other factors (reviewed in Plöckinger et al., 2016). In the  
147 case of D1, ALB3 appears to interact with factors acting downstream of D1 integration,  
148 namely HCF136, but also LPA1 and probably LPA2 and LPA3. HCF136 (and its  
149 homolog Ycf48 in cyanobacteria) is involved, along with the chloroplast-encoded  
150 PsbN, in the interaction between the D1 and D2 subcomplexes and thus the formation  
151 of the RC subcomplex (Komenda et al., 2008; Plücker et al., 2002; Torabi et al., 2014).  
152 Further downstream, LPA2 catalyzes a step leading to the formation of stable PSII

153 monomers, probably by facilitating the integration of CP43 into the RC47 subcomplex  
154 (Cecchin et al., 2021; Schneider et al., 2014; Spaniol et al., 2021).

155 Another factor that plays an important role in the early steps of PSII assembly is a  
156 rubredoxin-like protein (Calderon et al., 2013). Here, the cyanobacterial RubA, like the  
157 accessory factor Ycf48, has been shown to be a component of the initial D1 assembly  
158 module (Kiss et al., 2019). The homologous protein RBD1 from *Chlamydomonas* was  
159 proposed to participate in the protection of PSII intermediate complexes from  
160 photooxidative damage during de novo assembly and repair (García-Cerdán et al.,  
161 2019). Moreover, Calderon et al. (2022) propose a model in which RBD1 promotes the  
162 proper folding of D1, possibly via delivery or reduction of the non-heme iron during PSII  
163 assembly. Besides a function in the early steps of PSII assembly, RBD1 from  
164 *Arabidopsis* was additionally proposed to play a role in the translation of the *psbA*  
165 mRNA (Che et al., 2022). Chl insertion into nascent D1 is not only required during PSII  
166 biogenesis, but also during repair after photoinhibition. The D1 protein is the primary  
167 target of photodamage, which includes a reversible component, repaired in the  
168 absence of protein translation, and an irreversible component that requires  
169 degradation of D1 and *de novo* synthesis (for a review see e.g. Theis and Schroda,  
170 2016). FtsH is a multi-subunit ATP-dependent thylakoid membrane metalloprotease,  
171 combining chaperone and peptidase domains. It has been shown to be a major player  
172 in the degradation of D1 and is thus essential for repair after photoinhibition (Lindahl  
173 et al., 2000; Silva et al., 2003).(van Wijk, 2015)In *Chlamydomonas*, *ftsH1* mutants  
174 show light-sensitivity and defects in PSII repair (Malnoë et al., 2014). The mutants  
175 accumulate damaged D1 and its cleavage products generated by DEG-family  
176 endopeptidases. How D1 is extracted from PSII without its total disassembly, and how  
177 degradation is coupled with synthesis of a replacement subunit, is still largely unknown.

178 In this study, we show that a null mutant of OHP2 in *Chlamydomonas* is entirely  
179 devoid of PSII. The primary defect is a lack of D1 accumulation which can be fully  
180 restored by complementation of the mutant with *Arabidopsis* OHP2. Ribosome profiling  
181 and pulse labeling experiments show that this defect is not caused by an arrest of *psbA*  
182 translation, but by a reduced stability of the nascent D1 protein. Targeted proteomics  
183 show that HCF244 accumulates to ~25% of WT in the *ohp2* mutant, explaining this  
184 partial uncoupling between the association of D1 with Chl and the initiation of  
185 translation on *psbA*. In the mutant, D1 degradation appears in part mediated by the

186 FtsH protease. An intriguing specificity of *Chlamydomonas* OHP2 is the high-frequency  
187 suppression of the PSII-less phenotype in the mutant.

## 188 **RESULTS**

### 189 **Isolation of a *Chlamydomonas* photosynthetic mutant and identification of the** 190 **mutated locus**

191 By nuclear transformation of the antibiotic resistance marker *aphVIII*, conferring  
192 resistance to paromomycin (Pm), we previously generated a collection of insertional  
193 mutants of *Chlamydomonas* (Houille-Vernes et al., 2011). The primary screening of  
194 this collection combined analysis of fluorescence induction curves with sorting of  
195 photosynthetic versus non-photosynthetic (acetate-requiring) phenotypes to  
196 categorize the affected photosynthetic complex. The strain *10.1a* was identified as  
197 acetate requiring and deficient in PSII. This strain carries a single insertion of the  
198 *aphVIII*-cassette in gene *Cre11.g480000* as revealed by inverse PCR and Southern  
199 blotting (Supplemental Figure S1A). In a backcross of *10.1a* to the WT strain WT-S34  
200 (mt+), the PSII-deficient phenotype segregated 2:2, indicating a single nuclear  
201 mutation, but it was neither linked to Pm resistance nor to the cassette inserted in  
202 *Cre11.g480000* (Supplemental Figure S1B, top row). This suggested that a second  
203 nuclear mutation causes the observed PSII phenotype.

204 Strikingly, all of the 47 PSII-deficient progenies analyzed carried the mating type  
205 minus (mt-) gene *MID*, and none the mt+ specific *FUS1* (Supplemental Figure S1B,  
206 bottom). This indicates linkage to the mating type locus, a ~1 Mb region of  
207 recombinational suppression involved in sex determination, residing on chromosome  
208 6 and comprising about 35 genes, many unrelated to sexuality (Ferris et al., 2002).  
209 Illumina genome sequencing revealed near the MT locus a single structural variant at  
210 position chromosome\_6:305949 (v5 genome assembly), i.e. within exon 3 of the *OHP2*  
211 gene (*Cre06.g251150*, Figure 1A; Supplemental Figure S2). Analysis of reads flanking  
212 this site and their mate reads pointed to the long terminal repeat (LTR) sequence of  
213 *TOC1* (Transposon Of Chlamydomonas, Day et al., 1988), indicating a *TOC1* insertion  
214 (Supplemental Data S1). Accordingly, Southern blot analysis of the *10.1a* mutant  
215 showed integration of a large DNA fragment of ~6 kb into *OHP2* (Figure 1B) and none  
216 of the 47 PSII-deficient progenies described above gave rise to an *OHP2*-specific PCR-  
217 product (Supplemental Figure S1C). To exclude an impact of the initially identified

218 mutation in *Cre11.g480000* on the mutant phenotype, most analyses were performed  
219 with a strain of the backcrossed progeny, which we will term *ohp2*, possessing only the  
220 mutation in *OHP2* but not the one in *Cre11.g480000*.

221 The *OHP2* gene of *Chlamydomonas* encodes the One-Helix Protein 2 (OHP2)  
222 belonging to the Chl *a/b* binding protein superfamily (Engelken et al., 2010). OHP2  
223 consists of 144 amino acids with the N-terminal 26 amino acids predicted to represent  
224 a chloroplast transit peptide (cTP). The expected mature protein has a molecular mass  
225 of 13.4 kDa.

## 226 **The *ohp2* mutant reveals a complete loss of PSII activity and a reduced Chl** 227 **content**

228 As summarized in Supplemental Table S1, fluorescence induction kinetics and  
229 electrochromic shift (ECS) analysis of the *ohp2* mutant revealed a complete lack of  
230 PSII, while PSI and *Cytb<sub>6</sub>f* activities appeared normal. PSII quantum yield (Fv/Fm) was  
231 negative in the *ohp2* mutant because of the small initial drop in fluorescence, typical of  
232 mutants completely lacking PSII (Figure 1C, upper panel). This is due to chemical  
233 quenching by plastoquinone brought about by the oxidation of plastoquinol by PSI.  
234 Similarly, the PSII/PSI ratio, measured from the amplitude of the initial ECS after a  
235 flash in the absence and presence of the PSII inhibitors DCMU and hydroxylamine,  
236 indicated loss of PSII charge separation (Figure 1C, lower panel). The small effect of  
237 the inhibitor treatment on the *ohp2* mutant was ascribed to hydroxylamine and was  
238 also observed in PSII-null mutants such as *psbA-FuD7* (Bennoun et al., 1986).  
239 Together, these results revealed that PSII activity is completely abolished in the mutant.

240 When cultivated as liquid culture under photo-heterotrophic conditions, the *ohp2*  
241 mutant displayed a lighter-green coloration than the wild type (WT). Determination of  
242 the Chl *a* and *b* content showed that, on a per-cell basis, the *ohp2* strain accumulated  
243 only about 58% of the total Chl present in the WT. The Chl *a* content decreased more  
244 than the Chl *b* level, leading to a decreased Chl *a*/Chl *b* ratio in *ohp2* (Supplemental  
245 Table S1), typical of photosynthetic reaction center mutants.

## 246 **Complementation of the *ohp2* mutant with *Chlamydomonas* OHP2**

247 Complementation studies were carried out using a plasmid harboring a Pm resistance  
248 cassette along with the *OHP2* sequence. We used the full length *OHP2* CDS 3'-

249 terminally fused to a hemagglutinin (HA)-tag encoding sequence, placed under control  
250 of the strong nuclear *PSAD* promoter (construct 2 in Supplemental Figure S3).  
251 Transformants first selected for Pm resistance were tested for photoautotrophic growth  
252 on minimum medium under high or low light. Approximately three quarters of the  
253 analyzed clones exhibited restored photoautotrophic growth shown exemplarily for one  
254 clone (*ohp2:OHP2-HA* in Figure 1D). This strain was chosen for further experiments,  
255 with full restoration of photosynthetic capacities, as judged from its WT-like  
256 fluorescence induction and ECS kinetics as well as largely restored Chl accumulation  
257 (Figure 1C, Supplemental Table S1). These results confirm that the PSII deficient  
258 phenotype of the strain is due to inactivation of *OHP2*. They also show that *OHP2* can  
259 accommodate a small tag at its C-terminus with no deleterious effect on its  
260 functionality.

## 261 **Functional analysis of the OHP2 protein in *Chlamydomonas***

262 Mature *OHP2* from *Chlamydomonas* contains a single C-terminal transmembrane helix  
263 including highly conserved residues required for Chl binding (Figure 2A). Based on  
264 topology experiments of Li and colleagues (2019) for the *Arabidopsis* *OHP2* protein  
265 (*AtOHP2*), the less conserved N-terminus of *OHP2* should face the chloroplast stroma,  
266 while the C-terminus is located on the luminal side. In addition, multiple sequence  
267 alignment and hydropathy analysis indicate a well-conserved hydrophobic stretch (HS)  
268 of nine residues near the C-terminus which is too short to span the lipid bilayer and  
269 seems to be missing in cyanobacterial HLIPs like HliD or HliC (compare Figure 2A;  
270 Supplemental Figure S4). This HS might be an evolutionary leftover from a second  
271 helix of the ancestral two-helix Stress-Enhanced Proteins (SEP) from which *OHP2*  
272 potentially originates (Andersson et al., 2003; Beck et al., 2017; Heddad et al., 2012).

273 To test the functional role of this C-terminal hydrophobic region we transformed the  
274 mutant with a construct for the expression of a truncated *OHP2* protein (*pBC1-*  
275 *CrOHP2-ΔHS-HA*, construct 5 in Supplemental Figure S3) where the last 14 amino  
276 acids are replaced by the HA-tag. Thirty transformants selected on Pm were tested for  
277 photoautotrophic growth on minimal medium, but none showed restoration of  
278 photoautotrophy (Figure 2B, left panel). No accumulation of the truncated *OHP2-HA*  
279 could be detected in the transformants by immunoblots using an  $\alpha$ -HA-antibody (Figure  
280 2B, right panel). These results suggest that even though cyanobacterial orthologs do

281 not have the HS, the last 14 amino acids are essential for OHP2 stability in  
282 *Chlamydomonas*.

283 A construct for the expression of a fusion of full-length OHP2 with GFP, adapted to  
284 the codon usage of *Chlamydomonas* (CrGFP; construct 3 in Supplemental Figure S3),  
285 yielded Pm-resistant transformants, but none was photoautotrophic. Presumably, the  
286 large soluble GFP-tag interferes with the function or proper localization of OHP2.  
287 However, when we directly fused the predicted transit peptide of OHP2 to CrGFP  
288 (construct 4 in Supplemental Figure S3), we observed GFP protein accumulation in  
289 two out of eight immunologically analyzed clones upon transformation of the *UVM4*  
290 strain which is known for its high capacity to express nuclear transgenes  
291 (Supplemental Figure S5A, upper panel; Neupert et al., 2009). Confocal laser scanning  
292 microscopy of one transformant showed that the GFP fluorescence signal co-localized  
293 with the Chl autofluorescence in the single cup-shaped chloroplast (Supplemental  
294 Figure S5A, lower panel), clearly distinct from the cytosolic control (CrGFP). Moreover,  
295 immunological analysis of cellular subfractions of the strain *ohp2:OHP2-HA* described  
296 above revealed a membrane localization of the ~14 kDa OHP2-HA tagged protein  
297 comparable to that of the integral thylakoid membrane protein D2 (Supplemental  
298 Figure S5B).

299 Taken together, these results indicate that OHP2 is a thylakoid membrane-localized  
300 protein. Moreover, while a C-terminal fusion of a small tag to OHP2 still allows the  
301 protein to fulfil its native function, removal of its C-terminus or introduction of a large  
302 GFP tag severely impairs its function and /or stability.

### 303 **Reduced PSII subunit accumulation and synthesis in the *ohp2* mutant**

304 As judged from the analysis of photosynthetic parameters in the *ohp2* mutant, in  
305 particular the lack of effect on PSI, OHP2 appears exclusively involved in PSII  
306 biogenesis. Accordingly, immunoblot analyses revealed no change in the accumulation  
307 of the large subunit of Rubisco (RbcL), the PSI reaction center protein PsaA or Cyt $f$  of  
308 the Cyt $b_6f$  complex (Figure 3A). In contrast, the PSII core proteins D1 and D2 were  
309 below the detection limit. Typical for mutants lacking D1 or D2, the core antenna protein  
310 CP43 accumulated to some extent, about 12% of the level in WT (Figure 3A).  
311 Complementation of *ohp2* with the *OHP2* cDNA fully restored protein accumulation in  
312 the strain *ohp2:OHP2-HA* (Figure 3A).

313 Northern blot analysis showed no significant alteration of *psbA* and *psbD* transcript  
314 accumulation in *ohp2* compared to the WT (Figure 3B). To determine whether the  
315 mutation instead affects translation or prevents stabilization of chloroplast-encoded  
316 PSII subunits, <sup>14</sup>C pulse chase labeling experiments were carried out in the presence  
317 of cycloheximide (Figure 3C). Two independently grown *ohp2* cultures revealed a  
318 reduction of radiolabeled CP47, and to a lower extent of CP43 and D2 proteins in  
319 comparison to the WT. However, the most significant effect was observed for the D1  
320 protein for which no incorporation of the radiolabel was detected, neither during the  
321 pulse nor after the chase. This suggests that the primary defect in the *ohp2* mutant is  
322 either a deficiency in D1 synthesis or a very fast degradation of newly synthesized D1.  
323 The effects on other PSII subunits may be ascribed to assembly-dependent  
324 translational control (CES; Choquet and Wollman, 2009) or reduced stability of the  
325 unassembled subunits.

326 Our efforts to raise a functional antibody against recombinant OHP2 having failed,  
327 we used the *ohp2:OHP2-HA* strain, along with the WT and *ohp2*, to examine the  
328 assembly of PSII and the location of OHP2. Crude membrane fractions were  
329 solubilized and subjected to 2D-BN/SDS-PAGE (Figure 3D). As anticipated from the  
330 unaffected accumulation of the PSI subunit PsaA (Figure 3A), all PSI-related  
331 complexes, including PSI core and supercomplexes, were assembled in the *ohp2*  
332 mutant. Remarkably, a higher ratio of PSI supercomplexes to PSI core complexes was  
333 observed in the *ohp2* mutant, suggesting that LHCI antenna in this strain is more tightly  
334 coupled to the PSI core than in the WT. This phenomenon was only partially reverted  
335 in the complemented strain. All PSII complexes formed in the WT were below the  
336 detection limit in the absence of the OHP2 protein. In agreement with the restored  
337 photoautotrophic growth and PSII protein accumulation in the complemented strains,  
338 all PSII related complexes were present in the analyzed *ohp2:OHP2-HA* strain.  
339 Interestingly, different from *Arabidopsis*, where AtOHP2 was detected as a distinct  
340 signal at the size of a ~150 kDa PSII RC-like complex (Li et al., 2019), the HA-tagged  
341 OHP2 protein in our complemented strain was mainly detected in the low molecular  
342 range, reaching up as a weak smear to the first monomeric CP43-less PSII RC47  
343 subcomplex (Figure 3D). This suggests that in *Chlamydomonas* the putative OHC  
344 does not withstand the solubilization conditions used for BN-PAGE. Accordingly, we  
345 were unable to pull-down OHP2 interactants using the HA-epitope as a bait.

## 346 **Ribosome profiling reveals that D1 is still translated in the absence of OHP2**

347 Based on the molecular characterization of the *ohp2* mutant, the major effect of the  
348 loss of OHP2 in *Chlamydomonas* seems to be the missing accumulation of the PSII  
349 reaction center protein D1, in line with observations in other organisms. However, our  
350 <sup>14</sup>C pulse labelling experiments (Figure 3C) can be interpreted either as an absence of  
351 translation of the *psbA* mRNA or as a very rapid or even co-translational degradation  
352 of newly synthesized D1. To address this question and obtain a quantitative image of  
353 chloroplast translation in the *ohp2* mutant, we employed a previously established  
354 targeted chloroplast ribosome profiling approach (Trösch et al., 2018). Ribosome  
355 footprints (FP) from WT and mutant were extracted and analyzed by hybridization of  
356 highly tiled microarrays covering all open reading frames of the chloroplast genome. In  
357 parallel, total RNA was isolated, fragmented and detected by the same approach.  
358 Direct correlation between the three biological replicates showed a high reproducibility  
359 both for determining RNA accumulation ( $r > 0.96$ ) and translation output ( $r > 0.93$ )  
360 (Supplemental Figure S6A, Supplemental Dataset 1).

361 General gene expression defects were examined by averaging all probe intensities  
362 for mRNA or ribosome FPs that covered individual chloroplast CDS, respectively.  
363 Direct plotting of mRNA intensities showed highly comparable RNA abundances  
364 between the WT and the mutant ( $r = 0.98$ , Figure 4A). This suggests that the lack of  
365 OHP2 is not causing any obvious transcription or RNA stabilization defects. In contrast,  
366 the translational output, as determined by relative FP abundance for each ORF,  
367 revealed clear differences between mutant and WT cells and hence, reduced  
368 correlation ( $r = 0.9$ , Figure 4A).

369 Unexpectedly, averaged FP intensities of *psbA* showed only a mild reduction in the  
370 mutant (Supplemental Dataset 1). FP abundance was reduced by 1.6-fold ( $\pm 0.16$ ),  
371 which combined with a slight reduction in mRNA abundance, led to a practically  
372 unchanged translation efficiency (Figure 4B). For a more detailed view, we also plotted  
373 ribosome occupancy ( $\log_2$  of the *ohp2*/WT ratio) over the *psbA* ORF (Figure 4C, left  
374 panel). A more pronounced reduction (up to 4-fold) in ribosome occupancy in *ohp2* can  
375 be observed over the first half of the ORF (~160 codons, up to the 3<sup>rd</sup> transmembrane  
376 segment of nascent D1). In the second half, only mild reduction or even a slight  
377 increase in ribosome occupancy could be detected. For other PSII subunits, the most  
378 pronounced effect was the reduced translation of CP47 (PsbB), PsbH, and PsbT, with  
379 up to 3-fold lower translation output for *psbB* (Figure 4B and Supplemental Figure

380 S6B). This observation agrees with the hierarchical CES cascade contributing to the  
381 biogenesis of PSII in *Chlamydomonas*, where the presence of D2 is required for high-  
382 level translation of D1, which in turn is a prerequisite for efficient translation of the core  
383 antenna protein CP47 (Figure 4D; reviewed in Choquet and Wollman, 2009). When  
384 assembly of D1 and CP47 subunits is compromised, the unassembled proteins repress  
385 the translation initiation of their encoding mRNA (Minai et al., 2006). Also, in  
386 accordance with previous investigations of PSII mutants (de Vitry et al., 1989), we  
387 observed no effect on translation of the second inner antenna protein CP43, encoded  
388 by the *psbC* mRNA, confirming that its rate of synthesis is not dependent on the  
389 assembly with other PSII subunits.

390 Interestingly, also PsbH, which only recently has been hypothesized to be part of  
391 the CES cascade downstream of D1 (Trösch et al., 2018), exhibited a clearly reduced  
392 translation output in *ohp2*, thus further supporting its role as a CES subunit (Figures  
393 4A, 4B, 4D). A comparable effect was seen for the translation of *psbT*. Remarkably, the  
394 three affected genes are encoded in the *psbB-T-H* operon. To obtain a more detailed  
395 view on the translation of this polycistronic transcript, we further looked at relative  
396 ribosome FP accumulation along this *psbB-T-H* operon, calculated as mean log<sub>2</sub> ratios  
397 to directly compare mRNA abundance and ribosome occupancy between mutant and  
398 WT. While the RNA of the operon was only marginally reduced, a clear and relatively  
399 even reduction of ribosome occupancy was seen for all ORFs (Figure 4C, right panel).  
400 It is noteworthy that also *psbN*, which is positioned between *psbT* and *psbH* in  
401 antisense orientation, exhibits a reduced translational efficiency while the mRNA levels  
402 appeared slightly increased. PsbN acts as an assembly factor for PSII but is not part  
403 of the final complex (Knoppová et al., 2022).

404 Complementary dynamics were observed for the synthesis of D2, which showed  
405 increased ribosome FP intensities over the entire ORF in *ohp2* while maintaining  
406 constant mRNA abundance (Figures 4A, 4B, Supplemental Figure S7). As D2 is the  
407 dominant CES subunit (Figure 4D), this likely points to a compensatory effect caused  
408 by the lack of functional PSII. Similarly, other translational alterations in *ohp2*, like the  
409 reduction of ribosome FPs for the large Rubisco subunit RbcL or an upregulation of all  
410 chloroplast-encoded ATP synthase subunits (Supplemental Figure S6), may rather be  
411 secondary effects caused by a diminished photosynthetic electron flow and/or energy  
412 limitation.

### 413 **Proteomic analysis reveals the presence of HCF244 in the *ohp2* mutant**

414 For a quantitative and comprehensive view on the proteome composition in the *ohp2*  
415 mutant, proteomic shot-gun analysis was conducted on whole cell fractions (Figure 5A,  
416 Supplemental Dataset 2). The analysis additionally included an *ohp2* strain in which  
417 the PSII phenotype was genetically suppressed. Such photoautotrophic suppressor  
418 strains occurred at a high frequency (approximate rate:  $5.5 \times 10^{-6}$ ) under conditions  
419 selecting for photoautotrophic growth. Illumina sequencing and Southern blot analysis  
420 of photoautotrophic *ohp2* derivatives showed that they still contained the *TOC1*  
421 transposon in *OHP2* (Supplemental Figures S2, S8A), indicating suppression of the  
422 PSII phenotype by a second site mutation. The suppressor strains showed partially  
423 restored photosynthetic parameters (Fv/Fm ratios of 0.46-0.64), a re-accumulation of  
424 PSII subunits indicated by restored D2 levels, and the ability to grow  
425 photoautotrophically in liquid medium under strong illumination (Supplemental Figures  
426 S8B-D). We proteomically analyzed suppressor strain *M-Su1* to distinguish the direct  
427 effects of the lack of OHP2 from those linked to PSII loss. In addition, LC-MS analysis  
428 was carried out on membrane fractions of WT, *ohp2* and the suppressed strain (Figure  
429 5B, Supplemental Dataset 3). These two analyses allowed reliable quantification of  
430 3065 and 1127 proteins, respectively, with very high reproducibility between the  
431 biological replicates (Supplemental Figure S9A). The complemented strain  
432 *ohp2:OHP2-HA* was included as a control, and showed that most of the alterations  
433 observed in mutant cells were reverted by complementation (Supplemental Figure  
434 S9B). The most obvious defect observed in the *ohp2* strain, apart from the expected  
435 absence of OHP2 itself, was a severe depletion of the majority of PSII proteins. The  
436 most dramatic effect was on D1 showing a ~200/640-fold reduction in whole cell and  
437 membrane fractions, respectively. However, the detection in the *ohp2* mutant of five  
438 D1 peptides arising from the C-terminal third of the protein supports the notion, based  
439 on the results of the ribosome footprinting, that the *psbA* CDS is translated over its  
440 entire length. For the other large PSII subunits D2 and CP47, the depletion was less  
441 pronounced. The least affected subunit was CP43, in accordance with results obtained  
442 with other PSII mutants (de Vitry et al., 1989). The small molecular weight subunits of  
443 PSII were difficult to assess in these experiments, but we noticed a massive depletion  
444 of PsbH in the mutant (Supplemental Datasets 2, 3), in agreement with the ribosome  
445 profiling results. Other PSII subunits were severely depleted, at least at the membrane  
446 level, including the newly-discovered algae-specific subunit PBAS1 (= PBA1,

447 Putatively Photosystem B Associated 1; Spaniol et al., 2021). Among the extrinsic  
448 luminal Oxygen-Evolution Enhancer (OEE) subunits, no significant change was  
449 observed in whole cells, but OEE1 was depleted in membranes, in line with previous  
450 reports (de Vitry et al., 1989).

451 Of particular interest for us was the presence in the mutant and suppressor strains  
452 of OHP1 and HCF244, the partners of OHP2 in the plant OHC. Probably due to its  
453 small size and hydrophobicity, OHP1 was detected neither in the whole cell extracts  
454 nor in the untargeted analysis of membrane fractions, even in the WT. To further refine  
455 our analysis, the same membrane fractions were also subjected to a targeted  
456 proteomics approach, whereby specific peptides were monitored at high resolution. By  
457 reducing interferences from the MS1 level, this allows a more reliable quantitation. Five  
458 proteins were selected: OHP2, OHP1, HCF244, HCF136 and HCF173, using a total of  
459 25 peptides (Figure 5C and Supplemental Dataset 4). This targeted analysis used two  
460 peptides for OHP1 and showed a low signal in membrane fractions of the *ohp2* mutant  
461 (18% of WT). This suggests that in *Chlamydomonas*, OHP1 is not completely  
462 dependent on OHP2 for its stabilization. The OHP1 signal was not detected in the  
463 suppressed strain.

464 In whole cell extracts, HCF244 accumulated to ~25% of the WT level in *ohp2* and in  
465 the suppressor strain (Supplemental Dataset 2). Like OHP2 accumulation, this was  
466 largely complemented by expression of the *OHP2-HA* transgene. Untargeted analysis  
467 of the membrane fractions revealed the presence of HCF244 in the WT but not in *ohp2*  
468 or the suppressed strain. However, the more sensitive targeted analysis, based on  
469 eight peptides, revealed the presence of HCF244 in the membrane fractions of the  
470 mutant and suppressor strains, with an abundance of 22% and 13% relative to WT  
471 levels, respectively (Supplemental Dataset 4, Figure 5C). Together, these results  
472 indicate, that in contrast to *Arabidopsis*, the presence of OHP2 is not absolutely  
473 necessary for HCF244 accumulation, nor for its association with the membrane.

#### 474 **Proteomic analysis of other PS subunits, assembly factors and chloroplast** 475 **enzymes**

476 We also examined the accumulation of other known PSII biogenesis factors, in search  
477 of secondary effects of the lack of OHP2, or possible mechanisms or consequences of  
478 suppression (Figure 5; Supplemental Datasets 2 - 4). At the whole cell level, the  
479 abundance of HCF173 was not affected in the mutant or suppressor, but targeted

480 analysis of the membrane fractions revealed a ~3-fold higher level in *ohp2*, compared  
481 to the WT or suppressed strains, suggesting a regulatory mechanism. Another known  
482 interactant of the *psbA* mRNA in plants is SRRP1: its abundance was significantly  
483 increased in the *ohp2* mutant cells and restored to normal by complementation. In the  
484 suppressor strain, SRRP1 was barely detectable. Similarly, the *psbA* translation  
485 activator TBA1 (Somanchi et al., 2005) overaccumulated in the mutant.

486 The luminal protein HCF136 is an early interactant of the D2/Cytb<sub>559</sub> pre-complex,  
487 necessary for assembly of pD1 into the RC (Plücken et al., 2002). Its abundance in  
488 whole cells was not significantly altered, but both the untargeted and targeted analyses  
489 of the membrane fractions revealed a significantly higher membrane association in the  
490 suppressor strain (~3-fold when compared to the WT; Figure 5; Supplemental Datasets  
491 2 - 4). This may reveal a more prolonged association of HCF136 with the RC when the  
492 absence of OHP2 slows down the biogenesis process.

493 PSB28 binds to CP47 on the stromal surface and distorts the Q<sub>B</sub>-binding pocket, as  
494 revealed by the structural analysis of a cyanobacterial PSII assembly intermediate also  
495 comprising PSB27, which binds to CP43 and the transmembrane protein PSB34 on  
496 the luminal side (Xiao et al., 2021; Zabret et al., 2021). Here, PSB28 and PSB27  
497 showed interesting contrasted behaviors. At the whole cell level, the accumulation of  
498 PSB28 showed no significant change between strains. However, it was undetectable  
499 in the membrane of the *ohp2* strain or of the suppressor. In contrast, PSB27 was  
500 overaccumulated in *ohp2* cells, with no change in its abundance in membrane  
501 fractions. PSB27 was also somewhat overaccumulated in the suppressor, this time  
502 accompanied by a marked increase in its association with the membrane. We also  
503 noticed an increased accumulation in the *ohp2* mutant of PSB33/TEF5, a factor  
504 involved in the interaction of photosystems and antenna complexes (Fristedt et al.,  
505 2015; Nilsson et al., 2020), suggesting stabilization by prolonged interaction with the  
506 antenna in the absence of RC. The mutant also overaccumulated APE1, initially  
507 described as involved in the adaptation to high light (Walters et al., 2003) and later  
508 found as a specific interactant of *Arabidopsis* OHP1 in pull-down assays (Myouga et  
509 al., 2018) and associated with the RCIIa complex isolated from a *Synechocystis* strain  
510 lacking YCF39/HCF244 (Knoppová et al., 2022). Overaccumulation was also  
511 observed for the homolog of Slr1470, another component of the RCIIa complex  
512 (Knoppová et al., 2022). This suggests that absence of the OHC in *Chlamydomonas*  
513 leads to prolonged association (and stabilization) of those assembly factors whose

514 interaction with the RC is antagonized by that of the OHC. Finally, we observed  
515 increased membrane association of PSB29/THF1 in the mutant, which was partially  
516 reverted in the suppressor strain. THF1 has been shown to be involved in the dynamics  
517 of PSII-LHCII supramolecular organization and to associate with the FtsH protease  
518 (Bečková et al., 2017; Huang et al., 2013; Wang et al., 2004).

519 In contrast to PSII, no significant effect was observed on the accumulation of PSI  
520 RC subunits: the slight overaccumulation of PSAE in whole cells of the mutant was  
521 also found in the suppressed and complemented strains and may therefore be unlinked  
522 to OHP2. More interestingly, however, all the LHCI subunits showed increased  
523 accumulation in the *ohp2* mutant as well as in the suppressor strain, suggesting a direct  
524 effect of the lack of OHP2 (Supplemental Dataset 2). While subunits of the *Cytb<sub>6</sub>f*  
525 complex appeared unaffected in the *ohp2* mutant, we noticed an overaccumulation of  
526 those of the ATP synthase (Figure 5A, left panel). This agrees with the increased  
527 translational output for the ATP synthase ORFs seen in our ribosome profiling analysis  
528 (Supplemental Figure S6B). Overaccumulation of the ATP synthase was fully reversed  
529 in the complemented strain (Supplemental Figure S9B). We note, however, that the  
530 suppressor strain, although photoautotrophic, still showed some overaccumulation of  
531 the ATP synthase in its membrane fraction (Figure 5B). The Rubisco subunits RbcL  
532 and RBCS2, in contrast, appeared to accumulate at lower levels in the mutant in whole  
533 cells fractions (Supplemental Dataset 2). The significance of this observation is  
534 unclear, as their accumulation was not restored to normal in the suppressor strain.

535 Several enzymes of the porphyrin biosynthesis pathway showed changes in their  
536 accumulation or association with the membrane (Supplemental Datasets 2, 3, Figures  
537 5A, 5B). The Chl synthase CHLG, whose cyanobacterial homolog interacts with  
538 HliD/Ycf39 (Chidgey et al., 2014), remained unaltered in the *Chlamydomonas* mutant,  
539 but other Chl synthesis enzymes showed increased abundance in the membrane  
540 fractions. This includes the two subunits of the Mg-chelatase (ChlH, ChlI), the Mg  
541 protoporphyrin IX S-adenosyl methionine O-methyl transferase ChIM, the cyclases  
542 CRD1 and CTH1 and the protochlorophyllide *a* oxidoreductase POR1. Except for  
543 POR1, this increase was not observed at the whole cell level, suggesting that the cell  
544 responds to the absence of the OHC complex by stabilizing the interaction between  
545 these Chl pathway enzymes and the thylakoid membrane. This effect was largely or  
546 completely reversed in the suppressor, pointing to a regulatory mechanism  
547 compensating for the impaired stabilization of D1. In the upstream part of the pathway,

548 shared with heme biosynthesis, several enzymes overaccumulated in whole cells of  
549 the mutant (Figure 5A): the delta-aminolaevulinic acid dehydratase (ALAD1), the  
550 uroporphyrinogen III synthase (UPS1), one of the three uroporphyrinogen III  
551 decarboxylases (UPD1), and the protoporphyrinogen oxidase (PPX1). PPX1  
552 abundance also increased in the membrane fraction. The protein FLUORESCENT  
553 (FLU), which is involved in the regulation of the whole pathway (Falciatore et al., 2005)  
554 and which was severely reduced in a virus-induced *OHP2* gene silenced *Arabidopsis*  
555 line (Hey and Grimm, 2018), was not significantly affected in the *Chlamydomonas*  
556 mutants.

### 557 **The *Arabidopsis* OHP2 complements the *Chlamydomonas ohp2* mutation**

558 The observed phenotypical differences between *ohp2* mutants from higher plants and  
559 *Chlamydomonas*, as well as the divergence of the amino acid sequences (Li et al.,  
560 2019; Myouga et al., 2018) led us to test whether the *Arabidopsis* protein could  
561 complement the *Chlamydomonas* mutant. The N-terminal region of the mature OHP2  
562 proteins is much less conserved than the transmembrane and C-terminal regions and  
563 similarity between the *Chlamydomonas* and *Arabidopsis* proteins starts only at residue  
564 65 of the *Chlamydomonas* OHP2 preprotein with an overall identity of 58% (compare  
565 Figure 2A).

566 We used a construct (pBC1-TP<sub>CrOHP2</sub>-AtOHP2-HA, construct 6 in Supplemental  
567 Figure S3) fusing the predicted cTP of OHP2 from *Chlamydomonas* with a codon-  
568 adapted sequence encoding the mature part of AtOHP2 (Supplemental Data S2). After  
569 transformation of the *Chlamydomonas ohp2* mutant and selection for Pm-resistance,  
570 17 of the 20 clones (85%) showed restoration of photoautotrophy and Fv/Fm values of  
571 0.62 – 0.74 (Supplemental Figure S10A). For all of the immunologically analyzed  
572 clones, a protein with the expected size of ~ 15 kDa was detected using the  $\alpha$ -HA-tag  
573 antibody and D1 proteins accumulated to WT levels (Supplemental Figure S10B).  
574 Taken together these data indicate WT level PSII accumulation and successful  
575 complementation of the *Chlamydomonas* mutant with the *Arabidopsis* protein.

576 **Slowing down D1 degradation in the *ohp2* mutant can partially restore light-**  
577 **sensitive PSII activity, but not photoautotrophy**

578 The repair of photodamaged PSII involves the selective proteolytic degradation and  
579 replacement of the damaged D1 polypeptide with a newly synthesized one (reviewed  
580 in Kato and Sakamoto, 2009; Nixon et al., 2010). The thylakoid protease FtsH plays a  
581 major role in the D1 degradation process. In *Chlamydomonas*, FtsH is composed of  
582 two subunits FtsH1 and FtsH2 and the *ftsh1-1* mutation, changing a conserved arginine  
583 residue, strongly impairs oligomerization and proteolytic activity (Malnoë et al., 2014).  
584 This mutation restores photosynthesis in a mutant lacking heme *c*<sub>1</sub> of *Cytb<sub>6</sub>f* (Malnoë et  
585 al., 2011). In addition, it impairs degradation of D1 in high light, slowing down repair of  
586 PSII during photoinhibition and leading to enhanced light-sensitivity in mixotrophic and  
587 photoautotrophic conditions (Malnoë et al., 2014). To address the question whether the  
588 FtsH protease is involved in the immediate post-translational degradation of D1 in the  
589 *ohp2* mutant, we crossed the original *10.1a* strain with a strain carrying the *ftsh1-1*  
590 mutation. In the progeny, the phenotypes associated with the WT, *ohp2*, *ohp2 ftsh1-1*  
591 and *ftsh-1* genotypes showed distinct fluorescence and growth patterns (Figure 6).  
592 When strains were analyzed by spot tests on agar plates, the *ftsh1-1* strains showed  
593 retarded growth and decreased Fv/Fm with increasing light intensity under  
594 photoautotrophic as well as under mixotrophic conditions (Figure 6A). This is typical of  
595 the *ftsh1-1* light-sensitive phenotype (Malnoë et al., 2014). As expected, the *ohp2*  
596 progeny did not grow photoautotrophically, except for occasional suppressor clones.  
597 Under mixotrophic conditions, they showed no deleterious effect of high light for  
598 growth, as is typical of PSII mutants, and had practically no Fv/Fm (Figures 6A, 6B).  
599 The *ohp2 ftsh1-1* strains showed interesting phenotypes. Like the *ohp2* parent, they  
600 were unable to grow on minimal medium at any light intensity tested (Figure 6A).  
601 However, their fluorescence induction curves were clearly indicative of the presence of  
602 a small amount of PSII, especially in dark-grown cells (Figure 6C). On TAP, we did not  
603 observe any stimulatory effect of low light on growth (Figure 6A). These results suggest  
604 that FtsH is involved in the degradation of the PSII units, possibly abnormal, produced  
605 in *ohp2* mutants. However, the activity of the low amount of PSII stabilized by  
606 attenuation of FtsH is either insufficient or too light-sensitive to allow photoautotrophy.  
607 Note that the unknown suppressor locus was not genetically linked to *FTSH1* in  
608 crosses, and the suppressor strains showed no sequence change in any of the *FTSH*  
609 genes, nor in any other known chloroplast protease gene.

## 610 DISCUSSION

### 611 In *Chlamydomonas*, OHP2 is not necessary for *psbA* translation.

612 The molecular analysis of an *ohp2* knockout mutant from *Chlamydomonas* revealed a  
613 major defect in PSII biogenesis, as indicated by its inability to grow  
614 photoautotrophically, the complete loss of PSII activity and the absence of the major  
615 PSII subunits, in particular D1 (Figures 1, 3). No effect was observed on the PSI RC  
616 (Figures 1C, 3A, 3D, 5), as reported for cyanobacterial HLIPs (Komenda and Sobotka,  
617 2016), but at slight variance with land plants, where reductions in PSI subunits and  
618 antenna proteins have been reported (Beck et al., 2017; Li et al., 2019; Myouga et al.,  
619 2018). In *Chlamydomonas*, we observed an increased accumulation of LHCl antenna  
620 proteins in the mutant (Supplemental Dataset 2), and a higher abundance or stability  
621 of the PSI-LHCl complex in non-denaturing PAGE (Figure 3D). Whether this is related  
622 to the observed decrease in Chl abundance and changes in expression of Chl pathway  
623 genes remains to be investigated. In the *Arabidopsis ohp1* and *ohp2* mutants, the level  
624 of PSI RC decreases more than that of LHCl, but this appears to be true for other PSII  
625 mutants as well (Li et al., 2019)..

626 In other organisms, a dimer of One-Helix Proteins (HliC/D in cyanobacteria,  
627 OHP1/2 in land plants) has been found to mediate the early association of Chl *a* to the  
628 nascent D1 polypeptide. In *Chlamydomonas*, the expression patterns of OHP2, OHP1  
629 and HCF244 in a variety of conditions are highly similar (Supplemental Figure S11)  
630 and on the Phytozome website (<https://phytozome-next.jgi.doe.gov/>) the latter two  
631 appear in each other's lists of best correlated genes, pointing to the existence of a  
632 similar complex in the alga. We will therefore assume that the *Chlamydomonas ohp2*  
633 mutant lacks the OHC, which prevents normal cofactor insertion into D1, resulting in  
634 destabilization of PSII. Note that, although Chl is usually mentioned as the essential  
635 cofactor missing in *ohp2*, Phe *a* is also a possible substrate for the action of the OHC.  
636 The main result of our study, compared with those in higher plants, is that in  
637 *Chlamydomonas* the absence of OHP2 only marginally affects translation of the *psbA*  
638 mRNA. While our <sup>14</sup>C pulse labelling experiments do point to the primary phenotype  
639 being a defect in D1 production, they can be interpreted either as a deficiency in *psbA*  
640 translation, or as a very fast degradation of the newly synthesized D1 polypeptide  
641 (Figure 3C). Our ribosome profiling experiments strongly support the latter hypothesis,  
642 as they show an almost normal abundance of FPs over the CDS (1.6-fold reduction of

643 the signal). Continued translation of the *psbA* mRNA is observed over its whole length  
644 (Figure 4C), further supported by the fact that the proteomic analysis identified several  
645 peptides in *ohp2* that lie in the C-terminal part of D1.

646 Note that the two techniques we have used have their limitations. Usually, the  
647 absence of a signal in  $^{14}\text{C}$ -pulse labeling in *Chlamydomonas* is interpreted as loss of  
648 translation (e.g. de Vitry et al., 1989). But the *ohp2* mutant must be analyzed differently,  
649 as the mutation is supposed to affect a co-translational or early post-translational step.  
650 In addition, the sensitivity of this approach is limited by the broadness of the D1 band  
651 (D originally stands for "diffuse") and the unavoidable presence of background  
652 radioactivity. Ribosome footprinting is more sensitive, even though it also is an indirect  
653 proxy for translation initiation. The abundance of RFs over a transcript may also be  
654 affected by the dynamics of translation, for example the stalling of ribosomes will  
655 increase the probability to generate RFs at this position. It is interesting to note that the  
656 reduction in ribosome occupancy in the *ohp2* mutant is more pronounced over the first  
657 half of the mRNA. Chl attachment to D1 has been proposed in barley to be associated  
658 with ribosome pausing (Kim et al., 1994a; Kim et al., 1991, 1994b) even though RF  
659 experiments in maize have failed to identify such Chl supply dependent pauses  
660 (Zoschke and Barkan, 2015; Zoschke et al., 2017). If interaction of the nascent chain  
661 with the Chl attachment machinery slows down elongation, then its absence in *ohp2*  
662 may reduce the density of RFs. Conservatively, we propose that in the absence of  
663 OHP2, *psbA* translation is maintained at a significant rate, over the whole CDS.  
664 However, the degradation of apo-D1 is so fast that full-length D1 remains below  
665 detection in pulse labeling experiments, as it is in Western blots (Figures 3A, 3C). This  
666 statement is backed by the observation that mutations in the FtsH protease and at an  
667 unknown suppressor locus can partially or fully restore PSII accumulation, which would  
668 be difficult to achieve if OHP2 was required for translation.

669 The situation is different in land plants, where OHP1, OHP2 and HCF244 are  
670 required for the recruitment of ribosomes to the *psbA* mRNA (Chotewutmontri et al.,  
671 2020). The abundance of RFs in the *Arabidopsis ohp2-1* knockout mutant was ~12  
672 times lower than in the WT, a figure probably underestimated by normalization since  
673 the *psbA* CDS itself contributes to a large fraction of the RFs in the WT. Chotewutmontri  
674 et al. convincingly explained the discrepancy with a previous study using polysome  
675 analysis (Li et al., 2019), by the difficulty in analyzing *psbA* polysomes profiles due to  
676 the high abundance of the mRNA. They found that the *Arabidopsis ohp1-1* and maize

677 *hcf244-1/-3* null mutants also fail to translate D1 (6-fold and 10-fold reduction,  
678 respectively). No change was observed in the pattern of RFs over *psbA* in any of the  
679 mutants, in line with a defect in translation initiation rather than elongation. All this was  
680 in accordance with the role of HCF244 as an essential translation initiation factor for  
681 *psbA* and its total absence in mutants lacking OHP1 or OHP2 (Li et al., 2019).  
682 Chotewutmontri and coworkers recently proposed a model for regulation of *psbA*  
683 translation by light, whereby the D1-bound OHC inhibits the ability of the stroma-  
684 exposed components (HCF244 and/or OHP2's stromal tail) to initiate *psbA* translation  
685 (Chotewutmontri and Barkan, 2018; 2020; Chotewutmontri et al., 2020). It must be  
686 noted that Chotewutmontri et al. (2020) quantified RFs by Illumina read counting, while  
687 we have used hybridization to oligonucleotide probes. But both methods are fully  
688 validated, and the results are so different that we cannot invoke a technical bias.  
689 Therefore, we conclude that *Chlamydomonas* lacks the OHP2-dependent control  
690 mechanism for *psbA* translation described in land plants.

691 We used proteomics analyses to explain this apparent discrepancy in the context  
692 of the high conservation of the proteins between algae and land plants (as exemplified  
693 by our observation that *Arabidopsis* OHP2 can complement the *Chlamydomonas*  
694 mutant; Supplemental Figure S10). Quantification of OHP1 was difficult because of its  
695 small size, but the results of our targeted analysis point to the presence of traces of  
696 this protein in the *ohp2* mutant (Figure 5C). This, however, should have no effect on  
697 the insertion of cofactors: an *ohp2* mutation in *Arabidopsis* is not complemented by  
698 overexpression of OHP1 (Beck et al., 2017) and OHP1 alone does not bind pigments  
699 *in vitro* (Hey and Grimm, 2020). Studies in land plants have found that while some  
700 OHP2 can accumulate in *ohp1* null mutants, OHP1 is usually undetectable in *ohp2*  
701 mutants (Beck et al., 2017; Li et al., 2019). Some variability is seen in these studies,  
702 possibly due to the fact that the T-DNA insertion in *OHP2* lies within an intron (Beck et  
703 al., 2017; Li et al., 2019; Myouga et al., 2018). Similarly, the level of HCF244 reported  
704 in *Arabidopsis ohp2* mutants varied from nil to very low (Li et al., 2019) and the  
705 accumulation of HCF244, even when overexpressed, was found to be limited by that  
706 of OHP2 (Hey and Grimm, 2020; Li et al., 2019). Here, our untargeted and targeted  
707 LC-MS/MS analyses concur to demonstrate not only that HCF244 accumulates to  
708 significant levels in the *Chlamydomonas ohp2* mutant (~25% of WT), but that it can  
709 interact with the membrane (Figure 5C). This might be explained by the remaining  
710 traces of OHP1 acting as an anchor for HCF244 and stabilizing it, or to an intrinsically

711 higher stability of *Chlamydomonas* HCF244 when not assembled. We propose that  
712 these remaining 25% HCF244 in the *Chlamydomonas ohp2* mutant are responsible for  
713 the maintenance of *psbA* translation at an appreciable rate. This does not exclude that  
714 an autoregulatory circuit exists in WT *Chlamydomonas* similar to that described by  
715 Chotewutmontri and Barkan (2020) in land plants. In this case, the absence of  
716 OHP1/OHP2 in the mutant may prevent the remaining HCF244 from sensing the  
717 presence of unassembled D1, leading to enhanced activation of *psbA* translation. In  
718 any event, the strong coupling observed in plants between Chl insertion and *psbA*  
719 translation initiation appears largely broken in *Chlamydomonas*. This coupling may be  
720 ancient in oxygenic photosynthesis, as judged from the existence of a similar complex  
721 in cyanobacteria. It might be interesting to examine whether the cyanobacterial  
722 complex also controls *psbA* translation. A major difference between *Arabidopsis* and  
723 *Chlamydomonas* is that Chl production depends entirely on light in the former, while  
724 the latter can produce it in the dark. Thus, the availability of Chl for integration into PSII  
725 could be used by Angiosperms to sense light, while algae (and cyanobacteria) would  
726 have to use other clues. In addition, the translation of *psbA* appears to mobilize a larger  
727 fraction of ribosomes in the plant than in the alga, as judged from RF and pulse-labeling  
728 experiments. Arguably, the alga can partly dispense of a regulatory circuit aimed at  
729 ensuring that *psbA* is translated only in the light. Detailed studies in other systems,  
730 such as land plants capable of producing Chl in the dark, are necessary to decide how  
731 strong the coupling needs to be to prevent deleterious effects of uncoordinated D1  
732 synthesis and assembly, in particular during high light stress when D1 needs to be  
733 repaired at high rate.

#### 734 **The absence of OHP2 affects translation of several chloroplast transcripts** 735 **encoding PSII subunits**

736 Interestingly, the *ohp2* mutant revealed large changes in the ribosome profiling pattern  
737 of PSII transcripts other than *psbA*. Most striking was the reduced translational output  
738 for the CES subunit CP47, encoded by *psbB* (Figure 4). This observation is in line with  
739 previous knowledge on translational regulation by assembly (the CES process) in  
740 *Chlamydomonas* (reviewed in Choquet and Wollman, 2009; Minai et al., 2006).  
741 Genetic studies indeed have demonstrated that in the absence of D1, the unassembled  
742 CP47 subunit feeds back onto the translation initiation of its own mRNA. Here,  
743 assuming that no assembly of CP47 and D1 can occur in the *ohp2* mutant, we can

744 quantify this effect. With a 3-fold reduction of RFs over the *psbB* mRNA in the mutant  
745 ( $\log_2 \text{ohp2/WT} = -1.58$ ), the CES effect appears to be rather dramatic (Figure 4B).  
746 Remarkably, the translation of *psbH* is also reduced in the mutant, in line with a  
747 complete lack of detection of PsbH peptides by proteomic analyses (Supplemental  
748 Datasets 2, 3). This supports the identification of PsbH as a component of the PSII  
749 CES cascade, as proposed recently by Trösch et al. (2018; Figure 4D). Early reports  
750 in *Chlamydomonas* describe a role of PsbH in assembly and/or stabilization of PSII  
751 (O'Connor et al., 1998; Summer et al., 1997) but the protein's function has not been  
752 studied in detail. In the cyanobacterium *Synechocystis* sp. PCC 6803, PsbH is  
753 associated with CP47 and facilitates D1 processing and incorporation into PSII  
754 (Komenda et al., 2005). Assuming a similar position of PsbH in the assembly pathway  
755 in *Chlamydomonas*, its CES-mediated downregulation in the absence of D1 is  
756 consistent with that of CP47. The mechanism remains to be determined: is this  
757 regulation dependent upon the accumulation of unassembled PsbH, unassembled  
758 CP47 or the CP47/PsbH sub-complex? Genetic studies using mutants lacking the  
759 CP47 or PsbH proteins will be necessary to address this question. In their study,  
760 Chotewutmontri et al. (2020) pointed to an increased, rather than decreased,  
761 translation of *psbB* in their *ohp1*, *ohp2*, *hcf244* and even in *hcf173* mutants. The  
762 significance of these opposite behaviors between *Chlamydomonas* and land plants  
763 remains to be explored. Noticeably, *psbH*, compared to other PSII subunits, also  
764 appeared as slightly increased in the RF abundancy plots of the *ohp2* and *hcf244*  
765 mutants in Chotewutmontri et al. (2020).

766 The comparable effect that we observed on *psbT* translation in our experiments  
767 (Figure 4) could be construed as evidence that this small polypeptide also belongs to  
768 the PSII CES cascade. To our knowledge, no study has examined the possibility that  
769 PsbT also is subject to translational regulation. To complicate the matter, RF-analysis  
770 of the PSII mutant *nac2-26*, which lacks the D2 protein, rather showed a slight increase  
771 in translation of *psbT* (Trösch et al., 2018). Because D1 itself (and hence all the genes  
772 it controls via CES) needs D2 for full translation, the putative role of PsbT in the CES  
773 cascade remains ambiguous. It should be noted that all three cistrons belong to the  
774 conserved *psbB-T-H* operon and that the tetratricopeptide repeat protein MBB1 is  
775 necessary for stabilization of the 5' end of the *psbB-psbT* mRNA as well as the  
776 processing and translation of the co-transcribed *psbH* mRNA (Loizeau et al., 2014;  
777 Vaistij et al., 2000). No monocistronic transcript has been described for PsbT, so its

778 CDS can only be translated from the *psbB-psbT* dicistronic mRNA (Cavaiuolo et al.,  
779 2017; Monod et al., 1992). It is striking that the three ORFs that depend on MBB1 for  
780 transcript stabilization or translation showed a strong reduction of ribosome occupancy  
781 in the *ohp2* mutant (Figure 4C). This suggests a role for MBB1 in the CES-regulation:  
782 it may work as a translation factor or recruit such a factor. Even if MBB1 does not bind  
783 directly upstream of *psbT*, the increased recruitment of the *psbB-psbT* transcript to  
784 ribosomes may enhance translation initiation on *psbT*. Unfortunately, MBB1 could not  
785 be quantitated in our proteomic analyses. The decrease in RF level observed over the  
786 *psbN* ORF which is positioned between *psbT* and *psbH* but in antisense orientation  
787 could also indirectly involve MBB1, if recruitment of the precursor *psbB-T-H* transcript  
788 to the polysomes affects the stability/translatability of the antisense *psbN* mRNA.  
789 Coupling between chloroplast mRNA stabilization and translation factors has been  
790 reported in *Chlamydomonas*, for the MCA1/TCA1 couple. Changes in the abundance  
791 of MCA1/TCA1 rapidly regulate *petA* mRNA accumulation and translation to meet the  
792 cell's demand for synthesis of the CES-regulated Cyt $f$  subunit of the Cyt $b_6f$  complex  
793 (Loiselay et al., 2008; Raynaud et al., 2007). Similarly, the abundance of TAA1,  
794 involved in stabilization and translation of the *psaA* mRNA, has been reported to be  
795 involved in the downregulation of PSI subunits upon iron deficiency (Lefebvre-  
796 Legendre et al., 2015).

797 An unexpected result for the *ohp2* mutant in the RF experiment was the 2.8-fold  
798 increase in RFs over *psbD*, encoding the main RC subunit D2 (Figures 4A, 4B;  
799 Supplemental Figures S6B, S7; Supplemental Dataset 1). Because of the high  
800 variability in the efficiency of  $^{14}\text{C}$  incorporation between strains and of the gel  
801 background, we could not verify this by the pulse-labeling experiments (Figure 3C).  
802 However, as all other D1 mutants studied thus far, like FuD7 or F35, completely lacked  
803 D1 translation (Bennoun et al., 1986; Girard-Bascou et al., 1992; Yohn et al., 1996), a  
804 stimulatory effect on D2 translation by the presence of unassembled D1 (as possible  
805 in *ohp2*) could not have been observed. We note that translation of *psbD* was not  
806 enhanced in any of the mutants studied by Chotewutmontri et al. (2020).

## 807 **OHP2 is not completely essential for PSII biogenesis**

808 Another intriguing characteristic of the *Chlamydomonas ohp2* mutant is the high  
809 frequency at which the non-photoautotrophic phenotype can be suppressed. A mere  
810 plating on mineral medium, without any mutagenic treatment, was enough to generate

811 dozens of extragenic suppressors. In spite of all our efforts, we were unable to identify  
812 a plausible causative variant, even though genetic analysis of three independent  
813 suppressors indicated that the suppressor mutation(s) were tightly linked, but unlinked  
814 to the *OHP2* or *FTSH1* loci. Furthermore, combination of *ohp2* with the *ftsH1-1*  
815 mutation that severely affects activity of the FtsH protease (Malnoë et al., 2014)  
816 allowed partial recovery of stable PSII charge separation, as revealed by  
817 measurements of variable fluorescence and ECS (Figure 6C). This indicates that  
818 whatever the exact action of the OHC on D1, it is not completely essential for  
819 biogenesis of a functional PSII. A by-pass reaction must be possible, that leads to  
820 partial restoration of PSII biogenesis. This pathway would be activated by the  
821 suppressor mutation, but it probably already exists in the *ohp2* mutant: by slowing  
822 down D1 degradation, the *ftsH1-1* mutation may allow a normally unstable form of D1  
823 to proceed through this alternative pathway. Such a pathway also operates in  
824 cyanobacteria, where deletion of all HLIP genes does not prevent PSII biogenesis (Xu  
825 et al., 2004).

826 It is probable that the PSII produced by the OHP2-independent pathway is not fully  
827 functional, because the suppressed strains showed some light-sensitivity, and the  
828 *ohp2 ftsH1-1* double mutant was not photoautotrophic (Figure 6A). Alternatively, the  
829 activity of this alternative pathway may be too low to allow efficient repair in high light.  
830 The *ftsH1-1* mutation by itself renders the cell highly sensitive to photoinhibition.  
831 Detailed studies comparing PSII purified from these strains may be necessary to  
832 answer this question.

### 833 **Consequences of the absence of OHP2 on other steps of the PSII assembly** 834 **pathway**

835 Based on the above, it was interesting to characterize the assembly intermediates in  
836 the mutant and suppressor. We therefore used our proteomics data to monitor the  
837 accumulation of the many PSII assembly factors described in the literature. Not only  
838 did they represent candidate genes for the suppressor mutation, but changes in their  
839 accumulation in the mutant and suppressor could reveal regulatory mechanisms at  
840 play in *ohp2* and the suppressor. Some of the changes described below could be  
841 brought about by regulation of gene expression, others by (de)stabilization of the  
842 proteins. By comparing whole cells and membranes, we may also detect changes in  
843 the association of these factors with PSII assembly intermediates, as recently

844 demonstrated by Spaniol et al. (2021) for the *lpa2* mutant. Figure 7 proposes a working  
845 model for the role of the OHC and associated proteins in D1 synthesis, combining  
846 previous reports from other organisms with data presented here for *Chlamydomonas*.  
847 In the WT (Figure 7, left panel), HCF173 binds the 5'-UTR of *psbA* mRNA to promote  
848 its translation (Link et al., 2012; Schult et al., 2007; Williams-Carrier et al., 2019), which  
849 is then likely triggered by HCF244. Nascent pD1 protein is co-translationally inserted  
850 into the thylakoid membrane where it is stabilized by membrane-associated luminal  
851 HCF136. The OHC is proposed to further stabilize pD1 by the insertion of Chl/Phe/Car,  
852 allowing its assembly in the RC complex and formation of functional PSII core, dimer  
853 and finally supercomplexes. A striking result of our targeted proteomics analysis was  
854 the two-fold overaccumulation of HCF173 in the membranes of the *ohp2* mutant  
855 (Figure 5C, Supplemental Dataset 4). It was not significantly overexpressed in the  
856 mutant at the whole cell level, so its increased abundance in membranes probably  
857 indicates that a larger fraction of the *psbA* mRNA is undergoing translation and thus  
858 found on membrane-associated polysomes. This effect is no longer seen in the  
859 suppressor, suggesting that it is not a direct consequence of the absence of the OHC,  
860 but rather a compensatory mechanism caused by the PSII deficiency. As proposed  
861 above, unassembled D1 can no longer regulate the activity of HCF244 in the mutant  
862 because the OHC cannot form (Chotewutmontri et al., 2020; Link et al., 2012). This is  
863 expected to lead to enhanced translation initiation on the *psbA* mRNA driven by the  
864 residual but fully active HCF244 (Figure 7, middle panel), unless a counteracting  
865 mechanism signals that PSII biogenesis proceeds, as in the suppressor. SRRP1,  
866 another known interactant of HCF173 and of the *psbA* mRNA, is believed to act as a  
867 repressor of translation (Watkins et al., 2020). It could not be quantitated in the  
868 membranes, but interestingly, while abundant in cells of the mutant and WT it was  
869 undetectable in the suppressor, even though the gene appeared unaltered.

870 HCF136 is a conserved luminal protein, known in cyanobacteria as Ycf48. Its role  
871 seems mostly to promote the association of the pD1/PsbI subcomplex with a distinct  
872 precomplex consisting of D2 and the heterodimeric Cytb<sub>559</sub> (formed by the PsbE and  
873 PsbF subunits), to form the RC (reviewed in Nickelsen and Rengstl, 2013; Zhang et  
874 al., 1999). It is required for the assembly of early PSII intermediates, possibly with the  
875 concurrent incorporation of Chl and has been found to bind pD1 (but not mature D1)  
876 in a split-ubiquitin assay (Hey and Grimm, 2020; Knoppová et al., 2014; Komenda et  
877 al., 2008; Li et al., 2019; Meurer et al., 1998; Myouga et al., 2018; Plücker et al., 2002).

878 It was therefore striking to observe that association of HCF136 with the membrane (not  
879 its overall abundance) was markedly increased in the suppressor, compared to the WT  
880 or mutant (Figure 5; Supplemental Datasets 2- 4). While genome sequencing rules out  
881 *HCF136* as being the suppressor locus, its increased abundance in the membranes of  
882 the suppressor may indicate the stabilization of a complex comprising pD1, allowing it  
883 to associate with Chl via the alternative pathway proposed above (Figure 7, right  
884 panel).

885 Assembly factors acting downstream of the formation of the RC can also be  
886 impacted by the absence of OHP2. For example, the mutant did not show any  
887 association of PSB28 with the membrane, consistent with the fact that this protein  
888 binds to the RC47 complex (Zabret et al., 2021), that cannot form in the mutant. This  
889 effect was reverted in the suppressor, in line with a restoration of the assembly  
890 pathway. PSB27 binds to CP43 on the luminal surface of the PSII monomer and dimer  
891 and contributes to the formation of the OEC (Avramov et al., 2020; Huang et al., 2021;  
892 Zabret et al., 2021). In the *ohp2* mutant, it overaccumulates at the whole-cell level,  
893 presumably through enhanced expression of the gene, but not in the membranes,  
894 probably because of the absence of the RC. In contrast, the suppressor showed  
895 enhanced binding of PSB27 to the membrane, suggesting that PSII formed in the  
896 absence of the OHC requires prolonged association with PSB27, for example because  
897 pD1 maturation, binding of the OEEs and/or photoactivation are delayed. Another  
898 interesting observation that links back to the role of FtsH is the increased association  
899 of PSB29/THF1 with the membrane in *ohp2*. The *Arabidopsis thf1* mutant is variegated  
900 (Wang et al., 2004), a phenotype suppressed by mutations affecting chloroplast gene  
901 expression (Hu et al., 2015; Zhang et al., 2009), just like that of the *ftsH* mutants. THF1  
902 is necessary for normal accumulation and function of FtsH in land plants and  
903 cyanobacteria (Bečková et al., 2017; Huang et al., 2013; Zhan et al., 2016). Its 3D  
904 structure has been solved (Bečková et al., 2017), but it is unclear yet how it stabilizes  
905 or activates the protease. In our proteomic analysis, PSB29/THF1 was not over  
906 accumulated in any of the strains. However, its increased association with the  
907 membrane in the mutant may reflect the increased activity of FtsH involved in  
908 degrading ill-conformed D1.

## 909 **Other effects of the mutation in *OHP2* on chloroplast biogenesis**

910 Our RF experiments showed an increased translation of all the chloroplast-encoded  
911 subunits of the ATP synthase in the *ohp2* mutant (Supplemental Figure S6,  
912 Supplemental Dataset 1). Similarly, our proteomics experiments revealed an increased  
913 abundance of all subunits of the complex (including the nucleus-encoded ATPC/D/G),  
914 with a striking quantitative coherence in membrane samples (Figures 5A, 5B;  
915 Supplemental Datasets 2, 3). Increased levels of chloroplast ATP synthase subunits  
916 were recently also observed in complexome profiling of the PSII-deficient *lpa2* mutant  
917 from *Chlamydomonas* (Spaniol et al., 2021). Overaccumulation of the ATP synthase  
918 could be involved in maintaining energy balance in the chloroplast: in the absence of  
919 PSII, linear electron flow is abolished, but cyclic electron flow remains possible  
920 between PSI and the *Cytb<sub>6</sub>f* complex and generates a proton gradient that can be used  
921 for ATP synthesis.

922 Our proteomics analysis showed increased accumulation of LHCl subunits in the  
923 *ohp2* mutant, as well as in the suppressor strain. This suggests that the lack of OHP2  
924 directly affects the biogenesis of PSI. Interestingly, BN-PAGE revealed a stronger  
925 association of the PSI core and antenna subcomplexes in the mutant. This may be  
926 related to the observation that the PSAN subunit, although not affected in whole cells,  
927 was undetectable in the membranes of the mutant. In maize, PSAN lies at the interface  
928 between the RC and LHCl (Pan et al., 2018), and this subunit was not found in the  
929 structure of the *Chlamydomonas* PSI-LHCl complex (Suga et al., 2019), suggesting  
930 that it modulates the association between the antenna and the RC in PSI.

931 Proteomics also revealed a general stimulation of the heme/Chl biosynthesis  
932 pathway in the *ohp2* mutant (Figures 5A, 5B; Supplemental Datasets 2, 3). To our  
933 knowledge, this is not a general feature of PSII deficient mutants and may be  
934 specifically linked to the main purported function of OHP2/OHP1, i.e. integration of Chl  
935 into a nascent polypeptide. Accumulation of the protochlorophyllide *a* oxidoreductase  
936 POR1 was increased both at the cell and membrane level, while for other enzymes like  
937 the Mg-chelatase, the Mg-protoporphyrin-IX methyltransferase CHLM and the cyclase  
938 CTH1, the observed effect was an increased association with the membrane. The OHC  
939 complex has been linked before with Chl synthesis, albeit in different manners. In  
940 cyanobacteria, the Chl synthase ChlG (found unaffected here in the *Chlamydomonas*  
941 *ohp2* mutant) co-immunoprecipitates with the homologs of OHP2 and HCF244  
942 (Chidgey et al., 2014). In the *Arabidopsis ohp2* mutant, Hey and Grimm (2018) reported

943 a decreased accumulation for most of the immunologically analyzed Chl biosynthesis  
944 enzymes. The authors proposed a posttranslational destabilization of these proteins in  
945 response to OHP2 deficiency, which could be beneficial when D1, a major sink for  
946 newly synthesized Chl, is not translated. In *Chlamydomonas*, the continued translation  
947 of D1, when the absence of OHP2 limits its ability to ligate Chl, may instead start a  
948 signaling cascade aimed at increasing production of Chl. The *ohp2* mutant may be  
949 missing a negative feedback loop normally operating in the WT. For example, a product  
950 of the interaction between the OHC and pD1, such as the Chl-associated but  
951 unassembled pD1, could negatively regulate association of the Chl synthesis enzymes  
952 to the membrane, signaling that more pD1 is made than can be assembled and the  
953 Chl flux should be lowered. Similarly, the activity of HCF244 as a translation activator  
954 has been proposed to be regulated by the interaction of pD1 with the OHC in the dark  
955 (Chotewutmontri and Barkan, 2020). Another striking observation is the marked Chl  
956 deficiency in the mutant (Supplemental Table S1). The PSII core contains 35 Chl *a*  
957 molecules, while its pigment bed harbors ~200 Chl, so loss of the core itself cannot  
958 fully account for the 40% decrease in Chl content that we observed. In the *Arabidopsis*  
959 *ohp2* mutants, a comparable or even stronger reduction in Chl accumulation was  
960 observed, with Chl *b* reduced to a lesser extent than Chl *a* (Hey and Grimm, 2018;  
961 Myouga et al., 2018). A regulatory role of the OHC on Chl synthesis might be worth  
962 further exploration.

## 963 MATERIAL AND METHODS

### 964 Strains and culture conditions

965 The *ohp2* mutant (*10.1.a*) was isolated in a screen and was generated by nuclear  
966 transformation of the mating type minus (mt-) cell-walled recipient strain Jex4 with the  
967 plasmid pBC1 cut with *SacI* and *KpnI*, described in Houille-Vernes et al. (2011). For  
968 backcrossing of *ohp2* strains, the wild type WT-S34 was used. For localization studies  
969 the cell wall deficient *UVM4* strain generated by Neupert et al. (2009) was used as  
970 recipient strain for glass bead transformation. Two PSII mutants, *nac2-26* which lacks  
971 the *psbD* mRNA stabilization factor NAC2, and the *psbA* deletion mutant FuD7, served  
972 as controls (Bennoun et al., 1986; Boudreau et al., 2000).

973 Algal strains were grown at 23°C under continuous white light on Tris-acetate-  
974 phosphate medium (TAP) at ~10-20  $\mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ , or on High Salt Minimum medium (MIN)

975 agar plates at  $100 \mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ , respectively (Harris, 2009). *ohp2* mutant strains were  
976 kept on plates at very low light ( $\sim 5 \mu\text{E}/\text{m}^2/\text{s}$ ) to limit inadvertent selection of suppressor  
977 mutants. Before usage of *ohp2* in experiments the absence of potential PSII  
978 suppressors was confirmed by Fv/Fm measurements. For experiments, liquid cultures  
979 of *Chlamydomonas* were grown in TAP medium supplemented with 1% sorbitol (TAPS)  
980 at  $\sim 20 - 30 \mu\text{E}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$  until a cell density of  $2-3 \times 10^6$  cells/mL was reached.

981 The reversion rate of *ohp2*, was determined according to Kuras et al. (1997). *ohp2*  
982 mutant cells were collected by centrifugation and resuspended in HSM to a density of  
983  $2 \times 10^8$  cells/mL. 0.5 mL was spread onto 10 HSM agar plates and maintained under  
984 continuous light for three weeks, at which time colonies were counted.

### 985 **Spectroscopy and fluorescence measurements**

986 In vivo spectroscopy measurement using a JTS-10 spectrophotometer (Biologic,  
987 Grenoble, France) was performed as described before in Jalal et al. (2015), using cells  
988 grown under low light conditions. Fv/Fm is presented as the average of four  
989 fluorescence induction measurements at 26, 56, 135 and  $340 \mu\text{E}/\text{m}^2/\text{s}$  followed by a  
990 saturating pulse. PSII/PSI ratio was calculated from the difference in ECS signal at 520  
991 nm, measured 160  $\mu\text{s}$  after a saturating single turnover flash, in the absence or  
992 presence of DCMU (10  $\mu\text{M}$ ) and hydroxylamine (1 mM) to measure (PSI+PSII) or PSI  
993 activity, respectively. In the latter condition, the b/a ratio is the ratio of amplitude of the  
994 second phase of the ECS signal (phase b, due to proton pumping by *Cytb<sub>6</sub>f*) to the  
995 initial phase a (PSI) and is a rough measure of *Cytb<sub>6</sub>f* activity.

### 996 **Analysis of nucleic acids**

997 For isolation of nucleic acids, cells were harvested by centrifugation at  $1,100 \times g$ ,  $4^\circ\text{C}$   
998 for 6 min. Genomic DNA was extracted using the DNeasy Plant Mini Kit (Qiagen,  
999 Hilden, Germany) according to manufacturer's protocol or CTAB buffer (2%  
1000 cetyltrimethylammonium bromide, 100 mM Tris-HCl, pH 8) followed by  
1001 phenol/chloroform/isoamyl alcohol extraction (25:24:1, Carl Roth GmbH, Mannheim,  
1002 Germany). DNAs were digested by restriction enzymes and separated on 0.8%  
1003 agarose gels set up in TPE-buffer (89 mM Tris-phosphate, 2 mM  $\text{Na}_2\text{EDTA}$ ). Total  
1004 cellular RNA was extracted by using the TRI reagent (Sigma-Aldrich, Saint Louis,  
1005 USA), according to the manufacturer's instructions. RNAs were separated on 1%

1006 denaturing formaldehyde agarose gels. After separation nucleic acids were transferred  
1007 to Roti Nylon<sup>+</sup> membrane (Roth, Karlsruhe, Germany), followed by UV light cross-  
1008 linking (UV Crosslinker, UVC 500, Hoefer Inc., San Francisco, USA). Dig-labeled  
1009 probes were synthesized by PCR from total DNA or cloned cDNA (*psbA*) by using  
1010 primers denoted in Supplemental Table S2. Hybridizations and detection of dig-labeled  
1011 probes were performed using standard methods.

## 1012 **Immunoblot analysis**

1013 For isolation of total protein extracts, cells from 20 mL liquid cultures were harvested  
1014 by centrifugation and cell pellets resuspended in 200  $\mu$ L 2 x lysis buffer (120 mM KCl,  
1015 20 mM tricine pH 7.8, 5 mM  $\beta$ -mercaptoethanol, 0.4 mM EDTA, 0.2% Triton X100)  
1016 supplemented with protease inhibitors (cOmplete<sup>™</sup> ULTRA Tablets, Mini, Roche,  
1017 Switzerland) and lysed via sonication on ice. To separate membrane proteins from  
1018 soluble proteins for cell subfractionation, cells from 50 mL culture were resuspended  
1019 in 500  $\mu$ L hypotonic solution (10 mM Tricine/KOH pH 6.8, 10 mM EDTA, 5 mM  $\beta$ -  
1020 mercaptoethanol, and protease inhibitors). Lysis was performed mechanically by  
1021 vortexing thoroughly with glass beads (0.5 mm diameter) two times for 1 min. After  
1022 centrifugation at 15,000 g for 10 min, the supernatant was considered as total soluble  
1023 protein extract. The pellet containing the membrane proteins was resuspended in 2  $\times$   
1024 lysis buffer.

1025 Protein concentration was measured by Bradford Protein Assay (Bradford, 1976).  
1026 SDS-PAGE, protein gel blotting and immunodetection were performed as described by  
1027 Sambrook and Russel (2001). Antibodies were as follows:  $\alpha$ -PsaA (Agrisera, #AS06  
1028 172),  $\alpha$ -D1 (Agrisera, #AS05084),  $\alpha$ -CP43 (Agrisera, #AS11 1787)  $\alpha$ -Cytf (Agrisera, #  
1029 AS08 306),  $\alpha$ -HA (Sigma-Aldrich, #H6908). The antiserum against the spinach Rubisco  
1030 holoenzyme used for the detection of RbcL was kindly provided by G. F. Wildner (Ruhr-  
1031 University Bochum). The antibody against the *Chlamydomonas* D2 protein was  
1032 generated by immunization of rabbits using recombinant D2-GST protein (BioGenes  
1033 GmbH, Berlin, Germany).

## 1034 **Complementation and localization studies**

1035 Constructs generated for the complementation of the *ohp2* mutant and OHP2  
1036 localization studies are shown in Supplemental Figure S3. All constructs were created

1037 by the insertion of a PCR-amplified sequence of interest into the pBC1-CrGFP vector  
1038 (= pJR38, Neupert et al., 2009; construct 1, Supplemental Figure S3) via NdeI or  
1039 NdeI/EcoRI restriction sites. Primers used for cloning are given in Supplemental Table  
1040 S2. For cloning details of the synthetic *Arabidopsis* gene (construct 6, Supplemental  
1041 Figure S3) please see Supplemental Data S2. For localization studies, the cell wall  
1042 deficient strain *UVM4* was transformed by the glass bead method (Kindle, 1990;  
1043 Neupert et al., 2009) and positive transformants were selected by growth on TAP plates  
1044 supplemented with 10 µg/mL paromomycin (Pm). For complementation, constructs  
1045 were integrated into the genome of the *ohp2* mutant strain by the electroporation  
1046 method (Shimogawara et al., 1998), selected for Pm resistance and subsequently  
1047 screened for photoautotrophic growth on HSM plates.

#### 1048 **2D blue native (BN) PAGE**

1049 500 mL liquid cultures of *Chlamydomonas* strains were harvested at 1,000 x g for 10  
1050 min at 4°C and resuspended in 1 mL TMK buffer (10 mM Tris/HCl, pH 6.8, 10 mM  
1051 MgCl<sub>2</sub>, 20 mM KCl) and protease inhibitors (cOmplete™ ULTRA Tablets, Mini, Roche,  
1052 Switzerland). Resuspended cells were lysed by sonication (ultrasound pulses of 10  
1053 sec for 3 times). Cells were centrifuged (1,000 x g, 1 min, 4°C) to remove cell debris  
1054 and unlysed cells. The supernatant was centrifuged for 10 min at 20,000 x g at 4°C.  
1055 The resulting pellet was washed twice with TMK buffer and finally resuspended in 500  
1056 µL TMK buffer. Chl concentration was determined by adding 20 µL of the sample to  
1057 980 µL methanol, 5 min incubation at RT and a 1 min centrifugation step to remove the  
1058 starch. OD<sub>652</sub> of the supernatant was measured and Chl concentration calculated  
1059 using the formula: Chl *a* (mg/mL) = A<sub>652nm</sub> \* 1.45. Aliquots containing 25 µg of Chl  
1060 were again centrifuged for 10 min, 20.000 g at 4°C and resuspended in 51 µL ACA  
1061 buffer (750 mM ε-aminocaproic acid, 50 mM Bis-Tris pH 7.0, 5 mM pH 7.0 EDTA, 50  
1062 mM NaCl) and solubilized for 10 min on ice after addition of n-dodecyl-β-D-maltoside  
1063 (β-DM) to a final concentration of 1.5%. Solubilized proteins were separated from  
1064 insoluble material by centrifugation for 20 min, 16,000 x g at 4°C and mixed with 1/10  
1065 volume of BN loading dye (750 mM ε-aminocaproic acid, 5% Coomassie G250 (w/v)).  
1066 First dimension native electrophoresis was carried out according to Schägger and  
1067 Jagow (1991) on a 5 – 12 % linear gradient. Protein complexes were further subjected  
1068 to 2D 12% SDS- PAGE and analyzed by immunoblotting.

## 1069 **Ribosome profiling and data analysis**

1070 For each replicate, 500 mL culture was supplemented with 100 µg/mL chloramphenicol  
1071 and 100 µg/mL cycloheximide, followed by rapid cooling using plastic ice cubes. The  
1072 cold (< 8°C) culture was then centrifuged at 3,000 g for 2 min to pellet the cells. The  
1073 pellet was washed once with ice-cold polysome buffer (20 mM Tris pH 8.0, 25 mM KCl,  
1074 25 mM MgCl<sub>2</sub>, 1 mM dithiothreitol, 100 µg/mL chloramphenicol, 100 µg/mL  
1075 cycloheximide) and then flash-frozen in liquid nitrogen. The pellet was resuspended in  
1076 8 mL of polysome buffer containing 1x Protease inhibitor Cocktail (Roche) and 1 mM  
1077 PMSF, and the cells were lysed using a French press (Avestin®) at 2 bar. Polysome  
1078 isolation from the lysate, RNA digestion and FP isolation have been performed as  
1079 described in Trösch et al. (2018). For the total RNA, a separate 8 mL of the culture  
1080 was centrifuged at 3000 g for 2 min, and 750 µL of Trizol was added to the pellet  
1081 directly. Total RNA extraction as well as RNA labelling and microarray analysis was  
1082 performed as described in Trösch et al. (2018).

1083 Data processing and analysis was conducted according to previous studies (Trösch  
1084 et al., 2018; Zoschke et al., 2013): Local background was subtracted from single  
1085 channels (F635-B635 and F532-B532, respectively) and probes located within  
1086 annotated and confirmed chloroplast reading frames were normalized to the average  
1087 signal of the compared datasets including all replicates of ribosome FPs and total  
1088 mRNA to remove overall differences introduced by technical variations. Probes  
1089 covering a respective ORF were averaged, and relative abundance of ribosome FPs  
1090 and total mRNA was calculated for each ORF by normalizing each ORF value to the  
1091 average of all ORF values. By this, expression of the individual ORF is considered in  
1092 relation to mean values of all plastid-encoded genes. Relative translation efficiency is  
1093 determined by comparing to values of average FP intensities relative to the average  
1094 RNA intensities, for each ORF. All average values and standard deviations are based  
1095 on three independent biological replicates. Significant differences in gene-specific RNA  
1096 and ribosome FP accumulation and translation efficiencies between WT and *ohp2*  
1097 mutant data were determined with a Welch's t-test and corrected for multiple testing  
1098 according to Storey's q-value method. Genes were marked as significant for q-value  
1099 of <0.05 and with expression changes more than two-fold (Supplemental Dataset 1).

1100 **Mass spectrometry**

1101 Whole cell proteomics: 50 µg of proteins were separated by SDS-PAGE. After a short  
1102 migration (< 0.5 cm) and Coomassie blue staining, gel bands containing proteins were  
1103 excised and destained. Gel pieces were subjected to a 30 min reduction at 56°C and  
1104 a 1 h cysteine alkylation at room temperature using 10 mM dithiothreitol and 50 mM  
1105 iodoacetamide in 50 mM ammonium bicarbonate, respectively. Proteins were digested  
1106 overnight at 37°C using 500 ng of trypsin (Trypsin Gold, Promega). Supernatants were  
1107 kept and peptides remaining in gel pieces were further extracted with 1% (v/v)  
1108 trifluoroacetic acid. Corresponding supernatants were pooled and dried. Peptide  
1109 mixtures were subsequently reconstituted in 200 µL of solvent A (0.1% (v/v) formic acid  
1110 in 3% (v/v) acetonitrile). Five microliters of peptide mixtures were analyzed in duplicate  
1111 on a Q-Exactive Plus hybrid quadrupole-orbitrap mass spectrometer (Thermo Fisher)  
1112 as described in Pérez-Pérez et al. (2017) except that peptides were separated on a  
1113 PepMap™ RSLC C18 Easy-Spray column (75 µm x 50 cm, 2 µm, 100 Å; Thermo  
1114 Scientific) with a 90 min gradient (0 to 20% B solvent (0.1% (v/v) formic acid in  
1115 acetonitrile) in 70 min and 20 to 37% B solvent in 20 min).

1116 Membrane proteomics: Cells were collected by centrifugation, resuspended in 25 mM  
1117 phosphate buffer at a density of app. 0.5 µg/µL total Chl and disrupted by three  
1118 consecutive freeze and thaw cycles. Soluble proteins were separated by centrifugation  
1119 at 4°C, 25,000 g for 15 min. 20 µg total protein from the pellet fraction were precipitated  
1120 in 80 % acetone, tryptically digested and desalted as described (Hammel et al., 2018).  
1121 Peptides were resuspended in a solution of 2% acetonitrile, 1% formic acid just before  
1122 the LC-MS/MS run. The LC-MS/MS system (Eksigent nanoLC 425 coupled to a  
1123 TripleTOF 6600, ABSciex) was operated basically as described for data dependent  
1124 acquisition (Hammel et al., 2018).

1125 For targeted MRM-HR data acquisition a list of 39 precursor m/z was chosen for  
1126 targeting the 5 different proteins. The precursors were selected from results of the data  
1127 dependent acquisitions and good responding peptides as predicted by a deep learning  
1128 algorithm for peptide detectability prediction, d::pPop (Zimmer et al., 2018). Collision  
1129 energies were calculated from the standard CE parameters of the instrument, dwell  
1130 time was set to 60 ms, fragments were acquired from 110 m/z – 1600 m/z, resulting in  
1131 a cycle time of 2.6 s.

1132 Data processing and label-free quantification. Shotgun proteomics raw data were  
1133 processed using the MaxQuant software package as described in Martins et al. (2020)

1134 with slight modifications. For protein identification and target decoy searches were  
1135 performed using a home-made *Chlamydomonas reinhardtii* protein database  
1136 consisting in the JGI Phytozome nuclear-encoded proteins database (v.5.6)  
1137 concatenated with chloroplast- and mitochondria-encoded proteins in combination with  
1138 the Maxquant contaminants.

1139 For whole cell proteomics the mass tolerance in MS and MS/MS was set to 10 ppm  
1140 and 20 mDa, respectively and proteins were validated if at least 2 unique peptides  
1141 having a protein FDR < 0.01 were identified. For quantification, unique and razor  
1142 peptides with a minimum ratio count  $\geq 2$  unique peptides were used and protein  
1143 intensities were calculated by Delayed Normalization and Maximal Peptide Ratio  
1144 Extraction (MaxLFQ) according to Cox et al. (2014). For membrane protein data  
1145 dependent runs MaxQuant software (v1.6.0.1) (Cox and Mann, 2008; Tyanova et al.,  
1146 2016) was used. Peptide identification, protein group assignment and quantification  
1147 was done with standard settings for ABSciex Q-TOF data except that 3 miss-cleavages  
1148 were allowed, minimum peptide length was set to 6 AA, maximum peptide mass 6600  
1149 Da.

1150 *Shotgun data analysis:* For statistical analysis, LFQ intensities obtained from  
1151 MaxQuant were further analyzed with the Perseus software package version 1.6.15.0  
1152 (Tyanova et al., 2016). In case of the whole cell data the two measurement replicates  
1153 were averaged first. For both datasets the biological replicates were grouped and the  
1154 LFQ intensities were  $\text{Log}_2$  transformed. The data were filtered to contain at least three  
1155 valid values (out of four biological replicates) or two valid values (out of three biological  
1156 replicates) in case of the whole cell or membrane fractions, respectively. Significant  
1157 changes in LFQ intensities compared to the WT were identified by a modified two  
1158 sample t-test of the Perseus software (permutation-based FDR=5%, artificial within  
1159 group variance  $S_0=1$ ). The *ohp2* mutant, the suppressor mutant *M-su1* and in case of  
1160 the whole cell samples the complemented line *ohp2:OHP2-HA*, were compared to the  
1161 WT independently. Data visualizations shown in Figure 5 and Supplemental Figure S9  
1162 were done using R (R Core Team, 2018). The  $\text{log}_2$  transformed LFQ intensities and t-  
1163 test results of the whole cell samples and membrane fractions are shown in  
1164 Supplemental Dataset 2 and Supplemental Dataset 3, respectively.

1165 *Targeted mass spectrometry data analysis* was done using Skyline software  
1166 (v20.1.0.155) (MacLean et al., 2010; Pino et al., 2020). The spectral library was built  
1167 by prediction using ProSIT (Gessulat et al., 2019), retention times were taken, where

1168 possible, from identification results of the respective peptides using MaxQuant. A  
1169 minimum of 2 precursors per protein and 4 transitions per precursor was used for  
1170 quantification. The integration borders were manually adjusted where necessary and  
1171 dotp values for the coeluting fragments as reported from Skyline was taken as a quality  
1172 measure for correct assignment (Supplemental Dataset 4). Retention times varied over  
1173 all runs by less than 0.5 min, dotp values were usually >0.85 across the replicates  
1174 where a correct assignment was assumed. Summed fragment peak areas were  
1175 normalized for unequal sample loading to the total intensities of proteins as identified  
1176 by MaxQuant in data dependent runs.

1177 The untargeted mass spectrometry proteomics data have been deposited to the  
1178 ProteomeXchange Consortium via the PRIDE (Perez-Riverol et al., 2021) partner  
1179 repository with the dataset identifier PXD031558 (Reviewer account details:  
1180 <https://www.ebi.ac.uk/pride/login>; Username: reviewer\_pxd031558@ebi.ac.uk;  
1181 Password: D3xVDFGU). The targeted mass spectrometry proteomics data have been  
1182 deposited to the ProteomeXchange Consortium via the Panorama Public (Sharma et  
1183 al., 2018) partner repository with the dataset identifier PXD031631 (Reviewer account  
1184 details: <https://panoramaweb.org/TIFMYs.url>; Username:  
1185 panorama+reviewer102@proteinms.net; Password: iEqMdcRZ).

1186

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1194

## 1195 **AUTHOR CONTRIBUTIONS**

1196 AVB, OVA, FWi, JN, and MS conceived and designed the research; FWa, KD, IM, FE,  
1197 RT, FS, OVA, AVB, and XJ performed the experiments; FS, LDW, and FWi analyzed  
1198 data; AVB, OVA, and FWi wrote the paper with inputs from all authors.

1199

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1202 **REFERENCES**

1203  
1204

1205 **FIGURE LEGENDS**

1206 **Figure 1. Identification and complementation of the mutation in *OHP2*.**

1207 **A)** Schematic representation of the genomic region of *OHP2* (*Cre06.g251150*) on  
1208 chromosome 6. Grey boxes indicate exons, smaller grey arrow and box 5' and 3' UTRs,  
1209 respectively, and light grey boxes introns. For simplicity, upstream and downstream  
1210 loci are not displayed. The probe used in B) is shown as black bar above the gene  
1211 model. The insertion site of a putative >5 kb *TOC1* transposon in exon 3 of the *OHP2*  
1212 gene in the mutant strain *10.1a* is indicated by an open triangle (compare  
1213 Supplemental Figure S2). The filled black triangle in intron 2 of the *OHP2* gene  
1214 indicates the position of a polymorphism caused by an additional putative insertion in  
1215 the mutant and its background strain as identified during Illumina sequencing (compare  
1216 Supplemental Figure S2). Given position of HindIII (H) and PstI (P) restriction sites are  
1217 based on the reference genome of *Chlamydomonas reinhardtii* v5.5 in Phytozome.  
1218 Note that this sequence is based on the mt+ genome assembly of strain *CC-503* and  
1219 does not necessarily reflect the restriction site positions in the investigated mt- allele.  
1220 **B)** Southern blot analysis of genomic DNA from the Jex4 recipient strain (WT) and the  
1221 *10.1a* mutant. 10 µg of genomic DNA were fractionated in a 0.8% agarose gel after  
1222 digestion by restriction enzymes HindIII or PstI, respectively, blotted onto a nylon  
1223 membrane, and hybridized with the *OHP2*-specific probe indicated in A). **C)**  
1224 Photosynthetic parameters of the Jex4 recipient strain (WT), the *ohp2* mutant and *ohp2*  
1225 complemented with pBC1-CrOHP2-HA (*ohp2:OHP2-HA*, compare construct 2 in  
1226 Supplemental Figure S3). Fluorescence induction kinetics (upper panel), measured  
1227 under illumination at 135 µE/m<sup>2</sup>/s, followed by a saturating pulse (arrow) and dark  
1228 relaxation. Fluorescence intensity is normalized to the F<sub>m</sub> value. Electrochromic shift  
1229 at 520 nm (lower panel), measured in the absence or presence of DCMU and  
1230 hydroxylamine. Values are normalized to the signal after the saturating flash. **D)**

1231 Growth test. Cells were resuspended in sterile H<sub>2</sub>O to a concentration of 10<sup>5</sup> cells/mL  
1232 and spotted onto acetate-containing (TAP) or High Salt Minimum media (MIN) and  
1233 grown for 6 d under higher light (HL) at 100 μE/m<sup>2</sup>/s or low light (LL) at 30 μE/m<sup>2</sup>/s.

1234 **Figure 2. Protein sequence alignment and complementation analysis.**

1235 **A)** Sequence alignment of eukaryotic One-Helix Proteins OHP1 and OHP2 and two  
1236 cyanobacterial HLIPs. OHP2 and OHP1 protein sequences from *C.r.* (*C. reinhardtii*,  
1237 v5.5, *Cre06.g251150* and *Cre02.g109950*), *A.t.* (*Arabidopsis thaliana*, TAIR10,  
1238 *AT1G34000.1* and *AT5G02120*), *P.p.* (*Physcomitrella patens*, v3.3,  
1239 *Pp3c2\_26700V3.1*), and *Z.m.* (*Zea mays*, PH207 v1.1, *Zm00008a032025\_T01*) were  
1240 obtained from Phytozome (<https://phytozome.jgi.doe.gov/pz/portal.html>). Amino acid  
1241 sequences of the High light-induced proteins HliC (*ss/1633*) and HliD (*ssr1789*) from  
1242 *Synechocystis sp.* PCC6803 were taken from CyanoBase  
1243 (<http://www.kazusa.or.jp/cyano/>). The multiple sequence alignment was performed by  
1244 using ClustalW (Thompson et al., 2002), manually edited and displayed with Genedoc  
1245 (Nicholas et al., 1997). Black shading represents 100% conservation, dark grey and  
1246 grey 60% and 40%, respectively. The positions of the predicted Chl binding region as  
1247 well as a hydrophobic stretch (HS) at the C-terminus are indicated (compare  
1248 Supplemental Figure S4A). Two residues described to be important for Chl binding in  
1249 LHCB from spinach are labeled with red asterisks (Kühlbrandt et al., 1994). The N-  
1250 terminal chloroplast transit peptides predicted by TargetP-2.0 (Emanuelsson et al.,  
1251 2000; Nielsen et al., 1997) for all eukaryotic proteins shown, are not included in the  
1252 alignment. The *Chlamydomonas* OHP2-based prediction of a transmembrane helix by  
1253 TMPred (Hofmann and Stoffel, 1993; [https://embnet.vital-](https://embnet.vital-it.ch/software/TMPRED_form.html)  
1254 [it.ch/software/TMPRED\\_form.html](https://embnet.vital-it.ch/software/TMPRED_form.html)), is indicated as a grey bar above the sequence  
1255 (compare Supplemental Figure S4B). The mutation identified in the *Chlamydomonas*  
1256 *OHP2* gene corresponds to residue M76 of the protein (indicated by a red triangle)  
1257 which lies within the fully-conserved stretch in the N-terminal part of the protein. **B)** The  
1258 C-terminal hydrophobic stretch is required for restoration of photoautotrophy. Growth  
1259 test (left panel). The *ohp2* mutant was transformed with the construct pBC1-CrOHP2-  
1260 ΔHS-HA (see construct 5 in Supplemental Figure S3). Pm-resistant transformants  
1261 (*ohp2:CrOHP2-ΔHS-HA*) were tested for photoautotrophic growth on HSM plates  
1262 (MIN). Immunoblot analysis (right panel) of Pm-resistant transformants. 40 μg of total  
1263 proteins were separated on 15% denaturing polyacrylamide gels and analyzed with

1264 the antibodies indicated on the left. The complemented strain *ohp2:OHP2-HA* was  
1265 used as positive control for the accumulation of HA-tagged proteins. Samples were run  
1266 on the same gel but not in adjacent lanes as indicated by a vertical black line.

1267 **Figure 3. Drastically diminished D1 protein accumulation and synthesis in the**  
1268 ***ohp2* mutant.**

1269 The following strains were subjected to analysis: Jex4 (WT), *ohp2* and the  
1270 complemented strain *ohp2:OHP2-HA* clone #53 (*ohp2:OHP2-HA*). **A)** Accumulation of  
1271 photosynthesis related chloroplast proteins. 30 µg of total proteins from indicated  
1272 strains were separated by 12% (α-D1, α-PsaA, α-Cytf, α-CP43, α-D2) or 15% (α-HA,  
1273 α-RbcL) denaturing polyacrylamide gels and analyzed by the antibodies indicated. **B)**  
1274 Accumulation of photosynthesis related chloroplast transcripts. 3 µg of total cellular  
1275 RNA from indicated strains were fractionated by denaturing agarose gel  
1276 electrophoresis and blotted onto a nylon membrane. Membranes were hybridized with  
1277 probes specific for *psbA* and *psbD*. For loading control, the same blot was hybridized  
1278 with a probe specific for *rbcL* transcripts. *nac2* and *FuD7* mutants were employed as  
1279 negative controls for *psbD* or *psbA* mRNA accumulation, respectively. *ohp2* was run  
1280 on the same gel but not in adjacent lanes as indicated by a vertical black line. **C)**  
1281 Synthesis of PSII subunits. <sup>14</sup>C labelling of chloroplast-encoded proteins was  
1282 performed as described previously in Spaniol et al. (2021). Autoradiogram of cells  
1283 pulsed for 5 min with <sup>14</sup>C-acetate in the presence of cycloheximide (left), then chased  
1284 for 45 min after washing and chloramphenicol addition. For *ohp2*, two independently  
1285 grown cultures (#1, #2) were analyzed. The complemented strain was run on the same  
1286 gel and exposed in the same conditions, but in non-adjacent lanes as indicated by a  
1287 vertical black line. **D)** 2D BN-PAGE analysis of photosynthetic protein complexes in  
1288 indicated strains. After solubilization of membranes with 1.5% (w/v) dodecyl-b-D-  
1289 maltopyranoside, thylakoid proteins were separated by 5 to 12 % BN gels in the first  
1290 dimension and 12% SDS-gels in the second dimension. Photosynthetic complexes  
1291 were detected using the indicated antibodies. The positions of major PSI and PSII  
1292 complexes are designated. sc: supercomplex

1293 **Figure 4. Targeted ribosome profiling of chloroplast translation reveals protein**  
1294 **synthesis defects in the *ohp2* mutant.**

1295 Ribosome profiling and transcript analysis of WT and *ohp2* mutant grown  
1296 mixotrophically under 30  $\mu\text{E}/\text{m}^2/\text{s}$ . **A)** The average mRNA (blue) and ribosome FP (dark  
1297 grey) abundances were calculated from three independent biological replicates and  
1298 plotted in  $\log_{10}$  scale for *ohp2* mutant versus WT, respectively. mRNAs encoding PSII  
1299 subunits, which display altered translation in the mutant are highlighted. *Pearsons's r*-  
1300 value and *p*-value are given in *nEm* non-superscript format for  $n \cdot 10m$ . **B)** The relative  
1301 average transcript abundances (RNA), translation output and translation efficiency  
1302 were calculated for each chloroplast reading frame in both *Chlamydomonas* strains,  
1303 normalized to overall signal intensities, and plotted as heat map (*ohp2* versus WT) in  
1304  $\log_2$  scale. Increased RNA accumulation, translation output or translation efficiency in  
1305 the mutant is shown in red, reduced levels are shown in blue (see scale bar). PSII  
1306 subunits that are further discussed are highlighted with an arrow. **DC)** For the Open  
1307 Reading Frames (ORFs) of *psbA* (introns not shown) and the *psbB* operon normalized  
1308 ribosome FP intensities were plotted as mean  $\log_2$  ratios between *ohp2* mutant and  
1309 WT. Error bars denote differences between three independent biological replicates.  
1310 Grey bars below the panels indicate respective ORFs with trans-membrane segments  
1311 (TMS in green). Related to Supplemental Figures S6, S7 and Supplemental Dataset  
1312 1. **C)** CES cascade of PSII. When newly synthesized CES polypeptides, like D1 and  
1313 CP47, cannot assemble, they repress the translation initiation of their encoding mRNA  
1314 as described before by Minai and coworkers (2006). Reduced translation of *psbH* may  
1315 indicate that it represents a further component of the CES cascade.

1316 **Figure 5. Altered proteome composition in the *ohp2* mutant and suppressor**  
1317 **strain.**

1318 Volcano plots representing the relative proteome changes of whole cell lysates (**A)** of  
1319 *ohp2* mutants versus WT (left), suppressor line 1 versus WT (right) and thylakoid  
1320 membrane proteins (**B)** as comparison between *ohp2* mutant versus WT (left) and  
1321 suppressor line 1 versus WT (right). For B), proteins of the membrane fraction were  
1322 enriched by crude fractionation. All experiments were performed in at least three  
1323 independent biological replicates. Mean fold change of LFQ values (in  $\log_2$ ) is plotted  
1324 on the x-axis, *p*-values (in  $-\log_{10}$ ) are plotted on the y-axis. Light grey dots represent

1325 proteins with no significant change, dark grey dots show proteins that are significantly  
1326 different with  $FDR < 0.05$  and  $S_0 = 1$ . Proteins of PSI, PSII, the chloroplast ATP synthase,  
1327 and of proteins involved in Chl biogenesis are marked in color as depicted in the legend  
1328 on the right. Large colored dots are significantly different. **C)** Abundance of selected  
1329 proteins in membrane protein extracts as determined by targeted mass spectrometry  
1330 in different strains as compared to WT. Error bars indicate standard deviation over  
1331 three biological replicates. Missing values indicate instances where no peptides could  
1332 be found or are present below baseline signal of app. 5% compared to WT as judging  
1333 from dotp values and fragment coelution profiles. Related to Supplemental Figure S9  
1334 and Supplemental Datasets 2-4.

1335 **Figure 6. Mutation of the FtsH protease partially restores PSII activity but not**  
1336 **photoautotrophy.**

1337 **A)** Growth tests of the progeny of a cross *10.1a* X *ftsH1-1*. Cells were grown on TAP  
1338 or MIN medium at the indicated light intensity. Spots shown are typical of the indicated  
1339 genotypes. **B)**  $F_v/F_m$  values for the four genotypes, recorded from the plates in A).  
1340 Values are average of 8, 7, 11 and 6 strains for the WT, *ftsH1*, *ohp2* and *ohp2 ftsH1*  
1341 genotypes, respectively. Error bars represent S.D. **C)** Typical fluorescence induction  
1342 curves for the four genotypes, recorded with the fluorescence camera on the TAP  
1343 plates described in A). Fluorescence is normalized to the  $F_m$  value (arrow indicates  
1344 saturating flash).

1345 **Figure 7. Schematic model for the role the OHC complex and associated proteins**  
1346 **in D1 synthesis and first steps of PSII de novo assembly.**

1347 In the WT (**left panel**), likely triggered by HCF244, HCF173 binds the 5'-UTR of *psbA*  
1348 mRNA to promote translation initiation. pD1 is co-translationally inserted into the  
1349 thylakoid membrane. Membrane associated luminal HCF136 stabilizes pD1 and is  
1350 involved in RC assembly. The OHC is required for early PSII assembly steps and  
1351 proposed to insert Chl into pD1 proteins. Recently reported negative autoregulatory  
1352 circuits of *psbA* translation initiation involving the OHC are not displayed here  
1353 (Chotewutmontri and Barkan, 2020; Chotewutmontri et al., 2020). In *ohp2* (**middle**  
1354 **panel**) and suppressor strains (**right panel**) residual levels of HCF244 may promote  
1355 *psbA* translation via interaction with HCF173. Ongoing *psbA* translation may be further

1356 supported by overaccumulation of HCF173. However, in the *ohp2* mutant synthesized  
1357 D1 is rapidly degraded and does not accumulate in the absence of the OHC complex.  
1358 In the suppressor strains, stabilization of nascent D1 may be accomplished by  
1359 increased membrane association of HCF136 or other mechanisms to allow assembly  
1360 of early PSII intermediates. HCF173, HCF136, and HCF244 protein amounts detected  
1361 by proteomics are indicated by differently sized ovals. Names of cyanobacterial  
1362 homologs are given in brackets.

## 1363 SUPPLEMENTAL MATERIAL

### 1364 SUPPLEMENTAL FIGURES

1365 **Figure S1.** Genetic analysis of the *10.1a* mutant.

1366 **Figure S2.** Identification of the *ohp2* mutation.

1367 **Figure S3.** DNA constructs used for complementation and localization studies.

1368 **Figure S4.** Hydrophobicity prediction for *Chlamydomonas* and *Arabidopsis* OHP2  
1369 amino acid sequences.

1370 **Figure S5.** OHP2 is a membrane localized chloroplast protein.

1371 **Figure S6.** Reproducibility of RF experiments and results for all chloroplast encoding  
1372 genes.

1373 **Figure S7.** Targeted ribosome profiling of chloroplast translation reveals enhanced  
1374 *psbD* translation in the *ohp2* mutant.

1375 **Figure S8.** Re-accumulation of D2 and restoration of photoautotrophy in suppressor  
1376 mutants.

1377 **Figure S9.** Heat maps representing the reproducibility of LC-MS experiments and  
1378 complementation of *ohp2*.

1379 **Figure S10.** Complementation of the *ohp2* mutant with the orthologous *Arabidopsis*  
1380 protein restores photoautotrophy and D1 accumulation.

1381 **Figure S11.** Transcript accumulation level of *OHP2*, *OHP1*, and HCF244 in  
1382 *Chlamydomonas* under various growth conditions.

### 1383 SUPPLEMENTAL TABLES AND DATA

1384 **Supplemental Table S1.** Photosynthetic parameters and Chl composition of  
1385 *Chlamydomonas* WT Jex4 (WT), the *ohp2* mutant and complemented strain.

1386 **Supplemental Table S2.** Primers used in this study.

1387 **Supplemental Data S1.** Flanking sequence tags in the *ohp2* mutant obtained by  
1388 Illumina sequencing.

1389 **Supplemental Data S2.** Synthetic *Arabidopsis OHP2* nucleotide sequences and  
1390 derived protein sequence used to complement the *Chlamydomonas ohp2* mutant  
1391 strain.

## 1392 SUPPLEMENTAL DATASETS

1393 **Supplemental Dataset 1.** Complete dataset from array-based ribosome profiling  
1394 experiments.

1395 **Supplemental Dataset 2.** Complete dataset from whole cell proteomics experiments.

1396 **Supplemental Dataset 3.** Complete dataset from membrane proteomics experiments.

1397 **Supplemental Dataset 4.** Targeted mass spectrometry data

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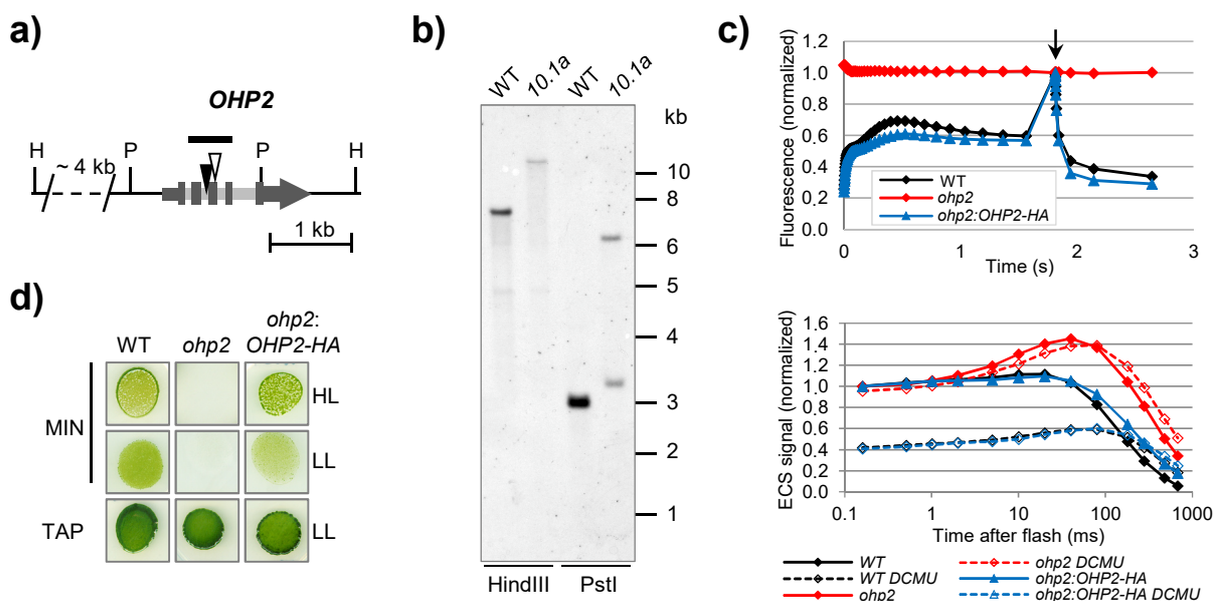
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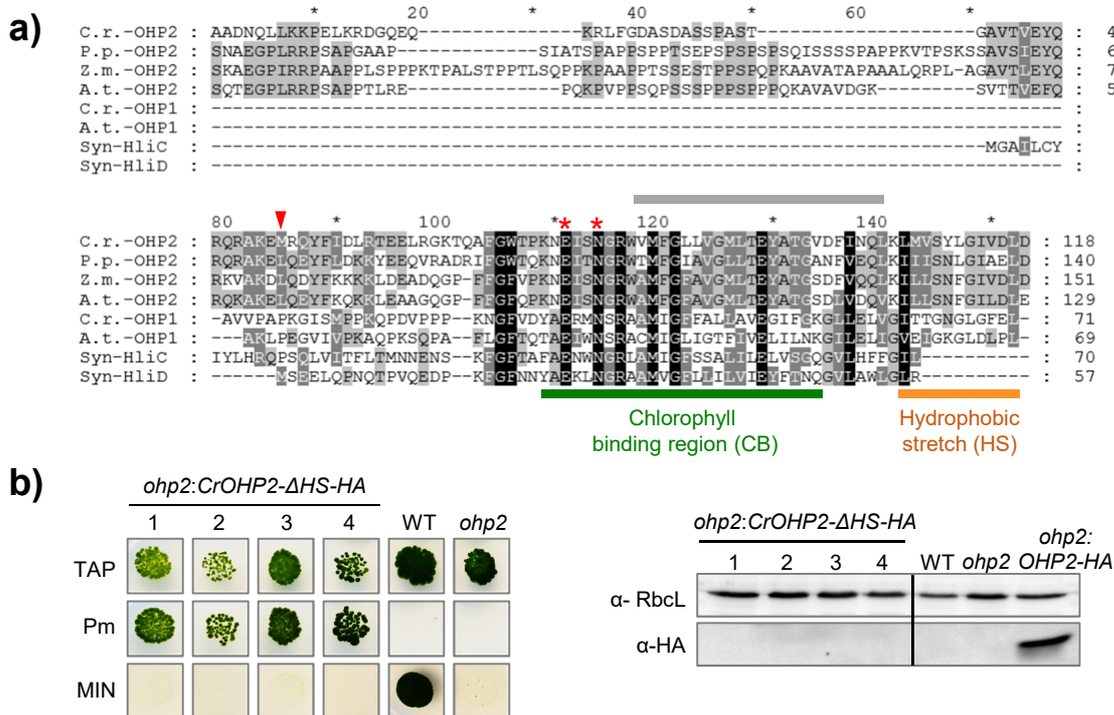
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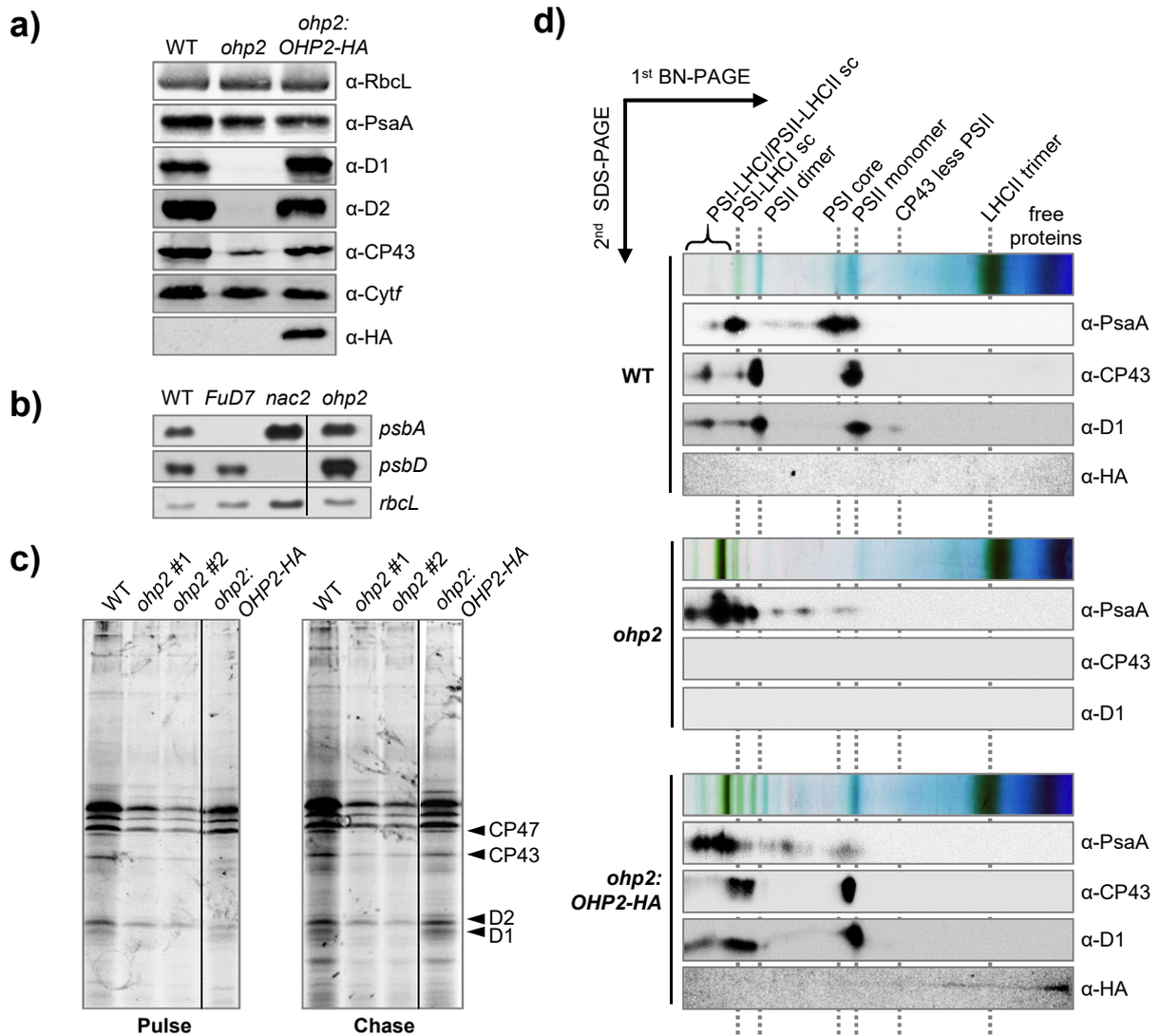
# Figures



**Figure 1. Identification and complementation of the mutation in *OHP2*.** a) Schematic representation of the genomic region of *OHP2* (*Cre06.g251150*) on chromosome 6. Grey boxes indicate exons, smaller grey arrow and box 5' and 3' UTRs, respectively, and light grey boxes introns. For simplicity, upstream and downstream loci are not displayed. The probe used in b) is shown as black bar above the gene model. The insertion site of a putative >5 kb *TOC1* transposon in exon 3 of the *OHP2* gene in the mutant strain *10.1a* is indicated by an open triangle (compare Supplemental Figure S2). The filled black triangle in intron 2 of the *OHP2* gene indicates the position of a polymorphism caused by an additional putative insertion in the mutant and its background strain as identified during Illumina sequencing (compare Supplemental Figure S2). Given position of HindIII (H) and PstI (P) restriction sites are based on the reference genome of *Chlamydomonas reinhardtii* v5.5 in Phytozome. Note that this sequence is based on the mt+ genome assembly of strain CC-503 and does not necessarily reflect the restriction site positions in the investigated mt- allele. b) Southern blot analysis of genomic DNA from the Jex4 recipient strain (WT) and the *10.1a* mutant. 10  $\mu$ g of genomic DNA were fractionated in a 0.8% agarose gel after digestion by restriction enzymes HindIII or PstI, respectively, blotted onto a nylon membrane, and hybridized with the *OHP2*-specific probe indicated in a). c) Photosynthetic parameters of the Jex4 recipient strain (WT), the *ohp2* mutant and *ohp2* complemented with pBC1-CrOHP2-HA (*ohp2:OHP2-HA*, compare construct 2 in Supplemental Figure S3). Fluorescence induction kinetics (upper panel), measured under illumination at 135  $\mu$ E/m<sup>2</sup>/s, followed by a saturating pulse (arrow) and dark relaxation. Fluorescence intensity is normalized to the Fm value. Electrochromic shift at 520 nm (lower panel), measured in the absence or presence of DCMU and hydroxylamine. Values are normalized to the signal after the saturating flash. d) Growth test. Cells were resuspended in sterile H<sub>2</sub>O to a concentration of 10<sup>5</sup> cells/mL and spotted onto acetate-containing (TAP) or High Salt Minimum media (MIN) and grown for 6 d under higher light (HL) at 100  $\mu$ E/m<sup>2</sup>/s or low light (LL) at 30  $\mu$ E/m<sup>2</sup>/s.

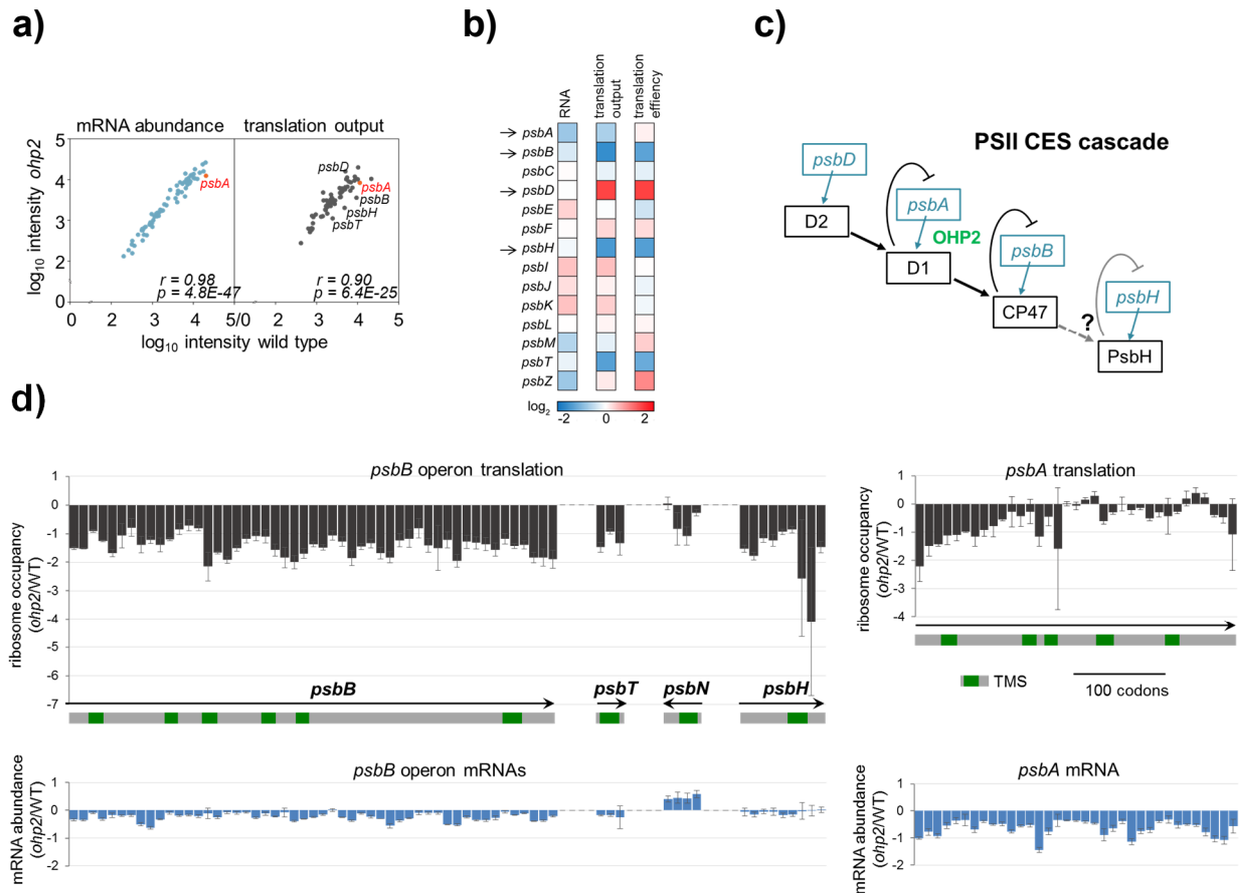


**Figure 2. Protein sequence alignment and complementation analysis.** a) Sequence alignment of eukaryotic One-Helix Proteins OHP1 and OHP2 and two cyanobacterial HLIPs. OHP2 and OHP1 protein sequences from *C.r.* (*C. reinhardtii*, v5.5, *Cre06.g251150* and *Cre02.g109950*), *A.t.* (*Arabidopsis thaliana*, TAIR10, *AT1G34000.1* and *AT5G02120*), *P.p.* (*Physcomitrella patens*, v3.3, *Pp3c2\_26700V3.1*), and *Z.m.* (*Zea mays*, PH207 v1.1, *Zm00008a032025\_T01*) were obtained from Phytozome (<https://phytozome.jgi.doe.gov/pz/portal.html>). Amino acid sequences of the High light-induced proteins HliC (*ssl1633*) and HliD (*ssl1789*) from *Synechocystis* sp. PCC6803 were taken from CyanoBase (<http://www.kazusa.or.jp/cyano/>). The multiple sequence alignment was performed by using ClustalW (Thompson et al., 2002), manually edited and displayed with Genedoc (Nicholas et al., 1997). Black shading represents 100% conservation, dark grey and grey 60% and 40%, respectively. The positions of the predicted Chl binding region (CB) as well as a hydrophobic stretch (HS) at the C-terminus are indicated (compare Supplemental Figure S4A). Two residues described to be important for Chl binding in LHCb from spinach are labeled with red asterisks (Kühlbrandt et al., 1994). The N-terminal chloroplast transit peptides predicted by TargetP-2.0 (Emanuelsson et al., 2000; Nielsen et al., 1997) for all eukaryotic proteins shown, are not included in the alignment. The *Chlamydomonas* OHP2-based prediction of a transmembrane helix by TMPred (Hofmann & Stoffel, 1993; [https://embnet.vital-it.ch/software/TMPRED\\_form.html](https://embnet.vital-it.ch/software/TMPRED_form.html)), is indicated as a grey bar above the sequence (compare Supplemental Figure S4B). The mutation identified in the *Chlamydomonas* OHP2 gene corresponds to residue M76 of the protein (indicated by a red triangle) which lies within the fully-conserved stretch in the N-terminal part of the protein. b) The C-terminal hydrophobic stretch is required for restoration of photoautotrophy. Growth test (left panel). The *ohp2* mutant was transformed with the construct pBC1-CrOHP2-ΔHS-HA (see construct 5 in Supplemental Figure S3). Pm-resistant transformants (*ohp2:CrOHP2-ΔHS-HA*) were tested for photoautotrophic growth on HSM plates (MIN). Immunoblot analysis (right panel) of Pm-resistant transformants. 40 μg of total proteins were separated on 15% denaturing polyacrylamide gels and analyzed with the antibodies indicated on the left. The complemented strain *ohp2:OHP2-HA* was used as positive control for the accumulation of HA-tagged proteins. Samples were run on the same gel but not in adjacent lanes as indicated by a vertical black line.



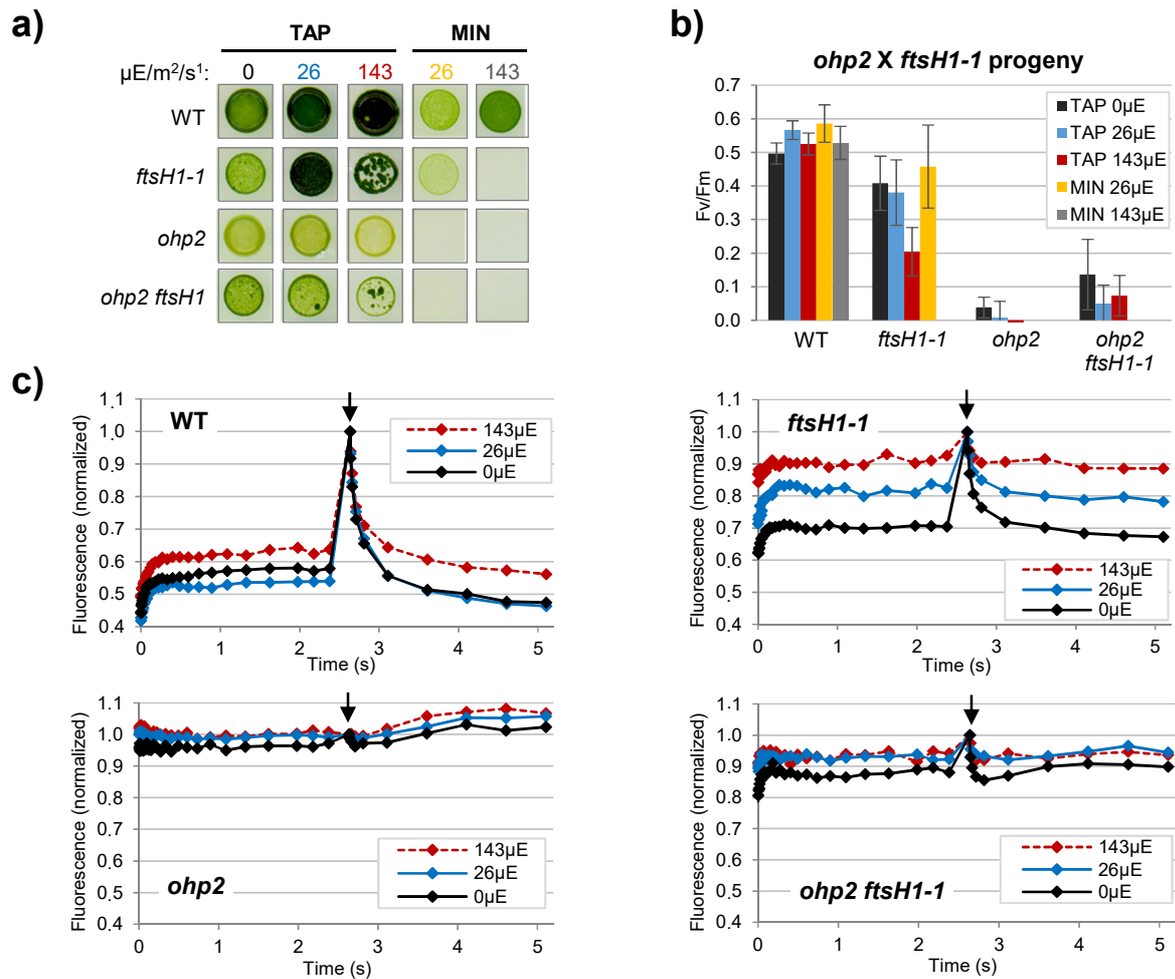
**Figure 3. Drastically diminished D1 protein accumulation and synthesis in the *ohp2* mutant.**

The following strains were subjected to analysis: Jex4 (WT), *ohp2* and the complemented strain *ohp2:OHP2-HA* clone #53 (*ohp2:OHP2-HA*). a) Accumulation of photosynthesis related chloroplast proteins. 30  $\mu\text{g}$  of total proteins from indicated strains were separated by 12% ( $\alpha$ -D1,  $\alpha$ -PsaA,  $\alpha$ -Cytf,  $\alpha$ -CP43,  $\alpha$ -D2) or 15% ( $\alpha$ -HA,  $\alpha$ -RbcL) denaturing polyacrylamide gels and analyzed by the antibodies indicated. b) Accumulation of photosynthesis related chloroplast transcripts. 3  $\mu\text{g}$  of total cellular RNA from indicated strains were fractionated by denaturing agarose gel electrophoresis and blotted onto a nylon membrane. Membranes were hybridized with probes specific for *psbA* and *psbD*. For loading control, the same blot was hybridized with a probe specific for *rbcL* transcripts. *nac2* and *FuD7* mutants were employed as negative controls for *psbD* or *psbA* mRNA accumulation, respectively. *ohp2* was run on the same gel but not in adjacent lanes as indicated by a vertical black line. c) Synthesis of PSII subunits.  $^{14}\text{C}$  labelling of chloroplast-encoded proteins was performed as described previously in Spaniol et al. (2021). Autoradiogram of cells pulsed for 5 min with  $^{14}\text{C}$ -acetate in the presence of cycloheximide (left), then chased for 45 min after washing and chloramphenicol addition. For *ohp2*, two independently grown cultures (#1, #2) were analyzed. The complemented strain was run on the same gel and exposed in the same conditions, but in non-adjacent lanes as indicated by a vertical black line. d) 2D BN-PAGE analysis of photosynthetic protein complexes in indicated strains. After solubilization of membranes with 1.5% (w/v) dodecyl-b-D-maltopyranoside, thylakoid proteins were separated by 5 to 12 % BN gels in the first dimension and 12% SDS-gels in the second dimension. Photosynthetic complexes were detected using the indicated antibodies. The positions of major PSI and PSII complexes are designated. sc: supercomplex

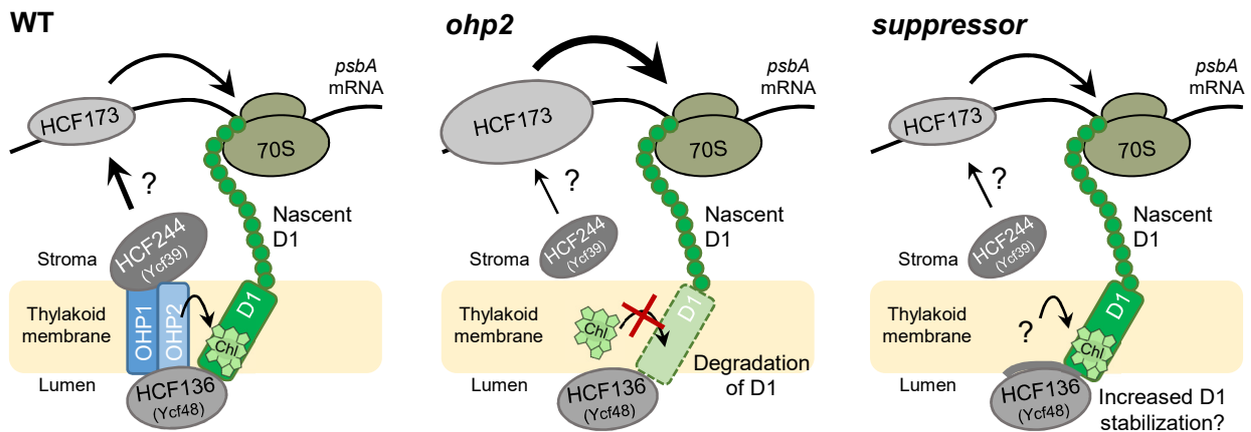


**Figure 4. Targeted ribosome profiling of chloroplast translation reveals protein synthesis defects in the *ohp2* mutant.** Ribosome profiling and transcript analysis of WT and *ohp2* mutant grown mixotrophically under 30  $\mu\text{E}/\text{m}^2/\text{s}$ . a) The average mRNA (blue) and ribosome FP (dark grey) abundances were calculated from three independent biological replicates and plotted in  $\log_{10}$  scale for *ohp2* mutant versus WT, respectively. mRNAs encoding PSII subunits, which display altered translation in the mutant are highlighted. *Pearson's*  $r$ -value and  $p$ -value are given in *nEm* non-superscript format for  $n=10m$ . b) The relative average transcript abundances (RNA), translation output and translation efficiency were calculated for each chloroplast reading frame in both *Chlamydomonas* strains, normalized to overall signal intensities, and plotted as heat map (*ohp2* versus WT) in  $\log_2$  scale. Increased RNA accumulation, translation output or translation efficiency in the mutant is shown in red, reduced levels are shown in blue (see scale bar). PSII subunits that are further discussed are highlighted with an arrow. c) CES cascade of Photosystem II. When newly synthesized CES polypeptides, like D1 and CP47, cannot assemble, they repress the translation initiation of their encoding mRNA as described before by Minai and coworkers (2006). Reduced translation of *psbH* may indicate that it represents a further component of the CES cascade. d) For the Open Reading Frames (ORFs) of the *psbB* operon and *psbA* (introns not shown), normalized ribosome FP intensities were plotted as mean  $\log_2$  ratios between *ohp2* mutant and WT. Error bars denote differences between three independent biological replicates. Grey bars below the panels indicate respective ORFs with trans-membrane segments (TMS in green). Related to Supplemental Figures S7-S9 and Supplemental Dataset 1.



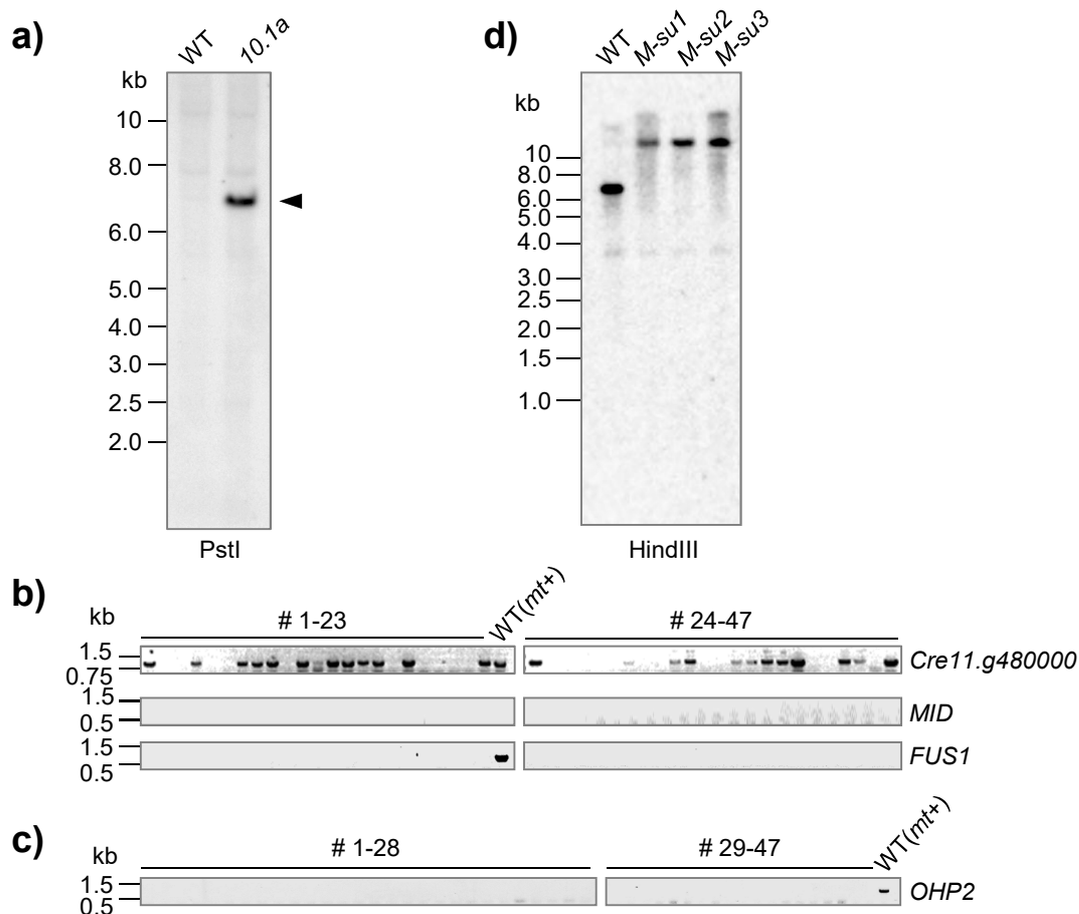


**Figure 6. Mutation of the FtsH protease partially restores PSII activity but not photoautotrophy.** a) Growth tests of the progeny of a cross *10.1a X ftsH1-1*. Cells were grown on TAP or MIN medium at the indicated light intensity. Spots shown are typical of the indicated genotypes. b)  $F_v/F_m$  values for the four genotypes, recorded from the plates in a). Values are average of 8, 7, 11 and 6 strains for the WT, *ftsH1*, *ohp2* and *ohp2 ftsH1* genotypes, respectively. Error bars represent S.D. c) Typical fluorescence induction curves for the four genotypes, recorded with the fluorescence camera on the TAP plates described in a). Fluorescence is normalized to the  $F_m$  value (arrow indicates saturating flash).

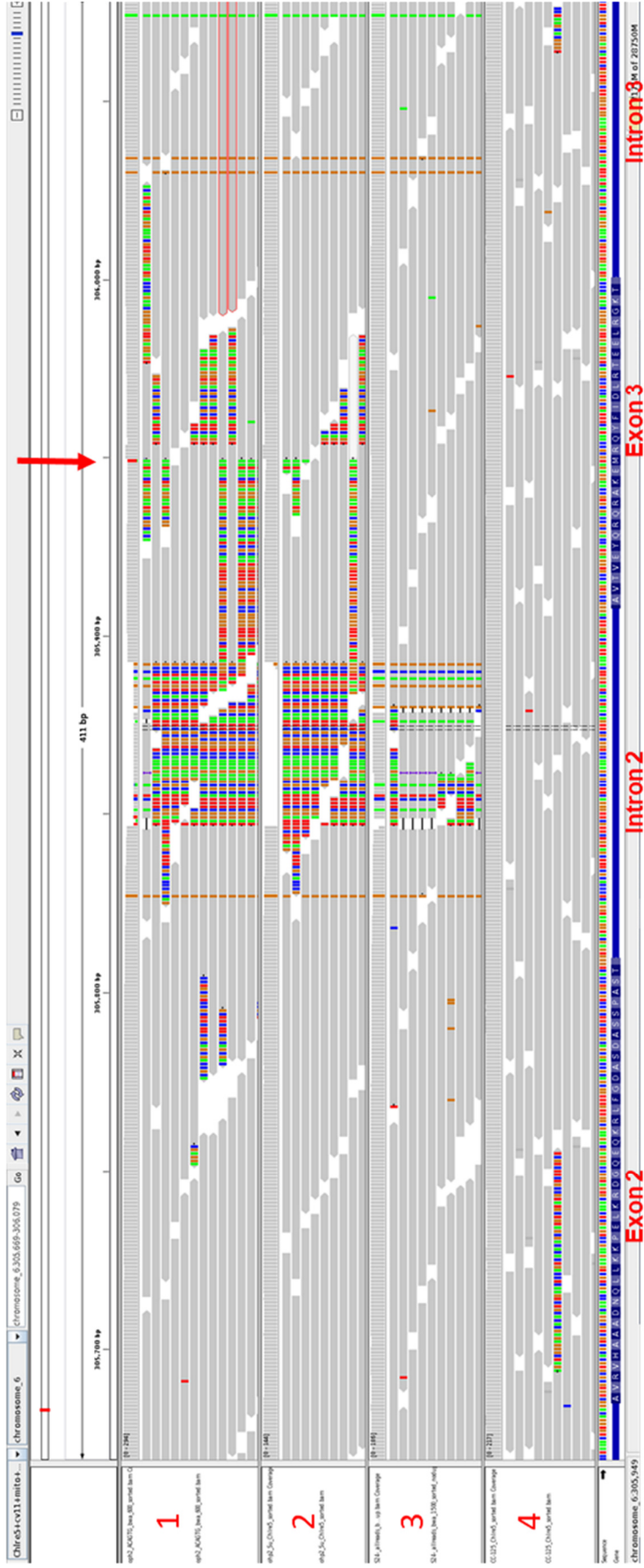


**Figure 7. Schematic model for the role the OHC complex and associated proteins in D1 synthesis and first steps of PSII de novo assembly.** In the WT (left panel), likely triggered by HCF244, HCF173 binds the 5'-UTR of *psbA* mRNA to promote translation initiation. pD1 is co-translationally inserted into the thylakoid membrane. Membrane associated luminal HCF136 stabilizes pD1 and is involved in RC assembly. The OHC is required for early PSII assembly steps and proposed to insert Chl into pD1 proteins. Recently reported negative autoregulatory circuits of *psbA* translation initiation involving the OHC are not displayed here (Chotewutmontri & Barkan, 2020; Chotewutmontri et al., 2020). In *ohp2* (middle panel) and suppressor strains (right panel) residual levels of HCF244 may promote *psbA* translation via interaction with HCF173. Ongoing *psbA* translation may be further supported by overaccumulation of HCF173. However, in the *ohp2* mutant synthesized D1 is rapidly degraded and does not accumulate in the absence of the OHC complex. In the suppressor strains, stabilization of nascent D1 may be accomplished by increased membrane association of HCF136 or other mechanisms to allow assembly of early PSII intermediates. HCF173, HCF136, and HCF244 protein amounts detected by proteomics are indicated by differently sized ovals. Names of cyanobacterial homologs are given in brackets.

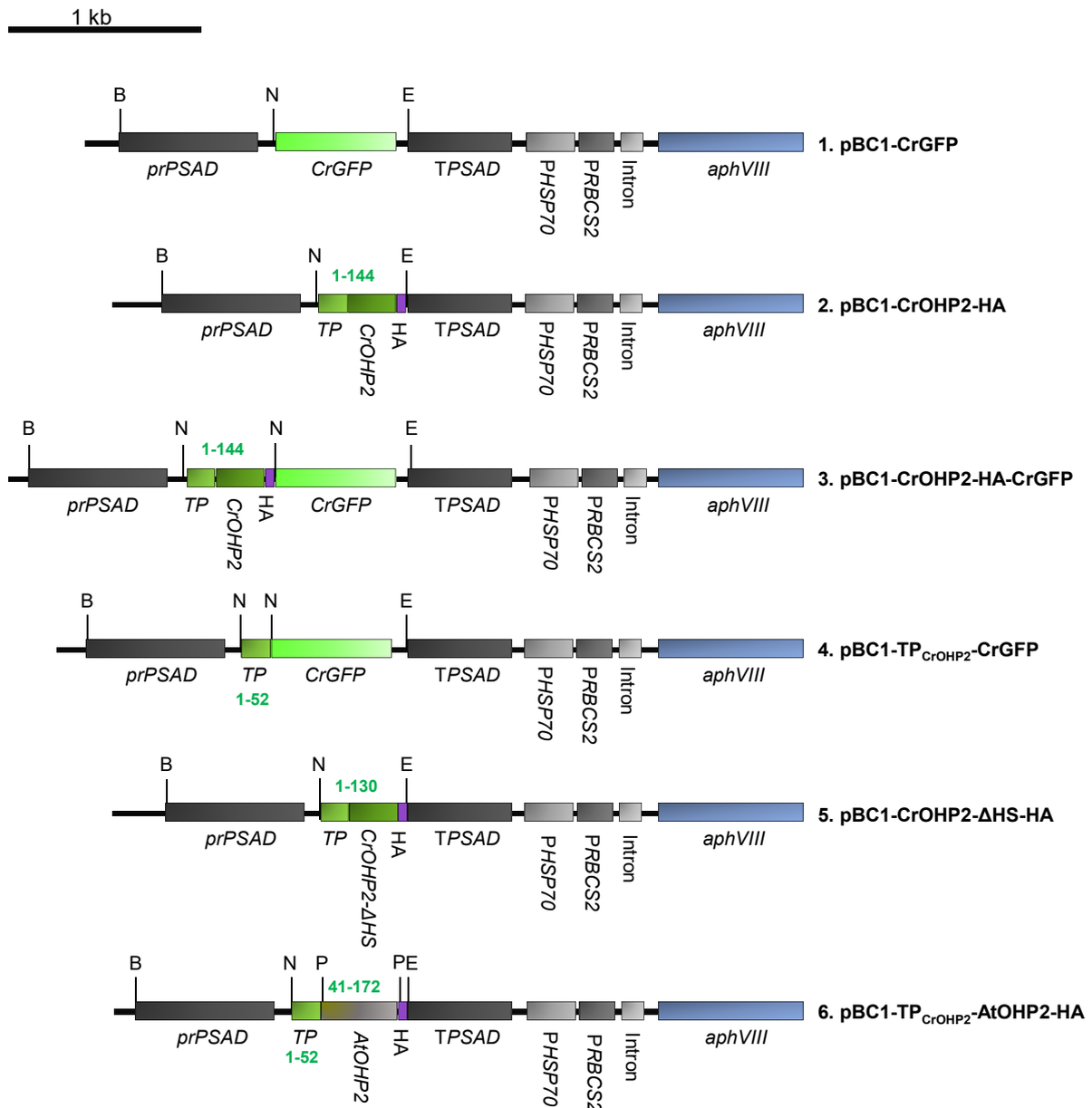
# **Supplemental Figures**



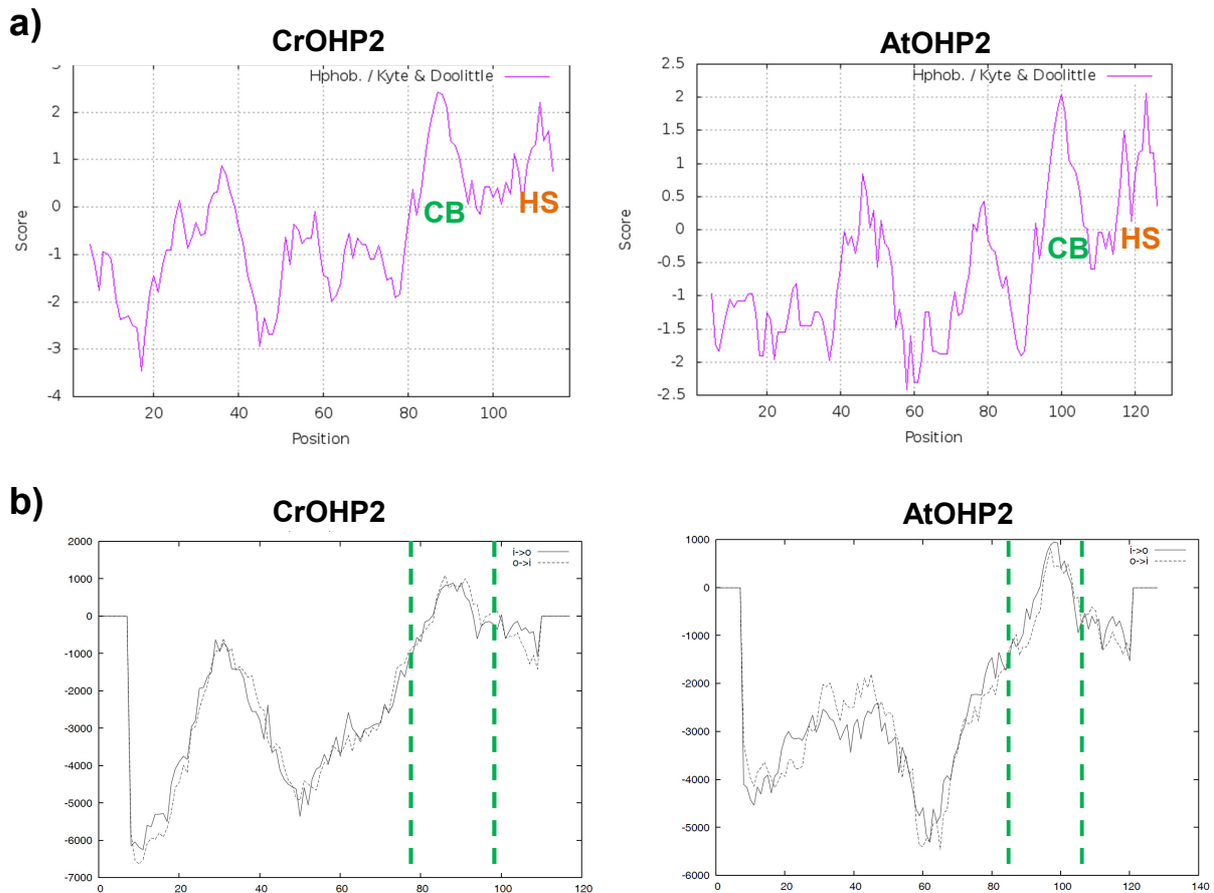
**Figure S1. Genetic analysis of the *10.1a* mutant and suppressor strains.** a) Southern blot analysis indicates a single insertion of the mutagenic vector pBC1 in the *10.1a* mutant. 10 µg of genomic DNA of the wild type Jex4 (WT) and the mutant strain *10.1a* were fractionated in an 0.8% agarose gel after digestion with PstI. The gel was blotted onto a nylon membrane, and hybridized with a dig-labelled probe specifically detecting the *aphVIII* gene. Molecular size markers are labeled at the left (M). Primers used to generate the probe were pBC1-APHV-fw / pBC1-APHV-rev. b), c) Genotyping of 47 non-photosynthetic clones resulting from a backcross of the *10.1a* mutant to the WT strain WT-S34 mt+. b) Analysis of PCR products using primers specific for the locus *Cre11.g480000* and mating type specific loci *MID* and *FUS1*. Upper panel: A *Cre11.g480000*-specific product of 829 bp is expected for the wild type. For clones possessing the insertion of the Pm-resistance cassette no product is expected due to the loss of the primer binding site. Only approximately half of the PSII deficient clones tested exhibited the identified insertion of the resistance cassette. The middle and lower panels show the detection of mating type specific *MID* (mt-) and *FUS1* (mt+) genes. Primers applied for PCR reactions were as follows: *Cre11.g480000* (hit2fw / hit2rev); *MID* and *FUS1* genes (mid-fw/mid-rev; fus1-fw/fus1-rev). Molecular size markers are indicated on the left. c) Analysis of PCR products using primers specific for the *OHP2* gene (*Cre06.g251150*). A product of the expected size could only be detected in the WT but not in the 47 non-photosynthetic clones investigated in b). Position of OHP1-fw2 (fw)/OHP1-rev2 (rev) primers used is indicated in Figure 1a. d) Southern blot analysis of genomic DNA from the Jex4 recipient strain (WT) and three independent *ohp2* suppressor mutants (*M-su1* to *M-su3*). 10 µg of genomic DNA were fractionated in a 0.8% agarose gel after digestion by HindIII, blotted onto a nylon membrane, and hybridized with the *OHP2*-specific probe indicated in Figure 1a. The signal pattern of all suppressor mutants analyzed, resembles that of the *ohp2* mutant indicating that they still harbor the large insertion of the putative *TOC1* transposon (compare strain *10.1a* in Figure 1b). For primer sequences see Supplemental Table S2.



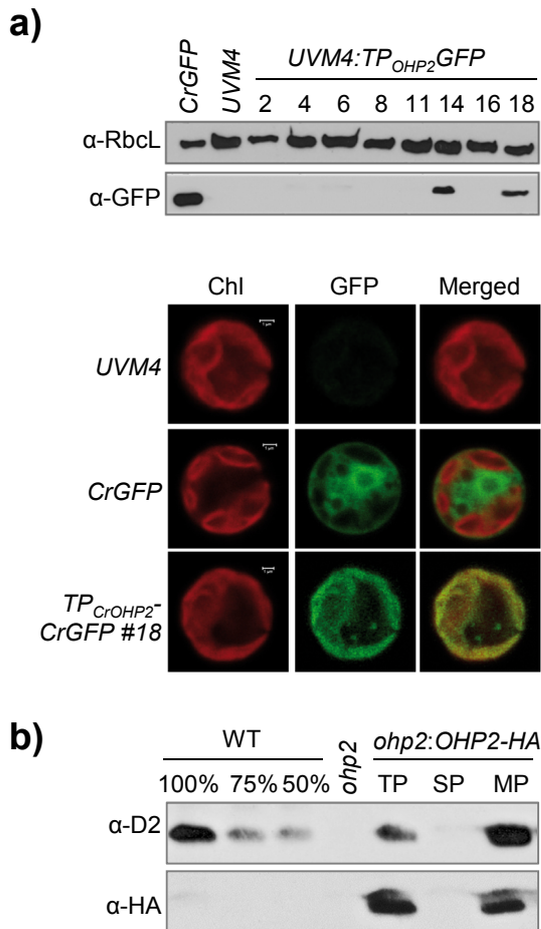
**Figure S2. Identification of the *ohp2* mutation.** Snapshot from the IGV browser showing the second and third exons of *OHp2* with mapping of genomic Illumina reads from (top to bottom) *ohp2* (1), a suppressed *ohp2* mutant (2) and the 137c WT strains WT S24- (CC-5100, Gallaher et al. 2015) (3) and CC-125 (4). The gene structure ranging from exon 2 to intron 3 is indicated in the bottom panel. Note presence of multiple polymorphisms common to all the strains from the Paris lab (top 3), including a major variant in intron 2. The *TOC1* insertion in *OHp2* is indicated by a red arrow.



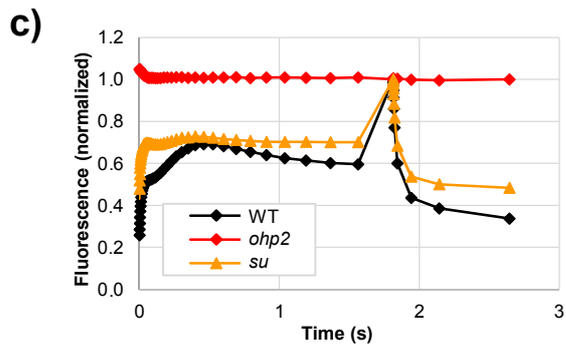
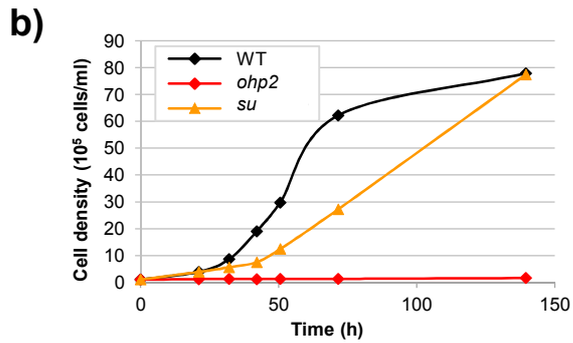
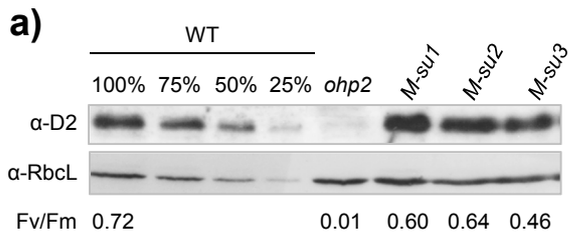
**Figure S3. DNA constructs used for complementation and localization studies.** Illustrations show the expression cassette of the pBC1-CrGFP vector used for cloning in 5' → 3' direction (= pJR38, Neupert et al. 2009). Sequences of interest were inserted downstream of the strong nuclear *PSAD* promoter (*prPSAD*) via *NdeI* or *NdeI/EcoRI* restriction sites. Pm-resistance "*aphVIII*" is under the control of a fusion promoter consisting of the promoter of the gene encoding the small subunit of Rubisco (*PRbcS2*) and the gene encoding the heat shock protein 70A (*PHSP70A*). *CrOHP2* was amplified from cDNA generated from the WT strain Jex4. All constructs contain the sequence encoding the *CrOHP2* transit peptide (TP) predicted by TargetP1.1 (Emanuelsson et al., 2007) to guarantee proper localization of the expressed proteins. Numbers given in green represent amino acid ranges encoded by the respective DNA insert. Important restriction sites are indicated (B: *BamHI*, N: *NdeI*, E: *EcoRI*, P: *PstI*). Primers used for cloning are given in Supplemental Table S2. For cloning details of the synthetic *Arabidopsis* gene in construct 6, see Supplemental Data S2.



**Figure S4. Hydrophobicity prediction for *Chlamydomonas* and *Arabidopsis* OHP2 amino acid sequences.** Predictions were performed for proteins lacking the transit peptides predicted by TargetP 2.0 (*C. reinhardtii*: CrOHP2 aa 27-144; *A. thaliana*: AtOHP2 aa 44-172) **a) Hydrophobicity plot.** Indicated is the C-terminal chlorophyll-binding region (CB) and a hydrophobic stretch (HS). Hydropathy was predicted using the *ProtScale* online software (<https://web.expasy.org/protscale/>) based on the model from Kyte et al. (1982). X-axis: position of amino acids; Y-axis: hydrophobicity value. **b) Prediction of membrane spanning segments.** The region between the green dashed lines is predicted to represent membrane-spanning regions (TMPred; Hofmann and Stoffel 1993; [https://embnet.vital-it.ch/software/TMPRED\\_form.html](https://embnet.vital-it.ch/software/TMPRED_form.html)). The X-axis represents the position of amino acids, the ordinate the TMPred score. Positive scores suggest hydrophobic regions.

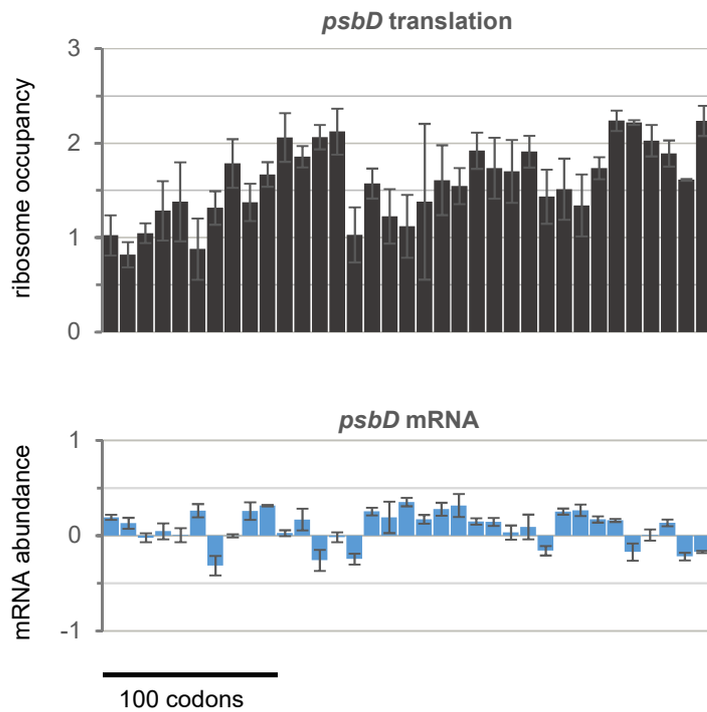


**Figure S5. OHP2 is a membrane localized chloroplast protein. a) GFP localization studies.** Upper Panel: Immunoblot analysis of *UVM4* strains transformed with pBC1-TP<sub>CrOHP2</sub>-CrGFP (construct 4, Supplementary Figure S3). 20  $\mu$ g of total proteins of eight Pm-selected transformants (*UVM4:TP<sub>OHP2</sub>-GFP*) were analyzed for GFP accumulation ( $\alpha$ -GFP) along with the untransformed recipient strain *UVM4* as negative control. 5  $\mu$ g total protein of pBC1-CrGFP (construct 1, Supplementary Figure S3) transformed *UVM4* cells previously shown to accumulate GFP in the cytosol (*CrGFP*) served as positive control (Jalal et al., 2015). The protein detected in *UVM4:TP<sub>OHP2</sub>-GFP* transformants migrated slightly slower than the signal detected in the strain expressing the unfused 27 KDa cytosolic CrGFP, suggesting partial retention of the fused transit peptide. RbcL was used as loading control ( $\alpha$ -RbcL). Lower panel: The GFP expressing clone #18 was used for GFP localization studies (*TP<sub>OHP2</sub>-CrGFP*, clone #18) by laser scanning confocal fluorescence microscopy using a TCS-SP5 system (Leica). The recipient strain *UVM4* as well as *CrGFP* served as controls. Chlorophyll auto fluorescence (Chl), GFP fluorescence (GFP), and the merged signals of *Chlamydomonas* cells are shown. Bar = 1  $\mu$ m. **b) Cell subfractionation.** Immunoblot analysis of cell fractions from the complemented strain *ohp2:OHP2-HA* clone #53 with a D2 (upper panel) or an HA specific antibody (bottom panel). 10  $\mu$ g of total (TP) and fractionated into soluble (SP) and membrane (MP) proteins of the complemented strain, as well as *ohp2* as negative control were analyzed. A dilution series of the wild type Jex4 (WT) served as reference.

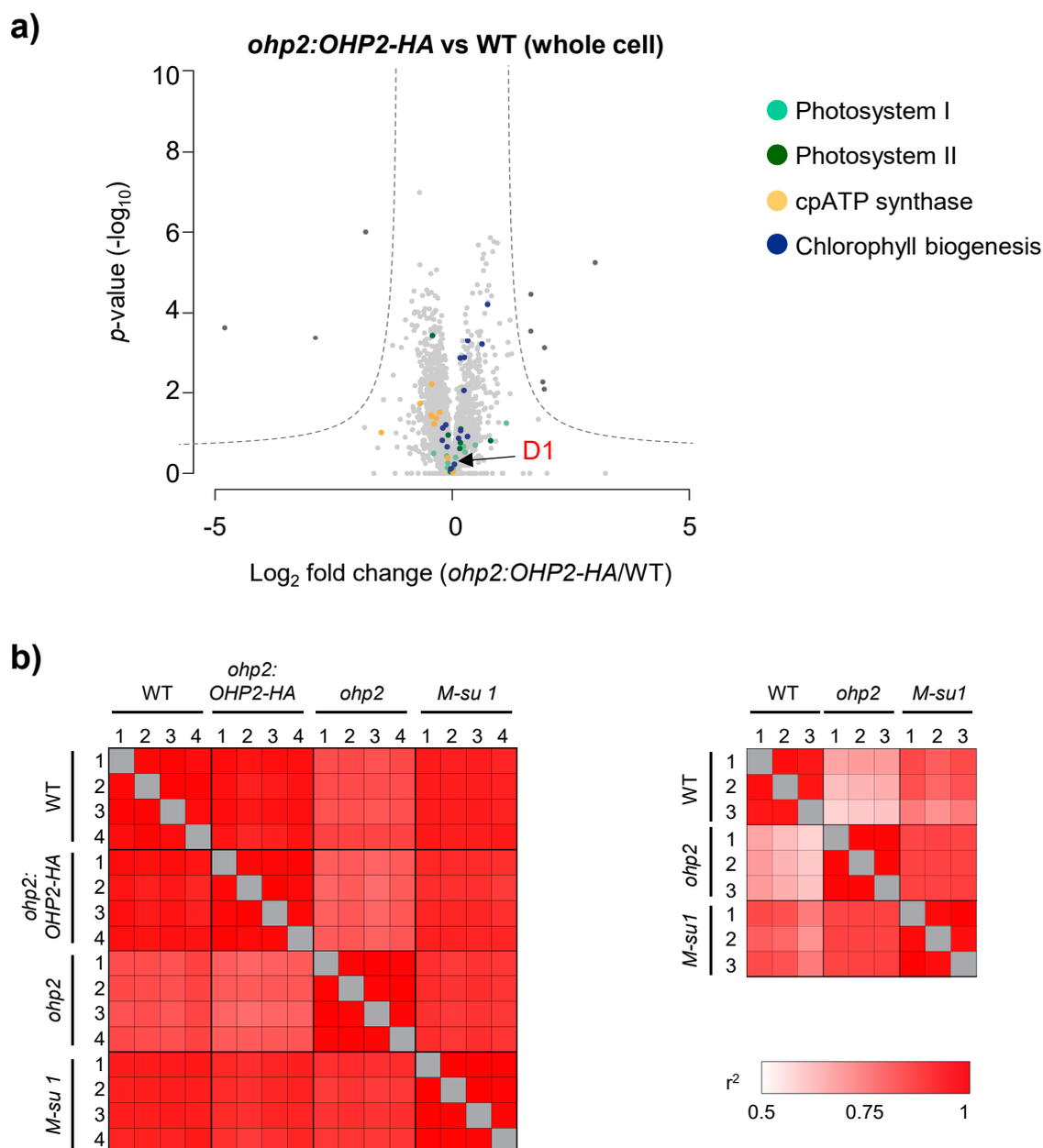


**Figure S6. Re-accumulation of D2 and restoration of photoautotrophy in suppressor mutants.** **a) Western blot.** 20  $\mu$ g of whole cell proteins of three suppressor mutants (*M-su1-3*), as well as the *ohp2* strain and a dilution series of the wild type Jex4 (WT) were immunodecorated with a D2 specific antibody. RbcL served as a loading control. To measure Fv/Fm ratios given below the panel, cells were resuspended in H<sub>2</sub>O up to a concentration of 10<sup>5</sup> cell/mL and 10  $\mu$ L were spotted onto TAP plates and grown for 7d under low light at 30  $\mu$ E/m<sup>2</sup>/s. **b) Growth curves under photoautotrophic conditions at 60  $\mu$ E/m<sup>2</sup>/s and c) fluorescence induction kinetics** measured under illumination at 135  $\mu$ E/m<sup>2</sup>/s of the WT, *ohp2*, and a suppressor strain (*su*).

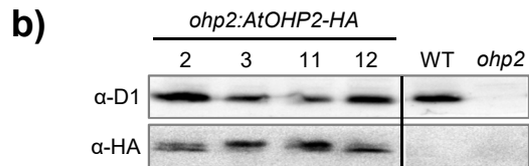
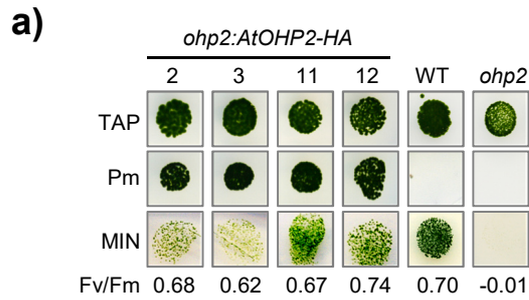




**Figure S8. Targeted ribosome profiling of chloroplast translation reveals enhanced *psbD* translation in the *ohp2* mutant.** For the Open Reading Frame of *psbD*, normalized ribosome footprint intensities were plotted as mean  $\log_2$  ratios between *ohp2* mutant and wild type. Error bars denote differences between three independent biological replicates.



**Figure S9. Complementation of *ohp2* and heat maps representing the reproducibility of LC-MS experiments.** **a) Complementation of *ohp2* with HA-tagged OHP2 fully reverts photosynthesis-deficient phenotype.** Volcano plots represent the relative whole cell proteome changes comparing lysates of *ohp2:OHP2-HA* versus wild type strains. All experiments were performed in four independent biological replicates. Mean fold change of LFQ values (in  $\log_2$ ) is plotted on the x-axis,  $p$ -values (in  $-\log_{10}$ ) are plotted on the y-axis. Light grey dots represent proteins with no significant change, dark grey dots show proteins that are significantly different with  $FDR < 0.05$  and  $S_0=1$ . Proteins of the PSI, PSII, the chloroplast ATP synthase and of proteins involved in chlorophyll biogenesis are marked in color. Large colored dots are significantly different. **b) Heat maps representing correlations between experiments of LC-MS of whole cell (left) and membrane fractions (right).** Correlations are given as  $r$ -squared LFQ values (Supplemental Datasets 2, 3). Proteins were identified through LC-MS as described in Methods.



**Figure S10. Complementation of the *ohp2* mutant with the orthologous *Arabidopsis* protein restores photoautotrophy and D1 accumulation.**  
**a) Growth test.** The *ohp2* mutant was transformed with the construct pBC1-TP<sub>CrOHP2</sub>-AtOHP2-HA (see construct 6 in Supplemental Figure S3). Transformants were first selected on paromomycin (Pm) and then tested for photoautotrophic growth on HSM plates (MIN). Four representative clones out of 20 are shown. Fv/Fm values are indicated below the panel. **b) Immunoblot analysis** of Pm-resistant transformants. 20  $\mu$ g of total proteins were separated on 15% denaturing polyacrylamide gels and analyzed with the antibodies indicated on the left. Jex4 (WT) and *ohp2* were run on the same gel but not in adjacent lanes as indicated by a vertical black line.



# **Supplemental Tables, Data and References**

## SUPPLEMENTAL TABLES, DATA AND REFERENCES

**Supplemental Table S1.** Photosynthetic parameters and Chl composition of *Chlamydomonas* WT Jex4 (WT), the *ohp2* mutant and *ohp2* complemented with OHP2-HA (construct 2, Supplemental Figure S3; strain *ohp2:OHP2-HA*, clone #53). Photosynthetic parameters were determined as described in Experimental Procedures. Chl determination was performed according to Arnon (1949).

Strain	WT	<i>ohp2</i>	<i>ohp2:OHP2-HA</i>
<b>Photosynthetic parameters:</b>			
Fluorescence: Fv/Fm	0.77	-0.04	0.79
ECS: PSII/PSI	1.40	0.05	1.50
ECS: phase b/phase a	0.40	0.47	0.50
<b>Chl determination (n= 3):</b>			
Chl <i>a</i> (pg/cell)	4.17 ± 0.12	2.31 ± 0.11 (55%)	3.32 ± 0.27 (80%)
Chl <i>b</i> (pg/cell)	2.25 ± 0.12	1.44 ± 0.08 (65%)	1.70 ± 0.10 (75%)
Chl <i>a + b</i> (pg/cell)	6.65 ± 0.24	3.75 ± 0.19 (58%)	5.00 ± 0.37 (78%)
Chl <i>a/b</i>	1.85	1.61	1.95

**Supplemental Table S2:** Primers used in this study.

Restriction sites in lower case letters. Sequence coding for an HA-tag is underlined.

Name	Sequence (5'→3')	Application
pBC1-APHV-fw	AGACTGCGATCGAACGGACA	Southern blot for detection of the <i>aphVIII</i> cassette
pBC1-APHV-rev	GCTGAAGAATTCGAAGCATGGAC	
hit2fw (Cre11.g480000)	ATTAACAGCACCTGCAACTACCA	Detection of gene <i>Cre11.g480000</i>
hit2rev (Cre11.g480000)	AACGTGATGCTGCCGTCATAAGGT	
mid-fw	ATGGCCTGTTTCTTAGC	Detection of mating type minus specific gene ( <i>MID</i> )
mid-rev	CTACATGTGTTTCTTGACG	
fus1-fw	ATGCCTATCTTTCTCACTTCT	Detection of mating type plus specific gene ( <i>FUS1</i> )
fus1-rev	GCAAATACACGTCTGGAAG	
OHP1-fw2 (fw)	GTGCACACGTATCACTTCTG	Screening of backcross progeny (OHP2), Southern blot
OHP1-rev2 (rev)	GCACACTTACCAGTCAGCAT	
OHP2-rev HA ( <i>NdeI</i> )	catatg <u>GGCGTAGTCCGGCACGTCGTACGGGTAGTCCAGGTCCACGATG</u>	Cloning of pBC1-CrOHP2-HA for complementation studies (construct 2, Supplemental Figure S3)
OHP2-rev HA ( <i>EcoRI</i> )	gaattcTTAG <u>GGCGTAGTCCGGCACGTCGTACGGGTAGTCCAGGTCCACGATG</u>	
OHP1-fw4 ( <i>NdeI</i> )	catatgTCGATTGCTGCACTCCG	Cloning of pBC1-CrOHP2-HA-CrGFP for complementation studies (construct 3, Supplemental Figure S3)
OHP2-rev HA ( <i>NdeI</i> )	catatg <u>GGCGTAGTCCGGCACGTCGTACGGGTAGTCCAGGTCCACGATG</u>	
OHP1-fw4 ( <i>NdeI</i> )	catatgTCGATTGCTGCACTCCG	Cloning of pBC1-TP <sub>CrOHP2</sub> -CrGFP for localization studies (construct 4, Supplemental Figure S3)
OHP2-TP rev ( <i>NdeI</i> )	catatgGGCGTCGCCAAACAACCGCTTC	
OHP1-fw4 ( <i>NdeI</i> )	catatgTCGATTGCTGCACTCCG	Cloning of pBC1-CrOHP2-ΔHS-HA for complementation studies (construct 5, Supplemental Figure S3)
OHP2-ΔMA rev ( <i>EcoRI</i> )	gaattcTTAG <u>GGCGTAGTCCGGCACGTCGTACGGGTACTGGTTAATGAAGTCG</u>	
rbcl Fw	AAGATTCAAGCAGCTACAGC	Northern blot analysis
rbcl Rv	CACTGCCTCTAATAAAGTCTAC	
psbD Fw	GCCGTAGGGTTGAATG	Northern blot analysis
psbD Rv	GTTGGTGTCAACTTGGTGG	
psbA Fw	TCTAGCCTATGGGCTCGT	Northern blot analysis
psbA Rv	ACCGAAACGGTAACCTTC	

**Supplemental Data S1. Flanking sequence tags in the *ohp2* mutant obtained by Illumina sequencing.** The *ohp2* mutant shows a putative TOC1 transposon insertion in the third exon of the *OHP2* gene between position chromosome\_6:305949 and 305950. Nucleotides shown in blue correspond to genome sequences (left side: chromosome\_6:305923..305949; right side: 305950..305975) while those in red are identical to LTR sequences described for *TOC1* retrotransposons.

The left side of the insertion can be read based on

>DE18INS652:189:C6M76ACXX:2:2215:12623:4634 Alignment start = 305923 (-) Cigar = 31M69S  
GTACCAGCGCCAGCGTGCCAAGGAGATGCGCTGACTTATCGCAGCGCAGTGGAGATAAGTCTAG  
TTATTGCGACGTAACCTGCCGTGTTGCGTTAGAGTCA

The right side of the insertion can be read based on

>DE18INS652:189:C6M76ACXX:2:1106:3608:11943 Alignment start = 305950 (+) Cigar = 74S26M  
CGGTTTGAGAGATTGGGATTATCGTTTGGGGCCGTGGCGTTTGTAAACGCTACACGGCAGTAAG  
GAGTCAATAAGCGCCAATACTTTATCGACCTCCGGA

**Supplemental Data S2. Synthetic Arabidopsis OHP2 nucleotide sequence and derived protein sequence used to complement the *Chlamydomonas ohp2* mutant strain.**

a) Sequence of codon adapted synthetic gene from *Arabidopsis* ordered from Integrated DNA Technologies (IDT) that was used to clone the construct pBC1-TP<sub>CrOHP2</sub>-AtOHP2-HA (construct 6 in Supplemental Figure S3). In green the *Chlamydomonas* sequence encoding the N-terminal region including the transit peptide (aa 1-52, *Cre06.g251150*) predicted by TargetP-1.0. In cyan, the sequence encoding the *Arabidopsis* protein OHP2 (aa 41-172, *AT1G34000*) adapted to the codon usage of *Chlamydomonas*. The HA-tag encoding sequence is shown in purple letters. Restriction sites introduced for cloning are displayed in lower case letters.

NdeI

catatgTCGATTGCTGCCCTCCGGAGCTCGCCGAGCCTGCCTGCGGTCCGCCACGGCCGC  
AAGGCTGTGCGCGTGCACGCTGCTGCGGACAACCAGCTGCTGAAGAAGCCTGAGCTCAAG

PstI

CGCGAGGGCCAGGAGCAGAAGCGGTTGTTTCGGCGACGCCctgcagATCAGGTGCTCCCAG  
ACCGACGGCCCTTTGAGGAGGCCGTTCGGCTCCTCCTACGCTCCGGGAGCCTCAAAAGCCT  
GTGCCTCCGTGCGAGCCGTCTCCTCGCCTCCTCCTTCGCCGCCCGCAGAAAGGCTGTG  
GCTGTGGAGGGCAAGAGCGTGACGACCGTGGAGTTCCAGAGGCAGAAAGGCTAAGGAGCTC  
CAGGAGTACTTCAAGCAGAAGAAGCTGGACGCTGCTGGCCAGGGCCCTTCTTCGGCTTC  
CAGCCCAAGAACGAGATCTCCAACGGCAGGTGGGCTATGTTTCGGCTTCGCCGTGGGCATG  
CTCACCGAGTACGCCACCGGCTCGGACCTGGTGGAGCAGGTGAAGATCCTGCTCTCGAAC

PstI

TTCGGCATTCTGGACTTGGAGctgcagTACCCGTACGACGTGCCGGACTACGCCTAAgaa

ttc

EcoRI

b) Chimeric protein sequence expressed from *Chlamydomonas* strains transformed with the vector pBC1-TP<sub>CrOHP2</sub>-AtOHP2-HA. Color code as in a). The fusion protein possesses the *Chlamydomonas* N-terminus fused to the C-terminally HA-tagged *Arabidopsis* protein lacking the predicted localization signal.

MSIAALRSSPSLPAVRHGRKAVRVHAAADNQLLKKPELKREGQEQKRLFGDALQIRCSQTDGPLRRPSAPPTLRE  
PQKPVPPSQPSSPPPSPPQKAVAVEGKSVTTVEFQRQKAKELQEYFKQKLDAAAGQGPFQPKNEISNGRW  
AMFGFAVGMLTEYATGSDLVEQVKILLSNFGILDLELQYPYDVPDYA

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