Novel insights into the transcriptional regulation of cell division in *Corynebacterium glutamicum*

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Abbreviations

®	registered trademark
С°	degree Celsius
μ	growth rate
μg	microgram
μL	microliter
А	nucleobase adenine
ag	attogram
ATCC	American Type Culture Collection
BHI(S)	Brain Heart Infusion (+ Sorbitol)
bp	base pairs
C-	carboxyl-terminal end
С	nucleobase cytosine
CA	California
CDC	Centers for Disease Control and Prevention
CGP3	Corynebacterium glutamicum prophage 3
CGXII	minimal medium for Corynebacterium glutamicum
ChAP-Seq	chromatin affinity purification with subsequent sequencing
ChIP	chromatin immunoprecipitation
Cm ^R	chloramphenicol resistance
CO ₂	carbon dioxide
CoA	coenzyme A
D	dexter, D-configuration (e.g. of an amino acid)
Da	Dalton
dcw	division cell wall
DNA	deoxyribonucleic acid
DNase	desoxyribonuclease
DSMZ	Deutsche Sammlung von Mikroorganismen und Zellkulturen
DTT	dithiothreitol
e.g.	exempli gratia, for example
EDTA	ethylenediaminetetraacetic acid
EMSA	electrophoretic mobility shift assay
et al.	et alii
etc.	et cetera
eYFP	enhanced yellow fluorescent protein
FHA	forkhead-associated
g	standard gravity, 9.80665 m/s²

G	nucleobase guanine
GDH	glutamate dehydrogenase
GmbH	Gesellschaft mit beschränkter Haftung
GOGAT	glutamate-2-oxoglutarate aminotransferase
GS	glutamate synthase
HPLC	high performance liquid chromatography
НТН	helix-turn-helix
IL	Illinois
Inc.	Incorporation
IPTG	isopropyl β -D-1-thiogalactopyranoside
k	kilo
Kan ^R	kanamycin resistance
L	<i>laevus</i> , L-configuration (e.g. of an amino acid)
L	liter
LB	lysogeny broth
Ltd.	private company limited by shares
MALDI	matrix-assisted laser desorption/ionization
MEME	Multiple Em for Motif Elicitation
mg	milligram
mL	milliliter
mM	millimolar
mRNA	messenger RNA
MS	mass spectrometry
MSG	monosodium glutamate
N-	amino-terminal end
NAD ⁺ /NADH	nicotinamide adenine dinucleotide, oxidized/reduced
NADP ⁺ /NADPH	nicotinamide adenine dinucleotide phosphate, oxidized/reduced
NCBI	National Ceter for Biotechnology Information
ng	nanogram
nm	nanometer
OD ₆₀₀	optical density at 600 nm
ODHC	2-oxoglutarate dehydrogenase complex
ori	origin of replication
PAGE	polyacrylamide gel electrophoresis
PBS	phosphate-buffered saline
PCR	polymerase chain reaction
pg	picogram

рН	potential of hydrogen
PhD	philosophiae doctor
pl	isoelectric point
pmol	picomol
psi	pound-force per square inch
qPCR	quantitative PCR
RBS	ribosome binding site
RNA	ribonucleic acid
rpm	revolutions per minute
SDS	sodium dodecyl sulfate
SNP	single nucleotide polymorphism
т	nucleobase thymine
t	ton
ТСА	trichloroacetic acid
TCA cycle	tricarboxylic acid cycle
TE	transposable element
TEV	tobacco etch virus
ТМ	Trademark
ToF	time of flight
TSS	transcriptional start site
UK	United Kingdom
USA	United States of America
V	Volt
v/v	volume per volume
w/v	weight per volume
WHO	World Health Organization
WT	wild type
Δ	delta/deletion
ΔΤ	temperature difference

Abbreviations not included in this section are according to international standards, as for example listed in the author guidelines of the American Society for Biochemistry and Molecular Biology.

1. Summary

1.1. English Summary

In the first part of this doctoral thesis the transcriptional regulation of the *odhl* gene (cg1630) of *Corynebacterium glutamicum* was analyzed. Odhl in its unphoshorylated state functions as inhibitor of the 2-oxoglutarate dehydrogenase complex (ODHC) by binding to the OdhA subunit. Phosphorylation of Odhl by serine/threonine protein kinases abolishes this effect. Inhibition of ODHC activity by Odhl was shown to be crucial for overproduction and secretion of L-glutamate, which is used as a flavour enhancer. Since downstream of *odhl* two genes presumably encoding transcriptional regulators (cg1631 and cg1633) are located, it was speculated that these could be involved in transcriptional regulation of *odhl*. However, transcriptome analysis of deletion mutants lacking cg1631 or cg1633 and DNA affinity chromatography with the *odhl* promoter did not support this hypothesis. Furthermore, no other potential transcriptional regulators of *odhl*.

The second part of this thesis addresses the regulation of cytokinesis in C. glutamicum. In contrast to e.g. Escherichia coli and Bacillus subtilis, knowledge about regulators of cytokinesis in Actinobacteria is very limited. In this study, the so far uncharacterized Cg1631 protein was discovered to be a transcriptional regulator of the *ftsZ* gene in *C. glutamicum* encoding the key player of bacterial cell division. Therefore, Cq1631 was named FtsR, standing for FtsZ regulator. Both deletion and overexpression of *ftsR* caused growth defects and an altered cell morphology, emphasizing an important function of FtsR in cell division or cell wall synthesis. The wild-type phenotype could be restored by plasmid-based complementation. Chromatin affinity purification with subsequent next generation sequencing (ChAP-Seg) identified a region in the *ftsZ* promoter as a major FtsR binding site, but revealed also additional potential target genes. With the ChAP-Seg results a putative DNA-binding motif could be identified for FtsR. Transcriptional activation of *ftsZ* expression by FtsR was underlined by DNA microarray experiments, electrophoretic mobility shift assays (EMSAs), and reporter gene studies. Analysis of strains expressing *ftsZ* under control of the gluconate-inducible *gntK* promoter revealed that the phenotype of the $\Delta ftsR$ mutant is not solely caused by reduced ftsZexpression but involves additional factors. In summary, FtsR was identified as the first transcriptional regulator of *ftsZ* in *C. glutamicum*. Furthermore, since FtsR and its DNA-binding site in the promoter region of *ftsZ* are highly conserved in Actinobacteria, it can be assumed that this regulatory mechanism is also relevant for the control of cell division in related Actinobacteria. This makes FtsR a promising target for the development of new antimicrobial drugs against pathogenic relatives of *C. glutamicum*.

1.2. Deutsche Zusammenfassung

Im ersten Teil dieser Doktorarbeit wurde die Transkriptionsregulation des *odhl*-Gens (cg1630) von *Corynebacterium glutamicum* analysiert. Odhl wirkt in seinem nichtphosphorylierten Zustand als Inhibitor des 2-Oxoglutarat-Dehydrogenase-Komplexes (ODHC) durch Bindung an die OdhA-Untereinheit. Die Phosphorylierung von Odhl durch Serin/Threonin-Proteinkinasen hebt diesen Effekt auf. Die Hemmung der ODHC-Aktivität durch Odhl ist entscheidend für die Überproduktion und Sekretion von L-Glutamat, das als Geschmacksverstärker verwendet wird. Da sich stromabwärts von *odhl* zwei Gene befinden, die vermutlich für Transkriptionsregulatoren (cg1631 und cg1633) kodieren, wurde spekuliert, dass diese an der Transkriptionsregulation von *odhl* beteiligt sein könnten. Die Transkriptomanalyse der Deletionsmutanten Δ cg1631 und Δ cg1633 sowie DNA-Affinitätschromatographie mit dem *odhl*-Promotor stützten diese Hypothese jedoch nicht. Darüber hinaus konnten keine anderen potenziellen Transkriptionsregulatoren von *odhl* identifiziert werden. Daher gibt es derzeit keine Hinweise auf eine Transkriptionsregulation von *odhl*.

Der zweite Teil dieser Arbeit befasst sich mit der Regulation der Zytokinese bei C. glutamicum. Im Gegensatz zu z.B. Escherichia coli und Bacillus subtilis ist das Wissen über Regulatoren der Zytokinese bei Actinobakterien sehr begrenzt. In dieser Arbeit wurde entdeckt, dass das bisher nicht charakterisierte Cg1631-Protein ein Transkriptionsregulator des ftsZ-Gens in C. glutamicum ist, das den Schlüsselakteur der bakteriellen Zellteilung codiert. Daher wurde Cg1631 als FtsR bezeichnet und steht für FtsZ-Regulator. Sowohl Deletion als auch Überexpression von ftsR verursachten Wachstumsdefekte und eine veränderte Zellmorphologie, was auf eine Funktion von FtsR bei Zellteilung oder Zellwandsynthese hindeutet. Der Wildtyp-Phänotyp konnte durch plasmidbasierte Komplementation der $\Delta ftsR$ -Mutante wiederhergestellt werden. Chromatin-Affinitätsreinigung mit anschließender Sequenzierung (ChAP-Seq) bestätigte die Bindung von FtsR an den ftsZ-Promotor und identifizierte weitere potenzielle FtsR-Zielgene. Mit den ChAP-Seq-Daten konnte ein mutmaßliches DNA-Bindungsmotiv für FtsR identifiziert werden. Die Aktivierung der ftsZ-Expression durch FtsR wurde durch DNA-Microarray-Experimente, elektrophoretische Retardationstests und Reportergenstudien bestätigt. Die Analyse von Stämmen, die ftsZ unter Kontrolle des Gluconat-induzierbaren *gntK*-Promotors exprimierten, ergab, dass der Phänotyp der $\Delta ftsR$ -Mutante nicht nur durch eine verringerte ftsZ-Expression verursacht wird, sondern zusätzliche Faktoren beinhaltet. Zusammenfassend wurde FtsR als erster Transkriptionsregulator von ftsZ in C. glutamicum identifiziert. Da FtsR und seine DNA-Bindungsstelle im ftsZ-Promotor in Actinobakterien hoch konserviert sind, wird angenommen, dass dieser Regulationsmechanismus auch für die Kontrolle der Zellteilung in verwandten Actinobakterien relevant ist. Dies macht FtsR zu einem vielversprechenden Ziel für die Entwicklung neuer antimikrobieller Wirkstoffe gegen pathogene Verwandte von C. glutamicum.

2. Introduction

2.1. Corynebacterium glutamicum

Corynebacterium glutamicum is a non-pathogenic, aerobic, Gram-positive soil bacterium which is used for the large-scale production of several L-amino acids and other industrially relevant compounds (Wendisch *et al.*, 2016, Eggeling & Bott, 2015, Eggeling & Bott, 2005). Moreover, it is a useful model organism for the *Corynebacteriales*, including pathogenic species such as *Corynebacterium diphtheriae* and *Mycobacterium tuberculosis*, which cause the fatal infectious diseases diphtheria and tuberculosis in humans, respectively (WHO, 2015, CDC, 2014, Lawn & Zumla, 2011, Hadfield *et al.*, 2000). Due to its industrial relevance and close relationship to pathogenic species, great efforts have been made in the last decades to unravel the regulatory network of *C. glutamicum*. About 140 transcriptional regulators were identified (Schröder & Tauch, 2010, Brinkrolf *et al.*, 2007) and approximately half of them have been characterized to date. This leads to a better understanding of the biomolecular circuits in this organism, which is of major importance in order to further improve biotechnological production strains or drug discovery processes.

In the past, C. glutamicum, which has been isolated in Japan by Kinoshita et al. (1957), became famous due to its native ability to produce the flavor enhancer monosodium glutamate (MSG). Originally, MSG was discovered in 1908 by Professor Kikunae Ikeda, as he isolated it from the seaweed Laminaria japonica because he wanted to unravel what constitutes the unique taste of Japanese seaweed broth (Lindemann et al., 2002). Generally, Ikeda had the vision that the findings of his research could result in a commercial application, for example in a seasoning that would help improve human nutrition and therefore lead to a longer life expectancy of the Japanese population (Sano, 2009). And indeed, it has been shown that MSG and its umami taste have various positive effects in relation to improving nutrition and health (Mouritsen, 2012). Back then, MSG has been produced by hydrolysis of plant proteins using hydrochloric acid or later on by direct chemical synthesis. The discovery and characterization of C. glutamicum opened up the possibility of bacterial MSG synthesis, which had the advantages of enantiopure production of the natural L-form of glutamic acid and to satisfy the increasing demand while reducing production costs and relieving the environment (Sano, 2009). Today, C. glutamicum is an important industrial amino acid producer which is used to produce amongst others several million tons of the above-mentioned flavor enhancer L-glutamate and the feed additive L-lysine (Becker & Wittmann, 2020).

By now, there are several conditions known which trigger glutamate overproduction and secretion: Biotin limitation (Shiio *et al.*, 1962), addition of detergents like Tween-40 or Tween-60 (Duperray *et al.*, 1992, Takinami *et al.*, 1966), addition of β -lactam antibiotics like penicillin G (Nunheimer *et al.*, 1970), or inhibitors of cell wall synthesis like ethambutol

(Radmacher *et al.*, 2005), glycerol, or fatty acid auxotrophy (Kimura *et al.*, 1997, Nakao *et al.*, 1972, Kanzaki *et al.*, 1967, Okazaki *et al.*, 1967), and high temperature cultivation of temperature-sensitive strains (Delaunay *et al.*, 1999, Momose & Takagi, 1978), which are all thought to influence cell wall composition or membrane permeability, but the actual molecular mechanism is not entirely unraveled to date (Kimura, 2002, Eggeling *et al.*, 2001). Indeed, a transporter involved in glutamate secretion, named YggB (Cg1434), has been identified (Nakamura *et al.*, 2007) and metabolic flux analyses confirmed a strongly reduced activity of the 2-oxoglutarate dehydrogenase complex (ODHC) during cultivation of *C. glutamicum* under producing *versus* non-producing conditions, indicating that regulation of the ODHC plays an important role in glutamate production (Kataoka *et al.*, 2006, Shirai *et al.*, 2005, Shimizu *et al.*, 2003, Kawahara *et al.*, 1997).

2.2. 2-Oxoglutarate dehydrogenase and its regulation in *C. glutamicum*

ODHC is a multimeric enzyme composed of three subunits that catalyzes the conversion of 2-oxoglutarate, NAD⁺, and coenzyme A to succinyl-CoA, NADH + H⁺, and CO₂ in the tricarboxylic acid (TCA) cycle (Usuda et al., 1996). The first subunit E1 is the OdhA protein encoded by the gene cg1280, harboring the actual 2-oxoglutarate dehydrogenase activity. The E2 and E3 subunits of the ODHC are called AceF (Cg2421) and LpdA (Cg0790) and encode a dihydrolipoamide acetyltransferase and a dihydrolipoamide dehydrogenase, respectively (Ikeda & Nakagawa, 2003, Kalinowski et al., 2003, Schwinde et al., 2001). The substrate 2-oxoglutarate marks an important branch point in the metabolism of C. glutamicum. Besides the conversion to succinyl-CoA by the ODHC in the TCA cycle it can also be reductively aminated to glutamate by the glutamate dehydrogenase Gdh (Cg2280), which is mainly active under nitrogen excess (Börmann et al., 1992). Glutamate is converted to glutamine by the glutamine synthetase GInA (Cg2429), which serves together with the glutamate-2-oxoglutarate aminotransferase GOGAT (encoded by gltBD, cg0229, and cg0230) for the assimilation of ammonium under nitrogen-limiting conditions (Jakoby et al., 1997). The 2-oxoglutarate dehydrogenase reaction and the nitrogen metabolism in C. glutamicum are schematically illustrated in Figure 1.



Figure 1: Schematic illustration of the 2-oxoglutarate dehydrogenase reaction and of the nitrogen metabolism in *C. glutamicum***.** Ammonium assimilation is mediated by the reactions carried out by the glutamate dehydrogenase (GDH) and glutamine synthetase (GS)/glutamate-2-oxoglutarate aminotransferase (GOGAT), whereas glutamine catabolism involves the glutaminase K (GIsK) and the glutamate-oxaloacetate transaminase reaction.

Because the GDH shows a 30 to 70 times lower affinity to the substrate 2-oxoglutarate compared to the ODHC (Shiio & Ozaki, 1970), inhibition of ODHC activity is necessary for efficient glutamate production. How this occurs has been unraveled by the discovery of a novel kind of regulation of the ODHC, which results in an increased flux of 2-oxoglutarate to glutamate (Bott, 2007). It was found out that the 15 kDa small inhibitor protein Odhl (Cg1630) is involved in the regulation of the ODHC, which in turn is regulated by a complex signalling cascade. The findings obtained in previous studies so far led to the assumption of the following mechanism, which is schematically illustrated in Figure 2: in the presence of L-glutamine as sole carbon and nitrogen source, it is bound by the membrane-associated lipoprotein GInH (Cg3045). The signal is then transmitted from GlnH via protein-protein interaction to the integral membrane protein GInX (Cg3044), consisting of four transmembrane helices with the N- and C-terminus located in the cytoplasm. GInX activates hereupon the soluble serine/threonine protein kinase PknG (Cg3046). Hence, PknG dissociates from the cytoplasmic membrane and phosphorylates the OdhA inhibitor protein OdhI at its threonine-14 residue. In its phosphorylated state, Odhl is inactive and not able to inhibit the ODHC, so that the complex is present in its active state. Under these conditions, a normal flux of glutamine through the TCA cycle towards succinyl-CoA occurs. The loss of interaction of phosphorylated Odhl with the ODHC is very likely caused by a conformational change which is triggered by the phosphorylation through PknG. Odhl consists of a non-folded N-terminal domain with two phosphorylation sites (Thr14 and Thr15) and a C-terminal forkheadassociated (FHA) domain. Phosphorylation through PknG at the threonine-14 residue causes a conformational change of the non-folded N-terminus which in turn can be bound by the FHA domain. This auto-inhibition leads to an inactive Odhl protein.



Figure 2: Model of the GlnX-GlnH-PknG-Odhl signal transduction cascade in *C. glutamicum*. Illustrated is the model of the GlnX-GlnH-PknG-Odhl signal transduction cascade in *C. glutamicum* as proposed based on the previous findings. A detailed description of the mechanism can be found in the text.

In the absence of the proteins of the cascade, the signal can neither be detected nor transmitted and consequently, Odhl can no longer be phosphorylated by the kinase PknG. In its unphosphorylated state, the inhibitor protein binds to the OdhA subunit of the ODHC, inhibiting its activity, which leads to a reduced conversion of 2-oxoglutarate to succinyl-CoA in the TCA cycle and an increased flux towards glutamate. Due to dephosphorylation of Odhl by the phospho-serine/threonine protein phosphatase Ppp (Cg0062), this process is reversible. The proposed model is based on the findings of Bosco (2011), Krawczyk *et al.* (2010), Schultz *et al.* (2009), Schultz *et al.* (2007), Gebel (2006), and Niebisch *et al.* (2006).

2.3. Cell division in C. glutamicum

Bacterial reproduction is usually characterized by cellular growth followed by binary fission of a mother cell into two daughter cells. During this process, the bacteria need to coordinate several distinct processes such as DNA replication, biogenesis of the new cell wall, and the division process itself. Although the overall process is similar, several different concepts have evolved in distinct bacterial groups. *Escherichia coli* and *Bacillus subtilis* represent two of the best studied species in this respect (Du & Lutkenhaus, 2017, Hajduk *et al.*, 2016).

Corynebacteria are rod-shaped with frequently engrossed cell poles during exponential growth, leading to the typical "club"-shaped morphology by which the name of this genus is

inspired (Cure & Keddie, 1973). Like for almost all other bacteria, the highly conserved FtsZ protein is also the key player of cell division in *C. glutamicum* (Donovan & Bramkamp, 2014, Adams & Errington, 2009). It is essential and the first protein that moves to the future division site, recruiting other proteins involved in this process (Margolin, 2005). Several FtsZ proteins together form the so-called Z-ring, which in rod-shaped bacteria is located in the middle of the cell, and constriction of this ring leads to a formation of two daughter cells (Adams & Errington, 2009, Margolin, 2005) (see Figure 3).



Figure 3: Assembly and disassembly of FtsZ and the Z-ring. On the left, a newborn cell with FtsZ proteins (blue) arranged in a spiral pattern is depicted, as known for *E. coli*. The cell in the middle already shows a formed Z-ring and the recruitment of other proteins of the divisome (violet, green, and orange). On the right site, a dividing cell with a contracting Z-ring is illustrated. Picture inspired by Margolin (2005).

Interestingly, C. glutamicum - as well as other Actinobacteria - is lacking homologs for several important cell division-related genes described for E. coli or B. subtilis (Table 1). Cell division in C. glutamicum is for example independent of the actin homolog MreB, which is essential for elongation of the lateral cell wall in E. coli, Caulobacter crescentus, and B. subtilis (Kruse et al., 2005, Figge et al., 2004, Jones et al., 2001). In contrast to these species, C. glutamicum inserts new cell wall material at the poles, representing an apical elongation mechanism (Daniel & Errington, 2003). Besides mreB, also other typically essential cell division genes are missing, such as *ftsL* and *ftsN* (Letek *et al.*, 2008). Most of the genes required for cell division and peptidoglycan synthesis are located in the so-called division cell wall (dcw) cluster, which is highly conserved throughout the bacterial kingdom (Mingorance et al., 2004). Several studies of this cluster in Actinobacteria revealed that its arrangement in Corynebacteria and Mycobacteria clearly differs from many other bacteria (Tamames et al., 2001). However, the most striking difference might be the complete absence of homologs of other known spatial and temporal, positive and negative regulators of cell division, as for example ftsA, ezrA, slmA, sulA, zipA, zapA, min, and noc are missing (Donovan & Bramkamp, 2014, Letek et al., 2007). FtsA is a positive regulator of FtsZ assembly and supports membrane association of the Z-ring (Pichoff & Lutkenhaus, 2005, Beall & Lutkenhaus, 1992). The integral membrane protein EzrA negatively influences Z-ring assembly and is involved in mid-cell localisation of the Z-ring (Haeusser et al., 2004, Levin et al., 1999). SIMA and NocA are nucleoid occlusion effector proteins, known to inhibit assembly of the Z-ring over the nucleoid (Bernhardt & de Boer, 2003, Wu & Errington, 2004). SulA also inhibits Z-ring formation and is induced upon DNA damage

(Bi & Lutkenhaus, 1993, Huisman *et al.*, 1984). ZipA and ZapA are positive regulators of Z-ring assembly (Slayden *et al.*, 2006, Gueiros-Filho & Losick, 2002, Hale & de Boer, 1997), and the Min system consists of several proteins preventing division occurring at the cell poles (Bramkamp *et al.*, 2008, Raskin & de Boer, 1999, Levin *et al.*, 1992, de Boer *et al.*, 1989). Taking all these differences into account, it is obvious that in Actinobacteria cell division and its regulation must differ profoundly from the processes described for other well-studied rod-shaped bacteria like *E. coli*, and *B. subtilis* (Letek *et al.*, 2007, Flardh, 2003, Ramos *et al.*, 2003).

Table 1: Putative cell division genes of *E. coli, B. subtilis, C. glutamicum,* and *M. tuberculosis.* Known positive regulators (green), negative regulators (red), and essential cell division genes (blue) which are missing in *C. glutamicum* are written in bold letters. The table is based on Letek *et al.* (2008) and Donovan and Bramkamp (2014). For an overview of the predicted protein functions, see Donovan and Bramkamp (2014).

gene name	E. coli	B. subtilis	C. glutamicum	M. tuberculosis
amiC	b2817	BSU09420 (<i>lytE</i>)	cg3424	rv3915
		BSU09370 (<i>lytF</i>)		
clpX	b0438	BSU28220	cg2620	rv2457c
crgA	?	?	cg0055	rv0011c
cwsA	?	?	?	rv008c
divIVA	?	BSU15420	cg2361	rv2154c (<i>wag31</i>)
divS	b0958 (<i>sulA</i>)	BSU17860 (<i>yneA</i>)	cg2133	rv2719 (<i>chiZ</i>)
ezrA	?	BSU29610	?	?
ftsA	b0094	BSU15280	?	?
ftsB (divIC)	b2748	?	cg1112	rv1024
ftsE	b3463	BSU35260	cg0914	rv3102c
ftsl	b0084	BSU15170	cg2375	rv2163c
ftsK	b0890	BSU29800 (<i>ytpT</i>) BSU16800 (<i>spoIIIE</i>)	cg2158	rv2748c
ftsL	b0083	BSU15150	?	?
ftsN	b3933	?	?	?
ftsQ (divB)	b0093	BSU15240	cg2367	rv2151c
ftsW	b0089	BSU14850	cg2370	rv2154c
		BSU15210 (spove)		
ftsX	b3462	BSU35250	cg0915	rv3101c
ftsZ	b0095	BSU15290	cg2366	rv2150c

(Continued)

Table 1. Continued.

gene name	E. coli	B. subtilis	C. glutamicum	M. tuberculosis
ftsX	b3462	BSU35250	cg0915	rv3101c
ftsZ	b0095	BSU15290	cg2366	rv2150c
minC	b1176	BSU28000	?	?
minD	b1175	BSU27990	?	?
minJ	?	BSU35220	?	?
minE	b1174	?	?	?
mreB	b3251	BSU14470 (<i>mreBH</i>) BSU28030 (<i>mreB</i>)	?	?
		BSU36410 (<i>mbl</i>)		
mreC	b3250	BSU28020	?	?
mreD	b3249	BSU28010	?	?
пос	?	BSU40990	?	?
pldP	?	?	cg1610	rv1708
rodA	b0634	BSU38120	cg0061	rv0017c
sepF	?	BSU15390	cg2363	rv2147c
sImA	b3641	?	?	?
sulA	b0958	?	?	?
ugtP	?	BSU21920	?	?
zapA	b2910	?	?	?
zapB	b3928	?	?	?
zapC	b0946	?	?	?
zapD	b0102	?	?	?
zipA	b2412	?	?	?

To date, *C. glutamicum* is an established model organism for industrial biotechnology and for pathogenic relatives, but we have a relatively narrow understanding about the exact regulation of its cytokinesis (Donovan & Bramkamp, 2014). However, it is known that accurate expression of FtsZ is critical for normal growth of *C. glutamicum* and *M. tuberculosis* and small variations lead to severe morphological changes and affect cell viability, suggesting that the intracellular levels of FtsZ are critical for proper cell division in these organisms (Letek *et al.*, 2007, Ramos *et al.*, 2005, Dziadek *et al.*, 2003). Thus, FtsZ levels must be subject to a tight regulation, which is supported by different studies of the *ftsZ* promoter region, showing a very complex

transcription of *ftsZ* in various bacteria (Letek *et al.*, 2007, Roy & Ajitkumar, 2005, Flardh *et al.*, 2000, Flardh *et al.*, 1997, Gonzy-Treboul *et al.*, 1992). A complex of the DNA-binding proteins WhiA (Cg1792) and WhcD (Cg0850) was reported to bind to the *ftsZ* promoter region (Lee *et al.*, 2018), however, the effect of this interaction on *ftsZ* expression remains to be elucidated. Previously, it was shown that the FtsZ protein can be phosphorylated *in vitro* by the serine/threonine protein kinases PknA, PknB, and PknL and is an *in vivo* substrate of the phospho-serine/threonine protein phosphatase Ppp (Schultz *et al.*, 2009), suggesting that the properties and activities of FtsZ can also be influenced by posttranslational regulation (Donovan & Bramkamp, 2014). However, a regulatory protein involved in direct control of *ftsZ* transcription has only been described for *Caulobacter crescentus* (Kelly *et al.*, 1998). For the *Corynebacteriales* order, the regulation of cell division processes remains largely undiscovered.

2.4. Aim of this work

Originally, the focus of this thesis lay on studying the role of GlnX, GlnH, PknG, and Odhl in C. glutamicum in order to attain a deeper understanding of the signal transduction cascade which consists of these proteins. More specifically, one major objective was to investigate whether the inhibitory protein of the ODHC is potentially regulated on the transcriptional level. This seemed sensible because odhl (cg1630) is located upstream of the two genes (cg1631 and cg1633), which are annotated to encode transcriptional regulators. However, the experiments performed in this direction (DNA affinity chromatography with the odhl promoter, DNA microarrays) did not provide any evidence to support transcriptional regulation of odhl. Nevertheless, since studying the GInH-GInX-PknG-OdhI-ODHC signal transduction cascade was the initial objective of this work, a part of this introduction and the first chapters of the results section are related to this topic. In these initial studies, deletion mutants of the genes cq1631, cq1632, and cq1633 were constructed and analyzed. Thereby it was observed that deletion of cg1631 led to a severe growth defect and drastic morphological changes. Because there is hitherto no transcriptional regulator of cell division known for C. glutamicum, this phenotype was highly interesting and let us redefine the aims of this thesis towards the functional analysis of the previously uncharacterized protein Cg1631. In this context, we searched for the target genes of Cq1631 by a variety of complementary approaches and identified *ftsZ* to be activated by Cg1631. Therefore, Cg1631 was renamed FtsR, standing for ftsZ regulator. We show that FtsR is critical for normal growth and cell morphology of C. glutamicum, suggesting an important role of this protein in the regulation of cell division.

3. Materials and Methods

3.1. Bacterial strains, plasmids, and growth conditions

Bacterial strains and plasmids used in this study are listed in Table 2. The *C. glutamicum* type strain ATCC13032 or its prophage-free variant *C. glutamicum* MB001 were used as wild type or reference strain, as indicated. The parental strain ATCC13032 (DSM No. 20300) was obtained from the Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSMZ, Braunschweig, Germany). The prophage-free strain MB001 was derived from an in-house stock but can also be obtained from the DSMZ (DSM No. 102070). First experiments were performed with ATCC13032, but to be able to discriminate between effects primarily caused by *ftsR* deletion or secondarily emerged due to phage-related effects, we later switched to the *C. glutamicum* MB001 strain. Deletion of the prophages did not lead to any negative effects on the physiology of *C. glutamicum* MB001 and due to its reduced complexity it is a useful platform organism for basic research (Baumgart *et al.*, 2013b).

For growth experiments, *C. glutamicum* was pre-cultivated for six to eight hours at 30 °C and 170 rpm in 5 mL BHI medium (BD BactoTM Brain Heart Infusion, Becton Dickinson and Company, Heidelberg, Germany). The cells were harvested by centrifugation, washed with phosphate-buffered saline (PBS, 137 mM NaCl, 2.7 mM KCl, 4.3 mM Na₂HPO₄, 1.4 mM KH₂PO₄, pH 7.3) and used as inoculum for a second pre-culture in 20 mL CGXII minimal medium (Keilhauer *et al.*, 1993) supplemented with 3,4-dihydroxybenzoate (30 mg/L) as iron chelator and, if not stated otherwise, 2% (w/v) glucose as carbon source. This second pre-culture was incubated overnight at 30 °C and 120 rpm. After harvesting by centrifugation and washing with PBS, the cells were used for inoculation of the main culture, using CGXII medium with 3,4-dihydroxybenzoate (30 mg/L) and 2% (w/v) glucose as carbon source, if not stated otherwise.

Growth experiments were either performed in 100 mL shake flasks with 20 mL medium (initial optical density at 600 nm (OD₆₀₀) of 1.0) that were shaken at 30 °C and 120 rpm or in 48-well FlowerPlates[®] (m2p-labs GmbH, Baesweiler, Germany) containing a final culture volume of 800 μ L (initial OD₆₀₀ of 0.5) that were shaken in a BioLector[®] system (m2p-labs GmbH, Baesweiler, Germany) at 30 °C and 1200 rpm. Growth was monitored as cell density by determining either OD₆₀₀ (shake flasks experiments, OD₆₀₀ measured with an Ultrospec 500 Pro UV/Vis spectrophotometer, Amersham Biosciences, Little Chalfont, United Kingdom) or as scattered light at 620 nm in the BioLector[®] (Kensy *et al.*, 2009), which is termed "backscatter" throughout this study.

For growth experiments with promoter exchange strains in which expression of the *ftsZ* gene was under control of the *gntK* promoter, the first pre-culture was supplemented with 0.1% (w/v) and the second pre-culture with 0.01% (w/v) gluconate and 1.99% (w/v) glucose as carbon source. The gluconate concentration of the main culture is indicated for each experiment.

For cloning purposes, *Escherichia coli* DH5 α was used and routinely cultivated at 37 °C in lysogeny broth (LB, (Sambrook & Russell, 2001)). When required, the media were supplemented with 25 µg/mL kanamycin or 10 µg/mL chloramphenicol for *C. glutamicum* or with 50 µg/mL kanamycin or 34 µg/mL chloramphenicol for *E. coli*.

Strain or plasmid	Characteristics	Source or Reference
Bacterial strains		
E. coli		
DH5α	F ⁻ Φ80 <i>dlac</i> ∆(<i>lacZ</i>)M15 ∆(<i>lacZYA-argF</i>) U169 endA1 recA1 hsdR17 (rκ⁻, mκ⁺) deoR thi-1 phoA supE44 λ⁻ gyrA96 relA1; strain used for cloning procedures	Hanahan (1983)
C. glutamicum		
ATCC13032	biotin-auxotrophic wild type.	Kinoshita <i>et al.</i> (1957)
ATCC13032∆ <i>ftsR</i>	ATCC13032 with in-frame deletion of <i>ftsR</i> (cg1631)	this work
ATCC13032∆cg1632	ATCC13032 with in-frame deletion of cg1632	this work
ATCC13032∆cg1633	ATCC13032 with in-frame deletion of cg1633	this work
ATCC13032∆ <i>ramB</i>	ATCC13032 with in-frame deletion of <i>ramB</i> (cg0444)	Gerstmeir <i>et al.</i> (2004)
ATCC13032::ftsZ-venus	ATCC13032 with a chromosomal insertion of pK18 <i>mob-ftsZ-venus</i> at the <i>ftsZ</i> locus; this strain has two chromosomal copies of <i>ftsZ</i> under control of its native promoter, one with and one without the fusion to <i>venus</i>	this work
ATCC13032∆ftsR::ftsZ- venus	ATCC13032 Δ <i>ftsR</i> with a chromosomal insertion of pK18 <i>mob-ftsZ-venus</i> at the <i>ftsZ</i> locus; this strain has two chromosomal copies of <i>ftsZ</i> under control of its native promoter, one with and one without the fusion to <i>venus</i>	this work
MB001	ATCC13032 with in-frame deletions of the prophages CGP1 (cg1507-cg1524), CGP2 (cg1746-cg1752), and CGP3 (cg1890-cg2071)	Baumgart <i>et al.</i> (2013b)
MB001∆ <i>ftsR</i>	MB001 with in-frame deletion of <i>ftsR</i> (cg1631)	this work
MB001::P _{gntK} -ftsZ	MB001 with a chromosomal promoter exchange of the native <i>ftsZ</i> promoter against the gluconate-inducible promoter of <i>gntK</i> (cg2732)	this work
MB001∆ <i>ftsR</i> ∷P _{gntK} -ftsZ	MB001::P _{gntK} -ftsZ with in-frame deletion of ftsR	this work
<i>C. diphtheriae</i> ATCC27010	genomic DNA of this strain was used as PCR template for amplification of the <i>ftsR</i> homolog CDC7B_1201	DSM 44123
<i>M. tuberculosis</i> H37Rv	genomic DNA of this strain was used as PCR template for amplification of the <i>ftsR</i> homolog rv1828	ATCC 25618

Table 2: Bacterial strains	and	plasmids	used in	this	study.
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(Continued)

Table 2. Continued.

Strain or plasmid	Characteristics	Source or reference
Plasmids		
pK18 <i>mob</i>	Kan ^R ; plasmid for insertion of a DNA fragment into the chromosome of <i>C. glutamicum</i> (pK18 <i>ori</i> V _{<i>E.c.</i>} , <i>lacZα</i>)	Schäfer <i>et al.</i> (1994)
pK18 <i>mob-ftsZ-venus</i>	Kan ^R ; pK18 <i>mob</i> derivative for chromosomal insertion of the coding sequence for a fusion protein of FtsZ and the fluorescent protein Venus under control of the native <i>ftsZ</i> promoter	this work
pK19 <i>mobsacB</i>	Kan ^R ; plasmid for allelic exchange in <i>C. glutamicum</i> (pK18 <i>ori</i> V _{E.c} ., sacB, lacZα).	Schäfer <i>et al.</i> (1994)
pK19 <i>mobsacB-∆ftsR</i>	Kan ^R ; pK19 <i>mobsacB</i> derivative for in-frame deletion of <i>ftsR</i> ; contains PCR product covering the fused up- and downstream regions of <i>ftsR</i>	this work
pK19 <i>mobsacB</i> -P _{gntK} - <i>ftsZ</i> Kan ^R ; pK19 <i>mobsacB</i> derivative for construction of MB001::P _{gntK} - <i>ftsZ</i> ; contains a PCR fragment encompassing 691 bp of the upstream region of cg2366 (<i>ftsZ</i>) covering its promoter, followed by a terminator sequence, the <i>gntK</i> (cg2732) promoter and 523 bp of the <i>ftsZ</i> (cg2366) coding region		this work
pK19-P2732- <i>lcpA</i>	Kan ^R ; pK19 <i>mobsacB</i> derivative used as template for the promoter exchange plasmid pK19 <i>mobsacB</i> -P _{gntK} -ftsZ	Baumgart <i>et al.</i> (2016)
pAN6	Kan ^R ; <i>C. glutamicum/E. coli</i> shuttle vector for regulated gene expression using P _{tac} (P _{tac} lacl ^q pBL1 <i>oriV_{Cg}</i> pUC18 <i>oriV_{Ec}</i>)	Frunzke <i>et al.</i> (2008)
pAN6- <i>ftsR</i>	Kan ^R ; pAN6 derivative for expression of <i>ftsR</i> (252 amino acids) under control of P _{tac}	this work
pAN6- <i>ftsR</i> -short	Kan ^R ; pAN6 derivative encoding an N-terminally shortened FtsR protein (224 amino acids) under control of P_{tac}	this work
pAN6- <i>ftsR</i> -Strep	Kan ^R ; pAN6 derivative for expression of <i>ftsR</i> under control of P _{tac} with a C-terminal Strep-tag [®]	this work
pAN6-CDC7B_1201	Kan ^R ; pAN6 derivative for expression of <i>C. diphtheriae</i> CDC7B_1201 (<i>ftsR</i> homolog) under control of P _{tac} .	this work
pAN6-rv1828	Kan ^R ; pAN6 derivative for expression of <i>M. tuberculosis</i> rv1828 (<i>ftsR</i> homolog) under control of P _{tac} .	this work
pEC-XC99E	Cm ^R ; <i>C. glutamicum/E. coli</i> shuttle vector for regulated gene expression using P _{trc} , (P _{trc} , <i>cat</i> _{<i>i</i>} , <i>lacl</i> ^q , <i>rrnB</i> , <i>oriV_{E.c}</i> , <i>per</i> and <i>repA</i> (pGA1) _{C.g.}).	Kirchner and Tauch (2003)
pEC- ftsR	Cm ^R ; pEC-XC99E derivative for expression of <i>ftsR</i> under control of P _{trc} .	this work
pJC1- <i>venus</i> -term	Kan ^R ; pJC1 derivative carrying the Venus coding sequence and additional terminators (pCG1 <i>ori_{Cg}</i> , pACYC177 <i>ori_{Ec}</i>).	Baumgart <i>et al.</i> (2013a)
pJC1-P _{ftsZ} -venus	Kan ^R , pJC1- <i>venus</i> -term derivative carrying the <i>ftsZ</i> promoter controlling expression of <i>venus</i> for promoter activity studies.	this work

Routine methods such as PCR, DNA restriction and ligation, Gibson assembly, and transformation were performed using standard protocols (Sambrook & Russell, 2001, Hanahan, 1983, van der Rest et al., 1999, Gibson, 2011). Phusion Green High Fidelity DNA Polymerase (Thermo Fisher Scientific Inc., Rockford, IL, USA) was used for cloning purposes. For all other PCRs, either the KAPA2G Fast ReadyMix PCR Kit (Kapa Biosystems, Wilmington, USA) or DreamTaq DNA Polymerase (Thermo Fisher Scientific Inc., Rockford, IL, USA) were used. The oligonucleotides used in this study (see Table 3) were purchased from Eurofins Genomics GmbH (Ebersberg, Germany) and DNA sequencing was also performed by this company. The $\Delta ftsR$ mutant of C. glutamicum and the promoter exchange strains carrying a DNA fragment with a transcription terminator sequence and the *gntK* promoter inserted into the chromosome between the native ftsZ promoter and the ftsZ coding region were constructed via a two-step homologous recombination protocol as described previously (Baumgart et al., 2016, Niebisch & Bott, 2001), using plasmids pK19mobsacB- Δ ftsR and pK19mobsacB-P_{antK}ftsZ. The strains ATCC13032::ftsZ-venus and ATCC13032

ftsR::ftsZ-venus were constructed by chromosomal insertion of pK18mob-ftsZ-venus by single homologous recombination. In all tested clones of both strains, the plasmid did not insert into the intergenic region between cg1121 and cg1122, but into the *ftsZ* region. As the insertion plasmid was constructed in such a way that these mutants also carry one native copy of *ftsZ* and one fused to *venus*, these strains were used for microscopy, anyway.

Oligonucleotide	Sequence (5' \rightarrow 3') and properties	Commentary			
DNA affinity purification with P _{odhl} and P _{ftsz}					
AP_PodhI_fw	TGATCAGTCCGTGAGGGAAC				
AP_Podhl_rv_bio	<i>GAGGAGTCGTCGATGTGGAGACC</i> CAGC GCGGAATACTGAGGTG	overlap (in italics) homologous to the biotin-oligo used as reverse primer in a second PCR to create a biotinylated fragment			
AP_PftsZ_fw	CATTAGCTCACCCTCAATGG				
AP_PftsZ_rv_bio	<i>GAGGAGTCGTCGATGTGGAGACC</i> GAGG CCTTCTTCAATCATGC	overlap (in italics) homologous to the biotin-oligo used as reverse primer in a second PCR to create a biotinylated fragment			
biotin_oligo	5'BIO- GAGGAGTCGTCGATGTGGAGACC	biotin-labelled sequence homologous to the overlap of the rv_bio primers for attachment of a biotin-tag to the resulting fragments; biotinylated molecules bind with high affinity to the Streptavidin-coated magnetic beads used for DNA affinity purification.			

 Table 3: Oligonucleotides used in this study for cloning, EMSAs, and DNA affinity purification.

 Restriction sites are underlined. Bold letters represent the overlapping sequences needed for Gibson

 Assembly. If not stated otherwise, *C. glutamicum* genomic DNA was used as template.

Table 3. Continued.

Oligonucleotide	Sequence (5' \rightarrow 3') and properties	Commentary
Construction of pK18mo	b- and pK19 <i>mobsacB</i> derivatives	
M13-fw	CGCCAGGGTTTTCCCAGTCAC	for sequencing of pK18mob-ftsZ-venus
M13-rv	AGCGGATAACAATTTCACACAGGA	and pK19 <i>mobsacb-∆ftsR</i>
Plasmid pK18mob-ftsZ-v	venus for chromosomal insertion of	ftsZ-venus
pK18_IRG-fw-V2	CCTGCAGGTCGACTCTAGAGGTTTACG CAGCACAAGACCCC	for amplification of a fragment covering about 450 bp of the intergenic region
IGR-rv	TCCTCCATAATTAGAGAGCGTAAGGCC C	between cg1121 and cg1122
PftsZ-fw	CGCTCTCTAATTATGGAGGA TGATGGT GACCATGTCATTGACACCG	for amplification of the promoter and the coding region of <i>ftsZ</i>
FtsZ-rv	TCCTCGCCCTTGCTCACCAT CTGGAGG AAGCTGGGTACATCCAG	
venus-fw	ATGGTGAGCAAGGGCGAGGAG	for amplification of the <i>venus</i> gene
pK18_venus-rv-V2	CAGCTATGACCATGATTACG TTACTTG TACAGCTCGTCCATGCC	encoding the fluorescent protein Venus
FtsZ-Seq1	CATTAGCTCACCCTCAATGGTG	for sequencing of pK18 <i>mob-ftsZ-venus</i>
FtsZ-Seq2	GAACCTGTCCATCATGGAAGC	
int-reg-fw	AGCACCTTCGGCAAGAAGTA	test of integration strains for integration
int-reg-rv	CATCGAAGGTGTCGCAAAC	into the intergenic region of cg1121-
ftsQ-rv	AGCAATAACCGCAGGAAGCAC	091122
M13-fw	CGCCAGGGTTTTCCCAGTCAC	
ftsQ-fw	ACAAGGCAGGACTAGCGTGAAC	test of integration strains for integration
ftsZ-downstream-rv	TCGTGAAGACCTTGCGGAC	into the chromosomal <i>ftsZ</i> -region
pK18-IGR-fw	CTTGGTTCGAATATGCAGTTCGG	
eYFP-int_rv	CGACCAGGATGGGCACCAC	
Deletion plasmid pK19m	obsacb-∆ftsR	
D_ftsR_1_fw	GATC <u>GGATCC</u> TCCGCACTCAACATCTA GAC	PCR product contains BamHI site for cloning in BamHI-cut pK19 <i>mobsacb</i>
D_ftsR_2_rv	TGTTTAAGTTTAGTGGATGGGGATGTT GCTGCTACCTGCTGTGAATTAAA	
D_ftsR_3_fw	CCCATCCACTAAACTTAAACATAGAAA AAATGAGTTTTGTTGAACTT	PCR product contains HindIII site for cloning in HindIII-cut pK19 <i>mobsacb</i>
D_ftsR_4_rv	GATC <u>AAGCTT</u> CGTCGCCTGAAGCAGAT TCC	
map_DftsR_fw	TCCGCACTCAACATCTAGAC	for verification of chromosomal <i>ftsR</i>
map_DftsR_rv	CAGGTAAAGCCATCTGGTTC	deletion
Deletion plasmid pK19m	obsacb-∆cg1632	
D_cg1632_1_fw	GATC <u>GGATCC</u> TCAGCCCGGAGAAGTTT CAG	PCR product contains BamHI site for cloning in BamHI-cut pK19 <i>mobsacb</i>
D_cg1632_2_rv	TGTTTAAGTTTAGTGGATGGGTTTTTC TATCAGTATCCAAGCTGCTCGCGAGTT GC	
D_cg1632_3_fw	CCCATCCACTAAACTTAAACAAATGTG ACTAATCACACCCTCAGATTTCAAC	PCR product contains HindIII site for cloning in HindIII-cut pK19 <i>mobsacb</i>
D_ cg1632_4_rv	GATC <u>AAGCTT</u> AGGTCGTTGGTGCCCAT	

Table 3. Continued.

Oligonucleotide	Sequence (5' $ ightarrow$ 3') and properties	Commentary				
Deletion plasmid pK19 <i>mobsacb</i> -∆cg1633						
D_cg1633_1_fw	GATC <u>GGATCC</u> TACATGGCTACCATCAC TAC	PCR product contains BamHI site for cloning in BamHI-cut pK19 <i>mobsacb</i>				
D_ cg1633_2_rv	TGTTTAAGTTTAGTGGATGGGTAGTTT CTCCGATTATTCAGCACCCATGTTTTA TTC					
D_ cg1633_3_fw	CCCATCCACTAAACTTAAACAAAACTC AGCGCAGTAAATCTTCAAGCC	PCR product contains HindIII site for cloning in HindIII-cut pK19 <i>mobsacb</i>				
D_cg1633_4_rv	GATC <u>AAGCTT</u> CAAGCCCTCGACCGACT TGG					
Promoter exchange plas	smid pK19 <i>mobsacb</i> -P _{gntK} -ftsZ					
ftsZ_upstream_fw	CAGGTCGACTCTAGAGGATC GAAGTGC TTCCTGCGGTTAT	for amplification of a fragment upstream of the <i>ftsZ</i> coding region with				
ftsZ_upstream_rv	CACTACCATCGGCGCTAC TGTCGATGT CTCGCCTTTCG	homologies to the pK19 <i>mobsacb</i> - backbone and to the terminator sequence				
term_fw	GTAGCGCCGATGGTAGTG	for amplification of a fragment harboring				
Pcg2732-rv	GTCTTATCCTTTCTTTGGTGGCG	the terminator and the P _{gntK} coding sequence; template: pK19-P2732- cg0847 (Baumgart <i>et al.</i> , 2016)				
PgntK_ftsZ_fw	CACCAAAGAAAGGATAAGACATGACCT CACCGAACAACTA	for amplification of a fragment of the first 523 bp of the <i>ftsZ</i> (cg2366) coding				
ftsZ_start_rv	AAAACGACGGCCAGTGAATTGTCGTTT GGAATAACGATGA	region (including start codon) with homologies to the P _{gntK} sequence and the pK19 <i>mobsacb</i> -backbone				
map_term- PgntK_insertion_fw	TGAGTGCGGAACCAGCTTCG	for verification of insertion of the terminator- P_{gntk} -fragment between P_{ftsZ}				
map_term- PgntK_insertion_rv	ACCATCACCGTGGAGCTGAC	chromosome				
Construction of pAN6 de	erivatives					
pAN6_check_fw	CATCGGAAGCTGTGGTATGG	for sequencing of pAN6 derivatives				
pAN6_check_rv	CCTGGCAGTTCCCTACTCTC					
pAN6-ftsR, pAN6-ftsR-s	hort & pAN6- <i>ftsR</i> -Strep					
pAN6_ftsR_fw	CTGCAGAAGGAGATATACATATGA GTG CACTCCGTAAAAC	PCR product with overlaps for cloning of <i>ftsR</i> in the Ndel/EcoRI-cut pAN6				
pAN6_ftsR_short_fw	CTGCAGAAGGAGATATACATATG TCAA TTGGTGTGGTACT	plasmid; with fw-primer, the <i>ftsR</i> start codon GTG is changed to ATG				
pAN6_ftsR_rv	AAAACGACGGCCAGTGAATTTCAGTAT CCAAGCTGCTCGC					
pAN6_ftsR_Strep_rv	CTGTGGGTGGGACCAGCTAGT GTATCC AAGCTGCTCGCGA	with pAN6_ftsR_fw amplification of a PCR product without stop codon with overlaps for in-frame cloning of <i>ftsR</i> upstream of a Strep-tag [®] coding region for expression of FtsR-Strep in the Ndel/Nhel-cut pAN6 vector				
pAN6-CDC7B_1201						
CDC7B_1201_pAN6_fw	GCCTGCAGAAGGAGATATACATATGAG TGCACTTCCGCAACG	PCR product with overlaps for cloning of the CDC7B_1201 gene (<i>ftsR</i> homolog)				
CDC7B_1201_pAN6_rv	AAAACGACGGCCAGTGAATTTTAACGG TTCAGCTCATCGCGC	IN THE NOEI/ECORI-CUT PAN6 vector; template: <i>C. diphtheriae</i> ATCC27010 genomic DNA				

(Continued)

Table 3. Continued.

Oligonucleotide	Sequence (5' \rightarrow 3') and properties	Commentary
pAN6-rv1828		
rv1828_pAN6_fw	GCCTGCAGAAGGAGATATACATGTGAG CGCACCCGATAGCC	PCR product with overlaps for cloning of the rv1828 gene (<i>ftsR</i> homolog) in the
rv1828_pAN6_rv	AAAACGACGGCCAGTGAATT TCAGCGG TGAAGAACGTCGCGAAC	Ndel/EcoRI-cut pAN6 vector; template: <i>M. tuberculosis</i> H37rv genomic DNA
Construction of pEC-ft	sR	
ftsR_BamHI_fw	GATC <u>GGATCC</u> GCCTGCAGAAGGAGATA TAC	PCR product contains BamHI and Xbal sites for cloning in pEC-XC99E vector
ftsR_Xbal_rv	GATC <u>TCTAGA</u> TCAGTATCCAAGCTGCT CGC	pAN6- <i>ftsR</i> , amplified fragment harbors RBS from pAN6
pEC-check_fw	TAATCATCGGCTCGTATAATGTGTG	for sequencing of pEC derivatives
pEC-check_rv	GCTTCTGCGTTCTGATTTAATCTG	
Construction of pJC1-	P _{ftsZ} -venus	
pJC1_PftsZ_fw	AGCGACGCCGCAGGGGGGATCTTCCTGC GGTTATTGCTGTA	amplification of the <i>ftsZ</i> -promoter region equipped with overlaps homologous to
PftsZ_venus_rv	CTCCTCGCCCTTGCTCACCATTGTCGA TGTCTCGCCTTTCG	the pJC1- <i>venus</i> -term vector backbone cut with SpeI and to the <i>venus</i> coding region
venus_fw	ATGGTGAGCAAGGGCGAGGAG	PCR product with overlap for cloning
venus_pJC1_rv	AAAACGACGGCCAGTACTAGTTACTTG TACAGCTCGTCCATGC	into BamHI-cut pJC1- <i>venus</i> -term; template: pJC1- <i>venus</i> -term
pJC1_check_fw	TGAAGACCGTCAACCAAAGG	for sequencing of pJC1-venus-term
pJC1_check_rv	TACGGCGTTTCACTTCTGAG	derivatives
30-bp oligonucleotides	for Electrophoretic Mobility Shift As	says
ftsZ_oligo	CGCTACCCTCAACCTTTACTTTAGGGT TGT	
ftsZ_oligo_compl	ACAACCCTAAAGTAAAGGTTGAGGGTA GCG	
cg1081_oligo	GAAGCCACATGACATATGTCATGAAAA TTA	
cg1081_oligo_compl	TAATTTTCATGACATATGTCATGTGGC TTC	
Competition-EMSA		
Cy3-ftsZ_prom_rv	Cy3*TCTTGGCGAGGTAGTTGTTC	5'-Cy3-label
ftsZ_prom_rv	TCTTGGCGAGGTAGTTGTTC	
ftsZ_prom250_fw	AGTTCCGGTTCACCCGTTTC	
ftsZ_prom500_fw	AGTGCTTCCTGCGGTTATTG	
ftsZ_prom250up_rv	CTACCCAGCCACTTTAGCGG	
qPCR for determination	n of the copy number of cg0834 and c	:g0840
qPCR_cg0834_fw	TTCGAATTTGCCTGCGGTTG	
qPCR_cg0834_rv	AAACCAACCTCGCATTCACC	
qPCR_cg0840_fw	TCTGACCAACCAGTTGCAAG	
qPCR_cg0840_rv	CCACAGTGCGTTTTTATGGC	
qPCR_recF_fw	GGCTATCTTGCGCATTTGTC	
aPCR recF rv	TCTCGGCCTTGATTAACAGC	

3.3. Fluorescence microscopy

The *C. glutamicum* cells were centrifuged and suspended in PBS containing 100 ng/mL Hoechst 33342 dye to stain DNA and and 300 ng/mL Nile red (Sigma-Aldrich Chemie GmbH, Taufkirchen, Germany) to stain membranes. After 10 minutes of incubation in the dark at room temperature, samples were spotted onto a glass slide covered with a thin agarose layer (1% (w/v) in TAE buffer (40 mM Tris, 1 mM Na₂EDTA, 20 mM glacial acetic acid, pH 8) and analyzed using a Zeiss Axio Imager M2 microscope equipped with a Zeiss AxioCam MRm camera and a Plan-Apochromat 100x/1.40-numerical aperture phase-contrast oil immersion objective and AxioVision 4.8 software (Carl Zeiss Microscopy GmbH, Jena, Germany). Hoechst 33342 fluorescence was visualized with filter set 49 and Nile red with filter set 63 HE (both Carl Zeiss AG). *C. glutamicum* strains producing the FtsZ-Venus fusion protein were directly used for microscopy without further staining.

3.4. Purification of FtsR

FtsR was purified using C. glutamicum ATCC13032 ftsR carrying the plasmid pAN6-FtsR-Strep. The preculture was prepared using a single colony from a fresh BHI agar plate to inoculate 20 mL BHI with 2% (w/v) glucose and 25 µg/mL kanamycin and incubated at 30 °C and 120 rpm for 6-8 hours. The main culture (500 mL of the same medium in a 2 L baffled flask) was inoculated with 1 mL of the preculture. Due to the growth defect of the strain, the culture was incubated overnight at 30 °C and 90 rpm. On the following morning, expression of ftsR was induced by addition of 50 μ M isopropyl β -D-1-thiogalactopyranoside (IPTG) and cultivation was continued for four hours at 30 °C and 120 rpm. Subsequently, the cells were harvested by centrifugation (4 °C, 30 min, 3399 g) and resuspended in buffer A (100 mM Tris-HCl, 100 mM NaCl, pH 7.5) supplemented with cOmplete Mini EDTA-free protease inhibitor cocktail tablets (Roche Diagnostics GmbH, Mannheim, Germany). Cell disruption was performed by five passages through a French® pressure cell using an HTU-Digi-F-Press (G. Heinemann Ultraschall- und Labortechnik, Schwaebisch Gmuend, Germany) at a pressure of 103.4 MPa (15,000 psi). Cell debris was removed by centrifugation for 20 minutes at 5300 gand 4 °C, followed by ultracentrifugation for one hour at 84,000 g and 4 °C. The supernatant was directly used for affinity chromatography using a Strep-Tactin[®]-Sepharose[®] column (IBA, Göttingen, Germany) with 1 mL bed volume, equilibrated with buffer W (50 mM Tris-HCl, 250 mM NaCl, pH 7.5). After the protein extract had passed, the column was washed three times with 15 mL buffer W and FtsR-Strep was eluted ten times with 1 mL buffer E (buffer W with 15 mM D-desthiobiotin (Sigma-Aldrich Chemie GmbH, Steinheim, Germany)). Aliquots of the elution fractions were analysed by SDS-PAGE (Laemmli, 1970) and Coomassie staining and the fractions containing FtsR-Strep were pooled and concentrated using Amicon® Ultra-4

Centrifugal Filter Devices with a 10 kDa cut-off (Merck Millipore Ltd., Carrigtwohill, Cork, Ireland). FtsR-Strep was further purified by gel filtration using a Superdex[™] 200 Increase 10/300 GL column attached to an ÄKTA[™] pure 25 system (GE Healthcare Bio-Sciences AB, Uppsala, Sweden). The column was equilibrated with buffer W and elution was performed with a flow rate of 0.6 mL/minute.

3.5. Electrophoretic mobility shift assays (EMSAs)

Two complementary single-stranded 30-bp oligonucleotides (see Table 3) covering the putative FtsR-binding motif in the *ftsZ* promoter region were annealed by heating the samples containing 10 µM Tris-HCl pH 8.0, 50 mM NaCl, 1 mM EDTA and 10 µM of each oligonucleotide to 95 °C for five minutes, followed by a slow cooling to room temperature. This double-stranded oligonucleotide (final concentration 1 µM) was incubated at room temperature for 30 minutes with increasing concentrations (0 to 8.5 µM) of freshly purified dimeric FtsR protein (see Figure 25) in binding buffer (10 mM Tris-HCl pH 7.5, 30 mM NaCl, 1.5 mM Na₂EDTA). For this purpose, FtsR protein was used directly after elution from the Strep-Tactin[®]-Sepharose[®] column, because its binding capability seemed to decrease during the purification process. A 30-bp fragment located in the cg1081 promoter region was used as negative control (see Table 3). After incubation, a suitable volume of 6-fold concentrated sample buffer (0.1% (w/v) xylene cyanol, 0.1% (w/v) bromophenol blue, 20% (v/v) glycerol in 1x TBE (89 mM Tris base, 89 mM boric acid, 2 mM Na₂EDTA)) was added and the samples were separated by native PAGE. using 15% (w/v) polyacrylamide gels in 1x TBE buffer. Electrophoresis was performed on ice at 180 V for one hour. Subsequently, the gel was stained with SYBR® Green I Nucleic Acid Gel Stain (Invitrogen, Ltd., Paisley, UK) and photographed.

The competition-EMSA was performed as described previously (Garcia-Nafria *et al.*, 2013). In brief, a 271-bp DNA fragment covering the *ftsZ* promoter region including the binding site was amplified using the oligonucleotides ftsZ_prom250_fw and Cy3-ftsZ_prom_rv to generate Cy3-labelled DNA or the oligonucleotides ftsZ_prom250_fw/ftsZ_prom_rv to generate unlabeled specific competitor DNA. A 260-bp DNA fragment further upstream in the *ftsZ* promoter was amplified using the oligonucleotide pair ftsZ_prom500_fw/ftsZ_prom250up_rv and used as unspecific competitor DNA. Purified FtsR protein was incubated with the DNA fragment(s) in a total volume of 10 µL binding buffer. After addition of loading buffer and separation by native PAGE on a 10% polyacrylamide gel, the gel was scanned using a Typhoon TrioTM scanner (GE Healthcare Bio-Sciences AB, Uppsala, Sweden).

3.6. Promoter studies with Pftsz fused to mVenus

In order to analyze transcriptional regulation of *ftsZ* by FtsR *in vivo*, a DNA fragment covering the *ftsZ* promoter region and the native ribosomal binding site extending from position -494 to +3 with respect to the *ftsZ* translational start site was fused to a DNA sequence coding for the fluorescent protein mVenus and cloned into the pJC1 vector backbone (for details see Tables 2 and 3). Strains carrying their chromosomal *ftsZ* gene either unter control of its native promoter or under control of the *gntK* promoter were transformed with the resulting plasmid pJC1-P_{*ftsZ*}-*venus*. For complementation studies, the strains were additionally transformed with pEC-*ftsR* or the empty plasmid pEC-XC99E as control. Growth experiments were performed in the BioLector[®] system as described above, with additional monitoring of the *ftsZ* promoter activity by online measurement of mVenus fluorescence at an excitation wavelength of 508 nm and an emission wavelength of 532 nm.

3.7. Chromatin affinity purification with subsequent Sequencing (ChAP-Seq)

For ChAP-Seq experiments, C. glutamicum ATCC13032 ftsR pAN6-FtsR-Strep and the negative control C. glutamicum ATCC13032 ftsR pAN6 were cultivated as described above for FtsR purification with 10 µM instead of 50 µM IPTG. FtsR-Strep was purified as described above with the following alterations: after harvesting (15 minutes, 6371 g, 4 °C), washing with 40 mL PBS, and a second centrifugation step (15 minutes, 5525 g, 4 °C), the supernatant was discarded and the cells were suspended in 20 mL PBS containing 1% (w/v) formaldehyde and incubated at room temperature for 20 minutes, followed by addition of glycine to a final concentration of 125 mM and incubation for another 5 minutes at room temperature. The cells were washed twice with 40 mL buffer A and suspended in 20 mL buffer A with a suitable amount of cOmplete Mini EDTA-free protease inhibitor cocktail tablets and 5 mg of RNase A (both Roche Diagnostics GmbH, Mannheim, Germany). After cell disruption using a French® press as described above, the crude extract was transferred to a sonication vessel, placed in an ice bath and sonified three times for 30 seconds using a Branson Sonifier 250 (G. Heinemann Ultraschall- und Labortechnik, Schwaebisch Gmuend, Germany) with a pulse length of 40% and an intensity of one to shear the genomic DNA. The sonified crude extract was used for Strep-Tactin[®] affinity chromatography as described above. The FtsR-containing elution fractions were pooled, 1% (w/v) SDS was added, and the sample was incubated overnight at 65 °C, followed by Proteinase K treatment (final concentration 400 µg/mL) for 3 hours at 55 °C. The DNA in the sample was purified by phenol-chloroform-isoamyl alcohol extraction (Chomczynski & Sacchi, 2006), precipitated with ethanol and sodium acetate, washed with 70% (v/v) ethanol, dried, and dissolved in 50 µL deionized water. Sequencing of the DNA fragments and data evaluation were performed as described previously (Pfeifer et al., 2016). Genomic binding sites of FtsR were defined to have a sequence coverage that is at least 50-fold above the average genome coverage determined for the negative control strain *C. glutamicum* ATCC13032 pAN6.

3.8. DNA microarrays

DNA microarray analysis was performed to compare the mRNA levels of the ATCC13032 Δ *ftsR* mutant with the wild type. Therefore, cells were cultivated in 50 mL CGXII medium with 2% (w/v) glucose and harvested on ice in the exponential growth phase. RNA sample preparation, labelling, hybridization, and comparative transcriptome analysis were performed as described previously (Vogt *et al.*, 2014). The full data set of this experiment has been deposited in the NCBI Gene Expression Omnibus and can be found under the GEO accession number GSE107921.

3.9. DNA affinity purification and MALDI-ToF-MS analysis

To identify putative regulatory proteins binding to the *odhl* or the *ftsZ* promoter region, DNA affinity purification was performed. The promoter region of odhl or ftsZ was amplified by PCR (odhl: primers AP Podhl fw and AP Podhl rv bio, product size 590 bp, DNA fragment extending from -201 to +62 with respect to the translational start site of odhl; ftsZ: primers AP PftsZ fw and AP PftsZ rv bio, product size 404 bp, DNA fragment extending from -285 to +96 with respect to the translational start site of ftsZ). The resulting fragments were tagged with biotin by a second PCR using the biotinylated oligonucleotide biotin oligo and the respective forward primer, AP_Podhl_fw or AP_PftsZ_fw. The PCR products were purified by size exclusion chromatography using a column with 8 mL Sephacryl[™] S-400 High Resolution Chromatography Media (GE Healthcare Bio-Sciences AB, Uppsala, Sweden) as bed material and TE buffer (10 mM Tris-HCl, 1 mM Na₂EDTA, pH 7.6). Fractions containing the desired DNA fragment were pooled, concentrated to about 500 µL using an Eppendorf concentrator plus (Eppendorf AG, Hamburg, Germany), and precipitated with ethanol. The precipitate was dissolved in 50 µL TE buffer and the DNA concentration was measured with a Colibri Microvolume Spectrophotometer (Berthold Detection Systems GmbH, Pforzheim, Germany) to assure a minimal amount of 220 pmol biotinylated DNA, which was required for the following steps. The DNA affinity chromatography was performed using Dynabeads[®] M-280 Streptavidin (Life Technologies AS, Oslo, Norway) as described in Pfeifer et al. (2016). In brief, the biotinylated DNA was coupled to the beads and incubated with crude cell extract of C. glutamicum ATCC13032 cells that had been cultivated in glucose minimal medium and harvested in the exponential growth pase. After several washing steps, the proteins were eluted using buffer containing 2 M NaCl. The elution fractions were subjected to protein precipitation with trichloroacetic acid (TCA, 10% (w/v) final concentration) and washed once with acetone. The precipitated protein was dissolved in 20 µL Tris-HCI buffer pH 7.5 and analysed by SDS-PAGE (Laemmli, 1970) using a 12% Mini-PROTEAN[®] TGX[™] gel (Bio-Rad Laboratories, Inc., Hercules, CA, USA). The gel was stained with GelCode[®] Blue Stain Reagent (Thermo Fisher Scientific Inc., Rockford, IL, USA). Proteins enriched with the promoter region of *odhI* or *ftsZ* were identified by peptide mass fingerprinting after tryptic in-gel digestion of the excised bands of the polyacrylamide gel, followed by MALDI-ToF-MS using an Ultraflex III TOF/TOF mass spectrometer (Bruker Daltonics, Bremen, Germany) as described previously (Schaffer *et al.*, 2001, Koch-Koerfges *et al.*, 2012).

3.10. Genome re-sequencing

For genome re-sequencing, C. glutamicum ATCC13032 and three independent clones of C. glutamicum ATCC13032 fts R were cultivated overnight in 20 mL BHI medium and the DNA was prepared as described previously (Pfeifer et al., 2017). Genomic DNA was purified using the NucleoSpin[®] Microbial DNA Kit (MACHEREY-NAGEL GmbH & Co. KG, Düren, Germany). 4 µg were used for library preparation and indexing with the TruSeq[®] DNA PCR-Free Sample Preparation Kit (illumina Inc., San Diego, CA, USA). Quantifications of the resulting libraries were conducted using KAPA Library Quantification Kits (PEQLAB Biotechnologie GmbH, Erlangen, Germany) and were normalized for pooling. A MiSeqTM sequencing device (illumina Inc., San Diego, CA, USA) was used for paired-end sequencing with a read-length of two times 150 bases. Data analysis and base calling were accomplished with the illumina® instrument software and stored as fastg output files. The sequencing data obtained were imported into CLC Genomics Workbench (Qiagen Aarhus A/S, Aarhus, Denmark) for trimming and base quality filtering. The output was mapped to accession BX927147 as the C. glutamicum ATCC13032 reference genome. The resulting mappings were used for the qualitybased single nucleotide polymorphisms (SNPs) variant detection with CLC Genomics Workbench. The detected SNPs were manually inspected for relevance.

3.11. Quantitative PCR

To elucidate how many copies of the genes cg0834 and cg0840 were present in the genome of different *C. glutamicum* strains, quantitative PCR (qPCR) was performed. For this purpose, genomic DNA was isolated as follows: selected strains were incubated overnight in 5 mL BHI medium and harvested by centrifugation. The cells were washed in TE buffer (10 mM Tris-HCl, 1 mM Na₂EDTA, pH 7.6), centrifuged again and suspended in 1 mL TE buffer with 15 mg lysozyme. After three hours incubation at 37 °C, 3 mL lysis buffer (10 mM Tris-HCl, 2 mM Na₂EDTA, pH 8.2), 220 μ L 10% (w/v) SDS and 150 μ L proteinase K

(20 µg/mL) were added to the suspension and mixed carefully, followed by an additional incubation for two hours at 60 °C. Two milliliter of saturated saline solution (about 6 M NaCl) was added and the samples were shaken vigorously until a white precipitate appeared, which was spun down by centrifugation at room temperature for for 30 minutes at 16000 g. The cleared supernatant was transferred into a fresh reaction tube and the DNA was precipitated by careful mixing with 2.5 volumes of ice-cold absolute ethanol. The DNA was removed from the tube with a Pasteur pipette whose tip was bent before using a Bunsen burner, dipped into 70% ethanol in a fresh Eppendorf tube for washing, and air-dried for a few seconds. Subsequently, the DNA was solubilized in 200 µL TE buffer by incubation at 4 °C overnight. For guality control, the isolated genomic DNA was analyzed by agarose gel electrophoresis. The DNA concentration was determined using a Colibri Microvolume Spectrophotometer (Berthold Detection Systems GmbH, Pforzheim, Germany) and adjusted to 50 ng/µL. For the qPCR experiment, PCR fragments for the generation of standard curves were amplified with the KAPA2G Fast ReadyMix PCR Kit (Kapa Biosystems, Boston, Massachusetts, United States) using chromosomal C. glutamicum DNA as template and the oligonucleotides given in Table 3. The gene recF was used as reference gene, as it has proven to be well suitable in previous projects. The primers were designed with the Primer3plus online tool (Untergasser et al., 2012), using the standard settings for qPCR and an annealing temperature of 60 °C. The amplified fragments were checked for purity by agarose gel electrophoresis. The standards were used in concentrations of 10 pg/µL to 100 ag/µL in a gradient cycle protocol to determine the optimal annealing temperature for the qPCR experiment. The setup of the cycling protocol was as following (see Table 4): three minutes pre-incubation at 95 °C (step 1), five seconds denaturation at 95 °C (step 2), and 25 seconds elongation at a temperaturegradient from 55.1 °C to 66.9 °C (step 3), 40 times repetition of steps two and three, followed by a melting curve analysis (step 4) from 60 °C to 95 °C, with $\Delta T = 1$ °C for every 6 seconds.

step #	°C	min:sec	go to step	loops	
1	95	3:00			
2	95	0:05			
3	55.1 to 66.9	0:30	2	39	
4		melting curve ana	alysis, 60 °C to 95 °C, 6	δ s with Δ T = 1 °C	

Table 4: Determination of the copy number of cg0834 and cg0840. The qPCR program used for the determination of the copy number of cg0834 and cg0840 is shown.

Again, agarose gel electrophoresis of the PCR products was performed. Based on this experiment, an annealing temperature of 59 °C was chosen for the qPCR experiment. The

qPCR reaction was performed using the innuMIX qPCR MasterMix SyGreen (Analytik Jena, Jena, Germany) and the qTOWER 2.2 (Analytik Jena, Jena, Germany), and the procotol stated in Table 4, but with a constant temperature of 59 °C in step three. The reaction mix contained 10 μL master mix (2x), 1 μL primer 1 (10 μM), 1 μL primer 2 (10 μM), 2 μL template (50 ng/μL) and 6 μL H₂O. The data were analyzed using the program qPCRsoft 3.1 (Analytik Jena, Jena, Germany) and the ΔΔCt method (Livak & Schmittgen, 2001).

3.12. Coulter counter measurements

Bacterial size distribution was determined *via* the Coulter principle using a MultiSizer 3 (Beckman Coulter, Krefeld, Germany) particle counter equipped with a 30 μ m capillary. Briefly, for each measurement the bacterial cells at an OD₆₀₀ of about 0.1 were 20-fold diluted in CASYton assay buffer (Schärfe System GmbH, Reutlingen, Germany). Each sample (100 μ L) was analyzed twice in the volumetric measurement mode. The data were visualized and extracted using the Beckman Coulter Multisizer 3 software package.

4. Results

4.1. Phylogenetic conservation of odhl (cg1630) and adjacent genes

The *odhI* (cg1630) gene and its immediate genomic vicinity including the three downstream genes *ftsR* (cg1631), cg1632, and cg1633 were found to be highly conserved within Actinobacteria (Figure 4).



Figure 4. Phylogenetic conservation of *odhl* **(cg1630) and adjacent genes.** The *ftsR* (cg1631) gene and its homologs in different actinobacterial species of various genera are shown in red. Neighboring homologous genes are colored alike, whereas grey arrows indicate genes that are not conserved in the gene cluster. The black arrows indicate transcriptional start sites determined with RNA-Seq for *C. glutamicum* by Pfeifer-Sancar *et al.* (2013). Note that the cluster comprising *ftsR*, its upstream gene *odhl/garA*, and the two downstream genes encoding a bifunctional nuclease and a MerR-type transcriptional regulator is strongly conserved. The figure was prepared based on data of MicrobesOnline (Alm *et al.*, 2005) and ERGO (Overbeek *et al.*, 2003). The amino acid sequence identity of the FtsR homologs to *C. glutamicum* FtsR is given on the right side and was derived from NCBI BLAST (Altschul *et al.*, 1990). For lack of space, some locus tag prefixes were omitted in short genes. Gene lengths are approximately to scale. The full species names are as following: *Corynebacterium glutamicum, Corynebacterium diphtheriae, Mycobacterium tuberculosis, Rhodococcus erythropolis, Nocardia farcinica, Tsukamurella paurometabola, Streptomyces coelicolor, Nakamurella multipartita, <i>Propionibacterium acnes, Acidothermus cellulolyticus, Actinosynnema mirum, Micrococcus luteus*.

The *odhl* gene encodes the 2-oxoglutarate dehydrogenase inhibitor protein Odhl, which acts as a phosphorylation-dependent switch, controlling the activity of the 2-oxoglutarate dehydrogenase complex (Niebisch *et al.*, 2006, Krawczyk *et al.*, 2010, Raasch *et al.*, 2014). The homologous protein of mycobacteria is GarA (Ventura *et al.*, 2013). The gene *ftsR* (cg1631) of *C. glutamicum* ATCC13032 encodes a protein of 252 amino acids (calculated mass 27.23 kDa), which was annotated as a transcriptional regulator of the MerR family due to the sequence similarity of the N-terminal region of FtsR to other members of the MerR family. The gene cg1632 encodes a putative bifunctional nuclease having both DNase and RNase

activity (PFAM PF02577). In *M. tuberculosis*, the homologous protein Rv1829 has been described as a carbon monoxide resistance gene (Zacharia *et al.*, 2013). cg1633 encodes another yet uncharacterized transcriptional regulator of the MerR family. Previous RNA-Seq analysis indicated that these neighboring genes are co-transcribed in various combinations: cg1629-cg1633 was annotated as primary operon and cg1630-cg1633 and cg1631-cg1633 as sub-operons (Pfeifer-Sancar *et al.*, 2013) (Figure 4). The genomic region upstream of *odhl* (cg1630) and downstream of cg1633 varies in different Actinobacteria.

An amino acid sequence alignment of FtsR homologs of different actinobacterial species revealed that especially the N-terminal part is highly conserved (Figure 5). Two features of the alignment are conspicuous, which are the variable length of the N-termini upstream of the highly conserved region starting with the sequence MSIG and the variable length of the region linking the N-terminal DNA-binding domain and the C-terminal regulatory domain. Notably, most of the other sequences are considerably shorter than *C. glutamicum* FtsR and indeed, a putative alternative start codon was found 28 amino acids downstream of the annotated start codon. However, complementation experiments with the two variants showed that the shorter protein version features a lower degree of complementation suggesting that the longer version is the protein natively synthesized *in vivo* (Figure 7).

MerR-like DNA-binding helix-turn-helix domain

	1	10	20	30	40	5 Q	60	7 <u>0</u>	вò	9Ò	100
C. glutamicum	MSALRKTS	PNGSIGA	SATRTVPV	KPTKTMSIGV	VLERLNAEFP	DVTVSKIRFLE	SEGLITPER	TA <mark>SGYRRFT</mark> ESI	VERLRYIL	TORDNYLPL	KVIREQLEAMDNG
C. diphtheriae	MSALPQRN	DSGSAGSYQP	RPQSSSLS	SSKKT <mark>MSIG</mark> V	VLAKITAEFP	DVTVSKIRYLE	SEGLITPOR	I S TGYRRFTQDI	VERLRYIL	TORDNYLPL	KVIREQLEAMD SG
M. tuberculosis	M		SAPDS	PALAGMSIGA	VLDLLRPDFP	DVTISKIRFLE	AEGLVTPRR	ASSGYRRFTAY	CARLRFIL	AQRDHYLPL	KVIRAQLDAQPDG
R. erythropolis	. м			QSQPGMSIGS	VLDRLRPDFP	DVTISKIRFLE	SEGLISPOR	T P SGYRRFS VEI	OCERLREVL	A ORDOYLPL A ORDOYLPL	KVIKAQIDALPDG KVIKEQLEAIDSG
N. farcinica	MT		GAAQQ	WARGG <mark>MSIG</mark> S	VLDLLRPDFP	DVTISKIR <mark>f</mark> le	SEGLIRPER	T P <mark>S G Y R F S</mark> V A I	CERLRFIL	T <mark>AQRDQYL</mark> PL	KVIKEQLEAIDKG
Dietzia sp. Williamsia sp.	MTA		AGHGA	AEEKHLSIGG MSIGS	VIALLSPDFP VLDOLRGDFP	DLTVSKVRFLE DVTISKIRFLE	SEGLITPOR	FASGYRRFSVE Spsgyrrfstgi	DRERLRYVL	AQRDRYLPL AORDRYLPL	KVIREELEALDSA KVIKEOLEAIDRG
Gordonia sp.	MTA		ASAGQ	RGAGVMSIGS	VLGRLRDDFP	DVTISKIRFLE	SEGLVTPER	A P <mark>S G Y R R F S</mark> D S I	YERLRFVL	T <mark>A</mark> QRDR <mark>YL</mark> PL	KVIKEQLDEMDAG
T. paurometabola	MTS		SCCACOCAAA	TAPSAMSIGA	VLEQLRAEFP. VLNALPDEEP	DVTISKIRFLE DTTTSKIRFLE	SEGLVEPDRS	S P S G Y R R F S P K I	DCERLRYVL	AQRDYYLPL MORDHYLPL	KIIKEQLDAHDRG
N. multipartita	MTTS		LAEQPDM	IPPNSTLSIGA	VMTRLKPEFP	EVSISKIRFLE	SEGLVTPHR	F P S G Y R Q F S P A I	VSRLRYVL	AQRDQYLPL	KVIKDHLDAIDRG
P. acnes				MPRSVRTIGQ	VMKTLKPDFP	DLSISKIRFLE	SEGLLSPER	A P S G Y R K Y S D S I	VERLRYIL	CORDHFOPL	RVIRDHLEMMDRG
A. cellulolylicus A. mirum				PORGGMGIGA	VLSQLRSEFP	GVTISKIRFLE	AEGLVRPAR	TASGYRRFSVSI	VERLRYVL	AORDRYLPL	KVIREQLDAADDG
M. luteus	МТ	I	DPRPRPDAVPTA	RPAGR <mark>LGIG</mark> E	V <mark>V</mark> AÃ <mark>le</mark> aefp	<mark>GVTA</mark> SK <mark>V</mark> RFLE	DRGLVLPER	<mark>F P A G Y R R F R</mark> P E I	DVDRLRFVL	ALQRDH <mark>FL</mark> PL	KVIADHLAALDRG
	110								120	130	140
C. glutamicum	SVTAILGS	. S		• • • • • • • • • • •				•••••	SEPLVSPE	KFQAP.AIT <mark>R</mark> KFRAP ALT <mark>R</mark>	LTD SDVAEKAGVN
C. diphtheriae	AVTPISEG	SD	 						NAPLISPES	SFRAS.VVT <mark>R</mark>	LSDSDVAARAQVS
M. tuberculosis	ELPPFGSP	YVLPRLV			PV.	AG	DSAGG	VGSI	TASVSL	TGI <mark>R</mark>	LSREDLLERS GVA
R. erythropolis	AATVSAEV	PAP			PV .	GL	RRI	ARVLSI	ATGDVSP.E	AFRVD.REV <mark>R</mark>	VGRQDLLARADID
N. farcinica	AATLGVRE	ARARAL		SSRAGG	AEAPA.	AG	AGSAGGRGA	PRKLGV	VPDEVSP.EI	ELRVD.HEI <mark>R</mark>	LTRADLLAQAGID
Dietzia sp. Williamsia sp.	IADGST.T ERIDPGTP	ALLPRHG							OGVPGSTPDI ARSWVAPAVI	DFRSD.TVL <mark>R</mark> DFER.SGG <mark>R</mark>	LTREDVVTQSGVD ISRETLLERTGAD
Gordonia sp.	SSTTGGQP			S				S2	ARGAVAPATI	DFGTRGS <mark>R</mark>	VSRDDLIERTGVD
T. paurometabola	ESVEGPAP	RAPRTL		T			• • • • • • • • •	AV	VSSKTAPATI	DFSAR.RQT <mark>R</mark>	ISRSDLLARSGAD
N. multipartita	LEPAVPQA	RLPAA						NO	GSSDTPLPRI	DLAA.GREVR	MTRNELLAHSGLT
P. acnes	EEPPVSEA	PPLPT.ENEG					T	PI	PQPQPTGQ	GIVRTRGPI <mark>R</mark>	MNRRELIRASGIT
A. cellulolyticus A. mirum	LSGPAPAG	PVPLSAERDVI		PWSTTRRPGG	ARSGDEVPAA	RDAEHPLPAGE	AGPAPDRTA	PARVAAPTAPAI	GEGMPGPD PRSELPTAAI	DLAPA.PGDR	LSREPLQAAGIA
M. luteus	ERPTGMPG	TAAVPASSDA							ERL0	GREVAGGRR <mark>S</mark>	WTRAELAAES <mark>GA</mark> G
	150	160	170	180	190	2 0 Q	21 Q	2 2 Q	230	24 Q	2 5 <u>0</u>
C. glutamicum C. efficiens	VELVVDLV	NARLIKPDAA SAKLIKPDAA	GEETHDDVAIA	STAASLKAMG	FDLRRLKSLG	NAASRQADLIS NTATROADLIT	QVASPIAQGI	KSDVARQQAEEN KSDVAROBAEEN	ACOMCSLV	SMHASLVKN	ATREQLGY
C. diphtheriae	ESEVSELA	EAGLIRPDSS	GFFTADDVQVV	SVAVQLKEFG	FDVRHLKSLR	NLASRHADLIS	RATTPVARS	QSES <mark>AR</mark> Q <mark>RAEE</mark>	ISQQLSALV	SLNATMLRS	MLRDELNR
M. tuberculosis	DELLTALL	KAGVITTGPG	. GFFDEHAVVIL	QCARALAEYG	VEPRHLRAFR	SAADRQSDLIA	QIAGPLVKA	GKAGARDRADD	LAREVAALA	TLHTSLIKS	AVRDVLHR
R. ervthropolis	DRFLTDLI	RTGLVVPGAA	. GEFDEHSVVIA	RTAKAMSEYG	LEVRHLRAFK	LAADROSDLIA	OIAGPVVKAG	RDAGARDRADD	IVRELAALSI	TLHTSLIKS	AVRGSLDR
N. farcinica	EPFLNDLI	RAGLITPGPA	. GFFDGEAVTLA	KT <mark>A</mark> RA <mark>M</mark> AEFG	LEARHLRAFK	LAADREAALLA	QIAAPIAKSI	RDAG <mark>AR</mark> A <mark>RAEE</mark>	TVRELA ALSI	LT LHTS LVKA	SVRHALGN
Dietzia sp.	AEFVGSLI	DAGLIVPGAG	. GFFDPEAVLVA	RTAHDLGGHG	VDVRHLRGFR	TAADROTGLII	QIAGPVARQO	GDADARDRAAE		ALHSTLVTV	AVRHALES
Gordonia sp.	TAFVAELQ SAFTTELM	RSGLLTPGAG RNGLLTAGPG	GFFDEDAVRLV	EAASALADYG	LEARHLRAFK	V SADREAGLVA V SADREAGLVA	QIANPIAKGE OTASPTAKGE	KGAGARDRAEEI	UREVAALS	TLHTQLVKA	
T. paurometabola	EAFLRELE	RSALLTSGKA	GFFDEDAVELV	RAAKALAEYG	LEARHLRAFK	TSADREAGLIA	QIANPVARGO	GDAGAAERAAE	LIRELAALS	TFHTQLVKA	AVKDAVR
S. coelicolor	EQELKEWE	SYGLLVPLPD	. GAYDAEAVTVA	SLVVQLGRFG	IEPRHLRVMK	AAADREAGLVI	QVVAPLKRHI	RNPQ TRAHAEA I	RTKELAGLA	KLHAALVQT	ALGVRLP
N. multipartita P acnes	ATSLAELE EAMLMELE	QFGLLSAGPG RHOLVEPKPC	SYFCOFALVIC	STSAELLAVG	LEARHLRSFR MDTRHMRATE	TAADREATLIT	QLVSAQAHQI	RDPDARERAGAI	SAAQLASTI	HAHTAMVYD	GLRRDLGR
A. cellulolyticus	DEELATLE	NFGLIAPR.G	. STYDGTALAIA	KTVKELTRFG	IEPRHLRAFR	AAADREVGLVE	QVITPLVRQ	RSPEARARAEE	AREIAALS	RLHAALVRS	GLRPLLSR
A. mirum	RAALAELE	RSGLLAPGPG	. GLHDPDALAVA	RTVRALARYG	VELRDLRPAR	AAADRELATVE	RVLAPLRPGI	RDPR <mark>TRARADE</mark>	GREVAALL	ALHTLLVRS	GVRELAGDRVSPR
M. IULEUS	EELLAELD	UISLLPVDPD	. GGIPSHAMDVA	RAAVVLAGHC	LEPKHLKPLR.	AACDRELGLVE	RAVAPLOARI	KDAGTRPRVARS	SARELAUASI	LHAALVAV	GLEQWDD

Figure 5. Alignment of *C. glutamicum* **FtsR and homologous proteins of other** *Actinobacteria.* The alignment was prepared using Clustal Omega (Sievers *et al.*, 2011) and edited using ESPript 3.0 (Robert & Gouet, 2014). Residues shown in yellow are at least 70% identical and residues indicated in red are fully conserved. The following proteins were aligned: *Corynebacterium glutamicum* FtsR (Cg1631); *Corynebacterium efficiens* CE1574; *Corynebacterium diphtheriae* DIP_1205; *Mycobacterium tuberculosis* Rv1828; *Mycobacterium smegmatis* MSMEG_3646; *Rhodococcus ruber* CS378_RS03820; *Nocardia farcinica* NFA_24870; *Dietzia* sp. H483_RS33310; *Williamsia* sp. ASG12_08425; *Gordonia effuse* GOEFS_105_00530; *Tsukamurella paurometabola* Tpau_2338; *Streptomyces coelicolor* SCO1383; *Nakamurella multipartite* NAMU_RS19840; *Propionibacterium acnes* PPA_RS05480; *Acidothermus cellulolyticus* ACEL_RS06295; *Actinosynnema mirum* AMIR_RS26540; *Micrococcus luteus* CRM77_RS01150.
4.2. Deletion of *ftsR* affects growth behavior

The possibility that *odhl* might be regulated on the transcriptional level seemed logical since it is located upstream of the two genes *ftsR* and cg1633 (Figure 4), which encode two transcriptional regulators. To elucidate if any of the three genes located downstream of *odhl* are involved in its regulation and to investigate their physiological role, in-frame deletion mutants of the genes *ftsR*, cg1632, and cg1633 were constructed and analyzed with respect to their growth behavior. Remarkably, the *ftsR* deletion strain grew significantly slower and to a lower final cell density (backscatter) compared to the wild type in CGXII minimal medium with glucose as carbon source, whereas the results for strains Δ cg1632 and Δ cg1633 were comparable with the wild type (Figure 6a).



Figure 6. Growth behavior of *C. glutamicum* ATCC13023 and *C. glutamicum* ATCC13023 with deleted *ftsR*, cg1632, or cg1633 in a BioLector microcultivaton system (a) and of strain $\Delta ftsR$ in shaking flasks (b). The cells were first cultivated in BHI medium followed by two consecutive cultivations in CGXII minimal medium with 4% (w/v) glucose as carbon source. Means and standard deviations of three biological replicates of the second CGXII culture are presented.

For further investigation of the growth defect of the $\Delta ftsR$ mutant, an additional growth experiment for the determination of the growth rate in comparison to the wild type was performed in shaking flasks (Figure 6b). The mutant showed a growth rate of 0.32 ± 0.01 h⁻¹, which equals an 20% slower growth in comparison to the wild type ($\mu = 0.40 \pm 0.00 \text{ h}^{-1}$).

4.3. Complementation experiments with C. glutamicum ATCC13032∆ftsR

The consequences of *ftsR* overexpression in the wild type strain ATCC13032 were tested using the plasmid pAN6-*ftsR*, in which *ftsR* expression is controlled by the *tac* promoter. The promoter is known to be leaky in *C. glutamicum* and allows basal expression of the target gene

also in the absence of the inducer IPTG (Baumgart et al., 2016). To confirm that the growth defect of the $\Delta ftsR$ mutant is caused by the *ftsR* deletion rather than by secondary mutations that might have occurred during construction of the mutants or polar effects on downstream genes, complementation experiments were performed with the plasmid pAN6-ftsR and two variants thereof. Without IPTG and with IPTG concentrations up to 10 µM, growth of ATCC13032 pAN6-ftsR was comparable to the reference strain ATCC13032 pAN6. However, when 100 µM IPTG was added, growth of ATCC13032 pAN6-ftsR was strongly inhibited (Figure 7). Thus, both the absence and the overexpression of *ftsR* had a negative effect on the growth performance. Plasmid pAN6-ftsR-Strep encodes an FtsR variant with a carboxyterminal linker sequence (AS) followed by a Strep-tag-II (WSHPQFEK). Plasmid pAN6ftsR-short encodes an FtsR variant shortened by 28 amino acids at the N-terminus, which now consist of the amino acids MSIGV. An alternative start codon was found here, based on the alignment reported above (Figure 5). As shown in Figure 7, the growth defect of ATCC13032∆ftsR pAN6 could partially be reversed by pAN6-ftsR and pAN6-ftsR-Strep, but not by pAN6-ftsR-short. This suggests that the longer FtsR variant composed of 256 amino acid residues represents the active protein and that its function is not disturbed by a C-terminal Strep-tag-II.



Figure 7. Growth of an *ftsR* overexpressing strain and complementation of the Δ *ftsR* mutant with FtsR variants including an N-terminally shortened protein and a protein with a C-terminal Streptag-II. The strains were pre-cultivated first in BHI medium and then in CGXII medium with 2% (w/v) glucose, followed by main cultivation in the same medium. All media were supplemented with kanamycin (25 µg/mL) and IPTG as indicated. Means and standard deviations of two biological replicates of the second CGXII culture are presented.

4.4. The morphological phenotype of *C. glutamicum* caused by *ftsR* deletion and overexpression

Phase-contrast and fluorescence microscopy with staining of DNA and membranes revealed strong morphological differences to wild type cells for both the $\Delta ftsR$ mutant and for the ftsR overexpressing strain (Figure 8, Figure 9).



Figure 8. Microscopic pictures of cells in the stationary phase. To visualize membranes and DNA, cells were stained with Nile red and Hoechst 33342, respectively. The arrow points towards a branched cell and the circle indicates a high DNA concentration at the cell pole. The scale bar represents 5 µm.



Figure 9. Morphology of *C. glutamicum* ATCC13032 (a), ATCC13032 Δ *ftsR* (b), and ATCC13032 pAN6-*ftsR* (c). The cells were first cultivated in BHI medium followed by two consecutive cultivations in CGXII minimal medium with 2% (w/v) glucose as carbon source. For the plasmid-based overexpression (c), kanamycin (25 µg/mL) and IPTG (100 µM) were added. Fluorescence microscopy of stationary cells was performed. DNA was stained with Hoechst 33342 (cyan) and membranes with Nile red (red) as described in the Methods section. The scale bar is 5 µm.



Figure 10. Morphology of *C. glutamicum* ATCC13032, ATCC13032∆*ftsR*, ATCC13032 pAN6, and ATCC13032 pAN6-*ftsR* in different growth phases. See description of Figure 9.

Different cell morphologies were apparent both in the exponential and stationary growth phase; while the phenotype of the $\Delta ftsR$ mutant did not change much along the cultivation, the phenotype of the overexpression strain became more severe over time (Figure 10). This could be due to an accumulation of FtsR in the cells resulting from the overexpression. The $\Delta ftsR$ cells were elongated and some of them were branched (Figure 8, black arrow). Several cells had multiple septa at different positions in the cell. Overexpression of *ftsR* led to enlarged cells with DNA accumulated at the poles (Figure 8, black circle). These cells also contained additional septa, which were mostly located in the middle of the cells. However, only a fraction of the cells was enlarged, whereas the others remained small and inconspicuous. It should be noted that the small cells appear to be DNA-free in some pictures because the brightness was adjusted to the cells with the high DNA concentration. All small cells of the *ftsR* overexpression strain seem to contain similar DNA amounts comparable to the wild type. The reason for this population heterogeneity is yet unknown. It could result from different FtsR levels in the cells due to heterogeneous induction of pAN6-based ftsR expression, as previously reported for eyfp expression by the pAN6 parent plasmid pEKEx2 (Kortmann et al., 2015). The observed morphological changes caused by *ftsR* deletion and overexpression, in particular the presence of multiple septa within a single cell that did not lead to a separation of the cells, hint toward a function of FtsR in cell division and to a potential role in the regulation of *ftsZ* expression.

To further evaluate the changes in cell size or rather cell volume, we analyzed the strains using a Coulter counter in the volumetric measurement mode. Figure 11 shows the size distribution at three different time points during cultivation (3 h, 6 h, 24 h). For the $\Delta ftsR$ strain in comparison to the wild type, the whole population was shifted to larger cell sizes at all time points and formed a single broad peak. The population of the *ftsR* overexpressing strain formed two peaks of different sizes at 24 h, differing weakly or strongly in size from wild type carrying pAN6. As mentioned above, this could be due to heterogeneous expression from plasmid pAN6-*ftsR*. Overall, these measurements confirmed that the *ftsR* deletion strain comprises a homogeneous population of cells with increased size in different growth phases. In contrast, the strain overexpressing *ftsR* showed a homogeneous population of enlarged cells in the exponential growth phase, and two populations with cells of different sizes in the stationary phase. These results coincide with the cell size distribution observed by microscopy.



Figure 11. Coulter counter measurements in the exponential phase at 3 h (a) and 6 h (b) and in the stationary phase at 24 h (c). The cells were diluted to an OD_{600} of ~0.1 and analyzed in the volumetric measurement mode. The cells were assigned to bins according to their size. The bin diameter is proportional to the cell size. Plotted are mean values of two technical replicates.

4.5. Transcriptome comparison of the $\Delta ftsR$ mutant with its parent wild type

In order to elucidate whether *odhl* is transcriptionally controlled by the regulator FtsR and to investigate global gene expression in general to get an impression of the FtsR regulon, transcriptome analysis of the ATCC13032 Δ *ftsR* mutant and the parent wild type was performed using DNA microarrays and RNA isolated from cells cultivated in glucose minimal medium and harvested in the exponential growth phase. In total, 52 genes had an at least two-fold altered mRNA level, of which 34 genes showed a ≥ 2-fold increased mRNA level and 18 genes a ≥ 2-fold decreased mRNA level in the Δ *ftsR* mutant. However, expression of *odhl*, *secA2*, cg1632, and cg1633 was not significantly altered in the Δ *ftsR* mutant (Table 5), argueing against transcriptional regulation of *odhl* by FtsR.

cg1633

-				
locus tag	gene name	annotated function	mRNA ratio	p-value
cg1629	secA2	preprotein translocase subunit SecA2	0.59	0.073
cg1630	odhl	oxoglutarate dehydrogenase inhibitor protein	1.09	0.332
cg1631	ftsR	putative transcriptional regulator, MerR family	0.02	0.004
cg1632		putative secreted protein	0.52	0.101

putative transcriptional regulator, MerR family

Table 5: mRNA ratios of the *secA2* operon in a $\Delta ftsR$ deletion mutant. The transcriptome analysis of ATCC13032 $\Delta ftsR$ versus ATCC13032 (three biological replicates) of the genes located together in an operon with *odhl* using DNA microarrays are shown.

Both the upregulated and the downregulated genes covered a broad and heterogeneous range of cellular functions (Table 6). Within the group of genes showing a decreased mRNA level, the *ftsZ* gene with an mRNA ratio of 0.35 was the most prominent one. The group of upregulated genes included *oppC* and *oppD* encoding components of a putative ABC-type peptide transport system, four *prp* genes involved in propionate or propionyl-CoA catabolism, seven genes located in the CGP3 prophage region, which in total extends from cg1890 to cg2071, and a cluster of eight neighboring genes (cg0830-cg0838) including those for an ABC-type trehalose uptake system.

Locus tag	Gene name	Annotated function	mRNA ratio	p-value
cg0027		putative transcriptional regulator, MarR-family	2.39	0.034
cg0078		putative membrane protein	2.06	0.029
cg0256		putative protein, conserved	2.32	0.047
cg0274	maoA	putative oxidoreductase, possibly involved in oxidative stress response	2.51	0.048
cg0292	tnp16a	transposase	2.00	0.032
cg0661		hypothetical protein, conserved	2.33	0.040
cg0760	prpB2	2-methylisocitrate lyase, propionate catabolism	4.20	0.017
cg0762	prpC2	2-methylcitrate synthase, propionate catabolism	4.22	0.007
cg0797	prpB1	2-methylisocitrate lyase	2.03	0.018
cg0798	prpC1	2-methycitrate synthase	2.01	0.007
cg0824	tnp5a	transposase	2.44	0.003
cg0830		putative membrane protein	3.22	0.045
cg0831	tusG	ABC-type trehalose uptake system, permease	2.89	0.029
cg0832	tusF	ABC-type trehalose uptake system, permease	2.90	0.026
cg0833		putative membrane protein involved in trehalose uptake	2.63	0.030
cg0834	tusE	ABC-type trehalose uptake system, binding protein	2.98	0.039
cg0836		hypothetical protein	3.30	0.027

Table 6: Transcriptome analysis of the *C. glutamicum* strains ATCC13032 Δ *ftsR* and ATCC13032 using DNA microarrays. Three biological replicates of the experiment were performed, and genes are listed whose average mRNA ratio was increased or decreased at least 2-fold and a p-value ≤ 0.05 .

0.434

1.03

(Continued)

Table 6. Continued.

Locus	Gene	Annotated function	mRNA	p-value
cg0837	name	hypothetical protein	3.28	0.021
cg0838		putative helicase	4.61	0.026
cg1942		putative secreted protein CGP3 region	2.05	0.015
cg1969		hypothetical protein CGP3 region	2.03	0.026
cg1971		hypothetical protein CGP3 region	2.41	0.025
cg1995		hypothetical protein CGP3 region	2.23	0.042
cg2009		putative CLP-family ATP-binding protease,	2.22	0.047
cg2065		CGP3 region putative superfamily II DNA or RNA helicase, CGP3 region	2.83	0.021
cg2066		putative low-complexity protein	2.02	0.036
cg2183	оррС	ABC-type peptide transport system,	8.82	0.015
cg2184	oppD	ABC-type peptide transport system, ATPase component	17.99	0.005
cg2461	tnp4a	transposase	2.23	0.031
cg2570	dctP	C4-dicarboxylate-binding protein, TRAP-family	2.10	0.018
cg2644	clpP2	ATP-dependent Clp protease proteolytic subunit	2.14	0.006
cg2915		hypothetical protein	2.61	0.021
cg3181		putative secreted protein	2.30	0.046
cg3266	tnp5c	transposase	2.13	0.001
cg0370		putative ATP-dependent RNA helicase, DEAD/DEAH box-family	0.50	0.015
cg0789	amiA	putative N-acyl-L-amino acid amidohydrolase	0.48	0.003
cg0812	dtsR1 (accD1)	acetyl/propionyl-CoA carboxylase, β chain, mycolic acid biosynthesis	0.47	0.025
cg0896	_		0.20	0.032
cg0980	терВ	putative secreted protein related to metalloendopeptidase	0.49	0.001
cg1037	rpf2	resuscitation promoting factor, secreted protein	0.42	0.015
cg1076	glmU	putative	0.49	0.014
cg1290	metE	5-methyltetrahydropteroyltriglutamate-homocysteine methyltransferase,	0.39	0.036
cg1336		putative secreted protein	0.46	0.037
cg1370		hypothetical protein, conserved	0.39	0.026
cg1631	ftsR	transcriptional activator of <i>ftsZ</i> , MerR-family	0.02	0.004
cg2118	fruR	transcriptional regulator of sugar metabolism,	0.36	0.002
cg2366	ftsZ	presumably fructose responsive, DeoR-family cell division protein FtsZ	0.35	0.019
cg2477		hypothetical protein, conserved	0.24	0.007
cg2519		hypothetical protein, conserved	0.49	0.02
cg2853		putative protein fragment, conserved	0.45	0.006
cg3186	cmt2	trehalose corynomycolyl transferase	0.49	0.049
cg3335	malE (mez)	malic enzyme (NADP ⁺)	0.49	0.002

4.6. Genome re-sequencing of the *ftsR* deletion mutant and amplification of the trehalose cluster

To investigate whether the phenotype of the $\Delta ftsR$ mutant and the inability to fully complement it is solely caused by the respective deletion and not partially the result of secondary mutations, genome re-sequencing of three individual clones and comparison with the ATCC13032 wild type strain (reference genome published under accession number BX927147) was performed. Only four prominent single nucleotide polymorphisms (SNPs) could be detected (Table 7): a silent mutation each in cg1158 and cg2704, a mutation in the intergenic region between cg2057 and cg2058 (which are located in the CGP3 region), and a mutation in cg2030 (also located in the CGP3 region), causing a change from methionine to threonine. Presumably, these SNPs are not responsible for causing the abovementioned effects.

Table 7. Single nucleotide polymorphisms in ATCC13032 Δ ftsR strains. The table shows the fourSNPs discovered by genome re-sequencing of three individual ATCC13032 Δ ftsR mutants.

position	locus tag	annotated function	original base	new base
1074171	cg1158	putative secreted protein	Т	С
1921443	cg2030	hypothetical protein in CGP3 region	А	G
1947024	cg2057-cg2058	intergenic region in CGP3 region	С	Т
2577235	cg2704	putative ABC-type sugar transporter, permease subunit	С	Т

The transcriptome comparison of ATCC13032∆ftsR with ATCC13032 revealed 2.6- to 4.6-fold increased mRNA levels of the genomic region encompassing cg0830-cg0838 in the $\Delta ftsR$ mutant (Table 8). Interestingly, two of the nine genes annotated in this region are oriented in the opposite direction than the other seven, but nevertheless also showed an increased mRNA level. Although this could be a consequence of a similar regulation of the nine genes, it might also result from an amplification event leading to an increased DNA copy number and therefore increased mRNA levels. We therefore analyzed the sequenced genome of the three clones of strain ATCC13032 AftsR in this respect. For all three clones, this analysis indeed revealed an amplification event. As shown in Table 8, the sequence coverage of the DNA region from cg0828 to cg0840 was 5- to 7-fold (mean value 6.36 ± 0.57) higher than that of the residual genome in all three clones. When looking at the DNA microarray data, it becomes evident that the mRNA levels of cg0828, cg0829, cg0839, and cg0840 were also 2.5- to 3.4-fold increased in the $\Delta ftsR$ mutant, but since the p-value was above 0.05, they were not included in Table 6. The amplified DNA region includes the genes cg0831 and cg0835, which enocde the trehalose uptake system TusFGK₂-E (Henrich, 2011). Trehalose is a component of mycolates, which are important building blocks of the cell envelope of the Corynebacteriales (Tropis et al., 2005).

Table 8. Amplification of the DNA region encompassing cg0828-cg0840 in the ATCC13032 Δ *ftsR* **mutants.** Gene coverage values obtained from genome re-sequencing as well as data from the abovementioned transcriptome analysis are presented.

Locus	Gene	Annotated function	Gene	Gene o	overage		mRNA	p-
tag	name		coverage ATCC 13032	ATCC1 clone 1	3032∆ <i>ft</i> s clone 2	sR clone 3	ratio <i>∆ftsR/</i> WT	value
cg0819		hypothetical protein	0.99	1.07	1.14	1.01	0.97	0.45
cg0820	purE	phosphoribosyl- aminoimidazole carboxylase, catalytic subunit	1.1	1.08	1.2	1.03	0.93	0.39
cg0821		hypothetical protein, conserved	1.07	1.07	0.92	1.01	0.90	<0.01
cg0822		hypothetical protein,	1.02	1.03	0.94	0.99	1.38	0.02
cg0823	ntaA	nitrilotriacetate mono- oxygenase component A	1.06	0.89	0.99	0.93	0.85	0.32
cg0824		tnp5a, transposase	0.98	1.5	1.57	1.5	2.44	<0.01
cg0825	fabG	3-ketoacyl-acyl-carrier- protein reductase	0.9	0.91	0.95	0.98	1.08	0.22
cg0826		putative membrane protein	1.1	1.32	1.34	1.21	0.91	0.33
cg0827		hypothetical protein	0.99	1.07	1.14	1.01		
cg0828		putative dihydrofolate reductase	0.97	5.24	6.49	6.22	3.41	0.05
cg0829		putative lactoylglutathione lyase or related lyase, dyoxylase-family, conserved	0.94	5.95	7	7.09	2.91	0.06
cg0830		putative membrane protein	0.99	5.6	6.42	6.48	3.22	0.05
cg0831	tusG	trehalose uptake system,	1.03	5.42	5.93	6.23	2.89	0.03
cg0832	tusF	trehalose uptake system, ABC-type, mombrane energing protein	0.96	5.46	5.68	6.18	2.90	0.02
cg0833		putative membrane protein, involved in trehalose uptake,	1.01	6.18	6.74	7.34	2.63	0.03
cg0834	tusE	trehalose uptake system, ABC-type, bacterial extracellular solute-	1.1	5.9	6.35	7.02	2.98	0.04
cg0835	tusK (msiK2)	trehalose uptake system, ABC-type component	1.03	5.51	6.28	6.72	2.27	0.06
cg0836		hypothetical protein	0.98	5.72	6.4	6.74	3.30	0.03
cg0837		hypothetical protein	1.03	6.12	6.28	6.73	3.28	0.02
cg0838		putative helicase	1.05	6.4	7.1	7.28	4.61	0.03
cg0839		hypothetical protein	1.12	6.46	7.04	7.33	2.45	0.16
cg0840		hypothetical protein, conserved	1.04	5.67	6.79	6.64	2.45	0.19
cg0841		hypothetical protein,	0.98	1.02	0.98	0.93	1.36	0.05
cg0842		putative DNA helicase	1.02	1.01	0.98	1.01	1.11	0.21
cg0843		putative helicase	0.98	0.95	0.99	0.95	1.10	0.21
cg0844		putative type II restriction enzyme, methylase subunit	1.06	1.03	1	0.96	1.57	0.05

(Continued)

Locus tag	Gene name	Annotated function	Gene coverage ATCC	Gene c ATCC1 clone	overage 3032∆ <i>fts</i> clone	R clone	mRNA ratio ∆ftsR/	p- value	
			13032	1	2	3	WT		
cg0845		putative superfamily II DNA/RNA helicase, SNF2-family	0.97	0.94	1.03	0.89	0.61	0.06	
cg0847		putative transcriptional regulator, conserved	0.82	0.97	0.93	0.86	0.64	<0.01	
cg0848	wbbL	putative rhamnosyl transferase WbbL	0.85	0.92	0.95	0.84	0.53	0.02	
cg0849	rmIA2	GDP-mannose pyrophosphorylase, mannose-1-phosphate guanylyltransferase	0.98	1.02	1.02	0.96	0.61	0.14	
cg0850	whcD (whiB2)	transcription factor, <i>whmD</i> homolg, not involved in oxidative stress	1.09	1	1.05	1.2	0.68	0.01	

Table 8. Continued.

4.7. DNA affinity purification for unraveling transcriptional regulation of odhl

Another approach for analyzing transcriptional regulation of *odhl* was to identify potential regulatory proteins which bind to the *odhl* promoter region by DNA affinity chromatography, using a 590-bp DNA fragment covering the *odhl* promoter region and the first 84 bp of the *odhl*-coding region. The immobilized DNA fragment was incubated with crude protein extract of *C. glutamicum* wild type cells grown in glucose minimal medium. The proteins eluting after several washing steps were separated by SDS-PAGE and stained with Coomassie (Figure 12). Subsequent MALDI-ToF-MS analysis resulted in the identification of seven proteins. They represent proteins that are often found in DNA affinity purifications with *C. glutamicum* cell extracts independent of the promoter region used. Unfortunately, neither FtsR nor any other transcriptional regulator which might be involved in the regulation of *odhl* was identified.



Figure 12. DNA affinity purification with the *odhl* promoter region and crude cell extract of *C. glutamicum* ATCC13032. Proteins binding to the *odhl* promoter were separated by SDS-PAGE, stained with Coomassie and analyzed by peptide mass fingerprinting using MALD-ToF-MS. Identified proteins are marked with arrows and labeled with names and corresponding Cg numbers. The protein ladder used (left lane) is the Precision Plus Protein[™] Dual Xtra (Bio-Rad Laboratories, Inc., CA, USA).

Kim *et al.* (2010) have described that addition of penicillin for induction of glutamate production leads to a ten-fold increased expression of *odhl*, which suggests that inhibition of OdhA by OdhI might take place under these conditions. Based on this report, an additional DNA affinity purification experiment was performed with the following alterations: the culture volume for preparation of the crude protein extract was increased to five liters, glutamate production was induced at an OD₆₀₀ of about five by addition of 10 μ M penicillin G and the cells were harvested after seven hours at an OD₆₀₀ of about seven. The DNA affinity purification was performed as described above. Again, identification of proteins typically found in these kind of experiments (polymerases, topoisomerase, helicase, etc.) confirmed that the experiment *per se* was successful (Figure 13). Unfortunately, it was once more not possible to identify FtsR or any other transcriptional regulator which might be potentially involved in the regulation of *odhl*. Growth monitoring during this experiment and subsequent HPLC analysis revealed that induction with penicillin was successful due to stagnated growth and detection of an increased glutamate secretion (for additional data, see protocol of the practical course of master student Michele Reindl).



Figure 13. DNA affinity purification with the *odhI* **promoter and crude cell extract of** *C. glutamicum* **ATCC13032 cells induced for glutamate secretion by penicillin addition.** Proteins binding to the *odhI* promoter were separated by SDS-PAGE, stained with Coomassie and subjected to peptide mass fingerprinting using MALD-ToF-MS. Identified proteins are marked with arrows and labeled with names and corresponding Cg numbers. The protein ladder used (left lane) is the Precision Plus Protein[™] Dual Xtra (Bio-Rad Laboratories, Inc., CA, USA).

4.8. Focusing on investigation of the uncharacterized regulator FtsR

The results shown so far did not support our hypothesis that FtsR or another transcriptional regulator is involved in the regulation of *odhl*, which was initially one major goal of this doctoral thesis. Since deletion of *ftsR* led to a severe growth defect under standard conditions and strong morphological changes, indicating an important function of FtsR, it seemed reasonable to shift the focus of this work on the investigation of this yet uncharactarized protein. As

described above, the *ftsR* gene and its immediate genomic vicinity were found to be highly conserved within the Actinobacteria (Figure 4) and an amino acid sequence alignment of FtsR homologs of different actinobacterial species revealed that especially the N-terminal part is highly conserved (Figure 5). Since the microscopy and microarray data presented led to the strong suspicion that FtsR might be involved in the regulation of *ftsZ* and no transcriptional regulator of this gene was known to date in *C. glutamicum*, it was considered worthwile to investigate this protein in more depth. Thenceforward, we decided to name it FtsR, standing for regulator of *ftsZ*. Moreover, we decided to switch from the ATCC13032 wild type to the MB001 strain for some of the upcoming experiments, because it cannot be excluded that the amplification of the trehalose cluster might provoke secondary effects, which makes interpreting the experimental data more difficult.

4.9. Deletion and overexpression of the *ftsR* gene in the MB001 background

MB001 differs from ATCC13032 by the deletion of the three prophage regions CGP1 (13.5 kb), CGP2 (3.9 kb), and CGP3 (187.3 kb) (Baumgart et al., 2013b). It has been reported previously that the MB001 strain behaved like its parent wild type in the majority of conditions tested but showed improved growth and fitness under SOS response-inducing conditions that trigger CGP3 induction in the wild type. Thus, MB001 is a useful strain background for the analysis of deletions causing stressed cells, as secondary effects resulting from prophage induction can be excluded. In the context of this thesis, working with MB001 makes it even more feasible to distinguish between effects primarily caused by the ftsR deletion and secondary effects. Therefore, the strain background was changed for some of the following experiments from ATCC13032 to MB001. The in-frame deletion mutant MB001 AftsR as well as an *ftsR*-overexpressing strain constructed in this background were analyzed with respect to their growth behaviour and the moprohological phenotype. Both, growth behavior (Figure 14a) and cell morphology of MB001 $\Delta ftsR$ (Figure 15) were comparable to those of ATCC13032∆ftsR. Overexpression of ftsR in MB001 also caused similar effects as in ATCC13032 (Figure 14b). Thus, the absence of the prophages CGP1, CGP2, and CGP3 had no obvious influence on the effects of *ftsR* deletion and overexpression on growth and cell morphology.



Figure 14. Growth behavior of *C. glutamicum* **MB001 with deleted (a) or (over)expressed (b)** *ftsR.* The cells were first cultivated in BHI medium followed by two consecutive cultivations in CGXII minimal medium with 2% (w/v) glucose as carbon source. For the plasmid-based (over)expression, kanamycin (25 µg/mL) and IPTG was added as indicated. Means and standard deviations of three biological replicates of the second CGXII culture are presented.

4.10. Complementation of the MB001∆*ftsR* phenotype with native FtsR and homologs of *C. diphtheriae* and *M. tuberculosis*

In the complementation experiments performed with strain MB001 Δ ftsR, a full reversal of the growth defect and of the morphological changes was observed for MB001 AftsR pAN6-ftsR without IPTG induction (Figure 15). These results confirm that *ftsR* deletion causes the growth defect and the morphological changes of the corresponding mutants. Due to the conservation of the ftsR locus in different actinobacterial species, we assumed a similar function of the homologous genes and proteins. To test this hypothesis, we expressed *ftsR* homologs of two pathogenic relatives, CDC7B_1201 of Corynebacterium diphtheriae C7 and Rv1828 of Mycobacterium tuberculosis H37Rv, in the C. glutamicum MB001 AftsR strain using pAN6based expression plasmids and monitored growth and cell morphology. While this thesis was in preparation, initial biochemical and structural studies of Rv1828 were published (Singh et al., 2018), which will be discussed alongside our results. With pAN6-CDC7B_1201, almost full reversal of the growth defect and of the morphological changes was achieved by basal expression without IPTG induction (Figure 15). As previously observed for overexpression of C. glutamicum ftsR, stronger induction of CDC7B 1201 gene expression had a negative effect on growth (data not shown). With pAN6-rv1828, the best complementation was achieved in the presence of 100 µM IPTG (Figure 15). Whereas a full reversal to wild type morphology was observed, growth was improved with pAN6-rv1828 but did not reach the wild type characteristics. These results strongly suggest that the FtsR homologs of Actinobacteria have similar or identical functions and target genes.



Figure 15. Complementation of the growth defect (left panel) and the morphological phenotype (right panel) of the MB001 Δ *ftsR* mutant by plasmid-encoded FtsR and homologous proteins of related species. The cells were first cultivated in BHI medium with kanamycin (25 µg/mL) followed by two consecutive cultivations in CGXII minimal medium with kanamycin (25 µg/mL) and 2% (w/v) glucose as carbon source. MB001 Δ *ftsR* was transformed with pAN6 encoding *ftsR*, CDC7B_1201, or rv1828 under control of the leaky *tac* promoter. IPTG was only added for expression of rv1828, where 100 µM were required to obtain the best complementation from the tested conditions. Means and standard deviations of three biological replicates of the second CGXII culture are presented. Microscopy was performed with stationary cells. The white scale bar represents 5 µm.

4.11. Quantitative PCR to further investigate trehalose cluter amplification

A qPCR experiment was performed to confirm the amplification of the trehalose cluster in ATCC13032 Δ ftsR with an alternative method and to investigate whether this amplification did also occur in MB001 Δ ftsR. To do so, the genomic DNA of three independent clones of ATCC13032 Δ ftsR, four independent clones of MB001 Δ ftsR, and the respective wild type strains was isolated and used as templates. Two primer pairs were designed to specifically amplify two genes located in the trehalose cluster, cg0834 and cg0840. With a third primer pair, the reference gene *recF* was specifically amplified, serving as a control fragment. The results of the genome resequencing of ATCC13032 Δ ftsR were confirmed by qPCR, which revealed a \geq 5-fold increased DNA level for cg0834 and cg0840 compared to the reference gene *recF* (cg0005) (Figure 16). The amplification mechanism of this cluster is unknown and the exact genomic structure of the amplification cannot be deduced from the short sequencing reads. In MB001 Δ ftsR, qPCR did not reveal an increased DNA level of cg0834 and cg0840,

indicating that the amplification event only occurred in ATCC13032 Δ *ftsR*, but not in the prophage-free strain MB001 Δ *ftsR*. Since *ftsR* deletion mutants were constructed only once for each strain, firm conclusions on a functional correlation between the observed amplification event and the presence of the prophages cannot be drawn. However, the amplification could be responsible for the observed differences in the complementation studies between the wild type and the MB001 strain.



Figure 16. Normalized DNA levels of cg0834 and cg0840 in different *ftsR* deletion strains and the corresponding reference strains. Chromosomal DNA of three independent clones of ATCC13032 Δ *ftsR*, four independent clones of MB001 Δ *ftsR*, and the respective wild type strains was isolated and the relative copy numbers of were determined. Means and standard deviations of two technical replicates of each clone are presented. The DNA level of recF was used as reference and was set to 1. Illustrated are the normalized DNA levels of the genes cg0834 and cg0840 determined by qPCR and calculated using the $\Delta\Delta$ Ct quantification method with the program qPCRsoft 3.1.

4.12. Effect of *ftsR* deletion on *ftsZ* promoter activity and FtsZ distribution

To investigate a transcriptional activation of *ftsZ* expression by FtsR *in vivo*, a transcriptional fusion of the *ftsZ* promoter to a reporter gene encoding the mVenus protein was constructed resulting in plasmid pJC1-P_{ftsZ}-venus. The reporter plasmid and the parent plasmid pJC1-*venus*-term that served as negative control were transformed into strains MB001 and MB001 Δ *ftsR*. Subsequently, *ftsZ* promoter activity was measured as cell-specific fluorescence (defined as ratio of fluorescence and backscatter) in a BioLector[®] growth experiment. As shown in Figure 17, the lack of *ftsR* caused a significantly lower specific fluorescence, supporting an activation of *ftsZ* expression by FtsR. Comparable results were obtained with the strains ATCC13032 and ATCC13032 Δ *ftsR* carrying pJC1-P_{ftsZ}-venus (Figure 18).



Figure 17. Influence of the *ftsR* deletion in *C. glutamicum* MB001 on the expression of the *venus* reporter gene under control of the *ftsZ* promoter. The indicated strains were cultivated in CGXII medium with 2% (w/v) glucose and 25 μ g/mL kanamycin in a BioLector[®] system with automated measurement of cell density as backscatter and of Venus fluorescence. (a) For measurement of backscatter, mean values and standard deviations of three biological replicates are shown. (b) Specific fluorescence is shown as ratio of fluorescence and backscatter after 20 h of cultivation. The results of six individual biological replicates is represented as scatter dot plot with the black lines indicating the mean value and the error bars depicting the standard deviation.



Figure 18. Influence of the *ftsR* **deletion on the expression of the** *venus* **reporter gene under control of the** *ftsZ* **promoter.** The indicated strains were cultivated in CGXII medium with 1% (w/v) glucose and 25 µg/mL kanamycin in a BioLector[®] system with automated measurement of cell density as backscatter at 620 nm and fluorescence (excitation at 510 nm, emission at 532 nm). Specific fluorescence represents the ratio of fluorescence and backscatter. (a) Growth data for the ATCC13032 background. (b) Growth data for the MB001 background. (c) Comparison of the specific fluorescence of all strains after 24 h. Means and standard deviations of three biological replicates are shown.

To test whether the lack of FtsR influences the distribution of FtsZ within the cell, we generated derivatives of ATCC13032 and ATCC13032 Δ *ftsR* with a chromosomal insertion of a second copy of *ftsZ* fused in-frame to the coding sequence of the fluorescent protein mVenus under control of the native *ftsZ* promoter. Both strains were cultivated in CGXII minimal medium with glucose as carbon source and analyzed by fluorescence microscopy (Figure 19). In the strain lacking FtsR, the frequency as well as the intensity of the FtsZ rings was reduced. This may be caused by a reduced availability of FtsZ within the cells, which likely contributes to the morphological phenotype of the *ftsR* deletion mutant.



ATCC13032:: ftsZ-venus

ftsZ-venus

Figure 19. Fluorescence microscopy of the C. glutamicum strains ATCC13032::ftsZ-venus and ATCC13032*dftsR*::*ftsZ-venus* carrying a second chromosomal copy of *ftsZ* fused in-frame to the coding sequence of the fluorescent protein Venus. The cells were first cultivated in BHI + 2% (w/v) glucose and afterwards transferred to CGXII + 2% (w/v) glucose, both with 25 µg/mL kanamycin. Samples for microscopy were taken after about 6 h of cultivation of the main culture, which is approximately in the middle of the exponential growth phase. Two representative pictures for each

4.13. Genome-wide profiling of in vivo FtsR binding sites

culture are shown. The experiment was performed with two biological replicates each.

Transcriptional regulators usually bind to specific DNA sequences within the promoter regions of their target genes. With the aim to identify the in vivo binding site of FtsR in the ftsZ promoter region and to identify further target genes of FtsR, a ChAP-Seq experiment was performed. For this purpose, in vivo formaldehyde-crosslinked FtsR-Strep-DNA complexes were purified by StrepTactin affinity chromatography and the isolated DNA fragments were sequenced. Nine peaks with a sequencing coverage above 2000 were detected, which is at least 50-fold higher than the background noise signal observed for the entire genome with a 40-fold coverage (Figure 20, Figure 21, Table 9). The nine peaks were analyzed with respect to their chromosomal location and the mRNA level of the neighboring genes (Table 9). Based on the mRNA ratios observed in the transcriptome comparison of C. glutamicum ATCC13032 AftsR *versus* ATCC13032, three potentially FtsR-activated genes (*ftsZ*, cg0852, cg2477) and five potentially FtsR-repressed genes (*cop1*-cg3181-cg3180, *phoC*, cg0838) were identified using mRNA ratio (Δ *ftsR*/wild type) cutoffs of 1.5 and 0.75. Indeed, the peak with the third highest coverage value was located in the promoter region of the *ftsZ* gene (for a separate picture of this peak, see Figure 21). A repetition of the ChAP-Seq experiment confirmed the results of the first one except that the overall coverage was lower, and the order of the peaks varied slightly (Table 9).



Figure 20. DNA regions showing the highest coverage (red peaks) in the ChAP-Seq experiment with FtsR-Strep (strain *C. glutamicum* $\Delta ftsR$ pAN6-ftsR-Strep). As negative control, the DNA enriched by StrepTactin affinity chromatography from strain *C. glutamicum* $\Delta ftsR$ pAN6 was used (blue background). The red peak in the *ftsZ* promoter region (between cg2365 and cg2366), which had the 3rd highest coverage, is depicted in Figure 21. Due to the high peaks in the negative control of sample peaks 1 and 2, an independent ChAP-Seq experiment was performed, which confirmed binding of FtsR between cg2477 and cg2478 and between *cop1* and cg3185 (see Table 9) without the high background peaks in the negative control.

Table 9. Results of ChAP-Seq analysis with *C. glutamicum* ATCC13032 Δ *ftsR* pAN6-*ftsR*-Strep. The nine DNA regions that showed a \geq 50-fold higher coverage than the entire genome (coverage 40) in the ChAP-Seq analysis with FtsR-Strep were analyzed with respect to the location of the peak within the genome and the mRNA levels of the genes in this region. Based on that, a prediction was made which genes could be potential FtsR targets. The arrows indicate whether the genes are encoded on the leading (\rightarrow) or lagging strand (\leftarrow) within the genome. Graphical representations of selected peaks as derived from the ChAP-Seq analysis are shown in Figure 20.

Noª	Coverage ^a		Peak position		DNA mic 13032∆ft	roarray s <i>R</i> /13032		Potential <i>ftsR</i> target genes
		in gene	downstream 3'- end of	upstream of	Gene	mRNA ratio	p- value	
1 (1)	28,811 (4781)	-	cg2478 (NCgl2178, penicillin binding protein) ←	cg2477 (NCgl2177, conserved hyp. protein) ←	cg2478	0.75	0.11	cg2477 potentially activated by FtsR
					cg2477	0.24	0.01	
2 (5)	22,757 (934)	-	-	cg3182 (NCgl2777, <i>cop1</i> , tre- halose coryno- mycolyl transferase) ←	cg3182	1.70	0.20	<i>cop1</i> -cg3181- cg3180 potentially repressed by FtsR
				cg3181 (NCgl2776, putative secreted protein) ←	cg3181	2.30	0.05	
				cg3180 (NCgl2775, putative secreted	cg3180	1.91	0.05	
				cg3183 (NCgl2778, putative transpo-	cg3183	n.a.	n.a.	
3 (2)	4437 (1556)	-	cg2367 (NCgl2076, <i>fts</i> Q, cell division septal protein) ←	cg2366 (NCgl2075, <i>ftsZ</i> , cell division GTPase) ←	cg2367	1.20	0.02	<i>ftsZ</i> activated by FtsR
				,	cg2366	0.35	0.02	
4 (11)	3899 (373)	cg0470 (NCgl0381, <i>htaB</i> , heme transport- associated protein) \rightarrow	cg0469 (NCgl0380, hmuV) →	cg0471 (NCgl0382, <i>htaC</i> , heme- transport associated protein) \rightarrow	cg0469	1.41	< 0.01	-
					cg0470	1.45	0.09	
					cg0471	1.55	0.03	
5 (12)	3836 (365)	-	-	cg0852 (NCgl0712, conserved hypothetical protein) ←	cg0852	0.62	0.01	cg0852 potentially activated by FtsR
				cg0853 (NCgl0713, conserved hypothetical protein) →	cg0853	1.03	0.29	
				. /				(Continued)

Table 9. Continued.

No ^a	Coverage ^a		Peak position		DNA mic 13032∆ft	roarray s <i>R</i> /13032		Potential <i>ftsR</i> target genes	
		in gene	downstream 3'- end of	upstream of	Gene	mRNA ratio	p- value		
				cg0854 (NCgl0714, pmmA, phospho- manno- mutase)	cg0854	0.89	0.25		
				cg0855 (NCgl0715, conserved hypothetical	cg0855	1.18	0.08		
				cg0856 (NCgl0716, mannose-6- phosphate isomerase)	cg0856	1.09	0.03		
6 (3)	3630 (1529)	-	cg3392 (<i>oxiE</i> , <i>idhA2</i> , NCgl2958) → (myo-inositol 2- dehydro- genase)	cg3393 (NCgl2959, <i>phoC</i> , secreted cell wall-ass. phos- phatase) →	cg3392	1.25	0.03	<i>phoC</i> potentially repressed by FtsR	
				priataco)	cg3393	1.60	0.03		
7 (15)	2871 (315)	cg2695 (NCgl2368, ATPase of ABC trans- porter) ←	cg2697 (<i>ssb</i> , NCgl2370, single-stranded DNA-binding protein) ←	cg2694 (NCgl2367, putative phospho- diesterase, nucleotide	cg2693	1.31	0.03	-	
				pyrophos- phatase) ← cg2693 (NCgl2366, putative phospho- diesterase, nucleotide pyrophos- phatase) ←	cg2694	1.29	0.05		
					cg2695	1.06	0.42		
					cg2697	1.04	0.24		
8 (6)	2355 (858)	-	cg2518 (NCgl2213, put. secreted	-	cg2518	0.81	0.18	-	
			protein) → cg2519 (NCgl2214, conserved hyp.		cg2519	0.49	0.03		
			cg2520 (NCgl2215, hyp.		cg2520	0.92	0.45		
			cg2521 (NCgl2216, <i>fadD15</i> , long chain fatty acid CoA ligase) ←		cg2521	0.91	0.43		
9 (8)	2248	cg0839 (NCgl0701) ← (hyp. protein)	cg0840 (NCgl0702, conserved hyp.	cg0838 (NCgl0700, ATP-dep. belicase) ←	cg0838	4.61	0.03	cg0838 potentially repressed by EtsB	
		protein			cg0839	2.45	0.16	I GIN	
					cg0840	2.45	0.19		

The genes cg0852 and cg2477 encode conserved proteins of unknown function. The putative operon *cop1*-cg3181-cg3180 encodes three secreted proteins (Brand *et al.*, 2003). The Cop1 protein (also termed Csp1) was shown to function as mycolyltransferase involved in the conversion of trehalose monocorynomycolate to trehalose dicorynomycolate (Brand *et al.*, 2003, Puech *et al.*, 2000). The protein encoded by *phoC* was proposed to function as cell-wall-associated phosphatase (Schaaf & Bott, 2007, Wendisch & Bott, 2005) and is induced under phosphate-limiting conditions (Ishige *et al.*, 2003). The cg0838 gene is part of the DNA region found to be amplified in the ATCC13032 Δ *ftsR* mutant but had a higher mRNA ratio than the neighboring genes that were also amplified (Table 8), pointing to a possible repression by FtsR. The large Cg0838 protein (179 kDa) is proposed to function as an ATP-dependent helicase and contains a unique C-terminal domain including a metal-binding cysteine cluster.



Figure 21. Binding of the *ftsZ* **promoter region by FtsR.** Sequencing coverage of the *ftsZ* promoter after ChAP-Seq with FtsR-Strep. The threshold was set at a sequence coverage value of 120, which corresponds approximately to 3 times above background level. Blue arrows represent the coding regions of the respective genes. The negative control is presented below the red peak in blue and points downwards.

The search for a common DNA sequence motif in the nine sequences with a coverage above 2000 using the MEME software (Bailey *et al.*, 2009) revealed the motif shown in Figure 22. The motif was identified in eight of the nine sequences except for the sequence with a coverage of 2871. When looking at the corresponding motif in the *ftsZ* promoter region, it forms an imperfect 25-bp inverted repeat: AACCCTAAAGTAAAGGTTGAGGGTA.



Figure 22. FtsR consensus DNA-binding motif. The motif was identified by the MEME software (Bailey *et al.*, 2009) using the nine DNA regions with the highest coverage in the ChAP-Seq experiment with FtsR-Strep.

The center of this motif was located at position -73 with respect to the transcriptional start site of *ftsZ* as determined by RNA-Seq (Pfeifer-Sancar *et al.*, 2013). This position is compatible with an activating function of FtsR for *ftsZ* expression (Figure 23). It should be mentioned that in *C. glutamicum* strain ATCC13689, five transcriptional start sites were identified for *ftsZ* by primer extension and RACE studies (Letek *et al.*, 2007), none of which corresponds to the one identified by RNA-Seq for strain ATCC13032 (Pfeifer-Sancar *et al.*, 2013).

	CAC	ccc	CGGT	GG	GAA	CAG	TA	GAA	GTT	TCTG	GCG	CGG	GA T G.	ΑA	GGA	AACI	CA	GAAG	GTGC	CTTC	CTC	GCGC	STTAT	TG	CTG	TAA	TC	AACG	CAA	TTA
124	> T	P	P	v	G	т	v	E	v	S	G	A	D	E	G	N	S	Е	v	L	P	A	v	I	A	v	I	N	A	> I
	AAG	CGC	AAGA	ΤG	CGC	AGA	TG	ACA	GAA	AGTA	TCC	AGG	GTAG	ΤG	GAA	GCTC	CCG	GATO	CAAI	TTTG	ATA	ATCI	TGCI	GA	AAA	TGA	AT	GATO	GCC	GGG
154	K	A	Q	D	A	Q	M	Т	E	S	I	Q	V	v	E	A	P	D	Q	F	D	I	L	L	K	М	N	D	G	R
	AAA	TCT	ACTG	GG	GAI	CCI	CG	GAA	AAC	AACC	ACG	ATA	AAG	C G	GTG	GCAR	ATG	TCGA	ACTO	GTTT	TG	AAGO	CGGGA	AG	GCC	AAC	GT	TGGA	ACA	TTA
184	E	I	Y	W	G	S	S	E	N	N	Н	D	K	A	v	A	М	S	т	v	L	K	R	Е	G	Q	R	W	N	I
214	GCT > S	CAC 	сстс Р	AA 	TGC fts M	TGA Q V	CA T	GTC	CGC	TAAA .>> -	GT0	GCI	GGG	T A	GTT	CCGG	GTT	CACC	CCGI	TTC	CAG	CAAC	CGCGI	TT	GCC	CAC	СТ	CA <u>AC</u>	AAC	CCT
	AAA	GTA	AAGG	TT	GAG	GGT	A G	CGA	CAC	GCCG	CTO	TAA	ATG	т с	CGA	CACA	ATT	GACO	GTGI	TTT	TCO	GCAC	CAAGA	TA	TAT	GTC	AG	CCGG	ATT	AAC
	ACT	TTT.	AGAT	AA	GCI	CTC	AC	AGT	GCT	GACI	GCI	TAT	CTG	A G	ATG	AAAA	ATA	AAAG	CAA	AA	TTT	FTTI	TAGI	AT	CAA	CTC	TT	CGAA	AGG	CGA
	GAC	ATC	GACA	AT >>	GAC	стс	AC	CGA	ACA	ACTA		CGC	CAA	G A	TTA	AGGI	rCG ft:	TCGO	GCGI	rggg	CGG	GCGC	GCGGA	GT	CAA	CGC	CG	ТСАА	CCG	CAT
1					М	т	S	P	N	N	Y	L	A	K	I	K	V	V	G	V	G	G	GG	;	V	N .	A	V	N	R

Figure 23. DNA sequence of the *ftsZ* promoter region including parts of the coding regions of *ftsQ* and *ftsZ* and the corresponding amino acid sequences. The ribosome binding site of *ftsZ* is double underlined. The transcriptional start site identified by RNA-Seq (Pfeifer-Sancar *et al.*, 2013) in strain ATCC13032 is highlighted in magenta, the transcriptional start sites identified by primer extension and RACE in strain ATCC13689 (Letek *et al.*, 2007) are indicated in blue. The deduced -10 regions are shown in grey boxes. The FtsR-binding site identified in this work is highlighted in yellow. The 30-bp region used for EMSAs with purified FtsR-Strep (Figure 26) is underlined.

The DNA-binding motif shown above was used to search for similar motifs in the *ftsZ* promoter regions of other actinobacterial species possessing FtsR homologs using Clustal Omega (Sievers *et al.*, 2011). All analyzed promoters contained a similar motif (Figure 24a), supporting the assumption that regulation of *ftsZ* expression by FtsR homologs is a conserved mechanism in FtsR-containing *Actinobacteria*. The motif generated from the proposed binding sites also represents a 25-bp inverted repeat (Figure 24b), which is very similar to the one derived from the *C. glutamicum* sequences enriched by ChAP-Seq with FtsR (Figure 22).

a

C. glutamicum	CAACAACCCTAAAGTAAAGGTTGAGGGTAGCGA-162bp-CGACAATG	cg2366
C. diphtheriae	TCAAAACCCTAAAGTAAAGGTTTAGGGTCTAGA-163bp-TCTTAATG	DIP1595
M. tuberculosis	CTCT AAGCCTATGGTTGAGGGTTGAGAGTT TGCC -42bp-GAACG ATG	rv2150
R. erythropolis	CTCT AACCCTGTGGTTGAGGTTTAGAGTT TATC-102bp-AGCCG ATG	RER_35490
N. farcinica	CGCT AACCCTATGGTTGAGGTTCAGGGTT TTCC -32bp-AGCCC ATG	NFA_17690
T. paurometabola	CTTT AAGCCTGTGGTTGAGGTTTAGGGTT CTGG -35bp-GAAAC ATG	Tpau_2643
S. coelicolor	TGGT AACCCTAAACTTCAGCGTTAGGGTT CGGG -98bp-TCGAC GTG	SCO2082
N. multipartita	CTAT AACCGTATGGTTGAGGGTTGAGGGTT CGAC-115bp-GGCGA ATG	Namu_3798
P. acnes	TGAG TAGTCTCAAGCTACGGTTGAGGGTC AAGG -87bp-CTCCA GTG	PPA0761
A. cellulolyticus	CTATAAGCCTCTAGTTGAGGGTGAGGGTTGCGA -34bp-TGCAGATG	Acel_1012
A. mirum	ACACAACTCTTGACCCAGCGTCGAGGGTTTCAA -65bp-GACCGATG	amir_5760
M. luteus	CTTGGAGCCTCACGTGAAGGTTGATGCGGAGGC-184bp-ACACCGTG	mlut 13570



Figure 24. Putative FtsR binding sites in the *ftsZ* promoters of various actinobacteria. **a** Proposed FtsR binding sites in the *ftsZ* promoters of several actinobacterial species identified by sequence alignment with the FtsR binding site in the *C. glutamicum ftsZ* promoter. The respective *ftsZ* locus tags are given on the right. The proposed binding motifs are shown as colored letters. The annotated start codons of the FtsZ proteins are indicated by bold letters. The distance between the proposed FtsR binding site and the start codon varies from 41 to 193 bp. **b** Consensus DNA binding motif generated by MEME from the sequences shown in **a** using default parameters.

4.14. *in vitro* binding of purified FtsR to the proposed binding motif in the *ftsZ* promoter region

To test whether purified FtsR is able to bind to the proposed binding motif in the *ftsZ* promoter region, FtsR-Strep was overproduced in *C. glutamicum* ATCC13032 Δ *ftsR* using plasmid pAN6-*ftsR*-Strep and purified using StrepTactin-Sepharose (Figure 25a). As described above, FtsR-Strep could complement the growth defect of the ATCC13032 Δ *ftsR* mutant to the same extent as native FtsR, indicating that the Strep-tag did not negatively influence FtsR activity. Size-exclusion chromatography of affinity-purified FtsR-Strep (calculated mass 28.5 kDa) and comparison to standard proteins indicated that the protein forms a dimer (Figure 25b). This is in line with the dimeric state of the homologous protein Rv1828 (Singh *et al.*, 2018).



Figure 25. Purification of FtsR-Strep. (a) SDS-PAGE and Coomassie staining of purified FtsR-Strep. On the left side, the elution fractions 1-4 of the Strep-Tactin[®] affinity purification are shown. The second elution fraction was subjected to gel filtration for further purification (see Materials and Methods for details) and the analyzed fractions marked in the elution profile of the gel filtration (b) are shown on the right side of the gel. Gel filtration was performed using a SuperdexTM 200 Increase 10/300 GL column integrated into an ÄKTATM Pure25 system at a flow rate of 0.6 mL/minute (GE Healthcare Bio-Sciences AB, Uppsala, Sweden) and a buffer composed of 50 mM Tris-HCl, 250 mM NaCl, pH 7.5. The molecular mass of FtsR was estimated by comparison with standard proteins of known molecular mass and elution volume, which were cytochrome *c* (12.4 kDa, 18,82 mL), carbonic anhydrase (29 kDa, 16.15 mL), albumin (66.0 kDa, 13.73 mL), alcohol dehydrogenase (150.0 kDa, 12.53 mL), and β-amylase (200.0 kDa, 11.58 mL). FtsR-Strep eluted at 13.70 mL.

Purified FtsR-Strep was able to completely shift a 30-bp double-stranded oligonucleotide covering the predicted FtsR-binding site in the *ftsZ* promoter region at a 2-fold molar excess of the dimeric protein (Figure 26). In contrast, a control 30-bp double-strand-oligonucleotide derived from the promoter region of cg1081 was incompletely shifted even at an 8.5-fold molar excess of dimeric FtsR-Strep.



Figure 26. *in vitro* **DNA** binding studies with FtsR-Strep. Purified FtsR-Strep was incubated in the indicated concentrations with a constant concentration $(1 \ \mu M)$ of a 30-bp double-stranded oligonucleotide covering the predicted FtsR-binding site in the *ftsZ*-promoter region. The mixture was then analyzed by electrophoresis using a non-denaturing 15% (w/v) polyacrylamide gel. As negative control, a DNA fragment of the promoter region of cg1081 was used $(1 \ \mu M)$.

The specificity of this interaction was further tested using a competition-EMSA (Figure 27). A Cy3-labelled 271-bp DNA fragment covering the *ftsZ* promoter region including the FtsR binding site was incubated with FtsR protein and increasing concentrations of either specific (same fragment as above) or unspecific unlabeled competitor DNA. Only the specific competitor DNA reversed the shift of the labelled DNA fragment by FtsR, supporting its specific binding to the *ftsZ* promoter region.



Figure 27. EMSA competition experiment with FtsR and the *ftsZ* promoter region. 2 nM Cy3labeled *ftsZ* promoter DNA (271 bp) was incubated with 0 or 1.4 μ M FtsR dimer and increasing concentrations (0.47 μ M, 0.94 μ M, 1.86 μ M) of either unlabeled specific competitor DNA (same DNA fragment as above) or unlabeled unspecific competitor DNA (260-bp DNA fragment further upstream in the *ftsZ* promoter).

4.15. DNA affinity chromatography with the ftsZ promoter

As a complementary approach to show transcriptional control of *ftsZ* by binding of FtsR to the ftsZ promoter region, DNA affinity purification was performed using a 417-bp DNA fragment covering the 3'-end of the ftsQ coding region (36 bp), the intergenic region between ftsQ and ftsZ, and the first 96 bp of the ftsZ coding region. The immobilized DNA fragment was incubated with crude protein extract of C. glutamicum wild type cells grown in glucose minimal medium and harvested in the exponential growth phase. The proteins eluting after several washing steps were separated by SDS-PAGE and stained with Coomassie. Subsequent peptide mass fingerprinting using MALDI-ToF-MS resulted in the identification of eight proteins (Figure 28, Table 10). Six of them represent proteins that are often found in DNA affinity purifications with C. glutamicum cell extracts independent of the promoter region used. These are the single-stranded DNA-binding protein Ssb (Cg3307), the restriction endonucleases CgIIR (Cg1997) and CgIIIR (Cg1998), the ε-subunit of DNA polymerase III (Cg2321), DNA polymerase I (Cg1525), and the transcription termination factor Rho (Cg1354). Most interesting were the remaining two proteins, both of which are annotated as transcriptional regulators, Cg0444 (RamB) and Cg1631 (FtsR). The identity of these two proteins was further verified by MS/MS analysis of selected peptides (Table 10).



Figure 28. DNA affinity chromatography with the *ftsZ* **promoter region.** Crude cell extract of *C. glutamicum* ATCC13032 cultivated in glucose minimal medium to the mid-exponential growth phase was incubated with an immobilized 417-bp DNA fragment covering the *ftsZ* promoter region and strongly binding proteins were eluted with a high-salt buffer. Proteins enriched with the *ftsZ* promoter were separated by SDS-PAGE, stained with colloidal Coomassie, and identified by peptide-mass fingerprinting using MALDI-ToF-MS analysis (Table 10). The band with an apparent mass of about 27 kDa was identified as FtsR (M: marker; S: sample).

Table 10. MALDI-TOF-MS analysis of the proteins enriched by DNA affinity chromatography with the *ftsZ* promoter region. ^aThe following parameters were used for the database search using the in-house database of *C. glutamicum* ATCC13032 proteins based on the genome sequence: missed cleavages, 1; global modifications, carbamidomethyl (C); variable modifications: oxidation (M), mass tolerance: 80 ppm; MS/MS tolerance, 0.5 Da. ^bGiven are the masses of the peptides identified by MS/MS and the corresponding ion scores.

Protein	Annotated function	Calculated mass (Da)	Mascot scoreª	Number of peptides matched	MS/MS⁵ (ion score)	Sequence coverage
Cg1354	transcription termination factor Rho	83985	161	19		39%
Cg1525	DNA polymerase I	96769	248	30		44%
Cg1998	restriction endonuclease CgIIIR	71053	181	23		43%
Cg2321	DNA polymerase III epsilon subunit	51383	68	10		24%
Cg0444	transcriptional regulator RamB	54117	202	23	1400.7 (35) 1559.7 (37) 1774.8 (31)	52%
Cg1997	type II restriction endonuclease CgIIR	39915	104	12	· · · ·	47%
Cg1631	transcriptional regulator FtsR	27267	52	7	1103.5 (21) 1389.6 (23) 1566.8 (47)	30%
Cg3307	single-stranded DNA- binding protein Ssb	23287	46	4		30%

The regulator of acetate metabolism RamB (Cg0444) has already been studied quite extensively and represses genes involved in acetate metabolism and alleviates glucose and sucrose uptake (Gerstmeir *et al.*, 2004, Auchter *et al.*, 2011). A function in *ftsZ* regulation has not been described for RamB and a well conserved RamB-binding site (AA/GAACTTTGCAAA (Auchter *et al.*, 2011)) is absent from the *ftsZ* promoter region. Analogous to the experiment described in chaper 4.12 and Figure 17, *ftsZ* promoter activity was additionally tested in a

 $\Delta ramB$ background, but it was only very slightly altered in comparison to the wild type (data not shown). Enrichment of the second transcriptional regulator FtsR with the *ftsZ* promoter fragment supports its role in *ftsZ* regulation.

4.16. Analysis of the transcriptome of a *∆ftsR* mutant in the MB001 background

The transcriptome of a $\Delta ftsR$ mutant has already been analyzed in the ATCC13032 background. 52 genes were regulated, with 34 genes showing a ≥two-fold increased and 18 genes a ≥two-fold decreased mRNA level in the $\Delta ftsZ$ mutant (mean value of 3 biological replicates, p-value ≤0.05). To reduce the complexity of the results and to be able to distinguish between effects primarily caused by the *ftsR* deletion and secondary effects (e.g. phage related effects), a second DNA microarray experiment was performed with the *C. glutamicum* MB001 strain and the corresponding deletion mutant MB001 $\Delta ftsR$. Here, the number of regulated genes was reduced to a set of eight (Table 12), with three genes showing a ≥two-fold increased and five genes a ≥two-fold decreased mRNA level in the $\Delta ftsR$ mutant (mean value of four biological replicates, p-value ≤0.05).

Table 11: Transcriptome analysis of the ∆ftsR mutant. Genes with a ≥two-fold increased or a ≥two-
fold decreased mRNA ratio in the comparison MB001 Δ <i>ftsR</i> /MB001 and a p-value of \leq 0.05 are shown.
The results shown are mean values of four biological replicates. Additionally, the result for ftsZ is shown
in the last row.

Locus tag	Gene name	Annotated function	mRNA ratio (MB001∆ <i>ftsR/</i> MB001)	p- value
cg1631	ftsR	transcriptional regulator, MerR-family	0.003	0.0027
cg2477		hypothetical protein, conserved	0.055	0.0143
cg2080		hypothetical protein, conserved	0.471	0.0003
cg0470	htaB	secreted heme transport-associated protein	0.494	0.0002
cg2893		putative cadaverine transporter,	0.498	0.0185
		multidrug efflux permease, MFS-type		
cg3181		putative secreted protein	2.066	0.0042
cg3182	cop1	trehalose corynomycolyl transferase	2.118	0.0028
cg2378	mraZ	putative MraZ protein	2.195	0.0014
cg2366	ftsZ	cell division protein FtsZ	0.755	0.0023

The downregulated genes include, besides the deleted cg1631 (*ftsR*) itself, the two conserved hypothetical protein-encoding genes cg2477 and cg2080, the secreted heme transport-associated protein-encoding gene cg0470 (*htaB*), and the putative cadaverin transporter-encoding gene cg2839. Among the upregulated genes are cg3181 and cg3182 (*cop1*), which

code for a putative secreted protein and a trehalose corynomycolyl transferase, respectively. The strongest upregulation was observed for cg2378 (*mraZ*). In the promoter regions of the genes cg2477 and cg3182, also peaks have been detected during the ChAP-Seq experiment (Figure 20 and Table 9). Compared to the transcriptome analysis of the $\Delta ftsR$ mutant in the wild type background (significantly decreased *ftsZ* mRNA ratio of 0.35, Table 6), the mRNA ratio of the *ftsZ* gene did not fulfil the set criteria (≥two-fold increased or ≥two-fold decreased mRNA level, p-value≤0.05) in the MB001 background. However, it showed 25% decreased expression with a p-value of 0.002 in the $\Delta ftsR$ mutant, which is in the same range as observed in the reporter gene studies (Figure 17) and might be sufficient for causing the specified phenotype because of the particular importance of FtsZ for cell division processes.

4.17. FtsR-independent expression of FtsZ

Previous studies showed that differences in *ftsZ* expression cause aberrant cell morphology and FtsZ localization in *C. glutamicum* (Letek *et al.*, 2006). In order to test whether the altered cell morphology and the growth defect of the Δ *ftsR* mutant are solely caused by the differences in *ftsZ* expression, strains were constructed in which the expression of *ftsZ* is independent of FtsR and controlled by the gluconate-inducible *gntK* promoter of *C. glutamicum* (Letek *et al.*, 2006, Frunzke *et al.*, 2008). A DNA fragment containing a terminator sequence and the *gntK* promoter was inserted between the native *ftsZ* promoter and the ribosome binding site of *ftsZ* within the chromosomes of *C. glutamicum* MB001 and MB001 Δ *ftsR* by double homologous recombination (Figure 29).



Figure 29. Promoter exchange of *ftsZ*. Strains with FtsR-independent *ftsZ*-expression were constructed using a DNA fragment with a terminator sequence and the gluconate-inducible *gntK*-promoter, which was inserted between the native *ftsZ* promoter and the ribosome binding site of *ftsZ* on the chromosome of MB001 and the MB001 Δ *ftsR* mutant.

The promoter exchange strains MB001::P_{gntK}-ftsZ and MB001 Δ ftsR::P_{gntK}-ftsZ showed no growth in presence of sucrose (repressive conditions, Figure 30a), confirming that the terminator prevented sufficient transcription of *ftsZ* from its native promoter. Under standard conditions with 2% (w/v) glucose as carbon source (Figure 30b), also no growth was observed, which was inconsistent with the results of similar experiments previously performed by Letek *et al.* (2006). The difference might be caused by residual *ftsZ* expression due to readthrough from the original promoter in the previous studies, which is absent in our strains due to an

efficient terminator inserted behind the native promoter. With gluconate as carbon source (inducing conditions, Figure 30c) the promoter exchange strains were able to grow, however, the wild type background and $\Delta ftsR$ background strains showed strong growth differences, which was a first hint that FtsR must have additional targets besides *ftsZ*.



Figure 30. Growth of promoter exchange strains and their parental strains. The growth of the promoter exchange strains was tested in comparison to strains with the native *ftsZ* promoter using either sucrose (**a**), glucose (**b**), or glucose plus gluconate (**c**) as carbon source(s). Cells were pre-cultivated in either BHI medium (**a** and **b**) or BHI medium supplemented with 0.1% (w/v) gluconate for P_{gntK} induction when gluconate was also used as carbon source in the main culture (**c**), followed by two consecutive cultivations in CGXII minimal medium supplemented with the indicated carbon sources. Means and standard deviations of three biological replicates are presented.

In order to analyze the consequences of different *ftsZ* expression levels on growth and morphology and to find conditions where both MB001 and MB001 Δ *ftsR* with *ftsZ* under control of P_{gntK} show similar growth behavior, cells were cultivated in CGXII medium with 2% (w/v) glucose supplemented with 0.005, 0.01, 0.05, and 0.1% (w/v) gluconate (Figure 31). The changes in growth and morphology observed in this experiment nicely demonstrated the potential of fine-tuning of gene expression with the P_{gntK}-system.



Figure 31. Effect of different gluconate concentrations on cell morphology (a) and growth (b, c) of the promoter exchange strains MB001::PgntK-ftsZ and MB001\deltaftsR::PgntK-ftsZ. The two strains were first pre-cultivated in BHI medium supplemented with 0.1% (w/v) gluconate to induce ftsZ expression by PartK. The second pre-cultivation was performed in CGXII medium with 2% (w/v) glucose supplemented with the indicated gluconate concentrations. The main cultures were then performed in media having the same composition as the ones for the second pre-cultivation. (a) Microscopic pictures of cells from the stationary phase. The white scale bar represents 5 µm. (b, c) The growth experiments show mean values and standard deviations of three biological replicates.

Strain MB001::P_{gntK}-ftsZ showed enlarged cells with thickened poles at the lowest gluconate concentrations. This phenotype was reverted to wild type morphology at 0.1% (w/v) gluconate (Figure 31a). Strain MB001 Δ ftsR::P_{gntK}-ftsZ displayed strongly branched cells at the two lowest gluconate concentrations and even at 0.1% (w/v) gluconate the cells were still enlarged compared to the *ftsR*-positive strain (Figure 31a). With respect to growth, the two strains showed a quite similar behavior at the lowest gluconate concentration. Increased gluconate concentrations strongly improved growth of the *ftsR*-positive strain (Figure 31a), whereas growth of the *ftsR*-negative strain was only marginally increased (Figure 31c).

The most similar growth of the two promoter exchange strains was achieved with a precultivation in BHI complex medium supplemented with 0.1% gluconate (w/v) and a main culture in CGXII medium supplemented with 0.01% gluconate and 1.99% glucose (Figure 32). Under these conditions, the influence of growth differences is strongly reduced, and it should be possible to distinguish between effects resulting directly from the deletion of *ftsR* and those that occur due to altered levels of FtsZ. Therefore, these strains are suitable for subsequent experiments such as reporter gene studies.



Figure 32. Growth conditions enabling comparable growth of the promoter exchange strains. The growth of the promoter exchange strains was tested in comparison to the corresponding strains with the native *ftsZ* promoter. Cells were pre-cultivated in BHI medium supplemented with 0.1% (w/v) gluconate for P_{gntK} induction, followed by two consecutive cultivations in CGXII minimal medium supplemented with 0.1% (w/v) gluconate and 1.99% (w/v) glucose. Means and standard deviations for three biological replicates of the second CGXII culture are presented.

Overall, these results demonstrate the importance of an appropriate FtsZ level for normal cell morphology and growth, but they also show that the branched cells of the $\Delta ftsR$ mutant are not solely caused by reduced *ftsZ* expression, but presumably also by altered expression of other genes as a consequence of the *ftsR* deletion. Similarly, the growth defect of the $\Delta ftsR$ mutant cannot be rescued simply by increased expression of *ftsZ* and thus involves further genes. These could either be direct target genes of FtsR or genes whose expression is indirectly influenced by the absence of FtsR.

4.18. Influence of FtsR on *ftsZ* promoter activity in strains with FtsR-independent *ftsZ*-expression

The ftsR deletion in C. glutamicum MB001 led to a significant reduction of ftsZ promoter activity (Figure 17). In order to confirm that the observed activation of the *ftsZ* promoter by FtsR is independent of the actual *ftsZ* expression, the strains MB001::P_{antK}-ftsZ and MB001 Δ *ftsR*::Pantk-ftsZ were transformed with the reporter plasmid pJC1-P_{ftsZ}-venus and either pEC-ftsR or pEC-XC99E as empty plasmid control (Figure 33). With ftsZ being under control of P_{antk} and addition of the same gluconate concentration to the medium, expression of the chromosomal ftsZ should be identical for all strains. In plasmid pEC-ftsR, the ftsR gene is expressed under control of the IPTG-inducible but leaky trc promoter (Figure 33). The growth conditions were chosen according to the previous experiment where the mixture of 0.01% (w/v) gluconate and 1.99% (w/v) glucose led to comparable growth of strains MB001::Pantk-ftsZ and MB001 AftsR:: Pantk-ftsZ. The latter strain carrying pJC1-PftsZ-venus and the vector pEC-XC99E showed a significant growth defect in comparison to the other strains tested (Figure 33a, red curve). This defect must be due to altered expression of genes besides ftsZ that are regulated directly or indirectly by FtsR. Accordingly, the growth defect was reversed in the presence of the *ftsR* expression plasmid (Figure 33a, blue curve). The activity of the native ftsZ promoter on plasmid pJC1-P_{ftsZ}-venus was much lower in the strain lacking FtsR (Figure 33b, red curve) and was strongly increased when ftsR was expressed via pEC-ftsR (Figure 33b, blue curve). Growth and specific fluorescence were comparable for the two strains with chromosomal *ftsR* expression (Figure 33, black and green curves). These results confirm transcriptional activation of *ftsZ* expression by FtsR and that the phenotype of *ftsR* deletion mutants is not solely caused by a reduced *ftsZ* expression.



Figure 33. Influence of *ftsR* expression on (a) growth and (b) plasmid-based *ftsZ* promoter activity in the *ftsZ* promoter exchange strains MB001::P_{gntK}-*ftsZ* and MB001 Δ *ftsR*::P_{gntK}-*ftsZ*. The strains were transformed with pJC1-P_{*ftsZ*}-venus for monitoring *ftsZ* promoter activity and either with the *ftsR* expression plasmid pEC-*ftsR* or the vector pEC-XC99E. Cells were cultivated first in BHI complex medium supplemented with 0.1% (w/v) gluconate followed by a second pre-culture and the main culture in CGXII minimal medium containing 0.01% (w/v) gluconate and 1.99% (w/v) glucose as carbon source. Kanamycin (25 µg/mL) and chloramphenicol (10 µg/mL) were added to all cultures. Means and standard deviations for three biological replicates of the second CGXII culture are presented.

5. Discussion

5.1. Regulation of Odhl in *C. glutamicum*

In the last decades, it was shown that the regulation of the 2-oxoglutarate dehydrogenase complex (ODHC) plays a pivotal role in glutamate production, as its activity is strongly reduced during cultivation of C. gutamicum under glutamate-producing versus non-producing conditions (Kataoka et al., 2006, Shirai et al., 2005, Shimizu et al., 2003, Kawahara et al., 1997). Furthermore, it was demonstrated that ODHC, which marks a very important branch point in the metabolism of C. glutamicum, is regulated by the small inhibitor protein Odhl in order to control the flux of 2-oxoglutarate towards efficient glutamate production (Bott, 2007). Regulation of Odhl itself as inhibitor of ODHC by phosphorylation or succinylation of the lysine-132-residue has been reported previously (Komine-Abe et al., 2017). The initial aim of this doctoral thesis was to analyze whether Odhl is not only controlled at the posttranslational level, but also at the transcriptional level. The chance that odhl might be transcriptionally regulated seemed reasonable due to its location upstream of the two genes cg1631 (ftsR) and cg1633, which are both annotated to encode putative MerR-type transcriptional regulators. Since these two genes were suspected to regulate *odhl* expression, they were investigated in more detail. Growth studies revealed that the absence of cg1631 (later designated as *ftsR*) caused a severe growth defect under standard conditions and an even stronger growth defect was observed when this gene was overexpressed. Microscopy revealed a strong morphological phenotype with cells containing multiple septa both for the *ftsR* deletion mutant and for the overexpressing strain. This showed that the FtsR level is highly critical in C. glutamicum and hinted towards a possible function of FtsR in the regulation of cell division. No growth defect or altered phenotype was observed for the Acg1633 mutant under these conditions. In fact, cg1633 had already been investigated in more detail during a practical course of Graziella Bosco. Her studies suggested that cg1633 is most likely involved in metal homostasis and no evidence was found that odhl expression is affected by Cg1633 (Bosco, 2007). Therefore, cg1633 was no longer investigated within this doctoral thesis.

An alignment with FtsR homologs of different bacterial species revealed that most of the other sequences are considerably shorter than *C. glutamicum* FtsR, implicating that maybe the annotated start codon is incorrect. However, complementation studies with the annotated FtsR protein and with the shortened variant indicated that the annotation is correct since the shorter variant complemented the phenotype to a lower degree. Nevertheless, the growth defect of the $\Delta ftsR$ mutant was not fully restored even with the presumably correct FtsR variant, as it showed a longer lag phase and did not reach the final backscatter values obtained with the wild type, indicating that there might be secondary effects of the *ftsR* deletion which are eventually responsible for this effect.

FtsR was further investigated regarding a possible regulation of *odhl*. However, neither transcriptome analysis with DNA microarrays of the $\Delta ftsR$ mutant nor DNA affinity purification with the *odhl* promoter region did provide any support for the idea that FtsR might be involved in transcriptional regulation of *odhl*. Also, no other potential proteins which might regulate *odhl* expression were identified by DNA affinity purification, although the experiment *per se* was reliable, since proteins that bind DNA unspecifically and are often found in DNA affinity purifications (e.g. Ssb, polymerase subunits, topoisomerase etc.) were enriched. The results of this thesis indicate that *odhl* is not regulated on a transcriptional level. However, the experimental set-up concerning cultivation or reaction conditions might not have been optimal for the identification of a transcriptional regulator of *odhl*. Thus further experiments should be performed to confim this preliminary assumption.

5.2. Do secondary effects contribute to the $\Delta ftsR$ mutant phenotype?

The DNA microarray experiment with C. glutamicum ATCC13032 AftsR versus the wild type revealed a total of 52 genes which showed an at least two-fold altered mRNA ratio. It cannot be excluded that the deletion of *ftsR* might have downstream effects on other genes which secondarily cause or contribute to the $\Delta ftsR$ phenotype. One striking difference in the transcriptome of the C. glutamicum ATCC13032∆ftsR mutant was the increased mRNA level of a set of genes, ranging from cg0828 to cg0840, which is referred to as "trehalose cluster". This cluster harbors the genes cg0831-cg0835 encoding the trehalose uptake system TusFGK₂-E. Intriguingly, trehalose is a component of mycolates, which are important building blocks of the cell envelope of Corynebacteriales (Tropis et al., 2005). Moreover, the disaccharide is known as a reserve carbohydrate and for its function as a stress protectant in many organisms and has also been reported to be associated with several more biological processes like sporulation, germination, virulence, and morphogenesis (Tournu et al., 2013). It is known that trehalose synthesis can occur via different pathways. Besides the abovementioned TusFGK₂-E that allows uptake of exogenous trehalose, three major pathways involved in trehalose synthesis, OtsAB, TreYZ, and TreS, have been described for C. glutamicum (Ruhal et al., 2013, Tzvetkov et al., 2003, Wolf et al., 2003). Trehalose synthesis varies in utilization of the metabolic routes in different bacterial species. However, severe growth defects as a result of impairment of trehalose production appear to be universal (Tournu et al., 2013).

Moreover, it cannot be excluded that the increased expression of the genes of the trehalose cluster might influence the expression of other genes. To further pursue any possible secondary mutations, genome re-sequencing of three independent *C. glutamicum* ATCC13032 Δ *ftsR* mutants was performed. The four detected prominent SNPs are presumably

not responsible for causing the abovementioned effects because they led to silent mutations or are located in an intergenic region or a hypothetical CGP3 region. However, the genome resequencing revealed that the trehalose cluster was present in an increased copy number in all three independent $\Delta ftsR$ mutants, which was additionally shown in a qPCR experiment. This strongly suggests that the increased mRNA level of the trehalose cluster revealed by transcriptome analysis is likely caused by a gene dosis effect and not by an upregulation on the transcriptional level. It remains unclear what may have triggered the amplification of the trehalose gene cluster. However, tnp5a (cg0824) which is located upstream in close proximity of the cluster, encodes a transposase that could have been involved in the process. In the $\Delta ftsR$ background, tnp5a showed a significantly increased coverage in genome resequencing and a significantly elevated mRNA ratio in transcriptome analysis, indicating an amplification or increased activity of the transposase. The three transposases tnp16a (cq0292), tnp4a (cg2461), and *tnp5c* (cg3266) are not in the immediate vicinity of the cluster, but also show a significantly elevated mRNA ratio in transcriptome analysis. The causal relationship between transposable elements (TEs) and adaptation to several environmental stresses has been reported previously for a variety of prokaryotes and eukaryotes. The molecular mechanisms underlying TE-induced mutations as well as the consequences of TE activation are highly diverse (Casacuberta & González, 2013). In the context of this doctoral thesis, it can be speculated whether the transposase tnp5a or one of the other transposases confers the trehalose cluster the ability to respond to stress through its amplification. However, the amplification could also have occured for other reasons or purely by chance and has not occurred in the MB001 background.

For trehalose being known as stress protectant and due to its role in cell envelope biosynthesis, the increased copy number of *tusFGK*₂-*E* observed for *C. glutamicum* ATCC13032 Δ *ftsR* could indicate a stress response to counteract the morphological phenotype of the Δ *ftsR* mutant. An imaginable consequence of the amplification is that the increased expression of *tusFGK*₂-*E* leads to an increased trehalose uptake capacity. However, the absorbed trehalose could only result from the degradation of for example trehalose mycolates of the cell envelope, since trehalose was not added to the medium. This increased uptake capacity could possibly allow a more efficient recycling of the trehalose split off from cell wall components. Additionally, the significantly decreased mRNA ratio of the trehalose corynomycolyl transferase encoding gene *cmt2* (cg3186) in the Δ *ftsR* mutant observed by transcriptome analysis could support this hypothesis, as Cmt2 is involved in trehalose dicorynomycolate synthesis.
5.3. The switch to MB001 as background strain

In addition to the abovementioned possibilities of secondary downstream effects of a ftsR deletion, it seemed reasonable that also phage-related effects might be triggered by stress induced by the ftsR deletion. Due to this, the background strain was switched from the ATCC13032 type strain to the MB001 strain for some experiments with the purpose to make it more feasible to distinguish between effects primarily caused by the ftsR deletion and secondary effects. The deletion and the overexpression of *ftsR* in the MB001 background led to the same severe growth defect and morphological phenotype as in the ATCC13032 background strain. Transcriptome analysis and the qPCR experiment to investigate the amplification of the trehalose cluster has also been performed with strain MB001 Δ ftsR. Neither increased expression nor amplification of the trehalose gene cluster were detected in the DNA microarray and in the qPCR experiment. This vitiated the previously elaborated "trehalose cluster hypothesis", since the same phenotype as for the ATCC13032 background was observed. So, it could no longer be assumed that the elevated copy number of the trehalose gene cluster is responsible for the observed phenotype, or it is at least not the main cause for the phenotype in the ATCC13032 background. Overall, the set of significantly regulated genes in DNA microarray experiments with deleted *ftsR* was strongly reduced in the MB001 background compared to the ATCC13032 strain, most likely due to the elimination of trehalose cluster amplification and abolished phage-related effects, which facilitates the analysis of FtsR in this strain. Moreover, in contrast to the ATCC13032 background, the complementation experiment worked readily in the MB001 background, as the growth as well as the morphology was completely restored by plasmid-based ftsR expression. This indicates once more that potential secondary effects of a *ftsR* deletion were strongly reduced or were no longer present in MB001∆ftsR.

5.4. FtsR, the first transcriptional regulator of FtsZ identified for the *Corynebacteriales* order

As already mentioned, the severe growth defect and drastic morphological changes caused by *ftsR* deletion and overexpression hinted to the encoded regulator playing a pivotal role in cell division. Because there is hitherto no transcriptional regulator of FtsZ known for *C. glutamicum* and the regulation of cell division in the *Corynebacteriales* order itself is largely unknown, the main focus of this work was shifted towards the investigation of this previously uncharacterized protein, FtsR. The obtained results revealed that FtsR is a transcriptional regulator of the tubulin-like GTPase FtsZ in *C. glutamicum* and the first transcriptional regulator of the key player of cell division discovered in this organism. Moreover, the conserved genomic locus in different *Corynebacteriales* species and successful complementation experiments of the $\Delta ftsR$ phenotype with FtsR homologs of *C. diphtheriae* and *M. tuberculosis* support the assumption that FtsR fulfils a similar function in related organisms. Antibiotics research in recent years has shown that the inhibition of bacterial cell division is a promising strategy against human pathogens, with particular focus on FtsZ itself. Most compounds identified so far mainly aim at its GTPase activity, leading to an impaired assembly of the Z-ring and therefore to cell death (Sass & Brötz-Oesterhelt, 2013). However, this antibiotic class is already subject to the emergence of resistance mechanisms. Contributing to this is the fact that most new antibiotics are merely variants of substances already in use, so that bacteria can adapt to them rapidly, which highlights the urgent need for antimicrobial therapies that follow new mechanisms or attack novel targets (Lock & Harry, 2008). The newly identified regulator FtsR could therefore serve as an interesting candidate for the development of new antimicrobial drugs against these pathogenic relatives of *C. glutamicum*.

5.5. The importance of fine-tuning

The overexpression experiments with increasing amounts of IPTG as well as the experiments with varying gluconate concentrations when *ftsZ* is under control of the *gntK* promoter showed strikingly the importance of accurate *ftsZ* expression for the maintanence of normal growth and morphology of *C. glutamicum*. Furthermore, evidence for the activation of *ftsZ* expression by FtsR came from transcriptome analysis, where the *ftsZ* mRNA level was reduced in the $\Delta ftsR$ mutant, and reporter gene studies showing reduced activity of the *ftsZ* promoter in the $\Delta ftsR$ background, representing a novel control mechanism of actinobacterial cytokinesis. In ATCC13032 $\Delta ftsR$ and MB001 $\Delta ftsR$, the *ftsZ* mRNA level was decreased by 65% and 25%, respectively. The latter reduction by a fourth seems to be minor, however, this slightly altered *ftsZ* expression might still be responsible for the impaired growth and severe morphological phenotype due to the particular importance of FtsZ concerning the cell division process.

5.6. FtsR's mode of action and binding site

Bioinformatic analyses revealed that FtsR belongs to the MerR superfamily of transcriptional regulators, which mostly function as activators, responding to a vast diversity of stimuli including oxidative stress, xenobiotics, and metal ion excess, with the latter forming a particular subgroup in this family of regulators (McEwan *et al.*, 2011, Brown *et al.*, 2003). Structures of already characterized MerR-type regulators show that they have very similar conformations, usually consisting of an N-terminal DNA-binding helix-turn-helix domain and a structurally more diverse effector-binding domain located in the C-terminus, which are linked by a helical domain forming an antiparallel coiled-coil required for dimerization (Kumaraswami *et al.*, 2010, Newberry & Brennan, 2004). MerR-type regulators typically bind as homodimers to inverted

repeats in the promoter region of their target genes. The binding sites are often located in the spacer regions between the -10 and -35 elements (Brown et al., 2003). However, there are also other cases reported where for example the recognition site is located upstream of the -35 region or where a C-terminal effector-binding region is missing or larger than usual, such as for the Gram-positive Lactobacilli, Clostridia, and Bacillus subtilis or for members of the Gram-negative genus Neisseria (Counago et al., 2016, Kidd et al., 2005, Ahmed et al., 1995). The spacer regions of MerR target promoters are often elongated, consisting of 19-20 bp instead of 16-18 bp, which hampers transcription because the -10 and -35 regions are located on opposite sides of the DNA strand (Brown et al., 2003). A well-described mechanism of transcriptional activation by regulators of the MerR superfamily is assumed to rely on a conformational change of the N-terminal DNA-binding region of the protein after recognition of a stimulus by the C-terminal effector binding domain, which causes a twist of the bound DNA molecule in a manner that the -10 and -35 regions are rearranged, enabling RNA polymerase to bind and initiate transcription (Philips et al., 2015, Newberry & Brennan, 2004, Ansari et al., 1992, Frantz & O'Halloran, 1990). FtsR eluted as dimer during gel filtration and the binding site identified in the *ftsZ* promoter region was found to be an inverted repeat, which was located at position -61 to -85 upstream of the *ftsZ* transcriptional start site identified by RNA-Seq (Pfeifer-Sancar et al., 2013) rather than between the -10 and -35 regions. This position is compatible with an activating function of FtsR but might involve a different mechanism of transcriptional activation than the one described above. Binding of FtsR to the *ftsZ* promoter was further confirmed by DNA affinity chromatography with the *ftsZ* promoter region. In vivo binding of FtsR to the ftsZ promoter was confirmed by ChAP-Seq experiments and purified FtsR was shown to bind to a 30 bp-DNA fragment covering the proposed DNA binding site of FtsR in EMSAs, which represents an imperfect 25-bp inverted repeat.

It should be mentioned that in the *C. glutamicum* strain ATCC13689 five transcriptional start sites were identified for *ftsZ* by RACE and primer extension experiments (Letek *et al.*, 2007), none of which corresponds to the one identified by RNA-Seq for the strain ATCC13032 (Pfeifer-Sancar *et al.*, 2013). Further studies are required to test for the presence of additional transcriptional start sites of the *ftsZ* gene in strain ATCC13032, in which the FtsR binding site might be located between the -10 and -35 regions. Multiple promoters contributing to *ftsZ* expression appear to be present in various bacteria such as *M. tuberculosis* (Roy & Ajitkumar, 2005, Kiran *et al.*, 2009), *E. coli* (Flardh *et al.*, 1997), *Streptomyces coelicolor* (Hajduk *et al.*, 2016, Flardh *et al.*, 2000), or *Streptomyces griseus* (Kwak *et al.*, 2001). They allow a dynamic adjustment of the FtsZ levels according to different stimuli and probably make the cells more robust with respect to spontaneous mutations in the *ftsZ* promoter region. Hitherto, an effector molecule which might trigger binding of FtsR remains to be discovered, as attempts for its identification by different GC-MS approaches were unsuccessful ((i) analysis after extraction

of fatty acid methyl esters from purified FtsR to detect a fatty acid as putative effector molecule, (ii) analysis after trypsin digestion of FtsR to detect a released effector molecule); data not shown.

FtsR was found to be conserved in many different families of Actinobacteria, including the genus Mycobacterium. In a ChIP-Seg experiment with the FtsR homolog Rv1828 of *M. tuberculosis*, the promoter region of the *ftsZ* gene was the top target (Galagan *et al.*, 2010, Minch et al., 2015, Reddy et al., 2009). Using the proposed binding motif of FtsR (AACCCTAAAGTAAAGGTTGAGGGTA) as template, we identified a very similar sequence (AACtCTAAgcctAtGGTTGAGGtTt) in the *ftsZ* promoter of *M. tuberculosis*, the region which also showed by far the highest coverage in the ChIP-Seq experiment (Minch et al., 2015). Here, the motif was located exactly between the -10 and -35 regions of the transcriptional start site P1, which is located at position -43 relative to the *ftsZ* translational start (Kiran *et al.*, 2009). The binding motif proposed for Rv1828 in a recent study is different but overlaps partially with the motif proposed in this thesis (Singh et al., 2018). Mutational analysis of this binding motif by EMSAs revealed that binding of Rv1828 is not influenced when the specific part of the motif is altered that does not overlap with our proposed binding motif (Singh et al., 2018). This, together with the strong conservation among many species, suggests that the motif proposed in this thesis is more likely to be correct. The results obtained for FtsR taken together with the earlier findings for and recently published characterization of Rv1828 (Singh et al., 2018) strongly support the notion that FtsR and its actinobacterial homologs are involved in transcriptional regulation of ftsZ. In line with this, the ftsZ promoter regions of various actinobacterial genera were found to contain DNA sequence motifs similar to the ones determined for FtsR and Rv1828.

5.7. FtsR must have additional targets besides FtsZ

The studies with the promoter exchange strains in which *ftsZ* expression was controlled by the gluconate-inducible *gntK* promoter revealed that growth behavior and morphology of the $\Delta ftsR$ mutant are not solely caused by altered *ftsZ* expression. Despite comparable *ftsZ* expression, the $\Delta ftsR$ strain grew much worse than the *ftsR*-positive reference strain and the morphology of the $\Delta ftsR$ strain differed from that of the *ftsR*-positive reference strain, in particular by the formation of branched cells. This suggests that (i) FtsZ alone is not responsible for the branched cell morphology, and (ii) that there must be some other, FtsR-regulated gene(s) that cause(s) enlarged cells even when FtsZ expression is fine-tuned by sufficient gluconate concentrations.

Potential further target genes of FtsR were deduced by combining ChAP-Seq and transcriptome results. The target with the most obvious relation to the altered cell morphology

of the $\Delta ftsR$ mutant was the *cop1*-cg3181-cg3182 gene cluster. These three genes encode two secreted proteins and Cop1 (Csp1), a mycolyltransferase converting trehalose monocorynomycolate to trehalose dicorynomycolate. This could possibly be linked to the morphological phenotype of $\Delta ftsR$ because of corynomycolic acids being important building blocks of the corynebacterial cell wall (Dover *et al.*, 2004). Together with the genera *Mycobacteria* and *Norcadia, Corynebacteria* belong to the mycolata group, whose cell envelope differs from other Gram-positive bacteria, as it additionally contains (coryno)mycolic acids, which are covalently linked to the peptidoglycan layer by arabinogalactan, surrounding the cell wall like an outer membrane, which provides resistance to antimicrobial compounds and other stress conditions (Favrot & Ronning, 2012, Hett & Rubin, 2008, Dover *et al.*, 2004).

The mycolyl transferase function was assigned to the N-terminal portion of Cop1 (Puech *et al.*, 2000). A *C. glutamicum cop1* deletion mutant had an altered cell morphology characterized by enlarged, club-shaped cells and it was speculated that the C-terminal part of Cop1 plays a role in cell shape formation (Brand *et al.*, 2003). In the case of *cop1*, the FtsR-binding site (AAGTCTAAAGTTGAACTTAAGATTG) starts downstream of the -35 region and ends downstream of the -10 region (TAAGAT). The transcriptome analysis data from DNA microarray experiments with MB001 Δ *ftsR* suggest a repression of *cop1* by FtsR and the morphology of the cells overexpressing *ftsR* resemble that of cells lacking functional *cop1* (Brand *et al.*, 2003). The peak located in the promoter region of *cop1* had the second highest coverage value in the ChAP-Seq experiment. The highest coverage value was obtained for the peak in the promoter region of cg2477, which encodes a putative protein that has not been characterized so far. The expression of cg2477 was significantly altered in ATCC13032 Δ *ftsR*, which makes both genes noticeable putative candidates for FtsR targets. DNA affinity purification experiments with promoter regions of *cop1* and cg2477 could be performed to further investigate this hypothesis.

Another interesting gene with significantly altered expression in the DNA microarray experiments with the MB001 Δ ftsR strain is mraZ. It was shown in previous studies that the encoded protein is a transcriptional repressor of ftsEX in *C. glutamicum* and that its overproduction causes an elongated cell shape (Maeda *et al.*, 2016, Eraso *et al.*, 2014). FtsEX are components of the early divisome forming an ATP-binding cassette transporter, with FtsE representing the ATPase and FtsX the membrane domain of the complex (Donovan & Bramkamp, 2014). The genomes of *C. glutamicum* and *M. tuberculosis* contain homologs of FtsEX, but relatively little is understood regarding their function. Different functions of FtsEX are proposed for different organisms: in *E. coli*, FtsEX regulates peptidoglycan hydrolases and location of FtsX in the middle of the cell is achieved by interaction of FtsE with FtsA and FtsQ, whereas FtsX interacts directly with FtsZ (Corbin *et al.*, 2007, Schmidt *et al.*, 2004); in *B. subtilis*, FtsEX is involved in spatio-temporal regulation of sporulation and is located around

the cell membrane (Garti-Levi *et al.*, 2008); in *B. subtilis* and *Streptococcus pneumonia*, FtsEX mediates coupling of peptidoglycan hydrolysis and cell division and it was shown to regulate the activity of a cell wall hydrolytic enzyme, CwlO (Domínguez-Cuevas *et al.*, 2013, Meisner *et al.*, 2013, Sham *et al.*, 2013); in *C. crescentus*, FtsEX is speculated to play a role in tethering of Z-rings to the membrane, because it is recruited to the nascent division site prior to FtsA, which is involved in membrane association of the Z-rings (Goley *et al.*, 2011). With the knowledge previously generated for these other bacteria and considering that for *C. glutamicum* and *M. tuberculosis* FtsA has not been described so far (Donovan & Bramkamp, 2014), it can be speculated that FtsEX plays a role in the formation of the Z-ring and the stabilization of the future cell division site, which in turn is regulated by MraZ-mediated repression. In the $\Delta ftsR$ mutant, the mRNA level of *mraZ* was elevated, which suggests that FtsR not only directly regulates FtsZ, but is also involved in the complex process of cell division by repressing MraZ and thus controlling FtsEX. However, in the DNA microarray experiments with the $\Delta ftsR$ mutant, no significantly altered expression of *ftsEX* was observed, suggesting that an effector may be needed to activate the repressive function of MraZ.

The presented results of the ChAP-Seq and transcriptome analysis as well as the proposed binding motif for FtsR suggest a direct influence of the regulator on the expression of the genes *cop1* and *mraZ*. However, further studies are required to investigate the exact mode of regulation by FtsR on these targets and to identify further candidate genes and their functions. Beyond that, it might also be possible that FtsR does not only play a role in cell division, but also in other yet undiscovered processes, which do not necessarily have to contribute to the described phenotype.

5.8. The physiological function of FtsR

Regarding the physiological function of FtsR, a possible role could be to serve as a regulatory mechanism that controls cell division in response to the nutritional status of the cell. In recent years, several studies have revealed metabolic enzymes to directly influence cell division by modulating the activity of FtsZ (Monahan & Harry, 2016). Transcriptional regulation, as shown here for FtsR, is another means to influence FtsZ activity. The fact that FtsR activates *ftsZ* expression suggests that FtsR signals favorable nutritional conditions with a high cell division rate and thus a high FtsZ demand. As the *ftsR* gene and its homologs in other Actinobacteria are always located downstream of the *odhl/garA* gene, a functional link to Odhl/GarA might exist. Odhl/GarA controls the metabolic flux at the 2-oxoglutarate node of the central metabolism, which is particularly relevant for nitrogen assimilation. Our preliminary attempts to find a potential ligand of FtsR were not successful and the identification of the stimulus controlling FtsR activity is ultimately required to understand the function of this regulator.

6. References

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Ich versichere an Eides Statt, dass die Dissertation von mir selbständig und ohne unzulässige fremde Hilfe unter Beachtung der "Grundsätze zur Sicherung guter wissenschaftlicher Praxis an der Heinrich-Heine-Universität Düsseldorf" erstellt worden ist.

Ort, Datum

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