

ASSEMBLY DYNAMICS OF THE BACTERIAL CELL DIVISION PROTEIN FtsZ: Poised at the Edge of Stability

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■ **Abstract** FtsZ is a prokaryotic tubulin homolog that assembles into a ring at the future site of cell division. The resulting “Z ring” forms the framework for the division apparatus, and its assembly is regulated throughout the bacterial cell cycle. A highly dynamic structure, the Z ring exhibits continual subunit turnover and the ability to rapidly assemble, disassemble, and, under certain circumstances, relocalize. These *in vivo* properties are ultimately due to FtsZ’s capacity for guanosine triphosphate (GTP)-dependent, reversible polymerization. FtsZ polymer stability appears to be fine-tuned such that subtle changes in its assembly kinetics result in large changes in the Z ring structure. Thus, regulatory proteins that modulate FtsZ’s assembly dynamics can cause the ring to rapidly remodel in response to developmental and environmental cues.

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INTRODUCTION

Seemingly static cellular structures can in reality be remarkably dynamic, undergoing continual regeneration of constituent parts. Subunit turnover in the eukaryotic cytoskeleton allows cells to rapidly reorganize in response to environmental and developmental signals. Populations of microtubules and actin filaments are maintained through continual and carefully balanced assembly and disassembly (24, 107). Bacteria have cytoskeletal elements that are related to actin and tubulin (137), and accumulating evidence shows that the prokaryotic cytoskeleton is also a dynamic system in which multiple inputs allow for precisely tuned and rapid responses to a changing environment.

The dynamic assembly of FtsZ, the prokaryotic homolog of tubulin, is fundamental to the regulation of bacterial cell division. In this review we focus on the molecular basis of FtsZ assembly and its modulation during cell growth and development. FtsZ is conserved throughout bacteria and archaea; it also exists in chloroplasts and in the mitochondria of certain eukaryotes (37, 82, 101, 120). In bacteria, FtsZ forms a macromolecular structure called the Z ring that provides the framework for the division apparatus and determines the site of cytokinesis. *In vitro*, the stability of FtsZ polymers, like that of microtubules, is precisely tuned. FtsZ polymerizes in the presence of GTP but tends to depolymerize again once the bound GTP has been hydrolyzed. *In vivo*, FtsZ assembly is also transient. Subunits in the Z ring are continually exchanged with those in the cytoplasm. The constant balance between assembly and disassembly means that small changes in FtsZ polymerization dynamics are able to trigger large and rapid changes in the location or morphology of FtsZ structures in the cell. Proteins that induce such changes dictate the timing and location of cytokinesis.

In this review we discuss (a) Z ring assembly and dynamics *in vivo*, (b) FtsZ polymer assembly *in vitro*, and (c) the counterbalancing effects of proteins known to regulate FtsZ assembly. Available data suggest that the mechanisms governing Z ring formation are fundamentally conserved. We have highlighted any differences in FtsZ assembly between species where they exist.

PART I: Z RING DYNAMICS THROUGHOUT THE CELL CYCLE

During different stages of the cell cycle FtsZ exhibits a series of different behaviors: assembly to form the Z ring at midcell, maintenance of the ring through continual subunit turnover, and constriction and disassembly of the ring. The transitions between Z ring growth, maintenance, and constriction are likely to be regulated by

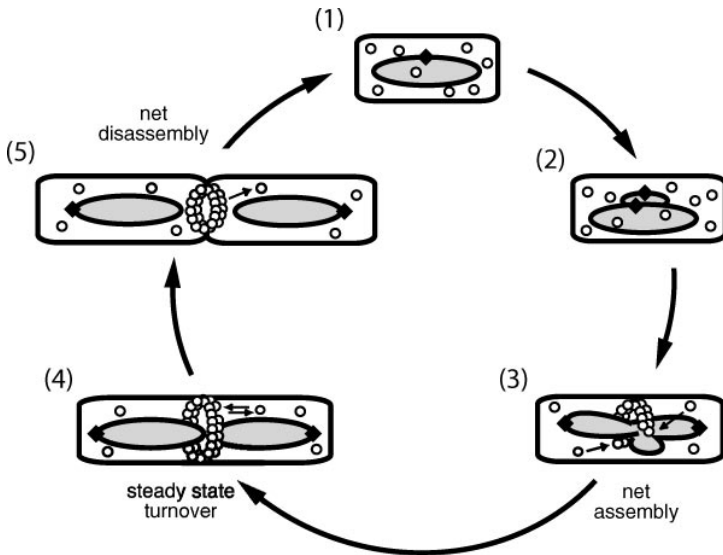


Figure 1 The bacterial division cycle. FtsZ assembly is coordinated with DNA replication and segregation. Step 1: In newborn cells FtsZ (O) is unassembled. A circular chromosome with a single origin of replication (♦) is located at midcell. Steps 2–4: After chromosome replication initiates, the polymerase machinery remains at midcell, but origins of replication separate and move to opposite poles of the cell. Once replication is complete, the condensed chromosomes separate, leaving a nucleoid-free space. Step 3: Z ring formation coincides with chromosome segregation. Assembly starts from a single point at midcell and extends bidirectionally. Step 5: During cytokinesis the Z ring constricts at the leading edge of the invaginating septum. Small arrows indicate the assembly and disassembly of FtsZ subunits.

factors that influence FtsZ's assembly dynamics. However, the precise nature of the signals driving these transitions remains unclear. In eukaryotes DNA replication, chromosome segregation, and cell division occur in strict succession, whereas in rapidly growing prokaryotes these processes often overlap (26). Nonetheless, in all organisms cytokinesis must be tightly coupled to the cell cycle so that each daughter cell receives a complete copy of the genome.

A strong correlation exists between Z ring behavior and cell cycle progression (Figure 1). Z ring formation begins after the initiation of DNA replication and the separation of replication origins, and it roughly coincides with nucleoid segregation (23, 47, 67, 110, 117). Data suggest that FtsZ assembly may start at a single point on the inner surface of the cell membrane and then rapidly extend bidirectionally to form an arc and then a closed circle (3). This ring serves as the framework for assembly of other cell division proteins (82, 120). Although the ring maintains its shape and size for a large portion of the cell cycle (23, 146), individual FtsZ

molecules in the ring are rapidly exchanged with cytoplasmic FtsZ (130). When an unknown signal triggers cytokinesis, new cell wall is deposited and the Z ring contracts along with the cytoplasmic membrane (1, 10, 62). By the end of the septation process the Z ring has disassembled (130). Below we discuss the dynamics of Z ring assembly and its coordination with events in the cell cycle.

Factors Affecting Z Ring Initiation and Localization

Z ring formation establishes the location of the future division site and is an integral part of the temporal regulation of cytokinesis. The initiation of FtsZ assembly must be tightly controlled both temporally and spatially to prevent aberrant septation. Z ring formation depends on at least three factors: FtsZ concentration, the initiation of DNA replication, and the presence or absence of the nucleoid mass.

A MINIMUM FtsZ CONCENTRATION IS NECESSARY BUT NOT SUFFICIENT FOR Z RING FORMATION FtsZ assembles *in vitro* in a concentration-dependent manner (see Part II: Biochemical Properties of FtsZ Protein, below), which suggests that FtsZ levels in a cell might control the timing of Z ring formation. In this model FtsZ concentration in newborn cells would be too low to support assembly but would gradually increase during the course of the cell cycle and reach a critical concentration immediately prior to Z ring formation. A minimum FtsZ concentration is in fact necessary for division in *Escherichia coli* and for Z ring assembly in *Bacillus subtilis* (17, 61), but under most circumstances fluctuations in FtsZ concentration are not sufficient to explain the timing of ring formation.

Both *E. coli* and *B. subtilis* can modulate transcription of the *ftsZ* operon (4, 35, 55, 140), but FtsZ protein levels remain constant throughout the cell cycle of both organisms during steady state growth (146). Altering FtsZ levels is not sufficient to change the timing of medial ring formation. Lowering *ftsZ* expression below wild-type levels does not change the timing of cytokinesis in *E. coli*, though cell size increases slightly (102). Increasing FtsZ levels twofold in *B. subtilis* does not alter the frequency or timing of medial Z ring formation (146), though it does lead to an increase in the formation of polar Z rings. Similarly, two- to sevenfold increases in FtsZ levels in *E. coli* lead to an increase in the frequency of polar septation (145). These data suggest that although high FtsZ expression levels do not change medial Z ring formation, they can overcome a division inhibitor that normally prevents assembly at the cell poles. (In *E. coli* and *B. subtilis* this inhibitor is most likely the MinCD complex; see Part III: Proteins that Modulate FtsZ Assembly.) In support of this model, during sporulation in *B. subtilis*, increased FtsZ expression is required to initiate Z ring formation at previously suppressed locations adjacent to the cell poles (7, 36, 38).

Unlike *B. subtilis* and *E. coli*, *Caulobacter crescentus* modulates FtsZ levels in a cell cycle-dependent manner. In this organism transcriptional regulation and proteolysis ensure that FtsZ is undetectable in swarmer cells, which do not initiate DNA replication or division (57, 111), but is highly expressed prior to cytokinesis

in the reproductively competent stalk cells (57, 109, 121). Following cytokinesis FtsZ is cleared from stalk cells (109). Nonetheless, ectopic expression of *ftsZ* in swarmer cells is not sufficient to drive medial Z ring formation (111). Ectopic FtsZ expression can only stimulate ring formation after cell differentiation and the subsequent initiation of DNA replication (110). Thus, it is ultimately cell cycle regulation that dictates the initiation of Z ring formation even when elaborate regulation of FtsZ levels exists.

REPLICATION INITIATION IS NECESSARY FOR POSITIONING THE Z RING AT MIDCELL Z ring formation initiates after the onset of DNA synthesis and the separation of newly replicated origin DNA, which suggests that assembly might be triggered by events at the start of chromosome replication (23, 47, 110, 117). However, blocking the initiation of DNA replication does not inhibit Z ring formation in either *E. coli* or *B. subtilis* (47, 131). The terminal phenotype of *B. subtilis* DNA replication initiation mutants is a cell in which the Z ring is located off-center, adjacent to the medially positioned, unreplicated nucleoid (47). Similarly, a Z ring forms adjacent to the nucleoid in *E. coli* cells in which either DNA replication or nucleoid segregation are defective (131, 132).

While DNA replication is not essential for FtsZ assembly in *E. coli* and *B. subtilis*, correct positioning of the Z ring at midcell does require replication initiation, but not elongation of the replication fork. Permitting DNA replication in germinating *B. subtilis* spores, but preventing elongation with a thymine auxotroph, leads to degradation of the origin region and the correct positioning of a medial Z ring over the unreplicated nucleoid (47, 117). Because the FtsZ ring forms proximal to the previous location of the origin of replication, and because replication initiation is required for proper ring localization, it is tempting to speculate that the same machinery that recruits the origin of replication and the polymerase machinery to midcell also helps to unmask a preferred FtsZ nucleation site at midcell. In support of this idea, the polar placement of the division septum during sporulation in *B. subtilis* is partially dependent on the anchoring of the origin region at the cell pole (41).

LOCAL INHIBITION OF Z RING FORMATION BY THE UNSEGREGATED NUCLEOID The correlation between Z ring formation and the separation of the bacterial nucleoid has led to the long-standing hypothesis that the bacterial nucleoid is a determinant in the timing and placement of the division septum [the nucleoid occlusion model (97, 149)]. According to this model, cell division can take place at any point along the length of a cell, but early in the cell cycle the presence of the unsegregated nucleoid prevents septation at midcell. Later, DNA segregation reveals a small, nucleoid-free zone at midcell that is able to support assembly of the division apparatus. Z ring formation in the nucleoid-free space adjacent to the cell poles is prevented by the presence of the MinCD division inhibitors (see Part III: Proteins that Modulate FtsZ Assembly, below) (155).

The nucleoid appears to play a role in preventing aberrant Z ring formation, but in both *E. coli* and *B. subtilis* its presence alone is not sufficient to dictate the

precise spatiotemporal regulation of FtsZ assembly. Z ring formation is clearly influenced by the location of the nucleoid under some circumstances. When DNA replication or partitioning has been blocked, the chromosome remains at midcell, and nonmedial Z rings form adjacent to the nucleoid (47, 131, 132). In addition, when chromosome partitioning is blocked in the absence of the Min proteins, the Z ring in *E. coli* is free to form at any position along the nucleoid-free space (155). Under certain circumstances, however, the position of the nucleoid coincides with that of the Z ring and of cytokinesis. During sporulation in *B. subtilis* the nucleoid is bisected by the asymmetrically positioned septum and the DNA must subsequently be pumped across the septum into the forespore (128, 150). Moreover, Z rings can form directly over the nucleoid in cells with mutations in chromosome condensation genes (*smc* in *B. subtilis* and *mukB* in *E. coli*) (14, 40, 156). One potential explanation is that Z ring formation and septation can occur in regions of the nucleoid that are relatively less dense. If this is indeed the case, this would explain why medial FtsZ ring formation occurs only after degradation of the DNA adjacent to the origin of replication in germinating *B. subtilis* spores blocked for DNA replication (117).

Maintenance of the Z Ring as a Dynamic Framework

Once formed, the Z ring is present for a significant portion of the cell cycle [e.g., 85% of the mass doubling period in *B. subtilis* cells growing in Luria broth medium] (23, 146). In *B. subtilis* the proportion of cells with Z rings increases with increasing growth rate, indicating that the Z period (the time the Z ring is present in a given cell) remains relatively constant even when doubling time decreases (67, 146). The prolonged duration of the Z period suggests there may be a minimum time required for proper assembly of the entire division apparatus.

During the Z period, although no gross changes in the Z ring can be detected by immunofluorescence, the ring is extremely dynamic, with subunit turnover on par with that of the eukaryotic cytoskeleton (103). For example, the Z ring can disappear and reassemble within minutes after shifting cells expressing a heat-sensitive *ftsZ* allele (*ftsZ84*) into and out of the nonpermissive temperature (2). More recently, the halftime for replacement of FtsZ subunits in the *E. coli* cytokinetic ring has been found to be between 8 and 30 sec, as indicated by fluorescence recovery after photobleaching (FRAP) experiments on cells expressing FtsZ-GFP (green fluorescent protein) [(130); H.P. Erickson, personal communication]. Similar fluorescence recovery rates are observed for GFP fused to ZipA, a protein associated with the Z ring in *E. coli* (130), which hints that the entire division machinery might also be continually turning over. The fluorescence intensity of the ring before and after FRAP shows that only ~30% of the FtsZ in a cell is in the ring at any given time, suggesting that a large pool of soluble FtsZ is available for exchange. In eukaryotic cells the detection of large pools of unassembled actin and tubulin prompted successful searches for proteins that regulate polymerization.

This rapid subunit turnover requires energy but does not appear to be critical for normal cytokinesis. Turnover rates of FtsZ in the ring appear to correspond to FtsZ's *in vitro* ability to hydrolyze GTP, but lowering the hydrolysis rate by an order of magnitude is not lethal to the cell. For example, FtsZ84 protein hydrolyzes GTP *in vitro* at rates 10 times slower than wild-type FtsZ (20, 115), and FRAP experiments conducted at the permissive temperature indicate that recovery rates for FtsZ in *ftsZ84* cells are also approximately 10 times slower than in wild type. Nonetheless, cell division still occurs with no apparent defect at these permissive temperatures. Many other FtsZ mutations also dramatically lower GTP hydrolysis rates without preventing cytokinesis (130). Thus, cells appear able to tolerate considerable variation in the subunit turnover rates of the Z ring.

Remodeling the Z Ring

If the rapid turnover of FtsZ subunits within the Z ring is not essential for cytokinesis under normal growth conditions, why does it occur? One function may be to allow cells to disassemble or remodel the division apparatus in response to DNA damage, starvation, or other environmental stresses. We review two examples below.

DNA damage induces the SOS response in bacteria, during which cell division is prevented until the DNA has been repaired (68). In *E. coli*, DNA damage results in the expression of Sula, which causes the Z ring to be dismantled (11, 39). Once the DNA lesions are repaired, Sula levels rapidly decline (88, 89), permitting FtsZ to once again assemble and division to resume.

Z ring remodeling also plays an important role during development in *B. subtilis*. In response to cell crowding and nutritional deprivation, *B. subtilis* produces environmentally resistant endospores [for reviews, see (15)]. A key step in sporulation is the switch from binary fission to polar septation (32, 63). Polar septation is preceded by the relocalization of FtsZ from midcell to positions adjacent to both cell poles (62). The shift in Z localization involves the transformation of the medial ring into a spiral that extends the length of the cell and then coalesces into rings at both cell poles (7). [The choice of which ring to use is largely stochastic (127).] Formation of the spiral depends both on an increase in FtsZ expression levels and on the expression of SpoIIIE, a protein that colocalizes with FtsZ during sporulation and may help stabilize the polymer (5, 58, 64, 77, 153). Intriguingly, if sporulating cells that have not yet divided are given fresh nutrients, FtsZ can relocalize again from the cell poles to midcell (7), indicating that the medial site is still available.

Z Ring Constriction

When a cell divides, the Z ring constricts at the leading edge of the invaginating septum (10). There is limited evidence as to FtsZ's precise function during this process. At minimum, FtsZ is needed to maintain the medial localization of the other division proteins. To do this, the FtsZ framework must be able to retain its

coherence despite an ever-shrinking structure. It has also been suggested that FtsZ might generate the force that directs cytokinesis, similar to the actomyosin ring in eukaryotes (31).

Cytokinesis occurs in two discrete stages: the formation of an initial, small invagination at the site of the septum, followed by the completion of cell division. FtsZ is necessary throughout both stages. If at any time during cytokinesis conditional alleles of *ftsZ* are shifted to nonpermissive temperatures, the Z ring disappears, all other proteins in the division apparatus delocalize, and the septum ceases to invaginate (2, 134). If invagination has not yet begun, cells are left with smooth walls, indicating that septum formation cannot begin without FtsZ. In contrast, when conditional alleles in later-acting genes such as *ftsI* (which encodes PBP3) are used to block division, cytokinesis ceases only after the initial invagination step (134). Qualitative changes appear to occur after constriction has begun. When heat-sensitive *ftsZ84* cells are raised to the nonpermissive temperature, Z rings rapidly disassemble. After returning to the permissive temperature, FtsZ can reoccupy some sites but not those where invagination of the cell wall is visible (2).

The molecular mechanism of Z ring constriction during septum formation remains unclear. One model suggests that as the diameter of the ring grows smaller, FtsZ subunits are released from the ring. Alternatively, filaments might be sliding against each other without disassembling, as occurs in the actomyosin ring in eukaryotes (13). Regardless of the mechanism of constriction, once cytokinesis is complete, the vast majority of FtsZ in *E. coli* and *B. subtilis* appears to be cytoplasmic, although FtsZ foci are occasionally observed at cell poles (62). These polar foci of FtsZ presumably reflect the remaining presence of potential FtsZ binding sites at these locations.

PART II: BIOCHEMICAL PROPERTIES OF FtsZ PROTEIN

Z ring initiation, maintenance, and constriction are regulated at least in part by the modulation of FtsZ assembly dynamics. The investigation of tubulin polymerization *in vitro* has provided invaluable insights into microtubule assembly dynamics and function *in vivo*. Similarly, understanding FtsZ behavior *in vivo* will require the detailed analysis of FtsZ polymer assembly and structure *in vitro*. Because FtsZ dynamics have been investigated and interpreted in the context of the eukaryotic cytoskeleton, we first review the mechanisms behind microtubule and actin assembly and disassembly. We next describe the structure of FtsZ subunits and polymers. Finally, we discuss the dynamic behavior that FtsZ exhibits *in vitro* and the possible molecular mechanism behind those dynamics.

Eukaryotic Cytoskeletal Polymers: Paradigms for Nucleotide-Dependent, Reversible Assembly

In eukaryotic cells, microtubules continually grow and shrink, probing the cytoplasm to locate disperse targets and to build and maintain the mitotic spindle.

Tubulin's GTP hydrolysis provides the energy for microtubule polymer dynamics; it is the continual input of energy that ultimately allows the cell to rapidly rearrange its cytoskeleton [reviewed in (24)].

The reversibility of microtubule polymerization is built into the tubulin subunits' inherent assembly and hydrolysis properties. Tubulin is a stable heterodimer consisting of highly homologous α and β subunits, both of which bind GTP (Figure 2*b*). The GTP on α -tubulin is sandwiched between the subunits of the dimer and does not hydrolyze or exchange with nucleotide in solution. The nucleotide bound to the β subunit can both exchange and hydrolyze. For the remainder of the paper, when referring to tubulin, only the β -tubulin nucleotide binding site is discussed. When β -tubulin is bound to GTP, the protein assembles into a microtubule comprised of 13 straight protofilaments. After assembly the GTP is hydrolyzed and the resulting guanosine diphosphate (GDP)-containing microtubule is destabilized. An analogous situation occurs for actin polymerization with adenosine triphosphate (ATP) [reviewed in (108)].

The coupling of nucleotide hydrolysis to polymer assembly also allows for complicated, nonequilibrium behaviors such as dynamic instability and treadmilling. The term dynamic instability is often misused and does not mean that polymers containing GTP are more stable than those containing GDP. Instead, it refers to the fact that at steady state, individual microtubules undergo stochastic and reversible transitions between growth and rapid shrinkage [reviewed in (24)]. This occurs because microtubules are only metastable structures: The GDP-bound protofilaments would prefer to curl outwards, but they are held together by the lateral interaction in the microtubule wall. When this balance of forces is disturbed, protofilaments separate and rapidly peel away.

Treadmilling is another nonequilibrium phenomenon. One end of a polymer steadily grows while the other end of the same polymer steadily shrinks. Treadmilling occurs in microtubules, but it has been best characterized in actin [reviewed in (108)]. In actin filaments, an asymmetric distribution of ATP, adenosine diphosphate (ADP)-Pi, and ADP-bound subunits develops. At steady state, ATP subunits associate at one end of the filament while ADP subunits dissociate from the other.

Currently there is no evidence whether FtsZ polymers exhibit either dynamic instability or treadmilling. Nonetheless, FtsZ and tubulin share many properties. Like tubulin, FtsZ polymerizes in the presence of GTP to form straight polymers that can curl and become more labile after hydrolysis to GDP. Much is still being learned about FtsZ polymerization. How does GTP hydrolysis affect polymer structure and dynamics? What higher-order structures does FtsZ form, and how do their dynamics differ from those of single-stranded protofilaments? How is a new polymer initiated? Do polymers assemble and disassemble as a population, or do they undergo dynamic instability or treadmilling? The most common techniques used to study these FtsZ assembly dynamics *in vitro*, along with the limitations of each, are described in Table 1. Unless otherwise noted, FtsZ proteins from all species behave similarly when tested.

TABLE 1 Assays for FtsZ's biochemical activities

Method	Purpose	Advantages	Disadvantages
Blot overlays or UV cross-linking ^a	Detecting bound nucleotide	Easy	<ul style="list-style-type: none"> • Not quantitative • Not practical for kinetic assays
Isolation of FtsZ and bound nucleotide by sedimentation or filter binding ^b	Detecting bound nucleotide	Quantitative	<ul style="list-style-type: none"> • Poor time resolution • May not distinguish between different hydrolysis intermediates (GTP, GDP-Pi, GDP) • Nucleotide may be hydrolyzed by FtsZ during centrifugation or filter handling
Conventional sedimentation ^c	Quantifying assembly	Easy	<ul style="list-style-type: none"> • Cannot distinguish increases in polymer mass from changes in polymer structure • Calculation of absolute polymer mass difficult due to heterogeneous length and width of FtsZ polymers • Bundling agents that enhance sedimentation artificially stabilize polymers^d
Analytical ultracentrifugation ^e	Quantifying assembly	Quantitative	<ul style="list-style-type: none"> • Not practical for kinetic assays • GTP can be depleted if magnesium is included in reaction
90° light scattering ^c	Quantifying assembly	Detection in real time	<ul style="list-style-type: none"> • Cannot distinguish increases in polymer mass from changes in polymer structure • Calculation of absolute polymer mass difficult due to heterogeneous length and width of FtsZ polymers
Electron microscopy ^f	Visualizing polymers	High resolution	<ul style="list-style-type: none"> • Cannot be used for real time observations • Polymer species may not be represented in proportion to their concentration in solution
Fluorescence microscopy ^g	Visualizing polymers	Visualization in real time	<ul style="list-style-type: none"> • Cannot resolve individual protofilaments or the lengths of small polymers

Assays were initially performed in ^a(20, 93, 115), ^b(87, 93, 115, 125, 126), ^c(13), ^d(94, 95, 123, 154), ^e(118, 129), ^f(13, 31, 94), and ^g(154).

FtsZ Structures

To understand the dynamic behavior of FtsZ, it is first necessary to review the data on its structure. Below we describe the structure of (a) the FtsZ subunit, (b) the GTP binding site, (c) the protofilaments assembled in GTP and GDP, and (d) the higher-order FtsZ structures. These static pictures of FtsZ subunits and polymers

will form the basis for discussing dynamic FtsZ polymer populations in the next section.

THE SUBUNIT Despite sharing minimal sequence homology (~10%) (94), FtsZ and tubulin have strikingly similar protein folds and GTP binding interactions (98) (Figure 2). FtsZ is a ~40-kDa polypeptide whose atomic structure was determined using FtsZ from the thermophilic archaeon *Methanococcus jannaschii* (70). The structure contains a major β sheet surrounded by α helices; perpendicular to this is a smaller subdomain, which also consists of a short β sheet surrounded by α helices (Figure 2a). The FtsZ and tubulin structures fall into a subclass different from that of small G proteins, dynamin, and all other GTP binding proteins possessing a P-loop (59, 98). Instead, they appear most closely related to that of the NADH binding protein, glyceraldehyde-3-phosphate dehydrogenase (98).

Not visible in the atomic structure is a flexible, highly charged C-terminal region. It is extremely short in *M. jannaschii* but is longer and essential for cytokinesis in *E. coli* and *C. crescentus* (25, 80, 81, 143). A ~15-amino-acid conserved sequence near the very C terminus is necessary for binding ZipA and FtsA, two essential Z ring proteins in *E. coli* (46, 72, 81, 90, 139, 143) and for binding FtsA in *C. crescentus* (25) and *S. aureus* (151). This sequence is also present in species that lack ZipA and FtsA (30), but otherwise, there is little conservation between the C termini of divergent organisms (6, 83). Tubulin's C terminus, like that of FtsZ, is unstructured, charged, and the binding site for many microtubule-associated proteins, but the sequences are unrelated.

THE GTP BINDING SITE The GTP binding residues are the amino acids that are best conserved between FtsZ and tubulin (70, 98) (Figure 2b), reflecting similarities in the way the two proteins interact with nucleotide during polymer assembly and disassembly. The nucleotide binds on the surface that will form the interaction site between subunits in a protofilament. In analogy with the microtubule structure, we call the protein bonds within an FtsZ protofilament longitudinal bonds and the GTP binding surface of the monomer the plus end of the protein. Unlike tubulin, unassembled FtsZ exists as a monomer rather than as a heterodimer, and all subunits can exchange nucleotide (93, 129). Once a longitudinal bond has formed, the nucleotide is likely to be almost completely buried in the interface between FtsZ subunits (98). Conserved sequences at both the plus and minus ends of a subunit are essential to FtsZ's ability to bind and hydrolyze GTP.

Monomeric FtsZ uses conserved residues at the plus end of the protein to bind GTP tightly [$\sim 5 \mu\text{M}$ K_d (93)]. Seven of these amino acids constitute the tubulin signature motif, a sequence conserved in all tubulins and originally used to identify FtsZ as a distant tubulin homolog (20, 93). *ftsZ84*, a mutation in a conserved glycine in this motif (G105S in *E. coli*) (Figure 2a), is the best-characterized mutation at the plus end of FtsZ. This mutation disrupts nucleotide binding, thereby decreasing both polymer assembly and GTP hydrolysis in vitro (13, 20, 115).

ftsZ84 renders *E. coli* cells heat-sensitive for division (79), but the FtsZ84 protein does not exhibit temperature-sensitive defects in vitro (13, 20, 116). Thus it is unclear how the defects in nucleotide binding and hydrolysis relate to FtsZ84 function in vivo.

Residues at the minus end of the subunit participate in both protofilament bond formation and GTP hydrolysis (29). Thus, FtsZ assembly is necessary for GTP hydrolysis (20, 129, 144). Several of the minus end residues are highly conserved in FtsZ and tubulin (98) (Figure 2). Mutations in many of these residues (Figure 2a) do not eliminate the in vitro ability of FtsZ to bind GTP or assemble, but they do reduce the protein's nucleotide hydrolysis rate (19, 75, 78, 123, 124, 136, 143).

PROTOFILAMENTS The conserved GTP binding residues reflect similarities in the structures that FtsZ and tubulin form with GTP and GDP. When GTP is present, protofilaments form (94) in which subunits are stacked linearly without any net twist or curvature (31, 71) (Figure 3a). FtsZ protofilaments can exist in isolation,

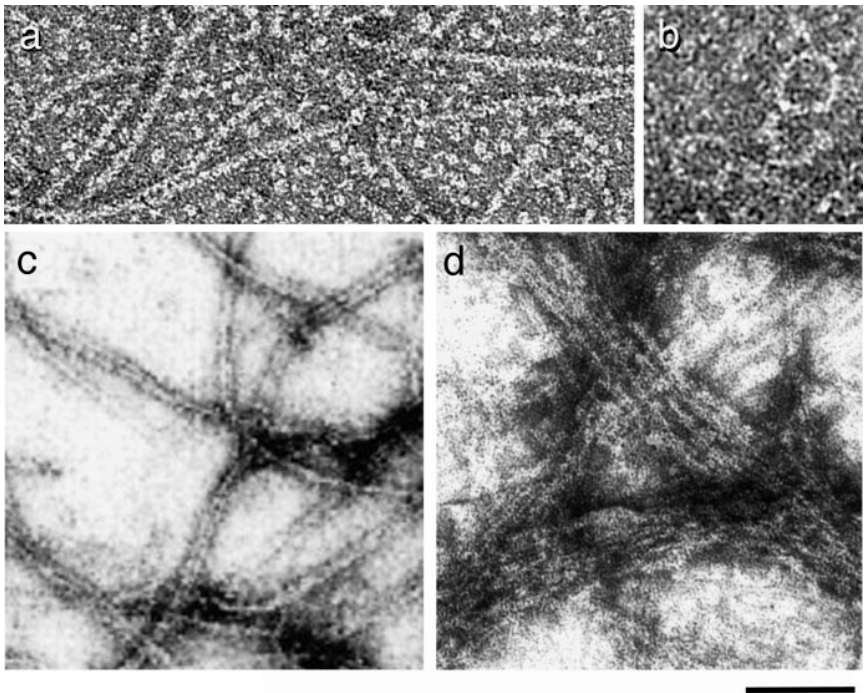


Figure 3 FtsZ polymers. (a) FtsZ protofilaments assembled with GTP. Monomeric FtsZ is also visible in the background (micrograph by L. Romberg). (b) FtsZ rings assembled with GDP (119). (c) Protofilament pairs and bundles assembled in reactions with GTP (96). (d) Large FtsZ bundles formed in the presence of ZipA and GTP (114). Bar is 50 nm in (a) and (b), 100 nm in (c) and (d). Images reprinted with permission.

whereas tubulin protofilaments do not form except in the context of a multi-stranded polymer. In the presence of GDP, the interaction between FtsZ subunits is substantially weakened (13, 95, 118, 129). In addition, GDP-bound protofilaments can curve sharply to form rings or spirals with a diameter of 15–25 nm (24, 31, 76, 114, 136) (Figure 3*b*). Because FtsZ protofilaments readily assemble under a wide variety of conditions and are structurally almost indistinguishable from those of tubulin, it is likely that FtsZ protofilaments are a genuine component of the Z ring *in vivo*.

The role of nucleotide hydrolysis in microtubule assembly has been studied using GMPCPP, a nonhydrolyzable analog of GTP. Equivalent studies of FtsZ have been hindered by the lack of GTP analogs that interact similarly with FtsZ. GMPCPP allows assembly of FtsZ polymers but is slowly hydrolyzed (76, 119). Chelating magnesium prevents GTP hydrolysis (20, 93, 115) and allows protofilament assembly (76, 96), but this assembly is not efficient at low protein concentrations (118, 129). GMPPNP does not bind FtsZ well (125), and GTP γ S, although it binds FtsZ, does not support assembly (13, 125, 154). Interestingly, GTP γ S has been reported to stabilize preformed FtsZ polymers (125), which suggests that subunits bound to this analog may cap the ends of filaments.

MULTISTRANDED POLYMERS The Z ring is thought to contain 6 to 7 protofilaments (130), but how these protofilaments are associated with each other is unknown. The filaments could interact directly with each other through their lateral surfaces or indirectly through cross-linking by other cell division proteins. The amino acids on the lateral surfaces of FtsZ and tubulin are not homologous; thus, microtubules do not provide a good model for the higher-order association of FtsZ protofilaments (98). For the moment the assembly of multistranded FtsZ structures *in vitro* provides the best information on the higher-order structures that might form *in vivo*.

Many different surfaces of the FtsZ protofilament can be used to form a wide variety of multistranded structures, including protofilament pairs and bundles (Figure 3*c,d*), sheets, spirals, hoops, and tubes (13, 31, 71, 74, 76, 95, 96, 143, 148). [FtsZ tubes are not analogous to microtubules: Protofilaments are paired and tightly spiraled (76), not straight as in the microtubule wall.] Lateral associations are affected by protein concentration, pH (31, 95), and the presence of multivalent cations including DEAE dextran (13, 76), calcium (76, 96, 154), and magnesium (96, 148). FtsZ's tendency to bundle also varies between species. For example, FtsZ from *Mycobacterium tuberculosis* appears to bundle more readily than that of *E. coli* (148).

Tubulin, in addition to forming microtubules, can also assemble *in vitro* into a wide array of other structures that are not observed *in vivo* (27). Thus, many of the lateral interactions described above for FtsZ may not be relevant *in vivo*. It will be important to determine what structures assemble in the context of the Z ring, with or without the help of other division proteins, because these structures will affect the pathway by which the Z ring assembles and turns over.

FtsZ Polymer Dynamics

During the cell cycle the Z ring goes through phases of net assembly, steady state turnover, and net disassembly. To understand these processes, we first need to understand the behavior of pure FtsZ polymers *in vitro*. At steady state, which nucleotide intermediates (GTP, GDP-Pi, or GDP) do the polymers contain, and how does this affect the assembly dynamics? Might FtsZ exhibit dynamic instability or treadmilling? Is the initial assembly of FtsZ polymers cooperative? Below we discuss what is known about FtsZ's *in vitro* steady state assembly dynamics and the initiation of FtsZ polymers.

THE NUCLEOTIDE CONTENT OF FtsZ POLYMERS AT STEADY STATE Turnover of the Z ring appears to be powered by FtsZ's GTP hydrolysis activity (130). Knowing the nucleotide intermediate that predominates in polymers at steady state may suggest how the energy from hydrolysis is used. For example, the net twist in an actin filament changes after phosphate release and alters the filament's affinity for proteins that sever or nucleate actin (107, 147). In addition, hydrolysis and phosphate release cause changes in the polymerization properties of actin and tubulin. These changes are fundamental to the mechanisms of dynamic instability and treadmilling and thus to the regulation of polymer turnover in eukaryotic cells.

For FtsZ each hydrolysis event requires many steps (GTP binding, hydrolysis, phosphate release, GDP release) (Figure 4), and the slowest step in the cycle

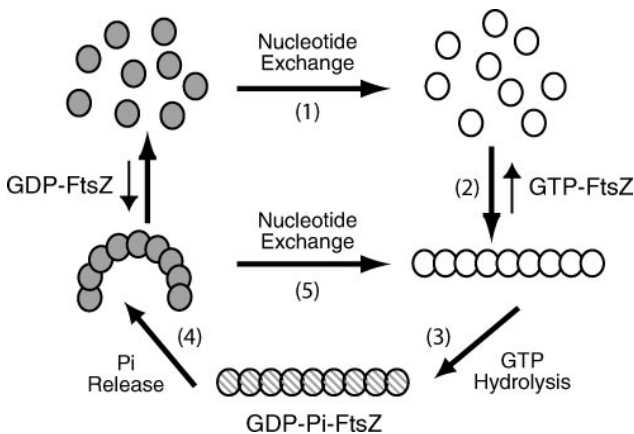


Figure 4 Model for FtsZ's GTP hydrolysis cycle. FtsZ's hydrolysis cycle is connected to polymer assembly and disassembly. 1: Monomers rapidly exchange nucleotide with solution. *In vivo*, more GTP than GDP exists in the cytoplasm. 2: When bound to GTP, FtsZ assembly is enhanced. 3: Once assembled, FtsZ can hydrolyze the GTP. 4: Phosphate release follows. GDP-containing polymers are more likely to be curved or to disassemble than GTP-containing polymers. 5: Nucleotide exchange might occur without complete disassembly of the polymer.

determines the intermediate that predominates. If hydrolysis is rapid and product release is slow, GDP or GDP-Pi will build up in the polymer. If product release is relatively fast, the polymers will contain GTP. Further complicating matters, during each hydrolysis cycle an individual subunit may go through a cycle of assembly and disassembly. Because the GTP is sandwiched between subunits in FtsZ polymers, the assembly state of a subunit may affect the rate at which the GDP and phosphate can be released into solution.

FtsZ polymers may not contain the same nucleotide as microtubules. Microtubules contain GDP-bound subunits because after tubulin assembly and GTP hydrolysis, the GDP is trapped between subunits in the microtubule wall. Since it is rare for longitudinal bonds in the middle of a microtubule to break, nucleotide exchange occurs only when subunits dissociate from the ends of a microtubule. Indirect evidence suggests that FtsZ polymers behave differently. FtsZ polymers have a much higher steady state GTP turnover rate than microtubules do: $\sim 5/\text{FtsZ}/\text{min}$ (95, 96, 129, 144) versus $0.06/\text{tubulin}/\text{min}$ (98a). This suggests either that FtsZ subunits release from the polymer at a high rate or that even without releasing, subunits in the center of the polymer can exchange nucleotide. If such reactions were more rapid than hydrolysis itself, steady state FtsZ polymers would contain GTP.

Two groups have addressed the steady state nucleotide content of FtsZ polymers (87, 126). Both groups found one equivalent of ^{32}P bound to FtsZ polymerized with $\gamma\text{-}^{32}\text{P}\text{-GTP}$, but it remains unclear whether hydrolysis or phosphate release is the rate-limiting step in the cycle. One group proposed that hydrolysis is rate limiting and polymers contain GTP. They found the exchange of nucleotide between polymer and solution to be faster than the steady state hydrolysis rate (87). The other group suggested that hydrolysis is rapid and phosphate release is slow (126). They determined that the vast majority of ^{32}P bound to FtsZ was inorganic phosphate rather than the terminal phosphate on a GTP. However, in these latter experiments it is unclear whether the hydrolysis occurred in solution or during experimental handling. Clearly, more experiments need to be done to resolve this tricky issue.

In vivo, the nucleotide content of FtsZ polymers may be affected by Z ring proteins that induce protofilament bundling. The kinetics of GTP hydrolysis changes when protofilament bundles are stabilized by calcium or DEAE dextran. Depending on reaction conditions, these cations can either increase or decrease the steady state hydrolysis rate (74, 96, 154). Calcium bundles exchange nucleotide only slowly (87) and may contain GDP-bound subunits (although as above, hydrolysis might have occurred during the time the polymers were sedimented) (125). Interestingly, the degree of curvature in a protofilament may also affect the ability of nucleotide to be exchanged. When curved protofilaments were assembled in DEAE dextran/GDP/Mg $^{++}$ and then excess GTP/EDTA was added, the curved protofilaments immediately straightened, whereas straight protofilaments could not rapidly exchange nucleotide and curve (76). In tightly associated, straight protofilaments multiple hydrolysis events may be needed in order for complete breakage of the bundle and release of the GDP (87).

Implications for dynamic instability and treadmilling The distribution of nucleotide intermediates in FtsZ polymers will help determine whether FtsZ turnover occurs through dynamic instability or treadmilling. In vivo, dynamic instability might trigger sudden depolymerization or force generation during ring contraction. Polymer treadmilling could participate in relocalization of the Z ring during *B. subtilis* sporulation. However, because individual FtsZ polymers are too small to be observed growing and shrinking under the light microscope (Table 1), no direct data currently exist. Neither phenomenon is likely if FtsZ polymers predominantly contain GTP. For dynamic instability to occur in microtubules, the energy from GTP hydrolysis must be stored as strain in the GDP-containing microtubule wall. It is unlikely that individual FtsZ protofilaments could exhibit dynamic instability by the same mechanism. They lack the lateral bonds that could prevent GDP-bound interfaces from breaking or relaxing to a lower-energy, curved conformation. In the various multistranded FtsZ polymers the ability to store the energy from GTP hydrolysis will be influenced by the relative rates of hydrolysis, nucleotide exchange, and protofilament disassembly. Likewise, if FtsZ polymers do not build up a region of GDP at one end of the filament, they could not treadmill with the same mechanism as actin filaments.

NUCLEATION AND COOPERATIVE ASSEMBLY Many biological polymers assemble cooperatively, meaning that it is more difficult for a new polymer to be established than for polymerization to continue from pre-established nuclei or polymer ends (100). Eukaryotic cells take advantage of nucleated assembly to control polymer localization [reviewed in (24, 147)]. The creation of new microtubules is largely limited to the microtubule organizing center, where γ -tubulin, a specialized tubulin isoform, is thought to form the nucleation sites. This localized nucleation allows interphase eukaryotic cells to form a single, radial array of microtubules that organizes the layout of the cell as a whole.

Similarly, cooperative assembly of FtsZ might help limit the number of Z rings in a cell by making polymer addition to a ring that already exists more favorable than initiation of a new ring elsewhere. Although a threshold level of FtsZ must be expressed before Z rings will form (61), potential nucleating factors in the bacterial membrane have remained elusive. In addition, it is unknown whether protofilaments preassemble in the cytoplasm and are only subsequently added to the Z ring or whether assembly of the Z ring occurs solely through direct monomer addition.

To understand the initiation of FtsZ assembly in vivo, it is necessary to determine the role of cooperativity in the assembly of both single and multistranded polymers in vitro. Below we describe the theoretical mechanisms behind cooperative assembly and the methods that can be used to detect it. We then review the evidence for cooperativity in (a) single-stranded polymers assembled in the absence of hydrolysis, (b) single-stranded polymers assembled with Mg^{++} and GTP, and (c) multistranded FtsZ polymers.

Cooperative assembly in multistranded polymers emerges by necessity (100). During polymer initiation, when two monomers associate, only a single new bond

is formed, whereas when a subunit adds to the end of a multistranded polymer, both longitudinal and lateral bonds can form simultaneously. As a result, subunit addition to the end of a multistranded structure is more favorable than de novo formation of a polymer. For microtubules the problem of nucleation is complicated by GTP hydrolysis. Microtubule nuclei may consist of two protofilaments aligned laterally (141). If the protofilaments hydrolyze GTP, they may curl and separate (16). In single-stranded polymers other mechanisms are needed to produce nucleated assembly. A nucleus is defined as the smallest polymer that is more likely to extend than to fall apart again (100). Cooperative assembly of individual protofilaments might therefore emerge if, following the association of a minimal number of subunits, a conformational change increased the affinity for binding of additional subunits.

Cooperative polymer assembly can be detected in a number of ways. The K_d for the formation of dimers should be dramatically higher than that for subunit addition to polymer ends. There is a sharp critical concentration for assembly, below which no polymer forms and above which all additional protein is converted into polymer. Dimers and other complexes smaller than the size of the minimal nucleus are extremely rare. Finally, at protein concentrations just above the critical concentration there are prominent, concentration-dependent lags in assembly because nuclei need to slowly accumulate before polymer extension can rapidly proceed.

Assembly of small polymers in the absence of nucleotide hydrolysis In the absence of nucleotide hydrolysis, FtsZ assembly does not appear to be highly cooperative. Two groups used analytical ultracentrifugation to show that FtsZ dimers are nearly as stable as small FtsZ polymers. Sossong et al. (129) studied assembly in both GDP and magnesium-free GTP and found that FtsZ dimers, trimers, and tetramers were prevalent. In addition, the K_d for association of FtsZ-GDP dimers was identical to that of trimers (31 μM). Rivas et al. (118) determined the dissociation constants for assembly of up to 12 GDP-FtsZ subunits. As polymer length increased, the K_d for binding of additional subunits decreased only by a factor of two. In contrast, there may be up to seven orders of magnitude difference between the K_d for association of actin dimers and that for the addition of actin subunits to filament ends (147).

Assembly with Mg^{++} and GTP Polymerization with Mg^{++} and GTP is more representative of in vivo assembly but is technically more challenging to study. Currently, data exist in support of both noncooperative and nucleated assembly models. In contrast to the hours required for low concentrations of tubulin to polymerize (106, 141), FtsZ assembly can reach steady state in 1 to 60 sec (13, 96, 119). Lags in initial FtsZ assembly have been irreproducibly observed, but they are not likely to be caused by nucleation reactions requiring multiple subunits to come together. Instead, they may be due to aggregated FtsZ needing to disperse before polymerizing or to subunits needing to bind nucleotide (20, 93, 119, 143, 144). Nonetheless, FtsZ polymerization and GTP hydrolysis often initiate rather abruptly at a critical concentration of $\sim 1 \mu\text{M}$ [(74, 95, 96, 118, 129, 144, 148) but see also (119)].

Perhaps polymerization of FtsZ requires a model for cooperative assembly different from those previously developed for microtubules and actin.

Assembly of multistranded polymers Data on the cooperativity of multistranded FtsZ polymers are also somewhat ambiguous. DEAE dextran and calcium appear to enhance FtsZ assembly (73, 94, 154). However, in one report the critical concentration did not appear to be lowered by calcium ($\sim 1 \mu\text{M}$) (154). While no lags have been detected in reactions with calcium and DEAE dextran (124, 125), *M. tuberculosis* FtsZ protofilaments bundle in the absence of these cations and assemble slowly, with brief lags in hydrolysis rates (148).

PART III: PROTEINS THAT MODULATE FtsZ ASSEMBLY

The polymerization properties described above are regulated *in vivo* by proteins that interact directly with FtsZ. Changes in FtsZ polymer structure or GTPase activity can alter the balance between assembly and disassembly. Positive-acting factors promote Z ring formation and stabilize the ring during assembly of the division apparatus. Negative-acting factors prevent FtsZ assembly at inappropriate locations and ensure that the ring is dynamic enough to respond to the signals governing cytokinesis. Together they form a regulatory network that dictates the spatial and temporal control of cytokinesis (Figure 5a).

Below we discuss the proteins known to affect FtsZ assembly. Several of these proteins are not essential for cytokinesis, and none are required for Z ring formation. Their role in regulating Z ring assembly is often confirmed by genetic experiments in which the expression of two proteins is simultaneously altered. Moreover, despite the dramatic evolutionary conservation of FtsZ itself, only two of these proteins, FtsA and ZapA, are themselves broadly conserved. Thus, the proteins that control FtsZ polymerization can have subtle effects that are tailored to the needs of individual species.

Stabilizing Proteins

FtsZ assembly is the first known event in bacterial cytokinesis and is required to localize all other components of the division machinery to the nascent septal site (82, 120). However, the Z ring is not sufficiently stable to support cytokinesis without the assistance of accessory division proteins. To date, the proteins FtsA, ZipA, and ZapA are known to interact directly with FtsZ to stabilize the Z ring.

FtsA FtsA is a widely conserved protein (120) that binds directly to FtsZ (25, 45, 46) and recruits other cell division proteins to the Z ring (82, 120). Although FtsA is not required for Z ring formation (143), the stability and function of the ring are affected by FtsA levels. FtsA is essential in *E. coli* and the ratio of FtsA to FtsZ must be maintained within a narrow range for cells to divide (18); this ratio is achieved by cotranscription of the two genes. FtsA is required for Z ring formation in situations where the Z ring is artificially destabilized (105).

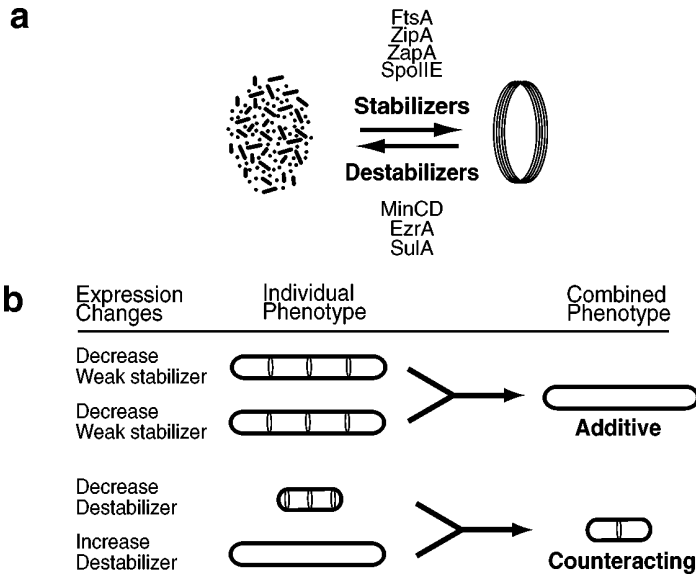


Figure 5 Synergistic interactions. (a) In a cell some FtsZ is in the cytoplasm while other FtsZ is assembled into a membrane-associated ring. The balance between these states is maintained by the combined actions of stabilizing proteins and destabilizing proteins. (b) Synergistic effects are seen when the expression levels of two regulatory proteins are changed simultaneously. (Bottom) A counteracting effect. Loss of a destabilizing factor leads to the formation of extra FtsZ rings at cell poles. Increasing expression of a destabilizing protein inhibits Z ring formation and leads to cell filamentation. Together these two changes can restore normal Z ring formation and cell viability. (Top) An additive effect. Removal of a single, weakly stabilizing protein allows FtsZ assembly but prevents division. Simultaneous removal of two weakly stabilizing proteins can block Z ring formation altogether.

The ability of FtsA to modulate FtsZ polymer stability *in vitro* has not yet been directly examined. Purified FtsA is phosphorylated and can bind and hydrolyze ATP (34, 122). The phosphorylation state of FtsA appears to correlate with ATP binding and membrane association, but a mutation that prevents phosphorylation and *in vitro* ATP binding still rescues conditional *ftsA* alleles (122). FtsA is part of the actin/hexokinase/HSP70 superfamily (139) and has been shown to dimerize (34, 152), but to date there is no evidence that it polymerizes. MreB, a protein required for maintaining cell shape, is a better candidate for a functional prokaryotic homolog of actin (54, 138).

ZipA Like FtsA, ZipA is a Z ring protein that is dispensable for ring formation but is required for recruitment of other proteins to the septal site (44, 69, 82, 120). While ZipA is essential in *E. coli*, it is only conserved in the γ subdivision of gram-negative bacteria. Genetic experiments suggest that ZipA stabilizes FtsZ polymers

in the cytokinetic ring. A twofold overexpression of ZipA suppresses the heat sensitivity of *ftsZ84*, a mutation that results in unstable Z rings at high temperatures (114). ZipA may be enhancing the association of FtsZ with the cell membrane, or it may be stabilizing the interaction between FtsZ protofilaments. The ZipA protein consists of a hydrophobic membrane anchor that is tethered to a cytoplasmic FtsZ binding domain via a flexible peptide (43, 46, 90, 91, 99). In vitro, the cytoplasmic domain of ZipA enhances the bundling of FtsZ protofilaments (45, 114) (Figure 3*d*).

ZapA ZapA is a Z ring protein that is conserved in a wide range of bacteria (42), but in contrast to both FtsA and ZipA, it appears to be dispensable for division under normal growth conditions. *B. subtilis zapA* null mutants are phenotypically unchanged, with wild-type frequency and positioning of Z rings (42). ZapA becomes essential for efficient division in the absence of EzrA or DivIVA, two proteins that normally modulate Z ring assembly, or in cells in which FtsZ concentrations have been artificially lowered (42). In vitro, ZapA interacts directly with FtsZ and induces protofilaments to associate into large bundles (42). The genetic experiments that originally identified ZapA (42) suggest that in vivo this bundling activity may help stabilize Z rings.

THE ADDITIVE EFFECTS OF STABILIZING FACTORS Although individual cell division proteins may be dispensable for Z ring formation, the simultaneous removal of two weakly stabilizing factors can disrupt FtsZ assembly (Figure 5*b, top*). Such synergistic interactions confirm that ZipA and FtsA promote Z ring stability. For example, whereas the loss of ZipA or FtsA alone does not prevent FtsZ assembly, depletion of both FtsA and ZipA in exponentially growing *E. coli* cells blocks the formation of new Z rings and renders previously formed ones unstable (105). Similarly, a mutation in FtsZ that renders it unable to interact with either ZipA or FtsA inhibits its ability to form rings (105).

Destabilizing Proteins

Despite FtsZ concentrations in *E. coli* cells that are well above the critical concentration for in vitro polymer assembly (74), much of the protein is cytoplasmic (130). To prevent aberrant assembly and maintain subunit turnover in the Z ring, destabilizing factors must counter the actions of the polymer-stabilizing proteins discussed above. Below we describe three sets of destabilizing factors: the MinCD complex, EzrA, and Sula.

THE MinCD COMPLEX In *E. coli* and *B. subtilis* the best-characterized proteins affecting FtsZ assembly are the Min proteins. These proteins were originally identified by *E. coli* mutations that caused aberrant septation at cell poles, which led to “minicell” formation (21). MinC and MinD form a division inhibitor (the MinCD complex) that is concentrated at the cell poles (127). Null mutations in either *minC*

or *minD* result in the formation of Z rings at polar positions as well as at medial positions (11, 66). Simultaneous overexpression of MinC and MinD prevents the formation of Z rings and cytokinesis (65, 104).

MinC interacts directly with FtsZ to inhibit Z ring formation (51). It may do so by altering FtsZ polymer assembly, although current data are ambiguous. When FtsZ was incubated with MinC *in vitro*, significantly fewer FtsZ polymers could be detected by electron microscopy (51). However, MinC had no effect on FtsZ's GTP hydrolysis rate (51), suggesting that MinC does not change the stability of protofilaments or their nucleotide exchange rates. In addition, overexpression of MinCD does not change the gel filtration profile of FtsZ polymers in *E. coli* lysates (56). It should be noted that these approaches are not equally sensitive to different aspects of polymer formation. These data would be consistent if MinC reduced the length or bundling of FtsZ polymers without completely inhibiting FtsZ self-association.

MinD is required for concentrating MinC at the membrane of cell poles (50, 84). It associates with the membrane via an N-terminal amphipathic helix (133) and has an ATPase activity that may play a role regulating this membrane localization (48, 52). The polar localization of MinD and hence MinC is dependent on a third protein that is not well conserved across evolutionary boundaries. Mutations in *minE* in *E. coli* or *divIVA* in *B. subtilis* result in the mislocalization of MinCD and extensive cell filamentation (21, 28, 85, 112, 113). MinE and DivIVA share no similarity in either their protein sequence or their mechanism of MinCD localization [reviewed in (127)]. MinE is concentrated at midcell and prevents MinCD from localizing there; instead the entire MinCD complex oscillates rapidly from pole to pole (112, 113). In contrast, DivIVA binds to MinD and statically tethers the MinCD complex to the cell poles (84, 85).

Although the Min proteins are important for preventing aberrant Z ring formation at cell poles, they are not required for establishing the nascent division site at midcell in either *E. coli* or *B. subtilis*. Septation occurs at midcell rather than at the cell poles ~90% of the time in *B. subtilis minCD* null mutants (66) and at a high frequency in *E. coli minCDE* null mutants as well (21). It is thus not surprising that the Min proteins are not conserved throughout the bacteria (120). The lack of conservation suggests either that cell poles are not competent for FtsZ assembly in certain species or that aberrant septation events are prevented through other means.

EzrA EzrA is a nonessential protein that is conserved throughout the low GC gram-positive bacteria (61). An *ezrA* null mutation results in the formation of extra Z rings, primarily at cell poles, a phenotype strikingly similar to that of a *minCD* null mutation (61). EzrA has an N-terminal transmembrane anchor and a large cytoplasmic domain that includes four coiled-coil motifs. An EzrA-GFP fusion distributes diffusely around the periphery of the cell but also concentrates at the cytokinetic ring in an FtsZ-dependent manner (61). The extra rings in EzrA null mutants suggest that EzrA acts throughout the plasma membrane to destabilize newly formed FtsZ polymers and prevent Z ring formation at inappropriate sites. EzrA

activity would need to be overcome at midcell by a factor promoting FtsZ assembly. The biochemical nature of the EzrA-FtsZ interaction has yet to be determined.

SulA SulA is a soluble *E. coli* protein that is induced as part of the SOS response. SulA binds directly to FtsZ and inhibits assembly of the Z ring (49, 53, 56, 92, 135, 136). In vitro, 1:4 ratios of SulA to FtsZ can inhibit polymer assembly without affecting FtsZ's GTP hydrolysis activity (136). At a 1:1 ratio of SulA to FtsZ, both assembly and GTP hydrolysis are inhibited (136), which suggests a model in which SulA binds to one end of FtsZ and prevents the addition of new subunits. At lower concentrations SulA would cap polymers, limiting their length, whereas at higher concentrations SulA would completely prevent FtsZ dimerization and formation of the active site. SulA's ability to prevent assembly at relatively low concentrations suggests that in vivo even limited induction of SulA may be sufficient to rapidly block cytokinesis.

EVIDENCE FOR OTHER PROTEINS To date, only two other proteins have been suggested to affect Z ring stability. The first, SpoIIE, colocalizes with FtsZ in polar rings during sporulation in *B. subtilis* (64). SpoIIE is a phosphatase that helps activate cell type-specific gene expression, but it may also play a structural role by directly stabilizing polar Z rings (7, 33, 58). The second, FtsW, is an essential cell division protein. Loss of FtsW was originally suggested to render Z rings less stable (12). However, more recent data indicate that FtsW is recruited to the Z ring late in the assembly process and functions primarily to localize its cognate transpeptidase FtsI (PBP3) (86).

THE COUNTERBALANCING EFFECTS OF STABILIZING AND DESTABILIZING FACTORS

The destabilization of FtsZ polymers by proteins that inhibit Z ring formation can be counteracted by a simultaneous stabilization of FtsZ assembly via different means (Figure 5b, bottom). Mutations that stabilize FtsZ polymers, such as *ftsZ2*, render FtsZ resistant to the destabilizing effects of both MinC and SulA (8, 9, 53, 136). Similarly, in *B. subtilis* a null mutation in the gene encoding EzrA, a destabilizing factor, suppresses the inhibition of Z ring formation and cell division associated with 15-fold overexpression of MinCD (65). Moreover, increasing MinCD levels can compensate for the loss of EzrA to some extent, as polar rings do not form in *ezrA* null mutant cells in which MinCD has been overexpressed (65). Synergistic effects such as these are useful for confirming the activities of nonessential genes.

PERSPECTIVES AND OUTLOOK

The forces governing Z ring assembly during bacterial cytokinesis are strikingly similar to those controlling the microtubule and actin cytoskeletons in eukaryotic cells. FtsZ polymer dynamics are modulated during the cell cycle by a precisely

balanced network of positive- and negative-acting factors. Assembly is exquisitely sensitive to these factors because of the intrinsic reversibility of FtsZ polymerization. Thus, developmental and environmental signals are integrated during Z ring assembly and maintenance to ensure the proper spatiotemporal control of cytokinesis.

A comprehensive understanding of FtsZ dynamics during cell division will likely depend on the extension of the experimental techniques currently being used. For example, polymerization assays with improved time resolution and linearity will help illuminate difficult enzymatic issues in FtsZ polymer dynamics. Furthermore, the relatively recent identification of ZapA and EzrA highlights the importance of nonessential proteins in cell division. The discovery of additional proteins that exert subtle effects on FtsZ assembly may require sophisticated biochemical or genetic approaches. Ultimately, the analysis of these proteins and their effects on FtsZ polymerization, both *in vivo* and *in vitro*, will pave the way for more precise models of FtsZ dynamics within the Z ring.

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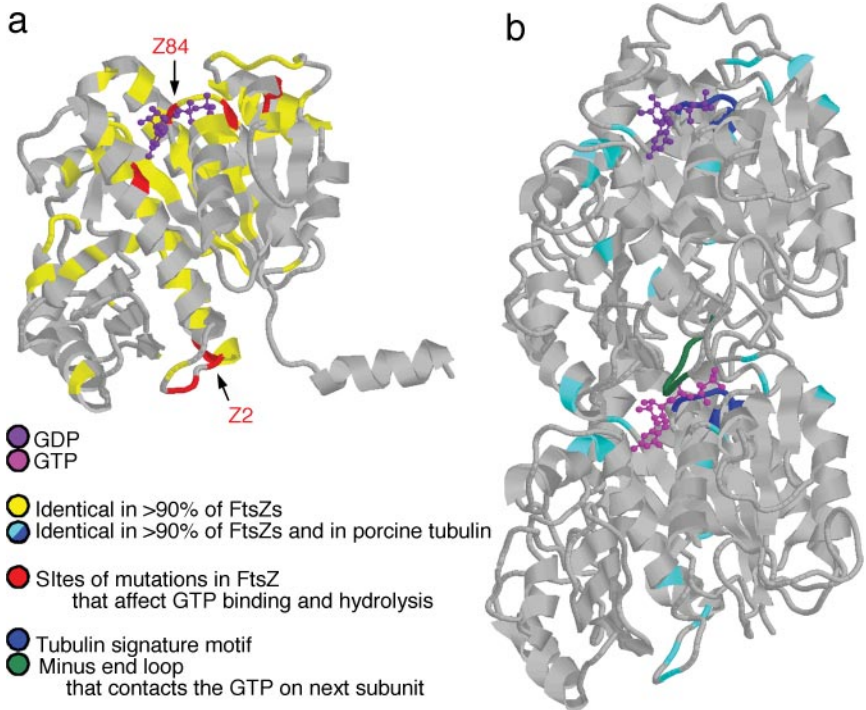


Figure 2 FtsZ and tubulin atomic structures. FtsZ and tubulin share minimal sequence homology but similar protein folds. (a) *Methanococcus jannaschii* FtsZ (70). Highly conserved residues (yellow) concentrate around the GTP binding site. Mutations that affect GTP binding and hydrolysis (red, e.g., Z84, Z2) are located at the plus end of the protein, where GTP binds, and at the minus end of the protein. (b) Porcine α/β tubulin dimer (98). A small number of residues in pig tubulin are also highly conserved in FtsZ (cyan). They include the tubulin signature motif (blue), which is important for GTP binding in both proteins, and a loop at the minus end of the subunits (green), which contacts the GTP in the neighboring subunit after assembly.