1	Xanthophyll carotenoids stabilise the association of cyanobacterial chlorophyll synthase with the
2	LHC-like protein HliD

3

Matthew S. Proctor^a, Marek Pazderník^{b,e}, Philip J. Jackson^{a,c}, Jan Pilný^b, Elizabeth C. Martin^a, Mark J.
Dickman^{a,c}, Daniel P. Canniffe^d, Matthew P. Johnson^a, C. Neil Hunter^a, Roman Sobotka^{b,e} and Andrew
Hitchcock^a

- 7
- ^aDepartment of Molecular Biology and Biotechnology, University of Sheffield, Sheffield S10 2TN,
 United Kingdom
- ^bInstitute of Microbiology of the Czech Academy of Sciences, Centre Algatech, 37981 Třeboň, Czech
 Republic
- ^cDepartment of Chemical and Biological Engineering, University of Sheffield, Sheffield S1 3JD, United
 Kingdom
- ^dInstitute of Systems, Molecular & Integrative Biology, University of Liverpool, Liverpool L69 7ZB,
 United Kingdom
- 16 ^eFaculty of Science, University of South Bohemia, 37005 České Budějovice, Czech Republic
- 17

18 **Authors for correspondence:** Roman Sobotka (sobotka@alga.cz) and Andrew Hitchcock 19 (a.hitchcock@sheffield.ac.uk)

20

Author ORCID IDs: 0000-0002-1484-850X (MSP); 0000-0001-9671-2472 (PJJ); 0000-0002-9236-0788
(MJD); 0000-0002-5022-0437 (DPC); 0000-0002-1663-0205 (MPJ); 0000-0003-2533-9783 (CNH);
0000-0001-5909-3879 (RS); 0000-0001-6572-434X (AH)

24

25 Abstract

Chlorophyll synthase (ChlG) catalyses a terminal reaction in the chlorophyll biosynthesis pathway, attachment of phytol or geranylgeraniol to the C17 propionate of chlorophyllide. Cyanobacterial ChlG forms a stable complex with high light-inducible protein D (HliD), a small single-helix protein homologous to the third transmembrane helix of plant light-harvesting complexes (LHCs). The ChlG- 30 HliD assembly binds chlorophyll, β -carotene, zeaxanthin and myxoxanthophyll and associates with the 31 YidC insertase, most likely to facilitate incorporation of chlorophyll into translated photosystem apoproteins. HliD independently coordinates chlorophyll and β -carotene but the role of the 32 xanthophylls, which appear to be exclusive to the ChIG-HliD assembly, is unclear. Here we generated 33 mutants of Synechocystis sp. PCC 6803 lacking specific combinations of carotenoids or HliD in a 34 35 background with FLAG- or His-tagged ChlG. Immunoprecipitation experiments and analysis of isolated 36 membranes demonstrate that the absence of zeaxanthin and myxoxanthophyll significantly weakens the interaction between HliD and ChIG. ChIG alone does not bind carotenoids and accumulation of the 37 chlorophyllide substrate in the absence of xanthophylls indicates that activity/stability of the 'naked' 38 enzyme is perturbed. In contrast, the interaction of HliD with a second partner, the photosystem II 39 40 assembly factor Ycf39, is preserved in the absence of xanthophylls. We propose that xanthophylls are required for the stable association of ChIG and HliD, acting as a 'molecular glue' at the lateral 41 42 transmembrane interface between these proteins; roles for zeaxanthin and myxoxanthophyll in ChIG-HliD complexation are discussed, as well as the possible presence of similar complexes between LHC-43 like proteins and chlorophyll biosynthesis enzymes in plants. 44

45

46 Introduction

47 Carotenoids are isoprenoid pigments that are categorized into two main classes, xanthophylls, which contain oxygen, and carotenes, which do not. In chlorophototrophs carotenoids are important for light 48 49 harvesting, photoprotection and structural stabilization of proteins and membranes (1, 2). For example, the trimeric photosystem I (PSI) from Synechocystis sp. PCC 6803 (hereafter Synechocystis) 50 contains 72 carotenoids and is dependent upon β -carotene for trimerization (3-6). β -carotene is also 51 52 a cofactor in photosystem II (PSII) and is required for its assembly (7, 8), is present in the cytochrome 53 $b_{6}f$ complex (9, 10) and photosynthetic complex I (11), and plays a role in the assembly of the 54 phycobilisome antenna complex (5). In plants, xanthophylls play important roles in structural 55 stabilization of light-harvesting complexes (LHCs), light harvesting and photoprotection, and act as lipid-soluble antioxidants (12-14). However, the role of xanthophylls in cyanobacteria, where they do 56 57 not participate in light harvesting, is less obvious. There is a pool of free xanthophylls in cyanobacterial 58 membranes that modulates membrane rigidity/fluidity (15, 16) and it is widely accepted that these 59 pigments provide protection against photooxidative stress and reactive oxygen and nitrogen species, especially under high light conditions (17, 18). Although not integral components of either 60 photosystem, xanthophylls stabilize oligomers of PSI and PSII (5, 19), while keto-xanthophylls are 61

specifically required for non-photochemical dissipation of excess energy by the orange carotenoidprotein (20).

64 Chlorophyll (Chl) synthase (ChlG) is an integral thylakoid membrane protein that catalyses the addition of a geranylgeranyl or phytyl tail to the chlorophyllide (Chlide) macrocycle as one of the terminal steps 65 66 of Chl biosynthesis (Figure 1A). In Synechocystis, tagged ChlG co-purifies with the carotenoids β carotene, zeaxanthin and myxoxanthophyll in a pigment-protein complex also containing Chl, high 67 light-inducible proteins (Hlips), the membrane insertase YidC and the PSII assembly factor Ycf39 (21, 68 69 22). ChIG binds tightly to HliD in a ChIG-HliD 'core', with YidC and Ycf39 present in sub-stoichiometric 70 amounts (21). Hlips are single helix transmembrane pigment-binding proteins that are thought to be the ancestors of LHCs in eukaryotic phototrophs (23). Synechocystis contains four Hlips (HliA-D), which 71 72 appear to be primarily involved in Chl biosynthesis/recycling and biogenesis and photoprotection of 73 Chl-binding proteins (24), although their exact roles remain enigmatic. An Hlip domain is also found fused to the C-terminus of cyanobacterial and plant ferrochelatases (25). YidC assists in the integration 74 75 of translated proteins into the membrane bilayer (26) and its association with ChIG is presumed to 76 facilitate insertion of Chl molecules into newly synthesized Chl-binding proteins (21). The function of 77 Ycf39 in the ChIG-HliD complex is unclear but it could play a regulatory role in re-modelling ChIG-Hlip assemblies in response to stress (27); Ycf39 and HliD interact in a separate complex that promotes the 78 79 synthesis and assembly of the core PSII subunits D1 and D2 during exposure to high light (28), 80 conditions under which Ycf39 dissociates from the ChIG complex (27) and HliD appears to be partially 81 replaced by HliC (22).

The approximate molar ratio of pigments in the ChIG complex is ChI (6): zeaxanthin (2.1-2.7): β -82 83 carotene (1): myxoxanthophyll (0.6-1) (21, 22). HliD likely binds to ChIG as a dimer as the predicted structure of Hlips shows that they cannot bind pigments as monomers (29); purified HliD dimers bind 84 85 6 Chl a molecules and 2 β -carotenes (22). Ultrafast transient absorption spectroscopy indicates that one of the β -carotenes in the HliD adopts a 'twisted' configuration that can quench excited states of 86 87 Chl, resulting in the safe dissipation of excitation energy as heat (22, 30, 31), leading to the suggestion 88 that HliD photoprotects Chl-binding proteins and Chl-biosynthesis enzymes (24, 28, 30, 31). Given that 89 neither isolated HliD nor the Ycf39-HliD complex binds xanthophylls (22; 28, 30), and specific removal 90 of Ycf39 does not alter the pigment composition of the larger ChIG complex (27), the presence of 91 zeaxanthin and myxoxanthophyll appears to be strictly dependent on interaction of ChIG and HliD, but their functional roles are unknown. 92

To investigate the requirement of xanthophylls in the ChIG-HliD complex, we generated a series of
 Synechocystis strains lacking either combinations of xanthophylls or HliD and performed

immunoprecipitations using tagged ChIG as bait. Our results demonstrate that zeaxanthin is required
for stable formation of the ChIG-HIID complex, both *in vitro* and *in vivo*. ChIG alone does not co-purify
with carotenoids and its function appears to be perturbed in the absence of the xanthophyll-mediated
association with HIID. Possible roles of HIID in ChI trafficking and photosystem assembly/repair, and
candidates which may perform analogous functions in higher phototrophs, are discussed.

100

101 Materials and methods

102 Growth of Synechocystis and strain generation

Synechocystis was grown at 30 °C with moderate light (30-50 µmol of photons m⁻²s⁻¹) in BG11 medium 103 104 (32) supplemented with 10 mM TES (Sigma Aldrich)-KOH pH 8.2 (BG11-TES). Liquid cultures were 105 shaken at approximately 150 rpm. For growth on plates, BG11-TES was supplemented with 1.5% (w/v) 106 agar and 0.3% (w/v) sodium thiosulphate. Antibiotics were included where appropriate (as detailed below). Cultures for purification of protein complexes were grown photoautotrophically with ~100 107 µmol photons m⁻²s⁻¹ illumination in 8 L vessels which were mixed by bubbling with sterile air and 108 maintained at 30 °C using a temperature coil connected to a thermostat-controlled circulating water 109 110 bath.

111 All reported mutant strains were prepared in the Synechocystis WT-P (WT) substrain (33); the FLAGchlG Δ chlG (FG/ Δ G) strain generated in this background has been reported previously (27). To 112 generate a strain producing 10xhistidine-tagged ChIG, the Ndel-Bg/II fragment encoding the 3xFLAG-113 114 tagged chIG in pPD-NFLAG::chIG (21) was replaced with sequence encoding 10xhistidine-tagged chIG 115 (synthesised as a gBLOCK by Integrated DNA Technologies; see Table S1 for sequence) and the resulting allele exchange construct (pAH97) was introduced into WT Synechocystis with selection and 116 117 segregation on kanamycin, as detailed below. The zeocin resistance mutagenesis construct described by Chidgey et al. (21) was used to delete the native chIG gene (slr0056) from the His-chIG strain, 118 119 generating the strain His-*chlG* Δ *chlG* (HG/ Δ G).

To generate a *crtR* null mutant, a linear mutagenesis construct was generated by OLE-PCR to replace the central 555 bp of the 939 bp gene (sll1468) with an erythromycin resistance cassette by allele exchange. Similar constructs were generated to replace 568 bp of the 912 bp *cruF* gene (sll0814) with the *aadA* gene (streptomycin resistance) from pCDFDuet-1 (Novagen), 710 bp of the 1629 bp *crtO* gene (slr0088) with the chloramphenicol acetyl transferase (*cat*) from pACYC184 (NEB) or 722 bp of the 1185 bp *cruG* gene (sll1004) with *aadA*. The erythromycin resistance mutagenesis construct described by Xu et al. (34) was used to delete *hliD* (ssr1789). Linear DNA constructs were introduced to *Synechocystis* by natural transformation and transformants were selected on BG11 agar with 7.5 μg ml⁻¹ kanamycin, 7.5 μg ml⁻¹ erythromycin, 5 μg ml⁻¹ streptomycin, 12.5 μg ml⁻¹ chloramphenicol or
2.5 μg ml⁻¹ zeocin, as appropriate. Segregation of genome copies was achieved by sequential plating
with increasing antibiotic concentration up to 20 μg ml⁻¹ (for zeocin), 30 μg ml⁻¹ (for
erythromycin/streptomycin/kanamycin) and 50 μg ml⁻¹ (for chloramphenicol) and was confirmed by
colony PCR. All primers used to generate constructs and screen *Synechocystis* strains/mutants are
provided in **Table S1**.

134

135 Purification of FLAG-tagged and His-tagged ChlG

136 FLAG-immunoprecipitations were performed as reported in our previous work (21, 27). Synechocystis cultures were grown to an OD₇₅₀ of ~0.7-1.0, harvested by centrifugation (17 700xg, 4 °C, 20 min), 137 resuspended in binding buffer (25 mM sodium phosphate pH 7.4, 10 mm MgCl₂ and 50 mm NaCl, 10% 138 139 (w/v) glycerol and EDTA-free Protease Inhibitor [Roche]) and lysed in a Mini-Beadbeater-16. The lysed 140 cells were collected from atop the glass beads and thylakoid membranes were pelleted by centrifugation (48 400xg, 4 °C, 30 min) and solubilised by incubation with 1.5% (w/v) n-dodecyl- β -D-141 142 maltoside (β-DDM; Anatrace) at 4 °C for 1 h. Following centrifugation (48 400xg, 4 °C, 30 min) to pellet 143 insoluble debris, the solubilised thylakoid fraction (the supernatant) was diluted 2-fold and applied to 144 a 300 µL anti-FLAG-M2 agarose (Sigma-Aldrich) column equilibrated in wash buffer (binding buffer with 0.04% (w/v) β -DDM). The resin was washed with 20 resin volumes of wash buffer to remove 145 contaminating proteins and the FLAG-tagged bait protein and associated interaction partners were 146 147 eluted in 400 µL of the same buffer containing 187.5 µg mL⁻¹ 3xFLAG peptide (Sigma-Aldrich). The 148 mixture was filtered through a 0.22 µm spin column (Sigma-Aldrich) to separate the resin from the 149 eluted protein. Eluates were analysed immediately or stored at -80 °C.

For purification of His-ChIG (H.ChIG), solubilised thylakoids were applied three times to a Ni²⁺ NTA 150 151 Agarose (Qiagen) immobilised metal affinity chromatography column that had been pre-equilibrated 152 in binding buffer (as above) supplemented with 5 mM imidazole. Approximately 250 μ L of resin was used per 8 L of cell culture from which the membrane fraction was isolated. The column was washed 153 with 20 column volumes of binding buffer followed by washes with binding buffer containing 154 progressively higher (20, 50 and 80 mM) imidazole concentrations, each containing 0.04% (w/v) β -155 DDM. H.ChIG was eluted by incubation of the resin with 400 µL of elution buffer (binding buffer 156 157 containing 400 mM imidazole and 0.04% (w/v) β -DDM) for 1 h with gentle agitation at 4 °C before the 158 solution was filtered through a 0.22 µm spin column, separating the resin from the eluted protein. 159 Eluates were analysed immediately or stored at -80 °C.

160

161 Pigment analysis by reverse phase high performance liquid chromatography (RP-HPLC)

Pigments were extracted from cell pellets or FLAG-/His-eluates in 100% methanol and separated by RP-HPLC on an Agilent 1200 HPLC system using a Discovery HS C₁₈ column (5 μ m, 250 × 4.6 mm) according to the slightly modified method of that of Largarde and Vermaas (35) described in Proctor et al. (27). Absorbance was monitored at 450 nm and 665 nm and carotenoid species and Chl *a* were identified by their known absorption spectra (**Figure S1**) and retention time (**Figures S2** and reference 27).

168

169 Quantitative proteomic analysis

Proteins were extracted from the FLAG eluates by precipitation using a 2-D clean-up kit (GE 170 171 Healthcare) and processed according to Hitchcock et al. (36) to generate tryptic peptides. Analysis was performed by nano-flow reverse phase chromatography coupled to a mass spectrometer using system 172 173 parameters described by MacGregor-Chatwin et al. (37) with the exception that peptides were 174 resolved with a 75 min gradient in this study. Proteins were identified and quantified using MaxQuant 175 v.1.5.3.30 **Synechocystis** (38) to search а proteome database 176 (http://genome.microbedb.jp/cyanobase/).

177

Biochemical Journal. This is an Accepted Manuscript. You are encouraged to use the Version of Record that, when published, will replace this version. The most up-to-date-version is available at https://doi.org/10.1042/BCJ20200561

178 Protein electrophoresis and immunoblotting

179 Protein electrophoresis was performed as reported in our previous work (21, 27). Proteins in the FLAG-180 and His-eluates were separated by SDS-PAGE on Invitrogen 12% Bis-Tris NuPage gels (Thermo Fisher 181 Scientific) and visualised by staining with Coomassie Brilliant Blue (Bio-Rad). For blue-native (BN)-PAGE, solubilised thylakoid membrane protein complexes, prepared as outlined above, were 182 separated on 8-16% BN-gels (25) to resolve ChIG-HliD and Ycf39-HliD complexes. Protein complexes 183 were further resolved by incubating the BN-gel strip in 2% (w/v) SDS and 1% (w/v) dithiothreitol for 184 185 30 min at room temperature followed by separation of individual protein components in the second 186 dimension by SDS-PAGE in a denaturing 12 to 20% (w/v) polyacrylamide gel containing 7 M urea. For immunoblotting, proteins were transferred onto polyvinylidene fluoride membranes (Thermo Fisher 187 188 Scientific) and incubated with specific primary antibodies against the 3xFLAG tag (Sigma-Aldrich), His₆-189 tag (Merck), HliD (Agrisera AS10-1615), ChlG (described in reference 21) or Ycf39 (21) followed by an 190 appropriate secondary antibody (anti-rat for 3xFLAG, anti-mouse for His₅ and anti-rabbit for HliD, ChIG and Ycf39) conjugated with horseradish peroxidase (Sigma-Aldrich) to allow detection using the
WESTAR ETA C 2.0 chemiluminescent substrate (Cyanagen) with an Amersham Imager 600 (GE
Healthcare).

194

195 Quantification of Chl and Chl precursors

196 Chl content was determined spectrophotometrically following extraction from cell pellets (from 1 mL 197 of culture at $OD_{750} \approx 0.4$) with 100% methanol according to Porra et al. (39). Chl precursors were 198 extracted from cell pellets (from 2 mL of culture at $OD_{750nm} \approx 0.4$) of WT and mutant *Synechocystis* 199 strains (five biological replicates per strain) and analysed by RP-HPLC with two fluorescence detectors, 200 as described previously (40). Equivalent peaks were integrated, summed, and calculated as a 201 percentage of the WT values, which were set as 100%.

202

203 Results

204 Generation of strains with altered xanthophyll content

205 Photosynthetic carotenoids are C_{40} molecules synthesized from 8 C_5 isoprene units (41). Synechocystis accumulates four major carotenoid species, β -carotene, zeaxanthin, myxoxanthophyll and 206 207 echinenone, and produces others in lesser amounts, including synechoxanthin, 3'-hydroxy-208 echinenone and β -cryptoxanthin (35, 42); the molecular structures of the major carotenoids are 209 presented in Figure 1B. An overview of carotenoid biosynthesis from all-trans-lycopene, the last 210 common precursor of all the mature carotenoids synthesized by Synechocystis, is given in Figure S3. 211 Briefly, lycopene is cyclized at one or both of its ψ -ends producing γ -carotene (one β -ionone ring) or β-carotene (two β-rings); the myxoxanthophyll biosynthesis pathway branches from γ-carotene, 212 213 whereas the other carotenoids are produced by modification of β -carotene. The carotenoid contents of WT Synechocystis and a strain containing an N-terminally 3xFLAG-tagged ChIG but lacking the native 214 215 chlG (FG/ Δ G) were analysed by RP-HPLC and the four expected major carotenoid species were identified (Figure S2). 216

217 Mutants unable to synthesize β -carotene display severe growth phenotypes because it is required for 218 assembly of PSII (4, 7), but xanthophylls are dispensable for photoautotrophic growth under our low-219 stress laboratory conditions (35, 42, 43). Mutants lacking xanthophyll biosynthesis genes were 220 generated in the WT and FG/ Δ G strains (**Table 1**; **Figure S4**), resulting in identical carotenoid 221 deficiencies in both backgrounds (**Figure S2**; summarised in **Table 2**).

Deletion of the *crtR* gene, which encodes the β -carotene hydroxylase acting at the 3/3' positions of 222 223 the ionone rings of β -carotene (7), prevented biosynthesis of both zeaxanthin and myxoxanthophyll 224 and resulted in the appearance of a new carotenoid species, previously identified as dehydroxy-225 myxoxanthophyll (myxoxanthophyll missing the hydroxyl group on the β -ring (35, 44)). It is not 226 possible to generate a knockout strain that produces myxoxanthophyll but not zeaxanthin owing to 227 the shared requirement of CrtR for synthesis of both carotenoids, however, myxoxanthophyll 228 biosynthesis is specifically halted at the first dedicated step, 1',2'-hydroxylation of lycopene/y-229 carotene, in the absence of the C-1'-hydroxylase CruF (Figure S3) (45, 46). Deletion of cruF generated 230 a mutant that did not contain any myxoxanthophyll or myxoxanthophyll-specific precursors but 231 otherwise had a normal carotenoid quota; cruF deletion in the Δ crtR strain resulted in a strain that 232 lacks 3-dehydroxy-myxoxanthophyll as well as zeaxanthin and myxoxanthophyll, accumulating only β carotene and echinenone in significant amounts (Figure S2). The only other confirmed enzyme in 233 234 myxoxanthophyll biosynthesis in cyanobacteria is the 2'-O-glycosyltransferase CruG, which adds a 235 sugar moiety to the carotenoid backbone (Figure S3; 45). Deletion of the Synechocystis cruG homologue from the WT and $\Delta crtR$ strains resulted in accumulation of myxol or 3-dehydroxy-myxol, 236 respectively (Figure S2). Finally, we constructed a $\Delta crtO$ mutant lacking the FAD-dependent β -ionone 237 238 ring ketolase, which produces myxoxanthophyll and zeaxanthin but is unable to synthesise the ketocarotenoids echinenone and 3'-hydroxy-echinenone (43). 239

241 The in vitro interaction of ChIG and HliD requires zeaxanthin

Parallel immunoprecipitations of FLAG-ChIG (F.ChIG) from the FG/AG strain and the carotenoid 242 243 mutants were performed to investigate the effect of xanthophyll deficiency on the ChIG-HliD 244 interaction in vitro (Figure 2). Retrieval of F.ChIG from each strain was confirmed by SDS-PAGE (Figure 2A) and immunoblotting with FLAG and ChIG specific primary antibodies (Figure 2B). F.ChIG from the 245 246 FG/ Δ G strain co-eluted with both HliD and Ycf39 as observed previously (21, 27); note that F.ChlG from strains lacking crtR migrated slightly further than expected on SDS-PAGE gels, which is discussed 247 248 below.

249 Only a residual immunoblot signal for HliD was observed in the FG/ Δ G/ Δ crtR and FG/ Δ G/ Δ crtR/ Δ cruF eluates, accompanied by the loss of signal for Ycf39; the level of HliD was comparable in thylakoids 250 isolated from each of the strains (Figure 2C), ruling out the possibility of any pleiotropic effect on the 251 252 production of HliD associated with deletion of crtR. Consistent with the absence of HliD, direct 253 measurement of the absorbance spectra of the eluates from the FG/ Δ G/ Δ crtR and FG/ Δ G/ Δ crtR/ Δ cruF 254 strains revealed a drastic reduction in pigmentation compared to the visibly orange eluate from the

FG/ΔG strain (**Figure 2D**). Qualitative analysis of the pigments in the eluates by RP-HPLC confirmed Chl, β-carotene myxoxanthophyll and zeaxanthin were present in the complex isolated from the FG/ΔG strain (**Figure 2E**). Small amounts of dehydroxy-myxoxanthophyll, Chl and β-carotene were present in the FG/ΔG/Δ*crtR* elution; the Chl and β-carotene likely originate from trimeric PSI which contaminates FLAG-tag pulldowns (21).

260 In contrast to those from strains lacking crtR, FLAG-immunoprecipitation eluates from the 261 $FG/\Delta G/\Delta cruF$ mutant were visibly orange and spectrally similar to those from the FG/ ΔG parent strain 262 (Figure 2D). Immunoblot analysis of the FG/ Δ G/ Δ cruF eluate gave clear signals for both HliD and Ycf39 and zeaxanthin was identified by RP-HPLC analysis of the extracted pigments (Figure 2B, E). Removing 263 264 the sugar group from myxoxanthophyll (producing myxol) or dehydroxy-myxoxanthophyll (producing dehydroxy-myxol) by deletion of cruG from the FG/ ΔG and FG/ ΔG / $\Delta crtR$ backgrounds, respectively, 265 266 did not alter the results compared to those of the respective parent strain (Figure S5). Finally, deletion 267 of crtO did not affect the composition of the F.ChlG complex (Figure S6), confirming that keto-268 carotenoids are not involved in the ChIG-HliD interaction.

269 Quantitative proteomic analysis by mass-spectrometry was also used to compare the levels of HliD and Ycf39 co-isolated with F.ChlG from the different carotenoid mutants (Figure 2F). When normalised 270 271 to the amount of bait protein, the absence of myxoxanthophyll (FG/ Δ G/ Δ cruF) did not significantly 272 change the amount of HliD co-purified with F.ChIG compared to the FG/ Δ G control (P = 0.35). 273 However, the absence of zeaxanthin (in the $FG/\Delta G/\Delta crtR$ strain) or both zeaxanthin and 274 myxoxanthophyll (in FG/ Δ G/ Δ crtR/ Δ cruF) drastically reduced the level of HliD to 2-3% of the control 275 level (Table S2). These effects on the HliD:F.ChlG stoichiometry were mirrored by that of Ycf39:F.ChlG, with no significant change after elimination of myxoxanthophyll (P = 0.67) and average decreases to 276 277 10-20% of the control level in the zeaxanthin-less strains.

Downloaded from http://portlandpress.com/biochemi/article-pdf/doi/10.1042/BCJ20200561/894140/bcj-2020-0561.pdf by UK user on 09 October 2020

278

279 The protein tag does not affect formation of the ChIG complex

As stated above, F.ChIG isolated from strains lacking *crtR* migrated slightly faster on SDS-PAGE gels (Figure 2A), suggesting the protein was somehow smaller. Sequencing of the *psbAll* locus in these FG/ Δ G/ Δ *crtR* strains confirmed the loss of one of the 3×FLAG-epitopes, leaving a 2×FLAG-tag in frame with the *chIG* gene (Figure S7). Using a freshly isolated FG/ Δ G/ Δ *crtR* mutant confirmed to have the 3×FLAG-tag and performing parallel co-immunoprecipitations alongside a strain with a 2×FLAG-tagged enzyme showed that the length of the tag did not affect the protein or pigment profiles of the eluates (Figure S8).

The 3xFLAG-tag is relatively long (24 amino acids) and highly positively charged. To confirm that this 287 288 extra amino-acid sequence does not generate artefacts regarding the interaction with xanthophylls, 289 the experiments were repeated with an N-terminally 10xHis-tagged ChlG (H.ChlG; Figure S4 and S9). 290 Purification of the His-tagged enzyme by immobilised nickel-affinity chromatography resulted in a 291 visibly pigmented eluate with very similar absorbance properties to the FLAG-immunoprecipitation 292 complex (Figure S10C). Analysis of the eluate by SDS-PAGE and immunoblotting showed a prominent 293 band corresponding to H.ChlG (Figure S10A-B); immunoblots also confirmed the presence of HliD 294 (Figure S10B), and zeaxanthin and myxoxanthophyll were identified by RP-HPLC (Figure S10D). Consistent with the results with the FLAG-tagged enzyme (Figure 2), HliD does not co-elute with 295 296 H.ChlG in the absence of zeaxanthin and myxoxanthophyll (Figure S10A-B) and the eluate contained 297 very low levels of pigments (Figure S10C-D).

298

299 Association of xanthophylls with ChlG is dependent on HliD

300 Previous studies have shown that Synechocystis $\Delta hliD$ mutants do not display a growth phenotype or 301 altered accumulation of photosystems under standard low-stress laboratory growth conditions (47-49). However, deletion of hliD did decrease the level of ChIG, resulting in a concomitant six-fold 302 303 increase in the level of its substrate Chlide a (21). We generated an independent hliD deletion in the 304 $FG/\Delta G$ background (Figures S2 and S4) in order to determine the inherent pigment binding properties 305 of isolated F.ChlG in the absence of HliD. In agreement with previous reports, there was a marked 306 reduction in the level of F.ChIG isolated from FG/ Δ G/ Δ hliD using the same amount of starting material 307 (solubilised thyloakoid membranes), although it was possible to isolate Coomassie-stainable 308 quantities of the protein (Figure 3A). As expected, Ycf39 was not detectable by immunoblot (Figure 309 **3B**) and the eluate lacked Chl and carotenoids, evident from both the absorbance spectra (Figure 3C) 310 and RP-HPLC analysis of extracted pigments (Figure 3D), confirming that ChIG alone does not bind 311 carotenoids. Consistent with the result obtained with the FLAG-tagged enzyme, isolation of H.ChlG 312 from a ΔhliD background also resulted in a decreased level of ChIG (Figure S10A-B) and an immobilised nickel-affinity chromatography eluate that lacked pigments (Figure S10C-D). 313

314

315 Restoration of the ChlG-xanthophyll-HliD interaction in isolated membranes

The results presented above indicate that xanthophylls are required to maintain the interaction between ChIG and HliD, and that ChIG does not co-purify with carotenoids in the absence of HliD. To determine whether the interaction could be restored in isolated membranes, solubilised membranes

from the FG/ Δ G/ Δ *hliD* strain, which synthesizes the normal complement of carotenoids but lacks HliD, 319 320 were incubated with those from the $FG/\Delta G/\Delta crtR/\Delta cruF$ strain, which lacks zeaxanthin and 321 myxoxanthophyll but produces HliD (schematically illustrated in Figure 4A). F.ChlG was subsequently 322 isolated from the individual or mixed membrane samples by immunoprecipitation (Figure 4B). Unlike 323 eluates from either individual sample, immunoblotting detected both HliD and Ycf39 in the elution 324 from the mixed membranes (Figure 4C). Although considerably less pigmented than that from the 325 FG/ Δ G strain, the eluate from the mixed sample was visibly coloured and the absorbance spectra 326 revealed a small but clear increase in pigmentation compared to the eluates from the two mutant strains (Figure 4D). Zeaxanthin and myxoxanthophyll were both present in the complex isolated from 327 the combined membranes (Figure 4E), confirming that the ChIG-HliD interaction can reform upon 328 provision of xanthophylls. Re-formation of the ChIG-HliD complex was also achieved in an analogous 329 experiment with membranes from the equivalent H.ChlG strains (Figure S11). 330

331

332 ChIG-HliD complexes in thylakoid membranes are affected by xanthophyll deficiency

Isolation of FLAG- or His-tagged ChIG complexes requires washing steps that might disrupt interactions 333 334 with weakly-binding proteins. To further ascertain the effects of the loss of xanthophylls on the ChIG-HliD complex we used an alternative approach, separating solubilised thylakoid membrane proteins 335 336 by two-dimensional BN/SDS-PAGE followed by immunoblotting (Figure 5). Most of the detectable ChIG in WT membranes appears to co-migrate with HliD in a ~100 KDa complex; this complex likely 337 forms the major fraction of our co-immunoprecipitated F.ChIG and it is likely to be composed of ChIG 338 339 associated with several copies of HliD (21). A larger ChIG-HliD complex is also detected; we refer to 340 the smaller ChIG-HID assembly as complex 1 and the larger one as complex 2, as indicated above the 341 figure. Small amounts of free ChIG and HliD, and the Ycf39-HliD complex (28), which migrates slightly faster than the ChIG-HliD complex 1, are also clearly observed. 342

Downloaded from http://portlandpress.com/biochemi/article-pdf/doi/10.1042/BCJ20200561/894140/bcj-2020-0561.pdf by UK user on 09 October

2020

Repeating the analysis with membranes of the $\Delta crtR/\Delta cruF$ double mutant lacking both zeaxanthin and myxoxanthophyll, the ratio between the ChIG-HliD complex 1 and unattached ChIG differed from the WT sample, with much less ChIG present in the complex with HliD and more in a free form. Along with the clear reduction in ChIG-HliD complex 1, the larger complex 2 is almost completely absent in the $\Delta crtR/\Delta cruF$ mutant. In contrast, the Ycf39-HliD interaction is unaffected by removal of xanthophylls. This 2D-BN/SDS-PAGE analysis supports the proposed requirement for xanthophylls for the stability of the ChIG-HliD complexes in the thylakoid membranes of *Synechocystis*.

350

351 Xanthophyll mediated association of ChIG and HliD promotes ChIG function

352 As discussed above, deletion of *hliD* results in a decreased cellular level of ChIG and accumulation of 353 its substrate, Chlide (21). To determine the effect of the absence of xanthophylls on Chl biosynthesis 354 the levels of Chl and its biosynthetic intermediates (see Figure S12 for an overview of the Chl 355 biosynthesis pathway) were compared in WT, $\Delta crtR$, $\Delta cruF$ and $\Delta crtR/\Delta cruF$ cells (Figure 6). Under standard growth conditions the whole cell absorbance spectra were similar for all four strains except 356 for the region where carotenoids absorb (450-500 nm; Figure 6A). The Chl level of the $\Delta cruF$ (4.8±0.3 357 μ g Chl per ml of 1 OD₇₅₀ unit cells) and $\Delta crtR/\Delta cruF$ (5.2±0.1 μ g ml⁻¹) strains were not significantly 358 359 different to that of the WT (5.1±0.2 μ g ml⁻¹), but the $\Delta crtR$ mutant did display a small but significant (P = 0.04) decrease in Chl (4.6±0.2 µg ml⁻¹) (Figure 6B). 360

Interestingly, the level of the ChIG substrate monovinyl (MV)-Chlide was 6-7 times higher in the $\Delta crtR$ 361 and $\Delta crtR/\Delta cruF$ mutants compared to the WT (set as 100%) (Figure 6C), indicating that ChIG activity 362 363 is affected by the loss of zeaxanthin. Similarly, divinyl (DV)-Chlide, the substrate of the preceding 364 enzyme in the pathway, 8-vinyl reductase (8VR), also accumulated ~4-fold in both strains compared 365 to the WT, suggesting the 8VR reaction is also affected, either directly or by the build-up of MV-Chlide. In contrast, the $\Delta cruF$ strain contained both Chlide species at levels comparable to the WT. The only 366 other statistically significant differences in precursor levels were the decrease in magnesium-367 protoporphyrin IX in the $\Delta crtR$ mutant and the increase in magnesium protoporphyrin monomethyl 368 369 ester in $\Delta cruF$. The reason(s) for these small variations in earlier pathway intermediates are likely due to pleiotropic effects resulting from the loss of xanthophylls in these strains. 370

371

372 Discussion

373 We have shown that zeaxanthin stabilises the ChIG-HliD interaction in Synechocystis membranes. The 374 alternating single-double carbon-carbon bonds in the polyene chain of carotenoids makes them much more rigid than other hydrophobic molecules, such as comparatively flexible lipids, and a scaffolding 375 role for carotenoids in LHC proteins in plants has been proposed previously (reviewed by 13). Although 376 377 HliD and ChIG still associate to a minor extent in the absence of both zeaxanthin and myxoxanthophyll, this complex appears to be considerably less stable than in WT cells since the HliD protein is almost 378 379 completely absent in F.ChlG immunoprecipitation eluates from a $\Delta crtR$ mutant. Therefore, neither β -380 carotene, echinenone nor dehydroxy-myxoxanthophyll (myxoxanthophyll with an unhydroxylated β -381 ring which accumulates in the $\Delta crtR$ mutant (35, 44)) can substitute for zeaxanthin/myxoxanthophyll 382 in stabilising the ChIG-HliD interaction. These results suggest that the hydroxyl groups on the β ring(s)

Biochemical Journal. This is an Accepted Manuscript. You are encouraged to use the Version of Record that, when published, will replace this version. The most up-to-date-version is available at https://doi.org/10.1042/BCJ20200561 389 390 391 392 393 394 395 396 397 398 399 400 401 402 403 404 405

of zeaxanthin and myxoxanthophyll, which allow xanthophylls to be held perpendicular to the membrane through strong hydrogen bonds, enhancing their structural function (13), are essential for their interaction with ChIG and HliD. Another Hlip (HliA/B) pigment-protein complex has also been reported to contain zeaxanthin and myxoxanthophyll (50), while zeaxanthin binds to the conserved Cterminal transmembrane Chl *a/b* binding (CAB) domains of dimeric ferrochelatase (25); thus, xanthophylls may play similar roles in other Hlip-/CAB-domain protein assemblies in *Synechocystis*.

The ChIG-HIID interaction was maintained in a $\Delta cruF$ mutant that produces zeaxanthin but not myxoxanthophyll, indicating that zeaxanthin alone can mediate association of the two proteins. However, reconstitution of the ChIG-HIID interaction in isolated membranes showed both myxoxanthophyll and zeaxanthin were incorporated into the re-formed complex; thus, promiscuity versus specificity of zeaxanthin and myxoxanthophyll binding sites in the ChIG-HIID complex requires further study. Insight into any specific role of myxoxanthophyll requires a strain that produces myxoxanthophyll in the absence of zeaxanthin, but this is not possible by deletion of native genes because of the shared requirement of CrtR for biosynthesis of both carotenoids. We attempted to generate such a strain using CrtR from the filamentous cyanobacteria *Nostoc* sp. PCC 7120 (alr4009), which produces keto-myxoxanthophyll species but does not synthesise zeaxanthin (51-53). However, our preliminary results indicate that neither myxoxanthophyll nor zeaxanthin biosynthesis was restored when alr4009 was expressed in place of the native *crtR* or at the *psbAll* locus in the $\Delta crtR$ mutant (data not shown).

Isolation of native ChIG complexes from other chlorophototrophic organisms has not yet been reported but given the high-level of sequence conservation of cyanobacterial ChIG enzymes (e.g., 84% identity in *Synechocystis* and *Synechococcus* sp. PCC 7002) we predict that ChIG-xanthophyll-HliD complexes will be conserved in cyanobacteria. In support of this, we previously reported that ChIG from *Synechococcus* sp. PCC 7002, produced in *Synechocystis*, interacts with HliD, zeaxanthin and myxoxanthophyll (27). The myxoxanthophyll species produced in *Synechocystis* is myxol-2' dimethylfucoside, whereas *Synechococcus* sp. PCC 7002 produces myxol-2' fucoside, which lacks two methyl groups on the sugar moiety (45, 54), indicating that the fucose group is not important for the interaction of myxoxanthophyll with the complex, as found with the $\Delta cruG$ mutant here.

Although we cannot rule out that altered thylakoid membrane carotenoid content in the xanthophyll mutants may affect the detergent-sensitivity of protein complexes, our data still supports a role for xanthophylls in stabilising the interaction of ChIG and HliD. Destabilising this association by removal of xanthophylls or deletion of HliD results in a significant build-up of Chlide and, in the latter case, the accumulation of ChIG is also reduced. However, consistent with previous results, there are no major phenotypic consequences relating to growth or chlorophyll biosynthesis upon the loss of HliD, at least
under our standard, low-stress laboratory growth conditions. It is possible that the ChlG-xanthophyllsHliD complex is particularly important under specific stress-conditions. For example, the xanthophylls
could quench singlet oxygen produced during photo-oxidative stress; work is underway to elucidate
how xanthophyll-mediated recruitment of HliD affects Chl metabolism in cyanobacteria *in vivo*.

In contrast to cyanobacterial ChIG, the enzymes from Arabidopsis thaliana (Arabidopsis) and the green 421 422 alga Chlamydomonas reinhardtii do not associate with HliD or xanthophylls when heterologously 423 produced in Synechocystis (27). These phototrophs lack Hlips, and decreased sequence identity 424 between plant and cyanobacterial ChIG (e.g., 63% for Arabidopsis and Synechocystis) may explain why 425 the eukaryotic enzymes do not interact with HliD. Plants do, however, contain single helix LHC-like 426 proteins that are homologous to Hlips called One-Helix Proteins (OHPs) (Figure S13) (55). Like HliD and 427 HliC, which are predicted to form heterodimers in cyanobacteria (22, 28), Arabidopsis OHP1 and OHP2 have been shown to dimerise in vivo (56). Furthermore, OHPs bind the plant homolog of Ycf39 428 429 (HCF244) and are suggested to function in pigment delivery to newly synthesized PSII subunits (56-430 58), as proposed for the cyanobacterial Ycf39-Hlip complex (28, 59). It is currently unclear whether 431 Ycf39-Hlip-dependent synthesis of PSII subunits relies on the additional interaction of Hlips with ChlG. If so, then OHPs (and HCF244) may associate with ChIG in plants as well, although evidence of such a 432 433 complex is yet to be reported. Plants also synthesise LIL3 (55), a two transmembrane helix protein 434 with a proposed role in Chl biosynthesis; LIL3 interacts with the Chl biosynthesis enzymes protochlorophyllide oxidoreductase (POR) and geranylgeranyl diphosphate reductase (ChIP), although 435 436 whether it forms a complex with ChlG is not clear and may be species-specific (60-62). Further work is 437 required to clarify the interactions between these LHC-like proteins and Chl biosynthesis enzymes in 438 plants, but it is feasible that zeaxanthin and/or other abundant plant xanthophylls, such as lutein, may 439 play stabilization roles in such complexes.

440

441 Competing Interests: The authors declare that there are no competing interests associated with the442 manuscript.

443

Author contributions: M.S.P., D.P.C., C.N.H., R.S. and A.H. conceived the study and designed the
experiments. M.P.J., C.N.H., R.S. and A.H. supervised the project. M.S.P., M.P., P.J.J., J.P., E.C.M.,
M.J.D., R.S. and A.H. performed the experiments and/or analysed the data. M.S.P., R.S. and A.H. wrote
the manuscript.

Funding: M.S.P. was supported by a University of Sheffield Faculty of Science PhD Studentship. M.P.,
J.P. and R.S. were supported by grant 19-29225X from the Czech Science Foundation. M.J.D.
acknowledges award BB/M012166/1 from the Biotechnology and Biological Sciences Research Council
(BBSRC) UK. M.P.J. acknowledges award RPG-2019-045 from the Leverhulme Trust. C.N.H.
acknowledges funding from the European Research Council (Synergy Award 854126) and the BBSRC
UK (award BB/M000265/1). A.H. acknowledges support from a Royal Society University Research
Fellowship (award number URF\R1\191548).

456

457 Abbreviations

β-DDM ,n-dodecyl-β-maltoside; BN-PAGE, blue native polyacrylamide gel electrophoresis; Chl(s),
chlorophyll(s); Chlide, chlorophyllide; ChlG, chlorophyll synthase; F.ChlG, FLAG-chlorophyll synthase;
H.ChlG, His-chlorophyll synthase; Hlips, high-light-inducible proteins; HliC, high-light-inducible protein
C; HliD, high light inducible protein D; LHCs, light-harvesting complexes; RP-HPLC, reverse phase highperformance liquid chromatography; SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel
electrophoresis; PSI, photosystem I, PSII, photosystem II; WT, wild-type.

464

Biochemical Journal. This is an Accepted Manuscript. You are encouraged to use the Version of Record that, when published, will replace this version. The most up-to-date-version is available at https://doi.org/10.1042/BCJ20200561

465 References

466 (1) Frank, H.A. and Cogdell, R.J. (1996) Carotenoids in Photosynthesis. *Photochem. Photobiol.* 63, 257467 264 https://doi.org/10.1111/j.1751-1097.1996.tb03022.x

468 (2) Hashimoto, H., Uragami, C. and Cogdell, R.J. (2016) Carotenoids and Photosynthesis. In: Stange C.
469 (eds) Carotenoids in Nature. Subcellular Biochemistry, vol 79. Springer, Cham.
470 https://doi.org/10.1007/978-3-319-39126-7_4

471 (3) Jordan, P., Fromme, P., Witt, H.T., Klukas, O., Saenger, W. and Krausz, N. (2001) Three-dimensional
472 structure of cyanobacterial photosystem I at 2.5 Å resolution. *Nature* 411, 909–917
473 https://doi.org/10.1038/35082000

474 (4) Sozer, O., Komenda, J., Ughy, B., Domonkos, I., Laczkó-Dobos, H., Malec, P. et al (2010) Involvement
475 of carotenoids in the synthesis and assembly of protein subunits of photosynthetic reaction centers
476 of *Synechocystis* sp. PCC 6803. *Plant Cell Physiol*. **51**, 823-835 https://doi.org/10.1093/pcp/pcq031

477 (5) Tóth, T.N., Chukhutsina, V., Domonkos, I., Knoppová, J., Komenda, J., Kis, M. et al (2015)
478 Carotenoids are essential for the assembly of cyanobacterial photosynthetic complexes. *Biochim.*479 *Biophys. Acta.* 1847, 1153-1165 https://doi.org/10.1016/j.bbabio.2015.05.020

(6) Malavath, T., Caspy, I., Netzer-El, S.Y., Klaiman, D. and Nelson, N. (2018) Structure and function of
wild-type and subunit-depleted photosystem I in *Synechocystis*. *Biochim. Biophys. Acta. Bioenerg.* **1859**, 645-654 https://doi.org/10.1016/j.bbabio.2018.02.002

(7) Masamoto, K., Misawa, N., Kaneko, T., Kikuro, T. and Toh, H. (1998) Beta-carotene hydroxylase
gene from the cyanobacterium *Synechocystis* sp. PCC6803. *Plant Cell Physiol.* **39**, 560-564
https://doi.org/10.1093/oxfordjournals.pcp.a029405

(8) Umena, Y., Kawakami, K., Shen, J.R. and Kamiya, N. (2011) Crystal structure of oxygen-evolving
photosystem II at a resolution of 1.9Å. *Nature* 473, 55–60 https://doi.org/10.1038/nature09913

(9) Kurisu, G., Zhang, H., Smith, J.L. and Cramer, W.A. (2003) Structure of the cytochrome b₆f complex
of oxygenic photosynthesis: tuning the cavity. *Science* **302**, 1009-1014
https://doi.org/10.1126/science.1090165

491 (10) Malone, L.A., Qian, P., Mayneord, G.E., Hitchcock, A., Farmer, D.A., Thompson, R.F. et al (2019)
492 Cryo-EM structure of the Spinach cytochrome b₆f complex at 3.6 Å resolution. *Nature* 575, 535-539
493 https://doi.org/10.1038/s41586-019-1746-6

494 (11) Schuller, J.M., Birrell, J.A., Tanaka, H., Konuma, T., Wulfhorst, H., Cox, N. et al (2019) Structural
495 adaptations of photosynthetic complex I enable ferredoxin-dependent electron transfer. *Science* 363,
496 257-260 https://doi.org/10.1126/science.aau3613.

497 (12) Havaux, M., Dall'Osto, L. and Bassi, R. (2007) Zeaxanthin has enhanced antioxidant capacity with
498 respect to all other xanthophylls in Arabidopsis leaves and functions independent of binding to PSII
499 antennae. *Plant Physiol.* 145, 1506–1520 https://doi.org/10.1104/pp.107.108480

(13) Ruban, A.V. and Johnson, M.P. (2010) Xanthophylls as modulators of membrane protein function. *Arch. Biochem. Biophys.* 504, 78-85 https://doi.org/10.1016/j.abb.2010.06.034

(14) Ruban, A.V. (2016) Nonphotochemical fluorescence quenching: mechanism and effectiveness in
protecting plants from photodamage. *Plant Physiol.* **170**, 1903–1916
https://doi.org/10.1104/pp.15.01935

(15) Gruszecki, W.I. and Strzałka, K. (2005) Carotenoids as modulators of lipid membrane physical
properties. *Biochim. Biophys. Acta.* **1740**, 108-115 https://doi.org/10.1016/j.bbadis.2004.11.015

507 (16) Domonkos, I., Kis, M., Gombos, Z. and Ughy, B. (2013) Carotenoids, versatile components of
508 oxygenic photosynthesis. *Prog. Lipid Res.* 52, 539-561 https://doi.org/10.1016/j.plipres.2013.07.001

509 (17) Zhu, Y., Graham, J.E., Ludwig, M., Xiong, W., Alvey, R.M., Shen, G. et al (2010) Roles of xanthophyll carotenoids in protection against photoinhibition and oxidative stress in the cyanobacterium 510 511 Synechococcus sp. strain PCC 7002. Arch. Biochem. Biophys. 504, 86-99 https://doi.org/10.1016/j.abb.2010.07.007 512

(18) Kusama, Y., Inoue, S., Jimbo, H., Takaichi, S., Sonoike, K., Hihara, Y. et al (2015) Zeaxanthin and
echinenone protect the repair of photosystem II from inhibition by singlet oxygen in *Synechocystis* sp.
PCC 6803. *Plant Cell Phys.* 56, 906–916 https://doi.org/10.1093/pcp/pcv018

(19) Vajravel, S., Kis, M., Kłodawska, K., Laczko-Dobos, H., Malec, P., Kovács, L. et al (2017) Zeaxanthin
and echinenone modify the structure of photosystem I trimer in *Synechocystis* sp. PCC 6803. *Biochim. Biophys. Acta. Bioenerg.* 1858, 510-518 https://doi.org/10.1016/j.bbabio.2017.05.001

(20) Kerfeld, C.A., Melnicki M.R., Sutter, M. and Dominguez-Martin, M.A. (2017) Structure, function
and evolution of the cyanobacterial orange carotenoid protein and its homologs. *New Phytol.* 215,
937-951 https://doi.org/10.1111/nph.14670

(21) Chidgey, J.W., Linhartová, M., Komenda, J., Jackson, P.J., Dickman, M.J., Canniffe, D.P. et al (2014)
A cyanobacterial chlorophyll synthase-HliD complex associates with the Ycf39 protein and the
YidC/Alb3 insertase. *Plant Cell* 26, 1267–1279 https://doi.org/10.1105/tpc.114.124495

(22) Niedzwiedzki, D.M., Tronina, T., Liu, H., Staleva, H., Komenda, J., Sobotka, R. et al (2016)
Carotenoid-induced non-photochemical quenching in the cyanobacterial chlorophyll synthase-HliC/D
complex. *Biochim. Biophys. Acta.* 1857, 1430-1439 https://doi.org/10.1016/j.bbabio.2016.04.280

(23) Dolganov, N.A., Bhaya, D. and Grossman A.R. (1995) Cyanobacterial protein with similarity to the
chlorophyll *a/b* binding proteins of higher plants: evolution and regulation. *Proc. Natl. Acad. Sci.* U.S.A. **92**, 636-640 https://doi.org/10.1073/pnas.92.2.636

(24) Komenda, J. and Sobotka, R. (2016) Cyanobacterial high-light-inducible proteins - Protectors of
chlorophyll-protein synthesis and assembly. *Biochim. Biophys. Acta.* 1857, 288–295
https://doi.org/10.1016/j.bbabio.2015.08.011

(25) Pazderník, M., Mareš, J., Pilný, J. and Sobotka, R. (2019) The antenna-like domain of the
cyanobacterial ferrochelatase can bind chlorophyll and carotenoids in an energy-dissipative
configuration. *J. Biol. Chem.* 294, 11131-11143 https://doi.org/10.1074/jbc.RA119.008434

(27) Proctor, M.S., Chidgey, J.W., Shukla, M.K., Jackson, P.J., Sobotka, R., Hunter, C.N. et al (2018) Plant
and algal chlorophyll synthases function in *Synechocystis* and interact with the YidC/Alb3 membrane
insertase. *FEBS Lett.* 18, 3062-3073 https://doi.org/10.1002/1873-3468.13222

(28) Knoppová, J., Sobotka, R., Tichý, M., Yu, J, Konik P, Halada P et al (2014) Discovery of a Chlorophyll
Binding Protein Complex Involved in the Early Steps of Photosystem II Assembly in *Synechocystis*. *Plant Cell* 26, 1200-1212 https://doi.org/10.1105/tpc.114.123919

545 (29) Shukla, M.K., Llansola-Portoles, M.J., Tichý, M., Pascal, A.A., Robert, B. and Sobotka, R. (2018)
546 Binding of pigments to the cyanobacterial high-light-inducible protein HliC. *Photosynth. Res.* 137, 29547 39 https://doi.org/10.1007/s11120-017-0475-7

(30) Staleva, H., Komenda, J., Shukla, M.K., Šlouf, V., Kaňa, R., Polívka, T. et al (2015) Mechanism of
photoprotection in the cyanobacterial ancestor of plant antenna proteins. *Nat. Chem. Biol.* 11, 287–
291 https://doi.org/10.1038/nchembio.1755

(31) Llansola-Portoles, M.J., Sobotka, R., Kish, E., Shukla, M.K., Pascal, A.A., Polivka, T. et al (2017)
Twisting a β-Carotene, an Adaptive Trick from Nature for Dissipating Energy during Photoprotection. *J. Biol. Chem.* 292, 1396-1403 https://doi.org/10.1074/jbc.M116.753723

(32) Rippka, R., Derueles, J., Waterbury, J.B., Herdman, M. and Stainer, R.Y. (1979) Generic
Assignments, Strain Histories and Properties of Pure Cultures of Cyanobacteria. *J. Gen. Microbiol.* 111,
1-61 https://doi.org/10.1099/00221287-111-1-1

(33) Tichý, M., Bečková, M., Kopečná, J., Noda, J., Sobotka, R. and Komenda, J. (2016) Strain of *Synechocystis* PCC 6803 with Aberrant Assembly of Photosystem II Contains Tandem Duplication of a
Large Chromosomal Region. *Front. Plant Sci.* 7, 648 https://doi.org/10.3389/fpls.2016.00648

(34) Xu, H., Vavilin, D., Funk, C. and Vermaas, W. (2004) Multiple deletions of small Cab-like proteins
in the cyanobacterium *Synechocystis* sp. PCC 6803: consequences for pigment biosynthesis and
accumulation. *J. Biol. Chem.* 279, 27971-27979 https://doi.org/10.1074/jbc.M403307200

(35) Lagarde, D. and Vermaas, W. (1999) The zeaxanthin biosynthesis enzyme β-carotene hydroxylase
is involved in myxoxanthophyll synthesis in *Synechocystis* sp. PCC 6803. *FEBS Lett.* 454, 247-251
https://doi.org/10.1016/s0014-5793(99)00817-0

569 570 571 572 573 574 575 (39) Porra, R.J., Thompson, W.A. and Kriedemann, P.E. (1989) Determination of accurate extinction 576 coefficients and simultaneous equations for assaying chlorophylls a and b extracted with four different solvents: verification of the concentration of chlorophyll standards by atomic absorption 577 578 spectroscopy. Biochim. Biophys. Acta. 975, 384-394 https://doi.org/10.1016/S0005-2728(89)80347-0 579 (40) Pilný, J., Kopečná, J., Noda, J. and Sobotka, R. (2015) Detection and Quantification of Heme and 580 Chlorophyll Precursors Using a High Performance Liquid Chromatography (HPLC) System Equipped

582 (41) Maresca, J.A., Graham, J.E. and Bryant, D.A. (2008) The biochemical basis for structural diversity 583 the carotenoids of chlorophototrophic bacteria. Photosynth. Res. 97, 121-140 in https://doi.org/10.1007/s11120-008-9312-3 584

with Two Fluorescence Detectors. Bio-Protocol 5, e1390 https://doi.org/10.21769/BioProtoc.1390

585 (42) Graham, J.E. and Bryant, D.A. (2008) The Biosynthetic pathway for synechoxanthin, an aromatic 586 carotenoid synthesized by the euryhaline, unicellular cyanobacterium Synechococcus sp. strain PCC 587 7002. J. Bacteriol. 190, 7966-7974 https://doi.org/10.1128/JB.00985-08

588 (43) Fernández-González, B., Sandmann, G. and Viogue, A. (1997) A new type of asymmetrically acting beta-carotene ketolase is required for the synthesis of echinenone in the cyanobacterium 589 590 Synechocystis sp. PCC 6803. J. Biol. Chem. 272, 9728-9733 https://doi.org/10.1074/jbc.272.15.9728

(44) Schäfer, L., Vioque, A. and Sandmann, G. (2005) Functional in situ evaluation of photosynthesis-591 592 protecting carotenoids in mutants of the cyanobacterium Synechocystis PCC6803. J. Photochem. Photobiol. B. 78, 195-201 https://doi.org/10.1016/j.jphotobiol.2004.11.007 593

(45) Graham, J.E. and Bryant, D.A. (2009) The biosynthetic pathway for myxol-2' fucoside 594 595 (myxoxanthophyll) in the cyanobacterium Synechococcus sp. strain PCC 7002. J. Bacteriol. 191, 3292-596 3300 https://doi.org/10.1128/JB.00050-09

581

(37) MacGregor-Chatwin, C., Jackson, P.J., Sener, M., Chidgey, J.W., Hitchcock, A., Qian, P. et al (2019) Membrane organisation of photosystem I complexes in the most abundant phototroph on Earth. Nat. Plants 5, 879-889 https://doi.org/10.1038/s41477-019-0475-z

(38) Cox, J. and Mann, M. (2008) MaxQuant enables high peptide identification rates, individualized p.p.b.-range mass accuracies and proteome-wide protein quantification. Nat. Biotechnol. 26, 1367-1372 https://doi.org/10.1038/nbt.1511

(47) Funk, C. and Vermaas, W. (1999) A cyanobacterial gene family coding for single-helix proteins
resembling part of the light-harvesting proteins from higher plants. *Biochemistry* 38, 9397-9404
https://doi.org/10.1021/bi990545+

(48) He, Q., Dolganov, N., Bjorkman, O. and Grossman, A.R. (2001) The high light-inducible
polypeptides in Synechocystis PCC6803. Expression and function in high light. *J. Biol. Chem.* 276, 306314 https://doi.org/ 10.1074/jbc.M008686200

(49) Wang, Q., Jantaro, S., Lu, B., Majeed, W., Baily, M. and He, Q. (2008) The High Light-Inducible
Polypeptides Stabilize Trimeric Photosystem I Complex under High Light Conditions in *Synechocystis*PCC 6803. *Plant Physiol.* 147, 1239-1250 https://doi.org/10.1104/pp.108.121087

609 (50) Daddy, S., Zhan J., Jantaro, S., He, C., He, Q. and Wang, Q. (2015) A Novel High Light-Inducible
610 Carotenoid-Binding Protein Complex in the Thylakoid Membranes of *Synechocystis* PCC 6803. *Sci. Rep.*611 5, 9480 https://doi.org/10.1038/srep09480

(51) Takaichi, S., Mochimaru, M., Maoka, T. and Masamoto, K. (2005) Myxol and 4-ketomyxol 2'fucosides, not rhamnosides, from Anabaena sp. PCC 7120 and Nostoc punctiforme PCC 73102, and
proposal for the biosynthetic pathway of carotenoids. *Plant Cell Physiol.* 46, 497-504
https://doi.org/10.1093/pcp/pci049

(52) Makino, T., Harada, H., Ikenaga, H., Matsuda, S., Takaichi, S., Shindo, K. et al (2008)
Characterization of cyanobacterial carotenoid ketolase CrtW and hydroxylase CrtR by
complementation analysis in *Escherichia coli*. *Plant Cell Physiol*. **49**, 1867-1878 https://doi.org/
10.1093/pcp/pcn169

(53) Mochimaru, M., Masukawa, H., Maoka, T., Mohamed, H.E., Vermaas, W.F. and Takaichi, S. (2008)
Substrate specificities and availability of fucosyltransferase and beta-carotene hydroxylase for myxol
2'-fucoside synthesis in *Anabaena* sp. strain PCC 7120 compared with *Synechocystis* sp. strain PCC
6803. *J. Bacteriol.* **190**, 6726-6733 https://doi.org/10.1128/JB.01881-07

(54) Takaichi, S., Maoka, T. and Masamoto, K. (2001) Myxoxanthophyll in *Synechocystis* sp. PCC 6803
is myxol 2'-dimethyl-fucoside, (3R,2S)-myxol 2'-(2,4-di-O-methyl-α-L-fucoside), not rhamnoside. *Plant Cell Physiol.* 42, 756-762 https://doi.org/10.1093/pcp/pce098

627 (55) Engelken, J., Brinkmann, H. and Adamska, I. (2010) Taxonomic distribution and origins of the
628 extended LHC (light-harvesting complex) antenna protein superfamily. *BMC Evol. Biol.* 10, 233
629 https://doi.org/10.1186/1471-2148-10-233

(56) Hey, D. and Grimm, B. (2020) ONE-HELIX PROTEIN1 and 2 Forms Heterodimers to Bind Chlorophyll
in Photosystem II Biogenesis. *Plant Physiol.* 183, 179-193 https://doi.org/10.1104/pp.19.01304

(57) Hey, D. and Grimm, B. (2018) ONE-HELIX PROTEIN 2 (OHP2) is required for the stability of OHP1
and assembly factor HCF244 and is functionally linked to PSII biogenesis. *Plant Physiol.* 177, 1453-1472
https://doi.org/10.1104/pp.18.00540

(58) Li, Y., Liu, B., Zhang, J., Kong, F., Zhang, L., Meng, H. et al (2019) OHP1, OHP2, and HCF244 Form a
Transient Functional Complex with the Photosystem II Reaction Center. *Plant Physiol.* 179, 195–208
https://doi.org/10.1104/pp.18.01231

(59) Knoppová, J. and Komenda, J. (2019) Sequential deletions of photosystem II assembly factors
Ycf48, Ycf39 and Pam68 result in progressive loss of autotrophy in the cyanobacterium *Synechocystis*PCC 6803. *Folia Microbiol. (Praha)* 64, 683-689 https://doi.org/10.1007/s12223-019-00736-w

(60) Tanaka, R., Rothbart, M., Oka, S., Takabayashi, A., Takahashi, K., Shibata, M. et al (2010) LIL3, a
light-harvesting-like protein, plays an essential role in chlorophyll and tocopherol biosynthesis. *Proc. Natl. Acad. Sci. U. S. A.* **107**, 16721-16725 https://doi.org/10.1073/pnas.1004699107

(61) Mork-Jansson, A., Bue, A.K., Gargano, D., Furnes, C., Reisinger, V., Arnold, J. et al (2015) Lil3
Assembles with Proteins Regulating Chlorophyll Synthesis in Barley. *PLos ONE* 10, e0133145
https://doi.org/10.1371/journal.pone.0133145

(62) Hey, D., Rothbart, M., Herbst, J., Wang, P., Müller, J., Wittmann, D. et al (2017) LIL3, a LightHarvesting Complex Protein, Links Terpenoid and Tetrapyrrole Biosynthesis in *Arabidopsis thaliana*. *Plant Physiol.* **174**, 1037-1050 https://doi.org/10.1104/pp.17.00505

Tables (1-2) and Figures (1-6)

Strain	Properties	Reference/source
Wild-type (WT)	Glucose tolerant WT-P substrain of Synechocystis sp. PCC 6803	33
56/46	N-terminally 3xFLAG-tagged Synechocystis sp. PCC 6803 chIG inserted in place of psbAll and deletion of native	27
FG/∆G	<i>chIG</i> gene; kanamycin resistant (Kan ^R) and zeocin resistance (Zeo ^R)	
	N-terminally 10xHis-tagged Synechocystis sp. PCC 6803 chIG inserted in place of psbAll and deletion of native	This study
HG/∆G	<i>chIG</i> gene; Kan ^R and Zeo ^R	
ΔcrtR	Deletion of <i>crtR</i> (sll1468); erythromycin resistant (Em ^R)	This study
∆cruF	Deletion of <i>cruF</i> (sll0814); streptomycin resistant (Sm ^R)	This study
∆crtO	Deletion of <i>crtO</i> (slr0088); chloramphenicol resistant (Cm ^R)	This study
∆cruG	Deletion of <i>cruG</i> (sll1004); Sm ^R	This study
∆crtR/∆cruF	Deletion of <i>cruF</i> in $\Delta crtR$ background; Em ^R and Sm ^R	This study
∆crtR/∆cruG	Deletion of <i>cruG</i> in $\Delta crtR$ background; Em ^R and Sm ^R	This study
ΔhliD	Deletion of <i>hliD</i> (ssr1789); Em ^R	This study

Table 1. *Synechocystis* strains used in this study. Mutations were generated in the WT, FG/ΔG and HG/ΔG backgrounds, as detailed in the text.

Mutant	Xanthophyll deficiency ^a	Intermediates accumulated
∆crtR	zeaxanthin, cryptoxanthin ^b , myxoxanthophyll, 3'-hydroxyechinenone ^b	dehydroxy-myxoxanthophyll
∆cruF	myxoxanthophyll	N/A
∆cruG	myxoxanthothyll	myxol
∆crtO	echinenone, 3'-hydroxyechinenone ^b , canthaxanthin ^b	N/A
ΔcrtR ΔcruF	zeaxanthin, cryptoxanthin ^b , myxoxanthophyll, 3'-hydroxyechinenone ^b	N/A
$\Delta crtR \Delta cruG$	zeaxanthin, cryptoxanthin ^b , myxoxanthophyll, 3'-hydroxyechinenone ^b	dehydroxy-myxol

Table 2. Carotenoid deficiencies of *Synechocystis* mutants lacking combinations of *crtR*, *cruF*, *cruG* and *crtO*.

^aThe xanthophyll deficiency was the same in the WT, FG/ΔG and HG/ΔG backgrounds. ^bThese carotenoids were not detected by RP-HPLC in this study;

deficiency is assumed based on literature, as detailed in the text and Figure S1.

chlorophyllide a chlorophyll a CH₂ CH₃ CH₃ GGPP H₃C H₃C CH₃ CH3 В or PPP ĺВ ChIG D H₃C H₃C E 0: 0 H₃CO H₃CO CH₃ CH₃ CH₃ CH₃ CH В β-carotene zeaxanthin echinenone

Figure 1. The reaction catalysed by chlorophyll synthase (ChlG) and the molecular structures of the four major carotenoids produced by *Synechocystis*. (A) ChlG esterifies C17 on ring D of chlorophyllide a with geranylgeranyl-pyrophosphate (GGPP) or phytyl pyrophosphate (PPP) to produce (GG-)chlorophyll a. (B) The molecular structures of the four major carotenoids produced by WT *Synechocystis*: β -carotene, zeaxanthin, echinenone and myxoxanthophyll. The hydroxyl groups on the β -rings of zeaxanthin and myxoxanthophyll are highlighted in red.

myxoxanthophyll

OCH₃

но

HC

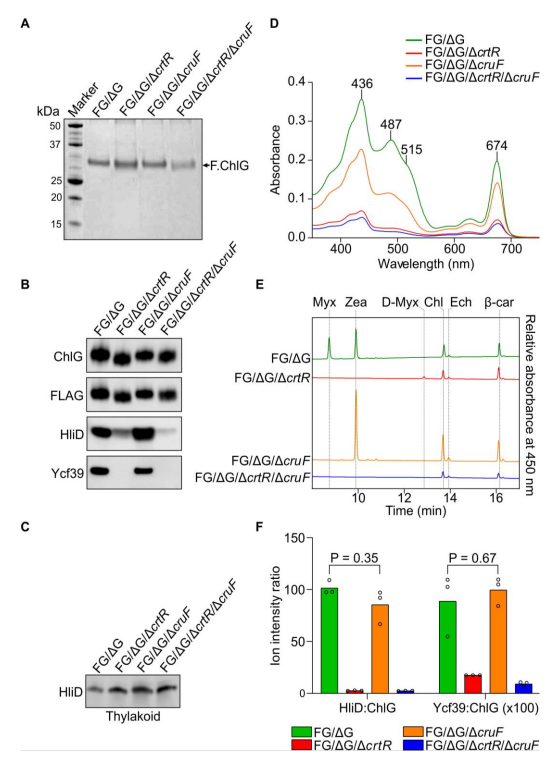


Figure 2. Analysis of pigment-protein complexes isolated by co-immunoprecipitation of F.ChIG from *Synechocystis* xanthophyll mutants. (A) Coomassie stained SDS-PAGE analysis of FLAG immunoprecipitation eluates obtained from the FLAG-chIG Δ chIG (FG/ Δ G), FLAG-chIG Δ chIG Δ crtR (FG/ Δ G/ Δ crtR), FLAG-chIG Δ chIG Δ cruF (FG/ Δ G/ Δ cruF) and FLAG-chIG Δ chIG Δ crtR Δ cruF (FG/ Δ G/ Δ crtR/ Δ cruF) strains of *Synechocystis*. (B) The presence of the FLAG-ChIG (F.ChIG) bait, HIID and Ycf39 was confirmed by immunoblotting with specific primary antibodies. (C) Anti-HIID

Biochemical Journal. This is an Accepted Manuscript. You are encouraged to use the Version of Record that, when published, will replace this version. The most up-to-date-version is available at https://doi.org/10.1042/BC.20200561

immunoblot of thylakoid membranes (30 µg of chlorophyll was loaded in each lane) from WT and mutant strains. (D) Absorbance spectra of the FG/ Δ G (green), FG/ Δ G/ Δ *crtR* (red), FG/ Δ G/ Δ *cruF* (orange) and FG/ Δ G/ Δ *crtR*/ Δ *cruF* (blue) eluates. (E) Separation of pigments extracted from immunoprecipitation eluates by RP-HPLC analysis monitoring absorbance at 450 nm. Myxoxanthophyll (Myx), zeaxanthin (Zea), 3-dehydroxy-myxoxanthophyll (D-Myx), chlorophyll *a* (Chl), echinenone (Ech) and β -carotene (β -car). In (A-E) data are representative of at least three independent experiments. (F) Quantification of F.ChlG, HliD and Ycf39 in immunoprecipitation eluates by mass spectrometry. The ion intensities shown in Table S2 were used to determine the ratios of HliD and Ycf39 to F.ChlG. The results of three technical repeats are presented with P values derived from a Student's t-test (paired, 2-tails).

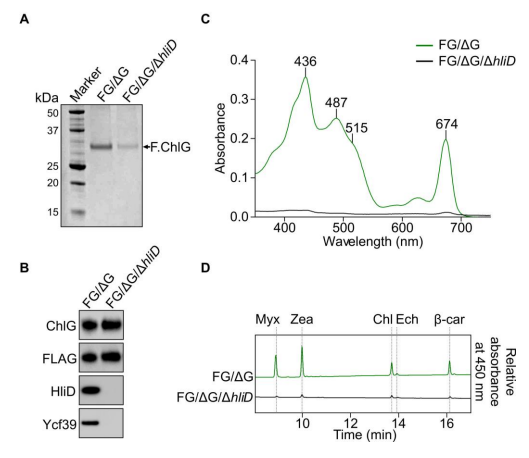


Figure 3. Analysis of pigment-protein complexes isolated by co-immunoprecipitation of F.ChIG from a *Synechocystis* $\Delta hliD$ mutant. (A) Coomassie stained SDS-PAGE analysis of FLAG immunoprecipitation eluates obtained from the FLAG-*chIG* $\Delta chIG$ (FG/ Δ G) and FLAG-*chIG* $\Delta chIG$ $\Delta hliD$ (FG/ Δ G/ $\Delta hliD$) strains. (B) The presence of the FLAG-ChIG (F.ChIG) bait, HliD and Ycf39 was identified by immunoblotting with specific primary antibodies. The FG/ Δ G/ $\Delta hliD$ protein concentration was normalised to that of FG/ Δ G. (C) Absorbance spectra of the FG/ Δ G (green) and FG/ Δ G/ $\Delta hliD$ (black) eluates. (D) Separation of pigments extracted from immunoprecipitation eluates by RP-HPLC analysis monitoring absorbance at 450 nm. Myxoxanthophyll (Myx), zeaxanthin (Zea), 3-dehydroxy-myxoxanthophyll (D-Myx), chlorophyll *a* (ChI), echinenone (Ech) and β -carotene (β -car). Data are representative of at least three independent experiments.

Biochemical Journal. This is an Accepted Manuscript. You are encouraged to use the Version of Record that, when published, will replace this version. The most up-to-date-version is available at https://doi.org/10.1042/BC.20200561

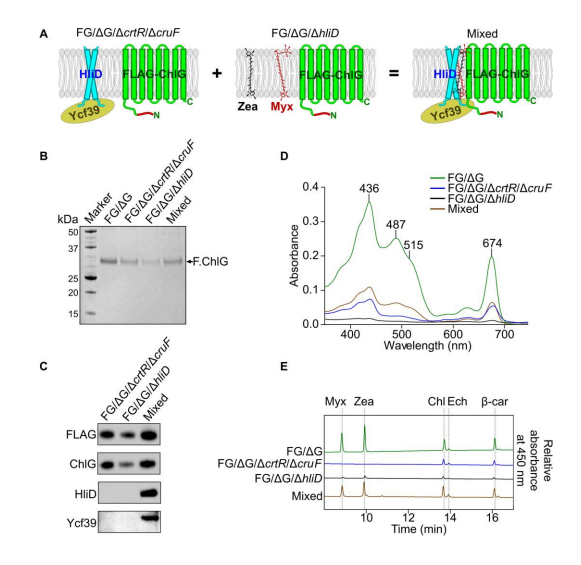


Figure 4. Reconstitution of F.ChIG with carotenoids and HliD in isolated membranes. (A) Solubilised thylakoid membranes from the FLAG-*chIG* Δ*chIG* strain lacking zeaxanthin and myxoxanthophyll (FG/ΔG/Δ*crtR*/Δ*cruF*) but containing FLAG-ChIG (green) and HliD (blue) were mixed with membranes from the FG/ΔG/Δ*hliD* strain, which lacks HliD but contains myxoxanthophyll (Myx, red) and zeaxanthin (Zea, black). (B) Analysis of FLAG immunoprecipitants by SDS-PAGE and Coomassie blue staining; the mixed membranes are labelled as 'Mixed'. (C) Immunodetection of FLAG-ChIG (F.ChIG), HliD and Ycf39 in the immunoprecipitation eluates. HliD and Ycf39 are restored in the mixed membrane sample. (D) Absorbance spectra of each immunoprecipitation eluate. (E) Qualitative RP-HPLC identification of myxoxanthophyll (Myx), zeaxanthin (Zea), chlorophyll *a* (ChI) echinenone (Ech) and β-carotene (β-car). Data are representative of at least three independent experiments.

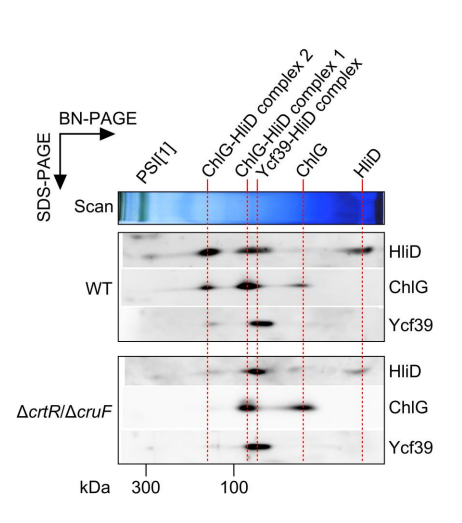
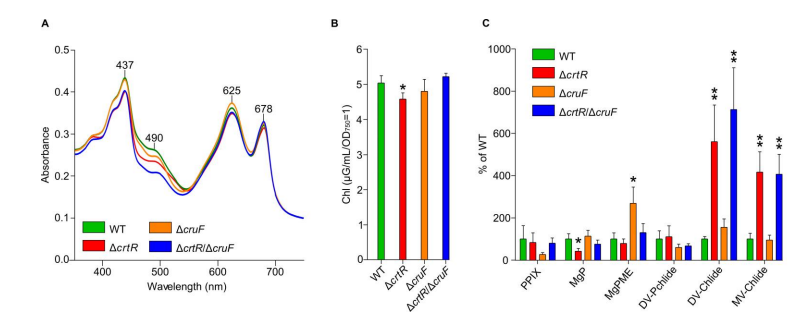


Figure 5. Formation of ChIG-HIiD complexes is altered by the loss of zeaxanthin and myxoxanthophyll. Analysis of thylakoid membrane proteins by 2D-PAGE and immunoblotting. Thylakoid membranes were purified from WT and $\Delta crtR/\Delta cruF$ cells and separated by BN-PAGE in the first dimension and denaturing SDS-PAGE in the second dimension. Proteins were transferred to a PVDF membrane and immunodetection of ChIG, HIiD and Ycf39 was performed using protein specific primary antibodies. Proteins and protein complexes assigned according to previous studies (28, 30) are indicated with red dashed lines and labelled above the gel slice/blots. The approximate molecular weight is indicated below the blots.



Biochemical Journal. This is an Accepted Manuscript

aged to use the

Version of Record that, when published, will replace this

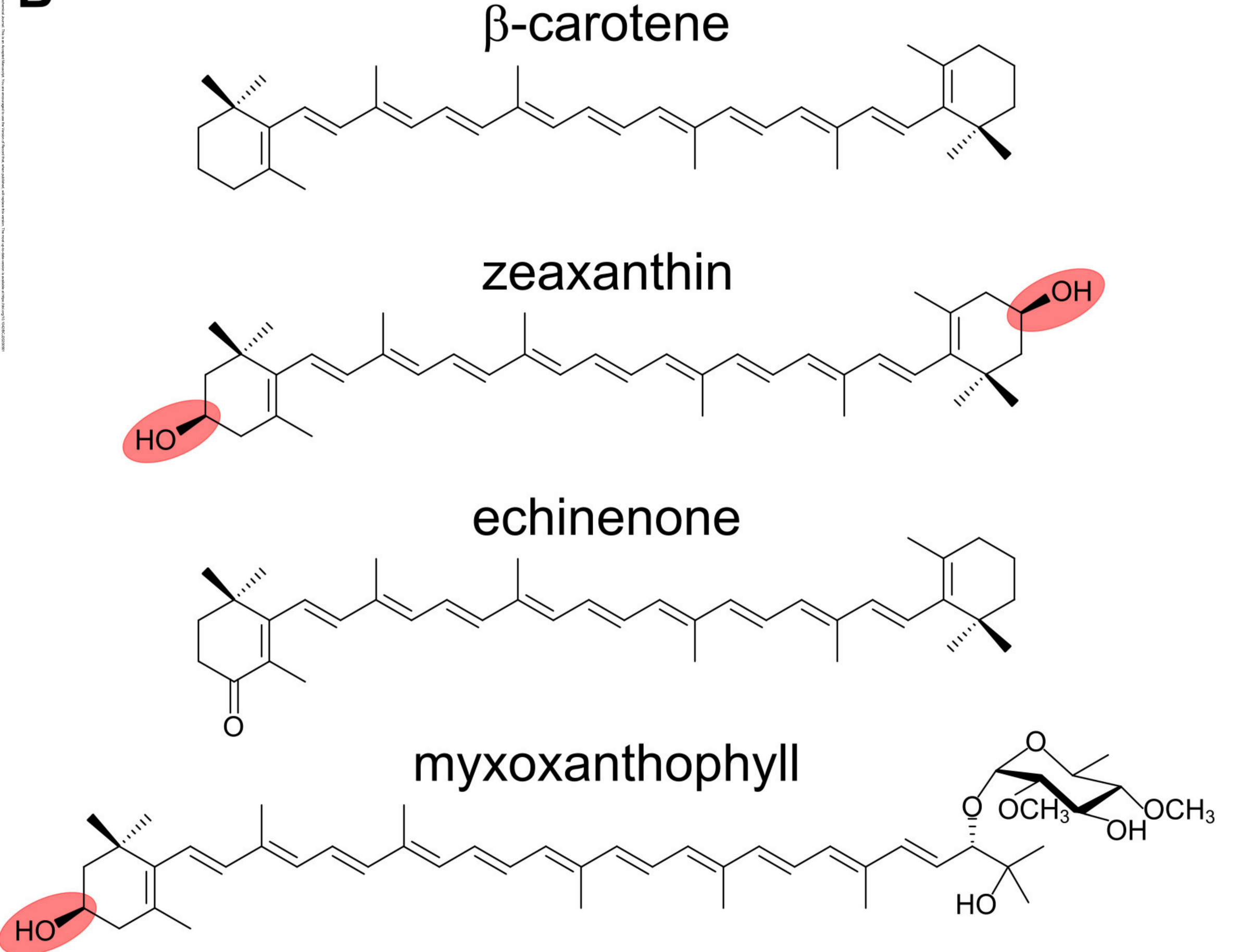
version. The most up-to

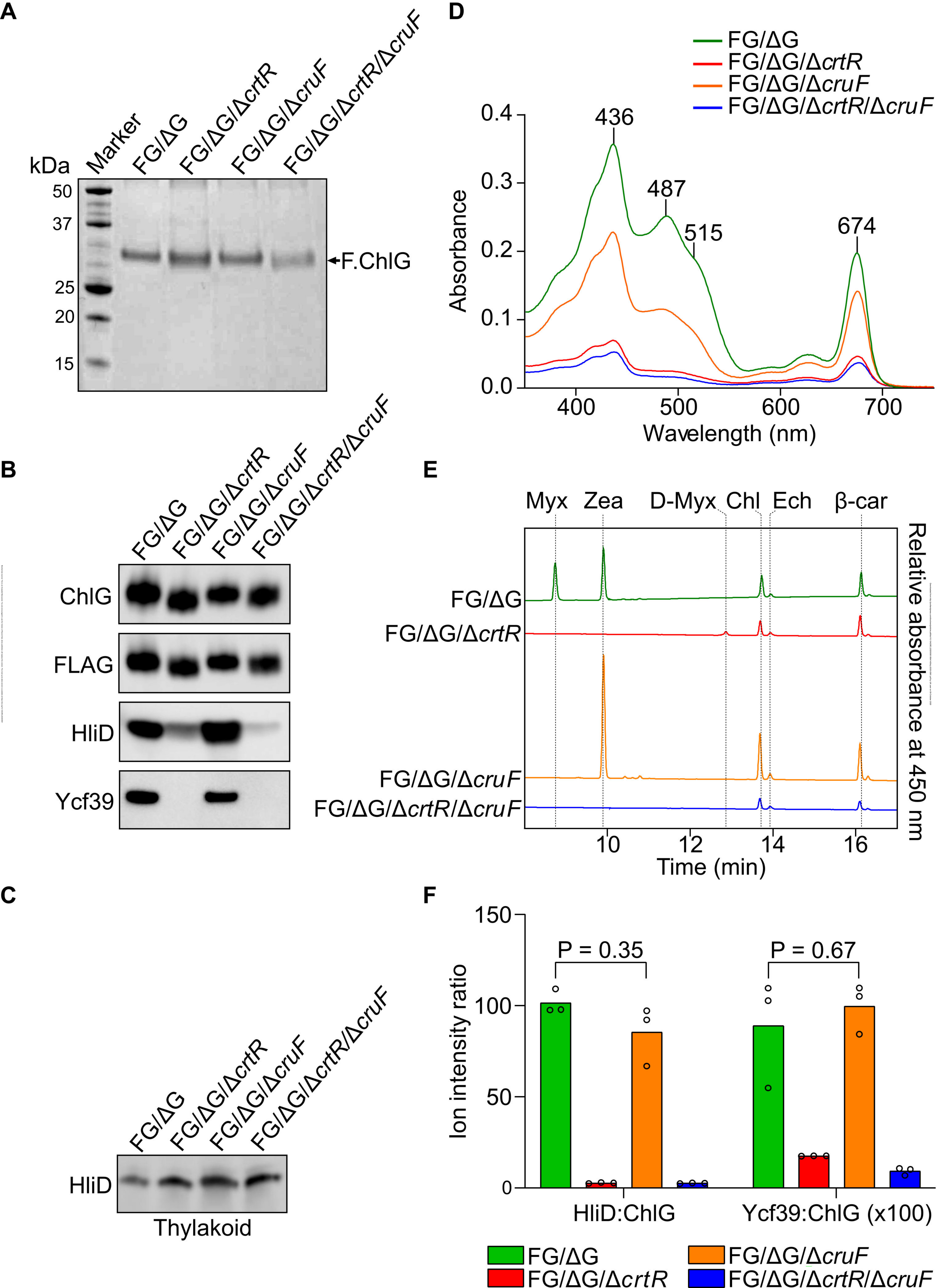
ersion is available at https://doi.org/10.1042/BCJ20200561

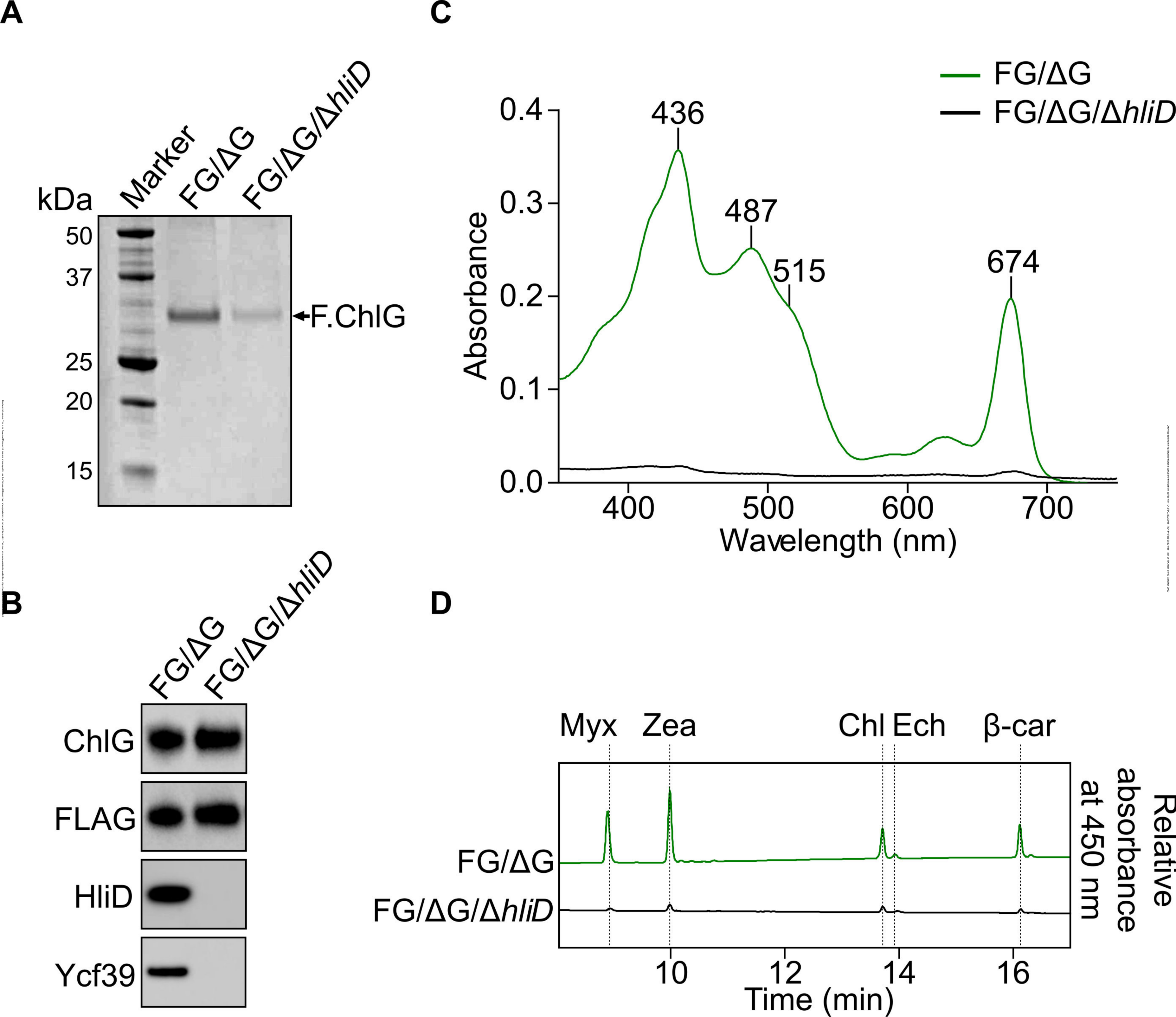
Figure 6. Quantification of Chl and Chl precursors in wild-type *Synechocystis* and xanthophyll deficient mutants. (A) Whole cell absorbance spectra of wild-type (WT, green), $\Delta crtR$ (red), $\Delta cruF$ (orange) and $\Delta crtR/\Delta cruF$ (blue). (B) Chl content of each strain. Error bars represent the standard deviation from the mean of 5 biological replicates. (C) Levels of Chl precursors extracted from $\Delta crtR$, $\Delta cruF$ and $\Delta crtR/\Delta cruF$ cells, relative to those in the WT, which was set at 100% for each pigment. PPIX = protoporphyrin IX; MgP = magnesium-protoporphyrin IX; MgPME = magnesium protoporphyrin monomethyl ester; DV-Pchlide = divinyl-protochlorophyllide; DV-Chlide = divinyl-chlorophyllide; MV-Chlide = monovinyl-chlorophyllide. Error bars represent the standard deviation from the mean of 5 biological replicates. In (B) and (C) statistical significance between the means for Chl/each precursor was determined using one-way analysis of variance (ANOVA); * = p < 0.05; ** = p < 0.001.

chlorophyllide a chlorophyll a **∠**CH₂ _CH₂ CH₃ CH_3 GGPP H₃C∽ H₃C-CH₃ CH₃ or PPP B В Mg Mg ChIG Ν Ν C \ D С D H₃C⁻ H₃C⁴ CH₃ CH₃ Е Е 0 **0**= 0= H₃CO-0 H₃CO CH₃ CH₃ CH₃ CH₃ ΟН 0 CH₃

В

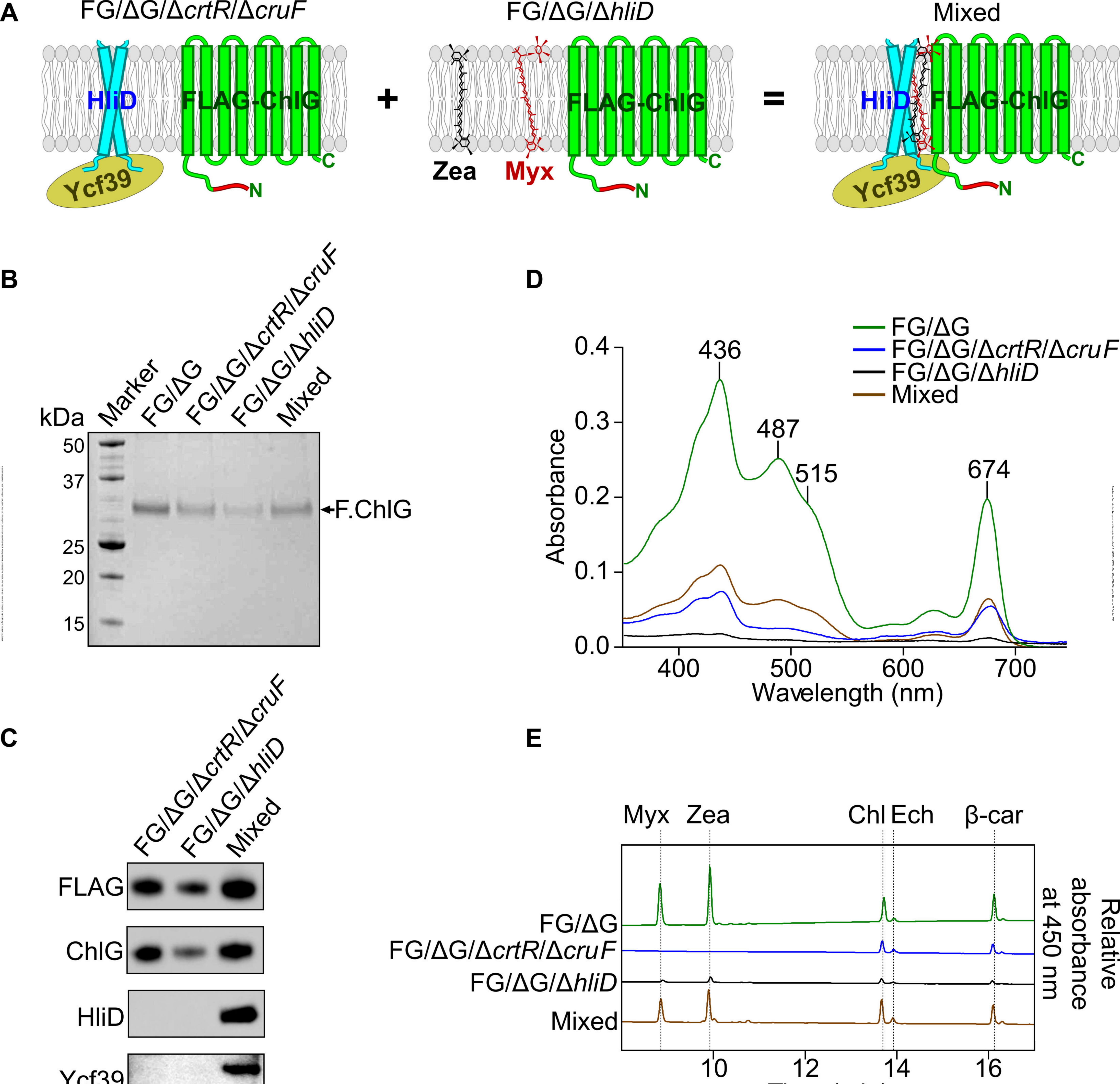


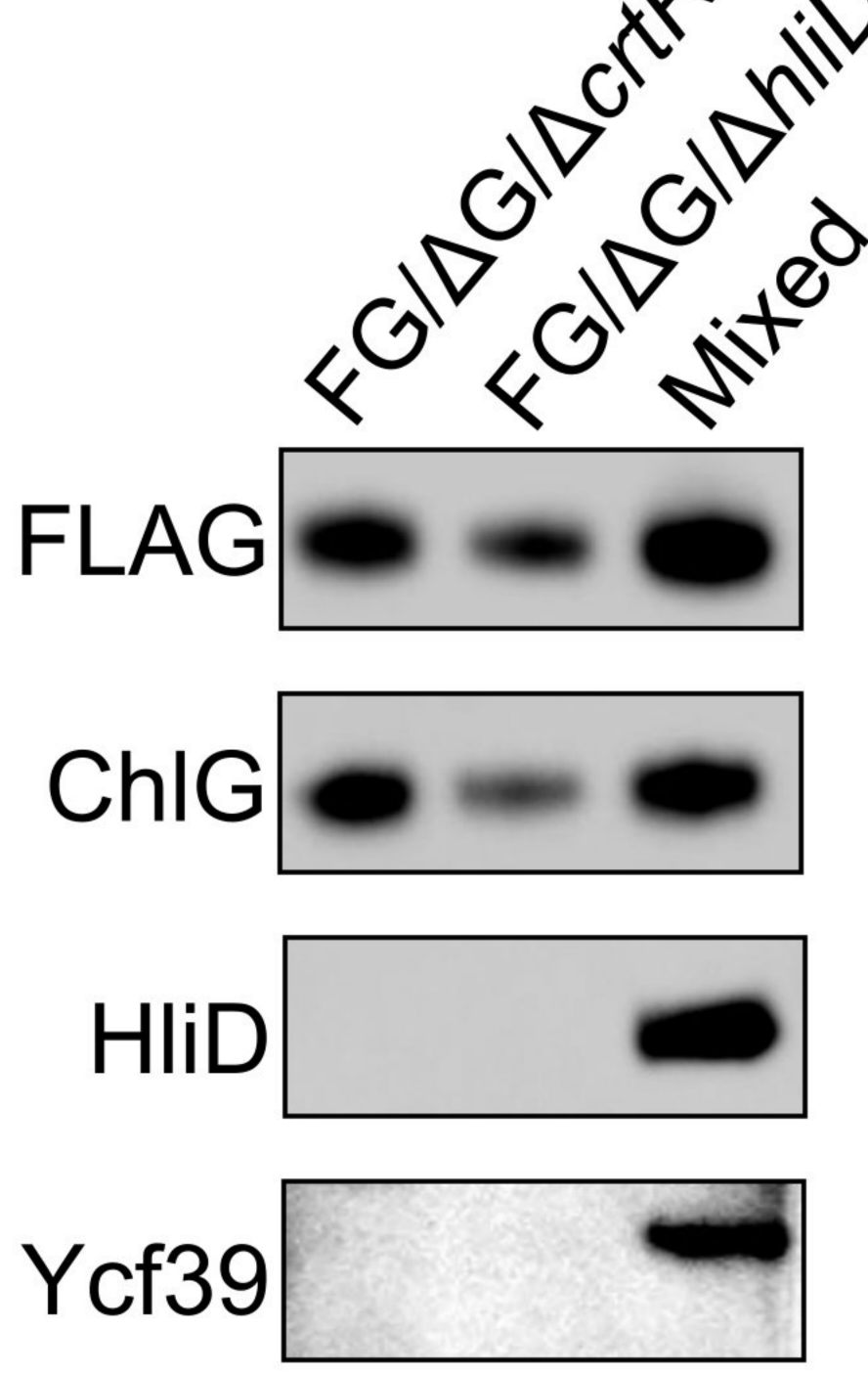




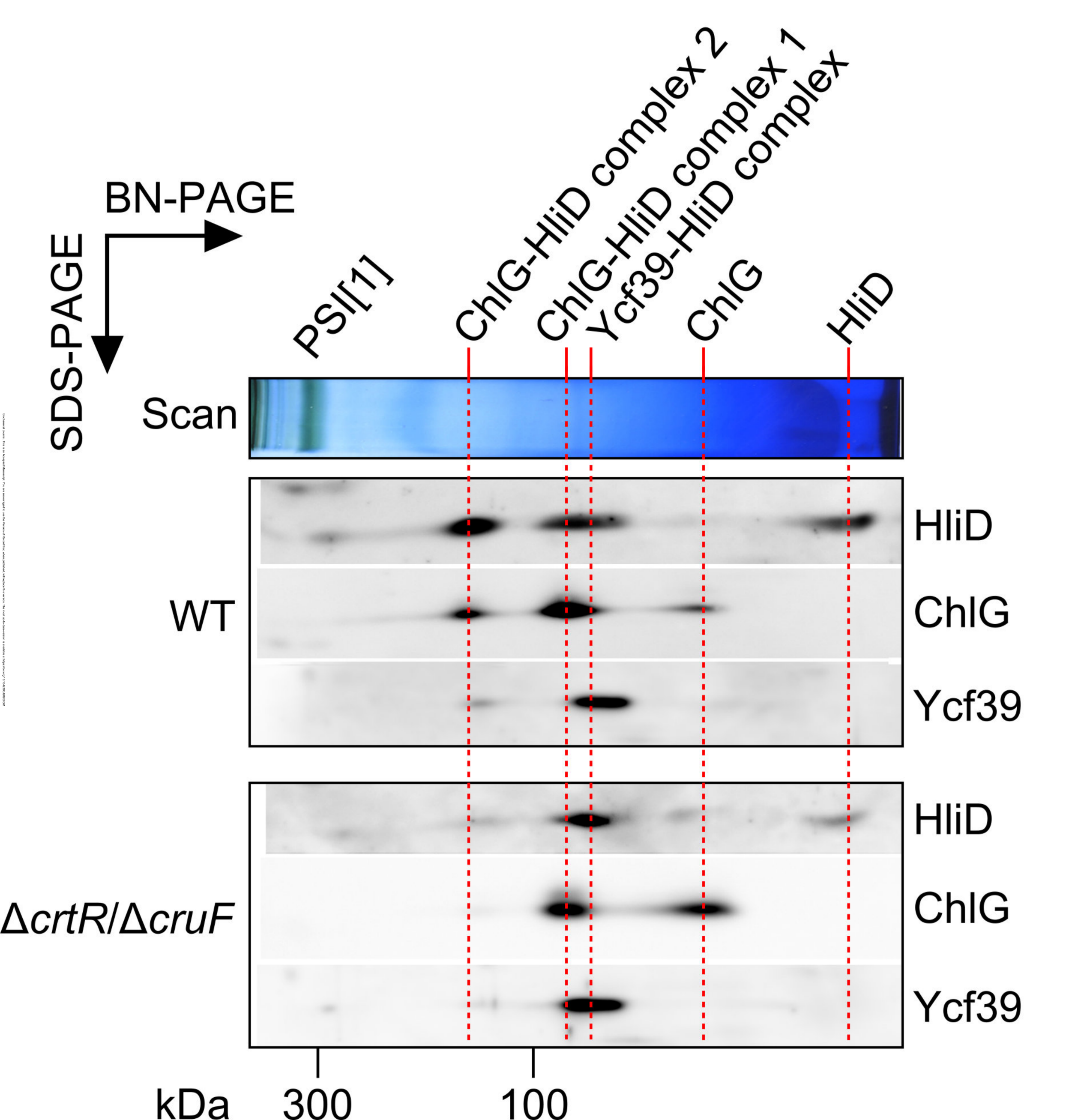








Time (min)



Downloaded from http://portlandpress.com/biochemj/article-pdf/doi/10.1042/BCJ20200561/894140/bcj-2020-0561.pdf by UK user on 09 October 202

