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Expression of the UVR8 photoreceptor in different tissues reveals tissue autonomous features of UV-B signalling

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	1	Expression	of the U	VR8 photo	receptor in	different	tissues	reveals	tissue-
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- 2 autonomous features of UV-B signalling
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- 6
- 7
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- 19
- 20 **Running Title**:
- 21 UVR8 signalling in different tissues

22 SUMMARY STATEMENT

23	This work analyses how the UV-B specific photoreceptor UVR8 regulates signalling,
24	development and growth when expressed only in specific tissues. We show that early
25	steps of UVR8-dependent signalling, such as accumulation of the key regulatory
26	transcription factor HY5, occur strictly in tissue-autonomous fashion. In contrast,
27	complex UV-B-induced changes, including proper acclimation of adult plants requires
28	simultaneous signalling in the epidermal and mesophyll cells and/or inter-tissue
29	signalling.
30	
31	
32	ABSTRACT (162 words)
33	
34	The Arabidopsis UV-B photoreceptor UV RESISTANCE LOCUS 8 (UVR8)
35	orchestrates the expression of hundreds of genes, many of which can be associated
36	with UV-B tolerance. UV-B does not efficiently penetrate into tissues, yet UV-B
37	regulates complex growth and developmental responses. To unravel to what extent
38	and how UVR8 located in different tissues contributes to UV-B-induced responses,
39	we expressed UVR8 fused to the YELLOW FLUORESCENT PROTEIN (YFP)
40	under the control of tissue-specific promoters in a <i>uvr8</i> null mutant background. We
41	show that (i) UVR8 localized in the epidermis plays a major role in regulating
42	cotyledon expansion, and (ii) expression of UVR8 in the mesophyll is important to
43	protect adult plants from the damaging effects of UV-B. We found that UV-B induces
44	transcription of selected genes, including the key transcriptional regulator
45	ELONGATED HYPOCOTYL 5 (HY5), only in tissues that express UVR8. Thus we

- 46 suggest that tissue-autonomous and simultaneous UVR8 signalling in different tissues
- 47 mediates, at least partly, developmental and defence responses to UV-B.

- 49
- 50 Key words:
- 51 Arabidopsis, ultraviolet-B, UVR8, signalling, tissue specificity

52 INTRODUCTION

53	Plants must adapt to the environment to optimize growth and development for
54	survival and successful reproduction. Light is an essential environmental factor and
55	necessary not only for photosynthesis but also as a signal for proper development and
56	growth. Plants evolved various photoreceptors that are able to monitor changes in the
57	quantity and quality of the ambient light environment. These include the blue/UV-A
58	light absorbing phototropins, cryptochromes and Zeitlupe family receptors; the
59	red/far-red absorbing phytochromes (phyA-phyE), as well as the UV-B photoreceptor
60	UV RESISTANCE LOCUS 8 (UVR8) (Galvao & Fankhauser, 2015).
61	UV-B radiation (280-315 nm) is an integral part of sunlight reaching the
62	Earth's surface, as it is only partially absorbed by the stratospheric ozone layer. UV-B
63	can damage several macromolecules (DNA, proteins etc.) (Hollosy, 2002). However,
64	UV-B also activates UVR8-dependent signal transduction pathways and triggers
65	responses that manifest as inhibition of hypocotyl elongation, reduction of leaf size,
66	entrainment of the circadian clock, modification of shade avoidance response,
67	alteration of phototropism, increased accumulation of photo-protective flavonoids and
68	increased survival under UV-B stress (Kliebenstein et al., 2002; Brown et al., 2005;
69	Favory et al., 2009; Feher et al., 2011; Morales et al., 2013; Hayes et al., 2014;
70	Jenkins, 2014; Vandenbussche et al., 2014).
71	At the cellular level the UVR8 photoreceptor can be detected both in the
72	cytoplasm and the nucleus in visible light, but irradiation with UV-B increases
73	accumulation of UVR8 in the nucleus (Kaiserli & Jenkins, 2007; Yin et al., 2016).
74	Nuclear localization of UVR8 is required but not sufficient for UV-B signalling
75	(Brown et al., 2005; Kaiserli & Jenkins, 2007; Yin et al., 2016). It is a matter of
76	debate whether or not UVR8 directly associates with chromatin to regulate UV-B-

	77	dependent transcription of target genes, including HY5 (Cloix & Jenkins, 2008;
	78	Binkert et al., 2016). The ELONGATED HYPOCOTYL 5 (HY5) transcription factor
	79	is a major positive regulator of photomorphogenesis both in visible (Lee et al., 2007)
	80	and UV-B light (Ulm et al., 2004; Brown et al., 2005; Oravecz et al., 2006; Binkert et
	81	al., 2014). hy5 mutants are largely impaired in UV-B-responsive gene expression and
	82	the accumulation of UV-B-protective flavonoid pigments, leading to reduced UV-B
	83	tolerance and survival (Brown et al., 2005; Oravecz et al., 2006; Stracke et al., 2010).
	84	UV-B irradiation was shown to rapidly induce HY5 gene expression (Ulm et al., 2004;
	85	Brown & Jenkins, 2008; Binkert et al., 2014; Binkert et al., 2016) and the
	86	accumulation of HY5 protein in the nucleus (Oravecz et al., 2006).
	87	UV-B penetrates rather poorly into tissues below the epidermis. Indeed, leaf
	88	epidermal transmittance of UV-B is less than 10 %, measured in many different
	89	species under various circumstances (Robberecht et al., 1980; Day et al., 1993;
	90	Markstadter et al., 2001; Qi et al., 2003; Nybakken et al., 2004). UVR8 is expressed
	91	ubiquitously in different organs of mature Arabidopsis (Rizzini et al., 2011), but the
	92	precise distribution pattern and the accumulation level of the photoreceptor in various
	93	tissues have not yet been investigated. It follows that it is not understood how the
	94	action of UVR8 in different tissues/organs is integrated to regulate complex
	95	physiological responses as hypocotyl growth inhibition or leaf size, and how the
	96	strongly varying UV-B intensities in different tissues modulate UVR8-dependent
	97	signalling.
	98	Here we characterized the spatio-temporal aspects of UV-B-induced, UVR8-
	99	mediated signalling to provide insight into the molecular mechanism mediating signal
1	00	integration between different tissues/organs. We first determined the distribution
1	01	pattern and level of YFP-UVR8 under the control of its own promoter. Next we

characterized to what extent UV-B-induced physiological and molecular responses are
mediated by tissue-autonomous and/or inter-tissue signalling in transgenic lines that
expressed the photoreceptor in a tissue-specific fashion. Our data suggest that UVR8
responses are mediated partly by tissue-autonomous signalling, but proper regulation
of hypocotyl growth inhibition and establishment of UV-B tolerance require either
UVR8 action in different tissues and/or inter-tissue signalling.

109 MATERIALS AND METHODS

110 Molecular cloning

- 111 The coding region of *YFP* and *UVR8* were cloned into the pPCV812 plasmid
- 112 (Bauer et al., 2004) as SmaI-EcoRI and EcoRI-SacI fragments, respectively. The
- 113 MERISTEM LAYER 1 (ProML1), SUCROSE /H⁺ SYMPORTER 2 (ProSUC2) and
- 114 CHLOROPHYLL A/B BINDING PROTEIN 3 (ProCAB3) promoter fragments were
- 115 cloned as described by Kirchenbauer et al. (2016) whereas the ProUVR8 was inserted
- as a 2569 bp *SalI-Bam*HI fragment including the 5` leader sequence. The coding
- 117 sequence of the β -glucuronidase (GUS) as a SmaI-XhoI fragment (Adam et al., 1995),
- 118 *GFP* as a *XhoI-ClaI* fragment and *NLS* as a *ClaI-SacI* fragment (Wolf *et al.*, 2011)
- 119 were cloned into the *pPCVB812* binary vector (Bauer *et al.*, 2004) resulting in *GUS*-
- 120 GFP-NLS pPCVB. This vector was digested with HindIII and SmaI restriction
- 121 enzymes and the ProHY5 (Oravecz et al., 2006) was inserted as a HindIII-StuI
- 122 fragment replacing the *Pro35S* promoter. *ProELIP2* and *ProPRR9* were cloned as
- 123 2772 bp (*Bam*HI-*Xba*I) and 1324 bp (*Bam*HI-*Sma*I) fragments including the 5` leader
- 124 sequences, respectively. Cloning of *ProHY5:HY5-GFP* was described in detail by
- 125 Kirchenbauer *et al.* (2016).

127 Plant material

128	Throughout the study we used the Arabidopsis thaliana L (Heynh.) uvr8-6 null
129	mutant (Favory et al., 2009), with the Columbia accession as wild type (WT) control.
130	We raised 10 independent transgenic lines per construct and selected those which
131	segregated the transgene as a single Mendelian trait. At least 3 independent lines were
132	studied and comparable results are presented. Arabidopsis transformation, principles
133	of selection and handling of transgenic lines were described earlier in detail
134	(Kirchenbauer et al., 2016).
135	
136	Seedling growth conditions and light treatments
137	Seeds were surface sterilized and subsequently stratified for 72 h in the dark (4
138	°C) on 1/2 Murashige and Skoog (MS) medium (Sigma-Aldrich, Budapest, Hungary)
139	containing 1% sucrose and 0.8% agar. For microscopic analysis, the seedlings were
140	grown in 12 h white light (WL, 80 $\mu mol~m^{-2}~s^{-1})/12$ h dark at 22 °C for 6 days (MLR-
141	350, Sanyo, Gallenkamp, UK) and then placed under continuous white light
142	supplemented with UV-B for 16 h at 22 °C. White light was produced by PHILIPS
143	TL-D 18W/33-640 tubes (10 μ mol m ⁻² s ⁻¹). Non-damaging photomorphogenic (low-
144	fluence) UV-B was produced by PHILIPS ULTRAVIOLET-B TL20W/01RS tubes
145	(1.5 μ mol m ⁻² s ⁻¹). To modulate UV-B light we used 3-mm thick transmission cut-off
146	filters of the WG series (Schott, Mainz, Germany), as described previously (Ulm et
147	al., 2004). UV-B treated seedlings (+UV-B) were covered with WG305 filter with
148	half-maximal transmission at 305 nm, whereas non-UV-B irradiated control seedlings
149	were covered with WG385 filter with half-maximal transmission at 385 nm (-UV-B)
150	as applied in work published earlier (Oravecz et al., 2006; Favory et al., 2009; Rizzini
151	et al., 2011). UV-B was measured with a VLX-3W UV light meter equipped with a

158	Microscopy techniques
157	
156	days.
155	before being exposed to continuous WL supplemented with UV-B for 4 days or 5
154	cotyledon measurements, seedlings were grown for 3 days in light/dark chambers
153	measured with an LI-250 Light Meter (Li-Cor, Lincoln, NE, USA). For hypocotyl and
152	CX-312 sensor (Vilber Lourmat, Eberhardzell, Germany) and the visible part was

159 Confocal laser scanning microscopy (CLSM) settings and quantification of

160 nuclear fluorescence were described in detail by Kirchenbauer *et al.* (2016)

161

162 Flavonoid detection using confocal laser scanning microscopy

163 Seeds were stratified and germinated as described above. Seedlings were 164 grown for 2 days in 12 h light/12 h dark chambers and were placed under 1.5 µmol $m^{-2}~s^{-1}$ WL supplemented with 1.5 $\mu mol~m^{-2}~s^{-1}$ UV-B light for 4 days. Seedlings 165 166 treated with UV-B were covered with a WG305 filter, whereas the negative controls 167 (-UV-B) were covered with WG385. Prior to microscopic analysis seedlings were 168 incubated in 0.1% (w/v) Naturstoffreagenz A (DPBA, Sigma-Aldrich) in 0.15 M 169 phosphate buffer (pH 6.8) in the dark. After 15 min incubation time DPBA was 170 removed by exchanging the buffer for fresh phosphate buffer twice. CLSM was used 171 to detect DPBA-flavonoid specific fluorescence (488 nm laser; pinhole: 200 µm; 172 spectral emission detector: 501-601 nm). 173 174 Hypocotyl length and cotyledon area measurements 175 Measurements of hypocotyl length and cotyledon area were performed as

176 described earlier (Adam et al., 2013). At least 40 seedlings (hypocotyl length) or 100

177 cotyledons were measured for each line and each treatment. Ratios of UV-B

178 treated/non-treated hypocotyl lengths and cotyledon areas were calculated in each

179 experiment. Experiments were repeated at least three times. The calculated ratio

180 values were averaged and the standard error values of the means were obtained and

181 plotted.

182

183 **Protein isolation and western blot**

184 Preparation of plant protein extracts and western blotting were described by

185 Bauer *et al.* (2004). Application of anti-UVR8, anti-ACTIN antibodies and signal

186 processing were also described earlier (Heijde & Ulm, 2013; Medzihradszky et al.,

187 2013). All protein extraction and western blotting was repeated three times and a

188 representative image is presented. Signal quantification was made using Image J

189 software (NIH).

190

Determination of transcript levels

192 Total RNA isolation, cDNA synthesis and quantitative RT-PCR analysis were

193 performed as described by Feher *et al.* (2011).

194

195 **Propagation and UV-B treatment of adult plants for phenotype analysis and**

196 chlorophyll determination

197 Arabidopsis seeds were sown on soil, stratified for three days at 4 °C and then grown

198 in a climate-controlled growth chamber (Grobank, CLF Plant Climatics, Wertingen,

199 Germany) in short days conditions (8 h light / 16 h dark) under WL (120 μ mol m⁻²

 200 s^{-1}) or WL supplemented with UV-B at 22°C. The visible part was measured with an

201 LI-250 Light Meter (Li-Cor). The light conditions in the chambers were set following

202	the general guidelines described by Aphalo et al. (2012) and the full spectra of the
203	applied light was analysed with a QE65000 spectrometer (Ocean Optics, Dunedin,
204	FL, USA) (Figure S1). We used white fluorescent tubes (Osram L18W) and the same
205	type narrowband UV-B tubes, what were used in the seedling irradiation treatments
206	(TL20W/01RS, Philips) without plastic filtering. The applied UV-B fluence rates (2
207	or 12 μ mol m ⁻² s ⁻¹) were comparable to the natural values measured in Szeged,
208	Hungary on an average sunny summer day (7-15 μ mol m ⁻² s ⁻¹ between 11:00-13:00
209	CET on 09.06.2010.). UV-B was measured with a VLX-3W UV light meter equipped
210	with a CX-312 sensor (Vilber Lourmat). Rosette diameter was quantified in images of
211	7-week-old plants using ImageJ. Three repetitions of each experiment were performed
212	using two independent lines for the tissue-specific lines. At least four plants were
213	measured in each repetition for each genotype and independent line. Determination of
214	chlorophyll levels were described earlier (Porra et al., 1989)

- 215
- 216
- 217 **RESULTS**
- 218 Expression of the *ProUVR8:YFP-UVR8* transgene is restricted to epidermal and
- 219 mesophyll cells

To address where UVR8 is expressed, we generated transgenic lines expressing the *YFP-UVR8* fusion protein under the control of its own promoter in a *uvr8* null mutant
background and determined its expression pattern by using CLSM. We found that the *UVR8* promoter drives the expression of YFP-UVR8 in the epidermal and, to a lesser
extent, the mesophyll/subepidermal cells of cotyledons and hypocotyls (Figures 1A-C,
S2-S4). Accumulation of the YFP-UVR8 fusion protein was below detection level in
the vascular bundles. But it should be noted that the YFP-UVR8 amount corresponded

227 to $\sim 10\%$ of the native UVR8 protein detected in WT seedlings (Figure 2A) and that 228 we did not identify any *ProUVR8: YFP-UVR8* line with higher YFP-UVR8 protein 229 amounts.

230

231 Characterization of transgenic lines expressing YFP-UVR8 in selected tissues 232 To assess the function of UVR8 located in different tissues we expressed YFP-UVR8 233 in the uvr8 mutant background under the control of ProML1, ProSUC2 and ProCAB3 234 promoters that have already been used in numerous studies to express proteins of 235 interest in epidermal, companion and mesophyll cells, respectively (Mitra *et al.*, 1989; 236 Sessions et al., 1999; Srivastava et al., 2008; Kirchenbauer et al., 2016). Figure 1D-I 237 and Figures S2-S4 demonstrate that the *ProML1* drives the expression of YFP-UVR8 238 selectively in epidermal cells, whereas *ProCAB3* in the sub-epidermal (mesophyll) 239 cells of cotyledons and hypocotyls. As expected, no activity of these promoters was 240 detected in the vascular bundles. By contrast, *ProSUC2* expressed YFP-UVR8 in the 241 vasculature and sub-epidermal cells of cotyledons and hypocotyls (Figure 1J-L and 242 S2-S4). Western blot analysis showed that the total amount of YFP-UVR8 in 243 ProML1:YFP-UVR8 was ~5%, in ProSUC2:YFP-UVR8 ~25% and in ProCAB3:YFP-244 UVR8 75% of the amount of endogenous UVR8 in WT seedlings (Figure 2A). To 245 facilitate direct comparison of the level of YFP-UVR8 in different cell types, we 246 monitored its accumulation by CLSM. The amount of YFP-UVR8 was (i) comparable 247 in the epidermal cells of ProUVR8: YFP-UVR8 and ProML1: YFP-UVR8, (ii) about 4-248 5-fold lower in the mesophyll cells of *ProUVR8:YFP-UVR8* as compared to 249 *ProCAB3:YFP-UVR8* and about the same in *ProSUC2:YFP-UVR8* (Figure S5). It was 250

not feasible to compare its accumulation by this method in the vascular bundles.

251

252 Complementation of seedling phenotypes of the *uvr8-6* mutant by tissue-specific

253 expression of YFP-UVR8

254 To assess the function of UVR8 in different tissues, we measured typical 255 photomorphogenic responses such as inhibition of hypocotyl elongation and 256 expansion of cotyledons, of the various transgenic seedlings exposed to UV-B 257 irradiation. Figure 2B shows that supplemental narrowband UV-B inhibited hypocotyl 258 growth in the wild-type seedlings, whereas the *uvr8* mutant seedlings were much less 259 responsive, in agreement with previous results (Favory et al., 2009). All transgenic 260 seedlings, except ProSUC2:YFP-UVR8, showed pronounced UV-B-induced 261 hypocotyl growth inhibition, but did not fully complement the phenotype of the *uvr8* 262 mutant (Figure 2B). We also measured the changes of cotyledon area caused by UV-B 263 irradiation. Figure 2C illustrates that UV-B irradiation decreased the cotyledon size of 264 the uvr8, ProCAB3: YFP-UVR8 and the ProSUC2: YFP-UVR8 seedlings, whereas the 265 same UV-B treatment slightly increased the cotyledon size in the ProUVR8:YFP-266 UVR8, ProML1:YFP-UVR8 and wild-type plants. 267 The above results indicate that (i) the YFP-UVR8 fusion protein is a functional 268 photoreceptor, confirming previous reports (Brown et al., 2005; Kaiserli & Jenkins, 269 2007; Huang et al., 2014; Binkert et al., 2016); (ii) UVR8 signalling contributes to 270 UV-B-induced inhibition of hypocotyl growth both in the epidermal and mesophyll 271 cells; (iii) UVR8 expression in the epidermis is necessary for proper cotyledon 272 expansion under UV-B light; and (iv) YFP-UVR8 expressed in vascular bundles plays 273 a very limited role, if any, in regulating hypocotyl growth and cotyledon expansion 274 (Table S1). 275

276 The UV-B-induced, UVR8-regulated induction of HY5 is tissue-autonomous

277	Increase in the mRNA level and nuclear accumulation of the key UV-B signal
278	transduction component HY5 are among the early steps of the UV-B-induced
279	signalling cascade initiated by UVR8 (Ulm et al., 2004; Brown et al., 2005; Oravecz
280	et al., 2006). To examine the tissue specificity of these responses, we introduced the
281	ProHY5:HY5-GFP (to determine the cell-specific accumulation of HY5 protein) and
282	ProHY5:GUS-GFP-NLS (to determine the cell-specific induction of HY5
283	transcription) reporters into transgenic uvr8 mutant lines expressing YFP-UVR8 in
284	different tissues. Figure 3 demonstrates that (i) the abundance of HY5-GFP was low
285	in seedlings grown in white light, and (ii) UV-B irradiation promoted accumulation of
286	HY5-GFP only in those cells which also contained detectable amounts of YFP-UVR8.
287	Similarly, we found that the UVR8-dependent induction of HY5 transcription is also
288	restricted to YFP-UVR8-containing cells (Figure S6). Thus our results indicate that
289	regulation of the expression of HY5 by UVR8 is a tissue-autonomous response.
290	
• • • •	

291 UV-B induction of the transcription of HY5-dependent and -independent genes is 292 controlled by UVR8 in a tissue-autonomous fashion

293

To get more insight into the tissue-related organization of UVR8 signalling,

294 we also introduced the ProELIP2:GUS-GFP-NLS and ProPRR9:GUS-GFP-NLS

295 transgenes into the ProUVR8: YFP-UVR8, ProML1: YFP-UVR8 and ProCAB3: YFP-

296 UVR8 expressing lines. The EARLY LIGHT-INDUCED PROTEIN 2 (ELIP2) is

297 involved in the photoprotection of thylakoid membranes (Hutin et al., 2003). UV-B

298 irradiation induces accumulation of ELIP2 mRNA (Ulm et al., 2004; Feher et al.,

299 2011), and this response requires functional UVR8 and HY5 (Figure S7) (Oravecz et

300 al., 2006; Favory et al., 2009). Figure 4 demonstrates that the activity of ProELIP2 is

301 low in white light, and that UV-B irradiation strongly enhances its activity only in

those cells which also contain detectable amounts of YFP-UVR8, indicating that the
photoreceptor regulates HY5-dependent expression of *ELIP2* in a tissue-autonomous
fashion.

305	PSEUDO-RESPONSE REGULATOR 9 (PRR9) is a component of the plant
306	circadian clock (Nakamichi et al., 2005). UV-B-induction of ProPRR9 depends on
307	UVR8 (Feher et al., 2011), but it is independent of HY5 (Figure S7). In contrast to the
308	HY5 and ELIP2 promoters, ProPRR9 was active in the sub-epidermal cells of
309	cotyledons in transgenic plants grown in white light. UV-B strongly induced
310	ProPRR9 activity only in those sub-epidermal cells that contained detectable amounts
311	of YFP-UVR8 (Figure S8). Elevated expression of <i>ProPRR9:GUS-GFP-NLS</i> was not
312	detectable in the epidermis of <i>ProML1:YFP-UVR8</i> and <i>ProUVR8:YFP:UVR8</i> lines,
313	although these cells express YFP-UVR8.
314	
315	UVR8-dependent flavonoid accumulation occurs in a tissue-autonomous fashion
315 316	UVR8-dependent flavonoid accumulation occurs in a tissue-autonomous fashion DPBA forms complexes with flavonoid compounds, which can be visualized by
315316317	UVR8-dependent flavonoid accumulation occurs in a tissue-autonomous fashion DPBA forms complexes with flavonoid compounds, which can be visualized by CLSM (Schnitzler <i>et al.</i> , 1996; Hutzler <i>et al.</i> , 1998; Peer <i>et al.</i> , 2001). We applied an
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 315 316 317 318 319 320 	UVR8-dependent flavonoid accumulation occurs in a tissue-autonomous fashion DPBA forms complexes with flavonoid compounds, which can be visualized by CLSM (Schnitzler <i>et al.</i> , 1996; Hutzler <i>et al.</i> , 1998; Peer <i>et al.</i> , 2001). We applied an irradiation protocol which allowed detectable accumulation of flavonoids under supplemental UV-B in wild-type but not in <i>uvr8</i> seedlings. We detected the highest level of UV-B-induced flavonoid accumulation on the inner side of the adaxial
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326 Adult plants require UVR8 in the mesophyll cells for proper acclimation and

327 survival under UV-B

in acclimation to
c-old plants
nts developed equally
rosette growth
reas the uvr8-6
oped light green
photoreceptor-
nutants are
genesis and
enic lines expressing
mparable rosette
6C), except for the
B) and accumulated
at, in contrast to wild
er these conditions
d. Next to wild type,
vived the higher UV-
R8 and
phenotype
the control of
Fasano <i>et al.</i> , 2014).
plants grown on soil
Siunts Brown

as compared with WT (Figure S9). Taken together, these results indicate that the
 expression of UVR8 in subepidermal or epidermal tissues efficiently facilitates

353 acclimation and survival under UV-B.

354

355 **DISCUSSION**

356 Analysis of transgenic ProUVR8: YFP-UVR8 plants revealed the presence of 357 the YFP-UVR8 fusion protein in the epidermal and sub-epidermal cells of cotyledons 358 and hypocotyls of seedlings exposed to UV-B. However, in these lines YFP-UVR8 359 accumulated to levels lower than endogenous UVR8, thus we cannot exclude the 360 presence of low amounts of UVR8 in the vascular tissues of WT seedlings. The fusion 361 protein was biologically active, since the *ProUVR8:YFP-UVR8* transgenic seedlings 362 and adult plants displayed partially or fully complemented UV-B responses. Thus we 363 assume that UVR8 signalling does not play a role in the vasculature, independently of 364 the developmental stage.

365 *ProML1:YFP-UVR8* displayed fully complemented UV-B-induced cotyledon

366 expansion and partially restored hypocotyl growth inhibition, suggesting that

367 epidermal UVR8 is critical for the regulation of these responses. ProCAB3:YFP-

368 UVR8 seedlings containing high levels of YFP-UVR8 in the subepidermal cells also

369 displayed a partially complemented hypocotyl growth inhibition but a non-

370 complemented cotyledon phenotype. The *ProSUC2:YFP-UVR8* line, despite the fact

that it contained a relatively high amount of fusion protein, failed to complement

372 cotyledon growth and displayed only a weak hypocotyl growth inhibition response

373 (Table S1, Figure 2). The latter could be the result of the UVR8 action in mesophyll

374 cells rather than in the vasculature. Based on these data we conclude that at the

375 seedling stage the primary sites of UV-B perception are the epidermis and, to a lesser

376	extent, the mesophyll/sub-epidermal cells. The low penetration of UV-B into deeper
377	layers of plant organs (Day et al., 1993), lends further support to the above
378	conclusion. The distinguished role of epidermis in regulating hypocotyl growth is not
379	unique to UVR8 action, as similar data were reported for phyA (Kirchenbauer et al.,
380	2016) phyB (Endo et al., 2005; Kim et al., 2016) and brassinosteroid signalling
381	(Savaldi-Goldstein et al., 2007). However, both Kirchenbauer et al. (2016). and
382	Savaldi-Goldstein et al. (2007) concluded that exclusive action of phyA or
383	brassinosteroid signalling in the epidermis is not sufficient to recapitulate full
384	regulation of this response. Therefore we assume that the UVR8-mediated inhibition
385	of hypocotyl growth is also mediated partly by the simultaneous action of UVR8 in
386	various tissues and/or inter-tissue signalling.
387	Adult ProCAB3: YFP-UVR8 and ProSUC2: YFP-UVR8 plants having high
388	levels of YFP-UVR8 in the mesophyll displayed an over-expression phenotype,
389	whereas the phenotype of ProUVR8:YFP-UVR8 plants was similar to WT when
390	exposed to strong UV-B (Figure 6). Although subepidermal/mesophyll cells also
391	contain flavonoids (Agati et al., 2011) we do not attribute the over-expression
392	phenotype directly to the accumulation of flavonoids in these cell types. However, we
393	assume that (i) UVR8 in the mesophyll is required for maintaining photosynthetic
394	efficiency under elevated UV-B (Davey et al., 2012) maybe by regulating the levels
395	of the D1 and D2 core proteins, as described recently in Chlamydomonas (Tilbrook et
396	al., 2016), and that (ii) this process needs UVR8 located in cells containing
397	chloroplasts. As for proper rosette development, the phenotypes of the ProML1:YFP-
398	UVR8 and ProUVR8:YFP-UVR8 lines suggest that together with the mesophyll
399	UVR8, the action of epidermal UVR8 is still required. As for the ProSUC2:YFP-
400	UVR8 plant, it remains to be seen whether the activity of UVR8 in the vasculature

401	contributes to the acclimation response, or it is due to <i>ProSUC2</i> promoter action in
402	subepidermal cells. Taken together, we conclude that in mature plants, simultaneous
403	signalling in the epidermal and mesophyll cells and/or inter-tissue signalling is
404	required to optimise growth and development under UV-B.
405	UV-B-induced flavonoid accumulation both in the epidermis (ProML1:YFP-
406	UVR8 or ProUVR8:YFP-UVR8) and mesophyll cells (ProCAB3:YFP-UVR8 or
407	ProSUC2:YFP-UVR8) appears to be regulated by YFP-UVR8 located in the same
408	tissue, i.e. in a tissue-autonomous fashion (Figure 5). However, at present the
409	contribution of inter-tissue signalling or transport of flavonoids (Buer et al., 2007) in
410	regulating their accumulation can not be ruled out.
411	To provide a mechanistic explanation for UV-B-induced developmental
412	responses, we examined the expression patterns of various genes shown to be
413	regulated by UVR8. UV-B-induced transcription and accumulation of the key
414	regulator HY5 was restricted to cells containing UVR8 (Figures 3, S6). The phyA
415	photoreceptor was also shown to regulate HY5 expression in a similar fashion
416	(Kirchenbauer et al., 2016). These data suggest that far-red and UV-B light regulated
417	expression of HY5, probably an early, rate-limiting step of both signal transduction
418	cascades, is mediated in a tissue-autonomous fashion by both photoreceptors.
419	Similarly to HY5, the UV-B-induced expression of <i>ELIP2</i> which requires functional
420	HY5 and that of <i>PRR9</i> whose expression is not regulated by HY5 occurs in a strictly
421	tissue-autonomous way (Figures 4, S8).
422	Taken together, we found no evidence at the molecular level that UVR8-
423	signalling initiates signal crosstalk between different tissues. However, it was reported
424	that UV-B irradiation of certain parts of the plants results in changes of gene
425	expression in shielded organs, indicating that UV-B-induced inter-organ signalling

426	can occur in higher plants (Casati & Walbot, 2004). Therefore we hypothesize that
427	inter-tissue signalling, mediated by yet unknown mobile compounds contributes to the
428	manifestation of UVR8-regulated responses. For example, it was reported that HY5
429	regulates auxin signalling under different light treatments including UV-B irradiation
430	(Cluis et al., 2004; Sibout et al., 2006; Hayes et al., 2014; Vandenbussche et al.,
431	2014). However, to unravel the molecular aspects of UVR8-modulated hormone
432	signalling requires the development of new cellular markers.
433	
434	
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- 634

635 FIGURES



649 whereas red arrows indicate nuclei/cells in the vasculature. Scale bar = $50 \mu m$.



651 Figure 2



- 653 A. Determination of endogenous and YFP-UVR8 protein levels.
- 654 Total protein extract was isolated from 4-day-old seedlings grown under constant WL
- 655 supplemented with UV-B. The proteins were detected using UVR8-specific antibody
- 656 (anti-UVR8). The blot was reprobed with anti-ACTIN antibody as loading control.

657 **B. Effect of UV-B on hypocotyl length.**

- 658 Hypocotyl lengths of seedlings irradiated with constant WL supplemented with (UV-
- B) or without (no-UV-B) UV-B for 4 days were measured and relative hypocotyl
- 660 lengths (UV-B/no-UV-B) were calculated. Each measurement was repeated 3 times;
- 661 error bars represent standard error of the mean. Lines: Col= Columbia wild type; uvr8
- 662 = *uvr8-6* mutant, *ProUVR8*= *ProUVR8*: *YFP-UVR8*; *ProML1*= *ProML1*: *YFP-UVR8*;
- 663 *ProCAB3= ProCAB3: YFP-UVR8; ProSUC2= ProSUC2: YFP-UVR8.* Each transgene
- 664 is expressed in the *uvr8-6* background. Asterisks mark lines that display significant
- 665 differences as compared with the *uvr8* mutant line calculated by the Student's t-test
- 666 (significance: *P < 0.05, ** P < 0.01, ***P < 0.005).

667 C. Effect of UV-B on cotyledon expansion.

- 668 Cotyledon areas of seedlings irradiated with constant WL supplemented with (UV-B)
- or without (no UV-B) UV-B were measured and relative cotyledon areas (UV-B/no
- 670 UV-B) are plotted here. Each measurement was repeated 3 times; error bars represent
- 671 standard error of the mean. Lines: Col= Columbia wild type; uvr8 = uvr8-6 mutant,
- 672 ProUVR8= ProUVR8: YFP-UVR8; ProML1= ProML1: YFP-UVR8; ProCAB3=
- 673 *ProCAB3:YFP-UVR8; ProSUC2= ProSUC2:YFP-UVR8*. Each transgene is expressed
- 674 in the *uvr8-6* background. Asterisks mark lines that display significant differences as
- 675 compared with the *uvr8* mutant line calculated by the Student's t-test (significance:
- 676 *P<0.05, ** P<0.01,***P<0.005).
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Figure 3

684 UV-B induction of *ProHY5:HY5-GFP* in the cotyledon cells of transgenic lines

685 expressing YFP-UVR8 in different tissues.

- 686 *ProHY5:HY5-GFP* was introduced into transgenic *uvr8* lines expressing
- 687 ProUVR8:YFP-UVR8 (UVR8), ProML1:YFP-UVR8 (ML1) or ProCAB3:YFP-UVR8
- 688 (CAB3). Localization of the HY5-GFP fusion protein was monitored by CLSM in the
- epidermis and mesophyll cells of the cotyledon of 7-day-old seedlings irradiated with
- 690 constant WL supplemented with UV-B (+UV-B) or not supplemented (-UV-B).
- 691 Identical microscope settings were used to allow determination of the difference
- 692 between the visual signals of the +UV-B and –UV-B image pairs. White arrows mark
- 693 the positions of selected nuclei in the epidermis; yellow arrows indicate nuclei in the
- 694 mesophyll. Scale bar = $50 \mu m$.





698 UV-B induction of *ProELIP2:GUS-GFP-NLS* in the cotyledon cells of transgenic

699 lines expressing YFP-UVR8 in different tissues.

- 700 *ProELIP2:GUS-GFP-NLS* was introduced into transgenic *uvr8-6* lines expressing
- 701 ProUVR8:YFP-UVR8 (UVR8), ProML1:YFP-UVR8 (ML1) or ProCAB3:YFP-UVR8
- 702 (CAB3). Localization of the GUS-GFP-NLS fusion protein was monitored by CLSM
- in the epidermis and mesophyll cells of the cotyledon of 7-day-old seedlings
- rradiated with constant WL supplemented with UV-B (+UV-B) or not supplemented
- 705 (-UV-B). Identical microscope settings were used to allow determination of the
- difference between the visual signals of the +UV-B and –UV-B image pairs. White
- arrows mark the positions of selected nuclei in the epidermis, yellow arrows indicate
- nuclei in the mesophyll. Scale bar = $50 \mu m$.





710

711 **Figure 5**

712 UV-B-induced flavonoid accumulation in the epidermis is regulated by UVR8

713 localised in both the epidermis and mesophyll cells.

- 714 3-day-old seedlings were grown under WL supplemented with weak UV-B for 4 days
- 715 and were covered with WG305 (+UV-B) or with WG385 (-UV-B) filter. After

- incubation with DPBA, flavonoids were visualized (green colour) using CLSM. All images were taken using the same microscope settings. The focal plane was set to the bottom zone of the adaxial epidermis, where the highest signal was obtained (see bottom panel). Scale bar = $50 \mu m$.
- 720





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721

722 **Figure 6**

723 Effect of YFP-UVR8 expressed in different tissues of adult Arabidopsis plants

724 A,B. Phenotypic characterization of adult plants grown under white light

725 supplemented with UV-B.

A. Phenotypic characterization of 7-week-old Arabidopsis plants grown under white

727 light (WL, 120 $\mu mol~m^{-2}~s^{-1}$), WL plus UV-B at 2 $\mu mol~m^{-2}~s^{-1}$ or WL plus UV-B at

- 12 μ mol m⁻² s⁻¹ in short-day conditions. Scale bar: 5 cm.
- 729 B. Rosette diameter of 7-week-old plants grown as described above. Bars represent
- the average values calculated from three independent experiments. Error bars indicate
- the standard error of the mean.

732 C. Chlorophyll content of UV-B irradiated adult plants.

- 733 Chlorophyll levels were determined from 7-week-old plants grown under white light
- 734 or white light supplemented with UV-B (1.5 μ mol m⁻² s⁻¹) under short day
- 735 conditions. Chl (mg/g) represents total chlorophyll content (mg/g fresh weight). Five
- plants were used as biological replicates for each line and light treatment. Error bars
- 737 indicate standard error of the mean. Asterisks indicate values that are significantly
- different from WL treatment in the same genotype (Student's t-test, *P < 0.05, **P < 0.05, *P < 0.05,
- 739 0.01, ***P < 0.005). NS, no significance.
- 740
- 741

742 SUPPORTING INFORMATION

Additional Supporting Information can be found in the online version of this article at

- the publisher's web-site:
- 745 **Table S1**
- 746 **Complementation of** *uvr8* **phenotype by different transgenes.**
- 747 Figure S1
- 748 Full spectra of the applied light in the GroBank growth cambers
- 749 **Figure S2**
- 750 **Detection of YFP-UVR8 in the cotyledon**
- 751 Figure S3

- 752 Detection of YFP-UVR8 in the upper part of the hypocotyl
- 753 Figure S4
- 754 Detection of YFP-UVR8 in the lower part of the hypocotyl.
- 755 **Figure S5**
- 756 Determination of YFP-UVR8 accumulation in certain tissues.
- 757 Figure S6
- 758 UV-B induction of *ProHY5:GUS-GFP-NLS* in the cotyledon cells of transgenic
- 759 lines expressing YFP-UVR8 in different tissues.
- 760 Figure S7
- 761 The UV-B-specific messenger accumulation of *ELIP2* does whereas the
- 762 accumulation of *PRR9* does not depend on *HY5*.
- 763 **Figure S8**
- 764 UVB induction of *ProPRR9:GUS-GFP-NLS* in the cotyledon cells of transgenic
- 765 lines expressing YFP-UVR8 in different tissues.
- 766 Figure S9
- 767 Determination of endogenous and YFP-UVR8 protein levels in adult plants.

768 SUPPORTING INFORMATION

769

- Table S1
- 771 Figures S1-S9
- 772
- 773 Expression of UVR8 photoreceptor in different tissues reveals tissue-autonomous
- 774 features of UV-B signalling
- 775
- 776 Péter Bernula, Carlos Daniel Crocco, Adriana Beatriz Arongaus, Roman Ulm, Ferenc
- 777 Nagy, András Viczián

line and transgene activity	seedling phenotype			adult phenotype	
	hypocotyl	cotyledon	flavonoid accumulation	rosette size	chlorophyll content
Col	+++	+++	+++	+++	+++
uvr8	-	-	-	-	-
<i>uvr8/ProUVR8:YFP- UVR8</i> (epidermis, subepidermis)	++	+++	++	++	++
<i>uvr8/ProML1:YFP- UVR8</i> (epidermis)	++	+++	++	++	++
<i>uvr8/ProCAB3:YFP-UVR8</i> (subepidermis)	++	-	++	+++	+++
<i>uvr8/ProSUC2:YFP- UVR8</i> (subepidermis, vasculature)	+	-	++	+++	+++

780

781 **Table S1**

782 Complementation of *uvr8* phenotype by different transgenes.

783 This table summarizes the results obtained from different phenotype analyses. Crosses

784 mark the level of *uvr8* mutant complementation. Wild type (Col) plants show fully

extended responses (+++) whereas *uvr8* mutant shows no UVR8-specific responses in

the assays (-). The first column also indicates the tissue types where the YFP-UVR8

787 was observed by CLSM.





Detection of YFP-UVR8 in the cotyledon. Localization of YFP-UVR8 fusion protein in *uvr8-6* mutant seedlings expressing ProUVR8:YFP-UVR8, or ProML1:YFP-UVR8 or ProCAB3:YFP-UVR8 or ProSUC2:YFP-UVR8 was monitored by confocal laser scanning microscopy in the cotyledons of seedlings grown for 4 days in constant white light supplemented with UV-B. Images were obtained using identical microscope settings and YFPspecific signal is presented. White arrows mark positions of selected nuclei in the epidermis, yellow arrows point at nuclei in the mesophyll whereas red arrows indicate nuclei/cells in the vasculature.

Figure S2



Figure S3

Detection of YFP-UVR8 in the upper part of the hypocotyl. A. Focus was set to the epidermis and subepidermis. **B.** Focus was set to the subepidermis and vasculature. Localization of the YFP-UVR8 fusion protein was monitored by CLSM in the upper part of the hypocotyls of seedlings grown for 4 days in constant white light supplemented with UV-B. Images were obtained using identical microscope settings, with the exception of images taken of the *ProSUC2:YFP-UVR8* line which is presented with enhanced signal for better visibility. White arrows mark positions of selected nuclei in the epidermis, yellow arrows point at nuclei in the mesophyll whereas red arrows indicate nuclei/cells in the vasculature.



Detection of YFP-UVR8 in the lower part of the hypocotyl. A. Focus was set to the epidermis and subepidermis. **B.** Focus was set to the subepidermis and vasculature. Localization of the YFP-UVR8 fusion protein was monitored by CLSM in the lower part of the hypocotyls grown for 4 days in constant white light supplemented with UV-B. Images were obtained using identical microscope settings. White arrows mark positions of selected nuclei in the epidermis; yellow arrows point at nuclei in the mesophyll whereas red arrows indicate nuclei/cells in the vasculature.

Figure S4



803 Figure S5

804 Determination of YFP-UVR8 accumulation in certain tissues.

805 Nuclear YFP-UVR8 signal was determined using confocal laser scanning microscopy.

- 806 uvr8-6 seedlings expressing (A) ProUVR8:YFP-UVR8 (UVR8) and ProML1:YFP-
- 807 UVR8 (ML1) transgenes in the epidermal cell layer of the cotyledon (**B**)
- 808 ProUVR8: YFP-UVR8 (UVR8) and ProCAB3: YFP-UVR8 (CAB3) ProSUC2: YFP-
- 809 UVR8 (SUC2) transgenes in the subepidermal cell layer of cotyledon were examined
- 810 using identical microscope settings. Error bars indicate standard error of the mean
- 811 (n>60).







UV-B induction of ProHY5:GUS-GFP-NLS in the cotyledon cells of transgenic lines expressing YFP-UVR8 in different tissues. ProHY5:GUS-GFP-NLS was introduced into transgenic uvr8-6 lines expressing ProUVR8:YFP-UVR8 (UVR8) ProML1:YFP-UVR8 (ML1) or ProCAB3:YFP-UVR8 (CAB3). Localization of the GUS-GFP-NLS fusion protein was monitored by confocal laser scanning microscopy in the epidermis and mesophyll cells of the cotyledon of 7-day-old seedlings irradiated with constant WL supplemented with (+UV-B) or without (-UV-B) UV-B (1.5 μ mol m⁻² sec⁻¹). Identical microscope settings were used to allow the visualisation signal difference between the +UV-B and -UV-B image pairs. White arrows mark positions of selected nuclei in the epidermis; yellow arrows indicate nuclei in the mesophyll. Scale bar = $50 \mu m$.

UVR8

epidermis











mesophyll

CAB3

mesophyll





ML1





820 accumulation of *PRR9* does not depend on *HY5*.

821 7-day-old seedlings were irradiated with a 90 min WL pulse mixed with UV-B. Half

822 of the seedlings were covered with WG305 filter (UV-B), the other half was covered

823 with WG385 filter (no UV-B). After the irradiation, seedlings were collected and

824 transcript level of *ELIP2* (A) and *PRR9* (B) was determined using quantitative RT-

825 PCR. The relative transcript level (UV-B/no UV-B) is plotted in the figure. Ws=

826 Wassilewskaya wild type; uvr8=uvr8-7 mutant; hy5=hy5-ks50 mutant (both in Ws

827 ecotype).



Figure S8 UVB induction of *ProPRR9:GUS-GFP-NLS* in the cotyledon cells of transgenic lines expressing YFP-UVR8 in different tissues.

ProPRR9:GUS-GFP-NLS was introduced into transgenic uvr8-6 lines expressing ProUVR8:YFP-UVR8 (UVR8) ProML1:YFP-UVR8 (ML1) or ProCAB3:YFP-UVR8 (CAB3). Localization of the GUS-GFP-NLS fusion protein was monitored by CLSM in the epidermis and mesophyll cells of the cotyledon of 7-day-old seedlings irradiated with constant WL supplemented with (+UV-B) or without UV-B (-UV-B). Identical microscope settings were used to allow the visualisation signal difference between the +UV-B and –UV-B image pairs. White arrows mark positions of selected nuclei in the epidermis, yellow arrows indicate nuclei in the mesophyll. Scale bar = $75 \mu m$.



831 Figure S9

832 Determination of endogenous and YFP-UVR8 protein levels in adult plants.

- 833 Total protein extract was isolated from 7-week-old seedlings grown under short day
- 834 conditions in the greenhouse. Proteins were separated by gel electrophoresis blotted
- 835 onto a membrane and hybridized with anti-UVR8-specific antibody (anti-UVR8). The
- 836 blot was reprobed with anti-ACTIN antibody to check the even loading.
- 837