- 1 Interplay between LHCSR proteins and state transitions governs the NPQ response in
- 2 intact cells of *Chlamydomonas* during light fluctuations.
- 4 Collin J. Steen *,1,2,3, Adrien Burlacot *,4,5,6, Audrey H. Short ^{2,3,7}, Krishna K. Niyogi ^{2,4,5} +,
- 5 Graham R. Fleming 1,2,3,7 +
- ¹ Department of Chemistry, University of California, Berkeley, CA 94720, USA
- 8 ² Molecular Biophysics and Integrated Bioimaging Division Lawrence Berkeley National
- 9 Laboratory, Berkeley, CA 94720, USA
- ³ Kavli Energy Nanoscience Institute, Berkeley, CA 94720, USA
- ⁴ Howard Hughes Medical Institute, University of California, Berkeley, CA 94720, USA
- ⁵ Department of Plant and Microbial Biology, University of California, Berkeley, CA, 94720
- 13 USA

21 22

26

29

3

- ⁶ Department of Plant Biology, Carnegie Institution for Science, Stanford, CA, 94305, USA
- ⁷ Graduate Group in Biophysics University of California, Berkeley, CA 94720, USA
- *: Authors have had an equally valued contribution to this work
- 17 +: correspondence to niyogi@berkeley.edu, fleming@berkeley.edu.
- 19 **ORCID IDs:** 0000-0002-7029-2892 (C.J.S.), 0000-0001-7434-6416 (A.B.), 0000-0003-4542-
- 20 1303 (A.H.S.), 0000-0001-7229-2071 (K.K.N.), 0000-0003-0847-1838 (G.R.F.)
- 23 Author contributions: C.J.S., A.B. and G.R.F. designed the research; C.J.S., A.B. and
- 24 A.H.S. performed research; C.J.S., A.B., A.H.S. and G.R.F. analyzed data; C.J.S. and A.B.
- 25 wrote the paper with inputs from A.H.S., K.K.N. and G.R.F.
- 27 One sentence summary: The roles of LHCSR and STT7 in NPQ vary with the light
- 28 fluctuation period and duration of light fluctuation.
- 30 **Key words:** photoprotection, non-photochemical quenching, chlorophyll fluorescence,
- 31 bioenergetics, microalgae, light-harvesting complex stress related, state transition.

Abstract

32

33

34

35

36

37

38

39

40

41

42

43 44

45

46

47

48

49

50

51

52 53

54

55

Photosynthetic organisms use sunlight as the primary energy source to fix CO₂. However, in the environment, light energy fluctuates rapidly and often exceeds saturating levels for periods ranging from seconds to hours, which can lead to detrimental effects for cells. Safe dissipation of excess light energy occurs primarily by non-photochemical quenching (NPQ) processes. In the model green microalga *Chlamydomonas reinhardtii*, photoprotective NPQ is mostly mediated by pH-sensing light-harvesting complex stress-related (LHCSR) proteins and the redistribution of light-harvesting antenna proteins between the photosystems (state transition). Although each component underlying NPQ has been documented, their relative contributions to the dynamic functioning of NPQ under fluctuating light conditions remains unknown. Here, by monitoring NPQ throughout multiple high light-dark cycles with fluctuation periods ranging from 1 to 10 minutes, we show that the dynamics of NPQ depend on the frequency of light fluctuations. Mutants impaired in the accumulation of LHCSRs (npq4, lhcsr1, and npq4lhcsr1) showed significantly less quenching during illumination, demonstrating that LHCSR proteins are responsible for the majority of NPQ during repetitive exposure to high light fluctuations. Activation of NPQ was also observed during the dark phases of light fluctuations, and this was exacerbated in mutants lacking LHCSRs. By analyzing 77K chlorophyll fluorescence spectra and chlorophyll fluorescence lifetimes and yields in a mutant impaired in state transition, we show that this phenomenon arises from state transition. Finally, we quantified the contributions of LHCSRs and state transition to the overall NPQ amplitude and dynamics for all light periods tested and compared those with cell growth under various periods of fluctuating light. These results highlight the dynamic functioning of photoprotection under light fluctuations and open a new way to systematically characterize the photosynthetic response to an ever-changing light environment.

Introduction 56 Most life on Earth is sustained by photosynthetic organisms that capture sunlight energy to 57 convert CO₂ and water into chemical energy. Light is captured by light-harvesting antenna 58 59 complexes that contain a network of pigments absorbing photons and funneling the energy 60 towards photosystems II and I that use it to perform photochemical reactions. Under light-61 limiting conditions, efficient light harvesting is crucial for maximizing the rate of CO₂ 62 fixation (Björkman and Demmig, 1987). However, high light (HL) intensities can saturate 63 reaction centers and lead to the build-up of excess excitation energy, which, if unchecked, can 64 lead to the production of reactive oxygen species and damage to both reaction centers 65 (Khorobrykh et al., 2020). In nature, light exposure rapidly fluctuates in intensity with periods 66 of HL ranging from milliseconds to hours (Graham et al., 2017), requiring photosynthesis to acclimate to different frequencies of HL fluctuations. For each period of HL acclimation, 67 68 photosynthetic organisms exhibit photoprotective mechanisms that regulate light harvesting 69 and safely remove excess energy (Erickson et al., 2015; Pinnola and Bassi, 2018; Roach, 2020). 70 71 Upon light absorption, the energy can be dissipated as heat in a process called non-72 photochemical quenching (NPQ). NPQ involves five components, each of which has been 73 distinguished by its time of induction and relaxation during transition between dark and HL 74 (Erickson et al., 2015). The fastest component, called energy-dependent quenching (qE), is 75 triggered by luminal acidification (Briantais et al., 1979) and is induced and relaxed within seconds. State transition (qT) occurs within minutes and involves the phosphorylation of 76 77 light-harvesting complexes (LHCs) (Allen, 1992) resulting in their detachment from 78 Photosystem (PS) II and subsequent aggregation in a quenched state and/or association to PSI 79 (Nagy et al., 2014; Unlü et al., 2014; Nawrocki et al., 2016). Zeaxanthin-dependent 80 quenching (qZ) requires the accumulation of zeaxanthin and probably involves quenching in 81 the minor LHCs of PSII (Dall'Osto et al., 2005; Wehner et al., 2006; Nilkens et al., 2010). On 82 longer time scales, two more sustained forms of NPQ occur: qH that takes places it the 83 antennae of PSII (Malnoë et al., 2018) directly in the LHCII trimers (Bru et al., 2021) and 84 photoinhibition (qI) that occurs when degradation of the PSII core protein D1 exceeds its 85 capacity for repair due to excess excitation energy (Aro et al., 1993). 86 In the green microalga Chlamydomonas reinhardtii, qE is mediated by pigment-binding LHC 87 stress-related (LHCSR) proteins (Peers et al., 2009; Rochaix and Bassi, 2019). LHCSRs contain protonatable residues, which sense the decreasing luminal pH generated under HL 88 89 conditions (Ballottari et al., 2016; Tian et al., 2019); the protonation of LHCSRs triggers

90 NPO within the protein (Liguori et al., 2013; Kondo et al., 2017; Troiano et al., 2021), allowing fast activation of qE. While there are two types of LHCSR proteins (LHCSR1 and 91 92 LHCSR3), both of which bind pigments (Bonente et al., 2011; Perozeni et al., 2020), 93 LHCSR3 (for which two homologs are present in *Chlamydomonas*) is thought to be the main 94 actor in qE (Peers et al., 2009; Truong, 2011). On the other hand, qT is activated by the 95 buildup of reducing equivalents in the thylakoid membrane, which activates a 96 serine/threonine-protein kinase (STT7) (Depege et al., 2003; Lemeille et al., 2009) that 97 phosphorylates LHCII, enabling it to detach from PSII and ultimately reattach to PSI (Iwai et 98 al., 2010a; Minagawa, 2011). While qZ has been described in Chlamydomonas (Niyogi et al., 99 1997), it does not seem to play a significant role in overall NPQ (Girolomoni et al., 2019; 100 Tian et al., 2019), and its potential mechanism of action remains to be determined. Finally, 101 while qH has not been described in Chlamydomonas, qI occurs upon continued excess 102 illumination (Aro et al., 1993; Erickson et al., 2015) at the level of the PSII center, where 103 oxygen-mediated sensitization creates the irreversible formation of a quenching site 104 (Nawrocki *et al.*, 2021). 105 The photophysical and biochemical bases for NPQ have been studied for decades (Erickson et 106 al., 2015), however the *in vivo* operation has mostly been assessed under a single dark-to-HL 107 transition (Nedbal and Lazár, 2021) leaving our understanding of photoprotection under more 108 complex light patterns limited. While LHCSR and STT7 activity are both known to be important for steady-state NPO under prolonged/continuous illumination (Allorent et al., 109 110 2013), their relative contributions to NPQ have not been quantified, and their response to faster-timescale fluctuating light remains unstudied. Recent work has started looking at the 111 112 response of NPQ to some specific light fluctuations in *Chlamydomonas* (Roach, 2020) and in 113 the moss Physcomitrella (Gao et al., 2021). However, the physiological role and the functioning of the NPQ components under the wide diversity of light patterns that are present 114 in the natural environment is unexplored. 115 Here we utilized two distinct methods to monitor chlorophyll fluorescence in intact cells of 116 Chlamydomonas that were exposed to varying frequencies of fluctuating light with HL/dark 117 118 periods ranging from 1 to 10 min (Fig. 1). The roles of qE and qT were investigated using 119 single or double mutants of LHCSRs and STT7. Our analysis of LHCSR mutants (npq4 120 (Peers et al., 2009), lhcsr1 (Truong, 2011), and npq4lhcsr1 (Truong, 2011)) revealed that 121 LHCSR3 is the main contributor to the NPQ response during the HL phase of light 122 fluctuations. Using mutants impaired in state transition (stt7 (Depege et al., 2003) and 123 stt7npq4 (Allorent et al., 2013)), we showed that qT quenching occurs primarily during the dark portion of the fluctuating light sequence and represents a significant part of NPQ during repeated light fluctuations. Our results showed that while qE and qT sustain most of the NPQ throughout light fluctuations, their relative importance varies during different phases of the fluctuating light response, with qT playing a larger role during dark periods and after repeated HL-dark fluctuations. Surprisingly, the light fluctuation period did not seem to have a major impact on the respective contributions of qE and qT although the contribution of qE during the dark phase was period dependent. Nonetheless, the various components of NPQ are not completely independent, and there may be an interplay between LHCSR- and STT7-mediated NPQ that enables the wild-type photoprotective response. We further show that while *stt7* mutants are not impaired in growth under light fluctuations, short time scale light fluctuations highly impair LHCSR mutants. These findings represent an important first step in investigating the photosynthetic response to the diversity of HL periods that occur in nature.

Results

124

125

126

127 128

129

130

131

132

133

134

135136

137138

Varying light fluctuation periods affect the dynamic NPQ response.

139 While the photosynthetic response of *Chlamydomonas* to some light fluctuations has been 140 reported (Cantrell and Peers, 2017; Roach, 2020), an analysis of NPQ for various periods of 141 light fluctuations is lacking. We therefore measured chlorophyll fluorescence during light-142 dark cycles with fluctuation periods ranging from 1 min to 10 min (Fig. 1). Chlorophyll 143 fluorescence yield was measured using pulse-amplitude modulated (PAM) fluorometry and 144 used to calculate NPQ (Klughammer and Schreiber, 2008). In tandem experiments, timecorrelated single photon counting (TCSPC) was used to measure the chlorophyll fluorescence 145 146 lifetime (Amarnath et al., 2012), which was used to calculate NPQτ (Sylak-Glassman et al., 147 2014). For all periods of light fluctuations in the wild-type strain, NPQ quickly turned on upon illumination but turned off more slowly (Fig. 2, Supp Fig. 1). The same trend was 148 149 observed in NPOτ (Fig. 2). The 1 min period light fluctuation led to a nearly square-like 150 response of NPQ and NPQ τ (**Fig. 2**). For the longer fluctuation periods ranging from 2 minutes to 10 minutes, after an initial burst 151 152 of NPQ in HL, the level of NPQ decreased with continued illumination, eventually reaching a 153 steady state for the 10 min fluctuating period (Fig. 2). A similar trend was also seen in NPQT (Fig. 2). Therefore, these kinetics are directly related to chlorophyll fluorescence quenching, 154 155 rather than other non-quenching processes that affect chlorophyll fluorescence. This phenomenon of decreasing NPQ during the light has been previously attributed to the 156

- consumption of the proton gradient by the activity of the CO₂ concentration mechanism
- 158 (CCM) (Burlacot et al., 2021). For fluctuating light periods longer than 4 minutes, upon a
- transition from HL to dark, the NPQ turned off rapidly but was then followed by a gradual
- rise in NPQ during further darkness, a trend that was also observed in NPQτ (Fig. 2).
- 161 However, compared to NPQ, NPQτ showed a larger magnitude of increase during the long
- dark periods (**Fig. 2C,D**). Differences between the NPQ and NPQτ traces are considered in
- the discussion.

169

185

186

- We conclude from these experiments that, when exposed to light fluctuations with periods
- ranging from 1 to 10 min, at least three components of NPQ are present: (i) a rapidly
- responding component, (ii) a slowly inducible component induced throughout the light
- fluctuations, and (iii) a component induced in the dark phases of light fluctuations.

The majority of NPQ during light fluctuations is mediated by LHCSR proteins.

- 170 It has been well established that LHCSR proteins are crucial for NPQ in Chlamydomonas
- during a single dark-to-light transition (Peers et al., 2009; Truong, 2011; Correa-Galvis *et al.*,
- 2016). To examine the relative importance of each LHCSR protein for the functioning of
- NPO during light fluctuations, we measured the chlorophyll fluorescence yield and lifetime
- during the same light-dark cycles on mutants impaired in the accumulation of LHCSR1
- (lhcsr1) (Truong, 2011), LHCSR3-1 and LHCSR3-2 (npq4) (Peers et al., 2009), or all three
- LHCSRs (npq4lhscr1) (Truong, 2011; Ballottari et al., 2016). While the npq4lhcsr1 mutant
- was highly impaired in its NPQ capacity for all light fluctuations (**Fig. 2, Supp Fig. 1**), single
- 178 npq4 and lhcsr1 mutants showed some NPQ in response to light fluctuation (Fig. 3, Supp
- 179 Fig. 2). Noticeably, for fluctuating periods longer than 4 minutes, the increasing NPQ
- observed during dark phases was more pronounced in the *npq4lhcsr1* mutant (**Fig. 2**). We
- conclude from these experiments that although LHCSRs are responsible for most of the NPQ
- during the light phase of all light fluctuations, a substantial portion of the NPQ in WT is
- nonetheless mediated by other biochemical processes, part of which is induced during the
- dark periods of light fluctuations.

The increasing quenching in the dark periods arises from state transition

188

189

190

191

192

193

194

195

196

197

198 199

200

201

202

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

218

exposure to darkness.

Induction of NPQ during darkness has been previously reported in chlorophytes (Casper-Lindley and Björkman, 1996; Allorent et al., 2013), and qT has been proposed to be involved (Allorent et al., 2013). Re-organization of light-harvesting antennae between PSII and PSI was thus followed throughout a light fluctuation by measuring 77K fluorescence emission spectra. The spectra for cells were compared at three time points: after acclimation to far-red light (cells in state 1 (Zhang et al., 2021)), after 10 min HL, and after 10 additional min dark (see arrows/lines in **Fig. 4**). In WT and *npq4lhcsr1*, an increase in the emission at 710 nm specific to PSI-bound LHCII was observed between the 10 min (after HL) and 20 min (after dark) time points, suggesting that some re-association of LHCII from PSII to PSI occurs during the dark portion of our measurements (Fig. 4, Supp Fig. 3). In contrast, mutants lacking the STT7 kinase responsible for qT (stt7 and stt7npq4) showed negligible changes in the 77K fluorescence emission spectra (Supp Fig 4) and only showed a minimal increase in NPQ or NPQτ during the dark periods of light fluctuations (Fig. 5, Supp Fig. 5) and with increasing duration of exposure to light fluctuations (**Supp Fig. 6**). To characterize the kinetics of qT occurring during the light-to-dark transition, we analyzed the response of the chlorophyll fluorescence lifetime for WT and npq4lhcsr1 mutant cells that were exposed to 10 min of HL followed by 30 min of dark. Interestingly, upon light-to-dark transition, both strains showed steadily decreasing lifetimes for the first 10 min of darkness, after which, the fluorescence lifetimes began to reverse, eventually reaching the starting lifetime after 30 minutes of darkness (Supp Fig. 7). Therefore, we conclude that the quenching observed during the dark phases of light fluctuations in *Chlamydomonas* arises from qT, which has an induction timescale of ~10 min and is reversible upon continued

Quantification of NPQ vs growth under fluctuating light.

It has been previously proposed that, while LHCSRs play an important role during short periods of illumination, state transitions are important for longer periods of high light acclimation (Erickson et al., 2015). To test this hypothesis and assess the scenario under fluctuating light conditions, we quantified the amount of NPQ that was mediated by each protein by comparing the remaining NPQ (or NPQ τ) in each mutant relative to the NPQ (or NPQ τ) in the WT reference strain. Surprisingly, the contribution of each protein to overall NPQ did not seem to depend on the period of the light fluctuation (**Supp Fig. 8**). While

LHCSR3 is responsible for the majority of overall NPO (72%, Fig. 6B), STT7 had a substantial contribution mediating 42% of the NPQ, with LHCSR1 having a smaller contribution at 22% of NPQ. LHCSRs were found to have a substantially larger contribution during light phases, where they are responsible for 94% of WT NPQ, while in the dark phases, their contribution declined to 57% (Fig. 6, Supp Fig. 9). On the other hand, STT7 played a significantly larger role in the NPQ during darkness (60%) than it does during illumination (36%). Interestingly, the amount of NPQ mediated by LHCSRs was more important during the beginning of light fluctuations while state transitions contributed more after 20 minutes of light fluctuation (Fig. 6, Supp Fig. 10), revealing increased relative contribution of qT and decreasing contribution of qE with increasing time of exposure to light fluctuations. Although mutants impaired in the accumulation of LHCSRs and STT7 have strongly impaired NPQ capacities, this does not seem to impair the growth of those strains under continuous high light conditions (Depege et al., 2003; Peers et al., 2009; Cantrell and Peers, 2017). Recent data have shown that the growth of npq4 and npq4lhcsr1 mutants is impaired under certain light fluctuation conditions (Cantrell and Peers, 2017; Roach, 2020). We therefore investigated whether the growth impairment of those strains could be dependent on the period of light fluctuation. While all the mutants grew as well as the control strain under continuous illumination (Supp. Fig. 11), npq4, lhcsr1, npq4lhcsr1 and stt7npq4 mutants exhibited impaired growth under fast light fluctuations with a 1-minute period (Fig. 7). In contrast, only npq4lhcsr1 and stt7npq4 mutants showed an impaired growth under slower light fluctuations with a period of 10 minutes, and the growth of all mutants was similar when the period was increased to 30 minutes (Fig. 7). We conclude from this experiment that qE mediated by LHCSR proteins is critical for growth under light fluctuations and that this role is more important for short light fluctuations.

Discussion

219

220

221

222

223224

225

226

227

228

229

230

231

232

233

234

235

236237

238

239

240

241

242

243

244

245

246

247

248

249

250

The involvement of pH-sensing LHCSR proteins and state transitions in the photoprotective response of *Chlamydomonas* has been previously described (Peers et al., 2009; Allorent et al., 2013). While mutants impaired in accumulation of LHCSRs were shown to have limited growth when light is provided in a day/night cycle (Cantrell and Peers, 2017) or fluctuating with a 10-minute period (Roach, 2020), our understanding of the contribution of LHCSRs and

state transitions to photoprotection during fluctuating light remains limited. Here, by measuring the NPQ levels during light fluctuations in a range of mutants impaired in the accumulation of LHCSR3, LHCSR1, and/or STT7 we have unraveled their relative contributions to NPQ. Varying the length of fluctuating periods from 1 to 10 min allowed us to assess the dynamics of rapid qE- and slower qT-type processes. Interestingly, we observe that qT builds up during the dark periods of the light fluctuations and continues to play a role in the NPQ response during subsequent light phases. It occurs even in the absence of LHCSR3 (see *npq4* and *npq4lhcsr1* mutant in **Fig. 2** and **Fig. 4**), has a timescale of 10 min, and is reversible (Supp **Fig. 7**), which is consistent with recent literature (Allorent et al., 2013; Zhang et al., 2021). During a transition between low-light and high-light stress, qE proteins take a few hours to be fully induced (Peers et al., 2009), and it was hypothesized that qT may substitute for qE during this delay (Allorent et al., 2013). Our results show that even when LHCSRs are fully activated (i.e., in high light-acclimated cells), the occurrence of qT remains substantial during light fluctuations (Fig. 2). State transition or qE mutants were previously shown to have high reactive oxygen species (ROS) production (Allorent et al., 2013; Barera et al., 2021). Thus, the substantial amount of qT induced during the dark periods of light fluctuations may enhance photoprotection and limit ROS production by "anticipating" the next exposure to high light. The combination of fast qE (turns on rapidly upon HL exposure due to ΔpH) and residual qT (from previous dark periods) could therefore provide effective photoprotection in an unpredictable fluctuating-light environment.

Since qE has long been ascribed as the fastest component of NPQ, directly responding to the thylakoid lumen proton concentration (Briantais et al., 1979), and qT as a slower component (Allorent et al., 2013), the contribution of qE to NPQ was proposed to be stronger for short periods of HL, with the contribution of qT becoming increasingly important during longer periods of high-light stress (Erickson et al., 2015). Our approach of systematically assessing the response of photosynthesis to various periods of light fluctuations has revealed nuances in this interpretation. Surprisingly, we found that the overall contributions of qE and qT were not dependent on the period of light fluctuations tested (**Fig. 6** and **Supp. Fig. 8**). However, the *npq4* mutant showed significantly reduced NPQ capacity compared to WT in the dark periods (by 74%) under fast light fluctuations (1-1 sequence), but only a 17% impairment under slower fluctuations (10-10) (see **Supp. Fig. 9**), showing that relaxation of qE (around 1 min) is mediated by LHCSR3 and contributes substantially to the response of NPQ to short periods of light fluctuations. Such relaxation kinetics may contribute to a faster response of NPQ to

the next illumination if the period of light fluctuations is shorter than 2 minutes. It is also worth noting that the relative importance of qE in NPQ decreased after 20 minutes of light fluctuations, while the opposite occurred for qT (**Fig. 6**), reflecting a build-up in qT throughout the 40 minutes of light fluctuations. Modeling the response of photosynthesis to complex light fluctuations has been done (Zaks *et al.*, 2012; Zaks *et al.*, 2013; Tanaka *et al.*, 2019; Steen *et al.*; Nedbal and Lazár, 2021) and would allow targeting specific mechanisms for increasing plant yields in the field. Our results show that such efforts should consider both the period of light and dark as well as the total time exposed to fluctuating light. In green microalgae it is tempting to speculate that in nature, where exposure to HL and dark occur repeatedly, qE may play a more important role in the beginning of light fluctuations while qT may provide a photoprotective response on a longer time scale.

Interestingly, when comparing NPQ and NPQt, the magnitude of the quenching decrease during HL was larger for NPQt than NPQ (Fig. 2). The energetic requirement (and thus its proton gradient consumption) of the CCM depends on the inorganic carbon (C_i) availability (Fei et al., 2021). Since the high cell concentration in the TCSPC sample leads to strong C_i consumption, this could deplete the C_i concentration even in the presence of bubbling, leading to a C_i level sensed by cells in the TCSPC sample being lower than what is experienced by cells in the PAM sample. This would lead to higher activity of CCM, and hence a larger decrease in quenching, under TCSPC sample conditions compared to PAM sample conditions. The decrease in both NPQ and NPQt was also stronger for longer HL periods as well as later in the sequence (Fig. 2). The slope of the initial decrease in NPQτ or NPQ during HL was similar for all four sequences (Supp. Fig. 12) and is likely dictated by C_i availability and its influence on the CCM kinetics. The magnitude of the decrease in NPQ and NPQt was larger for longer light periods (Supp. Fig. 12), likely due to simultaneous activation of the CCM that dissipates the proton gradient and the onset of slower forms of NPQ such as state transitions. Conversely, the differences in the magnitude of the increase in NPQ and NPQT during the dark periods could be due to differences in O₂ concentrations sensed by the cells in both conditions, which is known to affect the extent and rate of state transition (Forti and Caldiroli, 2005).

Although LHCSR3 plays the dominant role in photoprotection under constant and fluctuating light conditions, we also observe a role for LHCSR1 in our measurements. While the chlorophyll fluorescence dynamics of *lhcsr1* are similar to those of WT (**Fig. 3**), we observe a

~20% reduction in overall NPQ in this mutant under fluctuating light conditions (**Fig. 6**). This small amount of photoprotection afforded by LHCSR1 *in vivo* is consistent with previous *in vitro* investigations (Dinc *et al.*, 2016; Nawrocki *et al.*, 2020) in which LHCSR1 has been suggested to mediate energy transfer between LHCII and PSI (Kosuge *et al.*, 2018) and to compensate for the absence of LHCSR3 (Girolomoni et al., 2019). Interestingly, our results suggest that different from the case for LHCSR3, the qE that is mediated by LHCSR1 is largely frequency independent (**Supp. Fig. 8**). Both LHCSR1 and LHCSR3 are thought to generate NPQ in response to (i) the proton gradient and (ii) carotenoid composition (Kondo et al., 2017; Troiano et al., 2021), thus the frequency-dependent LHCSR3 and frequency-independent LHCSR1 could differ in their pH or carotenoid dependency.

The relationship between NPQ capacities and growth have remained puzzling in *Chlamydomonas*, because only some light fluctuation regimes have consistently been shown to impair growth (Peers et al., 2009; Truong, 2011; Cantrell and Peers, 2017; Roach, 2020). Here we show that all mutants lacking LHCSRs showed impaired growth under rapid light

fluctuations (1-1 sequence), and that this impairment was lower under slower fluctuations (10-10 sequence) and absent under an even slower 30-30 sequence or constant illumination (Fig. 7 and **Supp. Fig. 11**). There seems to be a good relationship between defect of NPQ and growth deficiency under short time-scale fluctuations when considering npq4lhcsr1 and stt7npq4 mutants (Fig. 7), clearly showing that LHCSR-dependent qE is critical for growth when light fluctuates with short period of time. However, surprisingly, the growth defect of *lhcsr1* seemed larger than npq4 under the 1-1 sequence. This suggests that the growth capacity of LHCSR mutants under light fluctuations may not depend only on the level of NPQ. Other factors may include activation of compensatory mechanisms that enable photoprotection at the expense of growth or the increased production of reactive oxygen species in npq4 mutants (Roach et al., 2020; Barera et al., 2021), which could be greater in the lhscr1 mutant. Note here that in our conditions, due to slightly different genetic background between stt7 and the WT control, we cannot make a conclusion on the mechanism by which stt7 grew better under short timescale fluctuations (Fig. 7). The WT background may be particularly sensitive to fast 1-1 and 10-10 fluctuations; another possibility could be that qT is detrimental for growth under medium to short time scale fluctuations.

Through 77K fluorescence emission spectra analysis, we have shown that the increasing dissipation observed in the dark requires the STT7 kinase responsible for state transition (**Fig.**

4 and Supp. Fig. 4). This effect, already described in *Chlamydomonas* (Allorent et al., 2013) and *Dunaliella salina* (Casper-Lindley and Björkman, 1996), greatly contributes to NPQ during light fluctuations. In plants, the occurrence of state transitions and its involvement in NPQ is thought to be minor (Allen, 1992; Minagawa, 2011) even if mutants of *Arabidopsis thaliana* impaired in state transition (*stm7*) exhibit impaired growth under light fluctuations (Bellafiore *et al.*, 2005). Interestingly, an increase of NPQ during the dark periods of fluctuating light was recently reported in *npq4* leaves of *A. thaliana* (Steen et al., 2020) for which about 53% of the WT NPQτ remained in the mutant after 40 minutes of exposure to light fluctuations despite the absence of the pH-sensing protein PsbS (Supp. Fig. 13). However, it remains unclear as to how much of the dark quenching in plants originates from qT as opposed to effects related to de-epoxidized xanthophyll pigments (Steen et al., 2020) or LHC protein conformation and/or aggregation (Goral *et al.*, 2012). In the future, periodic illumination experiments performed on *A. thaliana* mutants impaired in qE and/or qT could clarify the relative importance of each mechanism and allow a comparison of the *in vivo* functioning of NPQ in higher plants and green algae under light fluctuations.

Tuning the relaxation kinetics of NPQ in higher plants has been shown to improve crop plant productivity (Kromdijk et al., 2016), and recent modelling of the response of plant canopies to natural light fluctuations has shown that there remains ample room for improving photosynthetic efficiency under non-steady state conditions (Wang et al., 2020). Similar opportunities for improvement also exist for increasing biofuel production from microalgae (Benedetti et al., 2018; Perin and Jones, 2019; Vecchi et al., 2020). In both cases, such optimization will require detailed understanding of the dynamic activity of NPO mechanisms. Since the amount of LHCSR protein does not significantly differ between WT and stt7 (Supp. Fig. 14) and the sum of the NPQ contributions of each protein is greater than 100% (Fig. 6), this suggests that there is an interaction between qE and qT under fluctuating light. The possibility of an interaction between qE and qT has been previously suggested on the basis of a kinetic analysis of qT in the presence and absence of LHCSR3 protein (npq4 mutant) (Roach and Na, 2017). Our findings are consistent with a partial overlap of the functions of LHCSR3 and STT7 in both qE and qT. This partial overlap highlights the need for further investigations of the interactions occurring between proteins that underlie the in vivo NPQ response, not only in HL but also in dark, and more generally during light fluctuations. For example, in microalgae, the quenching mediated by LHCSR3 both at PSII and PSI level (Girolomoni et al., 2019) could be tuned by the movement of LHCSR3 from PSI to PSII during state transitions (Allorent et al., 2013). LHCSR3 is known to associate with LHCII trimers in the PSII supercomplex (Semchonok et al., 2017); therefore, a similar LHCSR3-LHCII interaction may also generate quenching in the trimers following the detachment of LHCII from PSII. It should be noted that in the thylakoid membrane, LHCII can exist in a range of different conformations and/or quenching states (Tian et al., 2015; Kawakami et al., 2019). At this point it is not possible to distinguish the relative contributions of different forms of LHCII in individual snapshot measurements. The ensemble fluorescence lifetime likely originates from some combination of at least three LHCII subpopulations: unphosphorylated and bound to PSII (state 1) (Drop et al., 2014) with an intermediate fluorescence decay component, phosphorylated and unbound (free LHCII) (Iwai et al., 2010b) which has been previously assigned to a long fluorescence decay component (Unlü et al., 2014), and phosphorylated and bound to PSI (state 2) (Huang et al., 2021) with a short fluorescence decay component. In intact algal cells, the relative abundance of each form of LHCII likely dynamically evolves throughout exposure to the fluctuating HL-dark sequences. Further development of in vivo spectroscopic tools will be required to disentangle the dynamics of LHCII conformations and correlate them with photoprotection. Overall, a deeper understanding of the protein interactions underlying NPQ dynamics will be highly valuable in finding new ways to improve plant and microalgal productivity.

Conclusions

LHCSR- and STT7-mediated nonphotochemical quenching processes (qE and qT) are known to underly the photoprotective response of the microalgae *Chlamydomonas*. Here, we have applied a new method to disentangle the involvement of qE and qT in real time by exposing intact algal cells to repetitive cycles of high light and darkness alternating at different frequencies. While both qE- and qT-type responses are present during all light fluctuations, LHCSR-dependent qE plays a larger role in the beginning of light fluctuations and during the HL periods. The contribution of STT7-dependent qT became more pronounced upon longer exposure to fluctuating light and especially during the dark periods of light fluctuations. Over the long term, rapid light fluctuations reduced the growth of mutants impaired in LHCSRs, demonstrating the importance of LHCSR proteins during abrupt changes in light intensity. Overall, our work suggests that a cooperativity between LHCSR proteins and STT7 may constitute an important regulatory feature of the photoprotective response in *Chlamydomonas*. These findings provide a valuable foundation for disentangling and modelling how the diverse molecular mechanisms involved in plant and microalgal acclimation to light fluctuations

- 421 interact and enable robust photosynthesis in nature. We envision that further knowledge on
- the response of photosynthetic mechanisms to various periods of light fluctuations will open
- new avenues for building a strong understanding of how photosynthetic organisms respond to
- 424 complex light fluctuations.

Accession numbers

425 426

436 437

438

449 450

- 427 Genes studied in this article can be found on https://phytozome.jgi.doe.gov/ under the loci
- 428 Cre08.g365900.t1.2 (LHCSR1), Cre08.g367500.t1.1 (LHCSR3.1), Cre08.g367400.t1.1
- 429 (LHCSR3.2), Cre02.g120250.t1.1 (STT7).
- 430 List of abbreviations. NPQ: non-photochemical quenching; qE: energy-dependent
- quenching; qT: state transitions; qI: photoinhibition; qZ: zeaxanthin-dependent quenching;
- 432 PAM: pulse-amplitude modulation; TCSPC: time-correlated single photon counting; LHCSR:
- light-harvesting stress related protein; STT7: serine/threonine-protein kinase; LHC: light-
- harvesting complex; PS: photosystem; CCM: CO₂ concentration mechanism; C_i: inorganic
- carbon (CO₂, HCO₃, CO₃²-); HL: high light

Materials and Methods

Strains and culture conditions

- 439 Chlamydomonas mutants and their respective wild-type 4A- were previously described (npq4)
- 440 (Peers et al., 2009), *lhcsr1* (Truong, 2011), *npq4lhcsr1* (Truong, 2011), *stt7* (Depege et al.,
- 441 2003), stt7npq4 (Allorent et al., 2013)). All strains were grown photoautotrophically under
- moderate light (50 µmol photons m⁻² s⁻¹) in minimal HS medium under air level of CO₂
- 443 (20°C). Except for the growth test, cell cultures (5-8 µg Chl mL⁻¹) were incubated overnight
- at high light (400 µmol photons m⁻² s⁻¹), for maximizing expression of LHCSR proteins
- 445 (Tibiletti et al., 2016). Prior to each measurement, cells were illuminated for at least 15 min
- with low intensity far-red light (3in1LED panel with far-red LED; 3LH series, NK system,
- Japan) to ensure a complete state 1 configuration (Bonaventura and Myers, 1969). All
- replicates shown are biological replicates from independent cultures.

Chlorophyll Fluorescence Measurements

- In this work, we employ two techniques to monitor the activation and deactivation of NPQ
- 452 throughout 40 minutes of exposure to repeated periods of high light and dark on the basis of
- changes in Chl fluorescence emission (see Fig. 1 for an illustration of the experimental
- design). Time-resolved Chl fluorescence was measured via time-correlated single photon

counting (TCSPC) while Chl fluorescence yield was measured in parallel experiments using pulse-amplitude modulation (PAM) fluorimetry. Although both methods can monitor NPQ, the fluorescence lifetime is not susceptible to a range of non-quenching processes that can impact the fluorescence yield (such as chromophore bleaching, changes in chlorophyll concentration, chloroplast movement, or enhanced light scattering (Zaks et al., 2013; Sylak-Glassman *et al.*, 2016). Therefore, fluorescence lifetime measurements provide insight into processes that directly quench chlorophyll fluorescence.

A) TCSPC measurement and fitting

The average chlorophyll fluorescence lifetime was measured by time-correlated single photon counting (TCSPC), as previously described (Sylak-Glassman et al., 2016; Steen et al., 2020). A diode laser (Coherent Verdi G10, 532 nm) pumped a Ti:Sapphire oscillator (Coherent Mira 900f, 808 nm, 76 MHz) and the output was subsequently frequency doubled using a β-barium borate crystal to obtain 404 nm light. These pulses were used for excitation of the sample with a power of 1.7 mW (20 pJ/pulse) and Chl fluorescence emission at 680 nm was detected via an MCP-PMT (Hamamatsu R3809U). A custom-built LABVIEW software was used to synchronize three shutters located in the laser path, actinic light path, and the path between the sample and detector. Every 15 sec, a fluorescence lifetime snapshot measurement was acquired by exposing the cells to the saturating laser (404 nm) for 1 second and detecting the emission. Fluorescence lifetime snapshots were measured by TCSPC using a Becker-Hickl SPC-850 data acquisition card and SPCM software. In between the snapshot measurements, high-light illumination of the cells was achieved by exposing the cuvette to white light set to an intensity of 620 µmol photons m⁻² s⁻¹ (Leica KL1500 LCD, peak 648 nm, FWHM 220 nm). The sample concentration was adjusted to ~80 µg Chl mL⁻¹ for TCSPC measurements. To control the gas composition of the culture and prevent cells from settling to the bottom of the cuvette, the sample was bubbled with air (ambient CO₂ concentrations) at a rate of 2-7 mL min⁻¹ throughout the entire 40 min experiment duration, although note that such bubbling increased the noise of the measurements.

For each fluorescence decay measurement, to ensure that PSII reaction centers were closed, we selected the 0.2 s step with the longest lifetime from the overall 1 s snapshot measurement duration (Sylak-Glassman et al., 2016). This longest step was then fit to a bi-exponential decay (Picoquant, Fluofit Pro-4.6) and the average amplitude-weighted fluorescence lifetime (τ_{avg}) was calculated for each snapshot measurement. The NPQ τ parameter is derived from the fluorescence lifetime snapshot measurements and is defined analogously to NPQ (Sylak-Glassman et al., 2014; Sylak-Glassman et al., 2016): NPQ τ (t) =

 $\frac{\tau_{avg}(0) - \tau_{avg}(t)}{\tau_{avg}(t)}$. The value of NPQ τ represents the magnitude of the quenching response based on the change in the average fluorescence lifetimes between time t=0 (after far-red acclimation but before HL exposure) and any other time t during the 40 min snapshot trajectory. Therefore, using NPQ τ removes confounding effects arising from any differences in the average chlorophyll excited state lifetime of the different strains following far-red acclimation. For all TCSPC measurements, each biological replicate represents the average of three technical replicates measured on the same day.

B) PAM measurements

Chlorophyll fluorescence yield was measured using a pulsed-amplitude modulation (PAM) fluorimeter (Dual-PAM 100, Walz GmbH, Effeltrich, Germany) with the red measuring head. Red saturating flashes (8,000 μ mol photons m⁻² s⁻¹, 600 ms, 620 nm) were delivered to measure F_M (maximal fluorescence yield in dark-acclimated samples) and then every 15 s or 30 s to measure $F_{M'}$ under actinic light exposure or dark phase respectively. Actinic light illumination (620 nm) was set to 620 μ mol photons m⁻² s⁻¹. Fluorescence emission was detected using a long-pass filter (>700 nm). NPQ was calculated as ($F_M - F_{M'}$)/ $F_{M'}$. The Chl concentration was ~5-8 μ g Chl mL⁻¹ and as for TCSPC, all PAM measurements and the sample was bubbled with air at a flux of 2-7 mL min⁻¹ for proper control of the gas concentrations of the sample throughout the entire 40 min experiment duration, note that such bubbling increased the noise of the measurements (but to a lesser extent than for TCSPC).

C) Quantifying the contributions of LHCSRs and STT7 to NPQ

To assess the relative contributions of LHCSR1, LHCSR3, and STT7 to overall NPQ, we analyzed the NPQ (PAM) and NPQτ (TCSPC) trajectories for WT and each mutant. The relative contribution of each protein was determined as the percent change in the integrated snapshot trajectory of NPQ or NPQτ for each mutant relative to the control WT strain. As the contribution of each actor was found to be overall independent of HL-dark fluctuation frequencies in the range of 1 min⁻¹ to 0.1 min⁻¹. (**Supp. Fig. 8**), the average contribution of each protein under the four light fluctuating sequences for both PAM and TCSPC was considered. Additionally, to characterize the involvement in activation or deactivation of NPQ, the quenching trajectories were integrated solely under HL or dark periods, respectively (**Supp. Fig. 9**). The contributions of each protein to the early vs. late responses were further

523

524 525

526

527

528529

530 531

532

533534

535 536

537

538

539

540

541

542

543

544

545

546

547 548

549 550

551

552553

554

assessed by integrating from 0-20 min and 20-40 min, respectively (Supp. Fig. 10). These results are summarized in Fig. 6. 77K Chlorophyll fluorescence emission Chlorophyll fluorescence emission spectra of Chlamydomonas cells at 77 K were obtained by freezing whole cells (~5-8 µg Chl mL⁻¹ final concentration) in liquid nitrogen. The emission spectrum was then measured between 600 and 800 nm (435 nm excitation wavelength, RF-5300PC spectrophotometer, Shimadzu). **Growth tests** The different Chlamydomonas strains were cultivated photoautotrophically under moderate light (50 µmol photons m⁻² s⁻¹) in minimal medium under air level of CO₂ (20°C). Cells were harvested during exponential growth and resuspended in fresh minimal medium to 0.1, 0.5, or 2 μg Chl mL⁻¹. Eight-microliter drops were spotted on minimal medium plates at pH=7.2 and exposed to the various light conditions. Homogeneous light was supplied by LED panels. Temperature was maintained at 25°C at the level of plates. **Acknowledgments:** We thank Dr. Setsuko Wakao for assistance in growing cells, Jacob Irby for performing immunodetection, and Dr. Guillaume Allorent and Dr. Giovanni Finazzi for providing the npq4stt7 strain. This work was supported by the U.S. Department of Energy, Office of Science, Basic Energy Sciences, Chemical Sciences, Geosciences, and Biosciences Division under the field work proposal 449B. K.K.N. is an investigator of the Howard Hughes Medical Institute. **Competing interests**: The authors declare that they have no competing interest. Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. List of Figures, Tables, and Supporting Material:

Figure 1. Experimental design for Chl *a* fluorescence measurements throughout exposure of *Chlamydomonas* cells to fluctuating light with various periods of HL-dark exposure. (**A, B**) Representation of the HL-dark cycles used for the 40 minutes of light fluctuation and their corresponding period, frequency, and name used throughout the main text. NPQ was measured using Pulsed Amplitude Modulation (PAM, C) and Time-Correlated Single Photon Counting (TCSPC, **D**). (**C**) Characteristics of the PAM measurement and representative data of fluorescence yield in WT cells. (**D**) Characteristics of the TCSPC apparatus. Shown are two decays representative of two snapshots taken in a quenched and unquenched state.

Figure 2. Quenching trajectories during light fluctuations in npq4lhcsr1 and its control strain. The response of NPQ and NPQ τ (upper and lower panel respectively) were measured in npq4lhcsr1 mutant and its control strain (red and blue curves respectively) during 40 minutes of light fluctuations with periods of 1, 2, 4 and 10 minutes (**A**, **B**, **C** and **D** respectively) as described in **Fig. 1**. Shown are average of three biological replicates. For TCSPC data, each biological replicate was averaged from three technical replicates. The fluorescence lifetime values used to calculate NPQ τ are shown in the Supplemental.

Figure 3. Quenching trajectories during light fluctuations in *lhcsr1* and *npq4*. The response of NPQ and NPQ τ (upper and lower panel respectively) were measured in *lhcsr1* and *npq4* (purple and orange curves respectively) during 40 minutes of light fluctuations with periods of 1, 2, 4 and 10 minutes (**A**, **B**, **C** and **D** respectively) as described in **Fig. 1**. Shown are average of three biological replicates. For TCSPC data, each biological replicate was averaged from three technical replicates. The fluorescence lifetime values used to calculate NPQ τ are shown in the Supplemental.

Figure 4. 77K chlorophyll fluorescence emission spectra during the first high light-dark cycle of light fluctuations. Cells were placed in a TCSPC cuvette as described in **Fig. 1** and both fluorescence lifetime snapshots and 77K chlorophyll fluorescence emission spectra were taken through 10 minutes of high light and 10 minutes darkness. (**A**) Fluorescence lifetime trajectory of *npq4lhcsr1* mutant (red dots) and its control strain (WT, blue dots). On the graph, dashed vertical lines depict the timepoints at which samples were taken for 77K fluorescence spectra measurement. (**B, C**) 77K fluorescence emission spectra of samples taken in **A** on the control strain (WT, **B**) and *npq4lhcsr1* mutant (**C**). Spectra were taken at 0, 10, and 20 min timepoints (blue, orange and grey spectra respectively). Shown are representative spectra. Three independent biological replicate spectra for WT and npq4lhcsr1 are shown in **Supp. Fig. 3**. 77K spectra for the stt7 and stt7npq4 strains are shown in **Supp. Fig. 4**.

Figure 5. Quenching trajectories during light fluctuations in *stt7* and *stt7npq4*. The response of NPQ and NPQ τ (upper and lower panel respectively) were measured in *stt7* and *stt7npq4* (green and magenta curves respectively) during 40 minutes of light fluctuations with periods of 1, 2, 4 and 10 minutes (**A**, **B**, **C** and **D** respectively) as described in **Fig. 1**. Shown are average of three biological replicates. For TCSPC data, each biological replicate was averaged from three technical replicates. The fluorescence lifetime values used to calculate NPQ τ are shown in the Supplemental.

Figure 6. Quantification of the contribution of LHCSRs and STT7 to wild-type NPQ under fluctuating light. (A) Example of quantification of the relative NPQ mediated by LHCSRs. The area under the NPQ curve of *npq4lhcsr1* mutant (red) was subtracted from that of the control strain (blue) and expressed relative to the area of NPQ of the control strain. (B, C)

- Overall contribution of LHCSRs (red), LHCSR3 (orange), LHCSR1 (purple) and STT7 (green) averaged over all 40 minutes (**B**) or specific periods of the light fluctuations (**C**). Each donut portrays the amount of wild-type NPQ that is lost in each mutant impaired in the accumulation of the given protein. Given that the contribution of each protein was largely independent of HL/dark period (**Supp. Fig. 6**), shown are the average of all 4 light fluctuation sequences. Distribution of individual replicates and estimates of error are presented in **Supp.**
- 611 **Fig. 8-10**.

623

631 632

- 613 **Table 1.** Average contribution of each protein to overall wild-type NPQ for each light 614 fluctuation sequence. Shown is the average value (n=6, evaluated from 3 TCSPC and 3 PAM 615 replicates) and standard deviation of all individual replicates. The contributions of LHCSR3 616 (orange) and LHCSR1 (purple) were determined from the single mutants npq4 and lhcsr1. 617 The contribution of LHCSRs overall (red) was evaluated from the npq4lhcsr1 mutant. The 618 contribution of qT was assessed from the stt7 mutant. Each error in the right column 619 represents the standard deviation of each protein's contribution across the 4 light fluctuation 620 sequences. For simplicity, only the average values (shown in the right column) were used to 621 generate **Figure 6** in the main text. Full distributions of the individual TCSPC and PAM data 622 points are shown in **Supp. Fig. 8**. [supports Fig. 6B]
- **Figure 7.** Growth of mutants impaired in qE and/or qT under various periods of dark/light cycles. *lhcsr1*, *npq4*, *npq4lhcsr1*, *stt7* and *stt7npq4* mutants and their control strain (WT) were diluted and spotted at different chlorophyll concentration and grown on plates under dark/light cycles with a period of 30 (30-30, upper panel), 10 (10-10, middle panel) or 1 minute (1-1, lower panel). Each row represents a different chlorophyll concentration. Shown are representative spots of three biological replicates. Growth under constant low light or high light are shown in Supporting Figure 7.

Supporting Materials:

- Quenching trajectories with error (standard deviation) for WT and npq4lhcsr1 (SI Fig 1)
- Ouenching trajectories with error (standard deviation) for *lhcsr1* and *npq4* (SI Fig 2)
- 635 77K emission spectra PAM replicates for WT and *npq4lhcsr1* (SI Fig 3)
- 636 77K emission spectra for *stt7* and *stt7npq4* (SI Fig 4)
- Ouenching trajectories with error (standard deviation) for stt7 and stt7npq4 (SI Fig 5)
- 638 Maximum quenching envelopes for WT and *stt7* (SI Fig 6).
- 639 Kinetics of qT in WT and *npq4lhcsr1* (SI Fig 7).
- Quantification of protein contributions: distributions, averages, errors (SI Fig 8-10).
- 641 Growth of cells under constant LL or HL (SI Fig 11).
- Kinetics of CCM-related decrease in WT NPQ during HL (SI Fig 12)
- 643 Comparison of integration results for WT and pH-sensing mutant in *Chlamydomonas* and
- 644 *Arabidopsis* (SI Fig 13)
- Immunodetection of LHCSR proteins (SI Fig 14)

References

Allen JF (1992) Protein phosphorylation in regulation of photosynthesis. Biochim. Biophys. Acta **1098**: 275-335 https://doi.org/10.1016/S0005-2728(09)91014-3

- Allorent G, Tokutsu R, Roach T, Peers G, Cardol P, Girard-Bascou J, Seigneurin-Berny D, Petroutsos D, Kuntz M, Breyton C, Franck F, Wollman FA, Niyogi KK, Krieger-Liszkay A, Minagawa J, Finazzi G (2013) A dual strategy to cope with high light in *Chlamydomonas reinhardtii*. Plant Cell **25**: 545-557 10.1105/tpc.112.108274
 - Amarnath K, Zaks J, Park SD, Niyogi KK, Fleming GR (2012) Fluorescence lifetime snapshots reveal two rapidly reversible mechanisms of photoprotection in live cells of *Chlamydomonas reinhardtii*. Proc. Natl. Acad. Sci. U.S.A **109**: 8405-8410 10.1073/pnas.1205303109
 - **Aro E-M, Virgin I, Andersson B** (1993) Photoinhibition of photosystem II. inactivation, protein damage and turnover. Biochim. Biophys. Acta **1143**: 113-134 https://doi.org/10.1016/0005-2728(93)90134-2
 - **Ballottari M, Truong TB, De Re E, Erickson E, Stella GR, Fleming GR, Bassi R, Niyogi KK** (2016) Identification of pH-sensing Sites in the Light Harvesting Complex Stressrelated 3 Protein Essential for Triggering Non-photochemical Quenching in *Chlamydomonas reinhardtii*. J. Biol. Chem. **291:** 7334-7346 10.1074/jbc.M115.704601
- **Barera S, Dall'Osto L, Bassi R** (2021) Effect of lhcsr gene dosage on oxidative stress and light use efficiency by *Chlamydomonas reinhardtii* cultures. J. Biotechnol. **328:** 12-22 https://doi.org/10.1016/j.jbiotec.2020.12.023
- **Bellafiore S, Barneche F, Peltier G, Rochaix J-D** (2005) State transitions and light adaptation require chloroplast thylakoid protein kinase STN7. Nature **433**: 892-895 10.1038/nature03286
- **Benedetti M, Vecchi V, Barera S, Dall'Osto L** (2018) Biomass from microalgae: the potential of domestication towards sustainable biofactories. Microb. Cell Factory **17:** 173 10.1186/s12934-018-1019-3
- **Björkman O, Demmig B** (1987) Photon yield of O₂ evolution and chlorophyll fluorescence characteristics at 77 K among vascular plants of diverse origins. Planta **170**: 489-504 10.1007/BF00402983
- **Bonaventura C, Myers J** (1969) Fluorescence and oxygen evolution from *Chlorella pyrenoidosa*. Biochim. Biophys. Acta **189:** 366-383 https://doi.org/10.1016/0005-2728(69)90168-6
- Bonente G, Ballottari M, Truong TB, Morosinotto T, Ahn TK, Fleming GR, Niyogi KK, Bassi R (2011) Analysis of LhcSR3, a protein essential for feedback de-excitation in the green alga *Chlamydomonas reinhardtii*. Plos Biol. **9:** e1000577 10.1371/journal.pbio.1000577
- **Briantais JM, Vernotte C, Picaud M, Krause GH** (1979) A quantitative study of the slow decline of chlorophyll a fluorescence in isolated chloroplasts. Biochim. Biophys. Acta **548:** 128-138 https://doi.org/10.1016/0005-2728(79)90193-2
- Bru P, Steen CJ, Park S, Amstutz CL, Sylak-Glassman EJ, Leuenberger M, Lam L,
 Longoni F, Fleming GR, Niyogi KK, Malnoë A (2021) Photoprotective qH occurs
 in the light-harvesting complex II trimer. BioRxiv: 2021.2007.2009.450705
 10.1101/2021.07.09.450705
- Burlacot A, Dao O, Auroy P, Cuiné S, Li-Beisson Y, Peltier G (2021) Alternative electron pathways of photosynthesis drive the algal CO₂ concentrating mechanism. BioRxiv: 2021.2002.2025.432959 10.1101/2021.02.25.432959

- **Cantrell M, Peers G** (2017) A mutant of *Chlamydomonas* without LHCSR maintains high 698 rates of photosynthesis, but has reduced cell division rates in sinusoidal light 699 conditions. Plos One **12**: e0179395 10.1371/journal.pone.0179395
- Casper-Lindley C, Björkman O (1996) Nigericin insensitive post-illumination reduction in fluorescence yield in Dunaliella tertiolecta (chlorophyte). Photosynth. Res. **50:** 209-222 10.1007/BF00033120

- Correa-Galvis V, Redekop P, Guan K, Griess A, Truong TB, Wakao S, Niyogi KK, Jahns P (2016) Photosystem II subunit PsbS is involved in the induction of LHCSR protein-dependent energy dissipation in *Chlamydomonas reinhardtii*. J. Biol. Chem. **291:** 17478-17487 10.1074/jbc.M116.737312
- **Dall'Osto L, Caffarri S, Bassi R** (2005) A Mechanism of Nonphotochemical Energy Dissipation, Independent from PsbS, Revealed by a Conformational Change in the Antenna Protein CP26. Plant Cell **17:** 1217-1232 10.1105/tpc.104.030601
- **Depege N, Bellafiore S, Rochaix JD** (2003) Role of chloroplast protein kinase Stt7 in LHCII phosphorylation and state transition in *Chlamydomonas*. Science **299:** 1572-1575
- Dinc E, Tian L, Roy LM, Roth R, Goodenough U, Croce R (2016) LHCSR1 induces a fast and reversible pH-dependent fluorescence quenching in LHCII in *Chlamydomonas reinhardtii* cells. Proc. Nat. Acad. Sci. U. S. A. 113: 7673-7678 10.1073/pnas.1605380113
- **Drop B, Webber-Birungi M, Yadav SKN, Filipowicz-Szymanska A, Fusetti F, Boekema EJ, Croce R** (2014) Light-harvesting complex II (LHCII) and its supramolecular organization in *Chlamydomonas reinhardtii*. Biochim. Biophys. Acta **1837**: 63-72 https://doi.org/10.1016/j.bbabio.2013.07.012
- **Erickson E, Wakao S, Niyogi KK** (2015) Light stress and photoprotection in *Chlamydomonas reinhardtii*. Plant J. **82:** 449-465 10.1111/tpj.12825
 - Fei C, Wilson AT, Mangan NM, Wingreen NS, Jonikas MC (2021) Diffusion barriers and adaptive carbon uptake strategies enhance the modeled performance of the algal CO₂ concentrating mechanism. BioRxiv: 2021.2003.2004.433933 10.1101/2021.03.04.433933
 - **Forti G, Caldiroli G** (2005) State transitions in *Chlamydomonas reinhardtii*. The role of the mehler reaction in state 2-to-state 1 transition. Plant Physiol. **137:** 492-499 10.1104/pp.104.048256
 - Gao S, Pinnola A, Zhou L, Zheng Z, Li Z, Bassi R, Wang G (2021) Light-harvesting complex stress-related proteins play crucial roles in the acclimation of *Physcomitrella patens* under fluctuating light conditions. Phot. Res. 10.1007/s11120-021-00874-8
 - Girolomoni L, Cazzaniga S, Pinnola A, Perozeni F, Ballottari M, Bassi R (2019) LHCSR3 is a nonphotochemical quencher of both photosystems in *Chlamydomonas reinhardtii*. Proc. Nat. Acad. Sci. U. S. A. **116**: 4212-4217 10.1073/pnas.1809812116
 - Goral TK, Johnson MP, Duffy CDP, Brain APR, Ruban AV, Mullineaux CW (2012) Light-harvesting antenna composition controls the macrostructure and dynamics of thylakoid membranes in *Arabidopsis*. Plant J. **69:** 289-301 https://doi.org/10.1111/j.1365-313X.2011.04790.x
- Graham PJ, Nguyen B, Burdyny T, Sinton D (2017) A penalty on photosynthetic growth in fluctuating light. Sci. Rep. 7: 12513 10.1038/s41598-017-12923-1
- Huang Z, Shen L, Wang W, Mao Z, Yi X, Kuang T, Shen J-R, Zhang X, Han G (2021)
 Structure of photosystem I-LHCI-LHCII from the green alga *Chlamydomonas* reinhardtii in State 2. Nature Commun. 12: 1100 10.1038/s41467-021-21362-6
- **Iwai M, Takizawa K, Tokutsu R, Okamuro A, Takahashi Y, Minagawa J** (2010a) 745 Isolation of the elusive supercomplex that drives cyclic electron flow in 746 photosynthesis. Nature **464**: 1210-U1134 10.1038/nature08885

- Iwai M, Yokono M, Inada N, Minagawa J (2010b) Live-cell imaging of photosystem II
 antenna dissociation during state transitions. Proc. Natl. Acad. Sci. USA 107: 2337-2342 10.1073/pnas.0908808107
- Kawakami K, Tokutsu R, Kim E, Minagawa J (2019) Four distinct trimeric forms of light-harvesting complex II isolated from the green alga *Chlamydomonas reinhardtii*. Photosynth. Res. **142:** 195-201 10.1007/s11120-019-00669-y
- Khorobrykh S, Havurinne V, Mattila H, Tyystjärvi E (2020) Oxygen and ROS in photosynthesis. Plants 9: 91

- **Klughammer C, Schreiber U** (2008) Complementary PS II quantum yields calculated from simple fluorescence parameters measured by PAM fluorometry and the Saturation Pulse method. PAM application notes **1:** 201-247
- Kondo T, Pinnola A, Chen WJ, Dall'Osto L, Bassi R, Schlau-Cohen GS (2017) Single-molecule spectroscopy of LHCSR1 protein dynamics identifies two distinct states responsible for multi-timescale photosynthetic photoprotection. Nature Chem. 9: 772-778 10.1038/nchem.2818
- Kosuge K, Tokutsu R, Kim E, Akimoto S, Yokono M, Ueno Y, Minagawa J (2018) LHCSR1-dependent fluorescence quenching is mediated by excitation energy transfer from LHCII to photosystem I in *Chlamydomonas reinhardtii*. Proc. Natl. Acad. Sci. USA 115: 3722-3727 10.1073/pnas.1720574115
- Kromdijk J, Głowacka K, Leonelli L, Gabilly ST, Iwai M, Niyogi KK, Long SP (2016) Improving photosynthesis and crop productivity by accelerating recovery from photoprotection. Science **354**: 857-861 10.1126/science.aai8878
- Lemeille S, Willig A, Depège-Fargeix N, Delessert C, Bassi R, Rochaix J-D (2009) Analysis of the Chloroplast Protein Kinase Stt7 during State Transitions. Plos Biol. 7: e1000045 10.1371/journal.pbio.1000045
- **Liguori N, Roy LM, Opacic M, Durand G, Croce R** (2013) Regulation of light harvesting in the green alga *Chlamydomonas reinhardtii*: the C-terminus of LHCSR Is the knob of a dimmer switch. J. Am. Chem. Soc. **135**: 18339-18342 10.1021/ja4107463
- Malnoë A, Schultink A, Shahrasbi S, Rumeau D, Havaux M, Niyogi KK (2018) The Plastid Lipocalin LCNP Is Required for Sustained Photoprotective Energy Dissipation in *Arabidopsis*. Plant Cell **30**: 196-208 10.1105/tpc.17.00536
- **Minagawa J** (2011) State transitions—The molecular remodeling of photosynthetic supercomplexes that controls energy flow in the chloroplast. Biochim. Biophys. Acta **1807:** 897-905 https://doi.org/10.1016/j.bbabio.2010.11.005
- Nagy G, Ünnep R, Zsiros O, Tokutsu R, Takizawa K, Porcar L, Moyet L, Petroutsos D, Garab G, Finazzi G, Minagawa J (2014) Chloroplast remodeling during state transitions in *Chlamydomonas reinhardtii* as revealed by noninvasive techniques *in vivo*. Proc. Natl. Acad. Sci. U.S.A 111: 5042 10.1073/pnas.1322494111
- Nawrocki WJ, Liu X, Croce R (2020) *Chlamydomonas reinhardtii* exhibits de facto constitutive NPQ capacity in physiologically relevant conditions. Plant Physiol. **182:** 472-479 10.1104/pp.19.00658
- Nawrocki WJ, Liu X, Raber B, Hu C, de Vitry C, Bennett DIG, Croce R (2021)
 Molecular origins of induction and loss of photoinhibition-related energy dissipation
 qI. Sci. Adv. 7: 2021.2003.2010.434601 10.1126/sciadv.abj0055
- Nawrocki WJ, Santabarbara S, Mosebach L, Wollman F-A, Rappaport F (2016) State transitions redistribute rather than dissipate energy between the two photosystems in *Chlamydomonas*. Nat. Plant **2:** 16031 10.1038/nplants.2016.31
- Nedbal L, Lazár D (2021) Photosynthesis dynamics and regulation sensed in the frequency domain. Plant Physiol. **187:** 646–661 10.1093/plphys/kiab317

- Nilkens M, Kress E, Lambrev P, Miloslavina Y, Müller M, Holzwarth AR, Jahns P 796 797 (2010) Identification of a slowly inducible zeaxanthin-dependent component of non-798 photochemical quenching of chlorophyll fluorescence generated under steady-state in Arabidopsis. Biochim. Biophys. 799 conditions Acta 1797: 466-475 800 https://doi.org/10.1016/j.bbabio.2010.01.001
- Niyogi KK, Bjorkman O, Grossman AR (1997) Chlamydomonas Xanthophyll Cycle 801 Mutants Identified by Video Imaging of Chlorophyll Fluorescence Quenching. Plant 802 Cell 9: 1369-1380 10.1105/tpc.9.8.1369 803

805

806

807

810

811 812

813

818 819

820

821

822 823

824

825

826

827

828

829 830

831

832

833

834

835 836

837

- Peers G, Truong TB, Ostendorf E, Busch A, Elrad D, Grossman AR, Hippler M, Niyogi **KK** (2009) An ancient light-harvesting protein is critical for the regulation of algal photosynthesis. Nature 462: 518-521
- **Perin G, Jones PR** (2019) Economic feasibility and long-term sustainability criteria on the 808 path to enable a transition from fossil fuels to biofuels. Curr. Opin. Biotechnol. 57: 809 175-182 https://doi.org/10.1016/j.copbio.2019.04.004
 - Perozeni F, Beghini G, Cazzaniga S, Ballottari M (2020) Chlamydomonas reinhardtii LHCSR1 and LHCSR3 proteins involved in photoprotective non-photochemical quenching have different quenching efficiency and different carotenoid affinity. Sci. Rep. 10: 21957 10.1038/s41598-020-78985-w
- 814 Pinnola A, Bassi R (2018) Molecular mechanisms involved in plant photoprotection. Biochem. Soc. Trans. 46: 467-482 10.1042/bst20170307 815
- 816 Roach T (2020) LHCSR3-Type NPQ Prevents Photoinhibition and Slowed Growth under 817 Fluctuating Light in Chlamydomonas reinhardtii. Plants 9: 1604
 - Roach T, Na CS (2017) LHCSR3 affects de-coupling and re-coupling of LHCII to PSII during state transitions in Chlamydomonas reinhardtii. Sci. Rep. 7: 43145 10.1038/srep43145
 - Roach T, Na CS, Stöggl W, Krieger-Liszkav A (2020) The non-photochemical quenching protein LHCSR3 prevents oxygen-dependent photoinhibition in Chlamydomonas reinhardtii. J. Exp. Bot. 71: 2650-2660 10.1093/jxb/eraa022
 - Rochaix J-D, Bassi R (2019) LHC-like proteins involved in stress responses and biogenesis/repair of the photosynthetic apparatus. Biochem. J. 476: 581-593 10.1042/BCJ20180718
 - Semchonok DA, Sathish Yadav KN, Xu P, Drop B, Croce R, Boekema EJ (2017) Interaction between the photoprotective protein LHCSR3 and C2S2 Photosystem II supercomplex in Chlamydomonas reinhardtii. Biochim. Biophys. Acta 1858: 379-385 https://doi.org/10.1016/j.bbabio.2017.02.015
 - Steen CJ, Morris JM, Short AH, Niyogi KK, Fleming GR (2020) Complex Roles of PsbS and Xanthophylls in the Regulation of Nonphotochemical Quenching in Arabidopsis thaliana under Fluctuating Light. J. Phys. Chem. B 124: 10311-10325 10.1021/acs.jpcb.0c06265
 - Sylak-Glassman EJ, Malnoë A, De Re E, Brooks MD, Fischer AL, Niyogi KK, Fleming GR (2014) Distinct roles of the photosystem II protein PsbS and zeaxanthin in the regulation of light harvesting in plants revealed by fluorescence lifetime snapshots. Proc. Natl. Acad. Sci. USA 111: 17498-17503 10.1073/pnas.1418317111
- 839 Sylak-Glassman EJ, Zaks J, Amarnath K, Leuenberger M, Fleming GR (2016) 840 Characterizing non-photochemical quenching in leaves through fluorescence lifetime 841 snapshots. Photosynth. Res. **127**: 69-76 10.1007/s11120-015-0104-2
- Tanaka Y, Adachi S, Yamori W (2019) Natural genetic variation of the photosynthetic 842 843 induction response to fluctuating light environment. Curr Opin. Plant Biol. 49: 52-59 844 https://doi.org/10.1016/j.pbi.2019.04.010

- Tian L, Dinc E, Croce R (2015) LHCII populations in different quenching states are present in the thylakoid membranes in a ratio that depends on the light conditions. J. Phys. Chem. Letters 6: 2339-2344 10.1021/acs.jpclett.5b00944
- Tian L, Nawrocki WJ, Liu X, Polukhina I, van Stokkum IHM, Croce R (2019) pH dependence, kinetics and light-harvesting regulation of nonphotochemical quenching in *Chlamydomonas*. Proc Nat. Acad. Sci. USA **116**: 8320-8325 10.1073/pnas.1817796116

854

855 856

857

858

859 860

861

862

863

864

865

866

867 868

869

- **Tibiletti T, Auroy P, Peltier G, Caffarri S** (2016) *Chlamydomonas reinhardtii* PsbS protein is functional and accumulates rapidly and transiently under high light. Plant Physiol. **171:** 2717-2730 10.1104/pp.16.00572
- Troiano JM, Perozeni F, Moya R, Zuliani L, Baek K, Jin E, Cazzaniga S, Ballottari M, Schlau-Cohen GS (2021) Identification of distinct pH- and zeaxanthin-dependent quenching in LHCSR3 from *Chlamydomonas reinhardtii*. eLife **10**: e60383 10.7554/eLife.60383
- **Truong TB** (2011) Investigating the role(s) of LHCSRs in *Chlamydomonas reinhardtii*. PhD thesis. University of California, Berkeley
- **Ünlü C, Drop B, Croce R, van Amerongen H** (2014) State transitions in *Chlamydomonas reinhardtii* strongly modulate the functional size of photosystem II but not of photosystem I. Proc. Natl. Acad. Sci. U.S.A **111:** 3460-3465 10.1073/pnas.1319164111
- Vecchi V, Barera S, Bassi R, Dall'Osto L (2020) Potential and challenges of improving photosynthesis in algae. Plants 9: 67
- Wang Y, Burgess SJ, de Becker EM, Long SP (2020) Photosynthesis in the fleeting shadows: an overlooked opportunity for increasing crop productivity? Plant J. 101: 874-884 https://doi.org/10.1111/tpj.14663
- Wehner A, Grasses T, Jahns P (2006) De-epoxidation of Violaxanthin in the Minor Antenna Proteins of Photosystem II, LHCB4, LHCB5, and LHCB6 J. Biol. Chem. 281: 21924-21933 10.1074/jbc.M602915200
- **Zaks J, Amarnath K, Kramer DM, Niyogi KK, Fleming GR** (2012) A kinetic model of rapidly reversible nonphotochemical quenching. Proc. Nat. Acad. U.S.A **109**: 15757-15762 10.1073/pnas.1211017109
- **Zaks J, Amarnath K, Sylak-Glassman EJ, Fleming GR** (2013) Models and measurements of energy-dependent quenching. Photosynth. Res. **116:** 389-409 10.1007/s11120-013-9857-7
- Zhang XJ, Fujita Y, Tokutsu R, Minagawa J, Ye S, Shibata Y (2021) High-Speed
 Excitation-Spectral Microscopy Uncovers In Situ Rearrangement of Light-Harvesting
 Apparatus in Chlamydomonas during State Transitions at Submicron Precision. Plant
 Cell Physiol. 62: 872-882 10.1093/pcp/pcab047

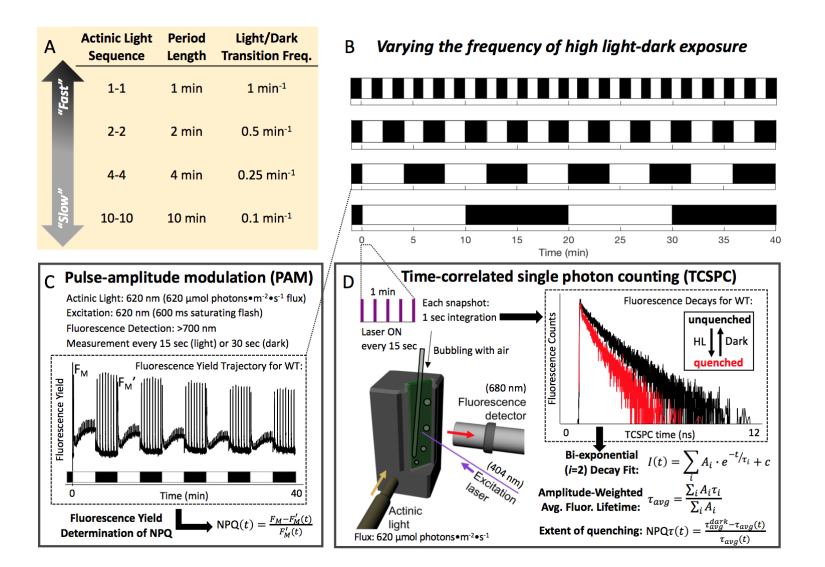


Figure 1. Experimental design for ChI *a* fluorescence measurements throughout exposure of *Chlamydomonas* cells to fluctuating light with various periods of HL-dark exposure. (**A**, **B**) Representation of the HL-dark cycles used for the 40 minutes of light fluctuation and their corresponding period, frequency, and name used throughout the main text. NPQ was measured using Pulsed Amplitude Modulation (PAM, **C**) and Time-Correlated Single Photon Counting (TCSPC, **D**). (**C**) Characteristics of the PAM measurement and representative data of fluorescence yield in WT cells. (**D**) Characteristics of the TCSPC apparatus. Shown are two decays representative of two snapshots taken in a quenched and unquenched state.

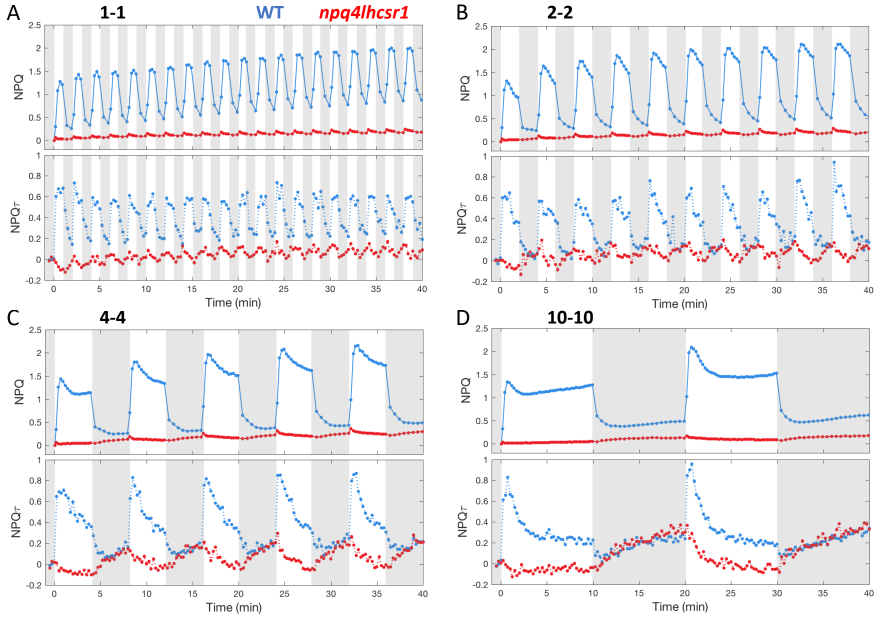


Figure 2. Quenching trajectories during light fluctuations in *npq4lhcsr1* and its control strain. The response of NPQ and NPQτ (upper and lower panel respectively) were measured in *npq4lhcsr1* mutant and its control strain (red and blue curves respectively) during 40 minutes of light fluctuations with periods of 1, 2, 4 and 10 minutes (**A**, **B**, **C** and **D** respectively) as described in **Fig. 1**. Shown are average of three biological replicates. For TCSPC data, each biological replicate was averaged from three technical replicates. The fluorescence lifetime values used to calculate NPQτ are shown in the Supplemental.

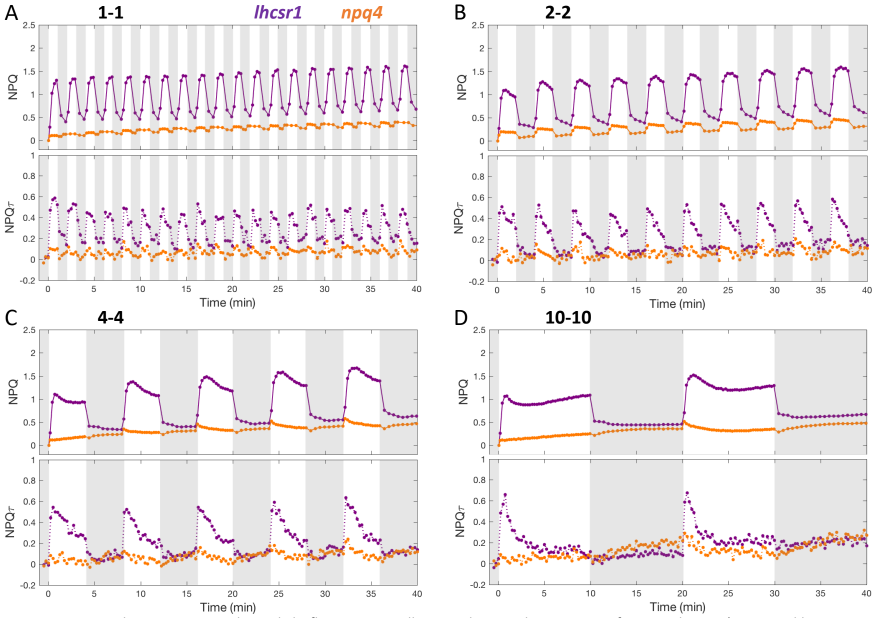


Figure 3. Quenching trajectories during light fluctuations in *lhcsr1* and *npq4*. The response of NPQ and NPQτ (upper and lower panel respectively) were measured in *lhcsr1* and *npq4* (purple and orange curves respectively) during 40 minutes of light fluctuations with periods of 1, 2, 4 and 10 minutes (**A**, **B**, **C** and **D** respectively) as described in **Fig. 1**. Shown are average of three biological replicates. For TCSPC data, each biological replicate was averaged from three technical replicates. The fluorescence lifetime values used to calculate NPQτ are shown in the Supplemental.

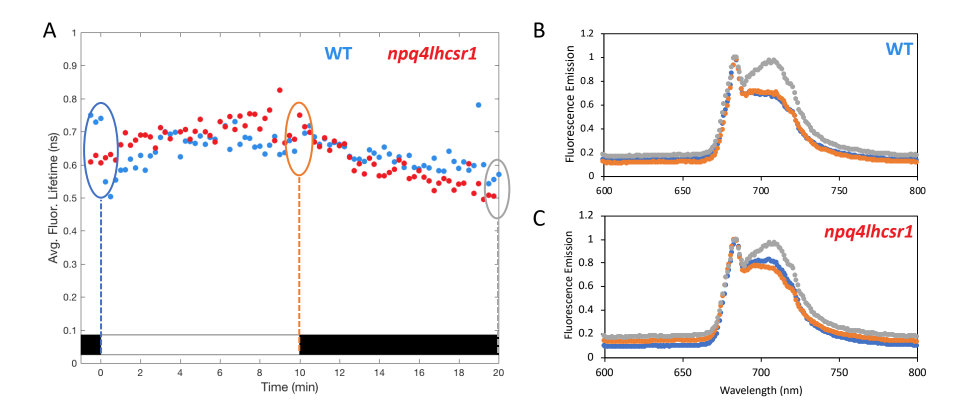


Figure 4. 77K Chlorophyll fluorescence emission spectra during the first high light-dark cycle of light fluctuations. Cells were placed in a TCSPC cuvette as described in Fig. 1 and both fluorescence lifetime snapshots and 77K chlorophyll fluorescence emission spectra were taken through 10 minutes of high light and 10 minutes darkness. (A) Fluorescence lifetime trajectory of npq4lhcsr1 mutant (red dots) and its control strain (WT, blue dots). On the graph, dashed vertical lines depict the timepoints at which samples were taken for 77K fluorescence spectra measurement. (B, C) 77K fluorescence emission spectra of samples taken in A on the control strain (WT, B) and npq4lhcsr1 mutant (C). Spectra were taken at 0, 10, and 20 min timepoints (blue, orange and grey spectra respectively). Shown are representative spectra. Three independent biological replicate spectra for WT and npq4lhcsr1 are shown in Supp. Fig. 3. 77K spectra for the stt7 and stt7npq4 strains are shown in Supp. Fig. 4.

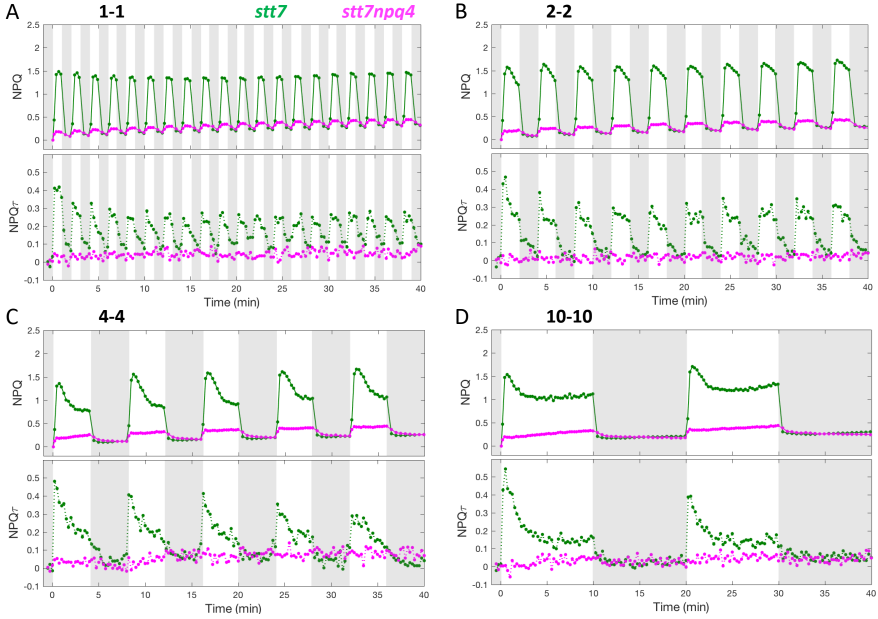


Figure 5. Quenching trajectories during light fluctuations in *stt7* and *stt7npq4*. The response of NPQ and NPQτ (upper and lower panel respectively) were measured in *stt7* and *stt7npq4* (green and magenta curves respectively) during 40 minutes of light fluctuations with periods of 1, 2, 4 and 10 minutes (A, B, C and D respectively) as described in Fig. 1. Shown are average of three biological replicates. For TCSPC data, each biological replicate was averaged from three technical replicates. The fluorescence lifetime values used to calculate NPQτ are shown in the Supplemental.

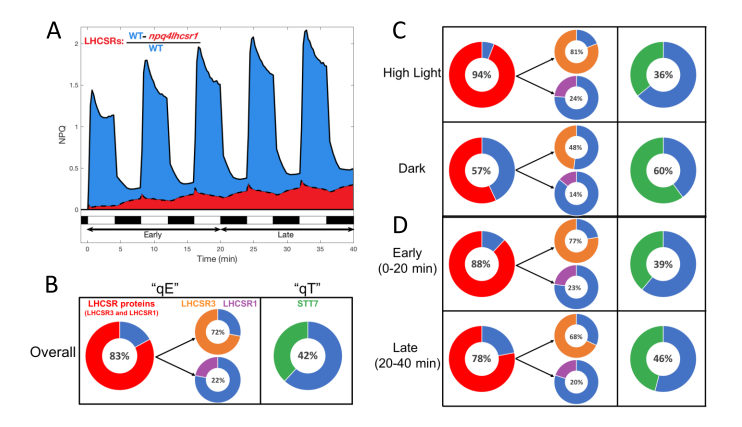


Figure 6. Quantification of the contribution of LHCSRs and STT7 to wild-type NPQ under fluctuating light. (**A**) Example of quantification of the relative NPQ mediated by LHCSR proteins. The area under the NPQ curve of *npq4lhcsr1* mutant (red) was subtracted from that of the control strain (blue) and expressed relative to the area of NPQ of the control strain. (**B**) Overall contribution of LHCSRs (red), LHCSR3 (orange), LHCSR1 (purple) and STT7 (green) averaged over all 40 minutes, (**C**) during HL and dark portions of the light fluctuations, or (**D**) during early (0-20 min) and late (20-40 min) portions of the experiment. Each donut portrays the amount of wild-type NPQ that is lost in each mutant impaired in the accumulation of the given protein. Given that the contribution of each protein was largely independent of HL/dark period (**Supp. Fig. 8**), shown here are the average of all 4 light fluctuation sequences. Distribution of individual replicates and estimates of error are presented in **Supp. Fig. 8-10**.

Table 1. Average contribution of each protein to overall wild-type NPQ for each light fluctuation sequence. Shown is the average value (n=6, evaluated from 3 TCSPC and 3 PAM replicates) and standard deviation of all individual replicates. The contributions of LHCSR3 (orange) and LHCSR1 (purple) were determined from the single mutants *npq4* and *lhcsr1*. The contribution of LHCSRs overall (red) was evaluated from the *npq4lhcsr1* mutant. The contribution of qT was assessed from the *stt7* mutant. Each error in the right column represents the standard deviation of each protein's contribution across the 4 light fluctuation sequences. For simplicity, only the average values (shown in the right column) were used to generate **Figure 6** in the main text. Full distributions of the individual TCSPC and PAM data points are shown in **Supp. Fig. 8**. [*supports Fig.* 6B]

Overall Contributions	1-1	2-2	4-4	10-10	AVERAGE
LHCSRs	89 ± 7%	84 ± 7%	83 ± 13%	77 ± 24%	83 ± 5%
LHCSR1	22 ± 12%	22 ± 10%	24 ± 16%	19 ± 30%	22 ± 2%
LHCSR3	81 ± 5%	78 ± 5%	73 ± 8%	58 ± 15%	72 ± 10%
STT7	46 ± 12%	38 ± 16%	46 ± 13%	39 ± 22%	42 ± 4%

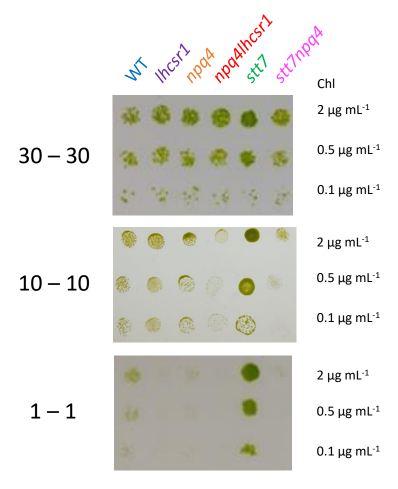


Figure 7. Growth of mutants impaired in qE and/or qT under various periods of dark/light cycles. *Ihcsr1*, *npq4*, *npq4lhcsr1*, *stt7* and *stt7npq4* mutants and their control strain (WT) were diluted and spotted at different chlorophyll concentration and grown on plates under dark/light cycles with a period of 30 (30-30, upper panel), 10 (10-10, middle panel) or 1 minute (1-1, lower panel). Each row represents a different chlorophyll concentration. Shown are representative spots of three biological replicates. Growth under constant low light or high light are shown in **Supp. Fig. 11**.