A TatABC-Type Tat Translocase Is Required for Unimpaired Aerobic Growth of Corynebacterium glutamicum ATCC13032

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Abstract

The twin-arginine translocation (Tat) system transports folded proteins across the cytoplasmic membrane of bacteria and the thylakoid membrane of plant chloroplasts. Escherichia coli and other Gram-negative bacteria possess a TatABC-type Tat translocase in which each of the three inner membrane proteins TatA, TatB, and TatC performs a mechanistically distinct function. In contrast, low-GC Gram-positive bacteria, such as Bacillus subtilis, use a TatAC-type minimal Tat translocase in which the TatB function is carried out by a bifunctional TatA. In high-GC Gram-positive Actinobacteria, such as Mycobacterium tuberculosis and Corynebacterium glutamicum, tatA, tatB, and tatC genes can be identified, suggesting that these organisms, just like E. coli, might use TatABC-type Tat translocases as well. However, since contrary to this view a previous study has suggested that C. glutamicum might in fact use a TatAC translocase with TatB only playing a minor role, we reexamined the requirement of TatB for Tat-dependent protein translocation in this microorganism. Under aerobic conditions, the misassembly of the Rieske iron-sulfur protein QcrA was identified as a major reason for the severe growth defect of Tat-defective C. glutamicum mutant strains. Furthermore, our results clearly show that TatB, besides TatA and TatC, is strictly required for unimpaired aerobic growth. In addition, TatB was also found to be essential for the secretion of a heterologous Tat-dependent model protein into the C. glutamicum culture supernatant. Together with our finding that expression of the C. glutamicum TatB in an E. coli ΔtatB mutant strain resulted in the formation of an active Tat translocase, our results clearly indicate that a TatABC translocase is used as the physiologically relevant functional unit for Tat-dependent protein translocation in C. glutamicum and, most likely, also in other TatB-containing Actinobacteria.

Introduction

The transport of proteins into or across biological membranes is catalyzed by membrane-bound, multi-component protein translocases. In most bacteria, the major route of protein export is the
TatB, TatC, (and also sometimes TatE) can be found. A notable difference to the situation in *um tuberculosis* components are expressed from a *coli* and other Gram-negative bacteria with TatABC-type Tat translocases, where the three Tat *tatB* that is located elsewhere on the chromosome [24]. Nevertheless, the presence of all three *tatC* are present in an operon and, in contrast to the situation mentioned above, the positive bacteria with high-GC-content genomes are differently organized. Here, only general secretion (Sec) pathway [1] that translocates its substrates as unfolded polypeptide chains [2]. In contrast, the alternative twin-arginine translocation (Tat) pathway transports fully folded proteins across the membrane [3]. Tat substrates are often proteins that have to recruit a cofactor in the cytosol and, therefore, must acquire their folded status prior to their export [4]. The signal peptides of Tat substrates contain a highly conserved twin-arginine motif (S/T-R-R-X-F-L-K) that has been shown to be important for the productive recognition and binding of the Tat substrates by the Tat translocase [5–7].

The Tat pathway has been most extensively studied in *Escherichia coli* and, in this organism (and also most other diderm-lipopolysaccharide (LPS) [8–10] Gram-negative bacteria), the respective Tat translocase is a TatABC-type translocase consisting of three inner membrane proteins, TatA, TatB, and TatC, that jointly are expressed from an operon [11]. Another tat gene, *tatE*, is located elsewhere in the genome of *E. coli* and encodes a paralogue of TatA. Due to the fact that *tatE* is expressed at a level 50–200 times lower than *tatA* [12], *tatE* is commonly regarded as a cryptic gene duplication of *tatA*. After folding and, if required, cofactor insertion, proteins containing a Tat signal peptide are recognized by a substrate receptor complex consisting of TatB and TatC [7, 13]. Subsequently, homo-oligomers of TatA are recruited to the substrate-loaded receptor complex in a proton motive force (pmf)-dependent manner [14], followed by substrate translocation across the inner membrane in a step that is still poorly understood. Whereas some models propose that TatA multimers form a substrate-fitted protein-conducting channel (e.g. [15]), an alternative model suggests that the TatA multimers might lead to a punctual weakening of the membrane in the vicinity of the TatBC-substrate complex and thereby allows a TatC-driven translocation of the substrate without the need of an actual translocation channel [16]. In TatABC-type Tat translocases, TatA and TatB carry out distinct functions and cannot substitute for each other [17].

A different scenario is found in monoderm [8–10] Gram-positive bacteria with low-GC-content genomes, such as the well-studied model organism *Bacillus subtilis*. In these bacteria, TatB is lacking and Tat-dependent protein translocation is performed by TatAC-type minimal Tat translocases in which the TatB function is carried out by a bifunctional TatA subunit [18–22]. However, in Gram-positive bacteria with high-GC-content genomes such as members of the phylum Actinobacteria (as exemplified by the monoderm [8–10, 23] bacteria *Streptomyces coelicolor* and *Streptomyces lividans*, and the diderm-mycolate [8–10, 23] bacteria *Mycobacterium tuberculosis* and *Corynebacterium glutamicum*), genes encoding homologues of TatA, TatB, TatC, (and also sometimes TatE) can be found. A notable difference to the situation in *E. coli* and other Gram-negative bacteria with TatABC-type Tat translocases, where the three Tat components are expressed from a tatABC operon, is the finding that the *tat* genes in the Gram-positive bacteria with high-GC-content genomes are differently organized. Here, only *tatA* and *tatC* are present in an operon and, in contrast to the situation mentioned above, the *tatB* gene is located elsewhere on the chromosome [24]. Nevertheless, the presence of all three *tat* genes in the respective genomes suggests that this class of bacteria, like *E. coli*, might use TatABC-type translocases as functional units of Tat-dependent protein translocation. A strong support for this view comes from investigations of the Tat system of *M. tuberculosis* which, in contrast to the Tat system of most other organisms tested so far, is essential for viability. Disruption of *tatA*, *tatC*, and importantly also of *tatB* on the chromosome of *M. tuberculosis* was not possible, strongly indicating that each Tat component is of equal importance (as typically is the case for TatABC-type Tat translocases) and arguing against the possibility of a TatAC translocase being the functional unit of the Tat system in this microorganism [25]. In line with these findings, results from the Actinobacteria *S. lividans* [26–27] and *S. coelicolor* [28–29] also showed that *tatB* mutants of these microorganisms are similarly affected in Tat-dependent protein translocation as their respective *tatA* or *tatC* mutant strains. A somewhat different situation
was described by Kikuchi et al. [30] for the Tat system of *C. glutamicum*. Also in *C. glutamicum*, a tatAC operon and a tatB gene located at different positions on the chromosome can be identified. Additionally, *C. glutamicum* also possesses a tatE gene located at another position in the genome that, like the situation in *E. coli*, presumably is a cryptic gene duplication of tatA [30]. The Tat system is not absolutely essential for the viability of *C. glutamicum* since all tat genes can be deleted. However, ΔtatA and ΔtatC mutant strains show drastic growth defects compared to the wild-type strain, indicating that the mislocalization of a physiologically important Tat substrate is responsible for this phenotype. Deletion of tatE had no noticeable effect on the growth of *C. glutamicum*, supporting the view that TatE is functionally redundant to TatA and does not play a significant role also in this microorganism. Strikingly however and in stark contrast to the ΔtatA and ΔtatC mutant strains, growth of a ΔtatB deletion mutant was described as being almost identical to that of the tat wild-type strain. Based on these findings it was concluded that TatB is dispensable and that *C. glutamicum* uses a TatAC-type minimal translocase as the basic functional unit for Tat-dependent protein translocation [30].

The major branch of the aerobic respiratory chain of *C. glutamicum* involves the cytochrome *bc*$_1$-aa$_3$ super-complex that includes the Rieske iron-sulfur component QcrA [31]. In Actinobacteria, the Rieske proteins are polytopic integral membrane proteins with three transmembrane segments. For the Rieske protein of *S. coelicolor*, it has been shown that the Sec and the Tat translocases cooperate in the assembly process [32]. The amino-terminal domain encompassing the first two transmembrane segments is integrated into the cytoplasmic membrane via the Sec system. Subsequently, the third transmembrane segment which corresponds to a non-cleavable internal Tat signal peptide mediates the Tat-dependent translocation of the fully folded carboxyl-terminal iron-sulfur cofactor-containing domain to the trans-side of the cytoplasmic membrane [32–33]. Deletion of the genes encoding the cytochrome *bc*$_1$ complex (i.e. *qcrABC*) causes a severe growth defect of *C. glutamicum* under aerobic conditions [31, 34]. It is therefore a likely possibility that the similarly severe growth defect caused by the absence of a functional Tat system might in fact be due to the mislocalization of the iron-sulfur domain of the Rieske protein QcrA.

Since the results described for the Actinobacteria *M. tuberculosis*, *S. lividans* and *S. coelicolor* are contradictory to the results described for *C. glutamicum* with respect to the suggested types of Tat translocases that operate in the respective organisms, we reexamined the role of TatB in the Tat-dependent protein translocation in *C. glutamicum* with a focus on the proper localization of the Rieske iron-sulfur protein QcrA. Evidence is presented that the mislocalization of QcrA is in fact a major reason for the growth defects of *C. glutamicum* strains defective in Tat-dependent protein translocation. Furthermore, our results make it very likely that, besides TatA and TatC, also TatB is required for efficient QcrA assembly and, as a consequence, for normal growth. In addition, the presence of TatB was found to be essential for the secretion of a heterologous Tat-dependent model protein. Taken together, our combined results clearly indicate that a TatABC-type translocase is the functional unit for Tat-dependent protein translocation in *C. glutamicum* and, most likely, also in other TatB-containing Actinobacteria.

### Results

Deletion of *tatAC*, *tatA/E*, or *tatB* abolishes Tat-dependent protein translocation and causes a severe growth defect in *C. glutamicum*

In Tat(A)EBC-type translocases, as exemplified by the *E. coli* Tat translocase, Tat(A), TatB, and TatC perform distinct mechanistically relevant functions and cannot substitute for each other. Mutant strains possessing deletions in either of the corresponding tat genes are therefore expected to possess similar phenotypes that ultimately are caused by a non-functional
Tat export machinery. To gain more insight in the type of Tat translocase that is used in the Actinobacterium *C. glutamicum*, we compared different *tat* deletion mutants with respect to their growth phenotypes and their ability to translocate a Tat-dependent model protein across the cytoplasmic membrane. Besides the previously described Δ*tatAC* mutant strain [35], a Δ*tatB* and a Δ*tatA/E* mutant were additionally constructed as described in the Materials and Methods section.

First, growth of *C. glutamicum* wild-type under aerobic conditions in BHIS medium was compared to the growth of Δ*tatAC*, Δ*tatA/E*, and Δ*tatB* mutant strains in 48-well FlowerPlates using a BioLector micro reactor cultivation device. In the BioLector, a high oxygen transfer rate and therefore growth under fully aerobic conditions can be achieved. Furthermore, in this device, the growth of bacterial strains can be determined online via light scatter measurements, thereby allowing a real time monitoring of biomass formation in each of the 48 growth chambers of the FlowerPlate [36]. As shown in Fig 1, the Δ*tatAC* and Δ*tatA/E* mutant strains showed severe growth defects when compared to the unaltered wild-type strain, being fully in line with
the results published by Kikuchi et al. [30] for ΔtatA and ΔtatC mutant strains. Strikingly however and in stark contrast to the previously published findings [30], we observed an identically severe growth defect also for the ΔtatB mutant strain. The growth defect of the ΔtatB mutant could almost completely be reverted by expressing TatB in trans from plasmid pEC-TatBCG, but not by introducing the pEC-X99E empty vector as control (Fig 2), confirming that the growth defect of the ΔtatB strain is indeed caused by the lack of TatB and not due to effects caused by the tatB deletion on the neighboring genes on the chromosome.

Next, Tat-dependent protein translocation was analyzed using a previously described model protein (PhoDCG-GFP [35]) consisting of the signal peptide of the C. glutamicum Tat substrate PhoD (an alkaline phosphatase) fused to green fluorescent protein (GFP). As shown in Fig 3, large amounts of mature GFP are present in the supernatant of C. glutamicum wild-type containing plasmid pCGPhoDCG-GFP (lane 4). Furthermore, hardly any detectable GFP-derived polypeptides were present in the cellular fraction of the same strain (lane 3). In contrast, no secreted GFP protein is found in the supernatant of the ΔtatAC, ΔtatA/E, or ΔtatB mutant strains.
and, in the cell fractions of the corresponding strains (lanes 5, 7, 9), an accumulation of PhoDCG-GFP precursor and degradation products of it can be observed. These results clearly indicate that TatA, TatB, and TatC are equally important for the translocation of PhoDCG-GFP across the cytoplasmic membrane of *C. glutamicum*.

The Rieske iron-sulfur protein QcrA is an important Tat substrate in *C. glutamicum*

The Rieske iron-sulfur protein QcrA is a component of the cytochrome *bc₁-aa₃* super-complex which constitutes the major branch of the aerobic respiratory chain of *C. glutamicum* [31]. Like other actinobacterial Rieske proteins, the *C. glutamicum* QcrA protein possesses three transmembrane segments. Preceding the third transmembrane segment, an internal Tat consensus motif (G-RR-K-L-I) can be identified suggesting that, like the situation in *S. coelicolor* [32], the C-terminal iron-sulfur cluster-containing domain might be translocated to the trans-side of the cytoplasmic membrane in a Tat-dependent manner. Furthermore, since deletion of the genes encoding the cytochrome *bc₁* complex (including *qcrA*) causes a severe growth defect to *C. glutamicum* [37], we thought that it is a likely possibility that the severe growth defect of the *C. glutamicum* tat mutants is in fact caused by the mislocalization of QcrA.

To experimentally address these assumptions, we first compared the growth of a *C. glutamicum* wild-type strain possessing the pEKEx2 empty vector with the growth of pEKEx2-containing mutant strains *C. glutamicum* ΔqcrA and *C. glutamicum* ΔtatB using the BioLector micro-reactor cultivation device. As shown in Fig 4, compared to *C. glutamicum* wild-type, both mutant strains show a severely impaired growth under aerobic conditions. Notably, the growth behavior of both mutant strains is almost identical, a finding that would be consistent with mislocalization of QcrA being the actual cause for the growth defect associated with *C. glutamicum* tat mutant strains.
To directly show that the correct localization of the QcrA iron-sulfur domain requires the Tat translocase, *Corynebacterium glutamicum* ΔqcrA was transformed with plasmids pEKEx2-QcrA (encoding unaltered QcrA), or pEKEx2-QcrAKK (encoding a mutated QcrA protein in which the two conserved twin arginine residues of the Tat-motif in front of the third transmembrane segment are replaced by a twin lysine pair). Replacement of the twin-arginines by a pair of lysines has been shown previously to significantly reduce or even completely abolish Tat-dependent translocation for a variety of different Tat substrates [38–39]. As shown in Fig 4, the growth defect of the ΔqcrA mutant could be almost completely reverted by expressing unaltered QcrA in trans from plasmid pEKEx2-QcrA, but not by the introduction of the pEKEx2 empty vector as the negative control. Importantly, no reversion of the growth defect of *C. glutamicum* ΔqcrA was observed when the QcrAKK variant was expressed from plasmid pEKEx2-QcrAKK, indicating that the Tat system is involved in the translocation of the iron-sulfur domain of the Rieske iron sulfur protein QcrA also in *C. glutamicum*.

As described in the previous paragraph, the growth defect of a *C. glutamicum* ΔqcrA mutant could almost completely be reverted by in trans expression of plasmid-encoded QcrA. Next,
we asked whether or not such an *in trans* complementation of the *qcrA* deletion in the chromosome can still be observed when the TatB component is missing. For this, a Δ*qcrA* Δ*tatB* double mutant was constructed and transformed with plasmid pEKEx2-*QcrA* or the pEKEx2 empty vector, respectively. In contrast to the situation found for the Δ*qcrA* single mutant strain, no reversion of the growth defect was observed in the Δ*qcrA* Δ*tatB* double mutant when *QcrA* was expressed from pEKEx2-*QcrA* *in trans* (Fig 5). One feasible, and in our opinion the most likely explanation for the observed results is that the presence of TatB is required for the correct membrane assembly of *QcrA* in *C. glutamicum*. However, at present we cannot completely rule out the possibility that, besides *QcrA*, one or more additional Tat substrate(s) exist whose mislocalization together has an identically severe negative impact on the growth of *C. glutamicum* as the mislocalization of *QcrA* alone and, if so, that TatB could specifically function in the correct localization of these additional proteins. In any case, our results nevertheless clearly demonstrate that the presence of TatB is of crucial importance for the correct localization of at least

**Fig 5.** The growth defect of a *C. glutamicum* Δ*qcrA* Δ*tatB* double mutant cannot be complemented *in trans* by plasmid-encoded *QcrA*. Cells were inoculated to an OD<sub>600</sub> of 0.5 in 750 μl BHIS medium containing 100 μM IPTG and cultivated in 48-well FlowerPlates in a BioLector system for 48 h at 30°C, 1100 rpm under constant 85% relative humidity. Growth of the respective *C. glutamicum* strains, as indicated in the figure, was monitored as backscattered light (620 nm; signal gain factor 20) in 15 min intervals. The growth curves show one representative experiment of three independent biological replicates. Standard deviations are given for 10 selected time points. a.u.: arbitrary units.

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one physiologically important Tat substrate and, as a consequence, for unimpaired growth of *C. glutamicum* under aerobic conditions.

**Heterologous expression of the *C. glutamicum* TatB in *E. coli* tat mutant strains**

*E. coli* tat mutants possess pleiotropic cell envelope defects that are caused by the mislocalization of two Tat-dependent periplasmic amidases (AmiA and AmiC) that are involved in cell wall turnover [40]. Due to this fact, the *E. coli* tat mutants are sensitive to the presence of sodium dodecyl sulfate (SDS), unless they are complemented with a *tat* gene that allows for the formation of a functionally active Tat translocase [41]. To investigate whether the *C. glutamicum* TatB can indeed perform the specialized functions associated with the *E. coli* TatB, we tested whether the *C. glutamicum* TatB can complement the *E. coli* ΔtatB mutant strain BØD [17]. Following the protocol established by Ize et al. [40], the respective cells were grown in the presence of various SDS concentrations and the % survival of each strain was determined for the SDS concentrations tested (Fig 6). In contrast to the MC4100 wild-type strain (black squares), the uncomplemented ΔtatB mutant (green circles) is highly sensitive to SDS. As expected, SDS resistance of BØD could be restored by complementation with plasmid-encoded *E. coli* TatB (pink triangles). Likewise, restoration of SDS resistance was observed when the *C. glutamicum* TatB was expressed in BØD (purple asterisks), clearly demonstrating that the heterologous TatB can fulfill the dedicated function of the *E. coli* TatB protein. In contrast, expression of the *C. glutamicum* TatB from plasmid pHSG-TatBCG in the ΔtatA/E mutant JARV15 [11] did not result in the formation of a functional Tat translocase, since the corresponding strain (red open pentagons) showed the same SDS sensitivity as the uncomplemented ΔtatA/E mutant (blue diamonds). These results strongly suggest that the *C. glutamicum* TatB protein in fact is a bona fide TatB protein that is functionally equivalent to the TatB proteins present in TatABC-type Tat translocases.

**Discussion**

In the present study, we have reexamined the requirement of TatB for Tat-dependent protein translocation in *C. glutamicum*. Our results suggest that TatB very likely is strictly needed for the correct inner membrane assembly of a physiologically important Tat substrate (i.e. the Rieske iron-sulfur protein QcrA). In addition, we have found that TatB, like TatA and TatC, is essential for the secretion of a Tat-dependent heterologous model protein (PhoDCG-GFP) into the culture supernatant. Our findings strongly support the view that a TatABC-type translocase operates in *C. glutamicum* and that the individual respective translocase subunits perform the same distinct functions as their corresponding subunits in the well characterized prototype TatABC translocase of *E. coli*.

So far, it was unknown what causes the severe growth defect of *C. glutamicum* strains that lack a functional Tat translocase. Our results clearly show that the mislocalization of the carboxyl-terminal iron-sulfur cofactor-containing domain of the QcrA Rieske protein is at least one of the major reasons for the poor growth of *C. glutamicum* tat mutants. Such a conclusion is supported by several findings: First, the *C. glutamicum* ΔqcrA mutant and all *C. glutamicum* mutant strains that lack a functional Tat translocase show an almost identical growth defect under the fully aerobic conditions in the BioLector micro reactor cultivation device. Second, the replacement of the twin-arginine residues of the internal Tat motif preceding the third transmembrane segment by a twin-lysine pair completely abolished the ability of plasmid-encoded QcrA to complement a ΔqcrA mutant strain, being strong evidence that the translocation of the carboxyl-terminal iron-sulfur domain to the trans-side of the plasma membrane is a
Tat-dependent step. Finally, only the growth defect of a ΔqcrA single mutant, but not that of a ΔqcrA ΔtatB double mutant could be complemented by plasmid-encoded QcrA, strongly sup-
porting the view that a functional Tat system is required for correct QcrA membrane assembly.

However, despite our data show that the defective main branch of the respiratory chain is a
major reason for the severe growth defect of C. glutamicum tat mutants, we cannot completely
exclude the possibility that the mislocalization of (an)other Tat substrate(s) also has negative
impacts on growth that is overshadowed by the dominant effect caused by the mislocalization
of QcrA.

In principle, the iron-sulfur cofactor-containing domain of a bacterial QcrA protein can be
exported also by a natural bona fide TatAC minimal Tat translocase, as has recently been dem-
onstrated for B. subtilis [42–43]. For C. glutamicum, our results however indicate that, under
fully aerobic conditions, a complete TatABC translocase is required for QcrA assembly and,
based on this, for unimpaired growth. Our suggestion that TatB very likely is required for QcrA localization is, to a great extent, based on our finding that a ΔqcrA ΔtatB double mutant cannot be complemented by plasmid-encoded QcrA. However, a hypothetical scenario that might have misled us with respect to such a conclusion cannot completely be excluded and is discussed in what follows. In such a scenario, besides QcrA one or more physiologically important Tat substrates would exist whose mislocalization has an identically severe negative impact on growth as the mislocalization of QcrA. Furthermore, such a scenario would require that both negative effects on growth, the one caused by the mislocalization of QcrA and the one caused by the mislocalization of the additional hypothetical Tat substrate(s), are not additive since, in the QcrA-complemented ΔqcrA ΔtatB double mutant, not even a partial improvement of its growth was observed in comparison to the uncomplemented strain. Our conclusion with respect to the TatB requirement for correct QcrA localization would only be incorrect if, against all expectations, QcrA would be special in that it solely relies on TatA and TatC for its membrane assembly, whereas in contrast the localization of the additional hypothetical physiologically important Tat substrate(s), and also the secretion of the heterologous TorA/PhoDCG-GFP model proteins, strictly requires the presence of TatB (since a tatB mutant shows the same severe growth defect as a tatAC mutant and no TorA/PhoDCG-GFP is secreted by the tatB mutant). To the best of our knowledge, such an extreme variation in the utilization of Tat components between different Tat substrates or, in other words, the co-existence and substrate-dependent differential use of a TatAC and a TatABC translocase, has never been observed in any TatABC-containing microorganism. With that said, we think that it is very likely that TatB (along with TatA and TatC) is required for proper QcrA assembly.

With respect to the effect of a tatB deletion on the growth of *C. glutamicum*, our results which show that TatB is of equal importance as TatAC for growth under aerobic conditions significantly differ from the results of Kikuchi et al. [30] which reported that deletion of tatA or tatC, but not of tatB resulted in a strong growth defect of the corresponding *C. glutamicum* tat mutant strains. The different behavior of the two ΔtatB mutants with respect to growth is also manifested in their different ability for the secretion of GFP model proteins. All our tat mutant strains tested (i.e. ΔtatB, ΔtatAC, and ΔtatA/E) showed a complete defect in the secretion of PhoDCG-GFP (Fig 3) and TorA-GFP (S1 Fig) into the culture supernatant. In contrast, a significant secretion of TorA-GFP (i.e. 9% of the wild-type strain level) into the culture supernatant of their ΔtatB mutant strain was observed by Kikuchi et al. [30]. Taken together, it seems that the two ΔtatB mutant strains possess different residual activities with respect to the Tat-dependent translocation of a native protein (i.e. QcrA) and also of heterologous model proteins (i.e. TorA-GFP/PhoDCG-GFP), respectively.

We can only speculate about the possible reason(s) for the observed differences. The TatA protein of *C. glutamicum* ATCC13869, the parent strain used in the study of Kikuchi et al. [30], is 12 amino acid residues longer than the TatA protein of *C. glutamicum* ATCC13032 which was used as the parent strain in our present study. The two TatA proteins show 88% identity in the overlapping regions and some of the amino acid differences can be found in the amino-terminal region [30]. Since previous results have shown that *E. coli* TatA can be converted by even a single amino acid alteration within the amino-terminal region into a bifunctional TatA protein that can take over the function of TatB to a significant degree [19], one possible explanation might be that the TatA proteins from the two *C. glutamicum* strains differ with respect to their evolutionary remnant bifunctionality. If so, then the *C. glutamicum* ATCC13869 ΔtatB mutant might possess a higher residual transport activity than *C. glutamicum* ATCC13032 ΔtatB because its TatA protein can more efficiently compensate for the absence of TatB compared to its *C. glutamicum* ATCC13032 counterpart. Another point that should be kept in mind is that our parent strain *C. glutamicum* ATCC13032 is an untreated *C.
glutamicum wild-type strain [44]. In contrast, the actual strain that has been used in the study of Kikuchi et al. [30] for the construction and subsequent characterization of the various tat deletion mutants was YDK010, an industrial C. glutamicum ATCC13869 derivative that has been extensively mutagenized by N-methyl-N-nitro-N-nitrosoguanidine (MNNG) followed by a screening for improved secretory production of proteins [45]. An alternative explanation therefore might be that in YDK010 a bifunctional TatA protein (TatA') has been created during MNNG mutagenesis of the ATCC13869 parental strain that, due to an MNNG-induced mutation, bypasses to some degree the essential requirement of TatB for unimpaired growth under aerobic conditions. Besides this, also mutations in one of the other remaining tat genes in the YDK010ΔtatB mutant (i.e. tatE or tatC) that cause a suppression of the growth defect of the YDK010ΔtatB mutant strain can also not be completely excluded.

Finally, our finding that the C. glutamicum TatB can functionally complement an E. coli ΔtatB, but not an E. coli ΔtatA/E mutant strongly suggests that the C. glutamicum TatB is in fact a bona fide TatB protein that, together with TatA(E) and TatC performs a distinct function in a TatABC-type translocase also in its native surrounding. Together with results from other Actinobacteria containing a tatAC operon and an additional tatB gene located elsewhere on the chromosome, such as M. tuberculosis [25], S. lividans [26–27], or S. coelicolor [28–29, 41], the combined data of our present study for C. glutamicum clearly strengthen the view that a TatABC-type translocase is the physiologically relevant functional unit for Tat-dependent protein translocation in most, if not all, members of this special class of microorganisms.

Materials and Methods

Bacterial strains, media, and growth conditions

The bacterial strains used in this study are listed in Table 1. E. coli strains were grown at 37°C in LB (lysogeny broth) medium [46]. The complementation of E. coli tat mutants by plasmid-encoded Tat proteins was analyzed in LB medium containing 0.1%, 0.5%, 1%, 2%, or 4% (w/v) sodium dodecyl sulfate (SDS). C. glutamicum strains were grown in BHIS medium, containing 37 g/L brain heart infusion (Difco) and 91 g/L sorbitol, at 30°C. If required, isopropyl-ß-D-thiogalactopyranoside (IPTG) was added at concentrations of 10 μM or 100 μM, as indicated. Antibiotic supplements were at the following concentrations: kanamycin 25 μg/ml (C. glutamicum), chloramphenicol 25 μg/ml (E. coli) and 12.5 μg/ml (C. glutamicum), respectively.

Growth curves of C. glutamicum strains were determined in 48-well FlowerPlates (mp2-labs, Aachen/D) using the BioLector micro reactor cultivation device (m2p-labs, Aachen/D) that allows the online monitoring of biomass formation [36]. 5 ml BHIS medium were inoculated with a fresh colony of the respective strains from a BHIS agar plate and grown overnight as the first preculture at 30°C and 170 rpm. Subsequently, 20 ml BHIS medium in a 4-chicane Erlenmeyer flask were inoculated from the first preculture to an OD600 of 0.5 and grown overnight at 30°C and 120 rpm. The OD600 of the respective second precultures was measured and equal amounts of cells from the different strains were centrifuged and subsequently resuspended in fresh BHIS medium to an OD600 of 0.5. 750 μl of the respective cultures were transferred into the wells of the 48-well FlowerPlates that were subsequently cultivated for 48 h in the BioLector at 30°C, 1100 rpm under constant 85% relative humidity. The production of biomass was determined as the backscattered light intensity of sent light with a wavelength of 620 nm (signal gain factor 20). Measurement intervals were set to 15 min. In each of the experiments shown in Figs 1, 2, 4, and 5, the growth curves of all strains that are shown in the respective figure were determined in biologically independent triplicates on the same FlowerPlate.
Table 1. Bacterial strains and plasmids used in this study.

| Strains or plasmids       | Relevant properties | Source or reference |
|---------------------------|--------------------|---------------------|
| **E.coli** strains        |                    |                     |
| MC4100                    | F' araD139 Δ(argF-lac)U169 rpsL150 relA1 fruA25 | [55]               |
| BØD                       | MC4100 ΔtatB        | [17]               |
| JARV15                    | MC4100 ΔtatA ΔtatE |                     |
| **C.glutamicum** strains  |                    |                     |
| ATCC13032                 | wild-type          | [44]               |
| ATCC13032 ΔtatAC          | ΔtatAC             | [35]               |
| ATCC13032 ΔtatB           | ΔtatB              | This study          |
| ATCC13032 ΔtatE           | ΔtatE              | This study          |
| ATCC13032 ΔtatA/E         | ΔtatA/E            | This study          |
| ATCC13032 ΔqcrA           | ΔqcrA              | This study          |
| ATCC13032 ΔqcrA ΔtatB     | ΔqcrA ΔtatB        | This study          |
| **E. coli** plasmids      |                    |                     |
| pHSG575                   | pSC101 replicon, lacZα+ CmR | [54]               |
| pHSG-TatBEC              | pHSG575 containing the E. coli tatB gene | This study          |
| pHSG-TatBCG              | pHSG575 containing the C. glutamicum tatB gene | This study          |
| **C. glutamicum** plasmids|                    |                     |
| pEKE2x                    | C. glutamicum/E. coli shuttle vector for regulated gene expression, pBL1 oriV<sub>E. coli</sub> P<sub>lac</sub>, lacI<sub>R</sub>, Km<sup>R</sup> | [52] |
| pCGP<sub>DG</sub>GFP     | pEKE2x containing a gene encoding PhoD<sub>DG</sub>-GFP         | [35]               |
| pCGTorA-GFP              | pEKE2x containing a gene encoding TorA-GFP         | [35]               |
| pEKE2x-QcrA              | pEKE2x containing the C. glutamicum qcrA gene       | This study          |
| pEKE2x-QcrA<sub>OK</sub> | pEKE2x-QcrA (R159K,R160K)                          | This study          |
| pEC-XC99E                | pGA1 mini replicon, P<sub>trc</sub>, lacI<sub>R</sub>, Cm<sup>R</sup> | [53]               |
| pEC-TatB<sub>C</sub>     | pEC-XC99E containing the C. glutamicum tatB gene   | This study          |
| pK19mobsacB              | Vector for allelic exchange in C. glutamicum, pK18 oriV<sub>E. coli</sub>, oriT, mob, sacB, lacZα, Km<sup>R</sup> | [49]               |
| pK19mobsacBΔqcrA         | pK19mobsacB containing the flanking regions of C. glutamicum qcrA | This study          |
| pK19mobsacBΔtatE         | pK19mobsacB containing the flanking regions of C. glutamicum tatE | This study          |
| pK19mobsacBΔtatA         | pK19mobsacB containing the flanking regions of C. glutamicum tatA | This study          |
| pCRD206                  | pCGR2 replicon (ts), sacB, Km<sup>R</sup>             | [51]               |
| pCRD206ΔtatB             | pCRD206 containing the flanking regions of C. glutamicum tatB | This study          |

aKm<sup>R</sup>, kanamycin resistance; Cm<sup>R</sup>, chloramphenicol resistance.

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**Strain constructions**

Routine methods such as PCR, DNA restriction, and ligation were performed using standard protocols [47]. Oligonucleotides used as PCR primers are listed in S1 Table. Chromosomal DNA of C. glutamicum was prepared as described previously [48]. The construction of C. glutamicum ATCC13032 ΔtatAC has been described previously [35]. C. glutamicum ATCC13032 ΔtatA/E was constructed by deleting tatE and tatA in a stepwise manner using the pK19mobsacB system which carries the counter-selectable marker sacB [49]. First, a deletion of tatE in C. glutamicum ATCC13032 wild-type was constructed by generating two DNA fragments containing approximately 480 bp of upstream and downstream sequences of the tatE gene using chromosomal DNA of C. glutamicum ATCC13032 as template together with primers del_tatEup-f and del_tatEup-r or primers del_tatEdn-f and del_tatEdn-r, respectively. Both PCR
products were purified and used as a template in a cross-over PCR reaction using primers del_tatEup-f and del_tatEdn-r to generate a fused DNA fragment that contained the up- and downstream sequences of tatE, but lacks the tatE structural gene. The corresponding PCR fragment was digested with Sall and BamHI and ligated into Sall/BamHI-digested pK19mobsacB, yielding pK19mobsacBtatE. For deletion of the chromosomal tatE structural gene, pK19mobsacBtatE was introduced into C. glutamicum ATCC13032 by electroporation [50]. Cells that had integrated the plasmid into the chromosome via homologous recombination were selected on plates containing kanamycin. A second homologous recombination event leading to the loss of the sacB gene (via excision of the integrated plasmid) was positively selected on BHIS agar plates containing 10% sucrose. Colonies were analyzed for the presence or absence of the tatE gene by colony PCR using primers deltaE-f and deltaE-r. One of the isolates that contained the desired chromosomal deletion of the tatE structural gene was designated C. glutamicum ATCC13032 ΔtatE. Next, a deletion of tatA in C. glutamicum ATCC13032 ΔtatE was constructed by generating two DNA fragments containing approximately 520 bp upstream and downstream sequences of the tatA structural gene using chromosomal DNA of C. glutamicum ATCC13032 as template together with primers del_tatAup-f and del_tatAup-r or primers del_tatAdn-f and del_tatAdn-r, respectively. Both PCR products were purified and used as a template in a cross-over PCR reaction using primers del_tatAup-f and del_tatAup-r to generate a fused DNA fragment that contained the up- and downstream sequences of tatA, but lacks the tatA structural gene. The corresponding PCR fragment was digested with Xbal and EcoRI and ligated into Xbal/EcoRI-digested pK19mobsacB, yielding pK19mobsacBΔtatA.

For deletion of the chromosomal tatA structural gene, pK19mobsacBΔtatA was introduced into C. glutamicum ATCC13032 ΔtatE by electroporation and further processed as described above. Colonies were analyzed for the presence or absence of the tatA gene by colony PCR using primers CgDel-for and CgDel-rev. One of the isolates that besides the tatE deletion also contained the desired chromosomal deletion of the tatA structural gene was designated C. glutamicum ATCC13032 ΔtatA/E.

C. glutamicum ATCC13032 ΔtatB was constructed by deleting the tatB gene using the pCRD206-based system that contains a temperature-sensitive replicon and also the sacB gene as a counter-selectable marker [51]. Two DNA fragments containing approximately 480 bp upstream and downstream sequences of the tatB gene were generated using chromosomal DNA of C. glutamicum ATCC13032 as template together with primers del_tatBup-f and del_tatBup-r or primers del_tatBdn-f and del_tatBdn-r, respectively. Both PCR products were purified and used as a template in a cross-over PCR reaction using primers del_tatBup-f and del_tatBdn-r to generate a fused DNA fragment that contained the up- and downstream sequences of tatB, but lacks the tatB structural gene. The corresponding PCR fragment was digested with Xbal and SbfI and ligated into Xbal/SbfI-digested pCRD206, yielding pCRD206ΔtatB. For deletion of the chromosomal tatB structural gene, pCRD206ΔtatB was introduced into C. glutamicum ATCC13032 by electroporation and further processed as described by Okibe et al. [51]. Colonies were analyzed for the presence or absence of the tatB gene by colony PCR using primers deltaB-f and deltaB-r. One of the isolates that contained the desired chromosomal deletion of the tatB structural gene was designated C. glutamicum ATCC13032 ΔtatB. The C. glutamicum ATCC13032 ΔqcrA ΔtatB double mutant was constructed in an identical way by applying the pCRD206ΔtatB deletion vector to the C. glutamicum ATCC13032 ΔqcrA mutant strain (see below).

For the deletion of the C. glutamicum qcrA structural gene from the chromosome, the pK19mobsacB system was used. Two DNA fragment containing approximately 490 bp of upstream and downstream sequences of the qcrA gene were generated using chromosomal DNA of C. glutamicum ATCC13032 as template together with primers del_qcrAup-f and
del_qcrAup-r or primers del_qcrAdn-f and del_qcrAdn-r, respectively. Both PCR products were purified and used as a template in a cross-over PCR reaction using primers del_qcrAup-f and del_qcrAdn-r to generate a fused DNA fragment that contained the up- and downstream sequences of qcrA, but lacks the qcrA structural gene. The corresponding PCR fragment was digested with XbaI and Xmal and ligated into XbaI/Xmal-digested pK19mobsacB, yielding pK19mobsacBΔqcrA. For deletion of the chromosomal qcrA structural gene, pK19mobsacBΔqcrA was introduced into C. glutamicum ATCC13032 by electroporation and further processed as described above, resulting in the strain C. glutamicum ATCC13032 ΔqcrA.

Plasmid constructions

The plasmids used in this study are listed in Table 1. Oligonucleotides used as PCR primers are listed in S1 Table. All DNA manipulations followed standard procedures [47]. The correctness of all newly constructed plasmids was verified by DNA sequencing.

To allow plasmid-based expression of C. glutamicum QcrA in C. glutamicum, the qcrA gene was amplified by PCR using chromosomal DNA of C. glutamicum ATCC13032 as template and primers SalI-qcrA-f and qcrA-KpnI-r. The resulting PCR fragment was digested with SalI and KpnI and ligated into SalI/KpnI-digested pEKEx2 [52], resulting in pEKEx2-QcrA. From this plasmid, a variant (pEKEx2-QcrAKK) was constructed that encodes a QcrAKK mutant protein in which the two conserved twin-arginine residues at positions 159 and 160 were replaced by a twin-lysine pair. The corresponding mutations were introduced into pEKEx2-QcrA via the Quick Change II site-directed mutagenesis kit (Agilent) using QC:QcrARR-KK-f and QC:QcrARR-KK-r as primers.

For plasmid-based expression of the C. glutamicum TatB in C. glutamicum, the tatB gene was amplified by PCR using chromosomal DNA of C. glutamicum ATCC13032 as template and primers SacI-tatB-f and tatB-BamHI-r. The resulting PCR fragment was digested with SacI and BamHI and ligated into SacI/BamHI-digested pEC-XC99E [53], resulting in pEC-TatBCG.

For plasmid-based expression of C. glutamicum TatB in E. coli tat mutant strains, the tatB gene was amplified by PCR using E. coli MC4100 chromosomal DNA as template and primers EcoRI-tatBEc-f and EctatB-SalI-r. The resulting PCR fragment was digested with EcoRI and SalI and ligated into the EcoRI/SalI-digested low copy number vector pHSG575 [54], resulting in plasmid pHSG-TatBEC.

For plasmid-based expression of C. glutamicum TatB in E. coli tat mutant strains, the tatB gene was amplified by PCR using C. glutamicum ATCC13032 chromosomal DNA as template and primers EcoRI-tatB-f and tatB-SalI-r. The resulting PCR fragment was digested with EcoRI and SalI and ligated into EcoRI/SalI-digested pHSG575, resulting in plasmid pHSG-TatBCG.

Miscellaneous procedures

For the analysis of PhoDCG-GFP secretion in C. glutamicum wild-type and tat mutant strains, the respective C. glutamicum cells containing plasmid pCGPhoD CG -GFP were grown in BHIS medium in 4-chicane Erlenmeyer flasks at 30°C and 120 rpm to an OD600 of 1.0 and gene expression was induced for 6 h by adding 100 μM IPTG. Subsequently, cellular and supernatant fractions were prepared as described previously [35]. The distribution of GFP-derived polypeptides in the corresponding fractions was analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and Western blotting using GFP-specific antibodies. Western blotting was performed using the ECL Western blotting detection kit (GE Healthcare) according to the manufacturer’s instructions. The chemiluminescent protein bands were recorded using the Fuji LAS-3000 Mini CCD camera and image analyzing system using the software AIDA 4.15 (Raytest).
E. coli tat mutants are sensitive to the presence of SDS unless they are complemented by functional homologous or heterologous tat genes [41]. The complementation of E. coli tat mutants by plasmid-encoded TatB proteins from E. coli or C. glutamicum was analyzed by the assay described by Ize et al. [40]. Overnight cultures of the strains were diluted to an OD<sub>600</sub> of 0.05 with LB medium containing either 0.1%, 0.5%, 1%, 2% or 4% SDS (w/v) and grown aerobically for 180 min at 37°C and 250 rpm. Subsequently, the OD<sub>600</sub> of each culture was measured. 100% survival is defined as the OD<sub>600</sub> of each strain after 180 min growth in LB medium without SDS.

**Supporting Information**

S1 Fig. Analysis of TorA-GFP secretion in C. glutamicum wild-type and tat mutant strains.
Cultures of C. glutamicum strains expressing the Tat-dependent TorA-GFP model protein [35] were fractionated into cells (C) and supernatant (S). Samples of the fractions corresponding to an equal number of cells (i.e. an OD<sub>600</sub> of 1.0) were subjected to SDS-PAGE and immunoblotting using GFP-specific antibodies. The following strains were analyzed: C. glutamicum wild-type (wt) containing the empty vector pEKEx2 as negative control (lanes 1 and 2), C. glutamicum wild-type (wt) containing plasmid pCGTorA-GFP (lanes 3 and 4) and the pCGTorA-GFP-containing C. glutamicum mutant strains ΔtatAC (lanes 5 and 6), ΔtatB (lanes 7 and 8), and ΔtatA/E (lanes 9 and 10). p: TorA-GFP precursor; asterisk: cytosolic degradation product; m: mature-sized GFP protein.

S2 Fig. Uncropped image of Western blot shown in Fig 3.

S3 Fig. Uncropped image of Western blot shown in S1 Fig

S1 Table. Primers used in this study.

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**Author Contributions**

Conceived and designed the experiments: DO RF. Performed the experiments: DO SS. Analyzed the data: DO SS RF. Contributed reagents/materials/analysis tools: DO SS. Wrote the paper: DO SS RF.

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