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2	brown alga, Mutiomo cylindricus (Cutleriaceae, Tilopteridales)
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27 Abstract

28 Mitochondrial morphology varies according to development and the physiological conditions of 29 the cell. Here, we performed electron tomography using serial sections to analyze the number, 30 individual volume, and morphological complexity of mitochondria in the cells across two 31 generations in the life cycle of the brown alga *Mutimo cylindricus*. This species shows a 32 heteromorphic alternation of generations between the macroscopic gametophyte and the 33 crustose sporophyte during its life cycle and displays anisogamous sexual reproduction. We 34 observed the mitochondria in the vegetative cells of gametophytes and sporophytes to mainly 35 show tubular or discoidal shapes with high morphological complexity. The morphology of the 36 mitochondria in the male and female gametes changed to a nearly spherical or oval shape from a 37 tubular or discoidal shape before release. In this species, degradation of the paternal 38 mitochondria was observed in the zygote 2 h after fertilization. Morphological changes in the 39 mitochondria were not observed until 6 h after fertilization. Twenty-four-hour-old zygotes 40 before and after cytokinesis showed a similar number of mitochondria as six-hour-old zygotes; 41 however, the volume and morphological complexity increased. The results indicated that the 42 maternal mitochondria did not undergo fission or fusion until this stage. Based on the analysis 43 results of the number and total volume of mitochondria before and after the release of the 44 gametes, it is possible that the mitochondria in the female gametes fuse immediately before 45 release. 46

47

48 Keywords

Brown algae · electron microscopy · life cycle · mitochondria · serial section tomography
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52 Introduction

Mitochondria are membrane-bound organelles with their own DNA and supply
adenosine triphosphate (ATP), which is used as a source of chemical energy in cell activity. It is
believed that morphological characteristics of the mitochondria, such as shape, size, and number,
change throughout the life of a cell. This suggests that, morphologically, mitochondria vary
depending on development, growth, differentiation, apoptosis, maturation, and aging (Chan
2012).

59 In unicellular organisms, following changes in mitochondrial morphology throughout 60 the life cycle is comparatively easier, and in some cases, the life cycle may directly indicate the 61 cell cycle. Cyanidioschyzon merolae possesses a single mitochondrion whose division, along 62 with chloroplast division, precedes nuclear division. The mitochondria change their morphology 63 into spherical, ovoid, and dumbbell-shaped structures before division and return to a spherical 64 shape after division (Kuroiwa 2010; Yoshida et al. 2017). In the unicellular green alga 65 Chlamvdomonas reinhardtii, mitochondria form a tubular network toward asexual reproduction, 66 and mitochondria in the liberated daughter cells exhibit a giant form and, subsequently, a thick-67 and thin-corded form (Ehara et al. 1995). However, during meiosis after conjugation, the 68 mitochondrial morphology exhibits a continuous change into spherical, tubule-, branch-, and 69 mesh-like structures (Aoyama et al. 2009).

70 In multicellular organisms, almost all studies relating to mitochondrial morphology 71 have focused on a specific developmental period, such as gametogenesis, fertilization, or early 72 embryonic development. Mitochondria in animal sperm exhibit characteristic features and 73 maternal inheritance (Sato and Sato 2013). Morphological changes and the transition of mtDNA 74 content during spermatogenesis can serve as a key to understanding the mechanism of maternal 75 inheritance of this organelle. In mammals, to prevent the transmission of paternal mtDNA to the 76 progeny, the number of nucleoid mitochondria gradually decreases during sperm formation 77 (Rantanen and Larsson 2000). In addition, a drastic morphological transformation of the

78 mitochondria are observed during spermatogenesis. The mitochondria in spermatogonia, 79 leptotene, and zygotene spermatocytes are generally small and spherical. They elongate in 80 pachytene spermatocytes and early spermatids, and fragment again in late spermatids (De 81 Martino et al. 1979). In maturing spermatids, small mitochondrial spheres are regularly arranged 82 across the midpiece, and they elongate as they wrap around the midpiece (Ho and Wey 2007). 83 In the early stage of spermatogenesis in rabbits, mitochondrial cristae have been observed to 84 become highly dilated during the transition phase between spermatogonia and primary 85 spermatocytes (Nicander and Ploen 1969).

86 In a previous study, we clarified paternal inheritance of mitochondria in the 87 anisogamous brown alga Mutimo cylindricus (Shen et al. 2020). We concluded that maternal 88 inheritance of mitochondria could be universal in brown algae. In the case of M. cylindricus, 89 observations of ultrathin sections indicated that the size of the mitochondria in the male and 90 female gametes was almost identical. However, the average DNA content of one mitochondrion 91 in male gametes was one-seventh of that in female gametes. Mitophagic digestion of paternal 92 mitochondria occurs in zygotes. One of the mechanisms in uniparental inheritance of 93 mitochondria in this species is the reduction of mitochondrial DNA in male gametes. However, 94 the mechanism underlying the formation of robust mitochondria in female gametes is unclear. 95 All species of brown algae show multicellularity. Some of them alternate between the 96 generation of diploid sporophytes and haploid gametophytes. M. cylindricus is an anisogamous 97 species with a heteromorphic life cycle alternating between macroscopic gametophytes and 98 microscopic sporophytes (Kitayama et al. 1992; Kawai et al. 2012). In this study, we examined 99 mitochondrial morphology in gametophytes, gametangia, gametes, zygotes, and sporophytes 100 through 3D analysis using transmission electron microscopy (TEM) on serial section images. In 101 brown algae, 3D analysis has been performed using electron tomography (Terauchi et al. 2012; 102 Fu et al. 2013; Nagasato et al. 2014; Kinoshita et al. 2016). Those studies revealed the structure 103 of the plasmodesmata, transient membrane configuration during cytokinesis, and flagella. The

104	present study is the first to demonstrate the 3D reconstruction of a structure using serial sections,
105	focusing on the structure of brown algal mitochondria. These data enabled quantification of
106	mitochondria in each target stage.
107	
108	Material and Methods
109	Culture
110	Mature male and female gametophytes of <i>M. cylindricus</i> (Okamura) H. Kawai and T.
111	Kitayama (Kawai et al. 2012) were collected at Odanohama, Toba, Mie, Japan (34°45' N
112	136°87' E), in March 2015 and April 2019. Male and female individuals were identified based
113	on the morphology of the plurilocular gametangia under a light microscope. Unialgal cultures of
114	M. cylindricus were established from gametes or trichothallic hairs of male and female
115	gametophytes (Shen et al. 2020) and incubated in sterilized seawater containing half-strength
116	Provasoil's enriched seawater medium (Provasoil 1968). Gametogenesis and maturation were
117	induced under the following conditions: 15 °C, using white-light-emitting diodes (LEDs, 20-40
118	μmol photons $m^{\text{-2}} \text{s}^{\text{-1}})$ and long-day conditions (14 h light:10 h dark). The release of gametes
119	was stimulated by exchanging the fresh culture medium the day before and by light the next day.
120	Male and female gametes were collected separately in microtubes on ice. Fertilization was
121	induced by adding male gametes to settled female gametes on a gel support film (ATTO, Tokyo,
122	Japan) in Petri dishes.
123	
124	Cryofixation and freeze-substitution for TEM samples
125	Rapid freezing and freeze substitution by the immersion method were applied to the
126	male and female gametes, 2-, 6-, and 24-h-old zygotes, and two-, four-, and seven-celled
127	sporophytes on the gel support film. The gametes were collected as pellets following
128	centrifugation at 500 \times g for 1 min using a KUBOTA model 4000 centrifuge (KUBOTA, Tokyo,
129	Japan). The pellets were then placed and spread on a formvar-coated gold ring immediately

130 before freezing. The gel support films containing the just fertilized and developing specimens 131 were cut into triangular shapes $< 1 \text{ cm}^2$ (Nagasato and Motomura 2002). The gametes on the 132 gold rings, or early zygotes on the gel support films, were rapidly immersed in a pre-cooled 133 liquid propane bath (-186 °C) and subsequently transferred into liquid nitrogen. Gametophytes 134 and 20-d-old crustose sporophytes were frozen using high-pressure freezing. Thalli fragments 135 were cut, placed into gold carriers, frozen using a high-pressure freezer (Leica Microsystems, 136 Vienna, Austria) at -196 °C and 2,100 bar, and then stored in liquid nitrogen. 137 Freeze-substitution was performed by transferring the frozen samples from liquid 138 nitrogen to a freeze-substitution medium (2% osmium tetroxide [OsO4] dissolved in anhydrous 139 acetone) pre-cooled in a liquid nitrogen cooling bath, and the samples were stored at -85 °C for 140 2 d. The samples were gradually warmed to -20 °C for 2 h, to 4 °C for 2 h, and then to room 141 temperature for approximately 30 min. Next, the samples were washed several times with 142 anhydrous acetone at room temperature and infiltrated with increasing concentrations (10%, 143 30%, 50%, 70%, 80%, 90%, and 100%) of low-viscosity Spurr's epoxy resin (Polysciences, 144 Warrington, PA, USA). Finally, the samples were embedded in Spurr's epoxy resin in 145 aluminum foil dishes. 146 Serial sections were cut using a NACC diamond knife (Micro Star Technologies, 147 Huntsville, TX, USA) with an ULTRACUT ultramicrotome (Reichert-Jung, Vienna, Austria), 148 mounted on formvar-coated slot grids, and stained using an EM stainer (Nisshin EM, Tokyo, 149 Japan). The sections were observed using a JEM-1011 electron microscope (JEOL, Tokyo, 150 Japan). Consecutive serial sections of each sample were examined. Images were acquired using 151 TEM films (Fujifilm, Tokyo, Japan). The developed films were scanned and converted into 152 digital data using an EPSON GT-X980 photo scanner (EPSON, Nagano, Japan) and images 153 were standardized (tiff format, resolution 300 dpi, 8-bit grayscale).

154

155 Three-dimensional reconstruction

156 Alignment of whole cell serial section images was performed using Fiji software 157 (http://fiji.sc/Fiji, Lowe 2004; Schindelin et al. 2012; Murtin et al. 2018). The alignment method 158 for the serial sections was modified from that described in previous studies. Stacks were made 159 using two adjacent (source and target) images, and feature extraction was automatically 160 performed using the scale-invariant feature transform (SIFT) algorithm. The images were 161 aligned using the "Linear Stack Alignment with SIFT" plugin. The stack was converted into 162 images, the transformed target image was treated as a new source image, and the next section 163 was aligned as the target image. 164 For the 3D reconstruction of whole-cell serial section images, the sequence of 165 transformed images was converted into stacks using the IMOD software package (version 166 4.9.12) (Kremer et al. 1996). The pixel sizes of several standardized section images were 167 obtained using the "set scale" Fiji plugin according to the scale bar in each image 168 (Supplementary Table S1). The thickness range of the sections was estimated based on the 169 interference color of the ultrathin sections (Peachey 1958). Therefore, it was impossible to set 170 the z-axis value, the ratio of section thickness (nm) to pixel size (nm), in the stacking of serial 171 section images as accurately as in electron tomography. For this reason, the thickness was 172 calibrated using spherical organelles, such as small vacuoles (Cui et al. 2019). Finally, the mean 173 thickness (approximately 100 nm) of each gold serial section was determined (Supplementary

174 Fig. S1a, b). The 3D models were generated from the outlines of each object in the serial

175 sections along the z-axis. The volumes and surface areas of individual intracellular structures

176 (nuclei, mitochondria, plasma membrane) were obtained from the IMOD software and

177 calculated using the program IMODINFO.

The mitochondrial complexity index (MCI) is a score between the mitochondrial
surface area (SA) and volume (V) and is associated with mitochondrial shape complexity
(Vincent et al. 2019). Based on 3D modeling of mitochondria, we calculated the index

181 according to a previously reported formula: $MCI = SA^3/16\pi^2 V^2$ (Vincent et al. 2019). MCI is

182	0.71 for a spherically shaped mitochondrion and > 0.71 for an ellipsoid or tubular shape. A
183	higher value indicates greater elongation or branching of the mitochondrion.
184	
185	Statistical analysis
186	All data shown are derived from three cells, with the exception of two- (24-h-old
187	zygotes) or four-celled structures (four daughter cells from two 24-h-old zygotes after
188	cytokinesis), from independent biological samples in each life-cycle stage (Supplementary
189	Table S1; the corresponding 3D images are shown in Supplementary Fig. S2). All results are
190	presented as mean \pm standard error of the mean (SEM). Significant differences among different
191	stages of the life cycle were determined using Tukey's multiple comparison test (Supplementary
192	Table S2). Statistical tests were performed using GraphPad Prism software (version 8.0.2,
193	GraphPad Software, Inc., San Diego, CA, USA).
194	
195	Results
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208 a 6-h-old zygote and a two-celled sporophyte, respectively. Two orange eyespots derived from 209 male and female gametes remained. One 20-d-old crustose sporophyte is shown in 210 Supplementary Figure S3i. 211 212 Mitochondrial morphology in gametophytes 213 We first observed the ultrastructure of uniseriate filamentous male and female 214 gametophytes (Fig. 1, Supplementary Movie S1, S2). The nucleus was generally located in the 215 middle part of the cell (Fig. 1a, b), and chloroplasts and mitochondria were positioned in the 216 perinuclear or pericytoplasmic regions by occupation of large vacuoles (Fig. 1a, b). 217 Reconstruction of 3D images was performed by tracing the outline of the nucleus, mitochondria, 218 and plasma membrane in the male and female gametophytes. Most mitochondria at this stage 219 were tubular, curved, V-, or Y-shaped (Fig. 1c-f). The mitochondria were not branched and did 220 not form interconnection networks. 221 We calculated the total number and total and mean volumes of mitochondria from three 222 cells in each of the male and female gametophytes (Supplementary Table S1). There were 50, 223 83, and 117 mitochondria in the male gametophytes and 42, 59, and 104 in the female 224 gametophytes. In the male gametophytes, the cells with the lowest number of mitochondria 225 exhibited the highest total mitochondrial volume in the cell ("Male Gametophyte 3" in 226 Supplementary Table S1) because of variation in the mean volume of mitochondria in each 227 male gametophyte (Supplementary Table S1). The mean values of the mitochondrial volume in 228 the female gametophytes were not significantly different in each cell (Supplementary Table S1); 229 therefore, the number of mitochondria reflected the total volume of the mitochondria in the cell. 230 In this study, mitochondrial complexity was calculated to understand mitochondrial morphology 231 (Viecent et al. 2019). The analysis showed that the mitochondria in each cell of the 232 gametophytes exhibited considerable differences from spherical or elliptical geometries 233 (Supplementary Table S1).

235 Mitochondrial morphology in mature gametangia

236 Next, we examined mitochondrial morphology in mature male and female gametangia 237 (Fig. 2, Supplementary Movie S3, S4). Mitochondrial morphology and distribution were 238 determined through 3D analysis using three cells each of the male and female gametangia. In 239 the mature male gametangia, two flagella were present and an eyespot was observed within the 240 chloroplast adhered to the nucleus (Fig. 2a). Most mitochondria in the mature male gametangia 241 showed a discoid shape (Fig. 2b, c). Similarly, the flagella and eyespots were also seen in the 242 cells of mature female gametangia, and most mitochondria were tubular in shape with some 243 discoid (Fig. 2d-f). In the cells of male and female gametangia, some mitochondria were 244 located around the nucleus, while others were scattered in the cytoplasm. The number of 245 mitochondria examined in this study is summarized in Supplementary Table S1. Although the 246 number of mitochondria is different in each cell, the variability in the numbers was not as large 247 as in the gametophytes.

The average number of mitochondria in the cells of male and female gametangia was approximately 6 and 38, respectively. The difference in the total volume of the mitochondria in male or female gametangium was less compared to that observed in the gametophytic cells. In addition, the average mitochondrial volume in the male or female gametangia was

approximately the same. The mitochondria in the cells of the male gametangia were slightly

253 larger $(0.162 \pm 0.013 \ \mu\text{m}^3)$ than those in the cells of the female gametangia $(0.133 \pm 0.004 \ \mu\text{m}^3)$;

254 however, there were no significant differences in the mitochondrial shape (Supplementary Table

255 S2), as indicated by the MCI, between the male (1.682 ± 0.091) and female (1.852 ± 0.074)

256 gametangia. The average MCI in the gametangia was lower than that in the gametophytes (male

257 gametophytes, 4.848 ± 0.282 ; female gametophytes, 6.190 ± 0.280 ; Supplementary Table S1),

258 indicating that the mitochondrial shape changed during gametogenesis.

260 Mitochondrial morphology in male and female gametes

261 3D reconstruction of the male and female gametes released from the gametangia was 262 performed next (Fig. 3, Supplementary Movie S5, S6). The shape of the mitochondria in the 263 male and female gametes were nearly spherical (Fig. 3a, b). Most mitochondria were distributed 264 in the anterior part of the gametes (Fig. 3c-f). The average mitochondrial number in three cells 265 each of the male and female gametes was 5 and 27, respectively (Supplementary Table S1). A 266 conspicuous decrease in the number of mitochondrial in female gametes prior to gamete release 267 was confirmed. However, the total mitochondrial volume did not change before and after the 268 release of gametes in both males and females. The average mitochondrial size was $0.173 \pm$ 269 $0.022 \ \mu\text{m}^3$ in male gametes and $0.191 \pm 0.006 \ \mu\text{m}^3$ in female gametes, indicating that while 270 female mitochondria were slightly larger than male mitochondria, the difference was not 271 significant (Tukey's multiple comparison P = 0.4593; Supplementary Table S2). The MCI of 272 the male and female mitochondria was 0.758 ± 0.007 and 1.034 ± 0.042 , respectively. 273 According to these results, the shape of mitochondria in released gametes underwent a drastic 274 change either during or after gamete release. Mitochondria in male gametes were nearly 275 spherical (MCI ≈ 0.71) while the shape of mitochondria in female gametes was only slightly 276 spherical (Supplementary Table S1). However, based on Tukey's multiple comparison, the 277 shapes of mitochondria in male and female gametes were significantly different from each other 278 (P < 0.0001; Supplementary Table S2).

279

280 Mitochondrial morphology in early sporophytes

The disappearance of parental mitochondria in 2-h-old zygotes was observed (Shen et al.
2020). Mitochondrial morphology was examined in 2- and 6-h-old zygotes to compare the
configuration of the maternal mitochondria during or after the disappearance of the paternal
mitochondria (Fig. 4, Supplementary Movie S7, S8). Fertilization was confirmed by the number
of eyespots on the chloroplasts derived from male and female gametes (Shen et al. 2020). Two

286	eyespots were maintained until the formation of seven-celled sporophytes. The number of
287	mitochondria in the three 2-h-old zygotes was 25, 26, and 35 (Supplementary Table S1). The
288	zygote with 35 mitochondria ("2-h-old_Zygote_2" in Supplementary Table S1) appeared to
289	have male mitochondria that had not yet been digested. In three 6-h-old zygotes, the number of
290	mitochondria in each cell was 25, 29, and 30. The mean volume per mitochondrion in the 6-h-
291	old zygotes was $0.243 \pm 0.007 \ \mu\text{m}^3$, which indicated that these had mitochondria increased in
292	size compared to those in the 2-h-old zygotes (0.190 \pm 0.006 μm^3), and this increase was
293	statistically significant (P < 0.0001; Supplementary Tables S1 and S2). However, the MCI
294	values between the 2- and 6-h-old zygotes and the female gametes were not significantly
295	different (P = 0.9997; Supplementary Table S2).
296	The first cell division occurred symmetrically 24–48 h after fertilization. The
297	morphology of mitochondria in 24-h-old zygotes was compared before and after cytokinesis
298	(Fig. 5, Supplementary Movie S9, S10). The two zygotes examined before cytokinesis had
299	either 32 or 33 mitochondria while the two zygotes examined after cytokinesis had either 31 or
300	34 mitochondria. Almost half of the mitochondria in each daughter cell were distributed from
301	the mother cell (Supplementary Table S1). Dumbbell-shaped and elongated mitochondria were
302	observed in the cells before cytokinesis (Fig. 5a-c). The number of mitochondria did not change
303	toward the first cell division (Supplementary Table S1); however, the mitochondria became
304	larger and adopted more complex shapes than those from 2- (Fig. 4a-c) and 6-h-old (Fig. 4d-f)
305	zygotes. After cytokinesis, mitochondria in two-celled sporophytes were oval and tubular in
306	shape, and the dumbbell-shaped mitochondria were no longer observed (Fig. 5d-f).
307	
308	Mitochondrial morphology in sporophytes

A cross-section of a 20-d-old crustose sporophyte is shown in Figure 6a. In the marginal
meristematic cell layer, longer cells with asymmetric cytoplasmic distribution and shorter cells
lacking large vacuoles were observed (Fig. 6b, c). We examined the mitochondria in a short cell

312	(Fig. 6c-e, "20-d-old_Sporophyte_1" in Supplementary Table S1; Supplementary Movie S11).
313	To compare the mitochondrial morphology among the cell layers, the cells positioned at the
314	subapical (Fig. 6f-h, "20-d-old_Sporophyte_2" in Supplementary Table S1; Supplementary
315	Movie S12) and central regions (Fig. 6i-k, "20-d-old_Sporophyte_3" in Supplementary Table
316	S1; Supplementary Movie S13) were examined. There were no significant differences in the
317	total number and volume of mitochondria among the three cells examined (Supplementary
318	Table S1). Tubular, curved, V-, and Y-shaped mitochondria were observed in all three cells (Fig.
319	6c–k). The MCI was highest in the subapical cell (4.180 \pm 0.250), followed by the cell in the
320	central region (3.363 ± 0.203). The marginal apical cell had the lowest MCI (2.463 ± 0.147)
321	(Supplementary Table S1). There was little difference between the shapes of mitochondria in
322	the central and subapical regions ($P = 0.3047$; Supplementary Table S2); however, there was a
323	statistically significant difference in the shape of the mitochondrial in the apical region from
324	those in the subapical ($P < 0.0001$) and central regions ($P = 0.0264$; Supplementary Table S2).
325	The volume of each mitochondrion in non-gametophyte cells are summarized in Figure
326	7. This analysis showed that the volume of mitochondria increased before and after cytokinesis;
327	however, mitochondria in other stages were stable in volume.
328	

329 Discussion

330 Ultrastructural studies through the generations of the life cycle in *M. cylindricus*

To understand the dynamics of mitochondrial morphology across generations in brown algae, we selected *M. cylindricus* because of the heteromorphic alternation of generations in its life cycles and its anisogamous sexual reproduction patterns. The mitochondria in the uniseriate filamentous gametophytes, mature gametangia, gametes, zygotes, and crustose sporophytes were examined through consecutive serial sections using TEM. Moreover, quantitative analysis based on the 3D images was performed to characterize the mitochondrial morphology at each stage of the life cycle. Observation of the ultrastructure throughout the life cycle of brown algae was previously performed in *Scytothamnus australis* and *Scytothamnus fasciculatus* (Clayton
1986) using cultured material. In that report, however, the mitochondrial structure was not
described. The ultrastructure of mitochondria in the gametangia, sporangia, and gametes of
brown algae have been reported (Brawley et al. 1976; La Claire and West 1978, 1979; Berkaloff
and Rousseau 1979; Henry and Cole 1982a, b; Clayton 1984; Katsaros and Galatis 1986; Maier
1997); however, our study was the first to report mitochondrial morphological observations

across generations and to quantify mitochondrial morphology in brown algae using 3D.

345

346 Fusion of mitochondria at the final stage of female gametogenesis

347 In this study, 3D image reconstruction allowed us to quantitatively analyze the volume 348 and complexity of the mitochondria. Mitochondria in gametes with two flagella and eyespots 349 within the locules of the male and female gametangia in M. cylindricus showed an elongated 350 tubular shape; however, they were almost spherical in the released male and female gametes. 351 The total number and volume of mitochondria contained in the released male gametes were 352 similar to those in the cells before release (Figs. 2a-c and 3a, c, e, Supplementary Table S1). 353 The mitochondrial morphology only changed immediately before release. In a similar analysis 354 of female gametes, the number of mitochondria diminished in the released gametes, and their 355 total volume did not change from the gametes before release (Figs. 2d-f and 3b, d, f). These 356 data suggest that mitochondrial fusion occurs in female gametes during the final stage of 357 gametogenesis.

358 Fission and fusion of mitochondria constitute the mitochondrial dynamics, which are
359 important processes for the maintenance of mitochondrial quality (Varuzhanyan and Chan
360 2020). In particular, fusion promotes the elimination of mitochondrial diversity, such as
361 mitochondrial DNA (mtDNA) and functional heterogeneity (Chan 2012). The fusion of
362 mitochondria is observed at a specific stage of spermatogenesis in *Drosophila melanogaster*363 (Hales and Fuller 1997) in which the mitochondria aggregate and fuse to form the nebenkern, an

364 onion-like giant mitochondrion. Mitochondria elongate during pachytene in meiosis I and 365 fragment again after meiosis in rats (De Martino et al. 1979, Varuzhanyan et al. 2019). 366 Mitochondrial fusion is considered particularly important for the production of healthy sperms. 367 The mitochondrial fusion gene fuzzy onions (Fzo) was isolated from mutants showing male 368 sterility in D. melanogaster (Hales and Fuller 1997). Mammalian homologs of FZO are 369 mitofusins (Mfns) (Santel and Fuller 2001), and defects in the MFN1 and MFN2 genes induce 370 the failure of sperm production (Varuzhanyan et al. 2019). Mitochondrial fusion-related genes 371 were not found in the brown algae. However, based on a study on female gametogenesis in M. 372 cylindricus, mitochondrial fusion occurs immediately before gamete release. In M. cylindricus, 373 mtDNA is derived from female gametes, as reported previously (Shen et al. 2020). The mtDNA 374 copy number in each mitochondrion in a male gamete is less than one-seventh of that in a 375 female gamete. In addition, male mitochondria are digested by mitophagy after fertilization. It is 376 possible that the fusion of mitochondria before the release of the female gametes could be a 377 necessary final quality control step to unify the matrix and ensure the success of the maternal 378 inheritance of mitochondria in this species.

379

380 Mitochondrial complexity and cell types

381 Mitochondria in the vegetative gametophytes and sporophytes showed high MCI scores 382 (Supplementary Table S1). Tubular, curved, V-, and Y-shaped mitochondria observed in these 383 cells were not observed in other cells. The high MCI score for the mitochondria in these cells 384 may be related to the development of large vacuoles within the cells (Figs. 1, 6). In a 20-d-old 385 crustose sporophyte, mitochondrial morphology was analyzed using three cells collected from 386 different regions of the alga. The results showed that the total and individual mitochondrial 387 volumes were similar in all the cells. However, the MCI scores in the subapical and central 388 regions were obviously higher ("20-d-old Sporophyte 2" and "20-d-old Sporophyte_3" in 389 Supplementary Table S1) than in the marginal apical region ("20-d-old Sporophyte 1" in

Supplementary Table S1). It was clear that the cytoplasm in the cell from the subapical region
was occupied by a large vacuole compared to cells in the other regions (Fig. 7a). The complex
morphology of mitochondria may thus be influenced to some extent by the development of
vacuoles.

394

395 Mitochondrial structure in the vegetative and motile reproductive cells

396 Generally, mitochondria in brown algae are tubular. Moreover, in the most motile 397 reproductive cells, the intercristal structures—high electron-density structures—in the 398 mitochondrial cristae are observed, for example, in the male and female gametes of *Cutleria* 399 hancockii (La Claire and West 1978, 1979), Scytosiphon lomentaria (Clayton 1984), and M. 400 cylindricus (Shen et al. 2020); the male gametes of *Ectocarpus siliculosus* (Maier 1997); 401 zoospores of Halopteris filicia (Katsaros and Galatis 1986) and several species belonging to 402 Laminariales (Henry and Cole 1982a); and sperm of Fucus serratus (Berkaloff and Rousseau 403 1979). The function of the intercristal structure has not vet been elucidated, and the presence of 404 this structure is limited in motile reproductive cells, not in eggs (Brawley et al. 1976). The 405 occurrence of this structure was observed in mature, but not immature, gametangia in Pylaiella 406 littoralis (Markey and Wilce 1975). The timing of the appearance of the tubular structure within 407 the mitochondrial cristae may indicate the preparation of a switch from the vegetative to the 408 reproductive type in the mitochondria. In this study, we did not examine the internal structure of 409 the mitochondria. The mitochondria in both male and female gametes changed from a dumbbell 410 or elongated shape to an almost spherical shape immediately before release. Morphological 411 changes at the final stage of gametogenesis may be linked to the completion of optimal 412 mitochondrial physiology in the motility cells. The timing of the disappearance of the 413 intercristal structure after fertilization or settlement of zoospores is unknown. In M. cylindricus, 414 the mitochondrial size and MCI score increased 6 h after fertilization. It is thought that the 415 physiological condition or function of the mitochondria may change from reproductive to

416 vegetative type during this transition.

417

418 Conclusion

419 To understand the morphological dynamics in mitochondria across generations, we 420 performed a quantitative analysis on mitochondria from M. cylindricus based on 3D 421 reconstruction of serial sections imaged by TEM. It became clear that mitochondria in 422 gametophytes and sporophytes showed high MCI scores compared to those in the gametes and 423 during early development of the sporophytes, and this appeared to be related to the development 424 of vacuoles in the cells. In the final stages of gametogenesis, a conspicuous reduction in the 425 number of mitochondria in female gametes was observed. By comparing the total volume and 426 the number of mitochondria in pre- and post-release female gametes, there is a strong possibility 427 that mitochondria in the female gametes fuse immediately before release. This phenomenon 428 suggests that mitochondria fuse to reduce heterogeneity and enhance maternal inheritance of 429 mitochondria upon fertilization in this species. To verify this hypothesis, the continuous 430 sequence of dynamic morphological changes in mitochondria needs to be further investigated. 431 432 433 Acknowledgements 434 We are grateful to Dr. Toyoki Iwao (Toba Fisheries Science Center) for collecting the fresh 435 Mie-strain of *M. cylindricus*, and to Dr. Toshiaki Ito (Electron Microscope Laboratory, 436 Research Faculty of Agriculture, Hokkaido University) for preparing the TEM samples of the 437 gametophyte stage using high-pressure freezing.

438

439

440 Author contributions

- 441 Yuan Shen analyzed the data and maintained the *M. cylindricus* strains. Chikako Nagasato and
- 442 Taizo Motomura designed the experiments and critically reviewed the manuscript. All authors
- 443 have written and edited the manuscript.
- 444

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447

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558	
559	
560	

561 Figure Legends

- 562 Fig. 1 Mitochondrial morphology in the somatic cells of gametophytes.
- **563 a**, **c**, **e** male gametophyte, **b**, **d**, **f** female gametophyte.
- **564 a** A thin-section image of male gametophyte.
- **b** A thin-section image of female gametophyte.
- **c** A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- serial thin sections in the same cell as **a**.
- 568 d A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- serial thin sections in the same cell as **b**.
- **571 f** A complete 3D model of **d**.
- 572 c, chloroplast; pm, plasma membrane; m, mitochondrion; n, nucleus; v, vacuole. Scale bar: 1
- 573 μm.
- 574
- 575 Fig. 2 Mitochondrial morphology in mature gametes from the plurilocular gametangia.
- 576 **a–c** male gamete in the gametangium, **d–f** female gamete in the gametangium.
- **577 a** A thin-section image of male gamete.
- 578 b A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- 579 serial thin sections in the same cell as **a**.
- **580 c** A complete 3D model of **b**.
- **581 d** A thin-section image of female gamete.
- **582** e A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- serial thin sections in the same cell as **d**.
- **584 f** A complete 3D model of **e**.
- 585 c, chloroplast; pm, plasma membrane; e, eyespot; f, flagellum; m, mitochondrion; n, nucleus; v,
- 586 vacuole. Scale bar: 1 μm.

- 588 Fig. 3 Mitochondrial morphology in gametes after release.
- 589 **a, c, e** male gamete, **b, d, f** female gamete.
- **590 a** A thin-section image of male gamete.
- **591 b** A thin-section image of female gamete.
- 592 c A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- serial thin sections in the same cell as **a**.
- **b d** A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- serial thin sections in the same cell as **b**.
- **6 e** A complete 3D model of **c**.
- **597 f** A complete 3D model of **d**.
- 598 Arrowheads indicate the anterior part of the cell.
- 599 c, chloroplast; pm, plasma membrane; e, eyespot; af, anterior flagellum; pf, posterior flagellum;
- 600 m, mitochondrion; n, nucleus; v, vacuole. Scale bar: 1 μm.

- **602** Fig. 4 Mitochondrial morphology in 2- and 6-h-old zygotes.
- **603 a**–**c** a 2-h-old zygote, **d**–**f** a 6-h-old zygote.
- **604 a** A thin-section image of a 2-h-old zygote.
- **b** A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- 606 serial thin sections in the same cell as **a**.
- 607 c A complete 3D model of b.
- **608 d** A thin-section image of a 6-h-old zygote.
- 609 e A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- 610 serial thin sections in the same cell as d.
- 611 **f** A complete 3D model of **e**.
- 612 c, chloroplast; pm, plasma membrane; e, eyespot; m, mitochondrion; n, nucleus; v, vacuole.

- 613 Scale bar: 1 μm.
- 614
- 615 Fig. 5 Mitochondrial morphology in 24-h-old zygotes.
- 616 **a–c** a 24-h-old zygote before cytokinesis, **d–f** a 24-h-old zygote after cytokinesis.
- 617 **a** A thin-section image of a 24-h-old zygote before cytokinesis.
- 618 b A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- 619 serial thin sections in the same cell as **a**.
- 620 c A complete 3D model of b.
- d A thin-section image of the 24-h-old zygote after cytokinesis.
- 622 e A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- 623 serial thin sections in the same cell as **d**.
- 624 **f** A complete 3D model of **e**.
- 625 c, chloroplast; pm, plasma membrane; e, eyespot; m, mitochondrion; n, nucleus; v, vacuole.
- **626** Scale bar: 1 μm.
- 627
- 628 Fig. 6 Mitochondrial morphology in a crustose sporophyte.
- **629 a** A thin-section image of a 20-d-old crustose sporophyte.
- **b** A thin-section image of a longer apical cell located in the marginal meristematic region in **a**.
- 631 c A thin-section image of a shorter apical cell in the marginal meristematic region in a.
- 632 d A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- 633 serial thin sections in the same cell as **c**.
- e A complete 3D model of d.
- 635 **f** A thin-section image of a cell in the subapical region in **a**.
- 636 g A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- 637 the serial thin sections in the same cell as f.
- 638 h A complete 3D model of g.

- 639 i A thin-section image of a cell in the central region in **a**.
- 640 j A partial 3D model of the nucleus, mitochondria, and plasma membrane reconstructed from
- 641 serial thin sections in the same cell as **i**.
- 642 k A complete 3D model of j.
- 643 c, chloroplast; pm, plasma membrane; h, hyaline hair; m, mitochondrion; n, nucleus; v, vacuole.
- 644 Scale bar: $10 \mu m$ in \mathbf{a} ; $1 \mu m$ in $\mathbf{b}-\mathbf{k}$.
- 645
- 646 Fig. 7 Box plot presenting quantitative analysis of each mitochondrial volume examined in the
- 647 mature gametangia, released gametes, one-day-old zygotes, and two-day-old zygotes before and
- 648 after cytokinesis. All mitochondria used for quantitative analysis originated from
- 649 Supplementary Table S1: MPG, Male Gametangium; FPG, Female Gametangium; Mg,
- 650 Male_Gamete; Fg, Female_Gamete; 2hZ, 2-h-old_Zygote; 6hZ, 6-h-old_Zygote; 24hZ, 24-h-
- old_Zygote; 24h2cS, 2-celled_Sporophyte; 20dS, 20-d-old_Sporophyte.
- 652 Within the box plot, the solid line represents the 50th percentile, the dashed line represents the
- mean, the box delimits the 25th and 75th percentiles, and bars indicate the 10th and 90th
- 654 percentiles. Within the line plot, data are mean \pm SEM.
- 655
- 656 Fig. S1 Serial ultrathin sections used for 3D reconstruction
- a Ribbon of thin serial sections generated using a diamond knife. b Enlarged image of a with
- 658 interference color as gold.
- 659
- 660 Fig. S2 Three-dimensional models used for quantitative analysis of mitochondrial morphology
- across gametophyte-to-sporophyte life cycle of *M. cylindricus*. All mitochondria used for
- quantitative analysis originated from Table S1. a1–3 Male_Gametophyte_1–3. b1–3
- **663** Female_Gametophyte_1–3. **c1–3** Male_Gametangium_1–3. **d1–3** Female_Gametangium_1–3.
- 664 e1-3 Male Gamete 1-3. F1-3 Female Gamete 1-3. g1-3 2-h-old Zygote 1-3. h1-3 6-h-

- old Zygote 1–3. i1–2 24-h-old Zygote 1 to 2. k1–4: 2-celled Sporophyte 1–2. l1–3 20-d-
- old_Sporophyte_1 to 3. The plasma membrane is indicated in gray. Nuclei are indicated in blue.
- 667 Mitochondria are indicated in orange. Scale bar: 1 μm.
- 668
- 669 Fig. S3 Gametophyte-to-sporophyte life cycle of *M. cylindricus*
- 670 **a**, **b** Male and female gametophytes. Arrowheads indicate released swimming gametes. **c**, **d**
- 671 Male and female plurilocular gametangia. Black arrowheads indicate orange eyespots in
- 672 gametangia small loci. White arrowheads indicate somatic cells of gametophytes. e, f Released
- 673 swimming male and female gametes with one eyespot (arrowhead) in each cell. g Zygote with
- 674 two orange eyespots (arrowheads). h Two-celled sporophytes with two orange eyespots
- 675 (arrowheads). i Crustose sporophyte.
- af, anterior flagellum; pf, posterior flagellum. Scale bar: 1 cm in **a**, **b**; 10 μm in **c**–**h**.
- 677
- 678 Movie S1 A 3D model of a male gametophyte
- 679
- 680 Movie S2 A 3D model of a female gametophyte
- 681
- 682 Movie S3 A 3D model of a male gametangium
- 683
- 684 Movie S4 A 3D model of a female gametangium
- 685
- 686 Movie S5 A 3D model of a male gamete
- 687
- 688 Movie S6 A 3D model of a female gamete
- 689
- 690 Movie S7 A 3D model of a 2-h-old zygote

691	
692	Movie S8 A 3D model of a 6-h-old zygote
693	
694	Movie S9 A 3D model of a 24-h-old zygote
695	
696	Movie S10 A 3D model of a two-celled sporophyte
697	
698	Movie S11 A 3D model of the marginal apical region in a 20-d-old sporophyte
699	
700	Movie S12 A 3D model of the subapical region in a 20-d-old sporophyte
701	
702	Movie S13 A 3D model of the central region in a 20-d-old sporophyte

	Pixel size	Total number		Total volume (µm ³)		Average volume	e (μm ³)	Average MCI	
Life-cycle stages and	of each	Each sample	Average	Each sample	Average	Each sample	Average	Each sample	Average
sample No.	TEM image		(mean ±		(mean ±	(mean ±	(mean ±	(mean ±	(mean ±
	(nm)		SEM)		SEM)	SEM)	SEM)	SEM)	SEM)
Male Gametophyte 1	3.318	117		11.114		0.095 ± 0.004		3.684 ± 0.172	
Male Gametophyte 2*	3.318	83	83 ± 19	9.123	10.925 ± 0.991	0.110 ± 0.004	0.131 ± 0.006	2.796 ± 0.132	4.848 ± 0.282
Male Gametophyte 3	3.318	50		12.539		0.251 ± 0.016		10.980 ± 0.911	
Female Gametophyte 1*	4.921	104		13.522		0.130 ± 0.005		5.708 ± 0.475	
Female Gametophyte 2	4.921	42	68 ± 19	7.536	9.479 ± 2.022	0.179 ± 0.011	0.139 ± 0.004	6.778 ± 0.380	6.190 ± 0.280
Female Gametophyte 3	4.921	59		7.379		0.125 ± 0.006		6.621 ± 0.408	
Male Gametangium 1*	3.318	7		0.840		0.120 ± 0.007		1.869 ± 1.682	
Male Gametangium 2	3.318	5	6 ± 1	1.024	0.918 ± 0.055	0.205 ± 0.024	0.162 ± 0.013	1.337 ± 0.133	1.682 ± 0.091
Male_Gametangium_3	3.318	5		0.890		0.178 ± 0.020		1.765 ± 0.152	
Female_Gametangium_1*	4.921	40		5.049		0.126 ± 0.007		2.563 ± 1.852	
Female Gametangium 2	4.921	38	38 ± 1	5.109	5.042 ± 0.041	0.134 ± 0.005	0.133 ± 0.004	1.448 ± 0.057	1.852 ± 0.074
Female_Gametangium_3	4.921	36		4.969		0.138 ± 0.006		1.489 ± 0.068	
Male_Gamete_1*	2.086	5		0.763		0.153 ± 0.022		0.744 ± 0.005	
Male_Gamete_2	2.086	5	5 ± 0	0.834	0.867 ± 0.072	0.167 ± 0.046	0.173 ± 0.022	0.745 ± 0.006	0.758 ± 0.007
Male_Gamete_3	2.086	5		1.004		0.201 ± 0.045		0.785 ± 0.014	
Female_Gamete_1	4.214	23		5.486		0.239 ± 0.013		0.980 ± 1.034	
Female_Gamete_2*	4.214	30	27 ± 2	4.937	5.144 ± 0.172	0.165 ± 0.009	0.191 ± 0.006	0.994 ± 0.106	1.034 ± 0.042
Female_Gamete_3	4.214	28		5.010		0.179 ± 0.008		1.121 ± 0.031	
2-h-old_Zygote_1	7.018	25		4.846		0.194 ± 0.010		0.867 ± 0.078	
2-h-old_Zygote_2	7.018	35	29 ± 3	5.995	5.472 ± 0.336	0.171 ± 0.007	0.190 ± 0.006	0.938 ± 0.034	0.901 ± 0.028
2-h-old_Zygote_3*	7.018	26		5.575		0.214 ± 0.013		0.884 ± 0.031	
6-h-old_Zygote_1*	4.243	29		7.859		0.271 ± 0.012		0.887 ± 0.069	
6-h-old_Zygote_2	4.243	30	28 ± 2	6.590	6.802 ± 0.559	0.220 ± 0.010	0.243 ± 0.007	0.857 ± 0.027	0.943 ± 0.031
6-h-old_Zygote_3	4.243	25		5.958		0.271 ± 0.012		0.887 ± 0.069	
24-h-old_Zygote_1*	5.255	33	33 ± 1	13.634	13.734 ± 0.100	0.413 ± 0.030	0.424 ± 0.018	1.308 ± 0.108	1.387 ± 0.065
24-h-old Zygote 2	5.255	32		13.834		0.432 ± 0.023		1.449 ± 0.078	
2-celled_Sporophyte_1*	5.255	31		12.304		0.397 ± 0.024		1.065 ± 0.050	
		(15 & 16)**	33 ± 2	(5.914 & 6.390)**	11.406 ± 0.899		0.370 ± 0.019		1.114 ± 0.047
2-celled_Sporophyte_2	5.255	34	(16±1)**	10.507	(5.703 ± 0.287) **	0.309 ± 0.014		1.575 ± 0.400	
		(18 & 16)**		(5.443 & 5.065)**					
20-d-old_Sporophyte_1*	7.918	36		6.349		0.176 ± 0.008		2.463 ± 0.147	
20-d-old Sporophyte 2*	7.918	39	36 ± 2	7.451	6.651 ± 0.404	0.191 ± 0.010	0.186 ± 0.006	4.180 ± 0.250	3.358 ± 0.138
20-d-old_Sporophyte_3*	7.918	32		6.153		0.192 ± 0.011		3.363 ± 0.203	

Table S1 Quantitative analysis of mitochondrial morphology across gametophyte-to-sporophyte life cycle of M. cyclindricus

*3D images are shown in Figure 1, 2, 3, 4, 5, 6 **The numerical values show the data from each daughter cell or from the zygote before fusion of male and female gamete nuclei (karyogamy)

Table S2 Partial pairwise multiple comparison using Tukey's multiple comparison test

Comparison	Difference of means	SE of difference	95.00% CIs (confidence intervals)	T- value	P- value	P < 0.050
Male_Gametangium vs. Female_Gametangium	0.02931	0.0132	(0.001637,0.05698)	2.221	0.0391	Yes
Male Gametangium vs. Male Gamete	-0.01134	0.0252	(-0.09738, 0.07470)	0.4502	0.6568	No
Female Gametangium vs. Female Gamete	-0.05784	0.007323	(-0.07233, -0.04335)	7.898	< 0.0001	Yes
Male Gamete vs. Female Gamete	-0.01719	0.2268	(-0.06514, 0.03077)	0.7577	0.4593	No
Female Gamete vs. 2-h-old Zygote	0.00192	0.008625	(-0.01511, 0.01895)	0.2226	0.8242	No
Female Gamete vs. 6-h-old Zygote	-0.05241	0.009791	(-0.07175, -0.03307)	5.353	< 0.0001	Yes
2-h-old Zygote vs. 6-h-old Zygote	-0.05433	0.009371	(-0.07284, -0.03582)	5.797	< 0.0001	Yes
6-h-old Zygote vs. 24-h-old Zygote	-0.1809	0.01962	(-0.2201, -0.1418)	9.223	< 0.0001	Yes
24-h-old Zygote vs. 2-celled Sporophyte	0.05434	0.2598	(0.002749, 0.1059)	2.091	0.0392	Yes
2-celled_Sporophyte vs. 20-d-old_sporophyte	0.1831	0.0194	(0.1442, 0.2219)	9.436	< 0.0001	Yes
Morphological complexity index of each mitochond	rion					
Male Gametangium vs. Female Gametangium	-0.1703	0.1172	(-0.5691, 0.2285)	1.453	0.9268	No
Male Gametangium vs. Male Gamete	0.9237	0.09103	(0.5859, 1.261)	10.15	< 0.0001	Yes
Female Gametangium vs. Female Gamete	0.818	0.08504	(0.5406, 1.095)	9.619	< 0.0001	Yes
Male Gamete vs. Female Gamete	-0.2759	0.04216	(-0.4154, -0.1364)	6.545	< 0.0001	Yes
Female Gamete vs. 2-h-old Zygote	0.1201	0.04942	(-0.04170, 0.2819)	2.43	0.3558	No
Female Gamete vs. 6-h-old Zygote	0.09075	0.05178	(-0.07855, 0.2601)	1.752	0.8054	No
2-h-old Zygote vs. 6-h-old Zygote	-0.02934	0.0409	(-0.1629, 0.1042)	0.7173	0.9997	No
6-h-old Zygote vs. 24-h-old Zygote	-0.4432	0.07155	(-0.6811, -0.2053)	6.194	< 0.0001	Yes
24-h-old Zygote vs. 2-celled Sporophyte	0.2731	0.08004	(0.008581, 0.5375)	3.412	0.037	Yes
2-celled_Sporophyte vs. 20-d-old_Sporophyte_1	-2.25	0.2088	(-2.969, -1.531)	10.77	< 0.0001	Yes
2-celled Sporophyte vs. 20-d-old Sporophyte 2	-3.067	0.2548	(-3.935, -2.199)	12.04	< 0.0001	Yes
2-celled Sporophyte vs. 20-d-old Sporophyte 3	-1.35	0.1546	(-1.875, -0.8241)	8.731	< 0.0001	Yes
20-d-old Sporophyte 1 vs.	-1.717	0.2904	(-2.688, -0.7459)	5.913	< 0.0001	Yes
20-d-old Sporophyte 2			, , , ,			
20-d-old Sporophyte 1 vs.	-0.9	0.2510	(-1.741, -0.05895)	3.585	0.0264	Yes
20-d-old Sporophyte 3						
20-d-old Sporophyte 2 vs.	0.8169	0.3225	(-0.2571, 1.891)	2.533	0.3047	No
20-d-old Sporophyte 3						
Data asymptotic componenting with Table S1.						

Data sourceis corresponding with **Table S1**; Overall significance level = 95.00%; When a P < 0.050, there is a statistically significant difference