1 BrphyB is critical for rapid recovery to darkness in mature Brassica rapa

2 leaves

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- 10

11 HIGHLIGHT

- 12 BrphyB plays a central role in recovery from darkness and return to carbon fixation
- by regulating photosynthesis and light response genes with those targeted to the
- 14 chloroplast especially affected.
- 15

16 **ABSTRACT**

17 Crop biomass and yield are tightly linked to how the light signaling network translates 18 information about the environment into allocation of resources, including 19 photosynthates. Once activated, the phytochrome (phy) class of photoreceptors 20 signal and re-deploy carbon resources to alter growth, plant architecture, and 21 reproductive timing. Brassica rapa has been used as a crop model to test for 22 conservation of the phytochrome-carbon network. B. rapa phyB mutants have 23 significantly decreased or absent CO₂-stimulated growth responses in seedlings, and 24 adult plants have reduced chlorophyll levels, photosynthetic rate, stomatal index, and 25 seed yield. Here, we examine the transcriptomic response of adult wild-type and 26 *BrphyB* leaves to darkening and recovery in light. Three days of darkness was 27 sufficient to elicit a response in wild type leaves suggesting a shift from carbon 28 fixation and nutrient acquisition to active redistribution of cellular resources. Upon a 29 return to light, wild-type leaves appeared to transcriptionally return to a pre-darkness 30 state restoring a focus on nutrient acquisition. Overall, BrphyB mutant plants have a 31 similar response with key differences in genes involved in photosynthesis and light 32 response which deviate from the wild type transcriptional dynamics. Genes targeted 33 to the chloroplast are especially affected. Adult *BrphyB* mutant plants had fewer,

34 larger chloroplasts, further linking phytochromes, chloroplast development,

35 photosynthetic deficiencies and optimal resource allocation.

36

37 **KEY WORDS:** Brassicaceae, chloroplast development, gene regulation, light

38 response, photosynthesis, phytochrome B.

39

40 **INTRODUCTION**

Light plays at least two distinct roles in shaping plant form and productivity. First, light is essential for photosynthesis, which allows plants to convert the energy held in photons into the high potential energy found in the chemical bonds of sugars. Second, light provides information on how a plant can optimize its architecture to maximize photosynthetic potential in a given environment. How these two light systems are coordinated remains largely unknown, especially in mature leaves.

47 Limited light supply by an established canopy triggers a rapid shade-avoidance 48 response that is characterized by increased elongation growth rate of stems and 49 petioles, decreased leaf surface area and thickness, and delayed leaf yellowing 50 (Casal, 2012; Franklin and Whitelam, 2005). On the other hand, partial plant shading 51 or darkening will induce a range of responses between acclimation to leaf 52 senescence (Weaver and Amasino, 2001; Brouwer et al., 2012). These processes 53 directly reduce the impact of shade or dark while additional responses such as 54 acclimation of the photosynthetic apparatus rather help to fine tune the use of 55 resources under shade/dark.

56 Plants use an array of photoreceptors to capture and transduce the light signal in

57 diverse responses known collectively as photomorphogenesis. Photoreceptors'

absorbance properties span most of the visible light spectrum, from the

59 phytochromes that absorb in the red (R)/far-red (FR) to the cyptochromes and

60 phototropins that absorb in the blue/near-ultraviolet to the UV-receptors. Among

61 these, the phytochromes (phys) are among the best characterized (Chen *et al.*,

62 2004). Upon illumination, phys undergo conformational changes from an inactive (Pr)

to an active (Pfr) form (Fraser *et al.*, 2016), which is subsequently translocated into

the nucleus and participates in transcriptional regulation (Chen *et al.*, 2005; Castillon

et al., 2007). Five PHY genes have been described in the A.thaliana genome

66 (*PHYA-PHYE*) with partial overlapping functions (reviewed in Chen *et al.*, 2004).

67 Phytochrome-dependent light signaling that initiates photomorphogenesis has been 68 extensively studied using the seedling model (reviewed in Arsovski et al., 2012). In 69 addition, it is clear from work in *A.thaliana* that phytochromes control chloroplast 70 gene expression, as well as nuclear-encoded factors involved in chloroplast 71 development (Oh and Montgomery, 2014; Nevarez et al., 2017). Recent studies in 72 A.thaliana and Brassica rapa showed that adult phyB mutants have reduced 73 chlorophyll levels, photosynthetic rate, and stomatal index. Work by a number of 74 groups has connected PhyB to biomass accumulation, carbon resource 75 management, seed yield and changes in metabolism across the plant life cycle 76 (Yang et al., 2016; Krahmer et al., 2018; Arsovski et al., 2018; Wies et al., 2019). 77 To date, most of our knowledge about the roles of phytochromes in the dark-to-light 78 transition primarily came from experiments focused on the de-etiolation process of 79 seedlings (Li et al., 2011). This has left a gap in our understanding about the role of 80 phytochromes in light-activated transcription of genes in mature leaves. This is 81 important because several light-regulatory mechanisms essential for photosynthetic 82 efficiency and adaptation occur only in mature leaves. For example, Chory et al. 83 demonstrated that the primary role of phytochrome in greening A. thaliana plants is 84 in modulating the degree rather than the initiation of chloroplast development (Chory 85 et al., 1989).

86

87 In this study, we investigated the effects of phyB on gene expression upon dark-to-88 light transition in the mature leaf of *B. rapa* by comparing the transcriptomic 89 responses between wild-type and a phyB mutant. B. rapa is closely related to 90 A.thaliana (Wang et al., 2011) but its leaves are significantly larger. Larger leaves 91 cause more self-shading, and, in combination with the longer life of *B. rapa* 92 compared to A.thaliana, there is more total demand for resources. As the B. rapa 93 genome contains only one *PhyB* ortholog and no likely ortholog for the closely 94 related AtPhyD, we took advantage of the BrphyB3 mutant allele described 95 previously (Arsovski et al., 2018). Wild-type and BrphyB leaves exhibited significant 96 overlaps in their transcriptomic response to dark and recovery; however, gene 97 ontology analyses pointed out important misregulations in *BrphyB* mutant for genes 98 involved in nitrogen metabolism, light harvesting and photosynthesis. Altogether 99 these results support a role for PhyB in chloroplast development and resource

100	allocation,	and have im	plications f	for increasi	ng the	resource-efficiency	of Brassica
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- 101 crops.
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103 MATERIALS AND METHODS

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105 Growth conditions of *B. rapa* adult plants

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107 The *B. rapa* wild-type R-o-18 and *BrphyB* mutants were originally from the John

108 Innes Center's RevGenUK resource. The BrphyB-3 previously described in Arsovski

109 et al., 2018 was used for RNAseq experiments. BrphyB-1 was also previously

110 described in Arsovski et al., 2018. Seeds were planted directly into our standard soil

- 111 mix of 1:1 Sunshine Mix #4 (SunGro Horticulture):vermiculite. Plants were grown in
- 112 2.6 liter square pots (McConkey Grower Products; Sumner, WA, USA) and bottom-
- 113 watered daily in long day conditions (16 h light, 8 h dark, ~115 μ mol.m⁻².s⁻¹ light

114 intensity) in a Percival E-30B growth chamber (<u>https://www.percival-scientific.com/</u>)

set to 20°C. Experiments were conducted at 3 weeks and the plants were then

- 116 moved to growth room until seed harvest.
- 117

118 Leaf sample preparation

119

120 Three weeks after sowing, two developmentally matched leaves from each wild-type 121 and BrphyB-3 plant were tagged that corresponded to the first and second true 122 leaves. Samples were collected at ZT 5 using a standard hole punch (28 mm²) 123 circular area of leaf blade tissue) with symmetrical harvest (a second hole punch on 124 the other side of the other side of the mid-vein of the same leaf) for chlorophyll 125 assay, chloroplast measurements and transcriptome analysis. Tissue from 3 126 individual plants was combined to make one biological replicate. At 3 weeks of age 127 the "Pre" sample was collected from the first leaf while the second one was covered 128 with tinfoil. 24 hours later the "24hr" sample was harvested from the uncovered leaf, 129 the same leaf that provided the "Pre" sample. Then, 48 hours later the tinfoil was 130 removed from the covered leaf and the "dark" samples were similarly collected. 131 Finally, 24 hours later the "recovery" samples were collected from this same leaf. 132 Samples were immediately flash frozen in liquid nitrogen (Fig.1). In total, three 133 biological replicates were collected in similar fashion.

134

135 Chlorophyll measurement

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137 For chlorophyll measurement, ethanol extractions were done as in (Yang et al., 138 2016). Determinations were run by measuring optical density at 645 nm and 665 nm 139 using an Epoch Microplate Spectrophotometer (www.biotek.com). Values were 140 the following formulas: Chl $a=5.21A_{665}-2.07A_{645}$; obtained using 141 Chl b=9.29A₆₄₅-2.74A₆₆₅, for Chlorophyll A and B, respectively. Three individual 142 biological replicates were used for this assay.

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144 RNAseq

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146 Leaf tissue was disrupted with Zirconia/Silica beads for 1 minute in a

147 MiniBeadbeater-96 (BioSpec Products, Inc.) while frozen. After adding 500 µL of

148 Lysis/Binding Buffer to each sample and vortexing until homogeneous, samples

149 were run on the MiniBeadbeater-96 for an additional minute. Following tissue

disruption samples were centrifuged at 16,000 x g for 10 minutes at 20 °C. For each

151 sample, a 50 μ L aliquot of the supernatant was added to 50 μ L of NEB RNA binding

buffer and mRNA isolated as per the NEBNext® Poly(A) mRNA Magnetic Isolation

- 153 Module manual.
- 154

RNA-seq libraries were prepared by using the Full Transcript mode YourSeq Dual
(FT & 3'-DGE) RNAseq Library Kit (Amaryllis Nucleics). A Bioanalyzer 2100 (Agilent,
High Sensitivity DNA Kit) was used for library quality control, to determine average
library size, and together with concentration data from a Qubit 2.0 Fluorometer (Life
Technologies, dsDNA High Sensitivity Assay Kit) to determine individual library
molarity and pooled library molarity. Pooled libraries were sequenced on a NextSeq
500 (Illumina, High Output v2 75 cycle kit) to yield single-read 80 bp reads.

163 FASTQ processing was performed by Amaryllis Nucleics (Oakland, CA). Sequence

164 files were preprocessed in two steps. A Python library (clipper.py,

165 <u>https://github.com/mfcovington/clipper</u>) was used to trim off the first 8 nucleotides of

166 each read to remove potential mismatches to the reference sequence caused by

- 167 annealing of a random hexamer required for library synthesis. Trimmomatic v0.36
- 168 (<u>http://www.usadellab.org/cms/?page=trimmomatic</u>) was used to remove adapter
- 169 sequences and trim or filter reads based on quality The parameters used for
- 170 Trimmomatic were ILLUMINACLIP:TruSeq3-PE-2.fa:2:30:10 LEADING:3
- 171 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:50.
- 172 Preprocessed reads were mapped to the Brassica rapa v2.5 genomic reference
- 173 sequence
- 174 (http://brassicadb.org/brad/datasets/pub/Genomes/Brassica_rapa/V2.0/V2.5/Chr/Bra
- 175 <u>paV2.5_Chr.fa.gz</u>) using bowtie2. Read counts for each gene in the gene annotation
- 176 file
- 177 (http://brassicadb.org/brad/datasets/pub/Genomes/Brassica_rapa/V2.0/V2.5/Chr/Bra
- 178 <u>paV2.5_Chr.gene.gff.gz</u>) were calculated using htseq-count (with the -s yes
- 179 parameter to enforce strand-specific alignment) from the HTSeq Python library
- 180 (https://academic.oup.com/bioinformatics/article/31/2/166/2366196;
- 181 http://htseq.readthedocs.io/en/master/index.html).
- 182

183 The package edgeR (Robinson *et al.*, 2010) was used to process the expression 184 matrix and identify differentially expressed genes between treatments and 185 genotypes. For the main analysis, the generalized linear model functionality of this 186 package, based on a negative binomial distribution model for gene expression was 187 used to identify differentially expressed genes. Genes were considered significantly 188 differentially expressed based on having a fold change greater than 2-fold up or down between conditions, and a q-value (adjusted p-value by Benjamini-Hochberg 189 190 procedure - (Benjamini and Hochberg, 1995) less than 0.01.

191

192 Contrasts between the Pre and 24hr timepoints were used to identify genes that 193 could be exhibiting differential expression caused by wound response from the tissue 194 sampling rather than response to darkness and recovery. Genes identified in these 195 wound control contrasts were tagged but not excluded from the rest of the analysis. 196 Dispersion was estimated independently for the wound control contrasts based on 197 the Pre and 24hr timepoints only. This was done because using the dispersion 198 estimates from the main analysis, including the Dark and Recovery timepoints 199 resulted in biased p-value distributions due to the significant change in expression of

200 most genes in the dark timepoint as compared to the other timepoints (68% of genes

201 between wild type-Dark and wild type-Pre).

202

203 Complete data can be accessed from the Gene Expression Omnibus (GEO) under

204 the entry GSE135955.

205

Venn diagrams and gene ontology (GO) analysis to prioritize the DEG

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To find differences between the wild-type and *phyB* response to darkness and reillumination we looked at sets of genes that were significantly changed in these contrasts (darkened vs. pre or recovery vs. darkened) in one genotype but not the other. For each up or down Venn-diagram used in the GO analysis, all genes which either the wild-type or *BrphyB* mutant were greater (or less for down regulated Venn) than the log fold change cutoff of 2 were considered. These genes were then put into 4 categories: wild-type significant only, *BrphyB* significant only, both wild-type and

- 215 *BrphyB* significant, and neither genotype significant.
- 216

To obtain a comprehensive list of all *B. rapa* presumptive orthologs in *A. thaliana* we

used each *B. rapa* protein as a query to perform local homology searches. Briefly, for

each protein in the *B. rapa* v2.5 proteome, the best *A. thaliana* hit was retrieved by

220 sequence similarity search using a local installation of the BlastP algorithm (Protein-

221 Protein BLAST 2.7.1+) of the NCBI tool BLAST

222 (<u>http://blast.ncbi.nlm.nih.gov/Blast.cgi</u>) against the A. thaliana ARAPORT11

proteome, with default parameters and the outfmt parameter set to 7 (to obtain a

tabular output with comment lines). This resulted in a collection of the best A.

thaliana hits for each *B. rapa* protein. This output was further processed to limit each

ortholog to the best hit, using a custom BASH script (available upon request). GO

227 term enrichment analysis was performed against the well annotated A.thaliana

228 genome using the PANTHER classification System (v.14.1 available at

229 http://pantherdb.org/; (Mi et al., 2019).

230 231

232 Law 2018 comparison

234 For the data in Table S3, we created a list by matching the *A. thaliana* gene IDs from 235 the (Law et al., 2018) analysis up to the B. rapa genes they were found to be the 236 nearest homolog of (see GO analysis section above). We considered the genes that 237 were differentially expressed after 3 days (D3) of darkness applied on an 238 individualized leaf (IDL) or a whole plant (DP). A. thaliana genes that did not match 239 to *B.rapa* genes in our set were dropped from the comparison, *A. thaliana* genes with 240 multiple brassica homologs were listed multiple times in this list. The elements of this 241 list were then broken down into a Venn diagram based on whether they were 242 considered significantly differentially expressed up or down in our darkened vs. pre 243 contrast and Law's IDL D3 and DP D3 contrasts. Each category list was then 244 reduced to only contain unique A. thaliana IDs. These counts are displayed in Table 245 S3.

246

247 RNA extraction and quantitative real-time PCR (qRT-PCR) analysis

248 Expression analysis was performed using 4 biological replicate samples collected

- 249 identically as described for RNA sequencing. Each sample was immediately frozen
- 250 in liquid nitrogen and stored at -80 °C until processing. Frozen tissue was ground in
- 251 liquid nitrogen and total RNA was extracted using the GE Illustra RNA kit (GE Life
- 252 Sciences), and 2 µg of eluted RNA was used for cDNA synthesis employing iScript
- 253 (Bio-Rad). Samples were analyzed using SYBR Green Supermix (Bio-Rad) reactions
- run in a C100 Thermal Cycler (Bio-Rad) fitted with a CFX96 Real-Time Detection
- 255 System (Bio-Rad). Relative expression levels were calculated using the formula
- 256 (E_{target})^{-CPtarget}/(E_{ref})-^{CPref} (PfaffI, 2001) and normalized to the *B.rapa PP2A*
- 257 (Brara.F00691) reference gene. qPCR primer sequences are as follows: *BrPP2A*
- 258 (forward 5'-TCGGTGGTAACGCCCCCGAT-3'; reverse 5'-
- 259 CGACTCTCGTGGTTCCCTCGC-3'); BrMGT6 (Brara.E02300) (forward 5'-
- 260 CAGCATCCGCCACCGCAAGA-3'; reverse 5'-GCCTTCGCAACAACCGCAGC-3');
- 261 BrRLK4 (Brara.100004) (forward 5'- TCCGCCGTCGCGATCTCTCT-3'; reverse 5'-
- 262 CCCGCTCCAAACGCTTGTCCA-3'); *BrPIFI* (forward 5'-
- 263 GCCACCACTCTTGCACCCCC-3'; reverse 5'- CCGCGGTTGGAGGAAGACCG-3').
- 264

265 **Code**

The R code used to generate all the analysis results is provided in supplement X and can also be found on <u>github.com/nemhauser-lab/brassica_rna_seq</u>.

268

269 Methods, Motif Enrichment

270 A set of binding motifs for 619 A.thaliana transcription factors was downloaded from 271 plantTFDB (http://planttfdb.cbi.pku.edu.cn/), (Jin et al., 2014, 2015, 2017). The RSAT 272 matrix clustering tool (http://rsat.eead.csic.es/plants/matrix-clustering_form.cgi 273 (Castro-Mondragon et al., 2017) was used with default parameters to group the 274 motifs into 56 clusters based on similarity of aligned position weight matrices. Genes 275 that were significantly differentially expressed from conditions pre to dark, and from 276 dark to recovery were divided into groups depending on 3 factors: genotype, 277 direction of regulation (up or down), and whether they are annotated with the GO 278 term "chloroplast" (GO:0009507). The package "motifmatchr" (Schep, 2019) was 279 used to count the number of genes in each set with promoter sequences with 280 matches to each of the 619 motifs. Promoters sequences were defined as the 1000 281 base pairs immediately upstream of the gene start position as defined by the 282 genome annotation file. The fisher exact test was then used to determine if there 283 was significant enrichment for each motif between genes only significantly regulated 284 in WT and genes only significantly regulated in *BrphyB* in these sets. The raw p-285 values from these tests were adjusted using the by Benjamini-Hochberg procedure 286 (Benjamini and Hochberg, 1995). Adjusted p-values of <0.01 were considered 287 significant.

288

All motifs found significant from these tests closely matched to three canonical motifs described in the literature: the G-box motif CACGTG, the Evening Element AGATATTTT, and the Telo-box motif AAACCCTAA. The proportions of promoters with one or more exact matches to three canonical motifs were found for the gene sets described above.

294

295 Chloroplast Measurement

Tissue was immediately cleared after collection and fixed using ClearSee solution as
described in (Kurihara *et al.*, 2015). Images were taken using a Leica TCS SP5 II

- 298 laser scanning confocal microscope (<u>https://www.leica-microsystems.com</u>).
- 299 Chloroplast number, area, and density were determined using ImageJ software.

300

301

302 **RESULTS**

303

304 Mature *BrPhyB* mutant leaves have significant transcript reductions of 305 chloroplast targeted genes.

306

307 Loss of phyB leads to significant reductions in both chlorophyll levels and rates of 308 photosynthesis in three-week-old *B. rapa* plants (Arsovski et al., 2018). To further 309 understand the link between phyB, chloroplast development, and photosynthesis we 310 examined the transcriptomic response of mature leaves that were subjected to three 311 days of darkness before being reintroduced into the light. As part of this experiment 312 we first compared the transcriptome of three-week-old *B.rapa* wild type and BrphyB 313 leaves. Genes were considered differentially expressed if the fold change between 314 timepoints or genotypes was greater than 2, and the significance (adjusted p-value) 315 was less than 0.01. 114 genes were significantly upregulated in BrphyB leaves 316 compared to wild type. Unsurprisingly, these include *B.rapa* orthologs to *A.thaliana* 317 LONG HYPOCOTYL IN FAR-RED(HFR1), PHYTOCHROME INTERACTING 318 FACTOR 3-LIKE 1 (PIL1), PHYTOCHROME-INTERACTING FACTOR 6 (PIF6), and 319 INDOLE-3-ACETIC ACID INDUCIBLE 29(IAA29). 79 genes were significantly 320 downregulated in BrphyB leaves compared to the wild type. Gene Ontology (GO) 321 analysis of cellular location annotations showed a strong enrichment for the 322 chloroplast envelope, stroma, and photosystem II. These include *B.rapa* orthologs 323 for A.thaliana PHOTOSYSTEM II SUBUNIT P, PHOTOSYSTEM II BY, LIGHT-324 HARVESTING CHLOROPHYLL B-BINDING PROTEIN 3, and CHLOROPHYLL A/B 325 BINDING PROTEIN 1 (Table S1).

326

327 Darkening of individual leaves for three days initiates resource reallocation 328

Samples were taken from the first or second true leaf of three-week-old wild-type
and *BrphyB*-3 plants (hereafter termed "pre"). As a wounding control, a second
sample was taken from the "pre" leaves 24 hours later (hereafter termed "24hr").
Leaves that were developmentally-matched with those selected for the "pre"
treatment were covered with foil. After three days, the foil was removed and the
"darkened" sample was collected immediately. The "recovery" sample was collected
from the same leaf 24 hours after this timepoint to capture the earliest stages of

336 recovery (Fig. 1A). Matching samples were collected from each leaf to assay 337 chlorophyll levels. At three weeks old, *BrphyB* mutants are visibly paler compared to 338 same aged wild-type plants, and have significantly reduced chlorophyll levels (Fig. 339 1B and Arsovski et al., 2018). Three days of dark resulted in a 23% reduction of 340 chlorophyll levels in wild type leaves while levels remained low in the mutant. This is 341 consistent with similar experiments performed on individually darkened A.thaliana 342 leaves where total chlorophyll levels and protein decline was observed after two days of darkness (Weaver and Amasino, 2001). The 24 hours of light exposure for 343 344 the recovery samples was not sufficient to restore chlorophyll levels in either wild-345 type or *BrphyB*-3 leaves (Fig. 1B).

346

347 Extended darkness of leaves acts as a signal to initiate the organized breakdown 348 and remobilization of valuable resources to growing vegetative and reproductive 349 tissues (Himelblau and Amasino, 2001; Buchanan-Wollaston et al., 2003; Lim et al., 350 2007). We performed RNAseg analysis on the pre, 24hr, dark and recovery samples 351 to assess the specific response to darkness and return to light of mature leaves in *B*. 352 rapa. We began our analysis with the response to darkness, as previous studies in 353 A. thaliana had already shown that dark stress is accompanied by dramatic 354 transcriptional changes, as well as depletion of chlorophyll and large-scale 355 degradation of proteins (Guo et al., 2004; Keech et al., 2007; Law et al., 2018). The 356 expression of 6852 *B.rapa* genes was significantly altered in leaves after three days 357 in darkness. Gene Ontology (GO) analysis of predicted A. thaliana orthologs showed 358 a pattern consistent with the overall expectations of metabolic reprogramming seen 359 in other species. The 3110 genes up-regulated in response to dark were mainly 360 involved with autophagy, catabolism, leaf senescence and vesicle fusion (Fig. 2A; 361 Table S2). Conversely, down-regulated genes were mainly involved in 362 photosynthesis, biosynthetic processes and plastid translation. Together, these data 363 suggest a shift from carbon fixation and nutrient acquisition to active redistribution of 364 cellular resources (Fig. 2B, Table S2).

365

A recent experiment in *A.thaliana* found that the effect of darkening individual leaves was substantially similar to the effect of darkening whole plants, albeit with distinct timing for peak differences in gene expression between the two treatments Law *et al.*, 2018). When we compared our transcriptomic response to this dataset, we found

370 a substantial overlap. The highest similarity between the *B.rapa* dark response was 371 to A.thaliana individually darkened leaves for 3 days (IDL_3D). Of the genes 372 significantly up or down regulated in *A. thaliana* individually darkened leaves after 3 373 days, 46.7% had *B.rapa* homologs also significantly up or down regulated (in the 374 same direction) (Table S3). Shared A. thaliana genes upregulated in response to 375 dark were significantly enriched for GO terms such as autophagy, catabolic process, 376 and leaf senescence (Table S4). The darkening response in *B.rapa* was more 377 similar to that of individually darkened leaves than whole darkened plants in 378 A.thaliana. Of the genes found to be up/down regulated in individual leaves (IDL_D3) 379 but not whole darkened plants (DP D3), 35.3% had *B.rapa* homologs with significant 380 change in the same direction, compared to 25.1% in whole darkened plant unique 381 genes (Table S3,4). In A. thaliana, 167 senescence-associated genes were shown to 382 change expression in response to darkening (Parlitz et al., 2011). The B.rapa 383 orthologs of 103 of these genes do not show significant changes in response to dark. 384 Of the remaining 57 senescence-associated genes, 42 show a reversible pattern of 385 upregulation in the dark and downregulation upon re-illumination. 15 are 'non-386 reversible', upregulated in the dark without significant changes upon a return to light 387 (Table S5).

388

389 Many of the genes that were regulated by returning the leaves to light were similar to 390 those already identified as light-responsive from experiments in seedlings. In A. 391 thaliana, expression of up to one-fourth of the whole genome is altered in seedlings 392 grown for 4 days in red light compared to those grown in the dark (Shi et al., 2018). 393 These changes are largely mediated by a small group of transcription factor families 394 which include the PHYTOCHROME INTERACTING FACTORS (PIFs). Nearly 60% 395 of PIF-dependent, red light induced genes in A. thaliana seedlings have Gene 396 Ontology (GO) annotations indicating functions related to photosynthesis and 397 chloroplast (Leivar et al., 2012). In B.rapa wild type leaves 3756 genes were 398 upregulated in the recovery condition when compared to the dark timepoint. The 399 most significantly enriched GO terms were response to light stimulus, 400 photosynthesis, translation and metabolism, suggesting a return to a pre-darkness 401 transcriptional state (Figure 3A, Table S2). The 3299 genes downregulated in 402 recovery compared to dark were mainly involved in catabolism, vesicle fusion and

403 transport, and protein degradation further supporting a shift from resource

404 remobilization towards nutrient acquisition (Figure 3B, Table S2).

405

406 *BrphyB* is critical for full recovery response

407

408 RNAseq analysis of *BrphyB* individually darkened leaves revealed an essentially 409 similar response to what was observed in wild-type plants. 7,994 genes were 410 differentially expressed in wild-type and *BrphyB*-3 leaves after 3 days of dark 411 treatment compared to the pre samples. The vast majority (81.8%) of this response 412 was shared between the genotypes (Fig. 2A, B). However, analysis of GO terms 413 significantly enriched in the uniquely wild-type or *BrphyB* differentially regulated gene 414 sets illustrated phyB-dependent responses to darkness. Unique wild-type enriched 415 terms were largely related to cellular responses to organic and inorganic 416 compounds, drugs and stress (Table S6). A closer look at these phyB-controlled 417 groups revealed orthologs to A. thaliana genes REVEILLE2, REVEILLE8, and 418 CIRCADIAN CLOCK ASSOCIATED1—all key transcriptional regulators of circadian 419 rhythm, auxin and stress response in A. thaliana and known to act downstream of 420 PhyB (Fig. 2A,C) (Alabadí *et al.*, 2001; Zhang *et al.*, 2007; Rawat *et al.*, 2011; 421 Farinas and Mas, 2011; Jiang et al., 2016). BrphyB unique up-regulated genes were 422 enriched in genes associated with response to light, including B.rapa orthologs of 423 PHYTOCHROME INTERACTING FACTOR4, PHYTOCHROME INTERACTING 424 FACTOR5, PHYTOCHROME KINASE SUBSTRATE1 and CRYPTOCHROME 1 425 (Table S6, Fig. 2C).

426

427 PhyB-repressed genes are enriched for categories such as response to light 428 stimulus and cellular biosynthetic process, while those activated by phyB are 429 enriched for categories related to protein synthesis such as cysteine metabolic 430 process, translational elongation, and peptide biosynthesis (Fig.2B). These include 431 *B. rapa* orthologs to three *A. thaliana* glutamate-ammonia ligases (GLUTAMINE) 432 SYNTHASE 1:2, 1:3 and 1:4) with roles in nitrogen remobilization and seed yield 433 (Guan et al., 2015), stress response and pollen viability (Ji et al., 2019). In B. rapa, 434 *BrphyB* mutants have up to a 90% decrease in seed yield (Arsovski *et al.*, 2018); 435 however, we did not observe a difference in area or weight of seeds compared to 436 wild type (data not shown). This would suggest that plants are re-calibrating the

amount of resources available, and maintaining quality by partitioning them into asmaller number of seeds.

439

440 While there is substantial overlap between the response of wild type and *BrphyB* 441 mutants to re-illumination (80.2% of the 7,765 genes are common to both 442 genotypes), there are also several key differences. To validate the RNAseq results, 443 we selected three genes whose expression in wild type was significantly 444 downregulated during darkening compared to pre followed by a significant 445 upregulation in recovery compared to darkening. qPCR of these genes was done in 446 wild-type, *BrphyB*-3, and an additional mutant allele *BrphyB*-1. Brara.E02555 is an 447 ortholog of the A.thaliana At3g15840 gene. In A.thaliana POST-ILLUMINATION 448 CHLOROPHYLL FLUORESCENCE INCREASE (PIFI) is a nuclear-encoded 449 chloroplast protein essential for NDH-mediated non-photochemical reduction of the 450 plastoquinone pool in chlororespiratory electron transport (Wang and Portis, 2007). 451 The A.thaliana orthologs RECEPTOR-LIKE PROTEIN KINASE 4 (RLK4) and 452 MAGNESIUM TRANSPORTER 6 (MGT6) are a Ser/Thr receptor-like protein kinase 453 expressed in the root and a magnesium transporter required for growth in 454 Magnesium limited conditions, respectively. The qPCR expression closely resembled 455 the RNAseq results for wild-type and *BrphyB*-3 and *BrphyB*-1 expression matched 456 that of BrphyB-3 for all three genes. BrPIFI expression decreases dramatically in 457 response to darkening in both wild-type and *BrphyB*-1 mutant leaves. However, 458 while BrPIFI expression increases in response to the leaf's return to light in wild-type, 459 it remains low in the mutant. Similarly, *BrRLK4* and *BrMGT6* expression increases in 460 recovery in wild-type leaves. However, in BrphyB-3 leaves expression of both 461 decreases 24 hours after the cover is removed from the leaf (Fig. S1) (Coello et al., 462 1999; Wang and Portis, 2007). 463

BrphyB is required for rapid return to the full photosynthetically-active transcriptional

program. There are 404 genes up-regulated in recovery of only wild-type leaves.

466 These genes are mainly involved in light harvesting in Photosystem I,

467 photosynthesis, cellular carbohydrate catabolism and generation of precursor

468 metabolites and energy and are not upregulated in *BrphyB* leaves 24 hours after

return to light (Fig. 3A, Table S6). We previously showed that the expression *B.rapa*

470 GOLDEN2-LIKE 1 (BrGLK1) increases 70% in response to high CO² in wild type

471 seedlings but decreases in BrphyB mutants (Arsovski et al., 2018). In A.thaliana 472 GLK1 is one of a pair of partially redundant transcription factors that affect the 473 expression of nuclear photosynthetic genes involved in chloroplast development 474 (Waters et al., 2008, 2009; Kobayashi et al., 2013). Here the B.rapa ortholog of 475 A.thaliana GOLDEN2-LIKE 2 is significantly upregulated upon return to light in wild 476 type but not BrphyB mutant leaves (Table S7). Closer examination of chloroplast 477 localized genes whose expression significantly changes in response to darkening 478 and recovery revealed a stark contrast in responsiveness between the two 479 genotypes. In response to darkening, 1861 chloroplast related genes were 480 significantly downregulated in either genotype. Of these, 94.7 % (1763 genes) were 481 unique to wild type leaves and were not significantly downregulated in the mutant. 482 Upon a return to light 1904 chloroplast localized genes were significantly upregulated 483 with 85.8% common to both genotypes. However, 131 localized genes were 484 upregulated only in wild type leaves while 140 were unique to the *BrphyB* mutant. 485 GO cellular component analysis identified 125 genes with predicted chloroplast-486 localization that are up-regulated in wild-type but not BrphyB leaves during recovery. 487 A.thaliana Photosystem II genes such as LIGHT-HARVESTING CHLOROPHYLL B-488 BINDING PROTEIN 3 (BraA10000990), LIGHT HARVESTING COMPLEX 489 PHOTOSYSTEM II SUBUNIT 6 (BraA09006187), LIGHT-HARVESTING 490 CHLOROPHYLL-PROTEIN COMPLEX II SUBUNIT B1(BraA05001183) and three 491 orthologs to A.thaliana CHLOROPHYLL A/B BINDING PROTEIN 1 (BraA04002510, 492 BraA07001020, BraA08002753) are upregulated on return to light only in wild type 493 leaves (Figure 3C, Table S7).

494

495

496 Downregulated genes unique to wild type are enriched in annotations associated 497 with nuclear transport, ribosome biogenesis, and rRNA processing, while terms 498 including phototropism and regulation of primary metabolism, cellular biosynthesis 499 and nitrogen compound metabolism are enriched in the BrphyB unique 500 downregulated genes (Fig. 3B). This overall pattern suggests that *BrphyB* may be 501 required for effective monitoring and switching between carbon- and nitrogen-502 demanding processes, and that this role may be essential for maximal reallocation of 503 resources to developing seeds. 504

505

506 BrphyB leaves have regulatory motif differences in chloroplast related genes 507 and fewer, larger chloroplasts

508

509 In the pre condition, the 79 genes significantly downregulated in *BrphyB* leaves 510 compared to the wild-type were enriched for localization to the chloroplast envelope, 511 stroma, and photosystem II (Table S1). The recovery condition created a sensitized 512 environment to detect the more immediate impacts of *BrphyB* on establishing or 513 maintaining the photosynthetic machinery. To investigate whether there were 514 regulatory differences in recovery between chloroplast and non-chloroplast genes in 515 wild-type and BrphyB leaves we examined the promoters (1Kb upstream for TSS) of 516 up and downregulated genes in recovery compared to dark. Genes with the GO 517 term 0009507: chloroplast were designated as 'chloroplast' and those without it 'non-518 chloroplast'.

519

520 The frequency of three major motifs appeared to change in response to dark and in 521 recovery and between genotypes (Fig.S2A). The G-box element (CACGTC) is a 522 focal point of light-regulated gene expression. In vitro gel-shift, random DNA-binding 523 selection, and chromatin immunoprecipitation (ChIP) assays in *A.thaliana* show that 524 four PIFs (PIF1, PIF3, PIF4, and PIF5) bound to either a G-box (CACGTG) and/or an 525 E box (CANNTG) (Martínez-García et al., 2000; Hug and Quail, 2002; Hug et al., 526 2004; Hornitschek et al., 2012). PIFs can also interact with other transcription factors 527 at the G-box, and these interactions modulate the PIF DNA-binding activity. PIF3 528 and PIF4 interact with BRASSINAZOLE-RESISTANT 1 (BZR1) and bind to the same 529 G-box DNA sequence element to regulate genes involved in the light and 530 brassinosteroid pathways (Oh et al., 2012; Zhang et al., 2013). PIF1 and PIF3 also 531 interact with the light-regulated activator ELONGATED HYPOCOTYL (HY5) at the 532 G-box where it can both promote PIF1/3 binding and compete for binding sites 533 (Chen et al., 2013; Toledo-Ortiz et al., 2014). When B.rapa leaves were returned to 534 light, 37% of chloroplast genes significantly upregulated in wild type but not BrphyB 535 leaves have a G-box in their promoter region compared to only 15% of non-536 chloroplast genes. This is not the case with chloroplast genes upregulated only in 537 BrphyB. For these genes, there was essentially no difference in the number of genes

538 a G-box whether or not they were annotated as chloroplast-associated (chloroplast 539 genes: 13%, non-chloroplast genes: 16%) (Fig.S2B).

540

541 Differences were also present in the frequencies of Evening Element (AAAATATCT) 542 and Telobox motif (AAACCCTAA) in chloroplast-annotated genes between wild-type 543 and BrphyB leaves in recovery as well. The Evening Element (EE) motif is central to 544 circadian clock function and environmental and endogenous signal coordination in 545 A.thaliana. Key regulators of the circadian clock CIRCADIAN CLOCK 546 ASSOCIATED1 (CCA1), LATE ELONGATED HYPOCOTYL (LHY) and REVEILLE 8 547 (RVE8) bind and regulate genes with EEs in their promoters (Harmer and Kay, 2005; 548 Hsu et al., 2013). Among chloroplast-annotated genes, the EE motif was present in 549 promoter regions of 37% of genes which were significantly downregulated in BrphyB 550 but not wild type leaves, while only 7% of those downregulated in wild type but not 551 *BrphyB* had the same motif (Fig.S2C).

552

553 Short interstitial telomere motifs (telo boxes) are short sequences identical to plant 554 telomere repeat units. In A.thaliana and O.sativa genomes telo boxes are associated 555 with genes involved in the biogenesis of the translational apparatus (Gaspin et al., 556 2010). Telo box motifs were enriched in the promoters of genes significantly 557 downregulated in wild type but not BrphyB, and genes significantly upregulated in 558 *BrphyB* but not wild type in recovery, 15% compared to 20%, respectively. Whereas, 559 of genes that were upregulated in wild type only, and genes downregulated in 560 BrphyB only in recovery, 5% and 7% respectively had Telo-box motifs in their 561 promoters. The differences between genes with and without the chloroplast GO 562 annotation was less noticeable for this motif than the other two. Together these 563 results point to a significant difference in the cis-regulatory landscape of BrphyB 564 leaves (Fig. S2D).

565

The chloroplast carries out many functions beyond photosynthetic carbon fixation
that are essential for metabolic homeostasis, including fatty acid synthesis and
fixation of nitrogen and sulfur (Lopez-Juez and Pyke, 2005). Mutants with reduced
phy function have significantly lower chlorophyll levels in *A.thaliana* and *B.rapa*(Ghassemian *et al.*, 2006; Strasser *et al.*, 2010; Hu *et al.*, 2013; Arsovski *et al.*,
2018). It has been suggested that phyA is primarily responsible for chloroplast

572 maturation during de etiolation in *A.thaliana*, although there are some reports that 573 phyB might also be involved (McCormac and Terry, 2002; Xu *et al.*, 2019).

574

575 Our results, in combination with our earlier findings that *BrphyB* mutants had 576 reduced chlorophyll levels and photosynthetic rates, led us to hypothesize that 577 BrphyB might be required for normal chloroplast development. We found that 578 chloroplast density was significantly decreased in the mature leaves of the BrphyB mutants. Wild-type leaves had an average of 466 chloroplasts per 0.5mm² compared 579 580 to 326 and 253 in BrphyB-1 and BrphyB-3, respectively (Fig. 4A, B ANOVA and 581 Tukey HSD multiple comparison test). Chloroplast area however was significantly 582 larger in BrphyB-3 and BrphyB-1 compared to wild type, 33.6 and 41.4 to 31.3 um² 583 respectively (Fig. 4C ANOVA and Tukey HSD multiple comparison test). In 584 A.thaliana, an investigation into photosynthetic, biochemical, and anatomical traits of 585 accumulation and replication of chloroplasts (arc) mutants found that fewer, enlarged 586 chloroplasts were less efficient at photosynthesis than more, smaller chloroplasts. 587 Photosynthetic rate and photosynthetic nitrogen use efficiency were significantly 588 lower in the mutants than their wild-types likely due to decreases in mesophyll 589 conductance and chloroplast CO₂ concentration (Xiong et al., 2017). These 590 functional differences could explain the reduced ability of *BrphyB* leaves to rapidly 591 switch metabolic functions when exposed to darkness and again with the return to

592

593

594 **DISCUSSION**

light.

595

596 Human-driven climate change, and the associated changes in temperature, 597 atmospheric CO₂, and precipitation, are an urgent challenge for plant life on Earth. 598 Crop yield and global food security will depend on how individual crop species 599 respond to new and potentially more variable conditions. *B.rapa* is a laboratory crop 600 model that has been successfully used to study the plant response to environmental 601 change. phyB is emerging as a key regulator of carbon response and supply, 602 metabolism and biomass production (Yang et al., 2016; Arsovski et al., 2018). In 603 addition to a diminished high CO₂ response we previously showed that *B.rapa phyB* 604 mutants have reduced chlorophyll levels and photosynthetic rate (Arsovski et al., 605 2018). In this work, we elicited a dark response in individual leaves and examined

the transcriptomic response of wild-type and *phyB* leaves as they are darkened andwere subsequently returned to light.

608

609 Wild-type *B.rapa* leaves darkened for three days have a significant upregulation of 610 genes involved in autophagy, catabolism and leaf senescence while large groups of 611 genes functioning in photosynthesis, metabolism and translation (Fig. 2A). This 612 senescence response and redistribution of cell resources is typical of the dark 613 response observed in various plant systems (Guo *et al.*, 2004; Brouwer *et al.*, 2012; 614 Song et al., 2014; Law et al., 2018; Sobieszczuk-Nowicka et al., 2018). Upon return 615 to light upregulated and downregulated functional groups are essentially reversed. 616 Genes acting mainly in response to light stimulus, photosynthesis, translation and 617 metabolism were upregulated in leaves, while those with roles in catabolism, vesicle 618 fusion and transport, and protein degradation were downregulated 24 hours following 619 return to light (Fig. 3B). In many plants, leaf senescence is reversible in a limited 620 time span after senescence initiation, leading to 'regreening' of the leaves. For 621 example, a return to light following a 2 day dark period initiates a reconstitution of 622 photosynthetic capability in A. thaliana (Parlitz et al., 2011).

623

624 While there is a significant overlap in the transcriptomic response to dark and 625 recovery between wild-type and BrphyB mutant leaves, there are important 626 categories of genes that may explain some of phenotypes associated with the 627 BrphyB mutant (Fig. 2,3). After three days of dark the orthologs of A.thaliana 628 PHYTOCHROME INTERACTING FACTORS 4 and 5 are only upregulated in the 629 *BrphyB* mutant suggesting a misregulation of the light response, as well as likely 630 impacts on hormone homeostasis. In A. thaliana PIFs/EIN3/HY5-regulated genes in 631 the dark were estimated to account for half of the light-directed transcriptome 632 changes (Shi et al., 2018). In recovery, BrphyB mutant leaves lack the increased 633 transcription of genes involved in light harvesting and photosynthesis, including 125 634 chloroplast-targeted genes, and the mutants uniquely down-regulate genes involved 635 in phototropism suggesting possible "crossed-wires" from mismatches between 636 different photoreceptor responses (Fig. 3A). Together these finding may connect the 637 observed reduction of chlorophyll and photosynthetic rate (Arsovski et al., 2018) and 638 fewer larger chloroplasts in *BrphyB* leaves compared to wild type. 639

640 Nitrogen is a critical resource that governs plant growth. The availability of nitrogen 641 to the roots plays a particularly significant role in constraining plant growth and crop 642 yield worldwide (Epstein and Bloom, 2005; Hirel et al., 2011; Alvarez et al., 2012). 643 Nitrogen deficiency is one of the endogenous and environmental factors that 644 regulates the onset of leaf senescence (Gregory, 1937; Mei and Thimann, 1984; 645 Masclaux-Daubresse et al., 2007; Koeslin-Findeklee et al., 2015). In B.rapa, nitrogen 646 availability limits growth increase in high CO₂ (Arsovski et al., 2018). In dark-induced 647 leaf senescence nitrogen from senescing leaves is mobilized and transported to still 648 growing vegetative tissues. Wild type *B.rapa* plants darkened for 3 days show a 649 significant upregulation of genes involved in senescence, catabolism, and vesicle 650 transport while downregulation of genes involved in protein synthesis indicating and 651 active export of nitrogen resources (Supplementary File 1, 2). While BrphyB mutants 652 largely share this response, there is evidence that this resource allocation is altered. 653 After 3 days of dark phyB unique downregulated genes are significantly enriched for 654 GO terms related to protein synthesis suggesting either a delayed or prolonged 655 response compared to wild type. In recovery, BrphyB uniquely upregulated genes 656 are significantly enriched for translation and peptide biosynthesis, while wild type 657 unique genes are photosynthesis-related. B.napus, a close relative of B.rapa has 658 poor nitrogen use efficiency (Masclaux-Daubresse et al., 2007; Xu et al., 2012). Only 659 50-60% of the applied nitrogen is recovered in the plants and at the time of harvest 660 a relatively low 80% of the total plant nitrogen is localized in the seeds (Schjoerring 661 et al., 1995; Jensen et al., 1997; Malagoli et al., 2005; Rathke et al., 2006). Here 662 BrphyB mutants had a misregulation of genes orthologous to A.thaliana 663 GLUTAMINE SYNTHASE 1;2, 1;3 and 1;4 that play a role in seed yield and size 664 (Fig. 3C). While seeds at harvest were not significantly different from wild type in size 665 and weight, seed yield is dramatically reduced in the mutant (Arsovski et al. 2018). A 666 more detailed understanding of the phyB-regulated network holds the promise of 667 improved plant growth models and identification of new targets for engineering more 668 resource-efficient crops. 669

- 670 Supplemental Figures
- 671
- 672 Figure S1: qPCR validation.
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674 Figure S2: Transcription factor motifs enriched in promoter regions. 675 676 Table S1: Gene expression comparison between BrphyB and wild type leaves in the 677 Pre condition. 678 Table S2: Gene Ontology comparison of differentially expressed genes between Pre 679 680 (P), Dark (D), and Recovery (R) in wild type and *BrphyB* leaves. 681 682 Table S3: A comparison of *A.thaliana* gene differentially expressed in whole 683 darkened plants or individually darkened leaves in response to 3 days of darkness 684 from Law et al., 2018 and A.thaliana orthologs of B.rapa genes differentially 685 expressed in Dark vs. Pre. 686 687 Table S4: Gene Ontology analysis of *A.thaliana* genes differentially expressed in 688 individually darkened leaves (IDL), or whole darkened plants (DP) from Law et al. 689 2018 and A.thaliana orthologs of B.rapa genes differentially expressed in Dark vs. Pre. Sheets show A.thaliana IDs, gene model names, MapMan bins and descriptions 690 691 of common up or down regulated genes followed by GO annotations of those genes. 692 693 Table S5: A comparison of genes differentially expressed in dark and re-illumination 694 from Parlitz et al. 2018 and A.thaliana orthologs of B.rapa genes in Dark vs. Pre and 695 Recovery vs. Dark. 696 697 Table S6: Gene Ontology enrichment for DEGs common and unique to wild type and 698 BrphyB in pre (P), dark (D), and recovery (R) conditions. 699 700 Table S7: Differentially expressed genes in pre, dark and recovery. Significance 701 column denotes whether the significance is common to both wild type and *BrphyB* or 702 unique to either. 703 704 Acknowledgements 705 We thank Prof. Mark Stitt and Dr. Virginie Mengin for sharing their insights and 706 expertise, as well as the other members of the PHYTOCAL consortium, Nemhauser 707 and Imaizumi labs for feedback and discussion. We also thank the undergraduate 708 researchers from Dr Arsovski's Spring 2018 Special Field Topics class who carried 709 out protocol optimization and preliminary measurements of phenotypes presented 710 here: Ericka Budinich, Jonas Hill, Sean Hoeger, Katrin Hosseini, Nikhil Kaza, Winnie 711 Kwong, Kellen Larsen, Andrew Lui, Rohan Menon, Claudia Moroney, Anita Nguyen, 712 Arthur Sargent, Emma Stevens, and Nanami Tsumura. This work was supported by 713 the National Science Foundation participation in the ERA-CAPS program (IOS-

714 1539834).

716 Figure 1: RNAseq experimental set-up (A). At 3 weeks of age the "pre" sample 717 was collected from the first of the two developmentally matched leaves. Symmetrical 718 samples from the same leaf was collected for chlorophyll measurement. The second 719 matched leaf was covered with tinfoil at this time. 24 hours later the "24" sample was 720 harvested from the uncovered leaf, the same leaf that provided the "pre" sample. 48 721 hours later the tinfoil was removed from the covered leaf and the "dark" samples 722 were similarly collected. 24 hours later the "recovery" samples were collected from 723 this same leaf. Samples were immediately frozen in liquid nitrogen. Three biological 724 replicated were similarly collected. Total chlorophyll in Pre, dark, and recovery 725 samples, error bars are SE.

726

727 Figure 2: Genes differentially expressed in dark. A Gene Ontology (GO) analysis

of genes uniquely differentially expressed in wild type and *BrphyB* mutant leaves

following 72 hours of dark. A) Upregulated genes. B) Downregulated genes. C)

730 Expression values of 3 biological replicates in exemplar genes in Pre(P), Dark (D)731 and Recovery (R).

731 a 732

733 Figure 3: Genes differentially expressed on return to light. A Gene Ontology

(GO) analysis of genes uniquely differentially expressed in wild type and *BrphyB*

mutant leaves 24 hours after return to light. A) Upregulated genes. B)

736 Downregulated genes. C) Expression values of 3 biological replicates in exemplar

737 genes in Pre(P), Dark (D) amd Recovery (R).

738

739 Figure 4 : BrphyB mutant plants have fewer and larger chloroplasts. A.

740 Fluorescent images of chloroplasts in 3 week old *B.rapa* leaves. B. Chloroplast

741 density in same leaves as A. Chloroplast area of individual chloroplasts in same

742 plants as A. Lower case letters in B and C indicate significant difference (ANOVA

and Tukey HSD multiple comparison test; p<0.001)

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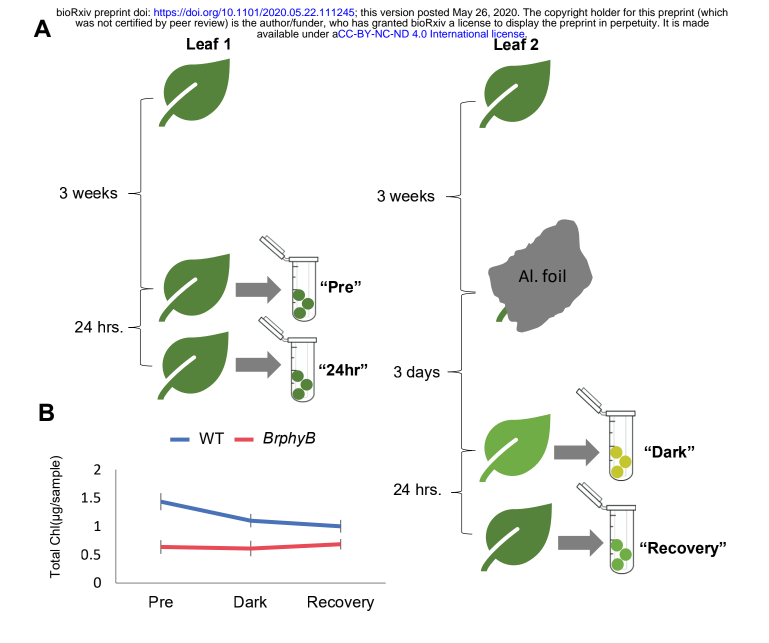


Figure 1: RNAseq experimental set-up (A). At 3 weeks of age the "pre" sample was collected from the first of the two developmentally matched leaves. Symmetrical samples from the same leaf was collected for chlorophyll measurement. The second matched leaf was covered with tinfoil at this time. 24 hours later the "24" sample was harvested from the uncovered leaf, the same leaf that provided the "pre" sample. 48 hours later the tinfoil was removed from the covered leaf and the "dark" samples were similarly collected. 24 hours later the "recovery" samples were collected from this same leaf. Samples were immediately frozen in liquid nitrogen. Three biological replicated were similarly collected. Total chlorophyll in Pre, dark, and recovery samples, error bars are SE.



Α



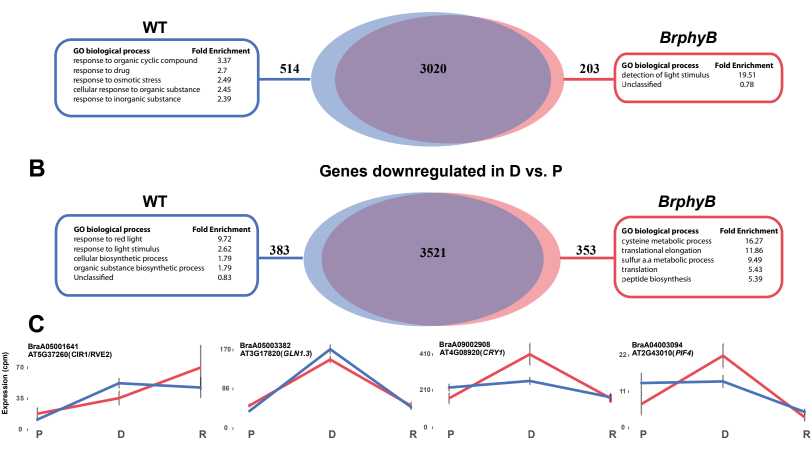


Figure 2: Genes differentially expressed in dark. A Gene Ontology (GO) analysis of genes uniquelly differentially expressed in wild type and BrphyB mutant leaves following 72 hours of dark. A) Upregulated genes. B) Downregulated genes. C) Expression values of 3 biological replicates in exemplar genes in Pre(P), Dark (D) and Recovery (R).

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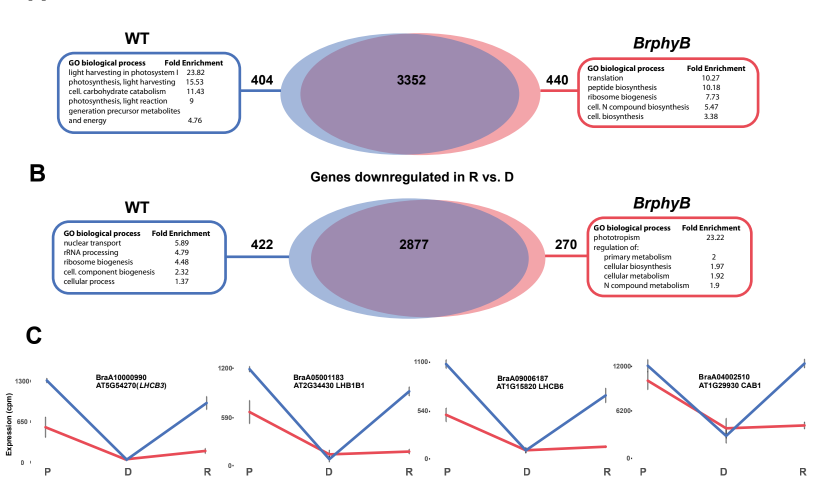


Figure 3: Genes differentially expressed on return to light. A Gene Ontology (GO) analysis of genes uniquelly differentially expressed in wild type and BrphyB mutant leaves 24 hours after return to light. A) Upregulated genes. B) Downregulated genes. C) Expression values of 3 biological replicates in exemplar genes in Pre(P), Dark (D) and Recovery (R).

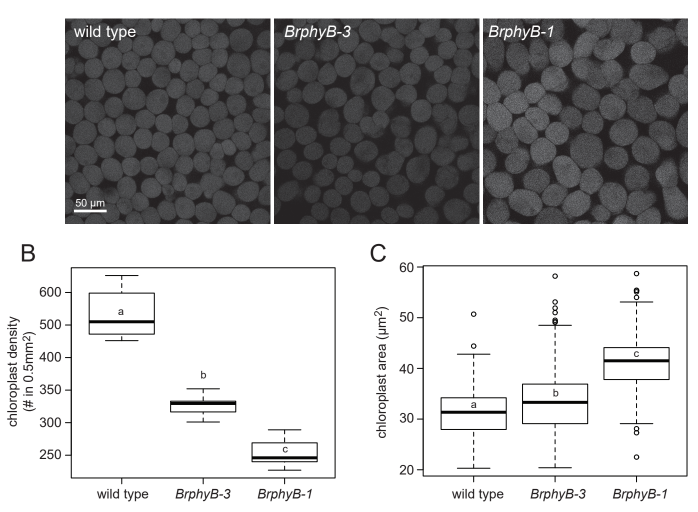


Figure 4 : *BrphyB* mutant plants have fewer and larger chloroplasts. A. Fluorescent images of chloroplasts in 3 week old *B.rapa* leaves. B. Chloroplast density in same leaves as A. Chloroplast area of individual chloroplasts in same plants as A. Lower case letters in B and C indicate significant difference (ANOVA and Tukey HSD multiple comparison test; p<0.001)