Proper levels of the Arabidopsis cohesion establishment factor CTF7 are essential for embryo and megagametophyte, but not endosperm development

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ABSTRACT

*CTF7* is an essential gene in yeast that is required for the formation of sister chromatid cohesion. While recent studies have provided insights into how sister chromatid cohesion is established, less is known about how specifically *CTF7* facilitates the formation of cohesion, and essentially nothing is known about how sister chromatid cohesion is established in plants. In this report we describe the isolation and characterization of *CTF7* from *Arabidopsis thaliana*. Arabidopsis *CTF7* is similar to *Saccharomyces cerevisiae* *CTF7* in that it lacks an N-terminal extension, exhibits acetyltransferase activity, and can complement a yeast *ctf7* temperature-sensitive mutation. *CTF7* transcripts are found throughout the plant with the highest levels present in buds. Seeds containing T-DNA insertions in *CTF7* exhibit mitotic defects in the zygote. Interestingly, the endosperm developed normally in *ctf7* seeds, suggesting that *CTF7* is not essential for mitosis in endosperm nuclei. Minor defects were observed in female gametophytes of *ctf7*+/− plants, and plants that over-express *CTF7* exhibit female gametophyte lethality. Pollen development appeared normal in both *CTF7* knockout and over expression plants. Therefore, proper levels of *CTF7* are critical for female gametophyte and embryo development, but not for the establishment of mitotic cohesion during microgametogenesis or during endosperm development.

**Key words**: Cohesin, mitosis, gametogenesis, embryo development
INTRODUCTION

The proper formation of sister chromatid cohesion and it’s subsequent release at the metaphase to anaphase transition is essential for the proper segregation of genetic material during cell division. It is critical for the compaction of chromosomes and their bipolar attachment to the spindle, DNA double strand break repair, and the regulation of gene expression (reviewed in (Nasmyth and Haering 2009)). Sister chromatid cohesion is controlled by the cohesin complex, which consists of a heterodimer of Structural Maintenance of Chromosome (SMC) proteins, SMC1 and SMC3, and two non-SMC proteins, SCC3 and an α-kleisin protein, either SCC1 or REC8. Perhaps the most widely accepted model of how cohesin functions has been referred to as the ring model where the cohesin ring encircles the replicated sisters holding them together until SCC1 cleavage by separase at the metaphase to anaphase transition opens the ring allowing the release of the sister chromatids (reviewed in Nasmyth and Haering 2009).

The establishment of sister chromatid cohesion in yeast involves a multi-step process that begins during telophase when cohesin complexes associate with chromatin in a process requires the Scc2/Scc4 complex (Ciosk et al. 2000, Gillespie and Hirano 2004), and appears to involve the binding and then opening of the SMC hinge domain (Gruber et al. 2006, Hirano and Hirano 2006). Cohesion is established during S-phase in a Ctf7/Eco1 dependent process (Skibbens et al. 1999, Toth et al. 1999). Ctf7 interacts with a number of replication factors and appears to facilitate cohesion formation in the context of the passing replication fork (Kenna and Skibbens 2003, Lengronne et al. 2006, Moldovan et al. 2006). An anti-establishment complex consisting of WAPL/Rad61 and Pds5 is required to maintain cohesion during G2/M by stabilizing the interaction between cohesin and the chromosomes (Gandhi et al. 2006, Losada et al. 2005, Panizza et al. 2000). While the specific details of how Ctf7, WAPL/Rad61 and Pds5 function together to first establish and then maintain cohesion still needs to be clarified, recent results indicate that Ctf7 acetylates conserved lysine residues in SMC3, which inhibits the anti-establishment function of the Wpl1-Pds5 complex and promotes cohesion establishment (Ben-Shahar et al. 2008, Rowland et al. 2009, Sutani et al. 2009, Unal et al. 2008, Zhang et al. 2008). Ctf7 is also involved in the postreplicative induction of cohesion induced by DNA double strand breaks (DSB’s) (Unal et al. 2007). In this instance acetylation of the α-kleisin...
subunit, Mcd1 by Ctf7 is required for cohesion establishment (Heidinger-Pauli et al. 2009).

The structure of CTF7 proteins along with the effects of ctf7 mutations vary considerably between organisms (Bellows et al. 2003, Hou and Zou 2005, Skibbens, et al. 1999, Tanaka et al. 2000, Toth, et al. 1999, Williams et al. 2003). Saccharomyces cerevisiae ctf7 mutations cause a delay in mitosis and cells with a 2C DNA content, separated DNA masses, and partially elongated spindles (Skibbens et al. 1999, Toth et al. 1999). The fission yeast eso1-H17 mutant displays a spindle assembly checkpoint-dependent mitotic delay and abnormal chromosome segregation (Tanaka et al. 2000). Mutation of Drosophila DECO affects the distribution of cohesin proteins and cells progress prematurely into anaphase where the spindle checkpoint is activated (Williams et al. 2003). The siRNA directed depletion of ESCO1 or ESCO2 in HeLa cells caused defects in sister-chromatid cohesion, mis-alignment of chromosomes on the spindle, and abnormal chromosome segregation; it did not however appear to affect the distribution of cohesin proteins (Hou and Zou 2005). Therefore, it is possible that CTF7 proteins may have multiple roles in organisms and/or have unique properties in different organisms.

The Arabidopsis genome contains orthologs for many of the cohesion-associated genes that have been identified in yeast, and studies have shown that the cohesin machinery is generally conserved in Arabidopsis (Bai et al. 1999, Bhatt et al. 1999, Chelysheva et al. 2005, da Costa-Nunes et al. 2006, Dong et al. 2001, Jiang et al. 2007, Lam et al. 2005, Liu and Makaroff 2006). However, essentially nothing is known about how sister chromatid cohesion is established in plants. Therefore, we were interested in determining if Arabidopsis contains a functional ortholog of CTF7, and if so what role it plays in plant growth and development.

In this report we present the results of an analysis of Arabidopsis CTF7, the Cohesion Establishment Factor, and the characterization of plants containing mutations in CTF7. Seeds containing T-DNA insertions in CTF7 exhibit embryo lethality early in development, while the endosperm develops normally. Minor defects were also observed in female gametophytes and pollen germination of CTF7+/- plants; however over expression of CTF7 blocked female gametophyte development at FG1. Therefore, variations in CTF7 levels have very different effects in different cell types.
RESULTS

Arabidopsis contains a single *CTF7* gene that lacks an N-terminal extension

BLAST searches of the Arabidopsis protein database identified one gene (At4g31400) with high sequence similarity with both the human (E value = 3e-21) and *S. cerevisiae* (E value = 3e-05) Eco1 proteins. Other genes (At3g20350, At5g26190) were identified with significantly lower similarity levels (E values = 0.62, 1.7, respectively) that were missing one or more highly conserved residues found in all Ctf7/Eco1 proteins studied to date (see below). Therefore, the Arabidopsis genome appears to contain one putative *CTF7* gene (AT4g31400), hereafter called *CTF7*. A *CTF7* cDNA was isolated and compared with the published genomic sequence of At4g31400. *CTF7* consists of 6 exons and 5 introns (Fig. 1A). While the predicted and actual gene structures are similar, differences in the position of splice sites were found for exons 2, 3 and 4.

Arabidopsis *CTF7* is capable of encoding a 345-amino acid protein. Similar to *S. cerevisiae* Eco1 (281 amino acids), Arabidopsis CTF7 lacks a large N-terminal extension, which is found in most other Eco1 proteins (Fig. 1B). A PIP box (QxxL/I, QFHL) and C2H2 Zn finger motif are present at residues 82-86 and 92-130 respectively, while an acetyltransferase domain is present between amino acids 184-335 (Fig. 1B,C). Arabidopsis CTF7 exhibits between 24-58% similarity with CTF7 proteins from yeast, fruit fly, and vertebrates, which is the same level of similarity observed between other CTF7 homologues (Fig. 1C). Overall Arabidopsis CTF7 shows the highest sequence similarity to human Eco1 (22% identity, 58% similarity).

*CTF7* transcript levels were analyzed in different tissues using RT-PCR. Low, but detectible levels of *CTF7* transcripts were found in all tissues examined, including buds, roots, leaves, stems and young seedlings, with buds exhibiting the highest signal and mature leaves the lowest (Fig. 1E). Expression profiling data for CTF7 is consistent with these results and show that CTF7 transcript levels are highest in bolting plants, buds and young flowers. CTF7 transcripts are also elevated in E2FaDPa transgenic plants, which over-express S-phase genes (Vlieghe *et al.* 2003). Therefore, CTF7 expression patterns are consistent with a predicted role in nuclear
division.

Yeast, fly, and human CTF7 proteins exhibit acetyltransferase activity (Bellows et al. 2003, Ivanov et al. 2002, Williams et al. 2003, Zou et al. 1999). In order to determine if this is also the case for Arabidopsis CTF7, the protein was over-expressed in E. coli and characterized. Soluble CTF7 was produced when expressed as a maltose-binding fusion protein in the presence of pGroESL. Acetyltransferase activity was measured by autoacetylation in crude E. coli extracts and with purified CTF7 protein. Western blotting with an anti-acetyl-lysine antibody and MALDI-TOF-MS demonstrated that CTF7 is acetylated during the in vitro autoacetylation assays (Fig. 2A). Therefore, Arabidopsis CTF7 exhibits acetyltransferase activity in vitro and is able to undergo autoacetylation.

The question of whether the Arabidopsis protein can substitute for yeast Eco1 was investigated by transforming the yeast ctf7 temperature sensitive mutant, ctf7-203 (YBS514) with the Arabidopsis CTF7 cDNA. A pAS2-CTF7 construct was able to restore the ability of YBS514 cells to grow at 33°C, while YBS514 cells containing the pAS2 vector alone were not (Fig. 2B). Therefore, Arabidopsis CTF7 is able to substitute for S. cerevisiae Eco1. Taken together, these results confirm that CTF7 is the Arabidopsis Cohesin Establishment Factor.

**Genetic analysis of Arabidopsis ctf7 plants**

The role of CTF7 in plant growth and development was investigated by characterizing the SALK_059500 (ctf7-1) and SAIL_1214G06 (ctf7-2) T-DNA insertion lines. PCR amplification followed by DNA sequence analysis confirmed that the ctf7-1 and ctf7-2 T-DNA insertions are located in intron 4 and exon 3, respectively. Both mutations are predicted to disrupt the gene upstream of the acetyltransferase domain.

No homozygous ctf7-1 or ctf7-2 plants were identified in initial screens. Therefore, seed from self-pollinated, heterozygous ctf7<sup>+/−</sup> plants was collected, resown, and the progeny screened by PCR. Fifty nine percent (223/379) of the progeny from heterozygous ctf7-1 plants were heterozygous for the ctf7 T-DNA insertion, whereas 41% of the plants were wild type. No homozygous mutant plants were identified. Likewise no homozygous plants containing the ctf7-2 insertion were identified, with 54% (143/263) of the progeny being heterozygous for the
T-DNA insertion. These results indicated that CTF7 is essential, and suggested that its disruption may cause embryo lethality. Furthermore, the observation that self pollinated ctf7-1+/− and ctf7-2+/− plants consistently produced fewer heterozygous progeny than expected (combined genetic data for the two lines: 336:276 (1.2:1) heterozygous:wild type, \(x^2=0.03; P>0.05\), also suggested a gametophytic effect of the mutations.

To investigate this possibility, ctf7-1+/− plants were backcrossed with wild type plants as both the male and female parent. When ctf7-1+/− plants were used as the female parent, 144 wild type and 146 ctf7+/− plants were obtained, while 120 wild type and 91 ctf7+/− plants were obtained when ctf7-1+/− plants were used as the male parent. Similar results were obtained in backcross experiments with ctf7-2+/−. These results indicated that the mutation is transmitted through both the male and female gametophyte, but that the transmission efficiency is reduced through male gametophytes.

Similar phenotypes were observed for ctf7-1 and ctf7-2 plants in all of our analyses. Therefore, only the results obtained from our analysis of ctf7-1 plants are presented and the mutant will be referred to generally as ctf7 from here forth. A light microscope analysis of semi-thin sections through stage 11 and 12 anthers (Sanders et al. 1999) failed to identify developmental abnormalities in pollen development in ctf7+/− plants (data not shown). Pollen from ctf7+/− plants appeared uniform in size and normal when examined by SEM (Fig. 3A). The large majority, if not all, pollen from ctf7+/− plants contained two sperm nuclei and a vegetative nucleus, confirming that pollen mitosis is not disrupted by the mutation (Fig. 3B). No significant difference in pollen viability was observed between wild type and ctf7+/− plants (Fig. 3C). However, pollen from ctf7+/− plants did exhibit a slightly reduced in vitro germination frequency (73% of wild-type levels, Fig. 3C).

Further analysis of ctf7+/− plants identified no obvious morphological alterations with the exception that they exhibited a reduced seed set (32±4 versus 43±6 per silique in wild type plants, n=50, Fig. 3C). Approximately 25% of the seeds in the siliques of ctf7+/− plants at cotyledon stage were normal in size, but white; later these seeds became shrunken and orange (Fig. 3D). The presence of 25% aborted seeds in ctf7+/− siliques is consistent with a mutation
that affects embryo development.

**CTF7 mutations cause minor alterations in female gametophyte development**

Potential effects of the *ctf7* mutation on megagametogenesis were characterized by comparing ovules in wild type and *ctf7*+/- plants using Laser Scanning Confocal Microscopy (LSCM). A total of 36 pistils at various stages of development from three different *ctf7*+/- plants were analyzed. Female gametophyte development was defined according to (Christensen et al. 1998). No detectable abnormalities were identified in embryo sacs of *ctf7*+/- plants between stages FG0 and FG6 (Fig. 4A-D). However, embryo sac development was less uniform in the siliques of *ctf7*+/- plants than those of wild type plants. In contrast to the two or three developmental stages observed in wild type siliques (Shi et al. 2005), at least four different stages were observed in *ctf7*+/- siliques (Table 1). For example, both fertilized and FG5 embryo sacs were found in the same slique. This was never observed in wild type siliques suggesting that *ctf7* female gametophytes develop slower than wild type female gametophytes.

The first developmental abnormality was observed at stage FG7 when the three antipodal cells did not degenerate normally in *ctf7* embryo sacs. Typically, by the time the egg and synergid cells are mature, as evidenced by the opposite positions of nuclei and vacuoles, the three antipodal cells have degenerated (Fig. 4A). However, three antipodal cells were often found along with mature FG7 egg and synergid cells in embryo sacs of *ctf7*+/- siliques. Antipodal cells were also observed in recently fertilized embryo sacs, and in embryo sacs containing endosperm with 2 or 4 nuclei (Fig. 4E-G, I). Antipodal cells were never observed in fertilized embryo sacs of wild type plants (Fig. 4B-D). Approximately fifty percent (45.8±1.4%) of the embryo sacs in *ctf7*+/- siliques exhibited this delayed degeneration of antipodal cells, suggesting that it is linked with the *ctf7* mutation. This observation along with variations in the timing of female gametophyte development and the reduced pollen germination indicates that the *ctf7* mutation has minor effects on gametophyte development. While not severe, together they likely contribute to the reduced transmission of the mutation that we observe.
Inactivation of CTF7 blocks embryo development, but has no significant effect on endosperm development.

Analysis of seed development in ctf7+/− plants identified alterations in zygote formation in approximately 25% of the seed. In wild type Arabidopsis plants a single-celled zygote and a primary endosperm nucleus are normally observed along with degenerated and persistent synergid cells after fertilization of the egg and central cell. The endosperm nucleus then undergoes a series of syncytial divisions that produce an endosperm with three domains: the embryo-surrounding region or micropylar endosperm (MCE), the peripheral endosperm (PEN) in the central chamber, and the chalazal endosperm (CZE) (Olsen, 2004). After the endosperm nucleus has undergone three to four rounds of nuclear division the zygote becomes elongated and divides asymmetrically generating a smaller apical cell and a larger basal cell (Ingouff, et al., 2005). Subsequent divisions produce an embryo composed of the embryo proper and the suspensor, which is composed of an enlarged basal cell and a file of six to eight additional cells. The embryo proper undergoes successive rounds of division forming the two cell, quadrant, octant (Fig. 5A), dermatogen, globular (Fig. 5G), early heart (Fig. 5M), heart, torpedo (Fig. 5S), and mature embryo, respectively.

Fertilization appeared to occur normally (Fig. 4H,I), but alterations were observed in ctf7 zygotes soon after the first division when a number of embryos arrested and subsequently degenerated at the one or two cell zygote stage (Figs. 4J-L, 5D). Some zygotes reached the four-celled pro-embryo stage at which point both the suspensor and embryo proper exhibited altered division planes (Fig. 5J,P). A small number of embryos, which exhibited dramatic alterations in division, reached the dermatogen and early globular stages before arresting (Fig. 5V). All together 23.7±0.6% of the seeds in ctf7+/− siliques (n=7) exhibited defects in early zygote and embryo development, strongly suggesting that the mutation causes embryo lethality by either blocking or altering nuclear division.

In contrast, the endosperm in ctf7 seeds appeared relatively normal throughout development. In wild type seeds the endosperm contains approximately 26-50 nuclear cytoplasmic domains (NCD’s), by the time the embryo first begins to divide, while octant staged embryos are typically surrounded by approximately 100 uncellularized NCDs (Fig. 5B,C).
appearance of NCD’s were normal in ctf7 seeds containing arrested zygotes and pro-embryos (Fig. 5D-F). Cellularization of the MCE begins when the embryo is at the early globular embryo stage in wild type seeds (Fig. 5G). At this time the PEN showed a gradient of cellularization stages (Fig. 5I) and endosperm nodules as well as the chalazal cyst (cz) were observed in the CZE (Fig. 5H). The endosperm of ctf7 seeds showed a similar pattern of development with no obvious abnormalities, even though the embryo had already arrested (Fig. 5J-L). By late globular/early heart stage the MCE had completed cellularization and the PEN had started the cellularization process in wild type seeds (Fig. 5M-O). Again cellularization of the MCE and PEN, and formation of the chalazal cyst occurred normally in ctf7 seeds (Fig. 5Q,R). Finally, by the time the wild type embryo had developed to the torpedo stage, the PEN had completed cellularization and the endosperm began to show signs of degeneration (Fig. 5S-U). Complete cellularization, followed by degeneration of the endosperm was also observed in ctf7 seeds (Fig. 5W,X).

Subtle defects in the NCD’s of some ctf7 seeds were however detected beginning when the corresponding wild type seeds were at approximately the 32-cell embryo stage. In particular, a small number of NCD’s with enlarged nucleoli, NCD’s that contained multiple nucleoli and NCD’s that contained multiple nuclei were observed (<5%, data not shown). However, these alterations were generally rare and did not appear to have any significant effect on endosperm development. Therefore, while inactivation of CTF7 blocks embryo development beginning at the zygote stage it has little to no effect on endosperm development.

**Over expression of CTF7 blocks early ovule development**

We next tested the effect of over expression of Arabidopsis CTF7 on plant growth. Transgenic plants that over express CTF7 from the 35S promoter appeared normal during vegetative growth; however 10 out 13 independent lines exhibited varying degrees of reduced fertility, ranging from 5% to 52% aborted seed. Southern blot analysis of two lines (1 & 18), that exhibited the highest level of aborted seed (484:311; 568:627 normal:aborted) showed that they contain a single insert. A genetic analysis of one of these lines showed that the T2 generation segregated approximately 1:1 for BASTA resistance (47 BASTA⁺:53 BASTA⁻),
suggesting that the 35S:CTF7 construct causes gametophytic lethality.

Further analysis of the ten 35S:CTF7 reduced fertility lines showed that they all produced normal levels of viable pollen, but contained alterations in female gametophyte development. Detailed analyses of gametophyte development in lines 1 and 18 demonstrated that pollen development is unaffected (data not shown), while 46% (n=795) of the ovules were found to abort early in development. Specifically, LSCM analysis of female gametophyte development in lines 1 and 18 showed that approximately half of the ovules arrested at approximately FG1 (Fig. 6 I,J). In the ovules exhibiting wild type development, the functional megaspore (FG1, Fig. 6A) underwent mitosis to give rise to a two-nucleate embryo sac (FG2, Fig. 6B). A central vacuole formed between the two nuclei (FG3, Fig. 6C) and the two nuclei underwent a second division to give rise to a four-nucleate embryo sac at early FG4 (Fig. 6D). At this time, the division plane between the two chalazal nuclei was parallel to the chalazal-micropylar axis. Then, two of the four nuclei migrated so that the division plane between them was orthogonal to the axis. Another round of division gave rise to an eight-nucleate embryo sac. The two polar nuclei, one from each pole, then migrated toward the micropylar half of the developing female gametophyte (FG5, Fig. 6E) and eventually fused to form the central cell (FG6, Fig. 6F). The three antipodal cells degenerated just before fertilization giving a mature female gametophyte consisting of two synergid cells, one egg cell and one central cell (FG7, Fig. 6G).

In contrast, a significant number of the female gametophytes, which we predict over express CTF7, arrested at FG1 (Fig. 6I,J). The arrested female gametophytes did not exhibit any obvious abnormalities, but typically failed to divide to form a two-nucleate embryo sac. Occasionally we observed an arrested female gametophyte at FG2 (Fig. 6K); however this was quite rare. Obvious abnormalities were also not observed in the arrested FG2 female gametophytes, but ultimately all of the arrested female gametophytes degraded (Fig. 6L). The size of embryo sac continued to enlarge in seeds with arrested female gametophytes, indicating that integument growth was normal (Fig. 6I-L).

The possibility that over expression of CTF7 affects female gametophyte development was further investigated by analyzing transgenic plants that express CTF7-YFP from the 35S promoter. Of the ten transgenic lines analyzed, six exhibited varying levels of reduced fertility (9-41% aborted seed). Analysis of the lines showed that YFP is expressed from the 35S
promoter in the female gametophyte throughout ovule development and that similar to 35S:CTF7 lines, female gametophytes arrested at FG1 and FG2 (Fig. 7 A-H). Therefore, expression of CTF7 from the 35S promoter can block female gametophyte development beginning at FG1.

**CTF7 is abundant in the embryo, but present at very low levels in the endosperm.**

Our observation that endosperm development is not significantly affected by inactivation CTF7 suggested that CTF7 is not required for nuclear division in the endosperm. This raised the question of whether CTF7 is actually present in the endosperm. To investigate this question we analyzed CTF7 distribution and localization patterns in seeds using both immunolocalization and transgenic plants in which YFP was fused to the C-terminus of the CTF7. The CTF7-YFP fusion protein could be readily detected in the embryo, while signals from the endosperm were barely detectable (Fig. 7I-L). While weak fluorescence was detected in the MCE, PEN, CZE, the signals in the endosperm were very weak, not always uniform, and often not above background levels. Immunolocalization experiments using a polyclonal antibody to full-length Arabidopsis CTF7 again revealed strong CTF7 signal in the embryo at various stages of development, while cross-reactivity in the endosperm was typically similar to that observed in negative control experiments with pre-immune or secondary antibody alone (Fig. 7M-P). Strong auto fluorescence was observed in the integuments in all samples. Therefore, while CTF7 is present at high levels in the embryo, it is present at very low levels, if at all in the endosperm. Finally, an analysis of the subcellular distribution of CTF7 showed that it is present in both the nucleus and the cytoplasm with the nuclear always significantly stronger than the signal detected in the cytoplasm. A function for CTF7 outside the nucleus has not been described in any organism. Therefore, the significance, if any of cytoplasmic CTF7 is unknown.

**DISCUSSION**

Ctf7 proteins possess several highly conserved domains, including a PIP box, a C$_2$H$_2$ zinc finger and an acetyltransferase domain, which are all present in Arabidopsis CTF7. The PIP box is required for PCNA binding (Moldovan *et al.* 2006). The C$_2$H$_2$ zinc finger motif appears

CTF7 proteins in most organisms contain additional diverse N-terminal domains. In fission yeast the N-terminal two-thirds of ESO1 is similar to DNA polymerase η of budding yeast and deletion of this domain increases sensitivity to UV irradiation (Tanaka et al. 2000). Human ESCO1 contains an N-terminal extension with similarity to linker histone proteins, which facilitates its binding to chromosomes (Hou and Zou 2005). Arabidopsis CTF7 is significantly shorter than the S. pombe and human proteins, and similar to the S. cerevisiae protein in that it lacks an extended N-terminal domain. Predicted rice and maize CTF7 proteins also lack an amino terminal extension, suggesting that this is a common feature of plant CTF7 proteins. Arabidopsis CTF7 is able to complement a S. cerevisiae ctf7 temperature sensitive mutant, while the human protein is not (Bellows et al. 2003). Taken together, these results suggest that CTF7 may have evolved additional different roles or may function somewhat differently in different organisms.

Proper levels of CTF7 are essential for mitosis in the female gametophyte

Self-pollinated ctf7+/− plants consistently produce a 1.2:1 ratio of heterozygous:wild type progeny, indicating that the mutation has an effect on gametophyte development. Transmission through pollen was reduced slightly in ctf7+/− plants with the pollen exhibiting reduced germination frequencies. However no dramatic alterations in pollen development were observed and the generative nucleus and two sperm nuclei were present and appeared normal. Likewise, pollen development appeared normal in transgenic plants over expressing CTF7. Therefore, changes in CTF7 levels either by mutation or over expression do not have a dramatic effect on pollen development or function.
Inactivation of *CTF7* also did not have a dramatic effect on female gametophyte development, although again subtle alterations were observed. Specifically, inactivation of CTF7 slowed the progression of mitosis and overall development of female gametophytes (Table 1); in addition the three antipodal cells did not degenerate normally in *ctf7* embryo sacs (Fig. 4E,F,I). However, these alterations do not appear to directly affect the function of the embryo sac or fertilization as the mutation is transmitted normally through female gametophytes (Table 1). These results are consistent with a number of reports showing that inactivation of essential genes, including several involved in the formation or release of cohesion does not block gametophyte development but rather cause alterations in embryo development (Lam *et al.* 2005, Liu and Meinke 1998, Liu and Makaroff 2006). It has been suggested that gametophytes may have a pool of mRNA or protein that can be utilized for the successful completion of the limited number of nuclear the divisions required for gametophyte development (Liu and Meinke 1998). Evidence in support of this hypothesis comes from the observation that transgenic plants expressing an ESP RNAi construct from the meiotic DMC1 promoter exhibit defects much earlier than T-DNA insertion lines (Liu and Makaroff 2006).

In contrast, expression of CTF7 and CTF7:YFP from the 35S promoter significantly impacted female gametophyte development, typically resulting in arrest at FG1 or FG2 (Fig. 6I-k). The arrested female gametophytes did not exhibit any obvious abnormalities, but generally failed to divide to form a two-nucleate embryo sac. The observed effect was variable, with the number of aborted ovules/slique ranging from 5 to 52%. We believe that this variability is likely due to the relatively weak and variable expression of the 35S promoter in female gametophytes of the different transgenic lines. Our results showing that the 35S promoter is active in female gametophytes of 35S-CTF7-YFP plants (Fig. 7A-H) supports this hypothesis and is consistent with another report showing that the 35S promoter is active during female gametophyte development (Liu *et al.* 2008). An alternative possibility that the observed female gametophyte arrest results from chromosomal rearrangements associated with T-DNA insertion (Curtis *et al.* 2009, Ray *et al.* 1997) is less likely given the absence of pollen defects, the results of our expression and the multiple independent occurrences of the phenotype in the transgenic lines.

At this time it is not clear how over-expression of CTF7 affects female gametophyte
development. However, it has recently been shown that the acetylation state of SMC3 affects the rate of clamp-loader dependent fork progression in human cells (Terret et al. 2009) and that over expression of Ctf7 in yeast bypasses the requirement for DSBs to generate cohesion during G2/M (Heidinger-Pauli et al. 2009). Therefore, it is possible that over-expression of CFT7 may shift the balance of acetylated cohesin complex resulting in a slowdown or blockage of the replication fork. It is also possible that a noncohesin protein is aberrantly acetylated, causing gametophyte arrest. Our observation that CFT7 over expression only affects the female gametophyte suggests that female gametophytes are more sensitive to these perturbations than male gametophytes or that over expression of CFT7 in the transgenic plants may be higher in the female gametophytes. Further experiments are required to better understand the role of CFT7 in female gametophyte development.

**CTF7 is essential for mitosis in the embryo but not in pollen or the endosperm**

*CTF7* is essential for nuclear division in the embryo, with most *ctf7* embryos arresting soon after fertilization and prior to the first division (Fig. 5 J,K). In contrast, the syncytial nuclear divisions of the fertilized central cell proceeded normally in *ctf7* seeds. Furthermore, cellularization of the endosperm and further cell division appeared similar to the situation in wild type endosperm (Brown et al. 1999, Ingouff et al. 2005, Sorensen et al. 2002). Therefore, while the zygote is not able to undergo nuclear division in the absence of CTF7, multiple rounds of endosperm nuclear division occur and are relatively normal in the absence of CTF7.

Our localization studies show that while CTF7 is present at high levels in the embryo, it is absent/barely detectable in the endosperm (Fig. 7). These results, along with those from our phenotypic studies suggest that CTF7 is not be required for the establishment of sister chromatid cohesion in the endosperm. In *S. pombe* Eso1 becomes dispensable if *Pds5* is deleted (Tanaka et al. 2001), while in *S. cerevisae* the lethality of an *Eco1* deletion is suppressed by inactivation of *Rad61/Wpl1* (Ben-Shahar et al. 2008 #4445). Furthermore, cohesion defects in human cells depleted for Esco1 can be rescued by co-depletion of Wapl1 (Gandhi et al. 2006). Therefore it is possible that the endosperm lacks CTF7, WAPL1 and PDS5 and instead uses a different mechanism to establish sister chromatid cohesion.

The possibility also exists that a CTF-like protein may substitute for CTF7 in the endosperm.
Several predicted proteins (At3g20350, At5g26190, At2g44140) exhibit low level similarity to yeast, human, and Arabidopsis CTF7. However none of the proteins contain all of the elements conserved in CTF7 proteins (e.g. PIP box, C₂H₂ Zn finger, and acetyltransferase domain), and phylogenetic analyses of the Arabidopsis CTF7, At3g20350, At5g26190, At2g44140 proteins support the conclusion that Arabidopsis contains a single CTF7 ortholog (Fig. 1d). Finally, it is possible that the early embryo abortion and continued endosperm nuclear division we observe results from different cell cycle checkpoints operating in embryos and the endosperm. It is possible that endosperm nuclei either lack cell cycle checkpoints or have altered cell cycle checkpoints that tolerate the mis-segregation of chromosomes. Consistent with this hypothesis are several embryo lethal mutants that arrest at the one or two-cell zygote stage while the endosperm continues to divide. However, very few of the mutants characterized to date appear to exhibit normal endosperm development. For example, of the 70 MEE mutants that arrest very early in embryo development, only six were found to exhibit normal endosperm development (Pagnussat et al. 2005). Therefore our observation that inactivation of CTF7 blocks embryo development soon after fertilization, but has no apparent effect on endosperm development is quite unusual.

**EXPERIMENTAL PROCEDURES**

**Plant material and growth conditions**

Plants were grown in Metro-Mix200 soil (Scotts-Sierra Horticultural Products Co.; http://www.scotts.com) or on germination plates (Murashige and Skoog, Caisson laboratories, Inc.; www.caissonlabs.peachhost.com) in a growth chamber at 22°C with a 16-hour-light and 8-hour-dark cycle. *Arabidopsis thaliana* L. Heynh, ecotype Wassilewskija (WS) was used for transcript analysis and plant transformation studies. Crossing studies were conducted with the Columbia ecotype. The SALK_059500 and SAIL_1214G06 T-DNA insertion lines were obtained from the Arabidopsis Biological Resource Center (ABRC). Seedlings and roots were collected from seven day old plants grown on plates. Leaves were harvested from rosette-stage plants grown on soil. Approximately 20 days after germination, buds and stems were collected.
All samples were harvested, frozen in liquid N₂ and stored at –80 °C until needed.

**Molecular analysis of CTF7**

The *Arabidopsis CTF7* (At4g31400) genomic DNA sequence was identified by BLAST searching the Arabidopsis AGI Proteins Database (http://www.arabidopsis.org/blast/) with the *S. cerevisiae* Eco1, *S. prombe* Eso1 and human ECO1 proteins.

A *CTF7* cDNA (accession number EU077499) was obtained from total RNA isolated using Trizol (Invitrogen Inc) by performing reverse transcription with primer 695 followed by PCR using primers 693 and 695 (Fig. 1). *CTF7* transcript levels in buds, leaves, shoots, stems, roots and seedlings of wild type plants, were analyzed using RT-PCR. Reverse transcription was conducted on total RNA (4 µg) using a Thermoscript™ RT-PCR system (Invitrogen) with an oligo dT primer followed by 25 cycles of PCR with primer pair 741/696 (Fig. 1). Amplification products were analyzed by Southern blotting using a ³²P-dATP-labeled 741/696 PCR fragment. After hybridization and washing radioactivity was detected using a Molecular Dynamics Phosphorimager (Molecular Dynamics). RT-PCR was conducted using primers to *ACTIN 8 (ACT8)* to standardize the amount of cDNA (An *et al.* 1996).

Genomic DNA of T-DNA insertion plants was genotyped by PCR with primer pairs specific for the T-DNA and wild type loci. After confirmation of the presence of the T-DNA insertion, the SALK_059500 line was renamed *ctf7-1*. Southern blot analysis on *ctf7°/°* plants confirmed that there was only one T-DNA insert in the line. A 4275 bp genomic DNA fragment containing the *CTF7* coding region and 2422 bp of 5' flanking sequence in pFGC5941 was used to complement *ctf7-1* plants using the floral dip transformation method (Clough and Bent 1998). Transgenic plants were identified by BASTA resistance and PCR genotyped using primer pairs for the transgene, the genomic *CTF7* locus, and for the T-DNA insert.

Two CTF7-YFP fusion protein constructs were generated. The first was generated by cloning the 2082 bp coding sequence fragment into 35s::YFP pFGC5941. In the second construct a 4275 bp genomic DNA fragment containing the *CTF7* coding region and 2422 bp of 5' flanking sequence was cloned into a pFGC5941 vector containing YFP. After confirmation by DNA sequencing the clones were introduced into plants by Agrobacterium transformation and
screened by BASTA. Positive plants were confirmed by PCR and further analyzed by confocal microscopy.

**Analysis of CTF7 acetyltransferase activity**

The CTF7 cDNA was cloned into pET-24b as an NdeI/EcoRI fragment and into pIADL14 as an NdeI/HindIII fragment to generate a CTF7-MBP fusion (CTF7-MBP). After DNA sequencing, pET-24b-CTF7 was transformed into BL21(DE3) pLysS E. coli cells and the over expression and solubility of the protein was determined. The solubility of CTF7-MBP was improved by introducing pGroES into the cell line (Goloubinoff et al. 1989). CTF7-MBP was purified on a maltose-binding resin from whole cell extracts of cells that had been grown overnight at 30 °C. Fractions containing recombinant CTF7-MBP were analyzed by SDS-PAGE and confirmed by MALDI-ToF-MS. CTF7-MBP was assayed for acetyltransferase activity as previously described (Bellows, Kenna, Cassimeris and Skibbens 2003).

**Complementation of yeast ctf7-203 cells**

Yeast expression constructs were prepared by transferring the CTF7 protein-coding region into pAS2 (Clontech Laboratories, Inc. http://www.clontech.com). CTF7-pAS2 and pAS2 were transferred into the temperature-sensitive ctf7-203 line, YBS514, using Li acetate transformation (Skibbens et al. 1999). The ability of the constructs to complement the ctf7-203 mutation was tested by growing YBS255 (wild type), and YBS514 cells containing CTF7-pAS2 or pAS2 in the appropriate dropout media. After growth at 25°C for 48 hrs, 10 μl of a serial dilution of YBS255, ctf7-203/YBS514, YBS514-PAS2 and YBS514-CTF7-PAS2 were plated on the selection plates and grown at 25°C, 28°C, 33°C and 38°C for 48 hr.

**Morphological Analysis of ctf7+/− and 35S-CTF7 plants**

Ovule development was analyzed in ctf7+/− plants by CLSM essentially as described (Christensen et al. 1997). Inflorescences were collected and fixed in 4% glutaraldehyde under vacuum
overnight, dehydrated in a graded ethanol series (20% steps for 1 hr each) and cleared in a 2:1 mixture of benzyl benzoate:benzyl alcohol. The pistils were mounted under sealed coverslips. Images were collected, and projected with Olympus Flouvview 2.0 software (http://www.olympus-global.com/), analyzed with Image Pro Plus (Media Cybernetics; http://www.mediacy.com) and presented with PHOTOSHOP version 7.0 (Adobe; http://www.adobe.com).

Endosperm development was analyzed in siliques from 5 individual ctf7+/− plants using CLSM essentially as described (Braselton et al. 1996). Inflorescences were fixed in 3:1 ethanol:glacial acetic acid at 4°C for 48 hours, hydrolyzed with 5 mol l-1 HCl for one hour followed by staining using Schiff’s reagent (Sigma, www.sigmaaldrich.com) for 24 hours. After dehydration in a serial ethanol series, individual seeds were dissected in methyl salicylate, and viewed using an Olympus IX-81 fluorescence deconvolution microscope system. Whole mount clearing was used to determine the embryo and endosperm phenotype. Siliques from ctf7 heterozygous plants were dissected and cleared in Herr's solution containing lactic acid:chloral hydrate:phenol:clove oil:xylene (2:2:2:2:1, w/w). Embryo and endosperm development was studied microscopically with a Nikon microscope equipped with differential interference contrast optics.

Pollen morphology, viability, and germination were compared in newly opened flowers of ctf7+/− plants and wild type siblings from the same pot. Pollen morphology was analyzed using scanning electron microscopy (SEM). Freshly opened flowers were fixed in formalin/acetic acid/alcohol (FAA) overnight, rinsed three times in fixation buffer, and dehydrated in a graded ethanol series. Once in 85% ethanol, the anthers were dissected, further dehydrated up to 100% ethanol, critical point dried with CO2, mounted on SEM stubs with carbon tabs, sputter-coated with 21 nm of gold, viewed and photographed using a Zeiss Supra 35 FEG-VP scanning electron microscope (Carl Zeiss, Inc.; www.zeiss.com).

Pollen viability was analyzed by soaking the grains in 2,3,5-triphenyl tetrazolium chloride (TTC, 1.0 % by weight in 50 % sucrose) at 37°C for 30 min followed by observation by fluorescence microscopy (Huang et al. 2004). Pollen mitosis was analyzed by 4',6-diamidino-2-phenylindole (DAPI) staining (2 mg l−1) followed by observation by
fluorescence microscopy. Pollen germination was tested according to (Fan et al. 2001).

**CTF7 Immunolocalization**

Polyclonal antibodies were raised to Arabidopsis CTF7, which was over-expressed in *E. coli* and affinity purified using standard procedures (Harlow and Lane 1988). For immunolocalization experiments Arabidopsis ovules were fixed overnight in 4% paraformaldehyde in 1XPBS supplemented with 0.1% triton X-100. The material was then squashed using a cover glass, treated for 1 hour with 2% Driselase, washed briefly with PBS and then with PBS containing 3% IGEPAL CA-630 (Sigma) plus 10% DMSO for 1 hour. The slides were washed three times with 1xPBS, blocked with 3% BSA and then treated with primary antibody over night at 37°C. The slides were washed six times with 1XPBS treated with secondary antibody, for three hrs at 37°C. After washing the slides were stained with DAPI for 30 min., washed briefly, treated with antifade and sealed with a cover glass. The slides were analyzed using confocal microscopy as described above.

**ACKNOWLEDGMENTS**

We are grateful to Richard Edelmann and Matthew L. Duley for technical support with microscopy, Robert Skibbens (Lehigh University) for providing us the yeast strains YBS255 and *ctf7-203* (YBS514), Rich Jorgenson (University of Arizona) for providing the pFGC5941 plasmid, Michael Crowder (Miami University) for providing the MBP plasmid and John Hawes for providing the pGroESL plasmid. We would like to thank Meghan Holdorf and Lara Strittmatter for helpful discussions and comments on the manuscript.

**LITERATURE CITED**


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FIGURE LEGENDS

Figure 1  Arabidopsis CTF7 locus and protein structure.
A, Gene map of CTF7. The exon positions and T-DNA insertion site are shown. Primers used in this study are shown above the map.  B, Schematic representations of ECO1/CTF7 proteins from different organisms. The ECO1/CTF7 domain (gray box), C$_2$H$_2$ zinc finger domain (black box) and PIP box (thin gray line) are shown. Accession numbers are as follows: A. thaliana CTF7 (EU077499), O. sativa CTF7 (Q7XY81), S. pombe Eso1 (O42917), S. cerevisiae Eco1 (P43605), Homo sapien Esco1 (Q5FWF5), Drosophila Eco (Q9VS50), Homo sapien Esco2 (Q56NI9).  C, Alignment of ECO1/CTF7 proteins in different organisms. The conserved PIP box, C$_2$H$_2$ zinc finger motif and acetyltransferase domain are overlined. Identical and similar amino acids are shaded black and gray, respectively. D, Phylogenetic tree of characterized ECO1/CTF7 proteins.  E, CTF7 transcripts in different tissues of Arabidopsis. ACT8 transcripts were used as a control.

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Figure 4  Female gametophyte and seed development in wild type and ctf7+/− siliques.
The images represent a single 1.5μm optical section unless otherwise noted.  A-D, normal ovules.  E-G, ctf7 ovules.  A, Wild type female gametophyte at stage FG7 containing two synergid cells (SN), an egg cell (E) and a central cell (CC).  A projection of three 1.5 μm optical sections is shown.  B, Female gametophyte at stage FG8 with degenerated synergid cell (DS).  A projection of three 1.5 μm optical sections is shown.  C, Fertilized developing seed that contains an elongated zygote (ZN) and two endosperm nuclei (EN).  A projection of two 1.5 μm optical sections is shown.  D, Developing seed in which the zygote has formed the suspensor (S) and terminal (T) cells.  E, First optical section of an ovule with three un-degenerated antipodal cells.  A degenerated synergid cell, a persistent synergid cell (SN), and the egg cell (E) are at the micropylar pole.  One of the three antipodal cells (AN) is at the bottom of the embryo sac.  F, Second optical section showing two antipodal cells (AN) at the bottom of the embryo sac.  G, Third optical section with the central cell (CC) in the middle of the embryo sac.  H, Fertilized seed containing degenerated (DS) and persistent (SN) synergid cells and a zygote (Z).  The endosperm nucleus has divided into two nuclei (EN).  A projection of three (3 μm) optical sections is shown.  I, Optical section (3 μm) through the chalazal pole of the same seed in H.  The nuclear and cell membrane of three antipodal cell nuclei (AN) are still intact.  J, ctf7 seed containing a degenerated zygote (DZ) and 8 endosperm nuclei.  Only one of the endosperm nuclei (EN) is in the same section as the zygote.  K, ctf7 seed containing a degenerated zygote (DZ) and 16 endosperm nuclei.  Five of the endosperm nuclei (EN) are observed.  A projection
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**Figure 5** Embryo and endosperm development in wild type and ctf7/+/+ sliques.

Wild type (A–C, G–I, M–O and S–U) and ctf7/+ (D–F, J–L, P–R, and V–X) embryo and endosperm whole mount images are shown. A, Wild-type octant stage embryo. Micropylar endosperm (MCE) is still at the NCD stage. B, Chalazal endosperm (CZE) of embryo sac in A. C, Peripheral endosperm (PEN) of embryo sac in A. D, ctf7 embryo showing aberrant cell division. Arrow indicates division plane. E, CZE of embryo sac in D. F, PEN of embryo sac in D. G, Wild type globular embryo. MCE shows signs of cellularization (arrow head). H, CZE of embryo sac in G. Endosperm nodules (no) and chalazal cyst (cz) are visible. I, PEN of embryo sac in G. J, ctf7 embryo showing aberrant cell division. Arrow indicates the division plane. K, CZE of embryo sac in J. L, PEN of embryo sac in J. M, Wild type early-heart stage embryo. N, CZE of embryo sac in M. O, PEN of embryo sac in M. Arrowheads indicate NCD’s starting to cellularize. P, ctf7 embryo showing aberrant cell division. Arrow indicates abnormal cell shape and cell organization. Q, CZE of embryo sac in P. R, PEN area of embryo sac in P. Arrowheads indicate NCDs starting to cellularize. S, Wild type torpedo stage embryo. T, CZE of embryo sac in S. NCD’s are still present. U, PEN of embryo sac in S. Arrowheads show cells that have completed cellularization. V, ctf7 embryo with an abnormal shape and disorganized cell order. W, CZE of embryo sac in V. X, PEN of embryo sac in V. Arrowheads show cells that have completed cellularization. Scale bars = 10 μm.

**Figure 6** Female gametophyte development revealed by CLSM in wild type and CTF7-OE plants.

A–H, Wild-type female gametophyte development. I–L, Female gametophyte development in CTF7-OE lines. A, Female gametophyte stage 1 (FG1) ovule showing the functional megaspore (M). A trace of the degraded megaspore (DM) still visible. Ch, chalazal. B, FG2-stage ovule with a two-nucleate (N) embryo sac. The degraded megaspore is still visible in some cases. C, FG3-stage ovule showing a late two-nucleate embryo sac with an enlarged central vacuole (V) and a small chalazal vacuole. D, Ovule with a four-nucleate embryo sac at FG4 stage. E, Ovule with an embryo sac at FG5 stage. Cellularization and cell differentiation is complete with the
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35S-CTF7 ovule. No difference with wild-type FG1 ovules is apparent. The degraded megaspore
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at FG1. K, 35S-CTF7 ovule. The female gametophyte completed one round of mitosis. L,
35S-CTF7 ovule. The female gametophyte is degraded. Scale bars = 5μm.

**Figure 7.** Cellular distribution of CTF7 in wild type and 35S-CTF7::YFP plants.
A-D, 35S driven CTF7::YFP localization during female gametophyte development. A, FG1 ovule,
arrow indicates FG nucleus. B, Bright field of image A, position of FG is circled. C, FG2 ovule,
arrows indicate nuclei. D, FG7 ovule, arrows indicate nuclei. E-H, Aborted female gametophytes
in 35S-CTF7::YFP plants. E, Aborted FG1 stage ovule, arrow indicates degraded nucleus. F,
Bright field of image E. G, Aborted ovule at early FG1, arrow indicates degraded nucleus. H,
Aborted ovule, arrow indicates remnants of degraded nucleus. I, Seed with globular staged
embryo from genomic-CTF7::YFP plant. J, Seed with heart staged embryo from
genomic-CTF7::YFP transgenic plant. K, Seed with torpedo staged embryo from
genomic-CTF7::YFP transgenic plant. L, Seed from wild type, negative control plant. M-P,
CTF7 immunolocalization on whole mount cleared seed. M, torpedo stage embryo. N, single
embryo at early torpedo stage. O, cotyledon stage embryo. P, Negative control with no primary
antibody. Bar=5μm
Table 1. Female Gametophyte Development in *ctf7+/* plants

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Figure 6 Female gametophyte development revealed by CLSM in wild type and CTF7-OE plants and CTF7 YFP fusion protein OE pattern in female gametophyte.
A-H, Wild-type female gametophyte development. I-L, Female gametophyte development in CTF7-OE lines. A, Female gametophyte stage 1 (FG1) ovule showing the functional megaspore (M). A trace of the degraded megaspore (DM) still visible. Ch, chalazal. B, FG2-stage ovule with a two-nucleate (N) embryo sac. The degraded megaspore is still visible in some cases. C, FG3-stage ovule showing a late two-nucleate embryo sac with an enlarged central vacuole (V) and a small chalazal vacuole. D, Ovule with a four-nucleate embryo sac at FG4 stage. E, Ovule with an embryo sac at FG5 stage. Cellularization and cell differentiation is complete with the formation of two synergid nuclei (SN), an egg nucleus (EN), three antipodal nuclei (AN) and the two prominent polar nuclei (PN), which have not yet fused. F, Ovule with a mature seven-celled embryo sac at stage FG6. The polar nuclei have fused to form a diploid central nucleus. G, Ovule at FG7 in which the antipodal cells have begun to degenerate. H, Ovule after fertilization. One synergid cell is degraded and the endosperm has completed a several rounds of nuclear division. I, Ovule from a CTF7-OE line. It shows no difference with wild-type FG1 ovule. The degraded megaspore is visible. J, Ovule from a CTF7-OE line. The embryo sac is enlarged but the female gametophyte remains at FG1. K, Ovule from a CTF7-OE line. The female gametophyte is degraded. Scale bars = 5μm.
Figure 7. Cellular distribution of CTF7.
A-D, 35S driven CTF7::YFP localization during female gametophyte development. A, FG1 ovule, arrow indicates FG nucleus. B, Bright field of image A, position of FG is circled. C, FG2 ovule, arrows indicate nuclei. D, FG7 ovule, arrows indicate nucleus. E-H, Aborted female gametophytes in 35S-CTF7::YFP plants. E, Aborted FG1 stage ovule, arrow indicates degraded nucleus. F, Bright field of image E. G, Aborted ovule at early FG1 ovule, arrow indicates degraded nucleus. H, Aborted ovule, arrow indicates the trace of degraded nucleus. I, Ovule from genomic-CTF7::YFP transgenic plant; globular staged embryo. J, Ovule from genomic-CTF7::YFP transgenic plant; heart staged embryo. K, Ovule from genomic-CTF7::YFP transgenic plant; torpedo staged embryo. L, Ovule from wild type, negative control plant. M-P, CTF7 immunolocalization on seed sections; M, torpedo stage embryo; N, single embryo at early torpedo stage; O, nitrocellulose stage embryo. P, Negative control with no primary antibody added. Bar=5 μm