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Uniparental inheritance of cpDNA and the genetic control of sexual differentiation in *Chlamydomonas reinhardtii*

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Abstract
An intriguing feature of most eukaryotes is that chloroplast (cp) and mitochondrial (mt) genomes are inherited almost exclusively from one parent. Uniparental inheritance of cp/mt genomes was long thought to be a passive outcome, based on the fact that eggs contain multiple numbers of organelles, while male gametes, at best, contribute only a few cp/mtDNA. However, the process is likely to be more dynamic because uniparental inheritance occurs in organisms that produce gametes of identical sizes (isogamous). In *Chlamydomonas reinhardtii*, the uniparental inheritance of cp/mt genomes is achieved by a series of mating type-controlled events that actively eliminate the mating type minus (mt-) cpDNA. The method by which *Chlamydomonas* selectively degrades mt-cpDNA has long fascinated researchers, and is the subject of this review.

Key Words
cpDNA, uniparental inheritance, chlamydomonas, sexual differentiation

Introduction
The first report on non-Mendelian inheritance appeared in 1909, just 9 years after the rediscovery of Mendel’s laws. Carl Correns followed the inheritance of green, variegated, and white color patterns in reciprocal crosses of the four-o’clock (*Milabilis jalapa*), and observed that the seedling resembled the maternal parent in phenotype. Simultaneously, Baur found that the green and variegated patterns of the leaves were
inherited in a biparental fashion in *Pelargonium*. During the time of th Correns and Baur studies Correns and Baur, however, no one knew the nature of the genetic substances that defied Mendel’s law. Extensive studies over the last century, using numerous techniques, including electron microscopy, genetics, molecular biology and biochemistry, have revealed that the genetic substances are in fact genomes within chloroplasts and mitochondria (Kuroiwa 1991; Birky 1995).

Chloroplasts (cp) and mitochondria (mt) are thought to have arisen from ancestral endosymbiotic relationships between nucleated cells and free-living bacteria – cyanobacteria and alpha-purple bacteria, respectively. They contain their own genomes, which are presumably vestiges from their progenitors (Gray 1992). Today, we know that cp and mt genes are transmitted to the progeny solely from the maternal parent in the diverse taxa of higher plants, ferns, mosses, algae (Kuroiwa 1991), fungi (Mitchell and Mitchell 1952; Kawano et al. 1987), and animals (Hutchison et al. 1974), including humans.

The uniparental inheritance of cp/mt genomes was long thought to be a passive outcome based on the fact that eggs contain multiple numbers of organelles, while male gametes, at best, contribute only a few (Gyllensten et al. 1991). However, the process of uniparental inheritance is likely to be more dynamic. A classic and striking example would be the occurrence of non-Mendelian inheritance in the unicellular green algae, *Chlamydomonas reinhardtii*, which produces gametes of identical sizes (isogamous) (Sager 1954).

The life cycle of *C. reinhardtii* is exquisitely simple. There are two mating types of *C. reinhardtii*, mating type plus (mt+) and mating type minus (mt−), controlled by a single complex mating type loci on linkage group VI (Ferris et al. 2002). *C. reinhardtii* undergoes a sexual life cycle, which includes defined stages of differentiation (Figure 1).

Vegetative cells differentiate into gametes under conditions of nitrogen starvation and light irradiation (Pan et al. 1996). Within minutes of being mixed, gametes of opposite mating types adhere to each other by their flagella and fuse to form zygotes. After a mandatory period of dormancy, the zygote undergoes meiosis and germination to produce four haploid progeny. More than 90% of the progeny that are formed inherit chloroplast (cp) traits, preferentially from the mt+ parent, a phenomenon first described more than 50 years ago (Sager 1954).
The dawn of research on the uniparental inheritance of cpDNA.
Sager described the inheritance patterns of two UV-induced mutations, \textit{sr1} and \textit{sr2}. The \textit{sr1} mutation confers resistance to low levels of the antibiotic streptomycin, and the \textit{sr2} mutation confers resistance to high levels of streptomycin. When \textit{sr1} was crossed to a streptomycin sensitive strain, low levels of streptomycin resistance were inherited, following Mendel’s law. In contrast, when a \textit{mt+} parent carrying the \textit{sr2} mutation was crossed to a sensitive \textit{mt-} parent, all of the meiotic progeny were resistant to high levels of streptomycin. In the reciprocal cross, the meiotic progeny were all streptomycin sensitive (Sager 1954). In 1962, evidence for the existence of cpDNA came from light and electron microscopic work by Ris and Plaut (Ris and Plaut 1962). Just one year later, Sager and Ishida described the isolation of cpDNA by cesium chloride (CsCl) density gradient centrifugation (Sager and Ishida 1963). In 1989, the \textit{sr2} mutation was shown to localize within the \textit{rps12} gene of the cp genome (Liu et al. 1989).

Biochemical and molecular biological evidence for the preferential reduction of \textit{mt-} cpDNA
In 1972, one biochemical study showed that the amount of \textit{mt-} cpDNA decreased relative to \textit{mt+} cpDNA, 6-24 h after mating (Sager and Lane 1972). The DNA from \textit{mt+} and \textit{mt-} gametes was labeled with either $^{14}$N- or $^{15}$NH$_4$Cl, and CsCl density gradient centrifugation was used to monitor the fate of nuclear DNA and cpDNA in 6- and 24-h zygotes. Six hours into zygote development, the signal representing \textit{mt-} cpDNA was clearly lower than the \textit{mt+} cpDNA, indicating a preferential reduction of \textit{mt-} cpDNA relative to \textit{mt+} cpDNA. In 1980, the first molecular evidence for the preferential reduction of \textit{mt-} cpDNA was provided by Grant et al. (Grant et al. 1980). The authors monitored the behavior of \textit{mt+} and \textit{mt-} cpDNA, taking advantage of restriction fragment length polymorphisms (RFLPs), in a \textit{C. reinhardtii} mutant strain \textit{ac-u-g-23} that carries two small deletions in its chloroplast DNA. The authors found that both the deletions in cpDNA and the non-photosynthetic phenotype were uniparentally inherited.

Active digestion of \textit{mt-} cp nucleoids observed by fluorescent microscopy
The 203-kb chloroplast genome of \textit{C. reinhardtii} (Maul et al. 2002) is present at ~80–100 copies per cell, and is organized into 5–10 DNA–protein complexes, which are called chloroplast nucleoids (Kuroiwa et al. 1981). In 1982, Kuroiwa et al. found that
DAPI (dsDNA specific fluorochrome, 4',6-diamidino-2-phenyindole)-stained mt- cp nucleoids disappeared preferentially in young zygotes within 50 min of mating (Kuroiwa et al. 1982). In 1999, the preferential disappearance of mt- cp nucleoids was observed in a living zygote, using SYBR Green I (dsDNA specific fluorochrome that can permeate into living cells) (Figure 1) (Nishimura et al. 1999).

The interpretation of this dramatic phenomenon, however, was controversial because the preferential disappearance of fluorescent mt- cp nucleoids occurred well before DNA reduction was detected by biochemical or molecular biological methods (6–24 h after mating) (Sager and Lane 1972). Two primary possibilities were proposed to explain this. One possibility was that the disintegration of cp nucleoids might lead to the dispersion of cpDNA molecules, and the second possibility was that the rapid digestion of cpDNA molecules might lead to the disappearance of cpDNA nucleoids.

**Single cell analysis using the optical tweezers**

One problem in addressing this question is that the mating reaction is performed using millions of mt+ and mt- gametes (Figure 2). The cell population is inevitably a heterogeneous mixture of cells, such as unmated mt+ and mt- gametes, zygotes with or without mt- cp nucleoids, and exceptional meiotic zygotes (1~5%) that do not form meiotic zygotes (Ebersold 1967). Generally, molecular and biochemical methods require large amounts of homogeneous samples for precise analyses, and any heterogeneity in the samples would confuse the results. In other words, the “personalities” of individual cells or organelles within a population would be diluted and most likely lost in the process of analyses. On the other hand, microscopy can reveal the “personalities” at the morphological level, but not at the molecular level.

For studying the condition of cpDNA molecules during the disappearance of mt- cp nucleoids, it was necessary to collect zygotes based on the presence or absence of mt- cp nucleoids, and to analyze the individual zygotes using molecular biological techniques. To achieve this, optical tweezers were employed (Figure 3). The use of optical tweezers is a novel technique for manipulating living cells or organelles under direct microscopic observation (Ashkin et al. 1987). In this study, a single zygote with or without cp nucleoids was collected using optical tweezers, and the presence or absence of mt- cpDNA molecules was determined by nested PCR analysis. The individual fates of mt+ and mt- zygotic cpDNA were followed separately using a
chloroplast transformant LO3c, which harbors the bacterial gene aadA (aminoglycoside adenyl transferase).

The single zygotes that were obtained with the optical tweezers were subjected to highly sensitive nested-PCR analysis for aadA (Figure 4)(Nishimura et al. 1999). When LO3c mt+ gametes were crossed with wild-type mt- gametes, aadA gene sequences were detected in all of the zygotes that were examined. On the contrary, when the LO3c mt-gametes were crossed with wild-type gametes, the aadA sequences were only amplified in younger zygotes (10 and 30 min after zygote formation). After the fluorescent mt- cp nucleoids disappeared, the aadA sequences were no longer detected in the zygotes (90 and 120 min after zygote formation). These results indicate that the mt- cpDNA molecules are completely digested in 10 min, during which the mt- cp nucleoids disappear, and also that at least one highly effective nuclease is activated in the mt-chloroplast just after zygote formation. This active digestion of mt- cpDNA is probably the basis for maternal inheritance of cpDNA.

**Destroyer-protector model for the active digestion of mt- cpDNA**

The simplest model for the uniparental inheritance of cpDNA is that the process consists of two distinct events that are likely to occur at different stages of the life cycle: a “protection” of mt+ cpDNA, perhaps during gametogenesis, and a “destroyer” of unprotected mt- cpDNA during early zygote development.

A) Restriction–Methylation hypothesis

In 1972, Sager and colleagues proposed that mt- cpDNA was digested by the action of restriction enzymes, whereas the mt+ cpDNA was protected by methylation – a model analogous to the bacterial restriction-methylation system (Sager and Lane 1972). Sager and colleagues rapidly accumulated convincing evidence that shows an increase in the methylation level of mt+ cpDNA, which was detected 7 h after mating (Burton et al. 1979; Royer and Sager 1979; Sano et al. 1980). Furthermore, the purification of mt+ gamete-specific DNA methyltransferases, with molecular weights of 60 kDa and 20 kDa, has been reported (Sano et al. 1981). This mt+ gamete specific methylation event was apparently reversible, as would be expected for protection (Sano et al. 1984). The gene for the chloroplast-resident DNA methyltransferase was finally identified in 2002, and its mt+ gamete-specific expression and chloroplast localization were confirmed
(Nishiyama et al. 2002).

On the other hand, a series of papers from independent groups has subsequently argued that methylation of mt+ cpDNA could not explain protection adequately. Bolen et al. isolated a nuclear mutant mel that constitutively methylates cpDNA at a higher level in both mt+ and mt- cells (Bolen et al. 1982). When mel gametes were used for crosses, normal uniparental inheritance patterns were observed, inconsistent with the restriction-methylation hypothesis. In 1984, Feng and Chiang presented evidence that treatment of cells with the methylation inhibitors, L-Ethionine and 5-azacytidine, resulted in a hypomethylation of gamete cpDNA (Feng and Chiang 1984). The hypomethylation of cpDNA had no effect on the uniparental inheritance of cpDNA, which was also inconsistent with the restriction-methylation hypothesis.

In 2001, Umen and Goodenough reported that 5-aza-2’-deoxycytidine did alter the inheritance pattern of cpDNA (Umen and Goodenough 2001). The authors did not observe destruction of hypomethylated mt+ cpDNA, as a simple restriction–methylation hypothesis would predict, but instead observed that mt+ cpDNA persisted until zygote germination, at which point it failed to replicate at normal levels. The authors proposed that germination is the critical time-point at which unmethylated, or damaged cpDNA, is finally destroyed. According to their discussion, Feng and Chiang (1984) failed to observe the effects of the inhibitors because they were added just prior to gametogenesis or mating in their experiments. Umen and Goodenough presumed that the inhibitor treatment in their experiments might have been too short to disturb critical methylation patterns. In Umen’s experiment, the cells were grown on agar plates supplemented with inhibitors, which should have altered the methylation of the cp genomes effectively, although the methylation status of nuclear genes may also have been affected.

In 2004, Nishiyama et al. expressed chloroplast-resident methyltransferase ectopically in mt- cells, and crossed the cells to normal mt+ cells (Nishiyama et al. 2004). The frequency of paternal or biparental transmission of cpDNA increased from 5–10% (in a wild type cross) to ~23%. This observation indicates that the protection of mt+ cpDNA cannot be explained solely by the methylation status of cpDNA, although it is possible that methylation may play a role in guaranteeing the strict uniparental inheritance, such as by enhancing the replication of mt+ cpDNA.
B) Nuclease C

One possible explanation for the active digestion of mt-cpDNA is that a nuclease is either activated or synthesized during early zygote development, and subsequently degrades the unprotected mt-cpDNA. Kuroiwa and colleagues set out to identify nucleases that are specifically required for the uniparental inheritance of cpDNA. Rather than using whole cell extracts from vegetative cells, Ogawa and Kuroiwa began with zygotes. They identified a class of Ca\(^{2+}\)-dependent nucleases that displayed both endo- and exo-nucleolytic activity. The extracts were composed of six small polypeptides, which were collectively designated as nuclease C (Ogawa and Kuroiwa 1985b). Because Ca\(^{2+}\) is required for the preferential digestion of mt-cpDNA (Kuroiwa 1985), it was proposed that some of these nucleases might engage in preferential digestion. However, no differences in nuclease activity were detected between the gametes and zygotes, and it was not possible to identify the nucleases responsible for cpDNA digestion (Ogawa and Kuroiwa 1985b). On the other hand, an experiment with partially purified Nuclease C provided an important insight into the protection of mt+ cpDNA. Ogawa and Kuroiwa showed that the mt+ cp nucleoids of mature mt+ gametes were resistant to Nuclease C, whereas the cp nucleoids of mature mt- gametes and vegetative cells of both mating types were rapidly digested by treatment with Nuclease C (Ogawa and Kuroiwa 1985a). This result indicates that Ca\(^{2+}\)-dependent nucleases may be responsible for preferential digestion, and that mt+ cpDNA is specifically protected during mt+ gametogenesis by unknown mechanisms.

One problem of the previous studies on Nuclease C might be that nuclease activities were studied only by SDS-PAGE / in gelo assay, which may artificially activate or inactivate nucleases. To eliminate this problem, a native-PAGE was employed, and changes in the nuclease activities were reexamined in mt+ and mt- chloroplasts that were isolated from zygotes 30, 60, and 90 min after mating (Nishimura et al. 2002). As shown in Figure 5, a novel nuclease activity was identified that only appeared in the mt-zygotic chloroplasts; this nuclease activity would be a good candidate to act as a driving force for the active degradation of mt- cpDNA. Research to identify the gene that encodes this nuclease activity is now underway. Based on these observations, our current model is summarized in Figure 6.

C) A surveillance system may monitor the cpDNA copy number in uniparental
inheritance

Further insight into how the cpDNA content of the mt+ parent influences uniparental inheritance has been obtained from studies on the mat3 mutant. The mat3 mutation was isolated in a screen to identify mt+-specific genes required for the uniparental inheritance of cpDNA (Gillham et al. 1987). The mat3 mutation is linked to the mt+ locus, and when present in the mt+ parent, prevents the elimination of mt- cpDNA in zygotes, while it has no effect on mtDNA inheritance. The size of mat3 vegetative cells and gametes are much smaller than those of the WT, and contain a limited amount of cpDNA (Armbrust et al. 1995).

When chlamydomonas cells are treated with 5-fluorodeoxyuridine (FdUrd), a specific inhibitor of cpDNA replication, the amount of cpDNA is reduced dramatically (Wurtz et al. 1979). When FdUrd-treated mt+ gametes are mated to WT mt- gametes, a higher frequency of biparental inheritance is observed. Based on these two observations, the mat3 mutation is hypothesized to prevent the degradation of mt- cpDNA as a secondary consequence of the reduced amount of mt+ cpDNA (Armbrust et al. 1995). Armbrust et al. proposed that chlamydomonas might be able to count the intact cpDNA copy number, and inhibit mt- cpDNA elimination when the amount of mt+ cpDNA is below a threshold level.

The causative gene for the mat3 phenotype has been identified as a retinoblastoma protein homolog (Umen and Goodenough 2001). The retinoblastoma protein is the prototype tumor suppressor, and plays a key role in regulating cell cycle entry and cell size (Butitta and Edgar 2007). Intriguingly, a retinoblastoma related protein was identified as a gender specific gene that is likely be related to sexual differentiation in Volvox carteri (Kianianmomeni et al. 2008).

Active digestion of male cp/mtDNA as one of the universal mechanisms of uniparental inheritance in eukaryotes

Active digestion of mt nucleoids is observed not only in chlamydomonas but also in animals. Shortly after a mouse egg is fertilized, male mtDNA can be PCR amplified from egg cytoplasm, whereas later male mtDNA cannot be detected (Kaneda et al. 1995).
In 2006, using Japanese medaka (Oryzias latipes) as a model system, mt nucleoids were visualized during spermatogenesis and fertilization by vital double staining with the DNA-specific fluorochrome SYBR Green I and a mitochondrial membrane-specific dye MitoTracker (Nishimura et al. 2006). The number of mt nucleoids was reduced dramatically during spermatogenesis. Upon fertilization, rapid disappearance of sperm mt nucleoids was observed in apparently intact mitochondria. Furthermore, the nested PCR analysis using a single sperm, with or without mt nucleoids, collected by the optical tweezer revealed that mtDNA is degraded along with the disappearance of mt nucleoids (Figure 7) (Nishimura et al. 2006). In addition to the examples above, active digestion of uniparental mt (cp) DNA has been observed in various species, including higher plants (Nagata et al. 1999; Liu et al. 2004), algae (Kuroiwa 1991), and true slime mold (Moriyama and Kawano 2003), indicating that active digestion of male cp/mtDNA is one of the universal mechanisms to achieve uniparental inheritance in eukaryotes. Intriguingly, studies with other mammals show that this phenomenon involves the ubiquitin-dependent proteolysis of mitochondrial membrane proteins (Sutovsky et al. 1999; Thompson et al. 2003)

**Genetic control of sexual differentiation and uniparental inheritance in C. reinhardtii**

In 1967, Sager and Ramanis showed that brief UV irradiation of mt+ gametes just prior to mating prevents the selective elimination of mt- cpDNA in zygotes, and thus enhances the frequency of biparental inheritance of cpDNA. The same treatment of mt- gametes had no effect on the transmission of cpDNA (Sager and Ramanis 1967). On the other hand, if the UV irradiated mt+ gametes were allowed to recover in light prior to mating, the frequency of exceptional zygotes decreased. Sager and Ramanis hypothesized that UV irradiation blocked the synthesis of a mt+ gamete specific gene product that is necessary for the elimination of mt- cpDNA in zygotes. In this section, our current understanding of the molecular mechanism that control gametogenesis, and zygote maturation is summarized (Figure 8).

A) Genetic control of gametogenesis

Upon the removal of nitrogen, vegetative cells undergo two critical programs
(Goodenough et al. 2007). First, the cells acclimate to nitrogen starvation through a variety of metabolic changes, including up-regulation of proteins required for transport and reduction of nitrate (Fernandez and Galvan 2008), extra-cellular nitrogen-scavenging enzymes (Vallon et al. 1993), and a reduction in photosynthetic activity through a selective destabilization of $b_{nf}$ complexes (Bulte and Wollman 1992). Secondly, the cells express a gamete differentiation program, which include the formation of sex-specific agglutinin proteins, responsible for flagellar adhesion (Adair et al. 1982), and the formation of sex-specific mating structures (Detmers et al. 1983; Ferris et al. 1996). The blue light is thought to be important to complete gametogenesis (Huang and Beck 2003; Huang et al. 2004). Although a large number of gamete-specific genes have been reported so far (von Gromoff and Beck 1993; Merchan et al. 2001; Abe et al. 2004), only some of these genes are summarized in this review.

Mating type is under the control of the MT locus, a region of suppressed recombination that spans nearly ~1 Mb on linkage group VI (Ebersold 1967; Ferris et al. 2002). The central rearranged domain (R) is flanked by centromere proximal (C) and telomere proximal (T) sequences.

It has been shown that when a mt+/mt- heterozygous vegetative diploid is nitrogen-starved, the diploid mates as a mt-, which demonstrates the dominance of mt- over mt+ (Ebersold 1967). The MID gene, unique to region f of the mt- locus, is responsible for the minus dominance (Ferris and Goodenough 1997). The MID protein has a bZIP motif, and is related to the RWP-RK family that also includes NIT2, a transcription factor involved in nitrate uptake and metabolism (Camargo et al. 2007). Cells expressing a MID gene differentiate as mt-; a loss of function mutation or a deletion of MID prevents the cells from differentiating as minus, and instead results in a “pseudo-plus” phenotype, indicating that MID encodes a transcription factor that both turns on mt- specific genes and turns off mt+ specific genes (Ferris and Goodenough 1997). Recently, a MID ortholog has been identified in the oogamous volvocacean Pleodorina starrii (Nozaki et al. 2006) and the isogamous Gonium pectorale (Hamaji et al. 2008), indicating the significance of the MID gene in the evolutionary process of sex determination.

The autosomal SAD1 gene encodes the minus agglutinin at the mt- mating locus, and the autosomal (LGVIII) gene SAG1 encodes the plus agglutinin, whose expression is probably under the control of the MID gene. The agglutinins are enormous fibrous hydroxyproline-rich glycoproteins (HRGPs) that are responsible for the flagellar...
adhesion between mt+ and mt- gametes (Ferris et al. 2005). The initial adhesion of the flagella of mt+ and mt- gametes triggers a complex signaling pathway, which includes a protein tyrosine kinase (Wang and Snell 2003), a cGMP-dependent protein kinase (Wang et al. 2006), adenylyl cyclase(s) (Saito et al. 1993), and an increase in intracellular cAMP. Both of the gametes shed their cell walls through the action of a metalloprotease (Kubo et al. 2001). Then, the mating structures located on the apical cell membranes between the two flagella of both gametes are activated (Detmers et al. 1983). Continued flagellar adhesion brings the activated mating structures close to each other. The mt+ gamete-specific protein FUS1, present on the plasma membrane of the mating structure, is essential for pre-fusion attachment between the plus and minus mating structures (Misamore et al. 2003). Complete cell fusion requires an additional factor, GCSI (GENERATIVE CELL SPECIFIC) / HAP2, on the mt- mating type structure, which is conserved ubiquitously in algae, higher plants, and protists (Mori et al. 2006; Liu et al. 2008).

After the cell fusion, two homeoproteins, GSP1 (Kurvari et al. 1998) and GSM1, contributed by the mt+ and mt- gametes respectively, physically interact and translocate from the cytosol to the nucleus upon zygote formation, initiating the zygote development program (Lee et al. 2008). Intriguingly, GSP1/GSM1 dyads are similar and functionally related to the KNOX/BELL dyads that regulate meristem specification in land plants. This prompted Lee et al. to propose that combinatorial homeoprotein–based transcriptional control may have originated in a sexual context, and enabled the evolution of land-plant body plans (Lee et al. 2008).

B) Genetic control of zygote maturation

In 1988, Nakamura et al. showed that the expression of at least six zygote polypeptides was inhibited when the mt+, but not the mt-, parent was exposed to brief UV irradiation (Nakamura et al. 1988). This experiment showed, for the first time, that UV treatment of mt+ gametes could inhibit the expression of a subset of zygote-specific genes. Several independent attempts to identify zygote-specific genes have been made (Ferris and Goodenough 1987; Wegener and Beck 1991; Uchida et al. 1993; Kubo et al. 2008). The list of the zygote-specific genes that has been obtained has become an excellent resource for insight into the molecular mechanism of zygote formation and maturation.

B-1) EZY1 (Early Zygote 1):
Ezy1 is tandemly repeated 7-8 times at both the mt+ and mt- mating loci, and the repeats are transcribed immediately upon zygote formation (Armbrust et al. 1993). Ezy1 was found in a differential screen for zygote-specific genes (Ferris and Goodenough 1987). The EZY1 protein localizes specifically to both mt+ and mt- cp nucleoids. Brief irradiation of mt+ gametes prior to mating inhibited the transcription of ezy1, whereas the same treatment of mt- gametes had no effect on ezy1 transcription. Polypeptide 4 was originally identified by Nakamura et al. as a zygote polypeptide whose expression is UV-sensitive (Nakamura et al. 1988), and likely corresponds to EZY1. Therefore, based on this evidence, the ezy1 gene has been hypothesized to be involved in the selective elimination of mt- cpDNA. However, the detailed function of EZY1 is still unknown.

B-2) ZYS 1, 3

Zys1 is a pair of invertedly repeated genes (zys1A and zys1B) that are 98 % identical, and encode 22 kDa polypeptide with a Zn finger motif (Uchida et al. 1993; Uchida et al. 1999). The gene product localizes to the cell nucleus, and gene expression begins at 10 min and lasts until approximately 3 h after mating. Importantly, when mt+ gametes are UV-irradiated, accumulation of the ZYS1 protein is considerably reduced, suggesting its involvement in the uniparental inheritance of cpDNA. Zys3 was identified as an early zygote-specific gene by Uchida et al. (Uchida et al. 1993), and was then carefully characterized by Kuriyama et al. (Kuriyama et al. 1999). The ZYS3 bears two types of protein-protein interaction motifs: the ankyrin repeats and the WW domain. ZYS3 localizes to the ER from 10 min to 6 h after mating. Kuriyama et al. speculated that ZYS3 might regulate the ER functions in the processing, sorting and targeting of rigid, zygote-specific cell wall materials, and/or sticky extracellular matrices. In an extensive macro-array analysis for zygote-specific genes (Kubo et al. 2008), a gene (EZY9) which is highly similar to ZYS3 was identified, implying that there is an interaction between these two gene products.

B-3) Zygote cell wall genes

After 2-3 h of mating, the zygotes proceed to develop new cell walls that are denser and thicker than those of vegetative cells, and that are insoluble in chaotropes and SDS. Eventually, the zygotes become adhesive so that if they are maintained in liquid, they adhere in flat sheets, forming a structure called a “pellicle”. During this process, Chlamydomonas cells switch back and forth between two differential developmental
programs expressed by vegetative cells and by gamete/zygotic cells, which features hydroxyproline rich glycoproteins (HGRPs), which are the major proteins of the cell wall. The zygote-specific genes that are involved in this process are ZSP1 (Woessner and Goodenough 1989) and ZSP2 (Suzuki et al. 2000). Other HRGPs (GAS28, 30) and glycoproteins (GAS31) are expressed in the late phase of gametogenesis and are further up-regulated in zygotes (Hoffmann and Beck 2005). Kubo et al. identified candidate zygotic genes that may be related to cell wall synthesis thorough sugar metabolism (EZY4: UDP-glucose 4-epimerase like protein, EZY11: UDP-glucose protein: protein trans glycosylase, EZY12: UDP-glucose 6-dehydrogenase, EZY16: cell wall protein pherophorin-C15) (Kubo et al. 2008). Since ezy11, 12, and 16 were up-regulated without the cell fusion, the authors proposed that these genes may be involved in cell wall synthesis of both vegetative / gamete cells and zygotes.

Conclusion
In this review, our current mechanistic model for the uniparental inheritance of cpDNA, and the molecular mechanisms controlling the sexual differentiation of Chlamydomonas, have been summarized with a historical overview of previous studies. Despite the abundance of information, our understanding of the molecular mechanism of uniparental inheritance is still at a very preliminary stage. For the next several decades, it will be of crucial interest to understand the regulatory mechanism of uniparental inheritance in the context of the complex molecular system of sexual differentiation of Chlamydomonas.

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53–65.


Suzuki L, Woessner JP, Uchida H, Kuroiwa H, Yuasa Y, Waffenschmidt S,


Fig. 1

(a) The sexual life cycle of *Chlamydomonas reinhardtii* consists primarily of four critical stages – gametogenesis, zygote formation, zygote maturation (zygospore formation), and meiosis (zygospore germination). Gametogenesis is induced by the loss of a nitrogen source and light. When gametes of opposite mating types are mixed together, flagellar adhesion triggers gamete activation that leads to the cell-cell fusion (zygote formation). Zygotes will develop into zygospores with a thick cell wall, which is a dormant stage in the life cycle. The cycle begins again when the appropriate environmental conditions stimulate the dormant zygote to undergo germination, in order to produce new haploid mt+ and mt- haploid cells. During this process, cpDNA and mtDNA molecules are uniparentally inherited from mt+ and mt- parents, respectively.

(b) Preferential disappearance of mt- cp nucleoids was visualized in a living zygote of *C. reinhardtii*. The living zygote was stained with SYBR Green I. Phase-contrast (i) and fluorescent images of an identical zygote before (ii) and after (iii) the
preferential disappearance are shown. The mt+ (Left) and mt- (Right) chloroplasts emit red autofluorescence. The cell nucleus (N), cp nucleoids (big fluorescent spots; large arrow), and mitochondrial nucleoids (small fluorescent spots; small arrow). The cp nucleoids in the mt- chloroplasts (ii, white arrowheads) disappeared completely within 10 min (ii and iii). (Nishimura et al. 1999)
The discrepancy between the preferential disappearance of mt- cp nucleoids (Cytological observation: ~50 min after mating (Kuroiwa et al., 1982)) and the relative reduction of mt- cpDNA (6~24 h after mating (Sager and Lane, 1972)). Due to the time lag between these two independent observations, the interpretation of the preferential disappearance of mt- cpDNA was controversial. Two primary explanations were proposed: 1) disintegration of cp nucleoids caused diffusion of cpDNA molecules 2) destruction of the cpDNA molecule. In order to determine the correct explanation, a single zygote with or without mt- cp nucleoids must be collected for molecular analysis, since the time lag between 50 min and 6~24 h may have been caused by the heterogeneous nature of the cell population during the mating reaction.
Fig. 3

(a) The optical tweezer (i). By using the IR (infra-red) laser, one chlamydomonas cell in the cell population was selectively manipulated (ii-iv). BY-2 protoplasts arranged into “P” (v). Chlamydomonas cells arranged into “C” (vi). DAPI-stained
mitochondria isolated from true slime mold were arranged into “Y” (vii).

(b) The mechanism of laser trapping. When a cell refracts the laser beam (gray sphere), the cell will receive forces (F1 and F2) from the laser beam, and the resultant force is F (red arrow). As the F is directed toward the center of the focal point, the cell is eventually trapped at the center of the laser focus.

(c) The optical isolation process and the “isolation microchamber.” The microchamber is formed by separation of a large (5 × 30 × 0.15 mm, length × width × thickness) and small (9 × 3 × 0.15 mm) coverslip with two thin strips of adhesive tape (9 × 1 × 0.1 mm). The inside dimensions of the chamber were 9 × 1 × 0.1 mm. A small scratch was made on the chamber with a diamond knife. The bottoms of 5-cm diameter Petri dishes were cut out with a knife and replaced with thin, plastic film (~100-μm thick). Filter paper was cut and placed into the dishes, and roughly 200 μl of distilled water was added dropwise to keep the inside of the dish moist. The chamber was then placed inside the dish and attached to the plastic film by using one drop of distilled water. First, the chamber was filled with sterile buffer containing 1.5% sucrose and 0.1% BSA (i). The BSA was added to prevent cells from adhering to the glass chamber. Then, 0.5 μl of the cell suspension was carefully applied to one end of the chamber (ii). The cells were observed with a microscope, and a single cell was trapped with the optical tweezers and transferred to the opposite end of the chamber (iii). The transfer was processed automatically with a microscopic stage control system (MCU26 X, Y, Z-Axes Motor Control; Zeiss) at a velocity of 1–60 μm/sec. When the cell moved past the scratch in the chamber, the chamber was cut immediately, and the piece containing the cell of interest was placed in a PCR tube (iv). The PCR tube was centrifuged for 5 sec to drop the cell into the tube. (Nishimura et al. 1999)
Fig. 4
Nested-PCR amplification of *rbcL* (cpDNA), *cox I* (mtDNA), and *aadA* sequences from one optically isolated gamete or zygote 10, 30, 90, and 120 min after zygote formation. One zygote resulting from the crosses *L03c mt+ × 137c mt−* or *137c mt+ × L03c mt−* was isolated by using the optical tweezers, and was then immediately subjected to nested PCR analysis. One typical zygote was isolated from each of the cell cultures 10, 30, 90, and 120 min after mating. To ensure reproducibility, each experiment was repeated five times (I–V). The lanes marked M are loaded with marker. (Nishimura et al. 1999)
Fig. 5

(a) Phase-contrast (i, iii, v, vii, ix, xi, xiii, xv, xvii, xviii) and fluorescent (ii, iv, vi, viii, x, xii, xiv, xvi, xix, xx) images of SYBR Green I-stained gametes (mocA84 mt+ [i, ii, v, vi]; 137c mt- [iii, iv, v, vii]), zygotes (30 min [ix-xii]; 90 min [xiii-xvi] after mating), and mt+ (xvii, xix) and mt- (xviii, xx) chloroplasts isolated from zygotes at 30 min after mating. Gametes that were incubated for 10 d accumulated large quantities of starch granules in their chloroplasts (st+; i, ii, vii, viii), whereas gametes that were incubated for 2 d accumulated few starch granules (st-; iii, iv, v, vi). By crossing the two different kinds of gametes, zygotes were formed that accumulate large numbers of starch granules, only in mt+ chloroplasts (ix, x, xiii, xiv) or mt- chloroplasts (xi, xii, xv, xvi). The presence of a large amount of starch did not affect the active digestion of mt-cpDNA (xiii-xvi). Chloroplasts containing large amounts of starch granules were isolated and selectively separated by Percoll step-gradient centrifugation. The purity of separated chloroplasts was verified by moc-type single cp nucleoid (mt+) or WT cp nucleoids (mt-) (xvii-xix). (St) Starch granules. (b) Comparison of Ca\(^{2+}\)-dependent nuclease activity in mt+ and mt- chloroplasts using the native-PAGE/in gelo assay. The mt+ and mt- chloroplasts that were isolated from zygotes at 30, 60, and 90 min after mating were analyzed. Experiments were repeated twice to ensure reproducibility. As a control, CBB-stained RuBisCO bands are shown under each lane. (Nishimura et al.)
A model for the molecular mechanism of uniparental inheritance in *C. reinhardtii* that is based on the active digestion of *mt*- cpDNA by MDN. In vegetative cells, MDN is absent or inactivated in both mating types. During gametogenesis, MDN is synthesized or activated only in *mt*+ cells. At the same time, *mt*+ cpDNA becomes resistant to the action of MDN. During gamete fusion, MDN obtains access to unprotected *mt*- chloroplasts and digests *mt*- cpDNA, leading to the uniparental inheritance of cpDNA. Several factors may mediate the successful digestion of *mt*- cpDNA after zygote formation: (1) entry of MDN into *mt*- chloroplasts; (2) efficient access of MDN to cpDNA molecules; and (3) an increase in Ca$^{2+}$ inside *mt*- chloroplasts. Zygote-specific gene expression may be crucial to these processes. (Nishimura et al. 2002)
Fig. 7
Active digestion of sperm mtDNA after fertilization. (A) Phase-contrast images (a and d), SYBR green I-stained images (green) (b and e), and SYBR green I/MitoTracker CMTMRos double-stained images (red) (c and f) of sperm before (a–c) and 60 min after (d–f) fertilization. a and b are the identical sperm. d and e are also identical. Sperm mt nucleoids completely disappeared 60 min after fertilization (e and f). The mitochondrial structure visualized by phase-contrast microscopy (d) or by MitoTracker CMTMRos staining (f) remained intact, even after the disappearance of fluorescent mt nucleoids. (C) Single sperm, with (+) or without (-) mt nucleoids, were selectively extracted from fertilized eggs by using optical tweezers, and were analyzed by nested PCR. In this experiment, the sperm and eggs were derived from AA2, and 10^8 nmol of the HNI PCR product was added to each reaction as an internal control. (Nishimura et al. 2006)
Fig. 8
The gametogenesis, mating, and zygote maturation in *C. reinhardtii*.
Nitrogen starvation and light signals induce vegetative cells to differentiate into mt+ and mt- gametes, controlled by the two mating type loci (+ and -).
Upon mixing, agglutinins mediate flagellar adhesion. A rapid increase in cAMP triggers cell wall loss and activation of mating structures. Activated mating structures fuse to form zygotes, allowing the formation of the GSP1/GSM1 heterodimer, which subsequently translocates into nuclei and switches on the zygote program.