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Two Mechanosensitive Channel Homologs Influence Division Ring Placement in *Arabidopsis* Chloroplasts

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Chloroplasts must divide repeatedly to maintain their population during plant growth and development. A number of proteins required for chloroplast division have been identified, and the functional relationships between them are beginning to be elucidated. In both chloroplasts and bacteria, the future site of division is specified by placement of the Filamentous temperature sensitive Z (FtsZ) ring, and the Min system serves to restrict FtsZ ring formation to mid-chloroplast or mid-cell. How the Min system is regulated in response to environmental and developmental factors is largely unstudied. Here, we investigated the role in chloroplast division played by two *Arabidopsis thaliana* homologs of the bacterial mechanosensitive (MS) channel MscS: MscS-Like 2 (*MSL2*) and *MSL3*. Immunofluorescence microscopy and live imaging approaches demonstrated that *msl2 msl3* double mutants have enlarged chloroplasts containing multiple FtsZ rings. Genetic analyses indicate that *MSL2*, *MSL3*, and components of the Min system function in the same pathway to regulate chloroplast size and FtsZ ring formation. In addition, an *Escherichia coli* strain lacking MS channels also showed aberrant FtsZ ring assembly. These results establish MS channels as components of the chloroplast division machinery and suggest that their role is evolutionarily conserved.

INTRODUCTION

Chloroplasts are specialized organelles responsible for many essential metabolic reactions, most notably photosynthesis. Like all plastids, chloroplasts are not created de novo but are differentiated from a preexisting population of proplastids present in meristematic cells (reviewed in Lopez-Juez and Pyke, 2005). As plant cells expand, established chloroplasts must undergo a series of divisions to maintain the appropriate population density. Modern day chloroplasts arose from an endosymbiotic relationship between a heterotrophic protozoan and a cyanobacterium and, reminiscent of their bacterial origins, divide by binary fission (Cavalier-Smith, 2000; Miyagishima et al., 2003). Given this evolutionary lineage, bacterial cell division is often used as a framework for identifying the components and understanding the mechanisms of chloroplast division. This approach has led to the identification of many homologs of essential bacterial division proteins within the plant genome, and a strong evolutionary conservation of division mechanisms has now been documented (Yang et al., 2008; reviewed in Nakanishi et al., 2009b).

Cell division has been well characterized in the gram-negative bacterium *Escherichia coli*. Division at mid-cell is specified by the polymer-forming GTPase Filamentous temperature sensitive Z (FtsZ), which forms a structure known as the Z-ring on the cytoplasmic surface of the inner membrane (Bi and Lutkenhaus, 1994). The Z-ring is thought to act as a scaffold for the assembly of the division apparatus and contribute some of the force required for fission (Osawa et al., 2009; reviewed in de Boer, 2010). Two nonredundant families of FtsZ genes are found in seed plants (Osteryoung and Vierling, 1995; Osteryoung et al., 1998; Stokes and Osteryoung, 2003; Schmitz et al., 2009), and both FtsZ1 and FtsZ2 localize to a ring-like structure at the division site (McAndrew et al., 2001; Vitha et al., 2001; Fujiwara et al., 2008). Plant and bacterial lines under- or overexpressing FtsZ exhibit enlarged chloroplasts and filamentous cells, respectively, though plants with altered levels of FtsZ are otherwise phenotypically normal (Dai and Lutkenhaus, 1992; Stokes et al., 2000; El-Kafafi et al., 2005, 2008; Liu et al., 2007; Yoder et al., 2007; Schmitz et al., 2009; Karamoko et al., 2011). *Arabidopsis thaliana* FtsZ proteins have GTPase activity and are capable of forming filaments in vitro (El-Kafafi et al., 2005; Olson et al., 2010; Smith et al., 2010).

Proper placement of the Z-ring ensures segregation of cytoplasmic contents and is tightly regulated in *E. coli* by the Min system (Bi et al., 1991; reviewed in de Boer, 2010). Minicell C (MinC) and MinD form a complex that negatively regulates FtsZ polymerization, while MinE acts as a topological specificity factor by inhibiting MinCD function (de Boer et al., 1989, 1991, 1992b; Huang et al., 1996). All three proteins oscillate from pole to pole such that MinCD activity is restricted to the ends of the cell and FtsZ polymerization is allowed to progress only at mid-cell (Hu and Lutkenhaus, 1999; Raskin and de Boer, 1999a, 1999b; Rowland et al., 2000; Fu et al., 2001; Hale et al., 2001). *Arabidopsis* homologs of MinD and MinE have been identified and appear to have some conserved functions in regulating Z-ring placement. *Arabidopsis* MinD antisense lines and a loss-of-function allele of MinD, *accumulation and replication of..."
chloroplasts11 (arc11), exhibit enlarged chloroplasts with multiple Z-rings, suggesting that At-MinD behaves as a Z-ring assembly inhibitor (Colletti et al., 2000; Kanamaru et al., 2000; Vitha et al., 2003; Fujiwara et al., 2004). Conversely, At-MinE antisense suppression lines and two mutant alleles of At-MinE, minE1 and arc12, exhibit grossly enlarged chloroplasts containing numerous short FtsZ filaments (Glynn et al., 2007; Fujiwara et al., 2008). In summary, multiple lines of evidence indicate that, similar to their bacterial homologs, At-MinD and At-MinE antagonistically regulate Z-ring assembly (Itoh et al., 2001; Maple et al., 2002; Reddy et al., 2002; Fujiwara et al., 2004; Glynn et al., 2007; Maple and Møller, 2007b; Fujiwara et al., 2008).

Several aspects of FtsZ ring assembly in chloroplasts are specific to plants (Nakanishi et al., 2009b). Arabidopsis ARC3, a stromal protein that contains an incomplete FtsZ-like region, has been reported to interact directly with MinD, MinE, and FtsZ and copurify with FtsZ1, FtsZ2, and ARC6 (Shimada et al., 2004; Maple et al., 2007; McAndrew et al., 2008). arc3 mutants resemble arc11 mutants in that they exhibit enlarged chloroplasts and multiple FtsZ rings (Pyke and Leech, 1994; Shimada et al., 2004; Glynn et al., 2007; Maple et al., 2007). No MinC homolog has been identified in plants, though FtsZ assembly is sensitive to overexpression of E. coli MinC, suggesting that the system is capable of responding to MinC activity (Tavva et al., 2006). Instead, it has been proposed that ARC3 has taken on the role of MinC in land plants and directly inhibits FtsZ polymerization by actively competing with FtsZ monomers (Maple and Møller, 2007b; Maple et al., 2007; Yang et al., 2008; Nakanishi et al., 2009b). Other plant-specific genes required for normal FtsZ assembly and Z-ring placement are MscS-Like 2 (MSL2) and MSL3. MSL2 and MSL3 are redundantly required for normal chloroplast size and colocalize with MinE at the poles of plastids (Haswell and Meyerowitz, 2006). Single msl2-1 and msl3-1 mutants appear wild type, suggesting that MSL2 and MSL3 function redundantly. In this study, we extend our characterization of the msl2-1 msl3-1 double mutant and add another allele of MSL2 (msl2-3) to our analysis. The T-DNA insertion in the msl2-3 allele introduces a stop codon at amino acid 98 and is therefore likely to be a null allele, whereas the insertions in msl2-1 and msl3-1 alleles are located in the last exon of each gene and are likely to be partial loss-of-function alleles (Figure 1A; see Supplemental Figure 1 online). Though msl2-3 single mutants do not exhibit appreciably enlarged chloroplasts, msl2-3 msl3-1 mutants have enlarged chloroplasts similar to those observed in the msl2-1 msl3-1 double mutant (see Figure 3).

![Figure 1](image)

**RESULTS**

**MSL2 and MSL3 Are Required for Normal Z-Ring Placement**

We previously observed that msl2-1 msl3-1 double mutant plants exhibit heterogeneously sized chloroplasts, as well as other plastidic and whole-plant phenotypes (Haswell and Meyerowitz, 2006). Single msl2-1 and msl3-1 mutants appear wild type, suggesting that MSL2 and MSL3 function redundantly. In this study, we extend our characterization of the msl2-1 msl3-1 double mutant and add another allele of MSL2 (msl2-3) to our analysis. The T-DNA insertion in the msl2-3 allele introduces a stop codon at amino acid 98 and is therefore likely to be a null allele, whereas the insertions in msl2-1 and msl3-1 alleles are located in the last exon of each gene and are likely to be partial loss-of-function alleles (Figure 1A; see Supplemental Figure 1 online). Though msl2-3 single mutants do not exhibit appreciably enlarged chloroplasts, msl2-3 msl3-1 mutants have enlarged chloroplasts similar to those observed in the msl2-1 msl3-1 double mutant (see Figure 3).
The enlarged chloroplast phenotype of msl2-1 msl3-1 and msl2-3 msl3-1 double mutant plants suggested that chloroplast division might be altered in these mutants. To test this hypothesis, we characterized Z-ring placement in wild-type and mutant chloroplasts by immunofluorescence microscopy (IFM). The primary antibody used in these experiments was an anti-FtsZ antibody raised against FtsZ from Bacillus subtilis (a gift of P. Levin, Washington University). This antibody recognized FtsZ1, but not FtsZ2 or FtsZ2-2, in an immunoblot of Arabidopsis proteins (see Supplemental Figure 2 online). Protoplasts isolated from the rosette leaves of 3-week-old plants were fixed and chloroplasts released onto poly-l-lysine–treated slides by tapping on the cover slip. After incubation with anti-FtsZ primary and fluorescein isothiocyanate (FITC)-conjugated secondary antibodies, confocal laser scanning microscopy was used to capture images of representative chloroplasts from each genotype (Figure 1B). Most wild-type chloroplasts contained only one Z-ring, though occasionally more were observed (Figure 1C). By contrast, msl2 msl3 mutant chloroplasts frequently contained multiple Z-rings, some with a disorganized, forked appearance, similar to those observed in the arc11, arc3, mcd1, and parc6 mutants but distinct from those observed in other plastid division mutants, such as arc5, plastid division1 plastid division2 (pvd1 pvd2), or arc2 (Vitha et al., 2003; Miyagishima et al., 2006; Glynn et al., 2007, 2009; Fujiwara et al., 2008; Nakashima et al., 2009a; Suzuki et al., 2009).

Z-ring number, like chloroplast size, is highly heterogeneous in msl2 msl3 mutants (Figure 1B, bottom right panel). We therefore quantified the number of complete Z-rings per chloroplast in wild-type, msl2 msl3, and arc11 plants. After IFM, the number of intact Z-rings was counted; complete Z-rings were identified by adjusting the focal plane to visualize both the top and bottom of each ring. Figure 1C presents data compiled from three independent experiments. In wild-type plants, 83 to 87% of chloroplasts had only one Z-ring, whereas arc11 mutants were at the opposite extreme, containing 91% of chloroplasts with two or more Z-rings. The msl2-1 msl3-1 and msl2-3 msl3-1 mutants exhibited a phenotype similar to the arc11 mutant, with 59 and 75% of the chloroplasts containing two or more Z-rings, respectively. Thus, not only are MSL2 and MSL3 required for normal Z-ring assembly, but the defect in Z-ring placement in msl2 msl3 mutants is nearly as severe as that observed in the absence of functional MinD, a primary component of the plastid division machinery.

To characterize Z-ring formation in live cells, we obtained a transgenic line expressing FtsZ1-GFP (green fluorescent protein) under the control of the FtsZ1-1 upstream region (M. Fujiwara, RIKEN). This and other FtsZ-GFP fusion proteins have been used in several studies of chloroplast division in Arabidopsis petioles and pollen (Vitha et al., 2001; Fujiwara et al., 2008, 2009, 2010). In wild-type lines with low levels of FtsZ1-GFP expression, a single ring of GFP fluorescence was observed in caulisle leaf chloroplasts (arrows, left panel, Figure 2A). However, when the FtsZ1-GFP transgene was crossed into the msl2-3 msl3-1 background, grossly enlarged chloroplasts containing multiple Z-rings were observed (arrows, right panel, Figure 2A). Many of these enlarged chloroplasts also contained partial or branched rings.

To avoid the well-documented phenotypic effects of At-FtsZ overexpression (Stokes et al., 2000; Vitha et al., 2001; Fujiwara et al., 2008), live imaging analysis was performed only on lines expressing FtsZ1-GFP at low levels. To establish further that the increased Z-ring number observed in the msl2 msl3 mutant chloroplasts cannot be attributed to overexpression of FtsZ1-GFP, quantitative immunoblotting of protein extracts from the plants imaged in Figure 2A was performed. Similar levels of FtsZ1-GFP were present in both mutant and wild-type backgrounds (Figure 2B).

MSL2 and MSL3 Function in the Same Pathway as MinD, MinE, and ARC3

We next wished to determine whether MSL2 and MSL3 regulate Z-ring formation through the previously characterized Min pathway. To address this question, we generated and characterized triple msl2 msl3 arc11, msl2 msl3 arc3, and msl2 msl3 arc12 mutant plants. The arc11 and arc3 alleles can be considered null as MinD and ARC3 proteins are undetectable in arc11 and arc3 mutants, respectively (Shimada et al., 2004; Nakashima et al., 2009a). The arc12 allele is also likely to be null, as it introduces a stop codon upstream of the conserved regions required for dimerization and interaction with MinD (Itô et al., 2001; Glynn et al., 2007; Maple and Moller, 2007a). We hypothesized that if MSL2 and MSL3 function in the same pathway as components of the Min system to regulate Z-ring assembly, triple mutants should exhibit the phenotype of the most severe parental mutant. However, if MSL2 and MSL3 act in a separate pathway, the triple mutants should display an additive or even synergistic chloroplast phenotype.

Figure 2. Multiple Rings of FtsZ1-GFP Signal Are Observed in msl2 msl3 Chloroplasts.

(A) FtsZ1-GFP localization in cauline leaves. Arrowheads indicate Z-rings present in Col-0 and msl2-3 msl3-1 plants, respectively. GFP (green) and chlorophyll fluorescence (red) are pseudocolored. Bars = 10 μm.

(B) Quantitative immunoblot of FtsZ1-GFP protein levels in wild-type and mutant backgrounds. Three 1.5-fold dilutions of each protein extract were loaded. Top panel, GFP signal; bottom panel, Ponceau S staining of ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco). Estimates of FtsZ1-GFP protein levels in msl2-3 msl3-1 mutants relative to those in Col-0 are shown below each lane.
Fixed mesophyll cells from arc11, arc12, and arc3 mutants contain a small number of greatly enlarged chloroplasts, whereas msl2 msl3 mutants contain relatively more, heterogeneously sized chloroplasts (Figure 3A). At this level of analysis, all three triple msl2 msl3 arc mutants resembled their arc mutant parent with respect to chloroplast size and number. To compare these mutants in a quantitative manner, we counted the number of chloroplasts in optical sections of mesophyll cells from each genotype. Confocal images were taken of mesophyll tissue at the tips of fully expanded rosette leaves that had been exposed to strong light to induce chloroplast movement to the sides of the cell. Representative images are presented in Supplemental Figure 3 online. The number of chloroplasts per section at the widest circumference of each cell was counted, and the results from two independent experiments compiled (Figure 3B). Wild-type accessions had an average of 12 to 14 chloroplasts per optical section, while the msl2 msl3 double mutant had an average of 9. We found that msl2 msl3 arc11, msl2 msl3 arc12, and msl2 msl3 arc3 triple mutants closely resembled the single arc11, arc12, and arc3 mutants, with five, two, and four chloroplasts per optical section, respectively (Figure 3B). The small but statistically significant decrease (Student’s t test, P > 0.01; less than one chloroplast per optical section) in chloroplast number seen in the msl2 msl3 arc11 and msl2 msl3 arc3 triple mutants is likely due to the variability in cell size seen in the msl2 msl3 mutant background. Cell size is linearly correlated with chloroplast number (Pyke and Leech, 1992).

We next analyzed Z-ring formation in the chloroplasts of single, double, and triple msl and arc mutant plants. FtsZ IFM revealed multiple Z-rings in chloroplasts isolated from msl2 msl3 arc11 plants, similar to those observed in the msl2 msl3 and arc11 genotypes (Figure 4). Consistent with the number of chloroplasts per cell, no statistically significant difference in the percentage of chloroplasts containing two or more rings was observed in msl2 msl3 arc11 triple mutants compared with single arc11 mutants (Student’s t test, P > 0.2). Similarly, msl2 msl3 arc3 triple mutants exhibited no significant increase in Z-ring number over the arc3 single mutant (Student’s t test, P > 0.2; see Supplemental Figure 4 online).

In the analysis presented in Figure 4B, chloroplasts were put into two categories: one Z-ring or more than one Z-ring. As the latter category contained chloroplasts with Z-ring numbers ranging from 2 to 9, we analyzed the distribution of Z-ring number in the chloroplasts of single, double, and triple mutants to determine if a difference between arc11 and msl2 msl3 arc11 mutants could be observed (Figure 4C). Most chloroplasts from wild-type plants had one Z-ring; <18% contained two or three Z-rings and none had more than four. By contrast, 56 and 51% of the chloroplasts from msl2 msl3 double mutants contained two or three Z-rings, and numbers as high as 7 were observed. In the arc11 mutant, the distribution of Z-ring number was shifted further toward increasing numbers of rings per chloroplast, with 62% of the chloroplasts showing ring numbers of 3, 4, and 5. The distribution of ring number in both msl2 msl3 arc11 triple mutant lines was strikingly similar to that observed in the arc11 single mutants, suggesting that the introduction of msl2 and msl3 mutant alleles did not enhance the defect in Z-ring formation already present in the arc11 mutant.

**Figure 3.** Chloroplast Number in Single, Double, and Triple msl and arc Mutants.

(A) Chloroplasts from mature leaf mesophyll cells. Leaves were fixed with glutaraldehyde prior to imaging. Bars = 50 µm. Ler, Landsberg erecta.

(B) Quantitative analysis of chloroplast number per optical section in the indicated genotypes. The mean value ± SD is presented from data collected in two independent experiments. n ≥ 125 cells per genotype.

**MinE Is Required for Multiple Z-Rings to Form in the msl2 msl3 Background**

We next analyzed Z-ring assembly in msl2 msl3 arc12 triple mutant plants. MinE, a component of the Arabidopsis Min system, is thought to oppose the action of MinD and ARC3, acting as a positive regulator of FtsZ assembly. As previously shown (Glynn et al., 2007), Z-rings were absent in chloroplasts isolated from arc12 plants. Instead, small fragments of FtsZ polymers were detected throughout the enlarged chloroplasts (bottom left panel, Figure 5). The same phenotype was observed in chloroplasts isolated from both msl2 msl3 arc12 triple mutants (bottom middle and right panels, Figure 5). This result suggests that the formation of multiple Z-rings in msl2 msl3 double mutants requires MinE and lends further support to a model.
where MSL2 and MSL3 act in the same pathway as the Min system to regulate chloroplast division.

MS Channels Are Required for Normal Z-Ring Placement in E. coli

We next considered that MS channels might also play a role in the regulation of Z-ring placement in bacteria. To test this hypothesis, IFM using an antibody raised against E. coli FtsZ was used to visualize Z-rings in the MJF465 strain, which lacks three MS channels, MscS, MscL, and MscK, and is unable to survive extreme hypoosmotic downshock (Levina et al., 1999). No apparent difference between the growth of the wild type (Frag-1) and MJF465 was observed under standard conditions, and IFM revealed Z-rings placed at mid-cell in both backgrounds (Figures 6A to 6C and 6G to 6I). However, when treated with the septation inhibitor cephalexin, MJF465 cells frequently exhibited increased numbers of Z-rings compared with the wild type (arrows, Figures 6D to 6F and 6J to 6L). Additionally, we consistently observed polar Z-rings and double Z-rings (purple arrow and inset, Figures 6J to 6L) in MJF465 cells treated with cephalexin. Approximately half of the MJF465 cells treated with cephalexin showed these types of alterations in Z-ring assembly; in these cells, the average distance between Z-rings was 1.7 ± 0.9 μm (n = 78) compared with 5.83 ± 2.0 μm (n = 55) in Frag-1. This phenotype resembles that of the $\Delta \text{minB}$ mutant (PB114), in which the Min system operon, including $\text{minC}$, $\text{minD}$, and $\text{minE}$, is deleted (de Boer et al., 1989; Yu and Margolin, 1999). Inhibition of mini-cell formation through treatment with cephalexin revealed that PB114 mutant cells exhibit multiple and polar Z-rings, as previously shown (Adler et al., 1967; Davie et al., 1984; Yu and Margolin, 1999; see Supplemental Figure 5 online). In PB114 cells treated with cephalexin, the distance between Z-rings was 2.1 ± 0.5 μm (n = 67). Thus, MS channels contribute to normal Z-ring placement in E. coli, and Z-ring placement in MJF465 cells resembles that of $\Delta \text{minB}$ cells.

DISCUSSION

MSL2 and MSL3 were first identified based on their homology to the well-characterized bacteria MS channel MscS (Martinac and Kloda, 2003; Pivetti et al., 2003; Haswell and Meyerowitz, 2006). Based on the observation that MSL3 can rescue the osmotic...
Defective Chloroplast Division Explains the Enlarged Chloroplasts of msl2 msl3 Double Mutants

The production of enlarged chloroplasts typically results from defects in the initiation or execution of chloroplast division. Many established components of the plastid division apparatus were first identified as mutants with enlarged plastids, including nine arc mutants, cdp1, mcd1, pdv1, and br04 (Pyke and Leech, 1992, 1994; Pyke et al., 1994; Aldridge et al., 2005; Miyagishima et al., 2006; Nakanishi et al., 2009a; Suzuki et al., 2009; Zhang et al., 2009). In this analysis of msl2 msl3 mutants, FtsZ ring formation was used as a cell-based marker for normal chloroplast division. The production of a Z-ring is an early step in division, so mutations that affect FtsZ ring assembly or placement could be distinguished from defects in division downstream of FtsZ ring formation or defects in the perception of size. Both IFM and confocal imaging of an FtsZ1-GFP transgene showed that the chloroplasts in msl2 msl3 mutant plants exhibited multiple Z-rings (Figures 1 and 2). Two independent methods of analysis thus show that the enlarged chloroplasts observed in msl2 msl3 double mutants can be explained by defects in chloroplast division and, more specifically, by overactive Z-ring assembly.

MSL2 and MSL3 Are Components of the Plastid Division Machinery

The multiple Z-ring phenotype of msl2 msl3 mutants closely resembles that of previously identified lesions in factors that negatively regulate FtsZ assembly: MinD, ARC3, MCD1, and PARC6 (Figures 1B and 2A; Glynn et al., 2007, 2009; Fujiwara et al., 2008; Nakanishi et al., 2009a). Analyses of genetic lesions, overexpression studies, and protein–protein assays have demonstrated that MinD, ARC3, MCD1, and PARC6 are dedicated components of the chloroplast Min system (reviewed in Maple and Möller, 2007b, 2010; Yang et al., 2008; Miyagishima and Kabeya, 2010). Given the specific nature of the multiple Z-ring phenotype, MSL2 and MSL3 can also be considered components of the chloroplast division machinery.

MSL2 and MSL3 Regulate Z-Ring Formation through the Min System

To determine if MSL2 and MSL3 regulate FtsZ assembly through the same pathway as known components of the plastid division apparatus, we compared single, double, and triple mutants. If MSL2 and MSL3 act in the same pathway as MinD, MinE, and ARC3, the phenotype of triple msl2 msl3 arc mutants would be the same as single arc mutants. Alternatively, if MSL2 and MSL3 affect Z-ring assembly in a manner independent of Z-ring assembly, both pathways would be impaired in msl2 msl3 arc triple mutants, and an increase in phenotypic severity should be seen. As shown in Figures 3 to 5 and Supplemental Figure 4 online, msl2 msl3 arc11, msl2 msl3 arc3, and msl2 msl3 arc12 triple mutants contained the same number of chloroplasts and displayed a Z-ring formation phenotype nearly identical to those observed in single arc11, arc13, or arc12 mutants. Importantly, the phenotypes of arc11 and arc3 mutants are not so severe that an additive or even synthetic phenotype would not be discernable in msl2 msl3 arc11 or msl2 msl3 arc3 triple mutants (Figures 3 and 4; see Supplemental Figure 4 online). The lack of enhancement of these phenotypes in the triple mutants suggests that MSL2, MSL3, ARC3, and MinD regulate Z-ring formation through the...
same pathway. Additional evidence that MSL2 and MSL3 function in the same pathway as the Min system is the observation that the multiple Z-ring phenotype of msl2 msl3 double mutants required the function of MinE. Whereas the msl2 msl3 mutants exhibited chloroplasts with multiple Z-rings, no complete Z-rings were present in the triple msl2 msl3 arc12 mutant, a phenotype identical to that observed in the single arc12 mutant (Glynn et al., 2007; Figure 5).

MS Channels Affect Z-Ring Assembly in Bacteria

MSL2 and MSL3, as well as several established components of the chloroplast Min system, are evolutionarily conserved (Haswell and Meyerowitz, 2006; reviewed by Pivetti et al., 2003; Nakanishi et al., 2009). It is therefore intriguing that MS channels also play a role in the placement of Z-rings in E. coli. As shown in Figure 6, in the absence of MscS, MscK, and MscL—the major MS channels of the E. coli plasma membrane—cells treated with cephalaxin exhibited increased Z-ring number, double rings, and rings located at the poles of cells. These results are consistent with the hypothesis that MS channels in E. coli affect Z-ring placement through the Min system, as they do in plants. As Z-rings are placed solely at the sites between nucleoids (cf. white arrows in Figures 6K and 6L), the defects in Z-ring placement we observed in MFJ465 cannot be attributed to a deficit of the nucleoid occlusion pathway (reviewed in Bramkamp and van Baarle, 2009).

MS Channels Provide a Functional Link between Chloroplast Division and Membrane Tension

These results raise the question of the molecular mechanism by which MSL2 and MSL3 affect Z-ring formation. They do not appear to act at the level of gene expression, as mRNA levels of FTSZ1, FTSZ2-1, FTSZ2-2, MinD, ARC3, and MCD1 were the same in msl2 msl3 and wild-type plants (see Suppl. Figure 6 online). Furthermore, quantitative immunoblotting showed that increased levels of FtsZ-GFP are not observed in msl2 msl3 mutants; alterations in FtsZ protein levels thus cannot explain the increased Z-ring number seen in this line (Figure 2B).

As MSL2 and MSL3 each contain an N-terminal chloroplast transit peptide, they are predicted to localize to the inner envelope of the chloroplast (Haswell and Meyerowitz, 2006; Li and Chiu, 2010). Topology predictions place their large C-terminal domain inside the stroma, (ARAMEMNON database; Schwacke et al., 2003), an arrangement analogous to that of MscS within the E. coli cell. This predicted topology raises the possibility that MSL2 and MSL3 might interact directly with stromal components of the plastid division machinery. However, we were unable to detect direct interactions between the C-terminal domain of MSL3 and FtsZ1, FtsZ2-1, MinE, MinD, ARC3, or PARC6 (A. Vijayaraghavan and E.S. Haswell, unpublished data; see Methods for details) and thus favor the hypothesis that MSL2 and MSL3 indirectly regulate the Min system. Like MscS, MSL2 and MSL3 may serve to release membrane tension, perhaps derived from chloroplast expansion, and constriction during division or other stresses. If so, the increased membrane tension present in msl2 msl3 mutant chloroplasts could affect the function of membrane-associated members of the Min system. Both ARC3 and MinD are predicted to interact with the inner chloroplast envelope, ARC3 through its membrane occupation and recognition nexus (MORN) motif and MinD through C-terminal amphipathic helices (Kanamaru et al., 2000; Hu et al., 2002; Szeto et al., 2002; Shimada et al., 2004). Membrane tension could also affect the function of PARC6, a chloroplast inner envelope protein that inhibits FtsZ assembly and interacts directly with ARC3 (Glynn et al., 2009). An alternative model is that MSL2 and MSL3 are required to maintain ion homeostasis in the chloroplast stroma; in their absence, changes in ion concentration could affect the function of chloroplast division proteins. For example, MinD is a calcium-dependent ATPase (Aldridge and Møller, 2005) and its function could be altered by abnormal stromal Ca²⁺ levels.

Taken together, the data presented here indicate that MS channels play an important role in the division of Arabidopsis chloroplasts and contribute to division site placement in E. coli cells, linking membrane tension to the assembly of FtsZ rings in chloroplasts and bacteria. Future work will uncover the nature of the mechanical signal that is relayed by MSL2 and MSL3 to the Min system to properly regulate FtsZ ring placement.

METHODS

Plant Growth and Mutant Analysis

All plants were grown on soil at 21°C under a 16-h light regime. arc11 (CS281) and arc3 (CS264) mutant lines were obtained from the ABRC (Ohio State University) and are in the Landsberg erecta background. The msl2-3 mutant line (GK-195D11) was identified in the GABI-Kat collection of T-DNA insertion lines (Rosso et al., 2003) and is in the Columbia-0 (Col-0) background. The ftsZ-KO and arc12 mutant lines were a gift of K. Osteryoung and are in the Columbia-0 background (Glynn et al., 2007; Schmitz et al., 2009), msl2-1 and msl3-1 mutants are in the Wassilewskija background (Haswell and Meyerowitz, 2006). Derived cleaved-amplified polymorphic sequence genotyping of the arc mutants was performed as described by Neff et al. (1998) using the following oligo pairs and restriction enzymes: arc11D-CAPS.F/arc11D-CAPS.R (SphI), arc12D-CAPS.F/arc12D-CAPS.R (NcoI), and arc3D-CAPS.F2/arc3D-CAPS.R6 (EcoRV). PCR genotyping of msl mutant alleles was performed using the following oligo pairs: JL-202/2.3’ (msl2-1), JL-202/3.3’ (msl3-1), and LB-GABI/msl2-R4 (msl2-3). Refer to Supplemental Table 1 online for oligo sequences.

Isolation and Immunofluorescence Staining of Chloroplasts

Chloroplast isolation and immunofluorescence staining was performed as described by Strawn et al. (2007) with the following modifications. Leaf fragments were digested in protoplast buffer (400 mM sorbitol, 20 mM MES/KOH, pH 5.2, and 0.5 mM CaCl₂) containing cellulase (2.5 mg mL⁻¹) and pectinase (0.8 mg mL⁻¹). Samples were blocked for 20 min in PBS, 5% BSA. The anti-FtsZ antibody (Levin and Losick, 1996) was used at a dilution of 1:5000. Slides were blocked for 20 min in PBS, 5% BSA, followed by a 1-h incubation with FITC-conjugated anti-rabbit secondary antibody (1:160; Sigma-Aldrich). Slides were mounted using the Slow-Fade Antifade kit (Invitrogen).

Light Microscopy of Chloroplasts

Tissue collection, fixation, and sample preparation were performed as described previously (Haswell and Meyerowitz, 2006). Images were...
captured with an Olympus DP71 microscope digital camera and pro-
cessed with DP-BSW software.

Confocal Microscopy
All confocal laser scanning microscopy was performed using a FLUO-
VIEW FV1000 (Olympus), and images were captured with FVIO-ASW
software (Olympus). For imaging of FtsZ-GFP–expressing lines (Fujiiwara
et al., 2008), FITC and GFP signals were excited at 488 nm and emissions
collected with a 505- to 606-nm band-pass filter. Chlorophyll autofluo-
rescence was excited at 635 nm and emissions collected with a 655- to
755-nm band-pass filter. For bacterial images, excitation wavelengths of
488 and 405 nm were used, and emissions were collected using 505- to
605-nm (FITC) and 430- to 470-nm (DAPI) band-pass filters.

Chloroplast Counting
The second true leaf was removed from 4-week-old soil-grown plants,
placed flat on Murashige and Skoog solid medium, and exposed to direct
light for 4 h, with an average intensity of 100 μmol m⁻² s⁻¹, to induce
chloroplast movement. Tissue removed from the tip of leaves was
mounted in water, and confocal scans were taken of mesophyll cells at
their largest circumference. After imaging, the number of chloroplasts per
optical section were counted using ImageJ version 1.44 software (http://
imagej.nih.gov/ij/).

Bacterial Growth Conditions and Immunofluorescence
Frag-1 and MUF465 strains (Levin et al., 1999) or PB103 and PB114 (de
Boer et al., 1989) were grown in Luria-Bertani medium to an OD600 of 0.4
and then diluted to an OD600 of 0.1 and treated with or without 10 μg mL⁻¹
cephalexin. After 1 to 2 h of growth, cells were fixed in 16% paraformal-
dehyde/4% glutaraldehyde for 30 min at room temperature followed by 1 h
on ice. Cells were washed and resuspended in GTE (50 mM Glc, 25 mM
Tris, and 10 mM EDTA) to an OD600 of 0.2. Immunofluorescence was
performed as described (Levin, 2002). Slides were incubated overnight at
4°C with rabbit anti-FtsZ antibody (1:4000; Mercer and Weiss, 2002),
followed by a conjugated anti-rabbit FITC antibody (1:200; Sigma-Aldrich).
Slides were treated with 1 μg mL⁻¹ 4',6-diamidino-2-phenylindole and
mounted with the SlowFade Antifade kit. Distance between FtsZ rings was
measured in pixels using ImageJ.

Quantitative Immunoblotting
One hundred milligrams of plant tissue was ground in liquid N2, and 200
μL of extraction buffer (150 mM NaCl, 100 mM Tris, pH 7.5, 0.1% Triton X-
100, 10% glycerol, and protease inhibitors) was added. After spinning at
4°C for 4 min, equal volumes of extract and 2× SDS sample buffer were
mixed and boiled for 5 min. Protein sample dilutions were separated on a
10% SDS-polyacrylamide gel and electroblotted to polyvinylidene fluo-
ride membrane. Membranes were blocked for 1 h with 5% nonfat milk in
TBST and probed with mouse anti-GFP (Clontech; 1:5000) or anti-FtsZ
antibody (1:5000) (Levin and Losick, 1996) followed by anti-mouse IgG-
HRP (Sigma-Aldrich; 1:5000). Chemiluminescent detection was per-
formed with the Thermo-Scientific SuperSignal West Femto detection
kit (Pierce). FtsZ1-GFP protein levels were quantified as described by
Schmitz et al. (2009) with the following modifications. The calculated
density value of msl2-3 msl3-1 bands was divided by the density value of
Col-0 bands to determine the FtsZ1-GFP protein levels present in msl2-3
msl3-1 plants relative to FtsZ1-GFP protein levels in Col-0 plants. Bands
imaged from lanes containing comparable total protein levels of Col-0
and msl2-3 msl3-1, as determined by Ponceau S staining of ribulose-
1,5-bisphosphate carboxylase/oxygenase, indicate similar total protein
concentrations at each dilution.

Quantitative RT-PCR
Primer mixes designed to amplify FtsZ1 (FtsZ1.QPCR.F and FtsZ1.
QPCR.R), FtsZ2-1 (FtsZ2-1.QPCR.F and FtsZ2-1.QPCR.R), FtsZ2-2
(FtsZ2-2.QPCR.F and FtsZ2-2.QPCR.R), MinD (MinD.QPCR.F and
MinD.QPCR.R), ARC3 (ARC3.QPCR.F and ARC3.QPCR.R) MCD1
(MCD1.QPCR.F and MCD1.QPCR.R), and ACTIN2/7/8 (ACTF-QPCR/
ACT17.R-QPCR and Actin8R-QPCR) were used to determine the
FtsZ1-GFP protein levels present in Col-0 plants. Bands were measured in
pixels using ImageJ.

Chloroplast Division Are Not Altered in
msl2 msl3 arc3 Mutants.
We tested for protein–protein interactions between MSL3 and compo-
ents of the plastid division machinery using the ProQuest two-hybrid
system (Invitrogen). cDNA sequence encoding the soluble C terminus of
MSL3 (amino acids 286 to 678) was cloned into the pDEST 32 bait vector,
while full-length cDNAs for MinD, MinE, FtsZ1, and cDNAs encoding the
stromal portions of ARC3 (amino acids 41 to 741) and PARC6 (amino acids
77 to 352) were cloned into the pDEST 22 prey vector. The assays were performed according to the manufacturer’s
instructions.

Accession Numbers
Sequence data from this article can be found in the Arabidopsis Genome
Initiative or GenBank/EMBL databases under the following accession
numbers: MSL2, At5g10490; MSL3, At1g58200; MinD, At1g24020; MinE,
At1g69390; ARC3, At1g75010; FtsZ1, At5g55280; FtsZ2-1, At2g36250;
FtsZ2-2, At3g52750; and MCD1, At1g20830.

Supplemental Data
The following materials are available in the online version of this article.

Supplemental Figure 1. Characterization of the msl2-3 Allele.
Supplemental Figure 2. An Antibody Raised against Bacillus subtilis
FtsZ Recognizes Arabidopsis thaliana FtsZ1.
Supplemental Figure 3. Representative Images of Confocal Scans
Used to Determine Chloroplast Number in Single, Double, and Triple
msl and arc Mutants.
Supplemental Figure 4. msl2 msl3 arc3 Triple Mutants Are Pheno-
typically Indistinguishable from Single arc3 Mutants.
Supplemental Figure 5. Z-Ring Placement in the ΔminB Strain of
E. coli.
Supplemental Figure 6. Transcript Levels of Genes Involved in
Chloroplast Division Are Not Altered in msl2 msl3 Mutants.
Supplemental Table 1. Oligos Used in Genotyping and Quantitative
RT-PCR.

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AUTHOR CONTRIBUTIONS

M.E.W. G.S.J. and E.S.H performed research and contributed new tools. M.E.W. and E.S.H designed research, analyzed data, and wrote the manuscript.

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