AtTFC B is involved in control of cell division

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1. ABSTRACT

Tubulin-folding cofactors play important roles in regulating plant development. Arabidopsis tubulin-folding cofactor B (AtTFC B) is an Arabidopsis homolog of mammalian tubulin-folding cofactor B, whose biological function in plant development remains poorly understood. Here we report that the homozygous attfc b (-/-) allele caused embryonic lethality. Embryogenesis was arrested at early embryo stage and the cells contained one or multiple nuclei. Plants carrying a heterozygous attfc b (+/-) allele exhibited enlarged mesophyll cells and leaf epidermal cells with bulged nuclei. Flow cytometry analysis showed increased ploidy in the leaves of the attfc b (+/-) mutant, as well as increased levels of Cdc2A and CycB1;1. In addition, immunofluorescence assay showed increased numbers of spindles and phragmoplasts in the attfc b (+/-) mutant. These results suggest that AtTFC B plays an important role in plant cell division.

2. INTRODUCTION

Microtubules control fundamental cellular processes, such as intracellular transport, cell division, cell expansion, cell polarity, and chromosome segregation. During cell cycle, plant cells display four kinds of microtubule arrays: the interphase cortical microtubule array, the preprophase band (PPB), the mitotic spindle, and the phragmoplast (1). Microtubules contain highly conserved alpha/beta-tubulin heterodimers, and different proteins are involved in the tubulin biogenesis, such as phosducin-like proteins required for beta-tubulin biogenesis (2, 3). The biogenesis of tubulin is a symmetrical process. First, the nascent alpha- and beta-tubulin heterodimers are captured and partially folded by prefoldin and chaperonins (4, 5). After release from the prefoldin and chaperonins, the partially folded alpha- and beta-tubulin chains are captured by tubulin-folding cofactors B and A, respectively. These cofactors are subsequently replaced by cofactors E and D,
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respectively, and form a quaternary complex containing alpha-tubulin/E and beta-tubulin/D. When cofactor C binds to the quaternary complex, the cofactors C, D, and E act as a GTPase-activating protein (GAP), triggering hydrolysis of GTP by beta-tubulin to release alpha/beta-tubulin heterodimers (6, 7).

Tubulin-folding cofactors not only regulate the folding of tubulin heterodimers, but are also involved in the regulation of cell division associated with microtubule function. In fission yeast, the alp1 and sto1 alleles, which encode the homologs of cofactor D and E, respectively, disturb the microtubule structure leading to condensed chromosomes or unequal segregation and mitotic division defects (8-11). In Arabidopsis, KIESEL (KIS) and PORCINO (POR), the orthologs of cofactor A and C, respectively, are required for the synthesis of microtubules; mutations in these two genes cause cell division defects (12, 13). Mutation in TITAN1 (TTN1), the ortholog of cofactor D, results in cell division defect attributable to the disruption of microtubule function and endosperm with condensed chromosomes (14, 15). The PILZ group genes include POR, CHO, and PFIFERLING (PFI), which encode the Arabidopsis orthologs of tubulin-folding cofactors C, D, and E, respectively (16). In pilz mutants, embryos show an absence of microtubule arrays and condensed chromosomes (16). Although mitotic division and cytokinesis are disturbed, the process of cell cycle is not affected in the pilz mutants (16, 17).

The function of tubulin-folding cofactor B is dependent on microtubules. In most cases, mammalian cofactor B is considered as a reservoir to sequester and deliver the alpha-tubulin folding intermediate to cofactor E (18). Overexpression of cofactor B can depolymerize microtubules (19-22). Mutation in Alp11, a fission yeast homolog of cofactor B, destroys microtubule structure and delayed mitosis with condensed chromosomes (23, 24), and is also essential for cell viability in Alp11, a fission yeast homolog of cofactor B, destroyed microtubule structure and delayed mitosis with condensed chromosomes (23, 24), and is also essential for cell viability in fission yeast (23, 24). Overexpression of Alp11, a budding yeast ortholog of mammalian cofactor B, caused depolymerization of microtubules; however, Alp11 is not essential for cell survival in budding yeast (25). Different roles of cofactor B in different species suggest a complexity of its function. AtTFC B (AT3G10220) is the Arabidopsis homolog of mammalian cofactor B (17), and it can interact with alpha-tubulin in vivo. Overexpression of AtTFC B in cowpea protoplasts reduced the number of microtubules (26). However, the function of AtTFC B in plant development remains unknown.

Here, we provide evidence that AtTFC B is essential for embryogenesis in Arabidopsis. Cellular and molecular data indicate that AtTFC B is also involved in the regulation of post-embryonic development. Our data suggest that AtTFC B is required for cell division.

3. MATERIALS AND METHODS

3.1. Plant growth, mutant screening, and plant transformation

All Arabidopsis materials used in this study are in the Columbia (Col) background. The T-DNA insertion lines attfc b (+/-) (SALK-019471) and attfc b-2 (SALK-002231) were obtained from the Arabidopsis Biological Resource Center (ABRC, Columbus, OH). Seeds were surface-sterilized with 10% sodium hypochlorite, sown on Murashige and Skoog medium (27) supplemented with 0.8% agar and 3% sucrose, and cold-treated for one day at 4°C. Seedlings were then grown in the medium under a 16 h light photoperiod at 21°C. Adult plants were grown in environmental chambers under a 16 h photoperiod at 21°C.

We genotyped the attfc b (+/-) mutant plants by PCR using the following specific primers, LBa1 (5’-T G T T T A C C T G T G C A C T C C -3’), BLB (5’-T G C T T A C A C T T G C A G G G T G A A -3’), and BRP (5’-G G G C T T T C G G G T T G T G C -3’). The primers were designed from SALK (http://signal.salk.edu/dnprimers2.html).

To generate transgenic plants ectopically expressing AtTFC B, we cloned the coding sequence of AtTFC B into the modified plasmid pCambia3301 under the control of CaMV 35S promoter, and transformed the construct into wild-type and attfc b (+/-) mutant plants using the floral dip method (28).

3.2. Cloning of AtTFC B coding sequence

Total RNA from wild-type Arabidopsis was extracted with RNeasy® Plant Mini kit (QIAGEN, Valencia CA). To obtain the first strand cDNA of the AtTFC B (AT3G10220) gene, total mRNA was used as a template for reverse transcription reaction with oligo-dT18 and SuperScript II™ RNase H-Reverse Transcriptase (Invitrogen, CA). The primers BFP (5’-C A C C C A T A T G A T G C C A A C T T C C G G T T T A C A T T G T G A -3’), and BRP (5’-G G G C T T T C G G G T T G T G C -3’) designed from SALK were used for PCR using the following specific primers, LBa1 (5’-T G G T T T A C C T G T G C A C T C C -3’), BLB (5’-T G C T T A C A C T T G C A G G G T G A A -3’), and BRP (5’-G G G C T T T C G G G T T G T G C -3’). The primers were designed from SALK (http://signal.salk.edu/dnprimers2.html).

3.3. Quantitative real-time PCR

To determine the expression levels of AtTFC B, Cdc2A, and CycB1;1, total RNA was isolated from different tissues using the RNA plant kit (Tiangen Biotech, Beijing). The RNA was quantified three times for each sample. To produce the first strand cDNA, 2 µg of total RNA pretreated with DNAase I (Promega, Madison, WI) was used as a template in reverse transcription reaction. At the end of the reaction, the cDNA was diluted with sterile distilled water. Real-time PCR was performed using the SYBR green-I qPCR kit (Finnzymes Oy). Fifty nanograms of the diluted cDNA and 0.75 µM gene-specific primers were used in the SYBR green-I qPCR kit (Finnzymes Oy). Fifty nanograms of the diluted cDNA and 0.75 µM gene-specific primers were used in the SYBR green-I qPCR kit (Finnzymes Oy).
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DNA Engine Opticon™ 2 machine (Bio-Rad, CA, USA). The 
Ubiquitin10 gene served as an internal control using specific primers UBQ10FP (5’-G A T C T T G C C G G A A A A C A T T G G G A T G G A T T G G T-3’) and UBQ10RP (5’-C G A C T T G T C A T T A G A A A G A A G A G A G A C A G G G-3’). PCR conditions were as follows: 5 min at 94°C for the denaturation step, 40 cycles followed with 10 s at 94°C, 20 s at 60°C, and 20 s at 72°C. Fluorescence of the PCR products was measured twice at the end of the extension step at 72°C and 84°C. Relative transcript levels of AtTFC B, Cdc2A, and CycB1;1 were calculated using the following equation: RE=2^{ΔΔCt}. For wild-type and attfc b (+/-) mutant plants, samples from three different individuals were measured.

3.4. Examination of ovaules

Siliques at different developmental stages were harvested and dissected in Hoyer’s solution (7.5 g of arabic gum, 100 g of trichloroacetaldehyde hydrate, and 5 mL of glycerol diluted with water to 100 mL) under a stereomicroscope. The ovaules were examined immediately under a Zeiss Axioskop2 plus Microscope and digital images were acquired using Carl Zeiss imaging systems (Carl Zeiss, Jena, Germany).

3.5. Leaf morphology and structure

Leaf epidermal cells were examined with a Field Emission Environmental Scanning Electron Microscopy (model QUANTA 200F, FEI, OR). To study the mesophyll structure, leaves were fixed in 4% (v/v) paraformaldehyde (Sigma-Aldrich, St. Louis, MO), washed, dehydrated in a series of ethanol, and embedded in LR White resin (London Resin Co., London, United Kingdom). Five micrometer-thick sections were obtained with a glass knife on a microtome (Model 1512, Leitz), mounted onto slides, and stained with 0.1% (w/v) toluidine blue-O. The slides were examined with a Leica microscope (Type 020-525.025, Leica Microsystems, Wetzlar, Germany) and photographed with a KX Series Imaging System (Model KX32E, Apogee Instruments Inc., Roseville, CA). The nuclei of leaf epidermal cells were detected as previously described by Boudolf and his colleagues (29). The nuclei and cell sizes of leaf epidermal cells and mesophyll cells were measured using Image J 1.36b software (NIH, Bethesda, MD).

3.6. Analysis of microtubule

Microtubule arrays were detected in roots based on previously described (30). The root tips were dissected from seven-day-old seedlings grown under light conditions and fixed in 4% (v/v) paraformaldehyde for immunolabeling, the root samples were first incubated with a mouse anti-alpha-tubulin monoclonal antibody (Sigma-Aldrich) at a 1:200 dilution, washed and then incubated with FITC-conjugated goat anti-mouse IgG (Molecular Probes, Eugene, OR) at a 1:500 dilution. The samples were examined under a Leica fluorescence microscope (Type 020-525.025, Leica Microsystems, Wetzlar, Germany).

3.7. Flow cytometry

The third leaf was chopped with a razor blade in 0.5 mL of cold Galbraith buffer (31). The extracts were filtered through a 60 µm pre-wet nylon mesh and collected in a 1.5 mL eppendorf tube. The filtrate was incubated with 10 µL of 10 mg/mL RNaseA for 20 min at 37°C, and then stained with 2 µL of 5 mg/mL propidium iodide (Sigma-Aldrich) and gently dispersed the nuclei. The sample was kept at 4°C in dark conditions for 10 min. Approximately 3500-5500 nuclei were analyzed with FACS Calibur flow cytometer, and DNA histograms and proportions were generated using Cellquest software (Becton, Dickinson and Company, Franklin Lakes, NJ).

3.8. Statistical analyses

Data were analyzed using student t-test for statistical significance and all images were processed with Photoshop™ 7.0.1 (Adobe Systems Inc., San Jose, CA).

4. RESULTS

4.1. The homozygous attfc b (-/-) allele caused embryonic lethality

To understand the function of AtTFC B in plant development, a T-DNA insertion line (SALK-019471) was obtained from ABRC (Figure 1A). The T-DNA was inserted at position +1846 of the 3’ region of AtTFC B genomic sequence (Figure 1B). To test if this insertion affects transcription of AtTFC B gene, the level of AtTFC B mRNA was determined using quantitative real-time PCR. As shown in Figure 1C, AtTFC B mRNA expression was reduced in the roots, stems, leaves and siliques in the T-DNA insertion line.

In screening for attfc b mutant plants, only heterozygous mutant plants were obtained in this T-DNA insertion line (Figure 1D). Progeny analysis using the X² test for heterozygous attfc b (+/-) plants showed a segregation ratio of 1:2 for wild-type (n = 188) to heterozygous plants (n = 321), suggesting the potential embryo lethality of the homozygous attfc b (-/-) plants. To test this hypothesis, we examined the phenotypes of the embryos using differential interference contrast microscope. In contrast to the wild-type embryos (Figures 2A, 2C, 2E and 2G), approximately 20% of mutant embryos were arrested at early two or four cells developmental stage, carrying multiple nuclei (Figures 2B, 2D, 2F and 2H). Given the possibility that some embryos may have stopped development or degenerated at an earlier stage, the percentage of defective embryos was close to what was expected for attfc b (-/-) genotype.

4.2. Plants carrying a heterozygous attfc b (+/-) allele had enlarged cell size

To test whether AtTFC B plays a role in plant growth beyond embryogenesis, we examined the phenotypes of heterozygous attfc b (+/-) plants. Heterozygous plants had different phenotypes from the wild-type: the plants were taller with bigger rosettes (Figures 3A, 3B), and the third true leaf was significantly longer and wider (Figure 3C) (Table 1). Consistent with enlarged leaf, the mesophyll cells in attfc b (+/-) mutant were significantly larger than those in the wild-type (Figures 3D, 3E; Table 1). We found that the leaf epidermal cells of attfc b (+/-) mutant were also larger than those in...
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Figure 1. T-DNA insertion line and the expression profiles of AtTFC B. (A) A T-DNA is inserted in the 3’ region of the AtTFC B genomic sequence in the attfc b (+/-) mutant. The open arrowhead indicates the left border of the T-DNA. Solid rectangles denote exons and the horizontal lines indicate the 5’ leader, 3’ trailer, and introns. The arrows indicate the positions of the BLP, BRP, and LBa1 primers. (B) The nucleotide position of T-DNA insertion site. One T-DNA is inserted at the +1846 position of the AtTFC B genomic sequence. The underlined nucleotides represent the T-DNA sequence. The AtTFC B genomic sequence is derived from the MIPS database (http://mips.gsf.de/proj/thal/db/index.html). (C) Relative expression levels of AtTFC B in different organs of wild-type and attfc b (+/-) plants. The accumulation levels of AtTFC B mRNA in different organs were reduced in attfc b (+/-) plants, as determined by quantitative real-time PCR. Each dataset represents the average from three biological replicates plus the standard error. (D) Genotyping of attfc b (+/-) mutant plants by PCR. Using gene-specific primers BLP and BRP for AtTFC B, and a T-DNA-specific primer LBa1, the PCR results confirmed a T-DNA insertion in the attfc b (+/-) plant as shown in (A). Col = Wild-type. M = λ DNA /EcoRI+HindIII molecular weight marker.

the wild-type (Figures 4A, 4B). The axial length in the mutant (52.2 µm) was significantly longer than that of the wild-type (44.8 µm) (Table 1), and the leaf epidermal cells contained enlarged nuclei (Figure 4D, Table 1).

To confirm that mutations in AtTFC B caused leaf morphological defects, we analyzed the phenotype of another attfc b mutant (SALK-002231) named attfc b-2. The attfc b-2 mutant also exhibited enlarged leaf sizes (Figure 5). These results indicate that AtTFC B play an important role in post-embryonic growth in Arabidopsis.

4.3. Overexpression of AtTFC B restored the attfc b (+/-) phenotype and inhibited growth in the wild-type background

We further tested if the expression of AtTFC B in attfc b (+/-) mutant background would rescue the attfc b mutant phenotype. We expressed the AtTFC B coding sequence (CDS) under the control of CaMV 35S promoter (Figure 6A). Phenotypic analysis showed that the expression of AtTFC B restored attfc b (+/-) phenotype to the wild-type phenotype (Figures 6B, 6C, Table 2).

The phenotypes of attfc b (+/-) plants suggested that AtTFC B functions to inhibit cell and organ growth in post-embryonic development. In this scenario, we would expect that overexpression of AtTFC B in the wild-type background would lead to inhibition of plant growth. Indeed, overexpression of AtTFC B in the wild-type background displayed reduction in both rosette size and leaf size (Figures 6B, 6D, Table 2).

4.4. Mutation in AtTFC B altered ploidy levels and mRNA levels of Cyclin B1;1 and Cdc2A

Homozygous attfc b (-/-) embryos displayed impaired nuclear division (Figures 2B, 2D, 2F, 2H) and the heterozygous attfc b (+/-) plants showed enlarged nuclei in leaf epidermal cells (Figure 4D). To further test the
AtTFC B controls cell division

Table 1. Statistics of leaf structural measured in wild type and attfc b (+/-) mutant plants

<table>
<thead>
<tr>
<th>Plant lines</th>
<th>Rosette diameter (cm)</th>
<th>Leaves</th>
<th>Leaf epidermal cells</th>
<th>Mesophyll cells</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Length (mm)</td>
<td>Width (mm)</td>
<td>Axial length (mm)</td>
</tr>
<tr>
<td>Wild type</td>
<td>3.3 +/- 1.0</td>
<td>(n = 38)</td>
<td>4.5 +/- 1.3</td>
<td>3.0 +/- 0.8</td>
</tr>
<tr>
<td>attfc b (+/-)</td>
<td>5.1 +/- 1.3</td>
<td>(n = 34)</td>
<td>6.1 +/- 1.7</td>
<td>3.8 +/- 0.8</td>
</tr>
</tbody>
</table>

Rosettes of three-week-old seedlings, leaves of two-week-old seedlings, and nuclei of leaf epidermal cells in three-week-old seedlings were examined. All data represent means and standard deviation, n = number of samples measured. 1T-test shows statistically significant differences between wild type and attfc b (+/-) mutant plants, and P value is less than 0.01.

Table 2. Sizes of rosettes and leaves from wild-type, rescued attfc b (+/-), and AtTFC B-overexpression plants

<table>
<thead>
<tr>
<th>Plant lines</th>
<th>Rosette diameter (cm)</th>
<th>Leaf length (cm)</th>
<th>Leaf width (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>3.5 +/- 0.9</td>
<td>(n = 43)</td>
<td>4.1 +/- 1.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3.4 +/- 0.9</td>
</tr>
<tr>
<td>RB12</td>
<td>3.5 +/- 0.8</td>
<td>(n = 30)</td>
<td>3.7 +/- 0.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3.2 +/- 0.9</td>
</tr>
<tr>
<td>OB13</td>
<td>3.0 +/- 0.5</td>
<td>(n = 18)</td>
<td>3.0 +/- 0.9</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2.4 +/- 0.8</td>
</tr>
</tbody>
</table>

Rosettes of three-week-old seedlings and leaves of two-week-old seedlings were measured. All data represent means and standard deviation. n = number of samples measured. 1Rescued attfc b (+/-) mutant line 12. 2AtTFC B-overexpression line 13. 3T-test shows statistically significant differences between wild-type and AtTFC B-overexpression plants, and P value is less than 0.01.

Table 3. DNA polyploidy levels in leaves of wild-type, attfc b (+/-) mutant, and AtTFC B-overexpression plants.

<table>
<thead>
<tr>
<th>Plant lines</th>
<th>2C (%)</th>
<th>4C (%)</th>
<th>8C (%)</th>
<th>16C (%)</th>
<th>32C (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>34.7 +/- 4.5 (n = 32)</td>
<td>54.7 +/- 3.8 (n = 35)</td>
<td>6.5 +/- 1.3 (n = 15)</td>
<td>0.5 +/- 0.20 (n = 5)</td>
<td></td>
</tr>
<tr>
<td>attfc b (+/-)</td>
<td>15.1 +/- 1.1 (n = 5)</td>
<td>45.4 +/- 9.2 (n = 30)</td>
<td>32.9 +/- 9.1 (n = 26)</td>
<td>2.0 +/- 0.84 (n = 20)</td>
<td>0.13 +/- 0.08 (n = 15)</td>
</tr>
<tr>
<td>OB13</td>
<td>29.8 +/- 3.9 (n = 3)</td>
<td>59.4 +/- 2.1 (n = 32)</td>
<td>6.1 +/- 4.1 (n = 30)</td>
<td>0.33 +/- 0.21 (n = 15)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>52.1 +/- 10.6 (n = 417)</td>
<td>38.0 +/- 12.5 (n = 273)</td>
<td>3.7 +/- 1.3 (n = 18)</td>
<td>0.4 +/- 0.20 (n = 15)</td>
<td></td>
</tr>
</tbody>
</table>

Leaves of 21 DAG seedlings grown under a 16 h photoperiod were analyzed with flow cytometry. For each sample, 3 000-5 500 nuclei were analyzed for their ploidy levels. All data represent means and standard deviation. 1AtTFC B-overexpression line 13. n = number of samples examined.

Table 4. Percentages of the four-microtubule arrays in root tip cells of wild-type and attfc b (+/-) mutant plants

<table>
<thead>
<tr>
<th>Cell lines</th>
<th>Cortical microtubule</th>
<th>Prophragmoplast</th>
<th>Spindle</th>
<th>Phragmoplast</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>93.7%</td>
<td>4.0%</td>
<td>1.0%</td>
<td>1.3%</td>
</tr>
<tr>
<td>attfc b (+/-)</td>
<td>92.3%</td>
<td>4.1%</td>
<td>1.3%</td>
<td>2.3%</td>
</tr>
</tbody>
</table>

Seven-day-old seedlings that grew under a 16 h photoperiod were used to observe microtubule arrays. The microtubule arrays in root tip cells of over 80 roots from each line were studied by confocal microscopy. n = number of cells examined.

Consistent with the ploidy level, the levels of Cdc2A and CycB1;1 mRNA were upregulated in attfc b (+/-) mutant and reduced in AtTFC B transgenic plants (Figure 8).

4.5. Microtubule arrays were affected in the attfc b (+/-) mutant

We further tested if microtubule arrays were affected in the attfc b (+/-) mutant. We performed immunostaining for alpha-tubulin in root tip cells of attfc b (+/-) mutant and the wild-type. As shown in Table 4, the percentage of spindle and phragmoplasts in attfc b (+/-) mutant increased by 30% and 77%, respectively, relative to the wild-type. The PPB was slightly increased (2.5%) and the cortical microtubule arrays were reduced (1.5%) in the attfc b (+/-) mutant compared with those in wild-type plants (Table 4). These data suggested that cytokinesis might be defective in the attfc b (+/-) mutant.

5. DISCUSSION

Tubulin-folding cofactors are involved in the control of cell division. In this study, we showed that AtTFC B, the homolog of mammalian cofactor B, is...
Figure 2. Embryo development in wild type and *attfc b* (-/-) mutant plants. (A) and (B) Early-stage embryo. Embryos are shown in wild type (A) and the *attfc b* (-/-) mutant (B). Scale bar = 20 µm. (C) and (D) Early heart-stage embryos. The *attfc b* (-/-) embryo (D) is arrested at the early stage embryo in contrast to the wild-type plant (C). The *attfc b* (-/-) embryo has four cells, each containing more nuclei. Scale bar = 20 µm. (E) and (F) Torpedo-stage embryo. The *attfc b* (-/-) embryo (F) still remains at the early stage embryo in contrast to that of the wild-type plant (E). The arrowhead indicates the embryo. Scale bar = 20 µm. (G) and (H) Mature embryos. The *attfc b* (-/-) embryo (H) remains arrested at the four-cell stage, with each cell containing one or more nuclei, compared with that of the wild-type plant (G). The arrowhead indicates the embryo. Scale bar = 20 µm.
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Figure 3. Growth heights and leaf structures in wild-type and attfc b (+/-) mutant plants. (A) Morphogenesis of nine-week-old plants. attfc b (+/-) mutant plants (right) are taller than wild-type plants (left). Scale bar = 10 cm. (B) Three-week-old rosettes of wild-type (left) and attfc b (+/-) mutant (right) plants. The mutant plant is larger than the wild-type. Scale bar = 10 mm. (C) The third leaves from 14-day-old wild-type (left) and attfc b (+/-) mutant plants (right). The mutant leaf is longer and wider than the wild-type. Scale bar = 2 mm. (D) and (E) Transverse sections of leaves showing mesophyll cells in wild-type (D) and attfc b (+/-) mutant (E) plants. Asterisks point to some enlarged mesophyll cells in the attfc b (+/-) mutant plants. Scale bar = 50 µm.

Figure 4. Nucleus morphogenesis of leaf epidermal cells in wild-type and attfc b (+/-) mutant plants. (A) and (B) Leaf epidermal cells in 14-day-old wild-type (A) and attfc b (+/-) mutant plants (B). The mutant plant has enlarged epidermal cells. Scale bar = 25 µm. (C) and (D) Nucleus phenotypes in 21-day-old wild-type (C) and attfc b (+/-) mutant plants (D). Nucleus was stained with DAPI. The mutant plant has bulged nuclei in the leaf epidermal cells. An arrow head points to the nucleus of leaf epidermal cells, and an arrow shows the nucleus of guard cells. Scale bar = 25 µm.
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Figure 5. Leaf size in attfc b-2 mutant plants. The leaf length of the attfc b-2 (6.3 +/- 1.0 mm, n = 43) was longer than that of the wild-type plants (4.6 +/- 0.6 mm, n = 22), and the leaf width of attfc b-2 (4.2 +/- 0.7 mm, n = 43) was also wider than wild-type plants (3.6 +/- 0.5 mm, n = 22). Col = Wild-type. Asterisks showed statistically significant differences between attfc b-2 and wild-type (t-test, P value is less than 0.01). Bars represent the standard deviation.

Figure 6. Effects of ectopic AtTFC B expression on the growth of attfc b (+/-) and wild-type plants. (A) The expression levels of AtTFC B mRNA in the rescued attfc b (+/-) mutant and AtTFC B-overexpression plants. AtTFC B coding sequence (CDS) under the control of CaMV 35S promoter was transformed into the attfc b (+/-) mutant and wild-type background, respectively, to obtain the rescued attfc b (+/-) mutant and AtTFC B-overexpression transgenic plants. Each dataset represents the average of three independent experiments plus the standard deviation. Col = Wild-type. RB12 = Rescued attfc b (+/-) mutant line 12. OB13 = AtTFC B-overexpression transgenic line 13. (B) Rosettes of three-week-old wild-type plant. Scale bar = 10 mm. (C) Rosettes of three-week-old rescued attfc b (+/-) mutant line 12. The rescued plant shows a similar morphology to that of the wild-type plant (B). Scale bar = 10 mm. (D) Rosettes of three-week-old AtTFC B-overexpression transgenic line 13. The plant produce greatly reduced rosette compared with the wild-type (B). Scale bar = 10 mm.

required for cell division in Arabidopsis development. Specifically, embryos of homozygous attfc b (-/-) plants were arrested at early developmental stage and cells have multiple nuclei. In the PILZ group genes, mutant embryos contain varying numbers of nuclei and lack microtubules such as the spindle and phragmoplast (16, 17). Mutation in the EDE1 gene caused defective endosperm, which was consistent with the absence of mitotic spindles and phragmoplasts (32). In the attfc b (+/-) mutant, the number of spindles and phragmoplasts was increased in root cells (Table 4). Given the direct role of the phragmoplast in the formation of new cell plate during cytokinesis (33), the possibility that the cytokinesis arrest resulted from the phragmoplast defect in the attfc b (-/-) mutant and caused improper cell division can not be excluded.

Besides cell division defect in embryogenesis, the heterozygous attfc b (+/-) mutant caused larger leaf epidermal cells (Figure 4B, Table 1), while AtTFC B-overexpressing plants were smaller in leaf size (Table 2). The distinct leaf phenotypes between attfc b (+/-) mutant and AtTFC B-overexpressing plants, suggest that AtTFC B
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Figure 7. Ploidy levels in the leaves of wild-type, attfc b (+/-) mutant, and AtTFC B-overexpression plants measured by flow cytometry. The ploidy levels of the third leaves of 21-day-old plants are represented. The polyploidy levels were increased in the attfc b (+/-) mutant but reduced in AtTFC B-overexpression line 13, compared with those of wild-type. Col = Wild-type. OB13 = AtTFC B-overexpression line 13.

Figure 8. The expression levels of Cdc2A and CycB1;1 in the attfc b (+/-) mutant and AtTFC B-overexpression plants. (A) The levels of Cdc2A were increased in the attfc b (+/-) mutant but decreased in AtTFC B-overexpression line OB13. (B) The levels of CycB1;1 were increased in the attfc b (+/-) mutant but decreased in AtTFC B-overexpression line OB13. Each dataset represents the average of three independent experiments plus the standard deviation. Col = Wild-type. OB13 = AtTFC B-overexpression line 13.
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functions as a negative regulator of cell division, at least in the post-embryonic organs.

Cell division is tightly related to cell cycle. Although function of tubulin-folding cofactors could affect cell division, the roles of tubulin-folding cofactors in cell cycle are different. Mammalian cells transfected with cofactor A siRNA were arrested at G1 phase (34), and overexpression of Alf1, the budding yeast homolog of cofactor B, arrested cell cycle causing defects in microtubule function (25). However, in pilz group mutant plants, cell cycle was not arrested even when mitotic division was arrested (16). In this study, the formation of PPB was still present in the root meristem cells (Table 4), indicating that G2-to-M transition point of the cell cycle was not arrested in attfc b (+/-) mutant. However, the number of phragmoplasts was obviously increased in attfc b (+/-) mutant (Table 4), suggesting that cell cycle might be arrested at the mitotic telophase, which resulted in cytokinesis defect.

Mutation of AtTFC B showed enlarged nuclei in leaf epidermal cells and was consistent with higher ploidy levels in the leaves (Figures 4D, 7B). A previous study showed that enhanced levels of a non-destructible Cyclin B1 (also named CycB1;1) leads to doubled DNA content resulting from endomitosis. Weingartner and his colleagues concluded that the mitotic cyclins were required for reorganization of the phragmoplast and for proper cytokinesis (35). In this study, mutation in AtTFC B upregulated levels of Cdc2A and CycB1;1 (Figure 8), as well as the number of phragmoplasts (Table 4). In contrast, overexpression of AtTFC B decreased the levels of Cdc2A and CycB1;1 (Figure 8) and the polyplody levels (Figure 7C). Thus, endomitosis might have occurred in attfc b (+/-) mutants when cytokinesis was disturbed. Mutation in AtTFC B also increased the number of spindles (Table 4), and increased mRNA levels of Cdc2A (Figure 8A). Cdc2 activity is required for microtubule recruitment. It is located at the nucleus and is associated with spindles and phragmoplasts during mitosis and cytokinesis (36). Taking these phenotypes into account, our data suggest that AtTFC B plays a critical role in ensuring a normal cell division process, and AtTFC B mutation might promote cells to enter into the process of endomitosis when cell division is arrested.

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7. REFERENCES

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**Abbreviations:** AtTFC B: Arabidopsis tubulin-folding cofactor B; PPB: preprophase band; GAP: GTPase-activating protein; Col: Columbia; ABRC: Arabidopsis Biological Resource Center; KIS: KIESEL; POR: PORCINO; TTN1: TITAN1; PFI: PFIFFERLING

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