The Search for Antibacterial Inhibitors Targeting Cell Division Protein FtsZ at Its Nucleotide and Allosteric Binding Sites

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Abstract: The global spread of bacterial antimicrobial resistance is associated to millions of deaths per year, many of which were previously treatable. This, combined with slow antibiotic deployment, has created an urgent need for developing new antibiotics. A still clinically unexploited mode of action consists in suppressing bacterial cell division. FtsZ, an assembling GTPase, is the key protein organizing division in most bacteria and an attractive target for antibiotic discovery. Nevertheless, developing effective antibacterial inhibitors targeting FtsZ has proven challenging. Here we review our decade-long multidisciplinary research on small molecule inhibitors of bacterial division, in the context of global efforts to discover FtsZ-targeting antibiotics. We focus on methods to characterize synthetic inhibitors that either replace bound GTP from the FtsZ nucleotide binding pocket conserved across diverse bacteria or selectively bind into the allosteric site at the interdomain cleft of FtsZ from Bacillus subtilis and the pathogen Staphylococcus aureus. These approaches include phenotype screening combined with fluorescence polarization screens for ligands binding into each site, followed by detailed cytological profiling, and biochemical and structural studies. The results are analyzed to design an optimized workflow to identify effective FtsZ inhibitors, and new approaches for the discovery of FtsZ-targeting antibiotics are discussed.

Keywords: antibiotics; bacterial division; FtsZ; nucleotide-replacing inhibitors; allosteric inhibitors

1. Cell Division Protein FtsZ as an Antibacterial Target

Bacterial pathogens easily develop resistance to antibiotics used to treat infections. In some cases, strains can become resistant to all clinically employed antibiotics. This major threat to human health has not been met by the development of new antibiotics, meaning we face a post-antibiotic era where common bacterial infections are potentially untreatable and routine surgery or chemotherapy become too dangerous [1]. In fact, bacterial antimicrobial resistance (AMR) has been recently estimated to be associated to ~5 million deaths worldwide, of which ~1.3 million were directly attributable to AMR [2]. Among the top resistant pathogens, antibiotic resistant Escherichia coli was associated to ~0.8 million deaths, and meticillin-resistant S. aureus (MRSA) alone was estimated to cause more than 100,000 deaths and 3.5 million years of healthy life lost due to disability or premature death. There is thus an urgent need to discover new antibiotics and novel antibacterial targets.

Among the cellular processes essential for bacterial reproduction and infection, bacterial cell division [3] still remains clinically unexploited. The central protein organizing cell division in most bacteria is FtsZ, which together with membrane-tethering proteins forms a dynamic ring (Z-ring) at the division site, that recruits the other divisome proteins to coordinate cell wall remodeling and membrane constriction during cytokinesis [4–7]. FtsZ is an assembling GTPase whose structural core is shared by tubulin [8], the constituent...
of eukaryotic microtubules. Whereas tubulin is a classical target of numerous anticancer drugs, discovering new useful antibiotics targeting FtsZ remains a challenge. Over one hundred X-ray structures of tubulin-inhibitor complexes are available [9], but only a handful of FtsZ small-molecule complex structures have been reported to date [10–16], suggesting a lesser degree of druggability of FtsZ compared to tubulin. Nevertheless, considerable knowledge has been acquired about the structure, assembly dynamics, and function of FtsZ filaments and the Z-ring, the FtsZ partner proteins, and several small molecule chemotypes that specifically interact with FtsZ.

The bacterial cytokinetic Z-ring (Figure 1A) is formed by condensates of FtsZ filaments that move circumferentially to guide correct septal peptidoglycan synthesis [17–20]. FtsZ is a protein assembly machine forming translocating filaments that grow from one end while shrinking from the other, in a process called treadmilling (Figure 1B). Treadmilling is driven by GTP hydrolysis, which takes place in the complete GTPase active site formed at the association interface between consecutive subunits in the filament, and triggers the disassembly of GDP-bound subunits at the shrinking end. The nucleotide-dependent interfacial interactions and the GTP hydrolysis mechanism have been recently revealed at atomic resolution by filament structures of SaFtsZ in different nucleotide states [21]. FtsZ cooperatively assembles into long single-stranded filaments, rather than forming short oligomers, which is made possible by conformational switching between a relaxed protein form (R) with low association affinity in monomers and a tense (T) high association affinity form in filaments [22–26]. The defined treadmilling direction (kinetic polarity) of FtsZ filaments is also enabled by the structural switch between T subunits inside filament and R monomers irrespective of the nucleotide bound, which creates structurally different encountering interfaces for association at each filament end [26,27].

The core structure of FtsZ consists of an N-terminal nucleotide-binding domain (NBD) and a C-terminal GTPase activation domain (GAD), joined by the long central helix H7 (Figure 1C). A variable unstructured C-terminal linker connects the FtsZ core to a binding motif for several of the FtsZ membrane tethering and partner proteins [5], whose interactions are also potential antibiotic targets [28]. FtsZ proteins spontaneously fold without requiring chaperones [29]. However, guanine nucleotide binding into its NBD pocket is required to stabilize the SaFtsZ core, which otherwise unfolds in the solution [30] and can thus be preferentially degraded by ADEP-activated ClpP protease [31]. Upon filament assembly, the interfacial GTPase site is completed as the GAD’s co-catalytic loop T7 inserts into the next subunit’s nucleotide binding pocket that adapts to the contact. The GAD itself rotates in the T-FtsZ conformation, opening a side interdomain cleft, flanked by helix H7 and the GAD’s beta-sheet, where the allosteric inhibitor PC190723 and related benzamides bind [11–13,16] (Figure 1C). The FtsZ inhibitor protein SulA [32] and peptide MciZ [33,34] bind the bottom of the GAD of FtsZ in the R-conformation, exerting monomer sequestering and filament capping activities.

FtsZ was soon recognized as an attractive antibacterial target [35,36] that was validated with the effective antistaphylococcal inhibitor PC190723 [11,37]. However, in a decade since our previous review [38], FtsZ inhibitors have not reached the clinic. It has been pointed out that many inhibitors reported in the literature were not really proven to specifically target FtsZ, slowing down progress in this field [39]. Currently, there are close to 200 publications with numerous compounds associated with FtsZ inhibition. The more recent reviews on FtsZ small molecule inhibitors include Refs. [40–43].

Among other approaches to targeting FtsZ, regulatory peptide MciZ inhibits FtsZ function across the Bacillus species and it is also recognized by FtsZ from S. aureus (SaFtsZ) [34]. However, conjugation with an effective bacterial membrane-penetrating peptide would be required for the delivery of MciZ into S. aureus cells. On the other hand, intracellular MRSA surviving antibiotic treatment within host cells is thought to represent an important pathogen reservoir in invasive and recurrent infections, which is experimentally eliminated by an antibody–antibiotic conjugate [44]. In addition to small molecule and peptide inhibitors, antisense peptide-nucleic acid mimetic polymers against the ftsZ gene conju-
gated to cell-penetrating peptides have been employed to suppress bacterial growth [45]. CRISPRi approaches can also be employed to control ftsZ transcription and thus bacterial division [46]. Targeted protein degradation approaches include FtsZ degradation by Clp cellular protease activated by an acyldepsipeptide (ADEP) antibiotic [47].

In this review we focus on the two main FtsZ binding sites for small molecules, the nucleotide site and the allosteric site, and the methods developed to screen and study specific antibacterial inhibitors targeting these sites.

2. The FtsZ Nucleotide Binding Site

2.1. Selective Inhibition of FtsZ Versus Tubulin by C8 Nucleotide Analogs

In view of the structural similarity of the FtsZ and tubulin enzymatic cores [8], a potential problem for FtsZ inhibitors could be cross-inhibiting human tubulin. However, both proteins share limited sequence identity, and it was soon realized that typical tubulin inhibitors do not inhibit FtsZ, whereas FtsZ inhibitors infrequently act on tubulin. A difference between the interfacial active sites of FtsZ and tubulin was demonstrated with GTP analogs bearing small hydrophobic substituents at the C8 position of the guanine ring, such as 8-methoxy-GTP and 8-Br-GTP, which bind with high affinity and selectively inhibit FtsZ polymerization in vitro, while promoting microtubule assembly and being substrates for tubulin GTPase [10]. The FtsZ assembly inhibitory activity of C8-GTP derivatives correlates with their binding affinity [10,48]. The crystal structure of thermophilic Aquifex aeolicus FtsZ (AaFtsZ) in complex with 8-morpholino-GTP [10] revealed a binding
mode similar to GDP, with the morpholine ring oriented outward from the binding pocket towards the position of the next subunit in a protofilament (Figure 1D). A similar anti conformation of the glycosidic bond in the 8-morpholino-nucleotide bound to B. subtilis FtsZ (BsFtsZ) was deduced from NMR trNOESY analysis, whereas 8-pyrrolidino-nucleotide binds in the syn conformation, different from the natural nucleotide [48]. In this pose, the pyrrolidine ring is roughly in place of the guanine ring and the guanine ring points out from the binding pocket, as shown by a molecular dynamics (MD) model complex with thermophilic Methanocaldococcus jannaschii FtsZ (MjFtsZ, Figure 1E). Homology modeling BsFtsZ with bound C8-nucleotide analogs onto filaments of SaFtsZ (supported by a 80% protein sequence identity) showed that the inhibition of BsFtsZ assembly may be simply explained by steric clashes of the nucleotide analogs with the incoming FtsZ monomer [48]. The C8-GTP analogs were thus useful tools to demonstrate how to selectively inhibit FtsZ without inhibiting tubulin in vitro. However, they were devoid of antibacterial activity, possibly not passing the bacterial cell envelope. Initial attempts to design prodrugs based on C8-nucleotide analogs that would enter bacteria and be hydrolyzed in the cytosol to release C8-nucleotide inhibitors were unsuccessful [50]. Nevertheless, a different analog was recently reported, 3-amino-3-deoxyguanosine (ADG), which behaves as a prodrug mimic of GTP; ADG is phosphorylated to ADG-5′-triphosphate, a mimic of GTP that inhibits transcription and FtsZ-GFP localization to the Z-ring [51].

2.2. Fluorescent Nucleotide Analogs and Competition Assays for FtsZ Inhibitors

Among various fluorescent nucleotide conjugates widely employed to investigate nucleotide-binding proteins, we selected 2’/3’-O-(N-methylanthraniloyl)-GTP (mant-GTP), which bears the small fluorescent mant tag on the ribose ring, to probe the GTP-binding site of the nucleotide-devoid apo-MjFtsZ in monomers and filament polymers [52]. Both 2’- and 3’-substituted mant-GTP isomers are in equilibrium and both bind in the natural anti-glycosidic bond conformation, as deduced from NMR studies [53]. In both cases, the mant tag strongly interacts with FtsZ at an extension at the back of the GTP-binding pocket, as indicated by MD simulations. Mant-GTP can inhibit or induce FtsZ assembly and be hydrolyzed, depending of the FtsZ species employed [53].

On the other hand, a small water-filled cavity below the nucleotide-binding pocket is available for small ligand binding, such as co-solvent N-methyl-2-pyrrolidone in SaFtsZ crystals (Figure 1C) [13,16,30]. Together with the C-8 nucleotide analogs, these results underscore the plasticity of the FtsZ nucleotide binding site and support possible ligand extensions in the design of inhibitors.

The binding of mant-GTP to apo-FtsZ entails a reduction of the rotational diffusion of the probe that is detected by a marked increase of its fluorescence anisotropy. We employed this property to devise competition methods for any molecules able to replace the nucleotide in stable apo-MjFtsZ [54], in BsFtsZ [55,56], in apo-SaFtsZ, and also apo-FtsZ from E. coli (apo-EcFtsZ) stabilized by osmolytes [30]. These robust fluorescence polarization (FP) assays with stabilized apo-FtsZ monomers and a fluorescent GTP derivative can be manually performed or conveniently adapted for high-throughput screening (HTS) for nucleotide-replacing inhibitors. The assayed concentration of the compounds under study should be below their solubility limit. Insoluble compound precipitates are typically detected by interference from the scattering of polarized excitation light giving artifactually high anisotropy values. It is also important to exclude colloid-forming aggregators that may non-specifically bind and inactivate FtsZ [57]. Therefore, to determine ligand solubility and exclude FtsZ aggregation, we separately subject inhibitor and protein–inhibitor-mant GTP solutions to high-speed centrifugation, followed by measuring the inhibitor fraction and the probe fluorescence anisotropy remaining in the supernatant [55,56]. The mant-GTP FP competition method is thus useful for inhibitor screening and affinity measurements, although it is not meant to replace direct binding methods for inhibitor characterization, such as analytical co-sedimentation measurements [55] and isothermal titration calorimetry (ITC) when technically feasible [34].
2.3. Synthetic Nucleotide-Replacing FtsZ Inhibitors

We employed the mant-GTP FP assay to screen for GTP-replacing ligands among compounds from the literature [38], virtual screening (VS), and compounds selected from an in-house synthetic library by docking into the BsFtsZ GTP site. The latter yielded several polyhydroxy aromatic compounds that specifically bind to FtsZ, impair the localization of FtsZ to the Z-ring, inhibit B. subtilis cell division, and show antibacterial activity against clinical isolates of MRSA and levofloxacin-resistant Enterococcus faecalis [55]. A series of analogs with different aromatic scaffolds, linkers, and substitution patterns of the hydroxyl groups were then synthesized (Figure 2A) in iterative rounds guided by the binding affinity (K_D value in mant-GTP FP assays) and minimal division inhibitory concentration (MDC value on B. subtilis) below the MIC value. Among them, compound 1 binds into the GTP site with K_D = 0.5 μM (Figure 2B), impairs normal FtsZ assembly in vitro and FtsZ-GFP localization at the mid-cell (Figure 2C, 3 μM), and has a 7 μM MIC value on MRSA. In a BsFtsZ-1 model complex, obtained by ligand docking and MD simulations, this inhibitor replaces GDP in several of its interactions with key residues in the nucleotide binding pocket (Figure 2D) [56]. Nevertheless, the affinity of these nucleotide-replacing inhibitors for FtsZ monomers is still lower than that of GTP, which is present at a high concentration in the bacterial cytosol.

![Figure 2](image_url)

Figure 2. (A) General chemical structure of polyhydroxy aromatic GTP-replacing inhibitors and of the identified inhibitor 1 [55,56]. (B) Displacement of mant-GTP (1 μM) from BsFtsZ (0.9 μM binding sites) by compound 1, measured by fluorescence anisotropy. The line is the least squares best-fitted competition curve (K_D = 0.5 μM). (C) Upper panel, cells of B. subtilis 168 treated with 1 (4 μM) for 3 h and observed by phase-contrast microscopy. Mean and standard error of cell length measurements are indicated. Lower panel, cellular localization of FtsZ-GFP in B. subtilis SU570 following growth for 1.5 h in the absence and presence of 1 (3 μM). Scale bars: 10 μm. (D) BsFtsZ-1 model complex from docking and MD simulations. The ligand is shown in sticks represented with cyan carbon atoms. The amino acid residues that interact with the inhibitor and are shared key residues for nucleotide binding in the BsFtsZ-GDP complex [PBD 2RHL, [56]] are indicated and displayed in sticks. ((B–D) adapted with permission from [56]; copyright 2015 American Chemical Society).

It can be estimated that only a small fraction of the FtsZ molecules in a bacterial cell will be bound by compound 1 or its analogs. However, these inhibitors are able to effectively impair FtsZ assembly and inhibit cell division, which might be explained by the binding of the substoichiometric inhibitor by a few subunits perturbing the correct
assembly of a FtsZ filament. Alternately the inhibitor-binding affinity for the nucleotide site in the filament association interface would have to be substantially larger than the affinity for monomers that is measured with the mant-GTP FP assay [55].

Interestingly, natural crysopaentins and synthetic hemi-crysopaentins are other examples of competitive inhibitors of nucleotide binding by FtsZ, endowed with antibacterial activity on S. aureus, as well as on permeable E. coli envA1 cells, but not cytotoxic to mammalian cells [58,59]. Many of our nucleotide-replacing compounds selectively perturb FtsZ vs tubulin assembly, and their growth inhibitory concentrations on human cells are typically above those required for the inhibition of bacterial cell division [55,56]. It can thus be concluded that it is feasible to selectively inhibit bacterial division by targeting the FtsZ nucleotide binding site without cross-inhibiting mammalian cell tubulin. Curiously, despite the many microtubule targeting agents, to our knowledge, no tubulin nucleotide-replacing inhibitors have been discovered [9,60,61].

2.4. Antibacterial Activity of Nucleotide-Replacing FtsZ Inhibitors

The identified nucleotide-replacing inhibitors show activity on Gram-positive pathogens, particularly MRSA, over Gram-negative species [55,56,58], possibly related to an inability to cross the Gram-negative outer membrane [59] rather than to small differences in the FtsZ GTP-binding site across bacterial species. Interestingly, the antibacterial activity MIC values in our compound series correlate better with their affinity $K_D$ values, than with FtsZ in vitro polymerization inhibition [56]. We favor the use of target affinity, rather than FtsZ polymerization or GTPase assays, as a predictor of antibacterial activity. Figure 3 shows several representative GTP-replacing inhibitors along the optimization process, on an activity–affinity log plot. It can be observed that the MIC values on MRSA (dotted circles) are within one factor of two of B. subtilis MIC values (filled circles). Further development of antibacterial nucleotide-replacing FtsZ inhibitors will, in our opinion, require enhancing chemical diversity by means of fragment screening, HTS, and new approaches discussed in Section 6.

Figure 3. Correlation of antimicrobial activity (MIC) with binding affinity ($K_D$) of inhibitor binding to the BsFtsZ nucleotide site. Data from representative analog compounds are shown together with their chemical structures. Filled circles, B. subtilis MIC; dotted circles, methicillin-resistant S. aureus MIC. The dashed lines are the 95% predictive intervals for B. subtilis MIC vs $K_D$ values of 36 analogs (adapted with permission from [56]; copyright 2015 American Chemical Society).
3. The Filamenting Phenotype and the Cytological Profile of FtsZ Inhibitors

3.1. The Importance of Assessing Membrane Integrity

The inhibition of the divisomal protein function in bacilli results in characteristic cell elongation (filamentation), while cocci show an enlarged spherical size (ballooning). The name of the *ftsZ* gene comes from “filamenting temperature sensitive mutant Z” [62]. Thus, a convenient cell-based method to screen for FtsZ/divisomal inhibitors is the induction of a relevant filamenting phenotype, documented by images and cell length measurements (Figure 2C). However, similar filamentation effects may also be induced by a loss of the transmembrane potential, which results in FtsA and a secondary FtsZ release from the membrane [63]. In this context, several putative FtsZ inhibitors were shown to really act by impairing membrane potential and permeability [64]. Aminoglycoside antibiotics are known to have a second mode of action altering membrane integrity, but in the case of FtsZ, this would hamper target-based optimization. Therefore, any selected compound inducing bacterial cell filamentation should not be toxic at the MDC and must also be tested for membrane integrity (by exclusion of the DNA-binding fluorescent dye propidium iodide) and for membrane potential with a suitable probe (such as 3,3’-diethyloxacarbocyanine iodide, DiOC₂) [65]. Whereas filamenting *B. subtilis* cells treated with our main GTP-replacing inhibitors, such as 1 (MDC 4 µM), had intact membranes and membrane potential, a few other analogs caused membrane lysis [56,65]. FtsZ inhibitors, following target-based optimization, might be combined with membrane depolarizing compounds, for the dual inhibition of cell division. On the other hand, several physiological mechanisms and proteins can inhibit FtsZ and block cell division, including the DNA damage SOS response with the expression of SulA, the nucleoid occlusion machinery, nutrient availability through moonlighting enzymes, sporulation through MciZ, and certain phage-encoded polypeptides [65] (and references therein).

3.2. The Cytological Profile of FtsZ Inhibitors

In addition to cell filamentation measurements and FtsZ-GFP ring impairment observations (Figure 2C) typically employed for FtsZ inhibitor cell-based screening, cytological profiling methods for FtsZ targeting inhibitors in *B. subtilis* are available [65]. Bacterial cytological profiling uses quantitative fluorescence microscopy to rapidly identify the cellular mechanism of action of antibacterial agents, distinguishing different cellular pathways and targets [66]. Among several cellular effects of seven selected inhibitors binding to different FtsZ sites, principal component analysis and cluster maps of cell length combined with nucleoid length showed separate clusters for the FtsZ inhibitors (with one exception) and five known antibiotics with different mechanisms of action (Figure 4A). The distribution of distances between the remaining Z-rings in treated cells clearly distinguished six FtsZ inhibitors from known antibiotics and untreated control cells (Figure 4B). This study also revealed that exogenous synthetic polypeptide MciZ enters *B. subtilis* cells to effectively inhibit cell division [65], supporting the cytological profile approach for characterizing FtsZ inhibitors. MciZ modifies the treadmilling rate of FtsZ around the Z-ring in *B. subtilis* [17], and slows down *Bacillus* division, including pathogenic *B. cereus* and *B. anthracis* [34].
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**Figure 4.** Cytological profile of antibacterial FtsZ inhibitors. Representative examples of *B. subtilis* cells observed upon treatment with 1 (3.5 µM), PC190723 (5.6 µM) and nisin (10 mg/L). (A) Undivided individual cells of *B. subtilis* 168 cells (phase contrast) and nucleoid morphology (DAPI). (B) Effects of FtsZ inhibitors and nisin on FtsZ subcellular localization on *B. subtilis* SU570 (FtsZ-GFP, contrast inverted). Scale bars: 10 µm. Right graphs, dendrograms obtained after a hierarchical cluster analysis based on values from a principal component analysis (PCA) using unweighted variables cell length and nucleoid length (upper graph) or Z-rings distribution (lower graph) [65].

4. The FtsZ Interdomain Cleft

4.1. The Reference Antibacterial FtsZ Inhibitor PC190723

The small compound 3-methoxybenzamide was known to have weak antibacterial and filamentation activities, which are suppressed by mutations in the *ftsZ* gene of *B. subtilis* [67]. From this starting point, a large medicinal chemistry program guided by MIC and cell morphology [68] yielded the potent antistaphylococcal FtsZ difluorobenzamide inhibitor, PC190723 (Figure 5A) [11,37], which restores MRSA susceptibility to methicillin [11]. PC190723 is an inducer, instead of a suppressor, of FtsZ filament assembly and condensation [69] which makes FtsZ assemble into delocalized cellular foci rather than at the Z-ring [70]. By binding into the bottom of the open interdomain cleft, located between the GAD domain beta-sheet, helix H7, and loop T7 (Figure 1C), PC190723 stabilizes the open cleft T conformation of SaFtsZ and thus, filament formation, as shown by the crystal structure of the complex [11,12,71]. Unfortunately, PC190723 and its numerous benzamide analogs have not reached the clinic, possibly due to non-optimal pharmacological properties and to the relatively high frequency of resistance mutations in the *ftsZ* gene (FOR~10⁻⁸), although several improved derivatives are able to overcome certain resistance mutations [13,72] and reduce FOR in antibiotic combination therapy [73].
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**Figure 5.** (A) Chemical structure of fluorescent probes and allosteric inhibitors targeting the FtsZ interdomain cleft. (B) Several examples from a florescence anisotropy screen for allosteric FtsZ inhibitors. The displacement of probe (S)-9 (3 μM) from stabilized BsFtsZ polymers (8 μM binding sites) by two different concentrations of each compounds, 20 μM (gray dots) and 200 μM (black dots), is monitored. Control values with the solvent vehicle (DMSO), PC190723 and its 2,6-difluoro-3-methoxybenzamide (DFMBA) head and heterocyclic tail 6-(chloro [1,3] thiazolo [5,4-b] pyridin-2-yl)methanol (CTPM), are included. The dashed lines represent the fluorescence anisotropy of the probe partially bound by FtsZ polymers and of the free probe. (C) Cell length measurements of \textit{B. subtilis} cells treated with compound 11, 12, and 13 for 3 h. Raw data plots with mean values ($n$ $\geq$ 30) \cite{16}.

### 4.2. Fluorescent Probes for the FtsZ Interdomain Cleft and the Assembly Switch

In order to gain insight into the FtsZ interdomain cleft, we developed fluorescent PC190723 analogs based on the difluorobenzamide head, by replacing the heterocyclic tail with small fluorophores \cite{25}. Among the synthesized compounds, several nitrobenzoxadiazole-based analogs were found to be suitable probes in terms of FtsZ affinity and fluorescent properties. The probes were selected for specific fluorescence anisotropy changes upon binding to polymers of BsFtsZ and SaFtsZ versus resistance mutants and Gram-negative FtsZ which cancelled binding. Probe 8 was biochemically characterized and shown to weakly label the Z-ring in bacterial cells, whereas at higher concentrations, it induced the characteristic cell division inhibition phenotype. The binding of probe 8 ($K_D$ = 26 μM) was employed to monitor the opening and closing of the interdomain cleft during the BsFtsZ and SaFtsZ assembly–disassembly process, thus demonstrating the operation in solution of the FtsZ structural assembly switch \cite{25}, which had been inferred from the open cleft crystal structure.
of SaFtsZ filaments compared to other closed FtsZ monomer structures at that time. This was soon followed by the determination of closed structures of SaFtsZ monomers [13,26]. However, BsFtsZ or other filament structures have not been reported so far.

4.3. Fluorescence Polarization Assay for Allosteric FtsZ Inhibitors

Fluorescent benzamide probes with enhanced affinity (S)-9 and (R)-10 (Figure 5A; \( K_D \) values 8 \( \mu M \) and 2 \( \mu M \), respectively) were designed and employed in FP competitive methods to screen for ligands binding into the FtsZ interdomain cleft and determine their binding affinity [16]. These methods use FtsZ polymers stabilized by mild crosslinking with glutaraldehyde, which do not disassemble upon nucleotide hydrolysis or by the action of assembly inhibitors binding elsewhere in the FtsZ molecule. Raw (S)-9 probe fluorescence anisotropy values at two compound concentrations and solubility measurements can be employed to screen compounds binding into the FtsZ allosteric site, distinguishing low from higher affinity inhibitor candidates (Figure 5B). Combined with phenotypic screening (Figure 5C), even the action of relatively weak FtsZ-targeting compounds can be effectively detected, such as compound 11, which is ~20-fold more active than 3-methoxybenzamide [16].

4.4. Enhancing the Affinity of Simplified Benzamide Inhibitors

In a full competition assay using our FP method, the affinity of compounds replacing the probe (R)-10 can be determined (Figure 6A). Combining affinity for BsFtsZ polymers and cell division inhibition activity on B. subtilis, a series of benzamide-based inhibitors with halogenated aryl tails (Figure 5A) were iteratively designed, reaching \( K_D \) and MDC values comparable to those of PC190723, while keeping a relatively simple chemical structure. Compound (R)-17 has \( K_D = 0.2 \mu M \) and B. subtilis MDC = 1 \mu M. Derivatives 15, 16, (R)-17, and (S)-17, at a somewhat higher concentration, impair FtsZ-mCherry localization into the Z-ring (Figure 6B) and selectively inhibit the cell division and growth of S. aureus rather than the growth of human lung fibroblasts. FtsZ targeting was confirmed by analyzing spontaneous resistance mutations in MRSA, which predominantly mapped to the ftsZ gene, clustering around the FtsZ interdomain cleft [16]. Figure 6C shows the positions of the amino acids (in red) modified in the MRSA mutants, resistant to 15 around the interdomain cleft where the inhibitor binds.

4.5. Structural Insights into Allosteric Inhibitor Binding to S. aureus FtsZ

We have determined the high-resolution crystal structures of SaFtsZ in a complex with 2,6-difluoro-3-methoxybenzamide (DFMBA) and inhibitors 14 and 15. In stark contrast, several other specifically binding compounds and the fluorescent probes did not show in the electron density maps [16], although they rendered stable MD model complexes [25]. Crystal structures of SaFtsZ in a complex with a different probe, TXA6101-Bodipy (BOPF), and with the inhibitors TXA6101 and TXA707 are also available [13,14]. A comparison of these structures shows that the protein interactions with the superimposable difluorobenzamide rings of each compound are conserved in all cases, including the amide group establishing hydrogen bonds to V207, L209, and N263, and coordinating a K\(^+\) ion bound by loop T7, as well as hydrophobic interactions made by the difluorobenzene ring. The halogenated phenyl moieties of 14 and 15 bind between T309 and G196, with their benzamide rings, adopting a flat geometry. The T309 residue acts as a gate, connecting with G196 in a closed conformation with DFMBA. Consequently, both G196 and T309 are among the positions modified by resistance mutations (Figure 6C). The M226 side chain constitutes another gate that gives access to a ligand-induced hydrophobic pocket for TXA707 and flexible inhibitor TXA6101 (not shown; [13]). The PC190723/15 binding site connects through a pore, formed by the M226, I228, and G192 side chains, with a smaller pocket [16] and the upper large unfilled cavity above the interdomain cleft, which is flanked by the loop containing L249 (Figure 6C, surface representation in Figure 1 [16]). This interesting area, overlapping with
the Taxol binding site of β-tubulin in microtubules [74], has not been targeted in FtsZ, but remains available for potential ligand extension [16,25].

Figure 6. (A) Displacement curves of probe (R)-10 (3 µM) from stabilized BsFtsZ polymers (2.7 µM binding sites) by synthetic allosteric inhibitors 13 (black), 15 (blue), 16 (green), (R)-17 (red), (S)-17 (cyan), and by PC190723 (gray). Lines correspond to the best-fitted competition curve in each case. (B) S. aureus cells expressing FtsZ-mCherry grown for 1.5 h in the absence (control) or presence of 15, 16, (R)-, (S)-17, and PC190723, observed by fluorescence microscopy. Scale bar: 2.5 µm. (C) Scheme of the structure of the SaFtsZ-15 complex around the inhibitor (cyan) binding cleft. The positions of the MRSA inhibitor resistance mutations analyzed, G196S, L249V, M262I, and T309I, are indicated by the wild type side chains (red). Indicated as well are residues V207, L209, and N263 (green) that establish hydrogen bonds with the amide group of 15. The T309 side chain has rotated to lodge the trifluoromethylphenyl ring between T309 and G196. The side chains of M226, I228, and Q192 (salmon) form a pore connecting the binding site to the upper part of the cleft. A K+ ion stabilizing loop T7 is also coordinated by the amide carbonyl of 15. Not shown are additional residues making hydrophobic contacts with the inhibitor (PDB 6YD6; Table S4, Figures 6 and S6 in ref. [16]).

4.6. Antibacterial Activity of Benzamide-Based Allosteric FtsZ Inhibitors

PC190723 and its analogs typically have narrow-spectrum antibacterial activity on several Gram-positive organisms, related to differences in the amino acid side chains lining the FtsZ interdomain cleft in other bacterial species [37]. Although this cleft is thought to open as well in T-state EcFtsZ filaments [26], two charged residue pairs have been proposed to form salt bridges between helix H7 and the C-terminal beta sheet, preventing cleft access to PC190723 in resistant species [75]. An allosteric modulator equivalent to PC190723 is thus missing for E. coli FtsZ, in addition to the general problem in crossing the outer membrane of Gram-negative bacteria [76], although the analog TXA6101 is active on Klebsiella pneumoniae [77].

Similarly to the nucleotide-replacing inhibitors, we found that the inhibition of bacterial growth correlates with the binding affinity of allosteric inhibitors [16], as shown for several representative inhibitors along the optimization process (Figure 7). On the other hand, changes in FtsZ GTPase activity are potentially misleading indicators of the activity of benzamide inhibitors, unless a complete analysis at different inhibitor and FtsZ concentrations is performed, due to their mixed FtsZ assembly promoting (GTPase activation) and GTPase modulatory effects [65]. In contrast with the nucleotide-replacing inhibitors, the antibacterial activity on the MRSA of our benzamide inhibitor series did not closely follow
the activity of *B. subtilis*. The unexpectedly high MRSA MIC values of 17 (25 μM; PC190723 MIC = 5 μM) and 16 (100 μM) with an aminoethoxy linker are 10-fold larger than their MIC values on *B. subtilis* (Figure 7), perhaps due to permeability differences or to the action of staphylococcal efflux pumps. This shows the convenience of early testing for antibacterial activity on pathogenic species. Replacing the difluorobenzamide head of current inhibitors with other effective moieties remains a challenge. Alternately, allosteric FtsZ antibacterial inhibitors may be made more effective by ligand extensions predictively passing the pore to bind into the small pocket [78] and beyond, into the larger unfilled cavity between FtsZ domains (Figure 6), together with the optimization of their pharmacological properties.

Figure 7. Correlation of antimicrobial activity (MIC) with binding affinity (*K*<sub>D</sub>) of inhibitor binding to the BsFtsZ benzamide site at the interdomain cleft. Data from representative analog compounds are shown together with their chemical structures. Filled circles, *B. subtilis* MIC; dotted circles, methicillin-resistant *S. aureus* MIC. The dashed lines are the 95% predictive intervals for *B. subtilis* MIC vs *K*<sub>D</sub> values of 15 analogs [16].

5. Proposed Experimental Strategies for FtsZ Inhibitor Screening and Characterization

Following the development of FP binding and bacterial phenotype approaches during our search for FtsZ inhibitors, we would like to suggest two streamlined screening approaches for the identification and characterization of selective FtsZ inhibitors:

(A) To screen for FtsZ inhibitors, we currently favor the following workflow:

1. Phenotypic screening putative FtsZ inhibitors (1–10<sup>2</sup> compounds) of *B. subtilis* and *E. coli* (cell filamentation), with the determination of the division inhibitory concentration (MDC) and membrane integrity.

   For HTS of large compound libraries, robotized phenotypic screening of *B. subtilis* or *E. coli* plate cultures, based on automatic image analysis of bacterial filamentation [79] may be implemented, if available.

2. FP determination of binding affinity (*K*<sub>D</sub>) to the nucleotide and allosteric binding sites of FtsZ and compound solubility limit determination.

3. Antibacterial activity (MIC) on pathogens and frequency of resistance (FOR) mutations of susceptible pathogens.

(B) As an alternate HTS route, we suggest a binding site-directed approach starting from a robotized FP plate screen of compound binding to the FtsZ nucleotide or allosteric sites, and compound solubility limit determination, followed by phenotypic analysis.
on *B. subtilis* and *E. coli* (filamentation), the determination of MDC and membrane integrity, and a final assessment of MIC and FOR in pathogens.

Each screening approach should be complemented by more detailed cytological profiling, selectivity, and biochemical and structural studies of selected hits, to be followed by the design of optimized analogs and their characterization. At this stage, pharmacokinetic studies should also be considered as a critical feature for the development of viable efficacious antibacterial drugs.

### 6. Outlook for Other Methods, VS, and Machine Learning Approaches for Selecting Potential Antibacterial FtsZ Inhibitors

Conventional HTS of natural extract and compound libraries for FtsZ inhibitors have historically yielded a few molecules that were not further developed (for example, viriditoxin [80], PC170942 [81]). Further screening of smaller focused and drug repurposing libraries still appears worthwhile. Larger areas of the chemical space currently available remain to be explored in natural sources and synthetic libraries, employing cell-based, target-binding, and computer-aided screens, in order to discover new FtsZ inhibitor chemotypes.

New cellular screens may be devised, for example, yeast cells expressing bacterial FtsZ-GFP fusions have been recently employed to monitor the effects of inhibitors on FtsZ assembly while assessing their toxicity of eukaryotic cells in a single-step microscopy screen. This approach [75] has revealed how PC190723 can bind into the interdomain cleft of FtsZ from Gram-negative *Helicobacter pylori*, as opposed to the case of EcFtsZ.

From a structural perspective, both the FtsZ nucleotide and the allosteric binding sites show annex cavities available for fragment binding (Figures 1 and 6). This suggests the possibility of applying HT X-ray crystallographic screening of fragment libraries and further structure-guided fragment-based approaches [82] to develop new FtsZ inhibitors. Several NMR methods may be also applied to search for FtsZ-binding fragments, including ligand-based fluorine NMR screening [83].

Virtual screening (VS) may be employed to expand the chemical diversity of potential inhibitors, as well as to narrow down experimental inhibitor screening. However, early docking-based VS campaigns for FtsZ inhibitors were frequently met by a lack of success, or produced compounds that were not further developed [for example, Refs. [54,84,85]]. More recently, we selected a series of 41 compounds by flexible docking VS with ICM [86] of 4 million compounds from the Zinc library [87,88] against the BsFtsZ nucleotide-binding site. These were experimentally selected for specific binding in place of mant-GTP and the inhibiting division of *B. subtilis* cells, followed by studying 43 well-docking chemically similar compounds from MolPort and Mcule commercial libraries [89]. However, the more active compounds also impaired the membrane potential [unpublished results], which hampered further optimization and underscores the importance of testing membrane integrity. We also screened the Zinc library against an ensemble of seven open conformations of the SaFtsZ PC190723 binding site, selecting seven benzamide and thirty-eight non-benzamide compounds. Among them, five compounds were weak competitors of the specific fluorescent probe 8 for binding to BsFtsZ polymers, but only one benzamide derivative was a weak inhibitor of *B. subtilis* cell division [90]. Complementary to structure-based VS, the chemical features of well-characterized FtsZ small molecule ligands could be employed in ligand-based VS for FtsZ inhibitors.

Artificial intelligence (AI), employing a neural network trained with experimental *E. coli* growth inhibition screening, has been successfully employed to discover new antibiotics [91]. AI drug discovery companies are producing a growing number of discovery and preclinical stage compounds in several therapeutic areas, some of which are in clinical trials [92]. Currently, AI-enabled VS permits screen ultra-large, billion-molecule chemical libraries, combining structural docking with ligand-based prediction [93]. It is thus conceivable that FtsZ-focused AI approaches employing learning data sets from new cell-based and target binding screens, combined with available FtsZ-ligand complex structures and ADMET scoring, may yield new antibacterial FtsZ inhibitors.
7. Conclusions

Three decades after the discovery of the central role of FtsZ in bacterial division [4], FtsZ remains a challenging target for discovering new antibiotics. It may be asked whether the profound basic knowledge of the FtsZ structural assembly mechanism and function that has been achieved can be effectively employed to discover new antibiotics. To further address this problem, we propose combining target-specific screening methods such as those reviewed here with a knowledge-based design and new AI approaches in the quest for FtsZ-targeting antibiotics.

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