Friendly regulates membrane depolarization induced mitophagy in Arabidopsis

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Abstract

The oxidative environment within the mitochondria makes them particularly vulnerable to proteotoxic stress. To maintain a healthy mitochondrial network, eukaryotes have evolved multitiered quality control pathways. If the stress cannot be alleviated, defective mitochondria are selectively removed by autophagy via a process termed mitophagy. Despite significant advances in metazoans and yeast, in plants, the molecular underpinnings of mitophagy are largely unknown. Here, using time-lapse imaging, electron tomography and biochemical assays, we show that uncoupler treatments cause loss of mitochondrial membrane potential and induce autophagy in Arabidopsis. The damaged mitochondria are selectively engulfed by autophagosomes that are ATG5 dependent and labelled by ATG8 proteins. Friendly, a member of the Clustered Mitochondria protein family, is recruited to the damaged mitochondria to mediate mitophagy. In addition to stress, mitophagy is also induced during de-etiolation, a major cellular transformation during photomorphogenesis that involves chloroplast biogenesis. De-etiolation triggered mitophagy regulates cotyledon greening, pointing towards an inter-organellar cross-talk mechanism. Altogether our results demonstrate how plants employ mitophagy to recycle damaged mitochondria during stress and development.

1 Introduction

2 Mitochondria are highly dynamic double-membraned organelles that function as cellular 3 powerhouses. They generate energy via oxidative phosphorylation (OXPHOS) and mediate the synthesis of essential macromolecules such as iron-sulfur clusters^{1,2}. One of the by-4 5 products of the oxidative environment in mitochondria is generation of toxic reactive oxygen 6 species that damage mitochondrial DNA, lipids and proteins. In addition, although most of the 7 mitochondrial proteins are encoded by nuclear genes, 13 subunits of the oxidative 8 phosphorylation complexes are still encoded by the mitochondrial genome. As the inter-9 genome coordination could be disrupted, and one cell could have thousands of times more 10 copies of mitochondrial genome than the nuclear genome; imbalances in stoichiometries of these multi-subunit OXPHOS complexes trigger proteotoxic stress^{3,4}. To overcome these 11 challenges and maintain a healthy mitochondrial network, eukaryotes have evolved multi-12 13 tiered and interconnected mitochondrial quality control pathways³.

14

15 One of the major mitochondrial quality control pathways is mitophagy, the selective 16 removal of damaged or superfluous mitochondria via autophagy. As many players involved in 17 mitophagy have been associated with disease, and mitophagy allows us to visualize selective 18 engulfment of an organelle into an autophagosome, mitophagy is one of the best studied 19 signalling mechanisms in metazoans⁵⁻⁸. One of the hallmarks of damaged mitochondria is loss 20 of membrane potential³. Various chemical protonophores such as carbonyl cyanide p-21 trifluoro-methoxyphenyl hydrazone (FCCP) or 2,4-dinitrophenol (DNP) have been used to 22 induce mitochondrial membrane depolarization and mitophagy⁹. Loss of mitochondrial 23 membrane potential leads to the stabilization of PINK1 on mitochondrial outer membrane. 24 PINKI phosphorylates ubiquitin and activates Parkin on mitochondrial membrane for 25 polyubiquitination of various outer membrane proteins. This creates a positive feedback loop 26 that results in recruitment of various selective autophagy receptors such as p62, Optineurin 27 or NDP52 to recruit the damaged mitochondria into autophagosomes for their subsequent degradation^{7,10,11}. Although much has been learnt about mitophagy in metazoans, molecular 28 29 players that mediate mitophagy in plants is currently unknown 2,12 .

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Mounting evidence suggests plant mitochondria are also recycled by selective autophagy^{2,13}. However, likely influenced by harbouring another endosymbiotic organelle, plants lack homologs of known mitophagy receptors and regulators¹². Also, so far, most 34 studies used genetic and biochemical assays to analyse mitochondrial turnover in plants. Cell 35 biological tools that would allow us to visualize different stages of mitophagy have not been 36 established. Here, we studied uncoupler induced mitophagy in the model plant Arabidopsis 37 thaliana. We used live cell imaging and electron tomography to visualize the engulfment of the 38 damaged mitochondria by mitophagosomes. We supported our cell biological findings with 39 autophagic flux assays to show autophagy regulates recycling of damaged mitochondria. We 40 also showed that Friendly (FMT) protein that has been linked to the regulation of 41 mitochondria dynamics is recruited to mitochondria upon damage. Consistently, fmt mutants 42 have defects in formation of mitophagosomes and mitochondrial turnover. Finally, we 43 demonstrate that de-etiolation also leads to accumulation of compromised mitochondria and 44 induces mitophagy. Altogether, our findings establish a cell biological and biochemical platform 45 to further dissect mitophagy and reveal a molecular player that is essential for mitophagy in 46 plants.

47

48 **Results**

49 Uncoupler treatments induce accumulation of depolarized mitochondria in 50 Arabidopsis root cells.

51 Uncouplers such as DNP and FCCP perturb the electrochemical potential of inner 52 mitochondrial membrane, triggering mitochondrial recycling in mammalian cells¹⁴. To test 53 whether these compounds also affect mitochondria in plant root tip cells, we incubated 54 Arabidopsis seedlings expressing mitochondrion-targeted GFP (Mito-GFP) in liquid MS medium 55 containing DNP or FCCP (Fig. 1). For consistency, our live cell imaging and electron 56 microscopy/tomography of mitochondria were limited to cortex cells in the root elongation 57 zone. However, loss of membrane potential and mitochondrial recycling were observed in all 58 cell types. To differentiate depolarized mitochondria, we pre-stained root cells with 59 tetramethylrhodamine ethyl ester (TMRE), a fluorescent dye sensitive to membrane potential¹⁵. 60 Normal mitochondria are seen as yellow puncta from dual fluorescence emitted from GFP and 61 TMRE, while depolarized mitochondria will be green as TMRE will not fluoresce upon 62 membrane depolarization.

63 Under normal conditions, depolarized mitochondria were rare in Mito-GFP roots (Fig. 64 Ia). When Mito-GFP roots were incubated with DNP (50 μ M) for I hr, numbers of 65 depolarized mitochondria increased significantly (Fig. Ia,c). Inactivation of a core 66 macroautophagy gene, ATG5, in the Mito-GFP line (*atg5-1*::Mito-GFP) led to accumulation of 67 more depolarized mitochondria in DNP-treated as well as untreated roots, indicating that 68 removal of depolarized mitochondria requires ATG5 (Fig. 1b,d). Addition of Concanamycin A 69 (ConA), an inhibitor of vacuolar H^+ -ATPase that disrupts protein transport to the vacuole¹⁶, 70 led to further build-up of mitochondria lacking membrane potential in Mito-GFP roots, 71 indicating vacuole is the final destination for these depolarized mitochondria. Importantly, 72 ConA did not lead to a similar build-up of depolarized mitochondria in atg5-1::Mito-GFP roots 73 (Fig. 1d, Extended Data Fig. 1). These observations agree with the inhibition of autophagy by 74 ConA^{17,18} and suggest that depolarized mitochondria are recycled via the macroautophagy 75 machinery in Arabidopsis root cells.

FCCP was a more potent uncoupler than DNP, depolarizing almost all mitochondria at a lower concentration after I hr (Fig. I). In the following analyses, however, we employed DNP to trigger mitophagy, because its slower action facilitated the monitoring of the mitophagy dynamics via cell biological and biochemical assays.

80

81 Uncoupler treatments induce autophagy

82 To examine whether autophagosome formation is induced following the uncoupler stress, 83 we visualized a member of the Arabidopsis ATG8 family, ATG8e, fused with a YFP (YFP-84 ATG8e). ATG8 is widely used as a marker for autophagosomes in all eukaryotes^{19,20}. Under 85 normal conditions we rarely observed puncta; most of the ATG8 signal was diffuse. However, 86 The YFP-ATG8e foci multiplied in root cells after I hr of DNP treatment, and their numbers 87 continued to increase at later time points (Fig. 2a,b). Upon activation of autophagy, ATG8 88 becomes conjugated to phosphatidylethanolamine (PE) by a complex containing ATG5, and 89 affixed to the limiting membrane of autophagosomes¹⁷. The lipidated form of ATG8 runs faster 90 in western blots, and the ratio between lipidated and unlipidated ATG8 is used as a proxy to 91 measure autophagy¹⁶. Immunoblot analyses with ATG8 antibody revealed only a faint upper 92 band in untreated wild type (WT) cells (Fig. 2c). This band became more abundant by DNP 93 treatment, especially in the membrane fraction. Samples incubated with phospholipase D 94 (PLD) or samples from *atg5-1* mutants lacked the upper band (Fig. 2d). Altogether, these 95 results suggest that ATG8 lipidation is induced upon DNP treatment in an ATG5 dependent 96 manner.

To further confirm that the uncoupler stress induces autophagosome formation and vacuolar delivery, we performed GFP cleavage assay with the YFP-ATG8e line and *atg5-1* mutant line expressing an mCherry-ATG8e chimeric protein (mCherry-ATG8e::*atg5-1*). A 100 free YFP polypeptide was detected in YFP-ATG8e samples and its amount increased over 101 time with a concomitant drop in YFP-ATG8e (Fig. 2e,f). No free mCherry was discerned in 102 the immunoblot of mCherry-ATG8e::*atg5-1* by an anti-mCherry antibody (Fig. 2e). Excitingly, 103 DNP treatment did not affect the autophagic flux of an aggrephagy receptor, Neighbor of 104 BRCAI (NBRI)²¹ (Fig. 2e). These data suggest that uncoupler treatment activates a selective 105 autophagy pathway to recycle depolarized mitochondria.

106

107 Damaged mitochondria are selectively engulfed by autophagosomes in uncoupler 108 treated root cells.

109 We then examined if DNP induced autophagosomes were indeed engulfing mitochondria 110 by staining YFP-ATG8e root cells with MitoTracker Red (MTR). Under a confocal microscope, 111 YFP-positive puncta were seen in the vicinity of mitochondria (Fig. 3a). In higher magnification 112 micrographs, we were able to identify ATG8e-specific fluorescent structures resembling open 113 pouches that contain mitochondria (arrowheads, Fig. 3b). We were also able to observe 114 mitochondria that were entirely surrounded by YFP-ATG8e rings (white arrow, Fig. 3b). In 115 time-lapse live cell imaging, YFP-ATG8e pockets partially enclosing a mitochondrion were 116 seen to grow, eventually encapsulating the mitochondrion over 300 seconds (Fig. 3c). These 117 autophagic compartments (i.e., mitophagosomes) were approximately 1-2 μ m in diameters 118 and each carried a single mitochondrion. MTR stains depolarized mitochondria better than 119 TMRE but does not concentrate in mitochondria with no membrane potential²². In this vein, 120 empty autophagosomes matching the size of mitophagosomes (grey arrow in Fig. 3b) in our 121 micrographs could correspond to mitophagosomes carrying mitochondrial corpses.

122 A time-lapse movie revealed that a small YFP-ATG8e puncta arose near MTR stained 123 mitochondria and elongated to be a semicircle capturing a mitochondrion. This process took 124 about 10 mins (Extended Data Fig. 2a). Another video documented an incomplete 125 mitophagosome that expanded to enclose a mitochondrion fully (Extended Data Fig. 2b). 126 Elongating tips of phagophores stayed in contact with the mitochondrial surface throughout 127 their growth (Extended Data Fig. 3a). From our live cell microscopy data, we estimated that 128 it takes about 15 min for mature mitophagosomes to develop from an initial YFP-ATG8 spot 129 on a mitochondrion.

We then cryofixed root samples and performed transmission electron microscopy (TEM) analysis. Mitochondria in control samples had smooth cristae that are evenly dispersed in the matrix (Fig. 3d,g). By contrast, mitochondria in DNP treated cells had electron-dense

133 precipitates in their matrix, some of which were engulfed by double membraned 134 mitophagosomes (Fig. 3 e-k). Serial section TEM of mitophagosomes showed that one 135 mitochondrion was contained per autophagosome, and no other organelles were identified in 136 autophagosomes (Extended Data Fig. 3b,c), in agreement with the live cell imaging results (Fig. 137 3b,c). Electron tomography analysis revealed that mitochondria sequestered in 138 mitophagosomes have more dark precipitate but less cristae in the matrix than free 139 mitochondria (Fig. 3g-k). Arabidopsis atg5-1 root cells had many mitochondria exhibiting the 140 signs of the internal precipitates, but they were not associated with mitophagosomes (Fig. 3I). 141 Altogether these results show that plant cells selectively recycle damaged mitochondria via 142 autophagy that involves ATG5.

143 To further investigate how mitochondria are degraded during uncoupler induced 144 mitophagy, we used immunoblot assays to assess the levels of various mitochondrial proteins 145 in Arabidopsis WT and atg5-1 mutant lines. We treated Arabidopsis seedlings with DNP for one 146 to four hours (DI-D4). To distinguish between uncoupler induced mitophagy and bulk 147 autophagy, we used nitrogen starvation as control (Extended Data Fig. 4). In contrast to the 148 endoplasmic reticulum protein Cycloartenol-C24-methyl transferase (SMTI), levels of outer 149 mitochondrial membrane (OMM) proteins peripheral-type benzodiazepine receptor (PBR) and voltage dependent anion channel I (VDACI); inner mitochondria membrane (IMM) 150 151 proteins cytochrome oxidase subunit II (COXII) and L-galactono-1,4-lactone dehydrogenase 152 (GLDH), and mitochondria matrix (MM) protein isocitrate dehydrogenase (IDH) were all 153 reduced upon uncoupler treatment (Fig. 4a). The reduction in protein levels were due to 154 vacuolar degradation as addition of ConA prevented the degradation (Fig. 4b). Furthermore, 155 the degradation was dependent on ATG5, since protein levels did not change significantly in 156 DNP treated atg5-1 mutant specimens (Fig. 4a,c). Interestingly, OMM proteins showed higher 157 levels of degradation in contrast to IMM and matrix proteins. Since, previous studies in mammalian cells showed OMMs could also be degraded via the proteasome²³, we checked 158 159 OMM protein levels of DNP treated cells following proteasome inhibition by MG132. Adding 160 MGI32 indeed stabilized only OMM proteins but not IMM or matrix proteins (Fig. 4d). 161 Considered together with normalized quantification of protein levels (Fig. 4e,f), these 162 experiments suggest upon loss of membrane potential, proteasome and autophagy cooperate 163 to degrade OMM proteins, whereas IMM and matrix proteins are primarily degraded by 164 autophagy.

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166 *Friendly* is essential for uncoupler induced mitophagy.

167 We then wanted to identify molecular players that mediate mitophagy in plants. Previous 168 studies have shown that for both ubiquitin dependent and independent mitophagy pathways, mitochondrial network needs to be fragmented^{24,25}. In Arabidopsis, Friendly (FMT), a clustered 169 170 mitochondria (CLU) family protein, has been shown to play important roles in mitochondrial 171 dynamics²⁶. However, whether it plays a role in mitophagy wasn't addressed. YFP-FMT 172 exhibited a diffuse cytosolic pattern under normal conditions. Interestingly, upon DNP 173 treatment, YFP-FMT localized to puncta that colocalized with mitochondria (Fig. 5a). Pull 174 down experiments showed that uncoupler treatment led to specific association of YFP-FMT 175 with ATG8, suggesting that YFP-FMT localizes to mitophagosomes (Fig. 5b). This prompted 176 us to analyse mitophagy in *fmt* mutant. Consistent with our previous findings presented in Fig. 177 3, live cell imaging of GFP-tagged mitochondria in mCherry-ATG8e expressing WT cells 178 showed ATG8e labelled vesicles engulfed mitochondria upon uncoupler treatment. However, 179 in fmt mutant, although ATG8 puncta were associated with the mitochondria, we did not 180 observe engulfment of mitochondria within autophagosomes (Fig. 5c). Further analyses of 181 mitophagosomes in the *fmt* mutant using TEM revealed aberrant mitophagosomes with 182 disconnected edges (Fig. 5c).

183 To further test the role of FMT in mitophagy, we performed live cell imaging and 184 immunoblot based autophagic flux experiments. Staining of depolarized mitochondria with 185 TMRE upon uncoupler treatments revealed accumulation of damaged mitochondria in the 186 cytosol of fmt mutants (Fig. 6a). Quantification of depolarized mitochondria showed fmt 187 mutants accumulated significantly more mitochondria in contrast to WT cells (Fig. 6b). 188 Furthermore, morphometric analyses of mitochondria in TEM micrographs from WT, fmt and 189 atg5-1 mutants showed that mitochondria were significantly larger in fmt and atg5-1 mutants 190 (Extended Data Fig. 5). Finally, immunoblot analyses using mitochondrial compartment specific 191 antibodies also showed a delay of mitochondrial protein degradation in *fmt* mutant (Fig. 6c). 192 Comparative analyses of the polypeptide intensities indicated that although *fmt* mutant had a 193 significant defect in mitochondrial protein recycling (Fig. 6d), it was not as severe as the atg5-194 I mutant (Fig. 4d,e), suggesting, in the absence of FMT, compensatory pathways prevent 195 accumulation of the toxic damaged mitochondria in the cell. Altogether these experiments 196 suggest FMT is required for mitophagy in plants.

198 Cotyledon greening during de-etiolation is affected in *atg5-1* mutant seedlings.

199 In germinating seeds, under darkness, proplastids transform into etioplasts that are 200 characterized by paracrystalline arrays of prolamellar bodies in their stroma²⁷. The etioplast 201 quickly transforms into the chloroplast upon exposure to light, a major developmental transition 202 termed de-etiolation. Since mitochondrial and chloroplast functions are tightly interconnected, 203 we hypothesized that mitochondrial population may also undergo remodelling during de-204 etiolation. First, we monitored greening of cotyledons when dark grown seedlings were 205 exposed to light. Green pigment levels increased gradually over a 12 hr period, with a significant 206 rise at eight hours after illumination (Fig. 7a,b). We then measured amounts of mitochondrial 207 proteins in greening cotyledon cells and showed that their levels also drop at 8 hr time point, 208 suggesting an accelerated removal of mitochondria (Fig 7c). Consistently, mitophagosomes 209 were most frequently detected in cotyledon cell sections from 6 and 8 hr samples under TEM 210 (Fig. 7d). Cotyledon greening was severely affected in atg5-1 mutant seedlings and no 211 mitophagosomes were discerned in their cotyledon cells (Fig. 7 a,b,e). These results indicate 212 de-etiolation is a physiological stress condition for mitochondria where mitophagy is induced to 213 mediate mitochondrial turnover that underlies light activated cotyledon development.

214 **Discussion**

215 Research in the last decade has transformed autophagy from a bulk degradation system 216 to a highly selective cellular quality control pathway that rapidly removes toxic or superfluous macromolecules^{28,29}. Especially organelles that get damaged due to metabolic and physiological 217 218 stress conditions are mainly recycled via distinct selective autophagy pathways⁶. Consistently, 219 in all the eukaryotes tested so far, autophagy is essential for adapting to environmental changes^{12,30}. However, despite significant advances made in metazoan selective autophagy field, 220 221 how plants recycle their organelles are still mostly unknown. Although the core autophagy 222 machinery that mediates formation of the autophagosome is highly conserved in plants, 223 selective autophagy receptors and adaptors that are responsible for recognition and 224 recruitment of damaged organelles to the autophagosomes are not well conserved¹². For 225 example, there are up to eight different receptor proteins and dozens of accessory proteins 226 that have been shown to mediate various mitophagy pathways in mammalian cells³. Homologs 227 of most of those proteins are lacking in plant genomes, implying plant mitophagy have followed 228 a different evolutionary path, likely due to the presence of another endosymbiont in the cell.

Here, we have established a detailed toolbox to study mitophagy in plants. We have shown that protonophore uncouplers specifically induce mitophagy similar to metazoans. Our findings also revealed high levels of mitophagy during de-etiolation, a fundamental developmental step that allows plants to survive day light after germination. Excitingly, our studies also revealed a molecular player, the Friendly protein, that is essential for mitophagy.

234 Mitochondrial membrane potential is closely monitored by mitochondrial quality

235 control pathways

236 As the primary producer of ATP in eukaryotic cells, healthy mitochondria must maintain 237 the electrochemical potential. It is conceivable that the loss/reduction in the potential serves 238 as a mark for the mitophagy machinery to recognize malfunctioning mitochondria. For 239 example, in the PINKI/Parkin-dependent mitophagy pathway of mammalian cells, a 240 mitochondrion-localized protein kinase, Pinkl, is stabilized when the membrane potential 241 drops and this leads to the activation of downstream effectors³¹. In *C. elegans* sperm 242 mitochondria are rapidly removed from the oocyte via mitophagy upon fertilization^{32,33}. 243 Although the underlying mechanism of depolarization is still unknown, the paternal 244 mitochondria that is recycled by mitophagy loses its membrane potential prior to mitophagy¹⁵. 245 Our uncoupler treatments clearly demonstrated that membrane potential serves as a proxy 246 for mitochondrial health across different organisms and loss of membrane potential triggers 247 mitophagy. However, how plants tag depolarized mitochondria for mitophagy needs to be 248 further investigated.

249

250 Mitochondria in mammalian and yeast cells undergo cycles of fusion and fission. This 251 mitochondrial dynamics collaborate with mitophagy; fusion can rescue damaged mitochondria 252 by diluting their injuries while fission singles out aberrant mitochondria for mitophagy. It was 253 shown that mitochondria fusion is inhibited when the mitochondrial membrane potential is 254 dissipated by uncouplers³⁴. Mitochondria in Arabidopsis root cells are mostly round, indicating 255 that mitochondria fission dominates over fusion³⁵. Therefore, our system is not suited for 256 investigating the link between mitochondria's membrane potential and their fusion/fission 257 dynamics. Some plant cells including seed cells after germination or shoot apical meristem cells have elongated mitochondria constituting a network^{36,37}. These cells are better for testing 258 259 whether mitochondria fragment in response to uncoupler stresses and studying roles of 260 mitochondria fission in plant mitophagy.

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262 Ultrastructural features of compromised mitochondria and mitophagosomes in 263 Arabidopsis root cells.

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265 Mitochondria with dark aggregates and shrivelled cristae in the matrix appeared after 266 incubation with uncouplers. They were also abundant in de-etiolating cotyledon cells and fmt 267 mutant cells (Fig. 3 and Extended Data Fig. 1). These impaired mitochondria were rare in 268 DMSO control samples but frequently observed in *atg5-1* mutant samples in TEM images. 269 Considering that they are specifically targeted by and enclosed in mitophagosomes, they 270 correspond to depolarized mitochondria being recycled. Sperm mitochondria in cryofixed C. elegans oocytes also exhibited similar ultrastructural features^{15,38}. However, we did not 271 272 observe large ruptures in mitochondrial membranes reported in TEM analysis of mammalian 273 mitophagy where cells were preserved by chemical fixation³⁹.

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An intriguing feature that we noticed in our TEM analysis is that tips of the elongating phagophores were in contact with the mitochondrial surface (Fig. 3e, arrowheads), This observation is consistent with the YFP-ATG8e fluorescence that expanded tightly over mitochondria in the time-lapse recordings (Fig. 3c). The affinity between the autophagosome membrane and mitochondria explains why mitophagosomes usually have one damaged mitochondrion, and they do not capture other organelles. It also suggests specific proteinprotein interactions regulate the two membranes, which need to be investigated further.

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283 FMT is essential for mitophagy in plants.

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285 FMT is required for normal mitochondria distribution in the plant cell, possibly regulating association between individual mitochondria before fusion²⁶. We have shown that DNP-286 287 induced mitochondria recycling is affected in *fmt* mutant cells, and FMT associates with ATG8 288 upon mitochondrial damage. These data suggest that FMT associates with autophagosomes 289 during mitophagy. Because FMT is a protein shuttling between the cytosol and mitochondria, 290 it is tempting to speculate that FMT may participate in an autophagy receptor or adaptor 291 complex through which the core autophagy machinery is recruited to mitochondria. It is also 292 possible that FMT plays a role in the repair of damaged mitochondria by controlling their 293 interaction with the healthy mitochondria, and excessive accumulation of FMT triggers the 294 onset of mitophagy.

295

296 FMT is a highly conserved protein with orthologs in evolutionarily distant eukaryotes 297 including yeast and metazoans⁴⁰. *fmt* mutants exhibit similar phenotypes where mitochondria 298 form large clusters next to nucleus. The Drosophila FMT homolog Clueless positively 299 regulates PINK1/Parkin dependent mitophagy by suppressing mitochondrial fusion⁴¹. Recent 300 studies have shown that mammalian FMT homolog CLUH could bind RNA to form granules. 301 CLUH granules regulate translation of mRNAs linked to metabolic activity and regulate 302 mitophagy and metabolic reprogramming^{42,43}. Plants lack PINK I/Parkin homologs, so whether 303 FMT regulates mitophagy by forming stress activated granules or via a PINK/Parkin-like 304 pathway need to be investigated further. Identification of FMT interacting proteins and RNA 305 during nutrient starvation or uncoupler treatments could help us understand the role of FMT 306 in mitophagy and mitochondrial quality control.

307

308 Mitochondrial recycling mediate organellar reprogramming during de-etiolation. 309

Reprograming the mitochondrial functions in response to nutrient availability is critical for cell survival⁴⁴. In plant cells, photosynthesis in chloroplasts and respiration in mitochondria are coordinated for homeostasis of cellular energy levels and redox status^{45,46}. Our results from greening Arabidopsis cotyledons indicated that de-etiolation involves a wave of mitochondrial turnover, probably for rewiring mitochondrial metabolic network for adapting to light condition and salvaging raw materials for chloroplast biogenesis.

316

317 In skotomorphogenic seedlings, nutrients reserved in the seed are mobilized to sustain 318 growth, and mitochondria are required for the anabolic processes. When light is available, the 319 seedlings become photosynthetically active and capable of autotrophic growth. It was shown 320 that the electron transport chain in mitochondria is slowed down in de-etiolating wheat leaves 321 and chloroplasts and mitochondria are functionally more intertwined⁴⁷. Accumulation of 322 mitophagosomes and rapid changes in mitochondrial protein levels in the greening cotyledon 323 suggest a cannibalization of pre-existing mitochondria. We speculate that the developmentally 324 programmed mitophagy could facilitate modulation of the mitochondrial pool and biosynthesis 325 of macromolecules for constructing new organelles. The lack of mitophagosomes and the 326 delay in greening of the *atg5-1* mutant cotyledon agrees with the notion. Altogether, our 327 findings on greening cotyledons present a clear example of inter-organelle communication,

- 328 and how mitochondria-chloroplast crosstalk could underlie a major developmental transition
- in plants.
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331 Methods

332 Materials and plant growth conditions

333 All the chemicals were purchased from Sigma-Aldrich (http://www.sigmaaldrich.com) or Thermo-334 Fisher (https://www.thermofisher.com) unless specified. YFP-ATG8e, mCherry-ATG8e, atg5-1, Mito-335 GFP, Mito-GFP::fmt and FMT-YFP seeds were described previously^{26,48,49}. mCherry-ATG8e::atg5-1, 336 Mito-GFP x mCherry-ATG8e and Mito-GFP x mCherry-ATG8e::fmt was obtained by crossing the 337 previously established lines^{26,50}. All transgenic lines were genotyped by PCR and homozygous lines 338 were isolated before experiments. All Arabidopsis seeds were surface-sterilized and geminated on $\frac{1}{2}$ 339 Murashige and Skoog (MS) agar plate in a growth chamber at 21 °C with 16 h light–8 h dark except for 340 de-etiolation experiments where seedlings were geminated under darkness.

341 **Protein extraction and immunoblot analysis**

342 Isolation of mitochondrial membrane proteins and GFP cleavage assay were carried out as described 343 previously^{48,51}. Briefly, seven-day-old Arabidopsis seedlings were incubated in DMSO, 50 μ M DNP, 50 344 μm MG132 or 1 μm ConA with indicated times in liquid half MS medium when necessary. Seedlings 345 for the nitrogen starvation experiments were germinated on half MS medium agar plate and then 346 transfer to liquid half MS medium without nitrogen for I day. All the protein samples were subjected 347 to 15% SDS-PAGE. Primary and secondary antibodies were diluted in 1x phosphate buffered saline 348 (PBS). Antibodies against GFP (Abcam), YFP (Agrisera), mCherry (Abcam), ATG8 (Agrisera), voltage-349 dependent anion channel I (VDACI; Agrisera), peripheral-type benzodiazepine receptor (PBR; 350 PhytoAB), cytochrome oxidase subunit II (COXII; Agrisera), L-galactono-1,4-lactone dehydrogenase 351 (GLDH; Agrisera, PhytoAB) and isocitrate dehydrogenase (IDH; Agrisera), Cycloartenol-C24-methyl 352 transferase (SMT1; Agrisera), Coatomer subunit gamma (Sec21p; Agrisera) were obtained from the 353 indicated sources. We performed Student's t test (one-tailed and unpaired test) with the triple 354 replicate immunoblot data and the quantification of band intensities was performed using Image] 355 (National Institutes of Health) and Microsoft Excel 2016, the graphs were made by Prism8 (GraphPad 356 Software). Representative of at least three independant immunoblot results were shown in the figures.

357 **ATG8 delipidation assay**

358 Protein extraction methods for ATG8-delipidation were described previously⁵². Seven-day-old 359 Arabidopsis seedlings were incubated in 50 μ M DNP for 2 hours before protein extraction. The total 360 plant lysates were extracted in lysis buffer [50 mM Tris-HCl (pH 8.0), 150 mM NaCl, 0.5 mM

Ethylenediaminetetraacetic acid (EDTA) and 1x Complete Protease Inhibitor Cocktail] and then centrifuged at 14 000 rpm for 10 min at 4 °C. The supernatant was centrifuged at 100 000 g for 1 h, with the membrane pellet then solubilized in lysis buffer containing 0.5% (v/v) Triton X-100. The solubilized membrane samples were incubated at 37°C for 1 h with 250 unit/ml of phospholipase D (PLD) or an equal volume of its buffer. Protein samples were subjected to 15% SDS-PAGE in the presence of 6 M urea and analyzed by immunoblot with anti-ATG8 antibody.

367 Immunoprecipitation

- 368 Protein extraction and immunoprecipitation were performed as described previously⁴⁸. Seven-day-old 369 Arabidopsis seedlings were incubated in 50 μ M DNP for 2 hours before protein extraction. Total plant lysates were centrifuged at 14000 rpm for 10 min at 4 °C. The supernatant was prepared in lysis buffer 370 371 (10 mM Tris/HCl at pH 7.4, 150 mM NaCl, 0.5 mM EDTA, 5% glycerol, 0.2% Nonidet P-40, and 2 mM 372 dithiobis [succinimidyl propionate] containing Ix Complete Protease Inhibitor Cocktail) and then 373 incubated with GFP-TRAP agarose beads (ChromoTek) for 2 hours at 4°C. The beads were washed 374 five times (4°C) in wash buffer (10 mM Tris/HCl, pH 7.5, 150 mM NaCl, and 0.5 mM EDTA with 1x 375 Complete Protease Inhibitor Cocktail) and then eluted by boiling in 2x SDS sample buffer. Samples 376 were separated by SDS-PAGE and analyzed by immunoblot using indicated antibodies.
- 377

378 Confocal microscopy and image processing

379 Confocal fluorescence images were acquired using the Leica SP8 laser scanning confocal system with 380 a 63x water lens. Seven-day-old Arabidopsis seedlings were incubated in DMSO, 50 μ M DNP, 10 μ M 381 FCCP or 0.5 µM ConA with indicated times in liquid half MS medium when necessary before imaging. 382 Tetramethylrhodamine ethyl ester (TMRE) and MitoTracker Red (MTR) were used to stain Arabidopsis 383 root cell mitochondria at 500 nm for 10 mins. A sequential acquisition was applied when observing 384 fluorescent proteins. Images were processed with Photoshop CC (https://www.adobe.com) and 385 performed Student's t test (one-tailed and unpaired test) with Microsoft Excel 2016 386 (https://www.microsoft.com/). The graphs were prepared with Prism8 (https://www.graphpad.com).

387 **TEM** analysis, electron tomography, and 3d modeling

For TEM samples preparation, high-pressure freezing, freeze substitution, resin embedding, and ultramicrotomy were performed as described previously^{53,54}. In brief, Seven-day-old *Arabidopsis* seedlings were incubated in DMSO or 50 μ M DNP for indicated times and then rapidly frozen with an HPM100 high-pressure freezer (Leica Microsystems). The samples were freeze-substituted at -80°C for 72 h, and excess OsO4 was removed by rinsing with precooled acetone. After being slowly warmed up to room temperature over 48 h, root samples were separated from planchettes and embedded in Embed-812 resin (Electron Microscopy Sciences). Thin sections (100 nm thick) prepared

from sample blocks of each time point were examined with a Hitachi 7400 TEM (Hitachi-High Technologies) operated at 80 kV.

397 For dual-axis tomography analysis, semi-thick sections (250 nm) were collected on formvar-coated 398 copper slot grids (Electron Microscopy Sciences) and stained with 2% uranyl acetate in 70% methanol 399 followed by Reynold's lead citrate as described previously⁵⁵. Tilt series were collected from 60° to -400 60° (1.5° intervals) with a 200-kV Tecnai F20 intermediate voltage electron microscopy 401 (https://www.fei.com/). Tomograms were reconstructed as described⁵⁶. To generate models of 402 complicated thylakoid membranes, used the we autocontour command 403 (bio3d.colorado.edu/imod/doc/3dmodHelp/autox.html) of the 3dmod software package as explained 404 in Keith and Kang (2017)⁵⁷.

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414 **Author contributions**

J.M., Y.D., and B.-H.K. conceived and designed the experiments. J.M. performed the confocal
microscopy and stereomicroscopy. J.M., Z.L., and P.W. carried out electron
microscopy/tomography analysis. J.M. and J.F. prepared 3D tomographic models. J.M. and
W.M. performed immunoblot and pull-down experiments. J.Z., Y.Z., and N.G. did other
experiments. J.M., J.Z., Z.L., P.W., L.J., Y.D., and B.-H.K. analysed the data. J.M., Y.D., and B.H.K. wrote the paper.

421 **Competing interests**

422 The authors declare no competing interests.

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Main Figures

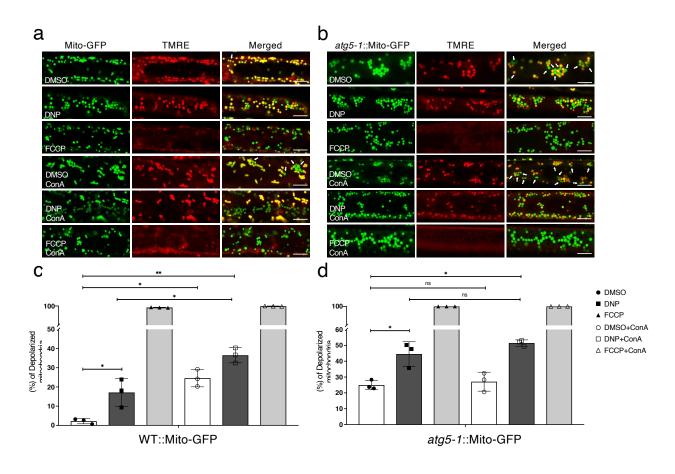
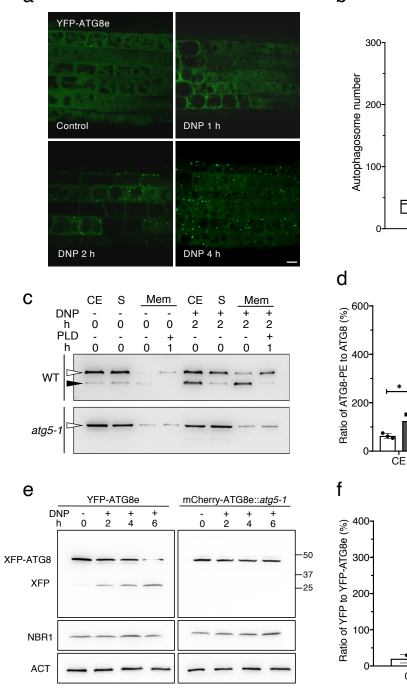


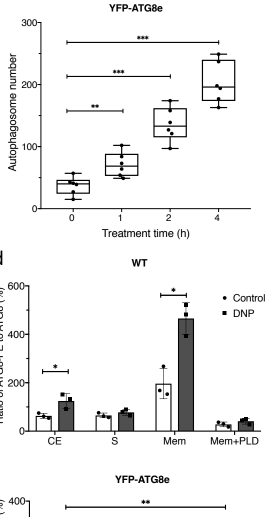
Figure I. Arabidopsis root cells accumulate depolarized mitochondria upon uncoupler treatments.

a,b, Uncoupler treatment induced mitochondria depolarization. Confocal micrographs of wild-type **(a)** and *atg5-1* **(b)** root cells expressing a mitochondrion-targeted GFP (Mito-GFP) after uncoupler treatment. Mitochondria were prestained with TMRE. Normal mitochondria exhibit yellow fluorescence while depolarized mitochondria exhibit green fluorescence (arrows) in DMSO or DMSO + ConA panels in the merged image columns. Note that most mitochondria are round. Scale bars, 8 μ m. **c,d**, Histograms illustrating the percentage of depolarized mitochondria in wild type (WT) and *atg5-1* root cells expressing Mito-GFP at each treatment conditions. Bars represent the mean (\pm SD) of three biological replicates, each generated with three technical replicates. About 500 mitochondria from 10 cells (five root samples) were counted per condition. Asterisks (*) denote significant differences in depolarized mitochondria percentages relative to DMSO control group under each condition (unpaired t-test, *p<0.05, **p<0.01, ns, no significant difference).

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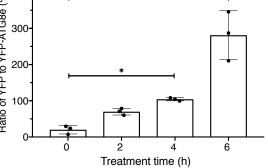


Figure 2. Uncoupler treatments induce autophagy.

a, DNP treatment induces autophagosome formation. *Arabidopsis* YFP-ATG8e seedlings were incubated in DNP solution for varying periods (0-4 h) before imaging. **b**, Quantification of the number of autophagosomes from more than five independent root samples (unpaired t-test, **p<0.01, ***p<0.001). Scale bars, 8 μ m. **c**, Uncoupler treatment activates ATG8 lipidation. Protein crude extracts (CE) were prepared from *Arabidopsis* root cells following incubation in DNP for 2 hours. Soluble (S) and membrane (Mem) fractions were separated and examined by immunoblot analysis with an anti-ATG8 antibody. White arrowheads mark ATG8. An additional polypeptide recognized by the antibody is enriched in the membrane fraction when cells are incubated with DNP (black arrowhead). **d**, Histograms illustrating polypeptide intensity ratios of lipidated ATG8 to free ATG8 in (**c**). Bars represent the mean (\pm SD) of three biological replicates. **e**, ATG8 cleavage assays of DNP treated WT and *atg5-1* seedlings expressing YFP-ATG8e or mCherry-ATG8e, respectively. Protein extracts were prepared from *Arabidopsis* seedlings exposed to DNP (50 μ M) for the indicated time periods and subjected to immunoblot analysis with anti-GFP or anti-mCherry antibodies. NBR1 and Actin were used as control. **f**, Histograms illustrating the polypeptide intensity ratios of free YFP to YFP-ATG8e in (**d**). Bars represent the mean (\pm SD) of three biological replicates. (unpaired t-test, *p<0.05, **p<0.01).

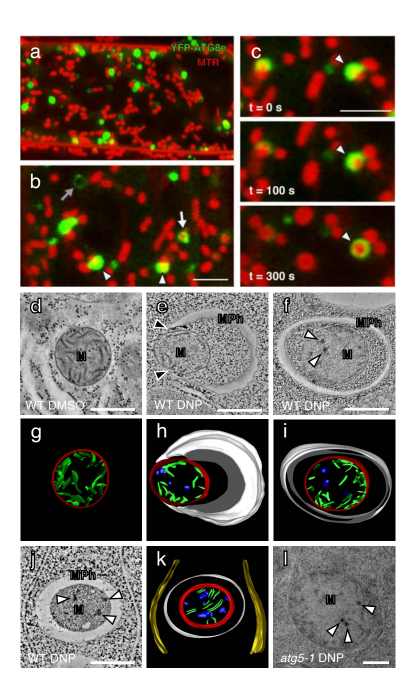


Figure 3. Depolarized mitochondria are selectively engulfed by autophagosomes in uncouplertreated Arabidopsis root cells.

a,b, DNP treatment induces mitophagy in *Arabidopsis* root cells. Confocal micrographs of *Arabidopsis* root cells expressing YFP-ATG8e stained with MitoTracker Red (MTR) and incubated with DNP for I hour. The mitochondria that associate with YFP-ATG8e are indicated with the arrowheads in panel (**b**). The mitochondria that are completely engulfed by ATG8e fluorescence is marked with a white arrow. Empty YFP fluorescence circles were also observed (grey arrow in panel **b**). Scale bars, 5 μ m. **c**, Time lapse imaging of a mitophagy event in an *Arabidopsis* root tip cell treated with DNP. The mitochondrion is engulfed by YFP-ATG8e over 5 minutes. Scale bar, 5 μ m. **d-f,j,l**, Transmission electron micrographs of mitochondria (M) in *Arabidopsis* WT or *atg5-1* root cells incubated with DMSO or DNP. Mitochondria phagophores (MPh) assemble in the vicinity of the mitochondria. Note that the phagophore tips (black arrow) are in contact with the mitochondrial cargo (M) based on the tomogram in (**d-f,j,l**). Mitophagosome (white), mitochondria outer membrane (red), mitochondria cristae (green), damaged cristae formed aggregates (blue) and ER (yellow) are modeled. White arrows indicate dark aggregates in the matrix of compromised mitochondria. Scale bars, 500 nm.

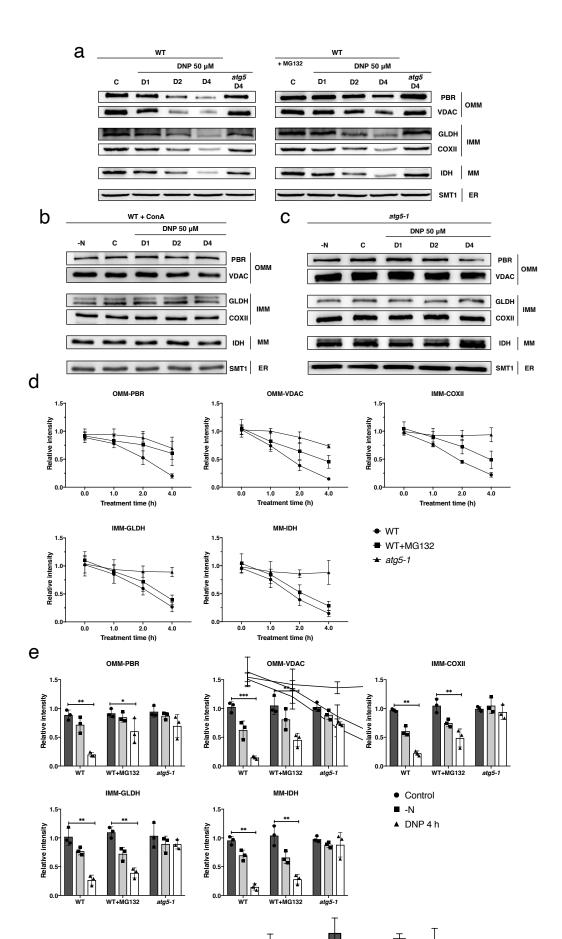
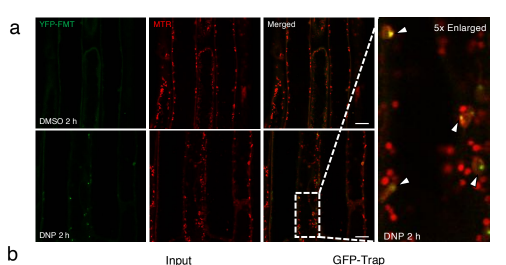
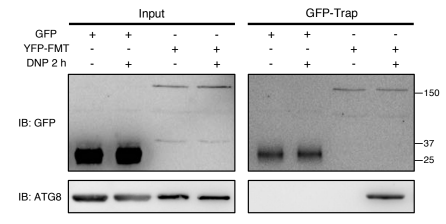
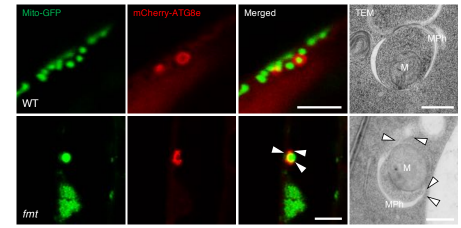


Figure 4. ATG5-dependent degradation of mitochondrial proteins in uncoupler-treated Arabidopsis root cells.

a-c, Immunoblot blot analyses of uncoupler treatment induced mitochondrial protein degradation in Arabidopsis WT and atg5-1 seedlings. WT and atg5-1 mutant Arabidopsis roots were incubated in DNP solution for I to 4 hours (DI-D4) or nitrogen starvation (-N) solutions for I day. Mitochondrial outer membrane, mitochondrial matrix, and endomembrane fractions were isolated and subjected to immunoblot analyses. For outer mitochondrial membrane (OMM) proteins, PBR and VDAC1, for inner mitochondrial membrane (IMM) COXII and GLDH, and for mitochondrial matrix IDH were analysed. Concanamycin A (ConA) was added to the treatment solution to test for vacuolar function in mitochondrial protein degradation. Proteasome inhibitor MGI32 was added to test the involvement of proteasomes in the recycling of mitochondrial membrane proteins (c). Note that an ER protein, SMTI was not affected by DNP treatment. Equal amounts of protein extracts were analysed in the immunoblots shown. d, Line charts illustrating degradation rates of OMM proteins (PBR and VDACI), IMM proteins (COXII and GLDH), and MM protein (IDH) in WT treated with DNP (WT), atg5-1 treated with DNP (atg5-1), and WT treated with DNP and MG132 (WT+MG132). e, Histograms illustrating the levels of mitochondria membrane proteins under the three treatment conditions, DMSO 4h (Control), DNP 4 h, and nitrogen starvation (-N) in WT and atg5-1 root cells. The polypeptide intensity values were normalized to that of the loading control (SMTI). Bars represent the mean (\pm SD) and the asterisks (*) indicate decreases in polypeptide readouts significantly from that of the control (C) point (unpaired ttest, *P < 0.05, **P < 0.01, ***P < 0.01).







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Figure 5. Friendly associates with damaged mitochondria and ATG8 upon uncoupler treatment.

a, DNP treatment induces recruitment of Friendly (FMT) to damaged mitochondria. Confocal micrographs of *Arabidopsis* root cells expressing a YFP-tagged FMT (YFP-FMT) prestained with MTR. FMT-YFP seedlings were incubated with DMSO or DNP for I hour prior to imaging. Scale bars, 8 μ m. **b**, FMT associates with ATG8 upon DNP treatment. *Arabidopsis* root cells expressing YFP-FMT were incubated with DMSO or DNP for 2 hours and then subjected to immunoprecipitation with GFP-trap followed by immunoblotting with indicated antibodies. **c**, Confocal micrographs of *Arabidopsis* WT and *fmt* root cells expressing mitochondria-targeted GFP (Mito-GFP) and mCherry-targeted ATG8e (mCherry-ATG8e). WT or *fmt* plants seedlings were incubated with DNP for I hour prior to imaging. Scale bars, 8 μ m. Transmission electron microscopy (TEM) photos show mitochondrial phagophores (MPh) assemble in the vicinity of the mitochondria (M) under the condition for the experiment in **(a)**. Arrowhead point to defective phagophores in confocal and TEM micrographs. Scale bars, 500 nm.

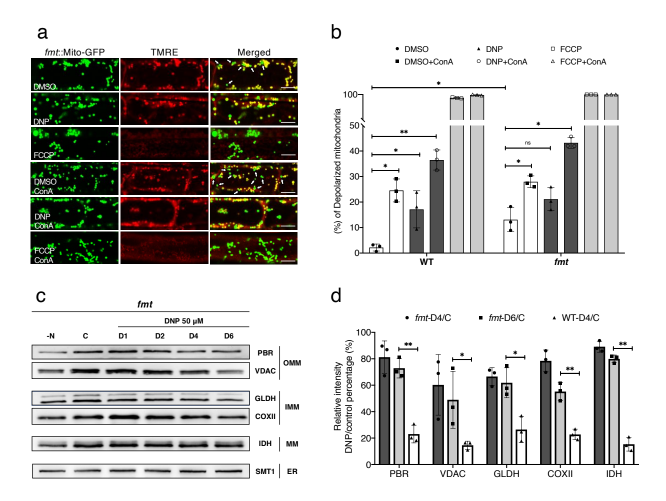
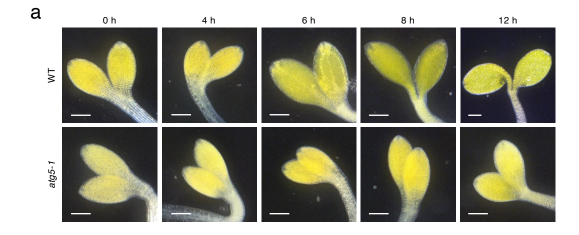
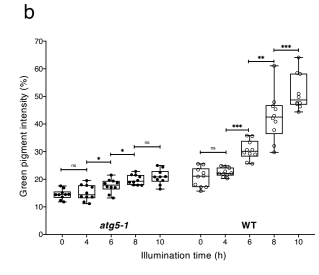
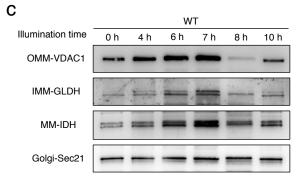


Figure 6. Arabidopsis friendly mutant have defects in clearance of depolarized mitochondria.

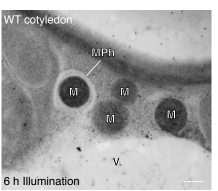
a, Uncoupler treatment induced mitochondria depolarization in Arabidopsis friendly mutant (fmt). Confocal micrographs of fmt root cells expressing a mitochondrion-targeted GFP (Mito-GFP), stained with TMRE. Normal mitochondria exhibit yellow fluorescence and depolarized mitochondria exhibit green florescence, denoted with arrows in DMSO or DMSO +ConA panels in the merged image column. Arabidopsis seedings were incubated with DMSO, DNP or FCCP with or without ConA for I hour prior to imaging. Scale bars, 8 µm. b, Histograms illustrating the percentage of depolarized mitochondria for each treatment conditions. Bars represent the mean (± SD) of three biological replicates. About 500 mitochondria from 10 cells (five root samples) were counted per condition. An asterisk (*) represents a significant difference of depolarized mitochondria percentage in each treatment relative to DMSO control group (unpaired t-test, *p<0.05, **p<0.01). c, Uncoupler treatment induced mitochondrial protein degradation in fmt mutant. Arabidopsis fmt mutant roots were incubated in DNP solutions for I hour to 6 hours (DI-D6) or nitrogen starvation (N-) solutions for I day. For outer mitochondrial membrane (OMM) proteins, PBR and VDACI, for inner mitochondrial membrane (IMM) COXII and GLDH, and for mitochondrial matrix IDH were examined with immunoblot analysis. Note that an ER protein, SMTI, was not affected by DNP treatment. d, Histograms illustrating the levels of mitochondrial membrane protein degradation for the DNP treatment (DNP 4 h, DNP 6 h) in WT and fmt root cells. The polypeptide intensity values were normalized with that of the loading control (SMTI) and the percentage of DNP treatment group to control group are quantitated. Bars represent the mean (\pm SD) and the asterisks (*) indicate the significantly difference in polypeptide readouts between fmt and WT groups (unpaired t-test, *P < 0.05, **P < 0.01).







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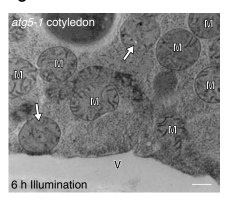


Figure 7. Mitophagy is triggered during de-etiolation of Arabidopsis seedlings.

a, Cotyledon greening after light exposure in WT and atg5-1 Arabidopsis. Darkfield stereo microscopy photos showing Arabidopsis WT and atg5-1 mutant cotyledon at multiple time points (0-12 h) after illumination. Arabidopsis seedlings were grown darkness before the experiment. Scale bars, 1 mm. b, Quantification of the green pigments in cotyledons after light exposure. The green colour of ten Arabidopsis cotyledons for each time points were calculated from their photos and normalized against the dark background. (unpaired t-test, *p<0.05, **p<0.01, ***p<0.001, ns, no significant difference). c, Immunoblot analyses of mitochondrial proteins in Arabidopsis seedlings during de-etiolation. For outer mitochondrial membrane (OMM) proteins, PBR and VDAC1, for inner mitochondrial membrane (IMM) COXII and GLDH, and for mitochondrial matrix IDH were analysed as representative proteins. An Arabidopsis Golgi protein, coatomer subunit gamma (Sec21), was employed as the loading control. d,e, TEM images of a cluster of mitochondria (M) near the vacuole (V) in Arabidopsis WT and atg5-1 cotyledon cell after 6 hours of illumination. Mitochondria were seen to be surrounded by mitophagosome (MPh) in WT. By contrast, compromised mitochondria with dark aggregates were abundant in atg5-1 (arrows in e) but no mitophagosomes were associated with them. Scale bars, 500 nm.