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Comparative study of wheat low-molecular-weight glutenin and alpha-gliadin trafficking in tobacco cells

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1 **Comparative study of wheat low-molecular-weight glutenin and**
2 **α -gliadin trafficking in tobacco cells.**

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33 **Abstract**

34 Wheat prolamins are the major proteins that accumulate in endosperm cells and are largely
35 responsible for the unique biochemical properties of wheat products. They are accumulated in
36 the endoplasmic reticulum (ER) where they form protein bodies (PBs) and are then
37 transported to the storage vacuole where they form a protein matrix in the ripe seeds. Whereas
38 previous studies have been carried out to determine the atypical trafficking pathway of
39 prolamins, the mechanisms leading to ER retention and PB formation are still not clear. In
40 this study, we examined the trafficking of a low-molecular-weight glutenin subunit (LMW-
41 glutenin) and α -gliadin fused to fluorescent proteins expressed in tobacco cells. Through
42 transient transformation in epidermal tobacco leaves, we demonstrated that both LMW-
43 glutenin and α -gliadin were retained in the ER and formed mobile protein body-like
44 structures (PBLs) that generally do not co-localise with Golgi bodies. An increased
45 expression level of BiP in tobacco cells transformed with α -gliadin or LMW-glutenin was
46 observed, suggesting the participation of this chaperone protein in the accumulation of wheat
47 prolamins in tobacco cells. When stably expressed in BY-2 cells, LMW-glutenin fusion was
48 retained longer in the ER before being exported to and degraded in the vacuole, compared to
49 α -gliadin fusion, suggesting the involvement of intermolecular disulphide bonds in ER
50 retention, but not in PBLs formation. Co-localisation experiments showed that gliadins and
51 LMW-glutenin were found in the same PBLs with no particular distribution, which could be
52 due to their ability to interact with each other as indicated by yeast two-hybrid assays.

53

54 **Keywords**

55 Wheat - Gliadin - Low-Molecular-Weight Glutenin - Tobacco cells - Trafficking -
56 Fluorescent proteins

57

58 **Key message**

59 Wheat low-molecular-weight-glutenin and α -gliadin were accumulated in the endoplasmic
60 reticulum and formed protein body-like structures in tobacco cells, with the participation of
61 BiP chaperone. Possible interactions between these prolamins were investigated.

62

63 **Abbreviations:**

64 BiP: Binding protein

65	BY-2:	Bright Yellow-2
66	ER:	Endoplasmic reticulum
67	GFP:	Green fluorescent protein
68	HMW:	High molecular weight
69	HMW-G:	High-molecular-weight glutenin
70	LMW:	Low molecular weight
71	LMW-G:	Low-molecular-weight glutenin
72	mRFP ₁ :	Monomeric red fluorescent protein
73	PDI:	Protein disulphide isomerase
74	PBLS:	Protein body-like structure
75	PB:	Protein body

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95 **Introduction**

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97 Wheat is the most widely cultivated cereal and the most widely consumed food crop in the
98 world. Wheat seed storage proteins, prolamins, are the major components that accumulate in
99 wheat endosperm cells, and are largely responsible for the viscoelasticity properties of gluten.
100 Prolamins are divided into two fractions, glutenins and gliadins, according to their
101 polymerisation state. Glutenins are polymers that have intra- and interchain disulphide bonds
102 composed of low-molecular-weight and high-molecular-weight glutenin subunits (LMW-G
103 and HMW-G, respectively) (Payne 1987). Gliadins are monomers with intramolecular
104 disulphide bonds (α and γ -gliadins) or without disulphide bonds (ω -gliadins) (Bietz et al.
105 1977). Gliadins account for about 40-50% of the total proteins in wheat seeds and play an
106 important role in the nutritional and processing quality of flour. γ -gliadins have been
107 considered to be the oldest proteins in the wheat prolamins family. HMW-glutenin subunits are
108 composed of a central repetitive domain flanked by two non-repetitive regions containing
109 cysteine residues critical for glutenin cross-linking. LMW-glutenin subunits are structurally
110 related to monomeric α and γ -gliadins except that they contain two additional cysteine
111 residues available for interchain disulphide formation, which would be responsible for the
112 polymeric feature of these proteins (Shewry and Tatham 1997).

113 In seed endosperm cells, prolamins are synthesized in the endoplasmic reticulum (ER), where
114 they accumulate. They form storage structures known as protein bodies (PBs) in the ER, and
115 are then transported to the storage vacuole (Parker and Hawes 1982; Levanony et al. 1992;
116 Altschuler et al. 1993; Galili et al. 1993). It seems that the major part of prolamins are
117 transported from the ER to the storage vacuole, bypassing the Golgi, via the PBs in
118 endosperm cells (Levanony et al. 1992; Galili et al. 1993; Hara-Nishimura et al. 1998; Vitale
119 and Galili 2001). Prolamins were also recently found to accumulate in maize aleurone cells in
120 a similar Golgi-independent pathway (Reyes et al. 2011). However, prolamins were also
121 detected in the Golgi stacks, especially during the early stage of seed development (Parker
122 and Hawes 1982; Galili et al. 1993; Shy et al. 2001; Loussert et al. 2008; Tosi et al. 2009).

123 PBs disappear during seed maturation to form the protein matrix around the starch granules
124 (Rubin et al. 1992). In wheat seeds, the distribution of the different prolamins within the
125 protein bodies is still unclear. No particular spatial organisation of gliadins and glutenins has
126 been observed in wheat PBs (Loussert et al. 2008), contrary to the prolamins found in maize
127 (Coleman et al. 1996, 2004) and rice PBs (Nagamine et al. 2011). However, a segregation of

128 protein type into a specific population of protein bodies can occur within the same endosperm
129 cell (Tosi et al. 2009). A segregation of gluten proteins occurred both between and within the
130 protein bodies during protein deposition. A recent study indicated that differential patterns of
131 distribution were found for the HMW glutenin subunits and γ -gliadin (more abundant in the
132 inner endosperm layers) when compared to the LMW-G and other gliadins (ω and α -gliadins
133 were more abundant in the subaleurone) (Tosi et al. 2011).

134 The mechanisms leading to the trafficking and accumulation of prolamins during seed
135 development are only partially known. Indeed, prolamins do not contain the classical ER
136 retention signals such as HDEL/KDEL, and the process leading to their retention in the ER is
137 not known. Several mechanisms have been suggested, including the involvement of the
138 repeated motif of prolamins for ER retention, as demonstrated for wheat gliadins expressed in
139 yeast (Rosenberg et al. 1993), xenopus oocytes (Altschuler et al. 1993) and, more recently, in
140 *Nicotiana tabacum* leaves (Francin-Allami et al. 2011). These repeated motifs could be used
141 as an anchor for the interaction of prolamins to the ER membrane, which would explain their
142 retention in the ER (Banc et al. 2009). However, the repeated sequence was not the only one
143 to contribute to the accumulation of prolamins in the ER and PBs. Disulphide bonds could
144 also play an essential role in ER retention and the accumulation of wheat prolamins (Shimoni
145 and Galili 1996; Lombardi et al. 2009). More recently, it was proposed that both cysteine-rich
146 C-terminal and repeated N-terminal domains of γ -gliadin were involved in γ -gliadin ER
147 retention and protein body-like structure formation (Francin-Allami et al. 2011). The
148 participation of chaperone proteins and, in particular, the ER luminal BiP, was also
149 demonstrated (Levanony et al. 1992; Li et al. 1993; Muench et al. 1997; Saito et al. 2009). In
150 rice, it has been suggested that BiP is necessary to maintain prolamins in a competent state for
151 subsequent assembly in the ER (Muench et al. 1997). It has also been suggested that the
152 disulphide isomerase (PDI) participates in the segregation of rice prolamins within the ER
153 lumen (Takemoto et al. 2002).

154 Trafficking of a wheat γ -gliadin and HMW-G subunit fused to a fluorescent protein was
155 recently investigated, demonstrating that tobacco cells are a good model for studying atypical
156 wheat prolamins trafficking (Francin-Allami et al. 2011; Saumonneau et al. 2011). In order to
157 further elucidate the trafficking pathway of wheat seed storage prolamins, we decided to study
158 other typical wheat prolamins components such as a LMW-G subunit and an α -gliadin, which
159 account for the majority of the storage proteins in the endosperm. To do this, we used a
160 fluorescent protein fusion strategy and a tobacco cell model. The results indicated that the two

161 prolamins were accumulated in the ER network and were able to form numerous protein
162 body-like structures in tobacco cells. We also investigated the potentiality of interactions
163 between the different prolamins. Taken together, these results provide global information
164 concerning the trafficking and assembly of wheat seed storage proteins.

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208 **Materials and methods**

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210 Plasmid constructs

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212 Gateway system cloning was used to perform the different constructs used in this study,
213 according to the manufacturer's instructions (Invitrogen). The binary vectors pMDC83
214 (Curtis and Grossniklaus 2003) and pK7RWG2 (Karimi et al. 2005) were used to construct C-
215 terminal GFP/mRFP₁ fusions with the different wheat ("Recital" variety) prolamin cDNAs.
216 The cDNA encoding α -gliadin (see nucleotide and amino acid sequences in supplementary
217 Fig. S1) was amplified using the primers AttB1-A4-PS (5'-aaa aaa gca ggc ggc ttc atg aag acc
218 ttt ctc atc) and AttB2-A4 (5'-caa gaa agc tgg gtc gtt agt acc gaa gat), followed by a second
219 PCR with the primers AttB1 (5'-ggg gac aag aag aag ttt gta caa aaa agc agg ct) and AttB2 (5'-
220 ggg gac cac ttt gta caa gaa agc tgg gt). The cDNA encoding the low molecular weight
221 glutenin (LMW-G) (see nucleotide and amino acid sequences in supplementary Fig. S2) was
222 amplified using the primers AttB1-LMWG-PS (5'-aaa aaa gca ggc ttc atg aag acc ttc ctc) and
223 AttB2-LMWG (5'-caa gaa agc tgg gtc gta ggc acc aac tcc ggt), followed by a second PCR
224 with the primers AttB1 and AttB2.

225 The constructs of γ -gliadin in fusion with GFP and mRFP₁ were previously described in
226 Francin-Allami et al. (2011).

227 In order to perform yeast two-hybrid experiments, the cDNAs encoding the α -gliadin and
228 LMW-G were cloned into pACT2 prey and pAS2-1 bait vectors (Clontech). The cDNAs were
229 amplified using the primers A4NcoIVRV-5' (5'-c cat gcc atg gca gtt aga gtt cca gtg cca caa
230 ttg-3') and A4BamHI-3' (5'-cg gga tcc tca gtt agt acc gaa gat gcc aaa tgg-3') for α -gliadin
231 amplification, and the primers L3NcoI-5' (5'-c cat gcc atg gca cag atg gag act aga tgc atc cct-
232 3') and L3BamHI-3' (5'-cg gga tcc tca gta ggc acc aac tcc ggt-3') for LMW-G amplification.
233 γ -gliadin cDNA was cloned into pAS2-1 vectors using the primers Gb5ncoI-NIQ-5' (5'-c cat
234 gcc atg gca aat ata cag gtc gac cct agc ggc-3') and Gb5bamHI-3' (5'-cg gga tcc tca ttg gcc acc
235 aat gcc ggc-3'). NcoI and BamHI were used as restriction sites.

236

237 Transient expression in plants

238

239 Transient expression in tobacco leaf epidermal cells was performed as described by Sparkes et
240 al. (2006). *Nicotiana tabacum* plants were grown in a growth chamber under a 16 h

241 photoperiod control at 21°C for 5-6 weeks prior to *Agrobacterium* infiltration. The
242 recombinant binary vectors were transferred into *Agrobacterium tumefaciens* (strain
243 GV3101::mp90) by heat shock transformation. Transformed *Agrobacteria* were inoculated in
244 a Luria-Bertani (LB) medium supplemented with the appropriate antibiotics at 28°C
245 overnight. Cultures were then centrifuged and washed with infiltration buffer (5 g l⁻¹ glucose,
246 50 mM MES, 2 mM Na₃PO₄·12H₂O, 100 μM acetosyringone) before being resuspended with
247 this same buffer at a DO600nm=0.1 and 0.05 for prolamin constructs and marker vectors (ER
248 and Golgi), respectively. The diluted bacteria were delivered to lower leaves of *N. tabacum*
249 by gentle pressure infiltration using a syringe. For experiments using co-expression, bacteria
250 containing the constructs were mixed in appropriate volumes before performing the
251 infiltration.

252

253 Stable transformation of BY-2 suspension cells

254

255 Suspension-cultured tobacco cells (BY-2, Bright Yellow 2) were provided by Chris Hawes
256 (Oxford Brookes University, UK), and cultured at 24°C in the dark in a Murashige and
257 Skoog-MES medium (Duchefa, no. MO254) supplemented with 20 mg l⁻¹ KH₂PO₄, 100 mg
258 l⁻¹ myo-inositol, 30 g l⁻¹ sucrose, 1 mg l⁻¹ thiamine and 0.2 mg l⁻¹ 2,4-D, pH 5.6. The binary
259 gateway-plasmid constructs were transferred into the GV3101::mp90 *Agrobacterium*
260 *tumefaciens* strain by heat shock. Transformed *Agrobacteria* were selected in LB medium
261 supplemented with appropriate antibiotics and were used to transform BY-2 cells as described
262 by Gomord et al. (1998). The transformed BY-2 calli were then screened by fluorescent
263 microscopy and selected calli were used to initiate transgenic suspension cell cultures. One ml
264 of cells was transferred to a fresh 30 ml of medium every week.

265 To compare the growth rate curves in the different BY-2 lines, the cells were filtered each day
266 from independent Erlenmeyer flasks on a 0.45 μm Millipore filter, and the fresh cell weight
267 determined.

268

269 Confocal laser scanning microscopy

270

271 Confocal microscopy experiments were performed at the Biopolymers-Structural Biology
272 platform, INRA Nantes (France). An inverted Nikon A1 confocal laser scanning microscope
273 was used to examine the subcellular localisation of GFP and mRFP₁ fluorescence. The
274 samples were examined with a water-immersion X40 objective. GFP and mRFP channels

275 were acquired by alternately scanning with line switching, using a 488 nm argon laser line for
276 excitation of the GFP and a 561 nm laser line for mRFP₁; emissions were collected via a
277 photomultiplier through a 515+/-30 nm and a 595+/-50 nm band-pass filters, with the same
278 gain settings for every experiments. Images were processed using NIS-Element (Nikon) and
279 Image J software.

280

281

282 Transmission electron microscopy

283

284 Infiltrated leaves were cut in 1 mm³ samples and placed in a fixation solution consisting of
285 2% (w/v) paraformaldehyde and 2.5% (v/v) glutaraldehyde in 0.1M phosphate buffer (pH
286 7.2). Air was removed from leaf samples with air vacuum cycles. Samples were then fixed at
287 4°C in the fixation solution overnight. Fixed samples were rinsed in phosphate buffer and de-
288 ionised water prior to postfixation with 1% osmium tetroxide during 1 hour. Dehydration
289 was performed in 30, 50, 70, 85, 95 and 100% ethanol solutions. Samples were successively
290 impregnated with 50/50 and 0/100 ethanol/propylene oxide (v/v) for 1 hour, then 50/50
291 propylene oxide/Epon (Epoxy Embedding Medium Kit, 45359-1EAF Sigma Aldrich) (v/v) 1
292 hour and 0/100 overnight. The impregnated samples were polymerised for 24 h at 55°C and
293 60 h at 72°C. A Leica EM UC7 Ultramicrotome (Leica) was used for making ultrathin
294 sections (80 nm) for electron microscopy. Sections were collected on copper grids,
295 counterstained with 2% uranyl acetate and examined on a JEOL 1230 TEM (JEOL;
296 accelerating voltage of 80 kV).

297

298 Treatments with Latrunculin B inhibitor

299 Tobacco leaf sections were incubated for 40 min in a 25-µM solution of Latrunculin B
300 (Calbiochem). Treated cells were then visualised by confocal microscopy. Each treatment
301 with inhibitor was repeated at least twice with similar results.

302

303 Protein extraction and immunoblot analysis

304

305 Tobacco BY-2 cells were harvested by filtration and resuspended in a SDS sample buffer
306 (100 mM TRIS-HCL, pH 6.8, 4% SDS, 20% glycerol, 5% 2-mercaptoethanol, 100 µg ml⁻¹
307 bromophenol blue) prior to boiling for 5 min, and then centrifuged. The supernatant was
308 subjected to SDS-PAGE according to Laemmli (Laemmli, 1970) with a separating gel

309 containing 4-20% acrylamide in gradient (mini-PROTEAN TGX gels, Biorad). Each protein
310 extract loaded on the gel corresponded to the same volume of filtered cells.

311 Infiltrated tobacco leaf sections were cut, crushed in liquid nitrogen, rinsed in acetone, and
312 dried. The leaf powders were weighed and an equal amount of powders was resuspended in
313 SDS sample buffer. 15 mg of leaf extracts were loaded to the each lanes of gel for
314 immunoblotting.

315 Proteins were then electrotransferred to a nitrocellulose membrane (IBlot Gel Transfer Stacks
316 Nitrocellulose, Invitrogen) before immunodetection by specific antibodies. Immunoblotting
317 was performed using mouse monoclonal anti-GFP antibodies (dilution: 1:2500) (Clontech), or
318 rabbit polyclonal anti-BiP antibodies (dilution: 1:4000) (Hatano et al. 1997). An alkaline
319 phosphatase-conjugated goat anti-mouse (dilution: 1:2000) or anti-rabbit (dilution: 1:3000),
320 respectively, were used for the detection. Proteins were stained with BCIP-NBT (Promega
321 kit) according to the manufacturer's instructions.

322

323 Yeast two-hybrid assays

324

325 Yeast two-hybrid assays were performed in *Saccharomyces cerevisiae* strain Y190 (*MATa*,
326 *gal4*, *gal80*, *his3*, *trp1*, *ade2*, *ura3*, *leu2*, *URA3::GAL1::lacZ*, *LYS2::GAL4(UAS)::HIS3*
327 *cyh^R*). Y190 was transformed with a pair of plasmids expressing Gal4BD fusion TRP1 (bait)
328 and Gal4AD fusion LEU2 (prey), plated onto selected media (YNBD medium (0.17% yeast
329 nitrogen base without amino acids, 2% glucose, 2% agar) supplemented with appropriate
330 amino acids and bases) for 4-5 days at 28°C, and tested for β -galactosidase production in an
331 overlay plate assay.

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344 **Results**

345

346 Prolamin constructs

347

348 Wheat α -gliadin has two main domains: a repeated N-terminal domain rich in glutamine and
349 proline residues, and a C-terminal domain containing six cysteines to form intramolecular
350 bonds. LMW-G has three domains: one repeated domain rich in glutamine and proline
351 surrounded by a small non-repeated domain with one cysteine at the N-terminal extremity,
352 and a larger non-repeated domain containing seven cysteines at the C-terminal extremity (Fig.
353 1). Both α -gliadin and LMW-G have a signal peptide that is cleaved off during protein
354 synthesis. In order to investigate the trafficking of wheat α -gliadin and LMW-G, the
355 corresponding genes were isolated from wheat endosperm cDNA at 250 degree days after
356 anthesis. They were then cloned into binary vectors containing a double 35S promoter of
357 cauliflower mosaic virus and the enhanced green fluorescent protein (eGFP) or monomeric
358 red fluorescent protein (mRFP₁) coding sequences downstream from the gateway cassette,
359 leading to a C-terminal GFP/mRFP₁ fusion of the prolamins (Fig. 1). These constructs were
360 used to transform tobacco leaf epidermal cells and BY-2 suspension cultured cells in order to
361 study the subcellular localisation of α -gliadin and LMW-G.

362

363 Gliadin and glutenin localisation in tobacco cells

364

365 The subcellular localisation of α -gliadin and LMW-G was first investigated after transient
366 transformation in epidermal tobacco leaf cells via *Agrobacterium* infiltration. Cells expressing
367 the α -gliadin fused to GFP presented fluorescence in the ER compartment, which was
368 confirmed by the co-localisation with an ER marker (mRFP₁-DPL1; Marion et al. 2008;
369 Tsegaye et al. 2007) (Fig. 2a, b). The fluorescence observed by confocal microscopy was
370 mainly represented by punctate structures instead of a typical ER network. These punctate
371 structures have already been described for γ -gliadin transformed tobacco leaves where they
372 were identified as protein body-like structures (PBLs) (Francin-Allami et al. 2011). α -gliadin
373 PBLs often form clusters that look like bunches of grapes, leading to large aggregated
374 structures. We also fused LMW-G with green fluorescent protein and used it to transiently
375 transform tobacco leaf cells. The observation of fluorescence by confocal microscopy

376 indicated that LMW-G was also expressed in ER and forms large PBLs (> 4 μm in diameter),
377 which tended to group together to form clusters (Figs. 2d, e). Like those formed by the
378 expression of gliadin, these PBLs seemed to be surrounded by the ER marker DPL1, which is
379 an ER membrane lyase, indicating that ER membranes encapsulate the bodies (Fig. 2e, high
380 magnification of the inset). α -gliadin and LMW-G seemed to be able to form more
381 systematically and well-defined PBLs compared to those previously obtained with γ -gliadin
382 constructs. Moreover, quantitative data indicated that whereas α -gliadin was accumulated as
383 numerous and relatively small PBLs, LMW-G formed less numerous but often larger PBLs
384 (Fig. 3). Western blot analysis of leaf cells expressing α -gliadin-GFP and LMW-G-GFP with
385 anti-GFP antibodies confirmed the integrity of the fused proteins, with an apparent molecular
386 mass of approximately 70 kDa (Fig. 4a). A fragment of lower mobility was also present at a
387 very low level, and corresponded to the mass of free GFP (27 kDa).

388 α -gliadin and LMW-G PBLs are highly mobile, and we demonstrated using Latrunculin
389 treatment that this mobility is actin-dependent (see movies S3 and S4 in supplementary data).
390 Because PBLs are very mobile and similar to Golgi bodies, we tested the co-localisation with
391 the Golgi marker, ST-RFP (sialyl transferase signal anchor-mRFP₁). As previously seen for
392 γ -gliadin-GFP, the result indicated that α -gliadin and LMW-G PBLs generally did not appear
393 to be co-localised with Golgi bodies (Figs. 2c, f), which was confirmed by a different pattern
394 of movement of PBLs and Golgi bodies when time-lapse series of confocal images were
395 analysed (see movie S3 in supplementary data).

396

397 PBLs morphology

398

399 In order to reveal the morphology of the gliadins and glutenin PBLs, the transformed tobacco
400 leaves were submitted to appropriate treatment (see the Materials and Methods section), and
401 examined by electron microscopy. The focalisation on the ER region of tobacco cells showed
402 relatively round shape structures next to conventional tubular ER when cells expressed α -
403 gliadin or LMW-G (Figs. 5a, b). These structures reflected the PBLs structures observed with
404 confocal microscopy. The PBLs structures were not seen to be continuous with the ER in
405 these images. It is however not excluded that PBLs were still linked to the ER network, or in
406 contrary they bud off the ER and localized in the cytosol. Their size (< 1 μm) corresponded to
407 the smallest PBLs observed with fluorescent microscopy. This was consistent with previously

408 studies describing PBLs formed in tobacco leaves after expression of maize β -zein (Randall
409 et al. 2005) or wheat γ -gliadin (Francin-Allami et al. 2011).

410

411 Gliadin and LMW-glutenin are degraded in the vacuole of BY-2 cells

412

413 In order to examine the dynamic trafficking of prolamins in a stable system, we expressed α -
414 gliadin and LMW-G fused to GFP in BY-2 suspension cultured cells. As previously seen after
415 transient infiltration in tobacco leaves, α -gliadin and LMW-G were localised in the ER
416 network (Fig. 6a). Numerous and well-defined PBLs were formed with both prolamins.
417 Quantitative data indicated that both prolamins could form PBLs with a diameter larger than
418 1 μ m, and that α -gliadin tended to include more cells with numerous and larger PBLs than
419 cells expressing LMW-G (Fig. 3). After 2-4 days of subculturing, no fluorescence was seen in
420 the vacuole (Table 1). After four days of subculturing, in addition to the ER and PBLs,
421 fluorescence was also seen in the vacuole of the α -gliadin transformed cells. At 7 days of
422 subculturing, almost all of the fluorescence was localised in the vacuole. These results were in
423 accordance with the changes previously observed in the case of γ -gliadin, where fluorescence
424 in the vacuole corresponded to the release of GFP core after transport of γ -gliadin-GFP to the
425 vacuole (Francin-Allami et al. 2011). In LMW-G expressing cells, fluorescence was also
426 clearly detected in the ER with the formation of numerous PBLs. The release of fluorescence
427 into the vacuole was slower since no or little fluorescence was observed in the vacuole of
428 cells at 7 days after subculturing, and even at 10 days after subculturing, there was still
429 fluorescence in the ER network in addition to the vacuole (Fig. 6a, Table 1). These results
430 were confirmed by immunoblotting analysis where LMW-G-GFP was still detected at 10 days
431 after subculturing, whereas only GFP is detected in the cells expressing α -gliadin as early as 7
432 days of subculturing (Figs. 7a, b). It is noteworthy that no cell growth difference was detected
433 between the two lines expressing α -gliadin or LMW-G (supplementary Fig. S5). The same
434 pattern of degradation was obtained for α - and γ -gliadin-GFP, whereas LMW-G showed a
435 slower degradation than gliadins. It is also interesting to note that the amount of PBLs
436 decreased with the stage of cell development for both transformed BY-2 cell lines (Table 1).
437 Whereas α -gliadin-GFP transformed cells tended to have more PBLs at 5-7 days of
438 subculturing but with rapid degradation in the vacuole, LMW-G-GFP transformed cells
439 presented a fluorescence mainly localised in the ER network with less PBLs at these stages
440 (Table 1, Fig. 6).

441

442 BiP expression in transformed tobacco cells

443

444 Because BiP is a typical ER-resident protein, and some studies have indicated that it is
445 involved in the folding and assembly of proteins in the ER, we decided to follow the
446 expression of this chaperone protein during the expression of gliadin and glutenin in tobacco
447 cells. Immunoblotting indicated an increased expression of BiP in epidermal tobacco cells
448 infiltrated with α -gliadin-GFP or LMW-G-GFP compared to the non-transformed plant (Fig.
449 4b). In the same way, expression of BiP is increased in the BY-2 suspension cultured cells
450 stably transformed with α -gliadin and LMW-G (Figs. 7a, b). Moreover, the expression of BiP
451 decreased as days of subculturing increased, which is in relation to the progressive
452 degradation of fused prolamins. At 10 subculturing days of cells expressing LMW-G or 7
453 subculturing days of cells expressing α -gliadin, the expression level of BiP was identical to
454 that of non-transformed BY-2. Besides, we demonstrated that the expression level of BiP was
455 relatively constant during subculturing of non-transformed BY-2 cells (Fig. 7c). In addition,
456 differential centrifugation indicated that a part of the BiP in BY-2 cells expressing α -gliadin-
457 GFP at four days of subculturing was found in the vacuolar fraction (Fig. 7d). This result is
458 consistent with those previously found (Tamura et al. 2004).

459

460 Co-localisation of prolamins in tobacco cells

461

462 The distribution of the different prolamins within the protein bodies in wheat endosperm is
463 not really clear. To determine their distribution in PBLs, gliadins and LMW-G were co-
464 expressed in tobacco leaf epidermal cells (Fig. 8). Confocal images indicated that γ -gliadin
465 and α -gliadin co-localised in protein bodies. All PBLs inside the tobacco cells contained both
466 gliadins. In the same way, we demonstrated that LMW-G was co-localised with α -gliadin or
467 γ -gliadin in all protein bodies. In addition, at the resolution provided by confocal microscopy,
468 all these combinations of proteins appeared to be homogeneously distributed inside PBLs.
469 These co-localisations within PBLs lead to the possibility that prolamins could interact with
470 each other. In order to test this possibility, a yeast two-hybrid experiment was performed
471 using the different combinations of prolamins. A yeast two-hybrid system was previously
472 used to detect interactions between maize prolamins (Kim et al. 2002) or wheat γ -gliadin
473 domains (Francin-Allami et al. 2011). For this, gliadins and LMW-G genes were cloned into

474 specific yeast two-hybrid vectors, in order to fuse the genes at their N-terminals with the
475 activation domain or binding domain of the yeast GAL4 transcription factor. The results
476 indicated that all prolamins could interact with each other in an *in vitro* yeast two-hybrid
477 system (Table 2). α -gliadin seemed to have a particularly good ability to interact with other
478 proteins, revealed by the fact that the detected interactions were strong when one of the
479 interacting proteins was α -gliadin. We also demonstrated that α -gliadin and LMW-G were
480 able to interact with themselves in this yeast two-hybrid context. Negative controls were
481 performed to check that proteins were not able to directly interact with the GAL4 activation
482 or binding domain. Thus, interactions tested by the yeast two-hybrid system reinforced the
483 possible interaction between gliadins and LMW-G in PBLs, as suggested by co-localisation
484 experiments with confocal microscopy.

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504 **Discussion**

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506 We demonstrated that wheat α -gliadin and LMW-G fused to GFP are stably expressed in non-
507 seed tissue, indicating that intrinsic molecular properties of these proteins are involved in ER
508 retention and PBLs biogenesis. This result was also found in the case of the expression of γ -
509 gliadin (Francin-Allami et al. 2011), HMW-glutenin (Saumonneau et al. 2011) and γ -zein
510 (maize storage protein) (Randall et al. 2005) in tobacco leaves. In addition, ER retention of
511 LMW-G was also demonstrated in tobacco protoplasts (Lombardi et al. 2009). Both α -gliadin
512 and LMW-G have a repeated proline-rich N-terminal domain and a more hydrophobic
513 disulphide-rich domain, like the majority of wheat prolamins. As previously suggested, these
514 particular domains are both probably involved in prolamins trafficking. It has been proposed
515 that the hydrophobicity and disulphide bond formation promote specific interactions that
516 result in the formation of large self-assemblies that are responsible for retention in ER and PB
517 formation (Kawagoe et al. 2005; Pompa and Vitale 2006). The isolated C-terminal domain of
518 wheat γ -gliadin in fusion with GFP was retained and was able to form PBLs when expressed
519 in tobacco leaves or BY-2 cells (Francin-Allami et al. 2011). Based on site-directed
520 mutagenesis of prolamins cysteine residues, the critical role of specific cysteines for PB
521 formation was demonstrated for maize γ -zein (Llop-Tous et al. 2010).

522 It has been shown that the N-terminal proline-rich repeated domain of prolamins participates
523 in ER retention and PB formation in maize (Geli et al. 1994; Llop-Tous et al. 2010) and wheat
524 (Francin-Allami et al. 2011), although the repeated motif sequences of γ -zein are different
525 from those of wheat prolamins. A minimal repeated motif sequence is required for PB
526 induction. For example, the deletion of 105 amino acids from the repeated sequence of the
527 HMW-glutenin fused to GFP reduces the ability of the glutenin to form PBLs in the ER
528 (Saumonneau et al. 2011). The ability of a hydrophobic elastin-like protein (ELP) fusion to
529 induce PB formation in plant leaves was recently shown, indicating the generally powerful
530 impact of repeated motif sequences on PB formation for heterologous expression (Conley et
531 al. 2009; Saumonneau et al. 2011).

532

533 In contrast to gliadins, glutenins form intermolecular disulphide bonds. The LMW-G
534 sequence is very similar to monomeric α and γ -gliadins, except that it contains two additional
535 cysteine residues available for interchain disulphide formation. In tobacco cells, gliadin and
536 glutenin can be retained in the ER and form PBLs. However, we showed that LMW-G-GFP

537 was less extensively and more slowly degraded when expressed in BY-2 cells. This could be
538 linked to the presence of these two supplementary cysteine residues and the possibility of
539 LMW-G to form intermolecular bonds. It was previously shown that a fusion between the
540 *Phaseolus vulgaris* vacuolar storage protein, phaseolin, and the N-terminal half of the *Zea*
541 *mays* prolamin, γ -zein, forms interchain disulphide bonds that facilitate the formation of ER-
542 located protein bodies (Pompa and Vitale 2006). Regarding the transient expression in
543 tobacco leaves or stable transformation of BY-2 suspension cells, the quantitative data do not
544 make it possible to distinguish a significant difference in PBLs formation between α -gliadin
545 and LMW-G. However, we noted that in BY-2 cells, the amount of PBLs tends to decrease
546 with the age of subculturing. Although α -gliadin was more rapidly degraded with the release
547 of GFP in the vacuole, more PBLs were observed at the exponential phase growth compared
548 to the LMW-G-GFP expressing cells, where fluorescence was preferentially in the ER
549 network. Although it seems difficult to directly link the quantity of PBLs to the degradation
550 of prolamins and the release of GFP in the vacuole, this result suggested that aggregated
551 prolamins in PBLs could be more easily degraded in the vacuole than prolamins localised in
552 the ER network, but this remains to be proven.

553 In wheat endosperm, prolamins are expressed in abundance. Protein concentration may play
554 an important role in prolamin accumulation and PB formation. This is probably the reason
555 why zein-GFP expressed in tobacco was not assembled into stable PBs, whereas when it was
556 expressed in a higher concentration, it was stably retained in the ER and forms PBs (Foresti et
557 al. 2008; Llop-Tous et al. 2010). In this case, a double 35S promoter was used to allow a high
558 expression of α -gliadin or LMW-G in tobacco cells in order to mimic the strong expression of
559 prolamin in wheat endosperm cells.

560
561 Molecular chaperones such as BiP and PDI are involved in ER quality control, and are known
562 to increase in cells under ER stress (Vitale and Ceriotti 2004). Several studies have
563 demonstrated that BiPs are involved during storage protein synthesis and stress response
564 (Hatano et al. 1997; Muench et al. 1997; Takaiwa et al. 2009). In addition to the intrinsic
565 prolamin properties and their expression level, molecular chaperones could play an important
566 role in ER retention and PB formation (Levanony et al. 1992; Muench et al. 1997; Saito et al.
567 2009). It was suggested that the interaction of the ER luminal BiP with zein proteins could
568 contribute to their stability and the ability to form protein bodies (Randall et al. 2005). In our
569 study, immunoblotting revealed an increase of the BiP expression level in tobacco cells

570 transformed with α -gliadin or LMW-G, compared to the basal BiP expression level of non-
571 transformed cells. Therefore, as previously reported, BiP seems to participate in the
572 accumulation of prolamins in the heterologous tobacco cells. In addition to assist in protein
573 folding, BiP participates in the protein degradation process known as ER-associated
574 degradation (ERAD). In this case, BiP expression decreased according to the days of
575 subculturing in transformed BY-2, which is concomitant to the degradation of the prolamins
576 fusions. Differential centrifugation indicated that a part of the BiP in BY-2 cells expressing α -
577 gliadin-GFP was found in the vacuolar fraction. These results imply that BiP could contribute
578 to the degradation process of α -gliadin or LMW-G-GFP constructs and their transport to the
579 vacuole of tobacco cells, as previously demonstrated for other ER-resident protein
580 degradation (Tamura et al. 2004; Pimpl et al. 2006). Further experiments would have to be
581 carried out to determine the role of BiP and other folding proteins such as PDI on the
582 retention and accumulation of prolamins. Moreover, it was demonstrated that ER protein
583 aggregates expressed in transgenic cultured cells are degraded by autophagy when cells reach
584 the stationary phase (Tamura et al. 2004; Toyooka et al. 2006). The degradation of α -gliadin
585 or LMW-G in BY-2 cells could also be the result of an autophagy mechanism. However,
586 neither ERAD nor autophagy processes involved traffic through the Golgi apparatus, in
587 contrast with the observation made in this study for the degradation of α -gliadin-GFP and
588 LMW-G-GFP in BY-2 cells. The processing of α -gliadin-GFP and LMW-G-GFP could
589 involve an alternative pathway of vacuolar delivery via a Golgi-dependent transport, which
590 also implies the BiP chaperone (Frigerio et al. 2001; Pimpl et al. 2006).

591
592 Through the co-localisation experiment, we demonstrated that α -gliadin and LMW-G were
593 found in the same PBLs after transient infiltration in epidermal tobacco cells, and seemed to
594 be homogeneously distributed in these protein bodies. This was consistent with results
595 previously reported in wheat seed by immunostaining (Loussert et al. 2008; Tosi et al. 2009).
596 However, this does not exclude the possibility that differential patterns of distribution of the
597 seed storage prolamins exist in endogenous wheat endosperm tissue, as was recently
598 demonstrated (Tosi et al. 2009, 2011). We demonstrated that all prolamins were able to
599 interact with each other in a yeast two-hybrid assay. However, in this yeast two-hybrid
600 system, protein interactions take place in a reducing environment, which is not totally
601 appropriate to permit the formation of intrachain disulphide bonds, which could lead to a
602 misfolding (Orsi et al. 2001). *In vivo* experiments should be assessed in order to confirm

603 these interactions in an appropriate environment context, in particular, to highlight the role of
604 inter- and intrachain disulphide bonds in the trafficking of gliadins and glutenins. It would
605 also be important to determine which part of the proteins contribute to the interaction in order
606 to better understand the molecular mechanisms involved in the prolamin assembly formation.

607

608 To our knowledge, this is the first time that a LMW-G was fused to GFP and expressed in a
609 plant model. Certainly due to the numerous repeated motifs and disulphide bonds along the
610 sequences, prolamins, and particularly glutenins, are difficult to manipulate at the genetic
611 level. We recently investigated the trafficking of high-molecular-weight glutenin fused to
612 GFP in tobacco cell suspension BY-2, and we demonstrated its retention in ER and its ability
613 to form PBLs (Saumonneau et al. 2011). The role of the repeated and cysteine-rich domains
614 of γ -gliadin was explored using a similar strategy, leading to the conclusion that tobacco cells
615 are a good model for studying atypical wheat prolamin trafficking with fluorescent protein
616 fusions (Francin-Allami et al. 2011). Our study was focused on wheat LMW-glutenin and α -
617 gliadin, which are two major types of wheat prolamins, and demonstrated their trafficking in
618 tobacco cells. Taken together, these studies provide general information on the trafficking
619 pathway of all wheat seed storage prolamins.

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622 **Supplementary material**

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624 Supplementary data are available at Plant Cell Reports online.

625 **Supplementary Fig. S1.** Nucleotide and amino acid sequences of α -gliadin used in this study.

626 **Supplementary Fig. S2.** Nucleotide and amino acid sequences of LMW-G used in this study.

627 **Supplementary Movie S3.** α -gliadin protein body-like structures (α -gliadin-GFP in green)
628 moved throughout the cell along the ER network and are not co-localised with Golgi bodies
629 (ST-mRFP₁ in red).

630 **Supplementary Movie S4.** Movie showing the absence of PBLs movement after Latrunculin
631 B treatment.

632 **Supplementary Fig. S5.** No cell growth difference was detected between the non-
633 transformed BY-2 and BY-2 expressing α -gliadin-GFP or LMW-G-GFP.

634

635

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637

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Figure legends

Figure 1 Schematic representation of the α -gliadin and LMW-glutenin constructs used in the study. The Gateway recombination system was used to replace the CmR/*ccdB* vector cassette by the α -gliadin or LMW-glutenin cDNA, in order to construct C-terminal GFP/mRFP₁ fusions. All fusion constructs were placed under the control of the cauliflower mosaic virus double 35S promoter (2X35S) and the nopaline synthase terminator (T). Repeats (light grey box): N-terminal repeated sequence; C-rich (dark grey box): C-terminal cysteine-rich sequence; GFP/mRFP₁: green fluorescent protein/monomeric red fluorescent protein; black square: ER signal peptide; B1/B2/R1/R2: recombination sites; CmR/*ccdB*: Chloramphenicol resistance gene/*ccdB* gene; short lines: cysteine residues. The diagram is not drawn to scale.

Figure 2 Subcellular localisation of α -gliadin and LMW-glutenin fluorescent fusions in tobacco epidermal cells. Confocal microscope images showing GFP fusion proteins (green) expressed alone or coexpressed with mRFP₁ fusion markers (red) in tobacco leaf epidermal cells, 2-3 days after infiltration with *Agrobacterium* suspension cultures. Images of cells expressing α -gliadin-GFP (green) (a); α -gliadin-GFP (green) coexpressed with an mRFP₁-DPL1 ER marker (red) and merged image (b); α -gliadin-GFP (green) coexpressed with a Golgi marker, ST-mRFP₁ (red), and merged image (c); LMW-glutenin-GFP (green) (d); LMW-glutenin-GFP (green) coexpressed with an mRFP₁-DPL1 ER marker (red) and merged image (e); LMW-glutenin-GFP (green) coexpressed with a Golgi marker, ST-mRFP₁ (red) and merged image (f). Scale bars = 10 μ m.

Figure 3 Classification of the transformed cells according to the protein body-like structures they contain.

Analysed cells were classified into three categories according to the density of prolamins protein body-like structures (PBLs) within the ER. The analysis was generated from confocal sections of transformed cells. The first category corresponds to cells containing a high PBLs density (more than 5 puncta/10 μ m²); the second category corresponds to cells containing a low PBLs density (less than 5 puncta/10 μ m²); and the third category corresponds to cells containing no apparent PBLs. The size of PBLs was taken into account in cell classification. Cells were provided from at least three individual transformation experiments in tobacco leaf

cells or BY-2 cells, transformed with each of the two constructs. A total of about 50 cells were examined for each transformation. The data were collected from transformed BY-2 cells comprised between 3 to 7 days of subculturing. The values are expressed as a percentage.

Figure 4 Immunoblotting from tobacco leaf epidermal cells expressing α -gliadin-GFP and LMW-glutenin-GFP. An immunoblot was performed with anti-GFP antibodies (a) or anti-BiP antibodies (b). (M): protein marker; (1): tobacco leaf expressing α -gliadin-GFP; (2): tobacco leaf expressing LMW-glutenin-GFP; (3): non-infiltrated tobacco leaf.

Figure 5 Electron micrographs of tobacco leaf epidermal cells expressing fused prolamins. Electron microscope images representing tobacco leaf epidermal cells 2-3 days after infiltration with *Agrobacterium* suspension cultures. Electron micrographs show areas in leaf epidermal cells expressing α -gliadin-GFP (a) or LMW-G-GFP (b) where PBLs (arrows) is demonstrated next to conventional rough ER. CW: cell wall; ER: endoplasmic reticulum; V: vacuole; C: cytosol. Scale bars = 0.5 μ m.

Figure 6 Subcellular localisation of α -gliadin and LMW-glutenin fused to GFP in stable transgenic BY-2 cells. Subcellular localisation of α -gliadin-GFP or LMW-glutenin-GFP expressed in BY-2 cells 3, 5, 7 days or 10 days after subculturing. Scale bar = 20 μ m.

Figure 7 Immunoblotting from tobacco cell suspension BY-2 expressing α -gliadin-GFP or LMW-glutenin-GFP. α -gliadin-GFP and GFP core, resulting from the same immunoblot, were revealed by an anti-GFP antibody from BY-2 cells expressing α -gliadin-GFP 2, 4 or 7 days after subculturing (a). LMW-glutenin-GFP and GFP core, resulting from the same immunoblot, were revealed by an anti-GFP antibody from BY-2 cells expressing LMW-glutenin-GFP 2, 4, 7 or 10 days after subculturing (b). BiP was revealed with an anti-BiP antibody (Hatano et al. 1997) (a, b). nt: non-transformed BY-2 cells. Immunoblotting performed with anti-BiP antibody from cells expressing α -gliadin-GFP 2, 4, 7 and 10 days after subculturing (c). Immunoblotting performed with anti-BiP antibody from 4-day-old BY-2 cells expressing homogenate subjected to differential centrifugation as previously described (Tamura et al. 2004) (d). P1: pellet fraction after a 1000 g centrifugation of the homogenate; P10: pellet fraction after a 10000 g centrifugation of the S1 supernatant; P100: pellet fraction

(microsomal fraction) after a 100 000 g centrifugation of the S10 supernatant; S100: 100 000 g supernatant (soluble vacuolar fraction).

Figure 8 α and γ -gliadins and LMW-glutenin fused to fluorescent protein locate to the same protein body-like structures in tobacco leaf epidermal cells. Confocal images of tobacco leaf epidermal cells transiently expressing LMW-glutenin-GFP (green), α -gliadin-mRFP₁ (red) and merge (a). Confocal images of tobacco leaf epidermal cells transiently expressing LMW-glutenin-GFP (green), γ -gliadin-mRFP₁ (red) and merge (b). Confocal images of tobacco leaf epidermal cells transiently expressing α -gliadin-GFP (green), γ -gliadin-mRFP₁ (red) and merge (c). Constructs were infiltrated and imaged 3 days after inoculation. Scale bars = 10 μ m.

Figures

Figure 1

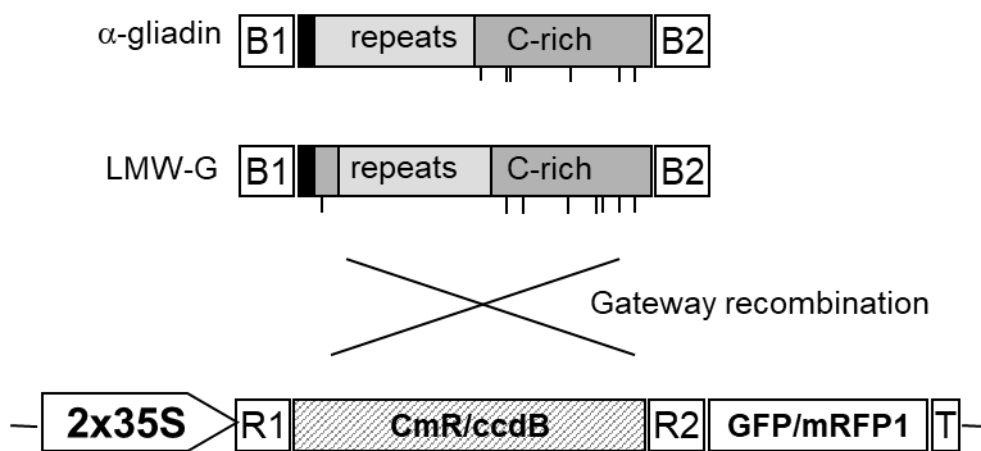


Figure 2

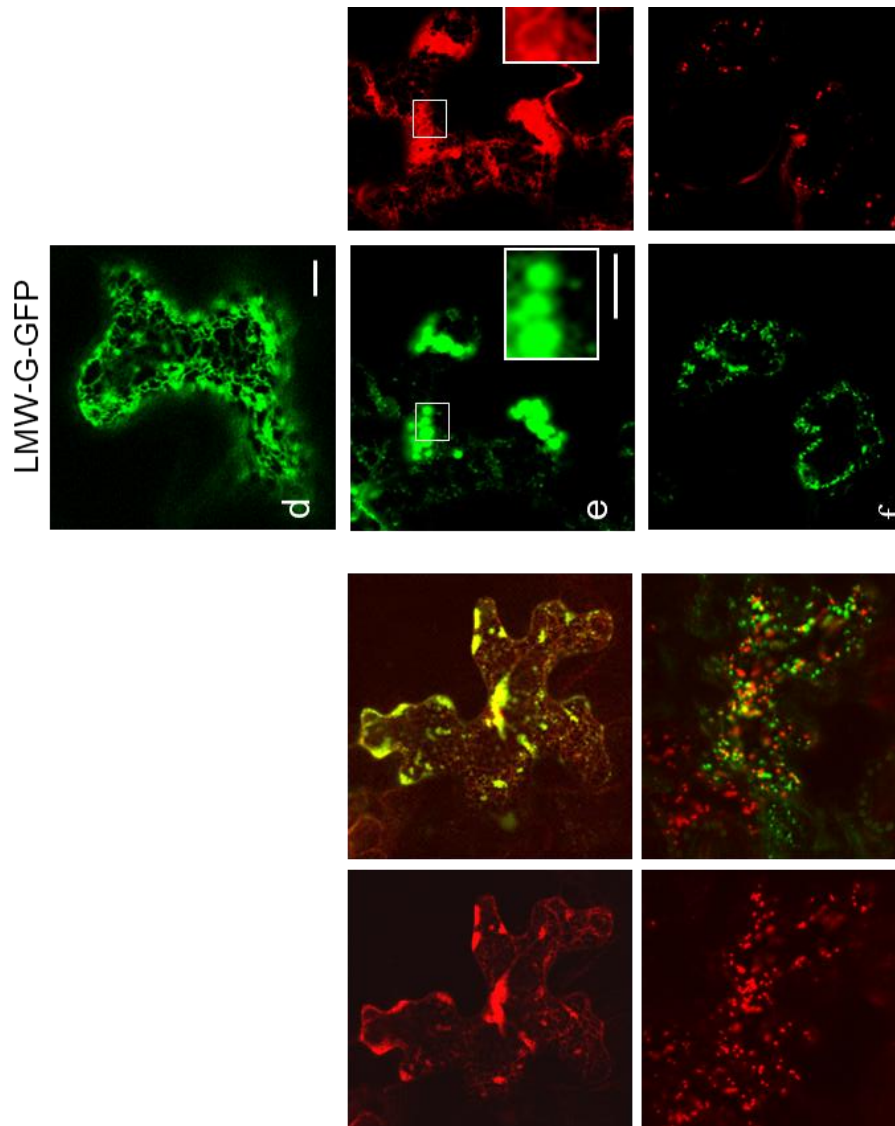


Figure 3

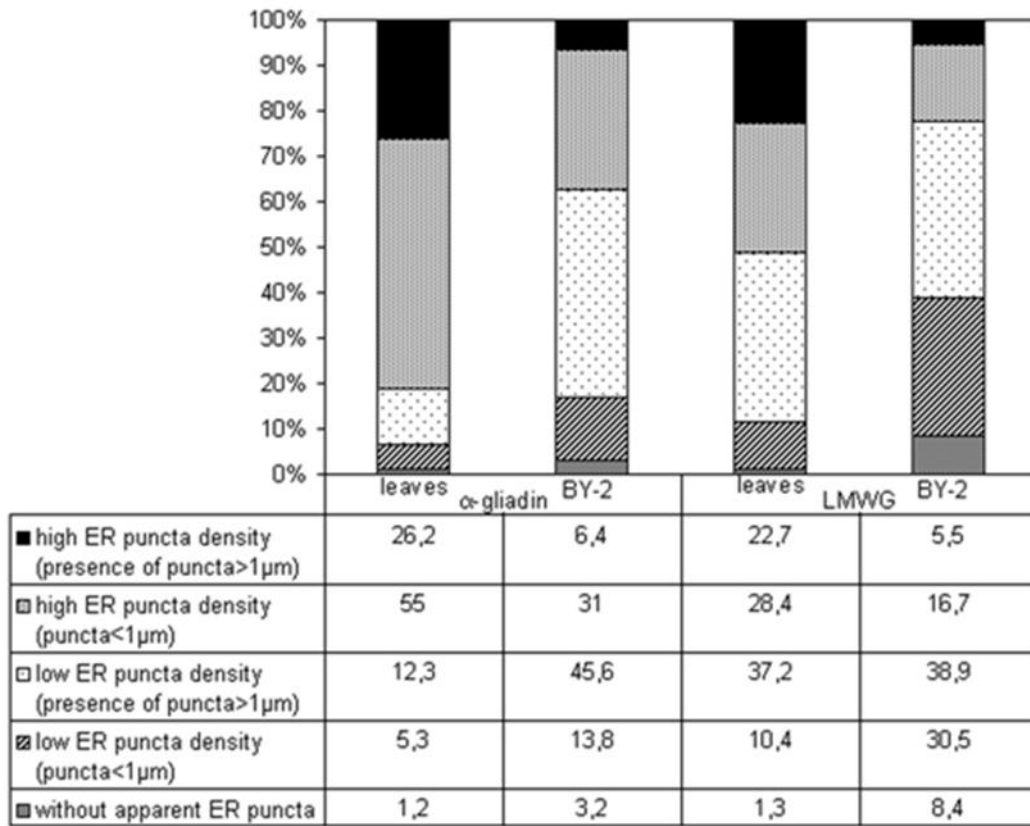


Figure 4

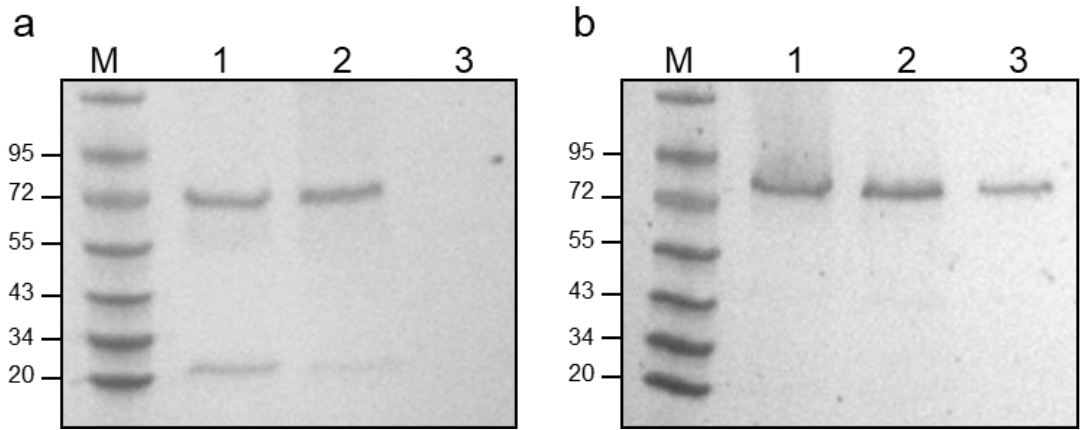


Figure 5

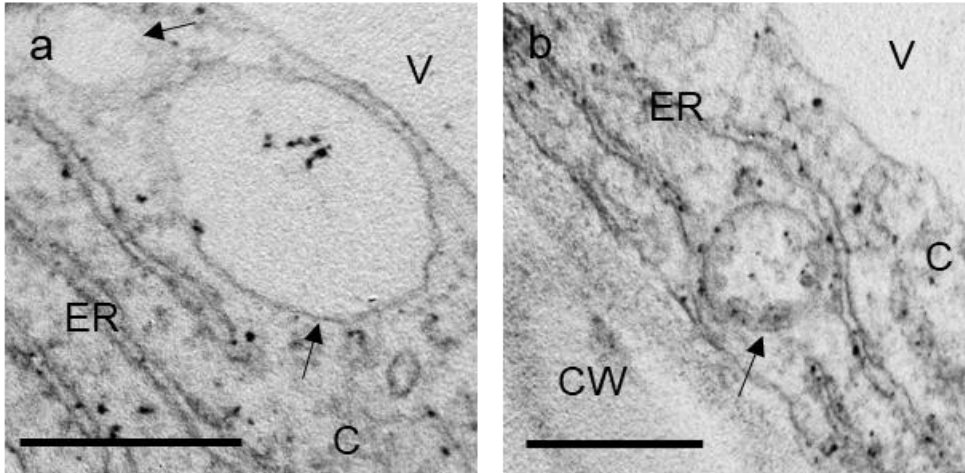


Figure 6

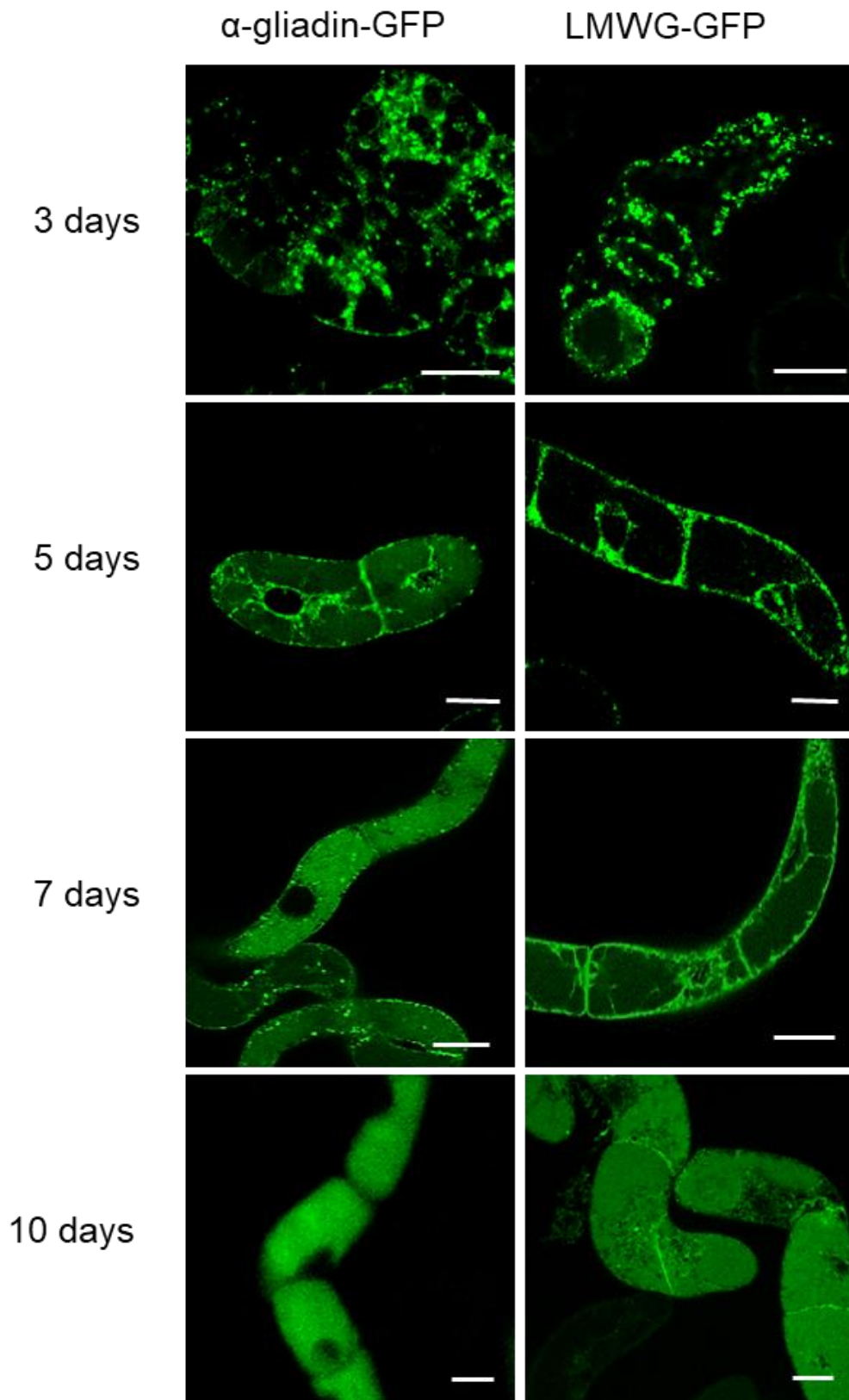


Figure 7

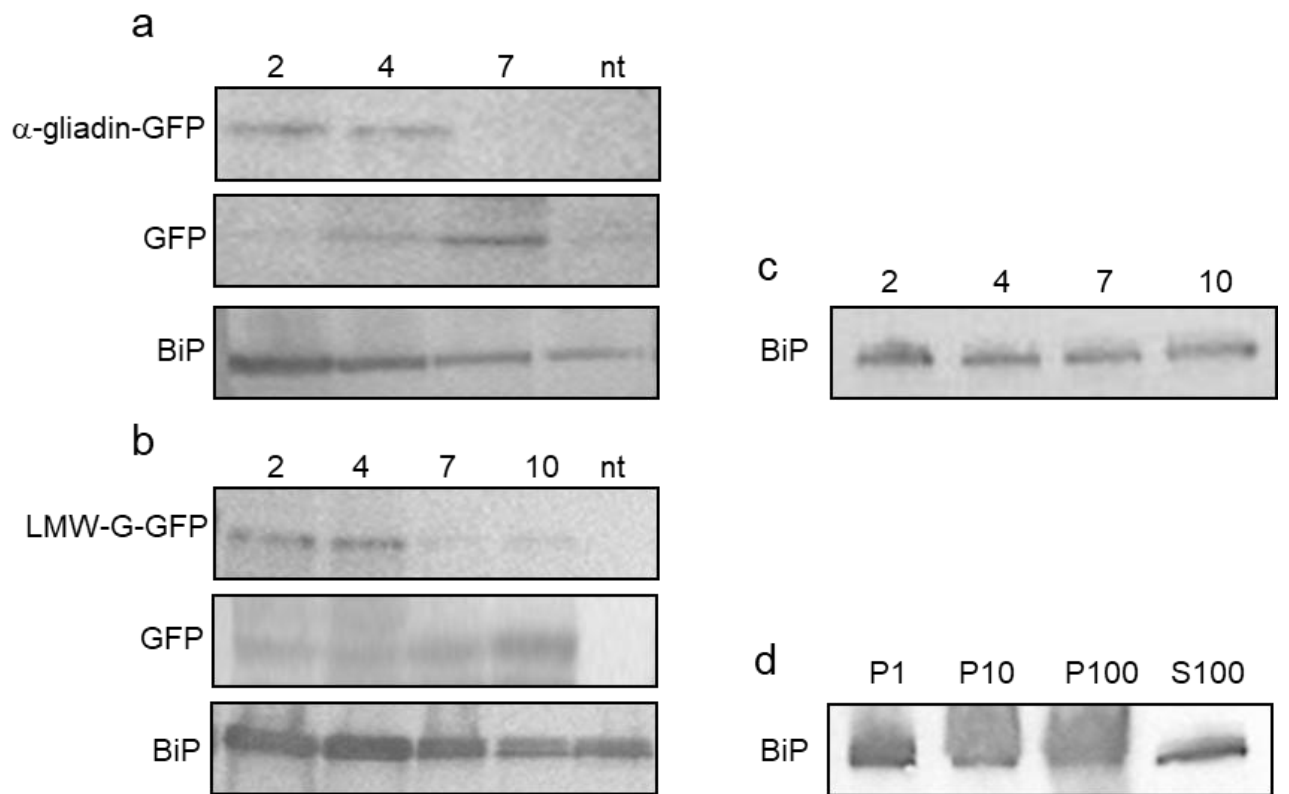
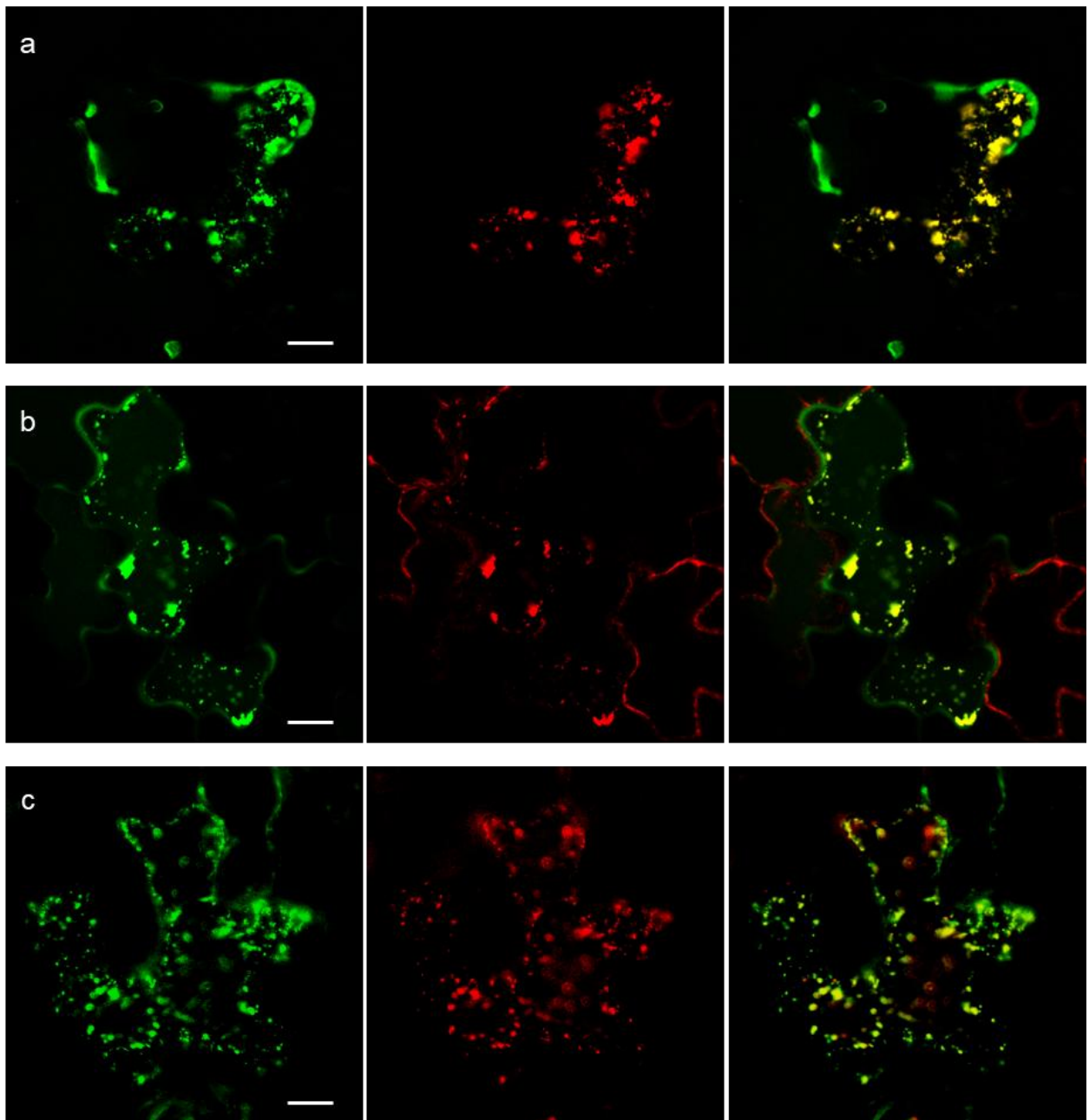


Figure 8



Tables

Table 1 Classification of transformed BY-2 cells according to their PBLs amount and vacuolar fluorescence during culture stages

Days of subculturing	cells with high PBLs density		cells with low PBLs density		cells without PBLs		vacuole fluorescence	
	(a)		(b)		(c)		(d)	
	α -gliadin	LMW-G	α -gliadin	LMW-G	α -gliadin	LMW-G	α -gliadin	LMW-G
3 d	85	63	14	25	1	12	0	0
5 d	15	13	77	72	8	15	73	2
7 d	2	5	62	43	36	52	98	56
10 d	0	0	0	9	100	91	100	91

Analysed cells were classified into three categories according to the density of PBLs at different stages of subculturing. The first category corresponds to cells with high PBLs density (more than 5 puncta/10 μm^2) (a); the second category corresponds to cells containing a low PBLs density (less than 5 puncta/10 μm^2) (b); and the third category corresponds to cells containing no apparent PBLs (c). In addition cells were also classified according to the vacuolar fluorescence they contain during subculturing stages (d). Cells provided from at least three individual transformation experiments in BY-2 cells transformed with each of the two constructs. A total of about 20 cells were examined for each transformation and culture stage. The values are expressed as a percentage.

Table 2 The interaction between α -gliadin, γ -gliadin and LMW-glutenin was performed using a yeast two-hybrid assay.

AD	BD	β -Gal
LMWG	α -gliadin	++
LMWG	γ -gliadin	+
LMWG	LMWG	+
LMWG	-	-
-	LMWG	-
α -gliadin	α -gliadin	++
α -gliadin	γ -gliadin	++
α -gliadin	LMWG	++
-	α -gliadin	-
α -gliadin	-	-

Column labelled “AD”: each prolamin was fused to the activator domain of GAL4 using the pACT prey vector; column labelled “DB”: each prolamin was fused to the binding domain of GAL4 using the pAS2.1 bait vector. Negative controls were performed by interacting constructs with empty prey or bait vectors (-). Results from β -galactosidase activity are summarised in the “ β -Gal” column, and marked with +, ++, or – in accordance with the blue coloration intensity.