1 A comprehensive toolkit for quick and easy visualization of marker proteins,

2 protein-protein interactions and cell morphology in Marchantia polymorpha

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13 Abstract

Even though stable genomic transformation of sporelings and thalli of Marchantia polymorpha is 14 15 comparatively straightforward and efficient, numerous problems can arise during critical phases of the process such as efficient spore production, poor selection capacity of antibiotics or low transformation 16 17 efficiency. It is therefore also desirable to establish quick methods not relying on stable transgenics to 18 analyze the localization, interactions and functions of proteins of interest. The introduction of foreign 19 DNA into living cells via biolistic mechanisms has been first reported roughly 30 years ago and has been 20 commonly exploited in established plant model species such as Arabidopsis thaliana or Nicotiana 21 benthamiana. Here we report the fast and reliable transient biolistic transformation of Marchantia 22 thallus epidermal cells using fluorescent protein fusions. We present a catalogue of fluorescent markers which can be readily used for tagging of a variety of subcellular compartments. Moreover, we 23 24 report the functionality of the bimolecular fluorescence complementation (BiFC) in M. polymorpha with the example of the p-body markers MpDCP1/2. Finally, we provide standard staining procedures 25 26 for live cell imaging in *M. polymorpha*, applicable to visualize cell boundaries or cellular structures, to 27 complement or support protein localizations and to understand how results gained by transient 28 transformations can be embedded in cell architecture and dynamics. Taken together, we offer a set of 29 easy and quick tools for experiments that aim at understanding subcellular localization, protein-30 protein interactions and thus functions of proteins of interest in the emerging early diverging land 31 plant model *M. polymorpha*.

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In the last decade the liverwort Marchantia polymorpha has emerged as a powerful model system to 34 study early land plant evolution due to its early evolutionary divergence in the land plant phylogenetic 35 36 tree (Shaw et al., 2011; Harrison, 2017; Morris, Puttick et al., 2017). Research deploying M. polymorpha has led to a series of insightful studies on the functional evolution of ABA (Lind et al., 2015; Eklund et 37 al., 2018) and JA signaling mechanisms (Monte et al., 2018; Monte et al., 2019; Peñuelas et al., 2019), 38 39 plant immunity (Carella et al., 2019; Gimenez-Ibanez et al., 2019), reproductive and vegetative 40 development (Flores-Sandoval et al., 2015; Proust et al., 2016; Jones and Dolan, 2017; Rövekamp et 41 al., 2017; Otani et al., 2018; Westermann et al., 2019) and cell division (Buschmann et al., 2015). It 42 offers the advantage of genetic and morphological simplicity in combination with its dominant haploid 43 vegetative life phase, allowing for fast generation of knockout mutants and subsequent phenotypic analyses, irrespectively of time-consuming homozygous mutant generation (Ishizaki et al., 2015 (B)). 44 45 Concomitantly, a plethora of molecular genetic tools was developed that include stable transformation of developing spores (Ishizaki et al., 2008) and regenerating thallus fragments (Kubota et al., 2013), 46 47 the suitability for genome editing via homologous recombination (Ishizaki et al., 2013) and 48 CRISPR/Cas9 (Sugano et al., 2014; Sugano et al., 2018), the cultivation in axenic conditions and on soil 49 and controlled crossing (Ishizaki et al., 2015 (B)).

50 Plant genetics and cell biological approaches generally rely on the efficient visualization of intracellular 51 features, including protein localization and organelle architecture or dynamics. In this regard, the 52 process of transient and stable transformation of plant cells is a powerful and commonly used 53 technique in molecular genetics and cell biology to study protein dynamics, as well as genetic and 54 physical (*i.e.* protein) interaction. It thus aids the elucidation of fundamental biological questions at 55 the (sub)cellular scale. While the performance of stable biolistic transformation of immature thalli and spores has been reported before (Chiyoda et al., 2008, Chiyoda et al., 2014), we describe here the 56 57 transient biolistic transformation of Marchantia thallus epidermal cells, a technique to study protein 58 localization that has commonly been used in other plant systems for 30 years (Sanford, 1990; Ueki et 59 el., 2009; Rasmussen et al., 1994). Importantly, we provide a comprehensive list of protein marker constructs that allows quick visualization of a variety of subcellular compartments within 24 hours and 60 61 the possibility for live-imaging. The marker list comprises constructs for visualization of the nucleus, 62 cytoplasm, plasma membrane, actin filaments, endosomes, peroxisomes, the Golgi apparatus and 63 processing bodies (Tab. S1).

64 Genetic interaction studies often rely on assessment of physical protein interactions to elucidate 65 intracellular signaling mechanisms. Therefore, the bimolecular fluorescence complementation 66 technique (BiFC; Hu et al., 2002; Walter et al., 2004) represents a time-efficient method to test for potentially interacting proteins *in vivo*. Hence, we also provide here evidence for the functionality ofBiFC in Marchantia epidermal cells.

In addition to transient expression, dye-based staining procedures represent a fast and reliable method to (co)visualize subcellular compartment architecture and dynamics. Therefore, we here provide a series of staining protocols for different organelles, both for Marchantia thallus epidermal cells and rhizoids and compare functionality regarding standard protocols used for the seed plant model *Arabidopsis thaliana*.

To further complement the here provided comprehensive cell biological marker toolkit, we compiled a list of additional Marchantia resources, methods, tools and databases (Tab. 1) that altogether will be useful for the young and growing community by complementing and supporting genetic / cell biological / biochemical approaches using *M. polymorpha* as a model system.

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79 Materials and Methods

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81 Plant material and growth conditions

The widely used *Marchantia polymorpha* Tak-1 (MpTak-1) ecotype was cultivated via propagation of vegetative propagules (gemmae) on solid Johnson's agar supplemented with 0.8 % micro agar under axenic conditions. Gemmae were grown under long day condition (16 h light/8 h darkness cycle) and white light irradiation (60 µmol m⁻¹ s⁻¹) at 21°C and 60 % humidity. After 2.5 - 3 weeks, a few thallus fragments of approximately 0.5 – 1 cm² were transferred onto a small petri dish (6 – 9 cm in diameter) containing fresh solid Johnson's medium on the day of transformation (Fig. 1A).

The Arabidopsis thaliana Col-0 ecotype, that was used for DAPI staining, was cultivated on soil and
 grown under long day conditions at 21°C and 120 μmol m⁻¹ s⁻¹ light intensity.

90

91 Cloning of DNA constructs

All constructs used in this study are summarized in Tab. S1, including their origin, promoter, fluorescent
 tag and oligonucleotide sequences used for PCR-based amplification of new constructs from
 Marchantia whole-thallus cDNA. The 35S promoter was used for all expression experiments (except
 for expression of AtSYP32, AtGot1p and Lifeactin) to guarantee comparability of subsequent analyses.
 The coding sequences of interest were cloned into Gateway (GW)-compatible entry vectors,
 pDONR201 and pDONR207 (Invitrogen), and then remobilized to be integrated in the respective GW

98 destination vectors (Tab. S1). The cloning procedure was as described before (Westermann et al.,99 2019).

100

101 DNA sample preparation for biolistic transformation

For a single shot, 300 ng of vector DNA were mixed with gold, serving as micro-carriers (30 mg/ml, 1 μ m), CaCl₂ (2.5 M), spermidine (0.1 M) and _{dd}H₂O under thorough shaking. Subsequently, micro-carriers were washed with 70% EtOH and 100% EtOH. The DNA-coated gold particles were suspended in 100% EtOH and placed onto macro-carriers. The EtOH was allowed to vaporize and the prepared macrocarriers were then used for biolistic transformation.

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108 Biolistic transformation procedure and efficiency of (co-)transformation

109 Marchantia thallus fragments were placed into a PDS-1000 / He Biolistic® Particle Delivery System (Bio-110 Rad). A vacuum of 25 in Hg vac was applied and the DNA-coated gold particles were shot at 900 psi from a distance of 10 cm. Finally, the bombarded plant material was allowed to recover for 24 h in 111 112 darkness while remaining in its humid environment, *i.e.* on the media in the closed petri dish (Fig. 1C). 113 Biolistic transformation generally yielded n > 50 transformed cells per sample shot. A representative 114 example for transformation efficiency is shown in Fig. 1B. Moderate to strong expression levels in each 115 individual cell could be observed irrespectively of the protein construct or promoter used (Fig. 1B; Tab. 116 S1). The use of strong promoters such as pro35S, proAtUBQ10 or proMpEF1 α can sometimes lead to 117 overexpression artefacts that may impede drawing secured conclusions. However, yielding a wide range of expression level in the same experimental round and plant sample allows for identification of 118 biologically meaningful protein localization patterns and to distinguish them from unwanted artefacts, 119 120 such as protein over-accumulations. In order to reliably assess the potential of transformed constructs 121 as single cell fluorescent markers, we co-bombarded all described vectors with either the nuclear 122 marker AtKRP1 or the plasma membrane markers AtNPSN12 or MpSYP13a fused to fluorescent tags 123 and subsequently created a collection of functional and useful Arabidopsis- and Marchantia-derived 124 fluorescent protein fusions (Tab. S1). In order to determine the efficiency of co-transformation, we counted cells expressing both markers in relation to the total number of transformed cells in 9 125 independent co-transformations of protein fusions used in this study. Successful biolistic co-126 127 transformation was on average 74% efficient (see Tab. S3).

128

129 Staining procedures

130 For FDA staining, young (2 – 5 days-old) gemmae were placed onto depression slides and covered with

an FDA solution (5mg/L FDA in ddH2O, diluted from a stock solution of 5 mg/ml FDA in acetone) for 5-

132 10 min. Afterwards, samples were rinsed in _{dd}H₂O.

For PI staining, young gemmae were placed onto depression slides and directly covered with a PI solution for 10 minutes (10 mg/L in $_{dd}H_2O$). Subsequently, samples were rinsed with $_{dd}H_2O$.

For FM4-64 staining, young gemmae were mounted onto depression slides in 2 μM FM4-64 diluted in
liquid Johnson's growth medium (Ishizaki et al., 2008) and allowed to incubate for 10 minutes prior to
imaging. For FM4-64 and FDA co-staining, gemmae were first stained in a FDA solution and then
mounted in a FM4-64 solution, both as described above.

139 For DAPI staining, several methods were used. Experiments were done using 0, 4, and 7 days old 140 gemmae. The DAPI staining solutions were composed of 10 – 100 mg/L DAPI in either 1xPBS-T 141 (phosphate buffered saline + 0.1% Tween-20) and 5% DMSO or ddH2O with 0.1 or 1% Tween-20 and 5% 142 DMSO. Different staining incubation times of 10, 30 or 60 minutes were tested. The staining was tested 143 with and without preceding or subsequent shaking of the samples in 70% EtOH at 80°C. To enhance 144 permeability of membranes, 10 or 50 mg/L digitonin was added to the aforementioned staining 145 solutions. As all attempts for staining living cells failed, the following fixation methods were tested. 146 Gemmae were fixed in a 3:1 EtOH:acetic acid mixture on ice for 1 h, washed 3 times in 100% EtOH and 147 stained in aforementioned DAPI solutions for 1 h. In another attempt, gemmae were fixed in 3% 148 glutaraldehyde in 1x PBS-T (phosphate buffered saline + 0.1% Tween-20) overnight, subsequently 149 washed in 1x PBS-T, and incubated in aforementioned DAPI solutions in darkness overnight. Furthermore, a modified version of a DAPI staining protocol published for gametophore leaves and 150 151 protonemata of Physcomitrella patens (Sato et al., 2017) was used. Gemmae were placed in 3.7% 152 formaldehyde in 1 x PBS for 30 min. Subsequently gemmae were immersed in 100% MeOH on ice for 153 10 min. Afterwards, gemmae were soaked in 1% Triton X-100 and then stained with the 154 aforementioned DAPI solutions for 30 min. Unfortunately, none of the experimental procedures 155 described here led to a reliable staining of nuclei by DAPI in viable or dead epidermal cells of M. 156 polymorpha gemmae.

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158 Confocal laser scanning microscopy

The transformed or stained plant material was transferred onto a depression slide supplemented with 300 μL _{dd}H₂O and covered with a 18x18 mm cover slip. Microscopic analysis was performed using a Leica SP8 CLSM with an argon gas laser intensity set at 20 %. Fluorophore excitation and fluorescence caption were performed at the wavelength spectra shown in Tab. S2. Images were taken using a digital

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163	gain of 100 % at a resolution of 1024 x 1024 pixels, a pinhole size of 1 AU, and a scan speed of 400-700
164	Hz using bidirectional confocal scanning and HyD hybrid detectors. For the caption of multiple
165	fluorophore types sequential or, if suitable, simultaneous scanning was performed.

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167 Data processing and analysis

168 Analysis of all microscopic captions was performed using ImageJ/FIJI (Schindelin et al., 2012), software 169 version 1.51n. Data manipulation included maximum projections from Z-stacks (\leq 20 frames, 1 µm slice 170 intervals) for some of the markers (as individually mentioned in the figure captions), as well as 171 generation of composite images from separate individual channels.

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173 Results

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175 A) Fluorescent protein markers to illuminate cellular compartments in Marchantia

To assess the potential capability of transiently transforming single Marchantia thallus epidermal cells,
 we first chose a set of proteins whose subcellular localization has been well studied in established
 model systems such as Arabidopsis or tobacco and thus could qualify as reliable subcellular markers in
 Marchantia as well.

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181 Nucleus

We first picked the *Arabidopsis thaliana* INHIBITOR OF CYCLIN-DEPENDENT KINASE 1 (AtICK1)/ KIP-RELATED PROTEIN 1 (AtKRP1), which localizes to the nucleus and functions in cell growth, differentiation and cell cycle progression (Wang et al., 1998; De Veylder et al., 2001; Schnittger et al., 2003; Weinl et al., 2005; Jakoby et al., 2006). Upon biolistic transformation of Marchantia thalli, we observed AtKRP1-CFP protein localization to the nucleus of epidermal cells (Fig. 2A). We therefore cotransformed AtKRP1 as a nuclear marker and indicator of successful cell transformation in subsequent experiments.

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190 Plasma membrane

191 As a second potential marker, we chose the Arabidopsis thaliana NOVEL PLANT SNARE 12 (AtNPSN12), 192 which represents a non-polar plasma membrane-localized protein commonly used as plasma 193 membrane marker (Alassimone et al., 2012; Kirchhelle et al., 2016). Biolistically transformed 194 Marchantia thallus epidermal cells showed AtNPSN12-mCherry fluorescence at the cell periphery 195 consistent with plasma membrane localization (Fig. 2B). To confirm this localization, we co-196 transformed AtNPSN12-mCherry with the known Marchantia plasma membrane marker mCitrine-197 MpSYP13a (Kanazawa et al., 2016; Fig. 2C). As single and co-bombardments with AtNPSN12 showed 198 (co)localization to the plasma membrane, we conclude that AtNPSN12-mCherry and mCitrine-199 MpSYP13a are both suitable plasma membrane markers for Marchantia epidermal cells (Fig. 2D).

200 Receptor-like kinases of the Malectin-like receptor kinase subfamily have been the subject of intensive 201 research in the past years given their multitude of functions in plant development and immunity 202 signaling (Franck et al., 2018 A). The Malectin-like receptors (MLRs) ANXUR1 and 2 (AtANX1/2) control 203 cell wall integrity during pollen tube growth (Boisson-Dernier et al., 2009; Miyazaki et al., 2009) and 204 negatively regulate plant immune responses in Arabidopsis (Mang et al., 2017) at the plasma 205 membrane. During pollen tube growth control, AtANX1/2 act genetically upstream of the cytosolic and 206 plasma membrane-attached receptor-like cytoplasmic kinase of the PTI1-like family, AtMRI, while the 207 AtANX1 homolog AtFERONIA (AtFER) acts upstream of AtMRI during root hair growth control (Boisson-208 Dernier et al., 2015). We showed recently that tip-growth control in Marchantia rhizoids relies on an 209 evolutionarily conserved signaling module comprised of the unique Marchantia MLR MpFER and its 210 downstream component and unique Marchantia PTI1-like MpMRI, both of which show plasma 211 membrane-localization comparable to their respective Arabidopsis homologs (Westermann et al., 212 2019). We transiently co-expressed the fluorescent protein fusions AtMRI-YFP, MpFER-YFP and 213 MpMRI-YFP with AtNPSN12-mCherry. While MpFER-YFP showed signal exclusive to the plasma 214 membrane, AtMRI and MpMRI displayed plasma membrane localization with traces in the cytoplasm 215 as reported before (Fig. 3; Boisson-Dernier et al., 2015; Westermann et al., 2019).

216 Noteworthily, we also wanted to test expressing the plasma membrane localized Arabidopsis MLRs in 217 Marchantia and thus co-transformed AtANX1-RFP with mCitrine-MpSYP13a and AtFER-Citrine with 218 AtNPSN12-mCherry. Intriguingly, while many cells expressed the plasma membrane markers mCitrine-219 MpSYP13a and AtNPSN12-mCherry, a great majority of them did not show expression of either AtANX1 220 or AtFER (Fig. S1A, B). This suggests that, unlike MpFER, the Arabidopsis MLRs fused to single 221 fluorescent tag cannot be expressed in Marchantia epidermal cells. Thus, we next tried to express 222 AtFER with a triple Citrine tag instead of a single one. It resulted in many Citrine-expressing cells but 223 mostly in the cytoplasm, with no hints of plasma membrane localization (Fig. S1C). These results 224 indicate that fusion of long protein tags may prevent transmembrane receptor kinases such as MLRs

225 to be correctly integrated into cellular membranes. To check if this was specifically due to Arabidopsis 226 proteins or to certain protein families, we co-expressed MpFER-3xCitrine with MpFER-TdTomato and 227 MpMRI-3xCitrine with MpMRI-RFP (Fig. S1D, E). Interestingly, the 3xCitrine tag didn't perturbate the cytosolic and plasma membrane localization of MpMRI, as MpMRI-3xCitrine co-localized with MpMRI-228 229 RFP at the cell periphery. However, while MpFER-TdTomato exhibited PM localization, MpFER-230 3xCitrine-derived signal was clearly present in the cytoplasm. Therefore, for some plasma membrane-231 localized protein families, fusion with a triple tag can lead to localization artefacts, and the use of single 232 tag is thus recommended by default. Why MpFER but neither AtFER nor AtANX1 can be expressed in 233 Marchantia thallus epidermis remains puzzling.

234

235 Cytoplasm

The *A. thaliana* type-one protein phosphatases (TOPP) AtATUNIS1/2 have recently been reported as negative regulators of cell wall integrity maintenance during Arabidopsis tip-growth (Franck et al., 2018 B). The nucleocytoplasmic localization of AtAUN1-YFP and AtAUN2-YFP was demonstrated in Arabidopsis pollen tubes and leaf epidermal cells (Franck et al., 2018 B). In Marchantia epidermal cells, expression of AtAUN1/2-YFP led to a comparable nucleocytoplasmic localization, as opposed to the co-expressed plasma membrane localized AtNPSN12-mCherry fusion (Fig. 4), therefore qualifying these phosphatases as reliable Marchantia nucleocytoplasmic markers.

243

244 Endosomes

245 As for endosomal compartments, we chose two Ras-related in brain (RAB) GTPases, the canonical 246 MpRAB5 and the plant-unique MpARA6, that were recently described in *M. polymorpha*. Both proteins 247 were successfully expressed in stably transformed lines and co-localized to endosomal punctate 248 structures stained by FM1-43 (Minamino et al., 2017). Upon biolistic co-transformation of the protein 249 fusions mCherry-MpRAB5 and MpARA6-YFP with the nuclear marker AtKRP1-CFP (Fig. 5A and B), we 250 found a comparable localization in punctate structures for both markers. Moreover, both GTPases 251 strongly co-localized with each other (Fig. 5C) showing that MpRAB5 and MpARA6 are suitable 252 endosomal markers also for transient transformation studies.

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254 Peroxisomes

255 The carboxyl-terminal amino acid sequence serine-lysine-leucine (SKL) is well known as the consensus 256 peroxisomal targeting sequence 1 (PTS1) and is sufficient to induce protein targeting and import to 257 peroxisomes. SKL was first shown to be able to signal protein import into peroxisomes of mammalian cells (Gould et al., 1989) but later was also found to be functional in yeast and plants (Keller et al., 258 259 1991). In Arabidopsis, SKL motif fused to fluorescent tags is frequently used as a peroxisomal marker (Mathur et al., 2002, Rodríguez-Serrano et al., 2009, Kim et al., 2013). In M. polymorpha, SKL targeting 260 261 was utilized for evaluation of CRISPR-Cas9 modules (Konno et al., 2018). In transiently transformed M. 262 polymorpha cells, we also found a clear and distinct localization of mCherry-SKL in punctate structures, 263 likely representing peroxisomes (Fig. 6A).

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265 Actin filaments

Both the Lifeact peptide - a short peptide of 17 amino acids – and the C-terminal 197 amino acids of mouse talin are known to bind to filamentous actin (Kost et al., 1998; Riedl et al., 2008). Therefore, to visualize the actin filaments in Marchantia epidermal cells, we used the Citrine-mTalin and LifeAct-Citrine reporters described previously (Kimura and Kodama, 2016). As in stably transformed Marchantia lines (Kimura and Kodama, 2016), both markers successfully revealed the actin filament networks around chloroplasts in epidermal cells (Fig. 6B, C).

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273 Golgi apparatus

274 As potential markers for the Golgi apparatus, we selected the Arabidopsis proteins SYNTAXIN OF 275 PLANTS 3 (AtSYP3) and the GOLGI TRANSPORT 1 p homolog (AtGot1p). Both proteins have been shown 276 to localize to the Golgi apparatus (Uemura et al., 2004; Conchon et al., 1999) and are reliable Golgi 277 markers for Arabidopsis, as being part of the Wave line multicolor marker set for membrane 278 compartments (WAVE22 and WAVE18, respectively; Geldner et al., 2009). Upon transient biolistic 279 expression of YFP-AtSYP3 and YFP-AtGot1p in *M. polymorpha*, a distinct and comparable localization 280 pattern of both proteins, likely representing the Golgi apparatus, was visible (Fig. 7A and B). 281 Furthermore, upon co-expression of CFP-AtSYP3 and YFP-AtGOT1p, we also found perfect co-282 localization (Fig. 7C) confirming that both markers are reliable to illuminate the Golgi in Marchantia.

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284 mRNA processing bodies

285 mRNA processing bodies (p-bodies), have been found to play a crucial role in mRNA processing comprising deadenylation, decapping, degradation, mRNA storage and mRNA guality control 286 287 (thoroughly reviewed for A. thaliang in Maldonado-Bonilla, 2014). As p-bodies markers, we chose the Arabidopsis thaliana DECAPPING PROTEIN 1 (AtDCP1) and AtDCP2, whose function has been well 288 289 studied in the past years (Xu et al., 2006; Xu and Chua, 2009; Steffens et al., 2015; Bhasin et al., 2017). 290 Upon transformation of the protein fusion AtDCP1-mCherry we found a comparable expression in dot-291 like structures, likely representing p-bodies (Fig. 8A). In contrast, transformation of mCherry-AtDCP2 292 revealed a diffused expression throughout the cytoplasm and in the nucleus (Fig. 8B), as reported in 293 Arabidopsis in the absence of stress (Motomura et al., 2014). Therefore, we assume that AtDCP2 is 294 also generally localized in the cytoplasm and nucleus in M. polymorpha and is only recruited to pbodies upon stress conditions (Motomura et al., 2014). 295

The similar localization of AtDCP1/2 in Arabidopsis (Iwasaki et al., 2007; Motomura et al., 2014) and Marchantia suggests that the function of DCPs in mRNA processing has been evolutionarily conserved. To assess whether the two Marchantia DCP-homologs MpDCP1/2 localize similarly as their Arabidopsis counterparts, we transformed different combinations of fluorescent fusions (MpDCP1-mCherry, MpDCP1-YFP, MpDCP2-mCherry, MpDCP2-YFP). As anticipated, MpDCP1 displayed a dot-like localization pattern similar to AtDCP1, while MpDCP2 exhibited an AtDCP2-like nucleocytoplasmic localization (Fig. 8C and D).

303

304 B) Bimolecular fluorescence complementation

305 Based on former reports of AtDCP1 to regulate mRNA decay and to recruit further functionally relevant 306 proteins, such as AtDCP2, to p-bodies (Iwasaki et al., 2007; Motomura et al., 2014), as well as our own 307 observations (see above), we selected MpDCP1/2 as promising candidates to assess the feasibility of 308 studying protein-protein interactions in *M. polymorpha* via bimolecular fluorescence 309 complementation (BiFC). The BiFC technique relies on the co-expression of two proteins fused to the 310 N- or C-terminal part of a fluorescent reporter (e.g. -YFP_N and -YFP_C, respectively. Upon physical 311 interaction of the two tagged proteins of interest, the N- and C-terminal parts of the reporter can 312 reconstitute a functional fluorescent protein. Capture of the respective fluorescent signal thus is used 313 as an indicator for protein-protein interaction. For BiFC to be meaningful, the co-transformation of 314 both reporter halves must lead to regular and frequent co-expression, which is the case for Marchantia 315 thallus transient biolistic transformation as it reaches, in our hands, 74% on average (see Material and 316 Methods section and Tab. S3). The physical interaction of AtDCP1/2 was foremost reported in in vitro

pull-down assays (Xu et al., 2006) and later independently confirmed by BiFC in tobacco mesophyll
protoplasts (Weber et al., 2008).

319 Interestingly, upon co-expression of YFP_N-MpDCP2 and YFP_c-MpDCP1, together with the nuclear 320 marker AtKRP1-CFP, we could observe a clear and specific YFP signal in dot-like structures, suggesting 321 that MpDCP1/2 are capable of interacting physically in p-bodies of Marchantia epidermal cells (Fig. 322 9A). To exclude the possibility of false positive signals (Kodama et al., 2012) in our experimental setup 323 we also transformed YFP_N-MpDCP2 and YFP_c-MpDCP1 with YFP_c-MpLYST interacting protein 5 324 (MpLIP5) and AtMYC related protein1 (AtMYC1)-YFP_N tags, respectively. Expression of both vector 325 combinations led to the absence of a YFP signals in cells expressing AtKRP1-CFP (Fig. 9B and C), 326 indicating that the observed interaction between MpDCP1 and MpDCP2 is specific. The integrity of 327 YFP_C-MpLIP5 was confirmed by co-expression with the Marchantia homolog of a known interactor of 328 LIP5 in Arabidopsis - MpSuppressor of K⁺ Transport Growth Defect1 (MpSKD1) (Haas et al., 2007), N-329 terminally fused to YFP_N showing a clear YFP signal in punctate structures consistent with localization to p-bodies (Fig. S2A). The integrity of AtMYC1-YFP_N was shown by a BiFC interaction in the nucleus 330 331 with its known interaction partner AtTRANSPARENT TESTA GLABRA1 (TTG1) (Zimmermann et al., 2014, Zhao et al., 2012; control used in Steffens et al., 2017), C-terminally fused to YFPc (Fig. S2B). In 332 333 conclusion, our results show that BiFC is functional in Marchantia and can be used to quickly assess 334 protein-protein interactions in vivo.

335

336 C) Staining intracellular structures in *M. polymorpha*

For the visualization of a cell and the investigation of cellular architecture and dynamics, it is crucial to have several quick and reliable staining methods for live cell imaging at hand. Therefore, we tested some standard staining procedures to label intracellular compartments and cellular structures (including the plasma membrane, cytoplasm, cell wall and nucleus) in *M. polymorpha* gemmae that have been established for other plants but lacking ready-to-use protocols for Marchantia.

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343 Fluorescein diacetate (FDA) for cytoplasm staining of living cells

FDA is a cell-permeable, per se non-fluorescent esterase substrate. As soon as it passes the plasma membrane, it is hydrolyzed by esterases in the cytoplasm of viable cells (Rotman and Papermaster, 1966). Thereby, FDA is converted to a negatively charged, green-fluorescent fluorescein unable to either cross back the plasma membrane or pass the tonoplast and thus it accumulates in the cytoplasm. Owing to these properties, FDA is suitable for cell viability assays and can be used as a negative stain for vacuoles. In Arabidopsis, FDA staining has been reliably used for testing root hair and
guard cell viability (Schapire at al., 2008; Hao et al., 2012) ,to visualize vacuoles in root hairs (Saedler
et al., 2009) and trichomes (Mathur et al., 2003), and to study pathogen response (Jones et al., 2016).

Here, we successfully utilized FDA to stain the cytoplasm of rhizoids and epidermal cells in young gemmae (Fig. 10). FDA showed a strong, green fluorescence already after a short incubation time of 10 minutes, demonstrating the viability of rhizoids and epidermal cells. We here present FDA as a tool to be readily used for visualization of the cytoplasm in *M. polymorpha*. As it is not able to pass the tonoplast, it can also be used to detect vacuolar architecture, especially in rhizoids, where vacuolar volume was clearly visible after staining with FDA (Fig. 10D).

358

359 Propidium iodide for cell wall staining

Propidium iodide (PI) is an intercalating, red-fluorescent cell dye. It penetrates damaged cell membranes and visualizes nuclei of dead cells by intercalating DNA with low base preference. However, PI cannot pass intact cell membranes and thus is excluded from viable cells, while remaining fluorescent. Therefore, PI can also readily be used to visualize cell wall of living cells. In Arabidopsis, PI is regularly utilized for counterstaining of cell walls (Takano et al., 2002, Ubeda-Tomás, 2008), such as for viability assays, frequently combined with FDA (Shahriari et al., 2010, Kong et al., 2018).

We here show successful PI staining of cell walls of *M. polymorpha* (Fig. 11). Strong fluorescence was observed already after short incubation times of 10 min. PI reliably stained the cell walls of living epidermal cells (Fig. 11A) and rhizoids (Fig. 11B) and thus can reveal cell shape and size. This staining was non-toxic as stained rhizoids kept elongating, thereby revealing the usefulness of PI staining for studying rhizoid tip-growth (Video S1).

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372 Nuclei of M. polymorpha cannot be reliably stained with 4',6-diamidino-2-phenylindole (DAPI)

373 DAPI is one of the most common DNA fluorochromes enabling staining and visualization of nuclei of 374 dead but also viable cells, as it is able to pass cell membranes – however often with weak effectiveness. 375 Upon excitation with ultraviolet light, DAPI emits blue fluorescence at a maximum of 461 nm. DAPI 376 binds stoichiometrically to adenine-thymine rich regions of DNA. DAPI also has a weak binding capacity 377 to RNA, however emission is then shifted to 500 nm. Thus, DAPI is frequently utilized not only to 378 visualize nuclei in trichomes, epidermal pavement cells or root cells (Kirik et al., 2001, Spitzer et al., 379 2006, Lee et al., 2006), but also to quantify DNA content in Arabidopsis, being a reliable tool to discover 380 endoreduplication (Schnittger and Hülskamp, 2007, Bramsiepe et al., 2010, Bhosale et al., 2018).

Kondou et al., 2019, report a functional DAPI staining of nuclei in wholemount samples of fixed 381 382 epidermal cells of *M. polymorpha*. In this study, we tested staining of fixed (i.a. after a modified version 383 of the protocol by Kondou et al., 2019) but also of viable thallus epidermal cells of Marchantia. 384 Surprisingly, despite usage of gemmae at different developmental stages, short to long DAPI incubation periods, preceding and subsequent de-staining steps using EtOH, different methods of 385 fixation (for more details see Material and Methods section), we were unable to stain and visualize 386 387 nuclei of Marchantia with DAPI (Fig. S3). In our hands, DAPI accumulated on cell walls and to a weaker 388 extent in the cytoplasm but did not enter the nucleus. To demonstrate functionality of the used DAPI 389 solution, we stained Arabidopsis leaves in parallel (Fig. S3), showing strong and distinctive visualization 390 of nuclei. Staining of DNA by PI after fixation also failed in our hands (data not shown). It remains to 391 be elucidated, why nuclei of *M. polymorpha* seem to be hardly accessible to DNA fluorochromes. Until 392 then, we either suggest to use a protein marker localizing in nuclei (*e.g.* AtKRP1) and to generate stably 393 expressing Marchantia lines if needed; or to visualize S-phase nuclei with 5-Ethynyl-2'-deoxyuridine 394 (EdU) staining, as reports show its functionality in *M. polymorpha* (Furuya et al., 2018, Busch et al., 395 2019)

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397 FM4-64 staining for visualization of plasma membrane and endocytic vesicles

The lipophilic steryl dye FM4-64 (3-triethylammoniumpropyl)-4-(6-(4-(diethylamino)-phenyl)hexatrienyl) pyridinium-dibromide) is commonly used as marker for the outer leaf of the cellular plasma membrane. Staining of young gemmae with FM4-64 resulted in a clear fluorescence signal at the cellular boundaries, likely representing the plasma membrane (Fig. 12A). Upon co-staining with FDA, the FM4-64-specific plasma membrane signal at the cell periphery was clearly distinct from the cytoplasmic FDA signal (Fig. 12B). Altogether, these findings support FM4-64 as a reliable marker dye to label the outer cellular membrane via single or co-staining in Marchantia.

405

406 Concluding remarks

We here present a comprehensive and reliable toolkit for visualization of intracellular architecture and dynamics in *M. polymorpha*, an emerging model system used to study land plant evolution. All methods described are based on standard techniques used in other systems and can be executed and analyzed within 1 - 2 working days, therefore allowing time-efficient analysis of basic intracellular traits, such as organelle organization and cell architecture, both in fixed and viable cells. The possibility to mark viable cells additionally allows their analysis in live-imaging setups, as we demonstrate with growing rhizoids stained with PI. A comprehensive list of transiently expressed markers covering the

majority of intracellular organelles and structures, allows fast assessment of aforementioned 414 415 intracellular dynamics in viable cells, but also provides a quick possibility for initial tests of functionality 416 and correct localization of cloned fluorescent constructs before committing to comparatively timecostly stable plant transformation. Finally, we demonstrate the BiFC system to be functional in 417 418 Marchantia epidermal cells, thus representing a quick and straightforward technique to test for protein-protein interactions in vivo, which should be confirmed with other protein-protein interaction 419 420 assays such as Yeast-2-Hybrid-like, FRET-FLIM and protein pulldown approaches. Altogether, we 421 provide a series of quick and useful techniques to exploit the potential of an emerging model system 422 to the maximum extent possible.

423

424 Acknowledgements

We thank Dr. Marc Jakoby for providing aliquots of the AtKRP, SKL motif, AtNPSN12, AtSYP32 and 425 426 AtGot1p homolog vectors. We thank Dr. Alexandra Steffens for providing aliquots of the AtDCP1 and 427 AtDCP2 expression vectors. We thank Dr. Lisa Stephan for providing aliquots of the AtMYC1 and 428 AtTTG1 BiFC vectors. We thank Dr. Clement Champion and the research group of Prof. Liam Dolan 429 (University of Oxford) for provision of an aliquot of the PM marker vector MpSYP13a. We thank Dr. 430 Joachim F. Uhrig for donation of pCL112/113 vectors. This research was partly funded by a short-term 431 stipend of the Deutscher Akademischer Austauschdienst (DAAD) to J.W.; the University of Cologne, 432 and grant from the University of Cologne Centre of Excellence in Plant Sciences to A.B.-D.

433

434 Author contributions

- 435 J.W., E.K., M.H. and A.B.-D. conceived the experiments.
- 436 J.W., E.K., R.L. and A.B.-D. performed the experiments.
- 437 J.W., E.K. and A.B.-D analyzed the data.
- 438 J.W. and E.K. wrote the manuscript with contributions of M.H. and A.B.-D.

439

440 **Declaration of interests**

441 The authors declare no competing interests.

442

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726 **FIGURE LEGENDS**:

727

Fig. 1: Biolistic transformation of Marchantia thalli. (A) Plant material used for the transformation, showing 2.5 weeks old thalli grown on solid Johnson's medium. Scale bars = 2 cm (left) and 5 mm (right). (B) Representative overview of transformation efficiency; arrowheads pointing at transformed cells expressing MpMRI-YFP; Scale bar = 100 μ m. (C) Schematic transformation procedure: Vectorial DNA was coupled to gold particles (1), attached to a macro-carrier (2), biolistically transferred into thallus fragments (3), plants were allowed to rest overnight (4) and the pieces expressing the construct of interest were analysed under a microscope (5).

735

Fig. 2: Confirmation of known nuclear and plasma membrane markers. (A) The Arabidopsis nuclear
 marker AtKRP1 localizes to the nucleus of *M. polymorpha* epidermal cells. (B) The Arabidopsis plasma
 membrane marker AtNPSN12 localizes to the plasma membrane in Marchantia epidermal cells. (C, D)
 The Marchantia plasma membrane marker MpSYP13a co-localizes with AtNPSN12. All scale bars = 20
 µm. BF = bright field.

741

Fig. 3: Plasma membrane markers for Marchantia research. MpMRI (A), MpFER (B) and AtMRI (C) all
localized to the plasma membrane of *M. polymorpha* thallus epidermal cells. All three constructs colocalized with the plasma membrane marker AtNPSN12. All scale bars = 20 μm.

745

Fig. 4: Nucleocytoplasm markers for Marchantia research. Both, AtAUN1 and AtAUN2 localized to the
cytoplasm and nucleus of *M. polymorpha* thallus epidermal cells, consistent with observations in *A. thaliana* (Franck et al., 2018). The constructs were co-bombarded with plasma membrane marker
AtNPSN12. All scale bars = 20 μm. Pictures show maximum projections of z-stack captions, hence the
appearance of the 'cytoplasmic noise' signal for AtNPSN12-mcherry (see Materials and methods
section for details).

752

Fig. 5: Endosomal markers for Marchantia research. Both, MpRAB5 (A) and MpARA6 (B) localized to
 punctuate intracellular structures of *M. polymorpha* thallus epidermal cells, likely representing

endosomes. The constructs were co-bombarded with nuclear marker AtKRP1. The endosomal markers
MpRAB5 and MpARA6 also show clear co-localization (C). All scale bars = 20 µm. Pictures show
maximum projections of z-stack captions (see Materials and methods section for details).

758

Fig. 6: Peroxisomal and actin filaments markers for Marchantia research. (A) The SKL-target sequence
tagged to mCherry localized to single intracellular foci of *M. polymorpha* thallus epidermal cells, likely
representing peroxisomes. mCherry-SKL was co-bombarded with nuclear marker AtKRP1. The actin
filament markers (B) Citrine-mTalin and (C) LifeAct-Citrine were co-bombarded with plasma membrane
marker AtNPSN12-Mcherry. All scale bars = 20 µm. Pictures show maximum projections of z-stack
captions (see Materials and methods section for details).

765

Fig. 7 : Golgi markers for Marchantia research. The Arabidopsis golgi markers AtGot1p (A) and AtSYP32
(B) localize to the golgi apparatus of *M. polymorpha* epidermal cells. The constructs were cobombarded with nuclear marker *At*KRP1. (C) The golgi markers AtGot1P and AtSYP32 show clear colocalization. All scale bars = 20 µm. BF = bright field.

770

Fig. 8: Marchantia p-bodies markers. Both, AtDCP1 (A) and MpDCP1 (C) localized to intracellular dotlike structures, that likely represent p-bodies. In contrast, AtDCP2 (B) and MpDCP2 (D) localized to the cytoplasm, consistent with former observations (Motomura et al., 2014). Additionally, both, AtDCP2 and MpDCP2 showed a nuclear localization, co-localizing with the nuclear signal of AtKRP1. The constructs were co-bombarded with nuclear marker AtKRP1. Scale bar = 20 µm. Pictures show maximum projections of z-stack captions (see Materials and methods section for details).

777

Fig. 9: Bimolecular fluorescent complementation assays showing interaction between MpDCP1 and MpDCP2. (A): Co-transformation of split-YFP fusion constructs of MpDCP1 and MpDCP2 result in a fluorescence signal in dot-like foci, indicating protein-protein interaction in p-bodies. (B) Cobombardment of split-fusions with MpDCP2 and the unrelated MpLIP5 protein were used as a negative control. (C) We also co-bombarded split-versions of MpDCP1 and unrelated AtMYC1, which also led to the absence of any fluorescence signal. The constructs were co-bombarded with nuclear marker AtKRP1. Scale bar = 20 μm. Pictures show maximum projections of z-stack captions (see Materials and
 methods section for details). See also Fig. S2 for other controls.

786

Fig. 10: Fluorescein diacetate staining of different *M. polymorpha* cell types. (A) Whole-thallus staining, scale bar = 100 μ m, with close-up captures of (B) a distal thallus fragment, scale bar = 30 μ m, and (C) a meristematically active apical notch, scale bar = 30 μ m. All three images show localization of FDA to the cytoplasm, as contrasted by absence of FDA-specific fluorescence in the vacuole and autofluorescent (AF) chloroplasts. Pictures show maximum projections of z-stack captions (see Materials and methods section for details). (D) FDA staining of a Tak-1 rhizoid of a 5 days-old gemmaling. BF = bright field. Scale bar = 50 μ m.

794

Fig. 11. Propidium iodide staining of different *M. polymorpha* **cell types.** Propidium iodide (PI) staining of Tak-1 rhizoids of a 2 days old gemma, staining the cell wall of both thallus epidermal cells (A; Pictures show maximum projections of z-stack) and rhizoids (B). Scale bars = $100 \mu m$ (A) and $50 \mu m$ (B). BF = bright field. AF = Autofluorescence (detected at an emission of 680 - 700 nm).

799

Fig. 12. FM4-64 staining of *M. polymorpha* thallus epidermal cells. (A) FM4-64 staining of a 2 days old
 Tak-1 gemmaling, staining the plasma membrane of thallus epidermal cells. (B) Co-staining of FM4-64
 and FDA showing opposing plasma membrane- and cytoplasm-localized fluorescence signal. BF =
 bright field. FDA = Fluorescein-diacetate. All scale bars = 100 μm.

804

Fig. S1: Co-expression of MLRs and MRI with single or triple tags in Marchantia epidermal cells. (A)
 and (B) Arabidopsis MLRs fused to single fluorescent tag are not expressed. (C) and (D) the 3xCitrine
 tag leads MLRs to localize to the cytoplasm. (E) the 3xCitrine tag leads to normal cytosolic and plasma
 membrane localization of MpMRI. Pictures show maximum projections of z-stack captions (see
 Materials and methods section for details). Scale bar = 20 µm.

810

811 Fig. S2: Bimolecular fluorescent complementation assay quality controls. (A): The functionality of the 812 negative control MpLIP5 was confirmed via co-bombardment of split-versions of MpLIP5 and the 813 Marchantia homolog of the known Arabidopsis LIP5 interactor MpSKD1, showing a clear protein 814 interaction in dot-like foci. (B): Split-YFP fusion constructs of AtMYC1 and AtTTG1, known interactors, 815 were co-bombarded and shown to physically interact in M. polymorpha thallus epidermal cells, supporting the functionality of AtMYC1-YFP_N. The constructs were co-bombarded with nuclear marker 816 817 AtKRP1. Scale bar = 20 μ m. Pictures show maximum projections of z-stack captions (see Materials and 818 methods section for details).

819

- 820 Fig. S3: Nuclei of *M. polymorpha* cannot be readily stained with DAPI. (A) DAPI staining of Tak-1
- epidermal cells of a 4 days-old gemmaling. (B) DAPI staining of leaf epidermal cells of a 2 weeks old A.
- 822 *thaliana* plant. Note the stained nuclei. All scale bars = $50 \mu m$.
- 823
- 824 Video S1: Growing rhizoids stained with propidium iodide.

825

Tab. 1: Important Marchantia resources.

Resource/method	Link	Reference
Marchantia genome	http://marchantia.info/	Bowman et
sequence and		al., 2017
database		- , -
Marchantia entry on	https://phytozome.jgi.doe.gov/pz/portal.html#!info?alias=Org Mpolymorpha	Bowman et
Phytozome (including		al., 2017
BLAST and Genome		
Browser)		
Marchantia		Ohyama et al
chloroplast genome		1988,
studies		Umesono et
		al, 1988,
		Fukuzawa et
		al., 1988,
		Kohchi et al.,
Mawapp	http://monup.dkia/mu.mu	1988 Delmone et
MarpoDB : gene- centric database for	http://marpodb.io/query	Delmans et
Marchantia		al., 2016
polymorpha genetic		
parts for purposes of		
genetic engineering		
and synthetic biology		
PlantTFDB : Plant	http://planttfdb.cbi.pku.edu.cn/index.php?sp=Mpo	Jin et al., 201
Transcription Factor		Jin et al., 201
Database		Jin et al., 2014
recombination- mediated genome		2013
editing		
Stable Agrobacterium-		Kubota et al.,
mediated thallus transformation		2013
Stable Agrobacterium-		Ishizaki et al.,
mediated sporeling		2008
transformation		2000
Design of Gateway-		Ishizaki et al.,
compatible vectors for		2015 (A);
expression in		Mano et al.,
Marchantia		2018
CRISPR-Cas-based		Sugano et al.,
genome editing		2014,
		Sugano et al.,
CDISDD direct torget	https://crispr.dbcls.jp/	2018 Naito et al
CRISPRdirect target search		Naito et al., 2015
Stable biolistic thallus		Chiyoda et al
transformation		2008
Comprehensive		This study
catalogue of		,
fluorescent cell		
compartment markers		
Protein-protein		This study
interaction studies via BiFC		

CELLULAR STAINING TECHNIQUES	
FM4-64 staining of epidermal cells and rhizoids	Kato et al., 2017 (gemmae cups); This study (whole thallus
	and rhizoids)
FM1-43 staining of epidermal cells	Minamino et al., 2017
PI staining of thallus epidermal cells and rhizoids	Fixed cells: Buschmann et al., 2015; Rövekamp et al., 2016
	Living cells: Jones and Dolan, 2017; this study
DAPI staining of epidermal cells	Kondue et al., 2019; this study
FDA staining of thallus epidermal cells and rhizoids	This study
FURTHER RESOURCES	
Expressed sequence tags (EST) sequencing	Nagai et al, 1999; Nishiyama et al., 2000
RNA sequencing of the gametophyte transcriptome	Sharma et al., 2014
3D imaging using micro-computed tomography and mathematical image- processing method	Furuya et al., 2019
Guidelines for Marchantia gene nomenclature	Bowman et al., 2015

Tab. S1: Comprehensive list of all marker constructs used for biolistic transformation. The list includes their origin (referenced publication or own generation), as well as the oligonucleotide sequences used as primers for amplification of new marker gene CDS. GW = Gateway-compatible cassette.

Construct	Generation / origin	Primers used for amplification of marker gene CDS
proMpEF1α:: Lifeactin- Citrine	Kimura and Kodama, 2016	-
pro35S::AtAUN1-YFP	AtAUN1 CDS from Franck et al., 2018 remobilized into pro355::GW-YFP	-
pro35S::AtAUN2-YFP	AtAUN2 CDS from Franck et al., 2018 remobilized into pro355::GW-YFP	-
pro35S::AtDCP1-mCherry	Cloning of CDS in Steffens et al., 2015 remobilized into pAUBERGINE (M. Jakoby, GenBank ID: FR695418), kindly provided by A. Steffens	-
pro35S::AtKRP1-CFP	Cloning of CDS in Weinl et al., 2005 remobilized into pEXSG-CFP (Feys et al., 2005), kindly provided by M. Jakoby	-
pro35S::AtMRI-YFP	CDS from Boisson-Dernier et al., 2015 remobilized into pro355::GW-YFP	-
<i>pro35S::</i> AtMYC1-YFP _N	Cloning of CDS in Pesch et al., 2013, remobilized into pSPYNE (Walter et al., 2004), kindly provided by L. Stephan	-
<i>pro</i> 35S::AtNPSN12- mCherry	Cloning procedure as described in Steffens et al., 2014; CDS amplified with primers containing attB sites on pUNI clone U60291 recombined in pDONR207 and introduced into pAMARENA (M. Jakoby, GenBank ID: FR695418), kindly provided by M. Jakoby	-
pro35S::AtTTG1-YFP _C	Cloning of CDS in Bouyer et al., 2008, remobilized into pSPYCE (Walter et al., 2004), kindly provided by L. Stephan	-
pro35S::Citrine-mTalin	Kimura and Kodama, 2016	-
<i>pro35S::</i> mCherry-AtDCP2	Cloning of CDS in Steffens et al., 2015 remobilized into pAMARENA (M. Jakoby, GenBank ID: FR695418), kindly provided by A. Steffens	-
pro35S::mCherry-MpRAB5	this study; CDS introduced into pAMARENA (M. Jakoby, GenBank ID: FR695418)	GGGGACAAGTTTGTACAAAAAAGCAGGCTCAATGGCCACCGCGGGAACGAA; GGGGACCACTTTGTACAAGAAAGCTGGGTATCMGACGCAGCACATGCTTGATT
pro35S::MpARA6-YFP	this study; CDS introduced into pEXSG-YFP (Feys et al., 2005)	GGGGACAAGTTTGTACAAAAAAGCAGGCTCAATGGGTTGTGCTGCCTCAGC; GGGGACCACTTTGTACAAGAAAGCTGGGTATCMAGGCTTCTGGGTTGGCTGTC
<i>pro35S::</i> mCherry-SKL	SKL motif introduced into pAMARENA (M. Jakoby, GenBank ID: FR695418), kindly provided by M. Jakoby	-
pro35S::mCitrine- MpSYP13a	Kanazawa et al., 2015; received from Clement Champion, Prof. Liam Dolan Lab	-
pro35S::MpDCP1-YFP	this study; CDS introduced into pEXSG-YFP (Feys et al., 2005)	GGGGACAAGTTTGTACAAAAAAGCAGGCTCAATGGCACAAAATGGCAAGCCGATGC; GGGGACCACTTTGTACAAGAAAGCTGGGTATCMTGTTGAATGTGCATTGAGCATCTCC
pro35S::MpDCP2-YFP	this study; CDS introduced into pEXSG-YFP (Feys et al., 2005)	GGGGACAAGTTTGTACAAAAAAGCAGGCTCAATGTCCGGCAACGCGCGTGC; GGGGACCACTTTGTACAAGAAAGCTGGGTATCMGACTTCCAACTTTTGTATTATGCTT

pro35S::MpFER-YFP	this study; CDS containing GW-	GGGGACAAGTTTGTACAAAAAAGCAGGCTTAATGAGGCGTTCGTCTTGTTT;
	compatible attB1/2-sites, cloned	GGGGACCACTTTGTACAAGAAAGCTGGGTTTAACCTTCCTT
	into GW entry vector and	
	introduced into pro35S::GW-YFP	
pro35S::MpMRI-YFP	MpMRI CDS from Westermann et	_
	al., 2019 remobilized into	
	pro35S::GW-YFP	
pro35S::YFPc-MpDCP1	this study; CDS introduced into	GGGGACAAGTTTGTACAAAAAAGCAGGCTCAATGGCACAAAATGGCAAGCCGATGC;
p	pCL113 (donated by J.F. Uhrig,	GGGGACCACTTTGTACAAGAAAGCTGGGTATCMTGTTGAATGTGCATTGAGCATCTCC
	unpublished data)	
pro35S::YFP _C -MpLIP5	this study; CDS introduced into	GGGGACAAGTTTGTACAAAAAAGCAGGCTCAATGGGGGAGACTGCGGATCCGAAGA;
p	pCL113 (donated by J.F. Uhrig,	GGGGACCACTTTGTACAAGAAAGCTGGGTATCMGTGAGCTTGTGATGAAGAAGAGGGTC
	unpublished data)	
pro35S::YFP _N -MpDCP2	this study; CDS introduced into	GGGGACAAGTTTGTACAAAAAAGCAGGCTCAATGTCCGGCAACGCGCGTGC;
,	pCL112 (donated by J.F. Uhrig,	GGGGACCACTTTGTACAAGAAAGCTGGGTATCMGACTTCCAACTTTTGTATTATGCTT
	unpublished data)	
pro35S::YFP _N -MpSKD1	this study; CDS introduced into	GGGGACAAGTTTGTACAAAAAAGCAGGCTCAATGTACAGCAATTTCAAGGA
	pCL112 (donated by J.F. Uhrig,	GGGGACCACTTTGTACAAGAAAGCTGGGTATCMACCCTCCTCACCAAATTCAC
	unpublished data)	
pUBQ10::YFP-AtSYP32 /	Cloning procedure as described in	-
pro35S::CFP-AtSYP32	Steffens et al., 2014; CDS	
	amplified with primers containing	
	attB sites on pUNI clone U20852,	
	recombined in pDONR207 and	
	introduced into pENSG-CFP/YFP	
	(Feys et al., 2005), kindly	
	provided by M. Jakoby	
pUBQ10::YFP-AtGot1p	Cloning procedure as described in	-
homolog	Steffens et al., 2014; CDS	
	amplified with primers containing	
	attB sites on pUNI clone U63080	
	recombined in pDONR207 and	
	introduced into	
	pENSG-YFP (Feys et al., 2005),	
	kindly provided by M. Jakoby	

Tab. S2: Excitation and captured emission wavelengths used for analysis of fluorescent markers.

Fluorophore	Excitation	Captured Emission
CFP	458 nm	470 nm – 480 nm
YFP	514 nm	524 nm – 530 nm
mCherry	561 nm	607 nm – 618 nm
FDA	514 nm	500 nm – 540 nm
PI	561 nm	610 nm – 630 nm
DAPI	405 nm	450 nm – 470 nm

Tab. S3: Quantification of co-bombardment efficiency in *M. polymorpha* **biolistic transformation.** Data from 9 independent co-transformation events of marker proteins used in this study. Scans of at least two transformed cells were used for the quantification.

Cells expressing both marker proteins	Total number of transformed cells	Efficiency of co-transformation [%]
2	2	100
8	12	67
2	2	100
1	2	50
2	2	100
2	2	100
3	3	100
1	2	50
1	2	50
5	8	63
4	5	80
1	2	50
1	2	50
3	3	100
4	6	67
7	7	100
1	2	50
2	2	100
1	2	50
1	2	50
3	4	75
2	2	100
3	5	60
SUM: 60	SUM: 81	Average: 74
		SD: 23

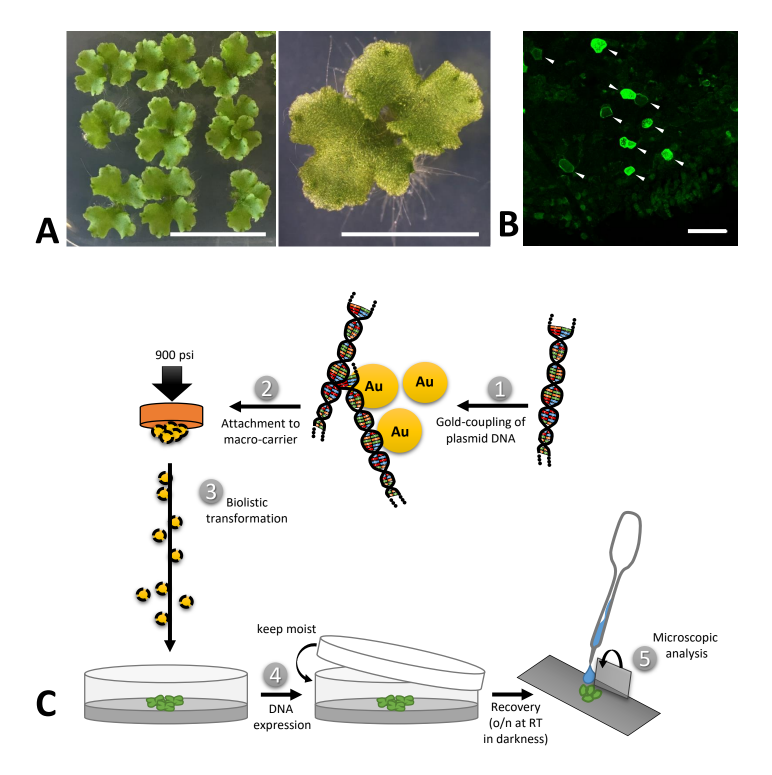


Fig. 1: Biolistic transformation of Marchantia thalli. (A) Plant material used for the transformation, showing 2.5 weeks old thalli grown on solid Johnson's medium. Scale bars = 2 cm (left) and 5 mm (right). (B) Representative overview of transformation efficiency; arrowheads pointing at transformed cells expressing MpMRI-YFP; Scale bar = 100 μ m. (C) Schematic transformation procedure: Vectorial DNA was coupled to gold particles (1), attached to a macro-carrier (2), biolistically transferred into thallus fragments (3), plants were allowed to rest overnight (4) and the pieces expressing the construct of interest were analysed under a microscope (5).

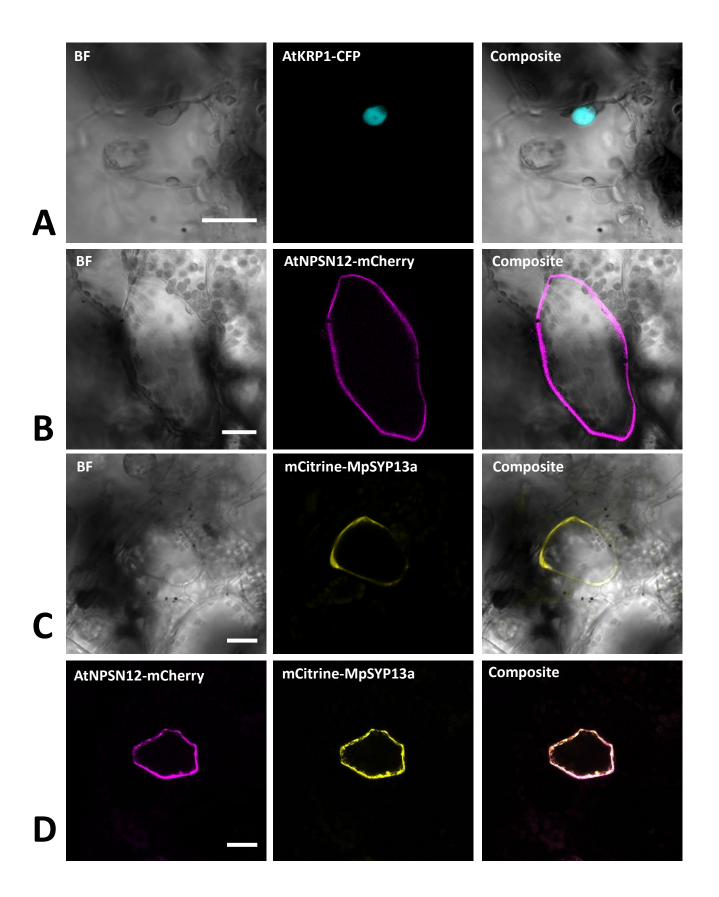


Fig. 2: Confirmation of known nuclear and plasma membrane markers. (A) The Arabidopsis nuclear marker AtKRP1 localizes to the nucleus of *M. polymorpha* epidermal cells. (B) The Arabidopsis plasma membrane marker AtNPSN12 localizes to the plasma membrane in Marchantia epidermal cells. (C, D) The Marchantia plasma membrane marker MpSYP13a co-localizes with AtNPSN12. All scale bars = $20 \ \mu$ m. BF = bright field.

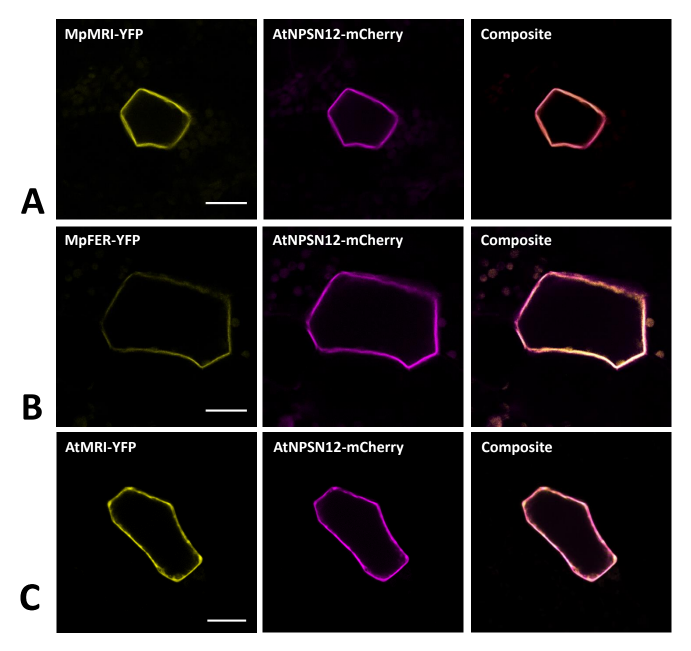


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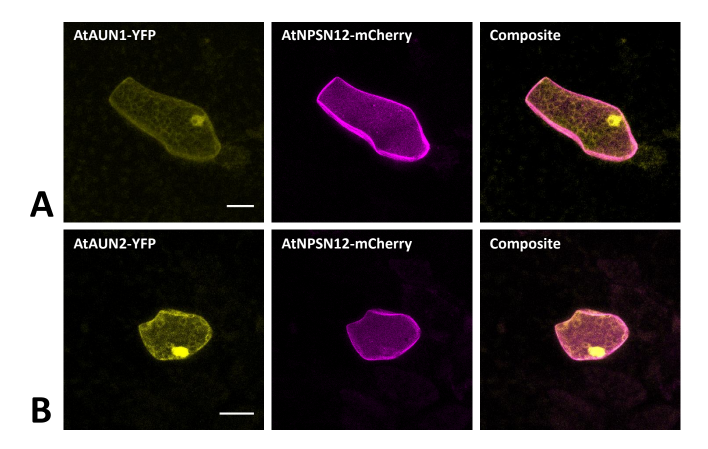


Fig. 4: Nucleocytoplasm markers for Marchantia research. Both, AtAUN1 and AtAUN2 localized to the cytoplasm and nucleus of *M. polymorpha* thallus epidermal cells, consistent with observations in *A. thaliana* (Franck et al., 2018). The constructs were co-bombarded with plasma membrane marker AtNPSN12. All scale bars = 20 µm. Pictures show maximum projections of z-stack captions, hence the appearance of the 'cytoplasmic noise' signal for AtNPSN12-mcherry (see Materials and methods section for details).

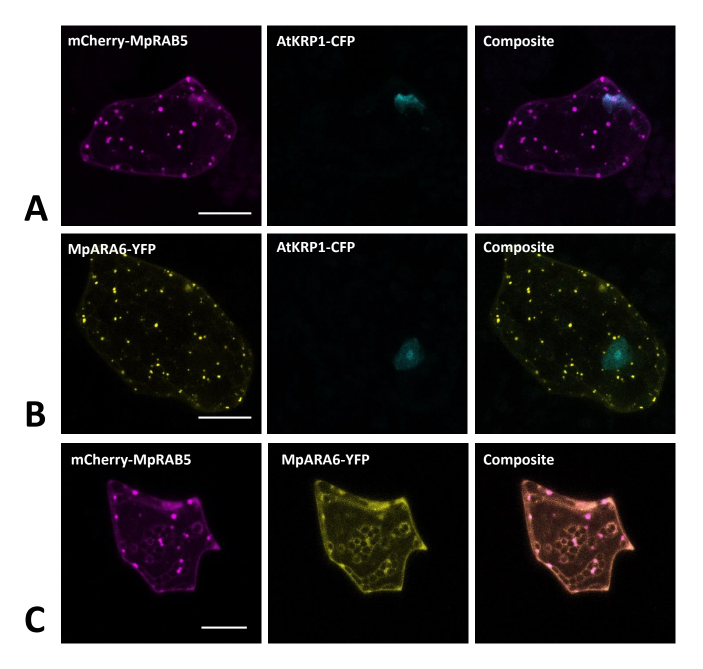


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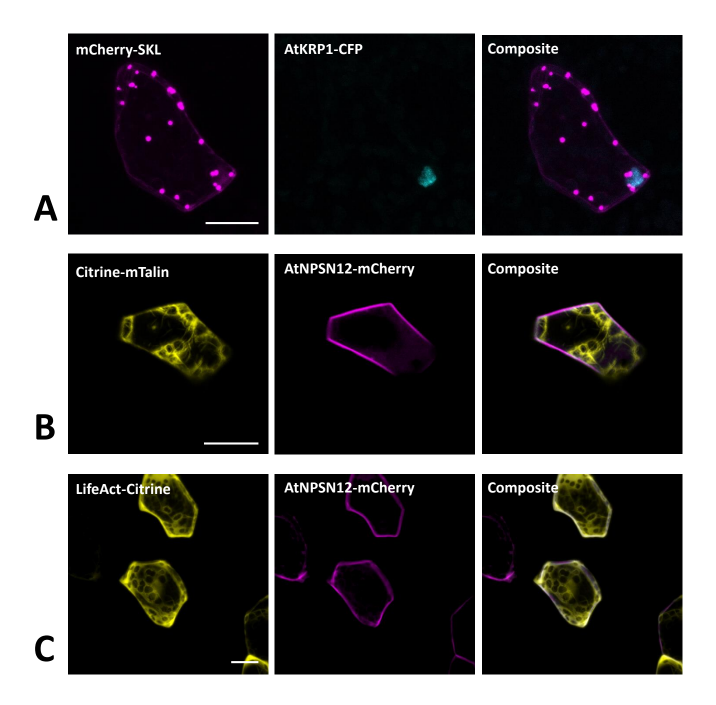


Fig. 6: Peroxisomal and actin filaments markers for Marchantia research. (A) The SKL-target sequence tagged to mCherry localized to single intracellular foci of *M. polymorpha* thallus epidermal cells, likely representing peroxisomes. mCherry-SKL was co-bombarded with nuclear marker AtKRP1. The actin filament markers (B) Citrine-mTalin and (C) LifeAct-Citrine were co-bombarded with plasma membrane marker AtNPSN12-Mcherry. All scale bars = 20 μm. Pictures show maximum projections of z-stack captions (see Materials and methods section for details).

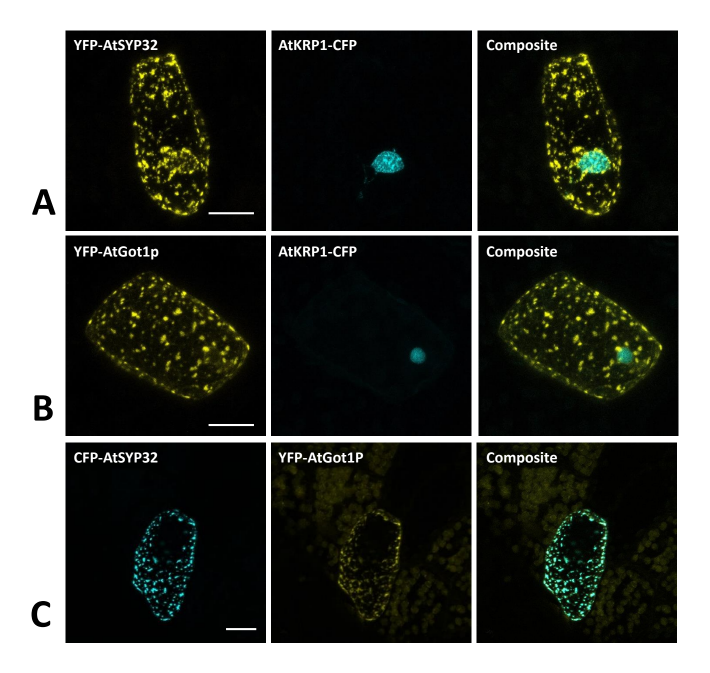


Fig. 7 : Golgi markers for Marchantia research. The Arabidopsis golgi markers AtGot1p (A) and AtSYP32 (B) localize to the golgi apparatus of *M. polymorpha* epidermal cells. The constructs were co-bombarded with nuclear marker *At*KRP1. (C) The golgi markers AtGot1P and AtSYP32 show clear co-localization. All scale bars = 20 μm. BF = bright field.

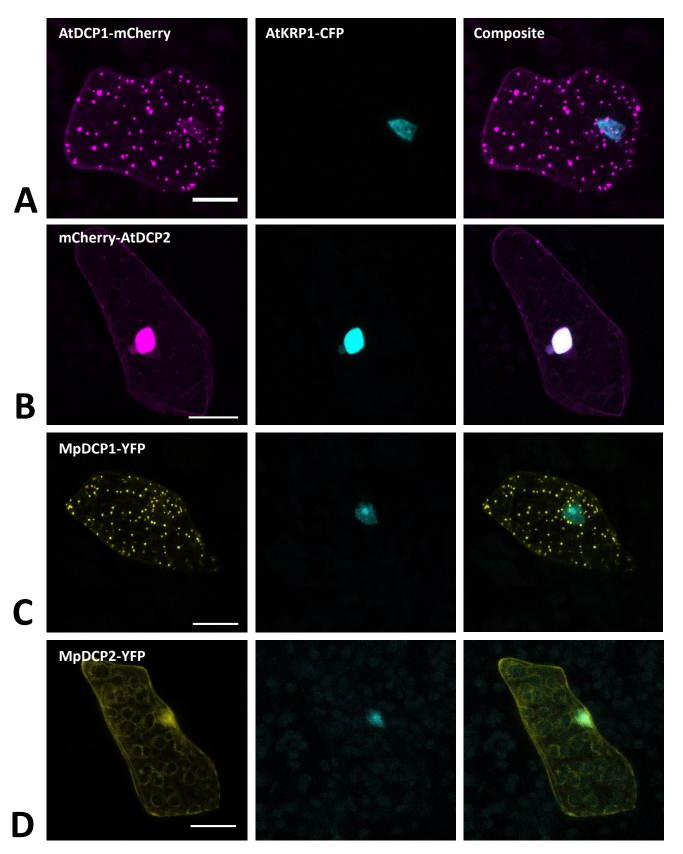


Fig. 8: Marchantia p-bodies markers. Both, AtDCP1 (A) and MpDCP1 (C) localized to intracellular dot-like structures, that likely represent p-bodies. In contrast, AtDCP2 (B) and MpDCP2 (D) localized to the cytoplasm, consistent with former observations (Motomura et al., 2014). Additionally, both, AtDCP2 and MpDCP2 showed a nuclear localization, co-localizing with the nuclear signal of AtKRP1. The constructs were co-bombarded with nuclear marker AtKRP1. Scale bar = 20 µm. Pictures show maximum projections of z-stack captions (see Materials and methods section for details).

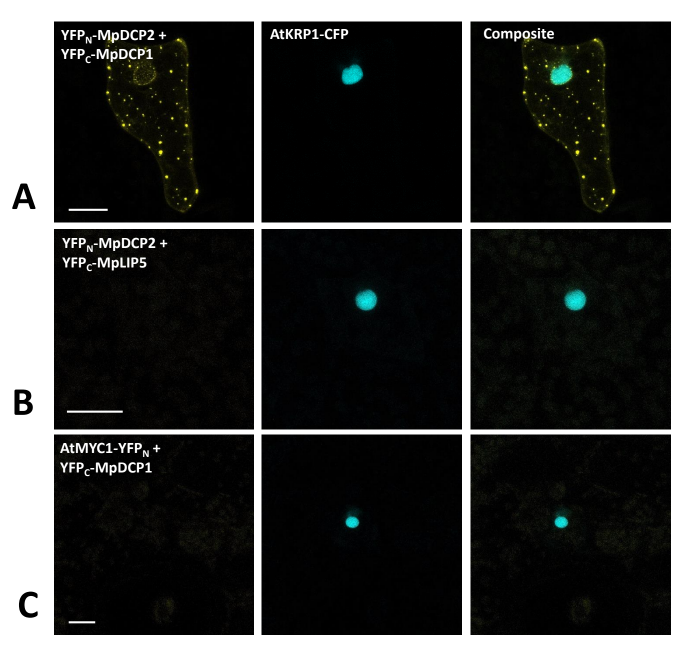


Fig. 9: Bimolecular fluorescent complementation assays showing interaction between MpDCP1 and MpDCP2. (A): Cotransformation of split-YFP fusion constructs of MpDCP1 and MpDCP2 result in a fluorescence signal in dot-like foci, indicating protein-protein interaction in p-bodies. (B) Co-bombardment of split-fusions with MpDCP2 and the unrelated MpLIP5 protein were used as a negative control. (C) We also co-bombarded split-versions of MpDCP1 and unrelated AtMYC1, which also led to the absence of any fluorescence signal. The constructs were co-bombarded with nuclear marker AtKRP1. Scale bar = 20 µm. Pictures show maximum projections of z-stack captions (see Materials and methods section for details). See also Fig. S2 for other controls.

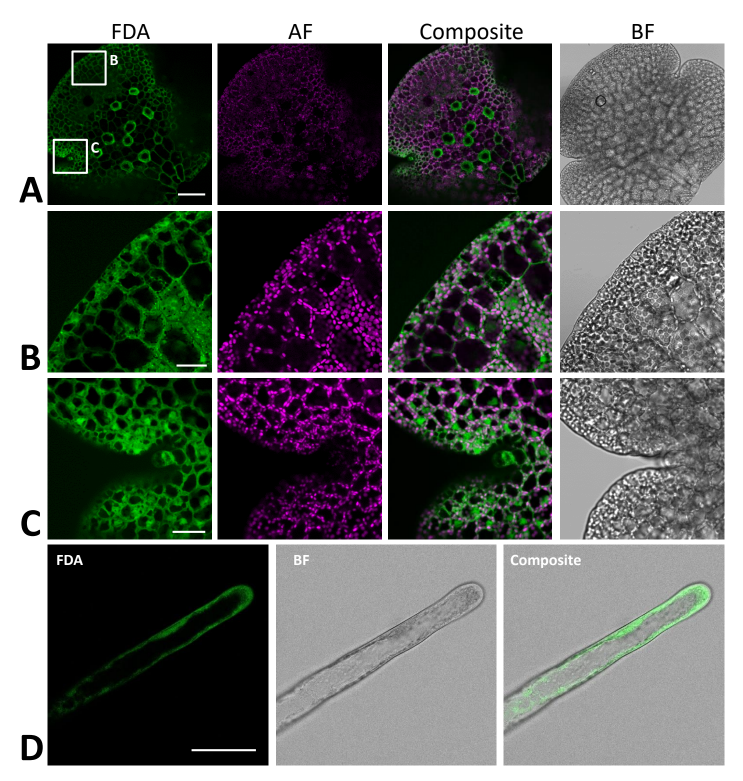


Fig. 10: Fluorescein diacetate staining of different *M. polymorpha* cell types. (A) Whole-thallus staining, scale bar = 100 μ m, with close-up captures of (B) a distal thallus fragment, scale bar = 30 μ m, and (C) a meristematically active apical notch, scale bar = 30 μ m. All three images show localization of FDA to the cytoplasm, as contrasted by absence of FDA-specific fluorescence in the vacuole and autofluorescent (AF) chloroplasts. Pictures show maximum projections of z-stack captions (see Materials and methods section for details). (D) FDA staining of a Tak-1 rhizoid of a 5 days-old gemmaling. BF = bright field. Scale bar = 50 μ m.

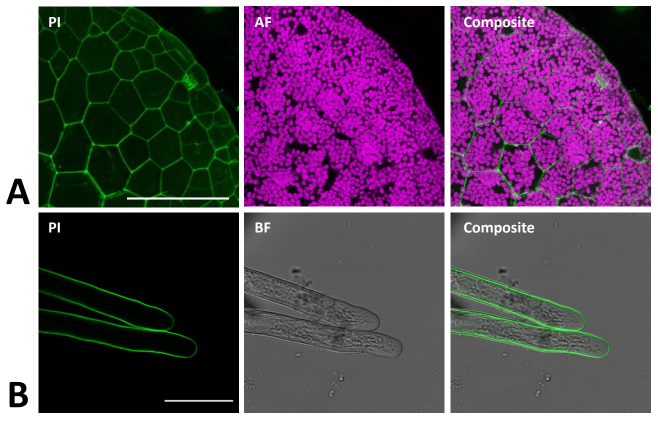


Fig. 11. Propidium iodide staining of different *M. polymorpha* **cell types.** Propidium iodide (PI) staining of Tak-1 rhizoids of a 2 days old gemma, staining the cell wall of both thallus epidermal cells (A; Pictures show maximum projections of z-stack) and rhizoids (B). Scale bars = 100 μ m (A) and 50 μ m (B). BF = bright field. AF = Autofluorescence (detected at an emission of 680 – 700 nm).

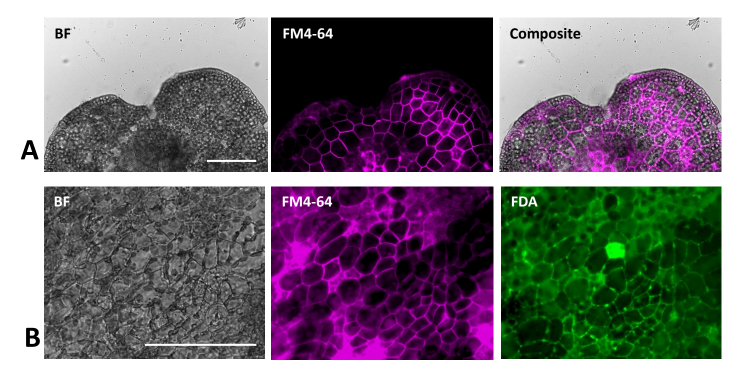


Fig. 12. FM4-64 staining of *M. polymorpha* **thallus epidermal cells.** (A) FM4-64 staining of a 2 days old Tak-1 gemmaling, staining the plasma membrane of thallus epidermal cells. (B) Co-staining of FM4-64 and FDA showing opposing plasma membrane- and cytoplasm-localized fluorescence signal. BF = bright field. FDA = Fluorescein-diacetate. All scale bars = 100 μ m.

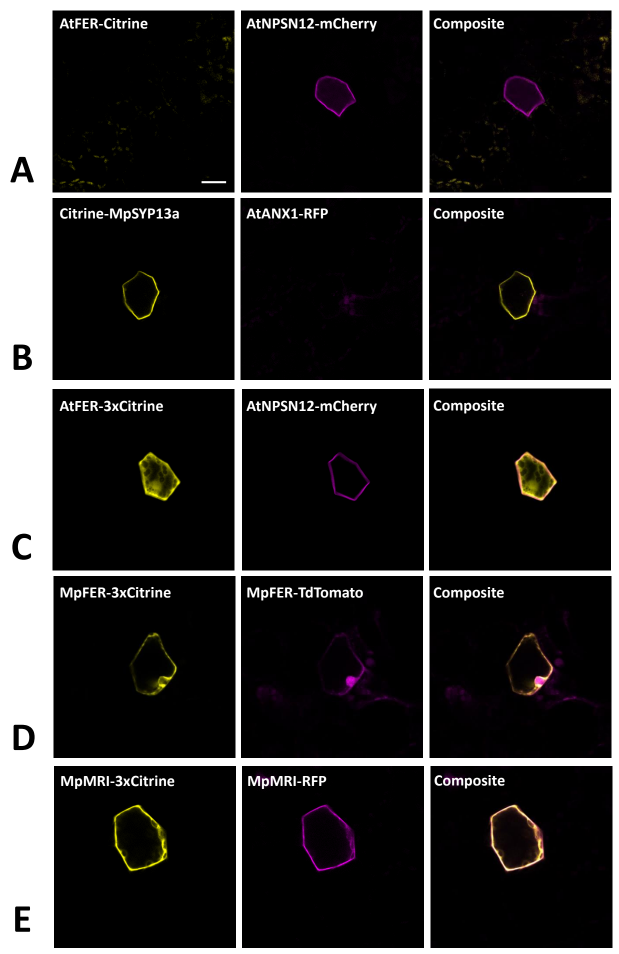


Fig. S1: Co-expression of MLRs and MRI with single or triple tags in Marchantia epidermal cells. (A) and (B) Arabidopsis MLRs fused to single fluorescent tag are not expressed. (C) and (D) the 3xCitrine tag leads MLRs to localize to the cytoplasm. (E) the 3xCitrine tag leads to normal cytosolic and plasma membrane localization of MpMRI. Pictures show maximum projections of z-stack captions (see Materials and methods section for details). Scale bar = 20 μ m.

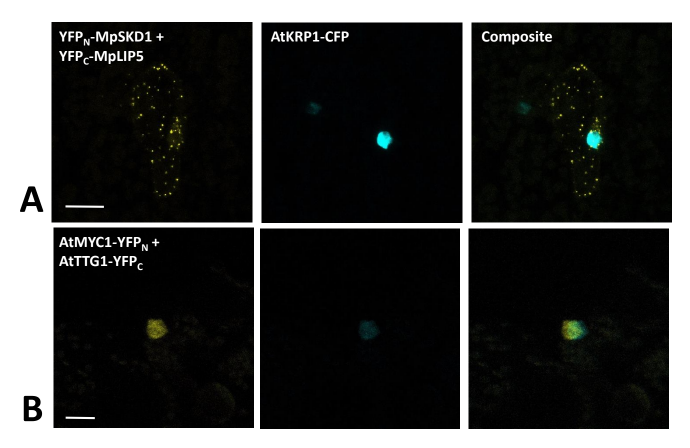


Fig. S2: Bimolecular fluorescent complementation assay quality controls. (A): The functionality of the negative control MpLIP5 was confirmed via co-bombardment of split-versions of MpLIP5 and the Marchantia homolog of the known Arabidopsis LIP5 interactor MpSKD1, showing a clear protein interaction in dot-like foci. (B): Split-YFP fusion constructs of AtMYC1 and AtTTG1, known interactors, were co-bombarded and shown to physically interact in *M. polymorpha* thallus epidermal cells, supporting the functionality of AtMYC1-YFP_N. The constructs were co-bombarded with nuclear marker AtKRP1. Scale bar = 20 μ m. Pictures show maximum projections of z-stack captions (see Materials and methods section for details).

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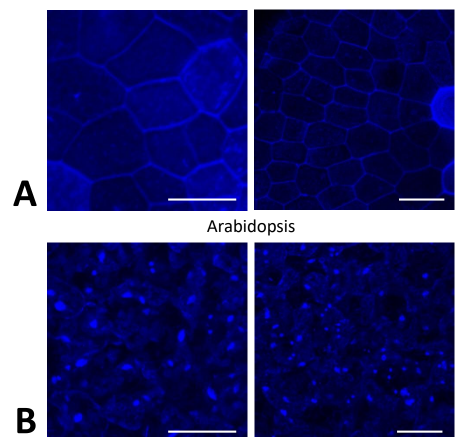


Fig. S3: Nuclei of *M. polymorpha* cannot be readily stained with DAPI. (A) DAPI staining of Tak-1 epidermal cells of a 4 days-old gemmaling. (B) DAPI staining of leaf epidermal cells of a 2 weeks old *A. thaliana* plant. Note the stained nuclei. All scale bars = $50 \mu m$.