**Yersinia enterocolitica** YopT and *Clostridium difficile* Toxin B Induce Expression of GILZ in Epithelial Cells

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**Abstract**

Glucocorticoid induced-leucine zipper (GILZ) has been shown to be induced in cells by different stimuli such as glucocorticoids, IL-10 or deprivation of IL-2. GILZ has anti-inflammatory properties and may be involved in signalling modulating apoptosis. Herein we demonstrate that wildtype *Yersinia enterocolitica* which carry the pYV plasmid upregulated GILZ mRNA levels and protein expression in epithelial cells. Infection of HeLa cells with different *Yersinia* mutant strains revealed that the protease activity of YopT, which cleaves the membrane-bound form of Rho GTPases was sufficient to induce GILZ expression. Similarly, *Clostridium difficile* toxin B, another bacterial inhibitor of Rho GTPases induced GILZ expression. YopT and toxin B both increased transcriptional activity of the GILZ promoter in HeLa cells. GILZ expression could not be linked to the inactivation of an individual Rho GTPase by these toxins. However, forced expression of RhoA and RhoB decreased basal GILZ promoter activity. Furthermore, MAPK activation proved necessary for profound GILZ induction by toxin B. Promoter studies and gel shift analyses defined binding of upstream stimulatory factor (USF) 1 and 2 to a canonical c-Myc binding site (E-box) in the GILZ promoter as a crucial step of its trans-activation. In addition we could show that USF-1 and USF-2 are essential for basal as well as toxin B induced GILZ expression. These findings define a novel way of promoter trans-activation mediated by bacterial toxins and differentiate it from those mediated by dexamethasone or deprivation of IL-2.

**Introduction**

*Yersinia enterocolitica* is an enteropathogenic bacterium which causes gastrointestinal disorders such as enteritis and enterocolitis, and extraintestinal manifestations such as lymphadenitis, reactive arthritis, erythema nodosum, uveitis and septicemia [1,2]. Host cells can sense *Y. enterocolitica* by recognizing bacterial factors like LPS, invasin, YadA and YopB and can react with a pro-inflammatory response [3,4,5].

In line with this, gene expression analysis of epithelial cells revealed that upon interaction with *Yersinia*, chromosomally encoded bacterial factors such as the adhesin invasin lead to gene expression of a large number of pro-inflammatory genes [6]. In cells infected with wildtype *Yersinia* this host response is suppressed by injection of virulence plasmid (pYV)-encoded factors into host cells [7]. In contrast, only a few host genes were found exclusively induced by pYV-encoded factors such as glucocorticoid induced leucine zipper (GILZ) and kruippel like factor (KLF) 2 [6,8].

GILZ or TSC22 domain family, member 3 (Tsc22d3), a member of the leucine zipper protein family, was identified by comparing mRNA species expressed in dexamethasone (DEX) treated and untreated murine thymocytes [9]. Furthermore, GILZ gene expression was also found to be induced by interleukin (IL) 10 signaling or by IL-2 deprivation in T-cells and macrophages [10,11]. GILZ expression protects T cells from apoptosis induced by treatment with anti-CD3 monoclonal antibodies, probably via down-regulation of Fas/FasL expression [9]. Most notably, GILZ has been demonstrated to be an important mediator of the anti-inflammatory and anti-proliferative effects of glucocorticoids [12]. For instance it has been shown that anti-inflammatory effects of DEX in lung epithelial cells are mediated by GILZ [13]. It prevents NF-κB activation by inhibition of NF-κB nuclear translocation and DNA-binding due to a direct protein-protein interaction with the NF-κB subunits [14]. By direct interaction of GILZ with Ras and Raf, the Ras/Raf/Erk signaling pathway is repressed [15,16]. In similar the activation of the transcription factor AP-1 can be inhibited by GILZ [17].

Here, we have investigated how *Yersinia* may induce GILZ expression. We identified YopT as the crucial toxin which mediates *Y. enterocolitica* induced GILZ expression. Additionally,
we present with *C. difficile* toxin B another bacterial toxin inducing GILZ expression. Moreover, we could demonstrate that GILZ promoter trans-activation and GILZ protein expression mediated by the Rho-inactivating toxin B and YopT depends on the binding of USF-1 and USF-2 to a canonical E-box element of the GILZ promoter, which represents a novel pathway of GILZ induction.

**Results**

**Yersinia Enterocolitica** Induces GILZ Expression in Epithelial Cells

Microarray experiments revealed that GILZ mRNA was upregulated in epithelial cells upon infection with the *Yersinia enterocolitica* patient isolate WA-314 carrying the pYV virulence plasmid (pYV+) [6]. To confirm the previously described microarray experiments, HeLa cells were infected with the pYV+ patient isolate or its virulence plasmid cured derivative pYV-. GILZ protein expression was analyzed in cell lysates by Western blotting using a GILZ specific polyclonal antibody preparation at different time points (Figure 1). An induced expression of an approximately 15 kDa immunoreactive band was observed 2, 4 and 8 h after infection with pYV+ but not with pYV-.

To investigate whether induction of GILZ mRNA expression is dependent on secretion or translocation of *Yersinia* outer proteins (Yops), HeLa cells were infected for 2 h with different *Yersinia* strains such as *Y. enterocolitica* pYV515, a lcrD mutant which does not secrete any Yops [18] and *Y. enterocolitica* WA-pTTSS [19] which expresses the type three secretion system and YadA but not YopE, YopP, YopT, YopO, YopM and YopH. GILZ mRNA expression was analyzed by real-time RT-PCR. As shown in Figure 2A, only infection with pYV+, but not infection with pYV515, pTTSS or pYV− induced GILZ mRNA expression, indicating that translocation of effector-Yops is a prerequisite for GILZ protein expression.

To identify those Yops involved in GILZ induction, HeLa cells were infected with different Yop deletion mutants for 4 h and cytosolic proteins were extracted. Western blot analysis revealed that deletion of yopT but none of the other yops tested diminished GILZ protein expression indicating that YopT is involved in the induction of GILZ expression (Figure 2B).

In line with this finding, also infection of the monocye cell line Mono Mac 6 infection of cells with *Yersinia* increased GILZ expression in a YopT dependent manner (data not shown). YopT, the most recently identified Yop [20] is a cysteine protease cleaving membrane-bound Rho GTPases from their isoprene membrane anchor [21]. To address whether the protease activity of YopT is essential for induction of GILZ, HeLa cells were infected with a *Yersinia* pYV+, ΔyopT deletion mutant complemented either with a plasmid encoding active YopT (pYV+ ΔyopT yopT) or with a protease-inactive YopT mutant (pYV+ ΔyopT yopT yopT(C139A)) [21] and GILZ protein and mRNA expression was analysed (Figure 2C, D). To ensure that yopT deletion does not affect secretion of other Yops, secretion of YopE and YopH was analyzed but no differences in secretion were found (data not shown). Complementation with yopT restored GILZ expression in contrast to complementation with yopT(C139A), providing evidence that cysteine protease activity of YopT is crucial for induction of GILZ expression (Figure 2C). In parallel, the effect of YopT on membrane bound and cytosolic RhoA was investigated. Reduction in the amount of membrane bound RhoA was found after infection with pYV+ or infection with pYV+ ΔyopT pypotP. A clear accumulation of RhoA was found in the cytosolic fraction after infection with pYV+ or pYV+ ΔyopT pypotP compared to infection with the other strains investigated.

Other virulence factors of *Yersinia* affect RhoGTPases as well. To investigate whether YopE or YopO also might induce GILZ expression in HeLa cells we used the *Yersinia* strain pTTSS complemented with plasmids encoding yopE3,5-ova, yopE3,5-yopT, yopE or yopO. Only infection with pTTSS yopE53-yopT or pYV+ induced GILZ expression (Figure 2E). This data provided no evidence that other bacterial factors beside Yop T contribute to induce GILZ expression.

**YopT Mediated GILZ Induction does not Silence Pro-inflammatory Gene Transcription by NFκB**

In response to various inflammatory stimuli, NFκB is released from cytoplasmic sequestration by IκB to the nucleus by ubiquitinilation and subsequent degradation of its binding partner, though it may even play a role in anucleated erythrocytes [22]. In *Yersinia* infection, this activates transcription of pro-inflammatory cytokines, e.g. IL-8 [23,24]. In line with previous reports showing massively reduced NFκB transcriptional activity in cells over-expressing GILZ [14], we could demonstrate strong inhibition of basal or induced activity of a NFκB promoter luciferase construct in HeLa cells over-expressing GILZ (Figure S1A). Examining the effect of YopT expression on basal and induced NFκB transcription we observed moderate suppression in both cases consistent with results yielded with *T. pseudobacterellosis* YopT [25]. Furthermore we could link NFκB suppression to YopT protease activity (Figure S1B). However, inhibition of GILZ expression with GILZ specific siRNA (Figure S1C) did not cause significant difference in NFκB transcriptional activity (Figure S1D).
Figure 2. *Yersinia* induced GILZ expression depends on YopT protease activity. HeLa cells were infected with (A) strains with (pYV') or without (pYV") pathogenicity plasmid or with a pathogenicity plasmid derivative coding for a functional translocation apparatus but not for effector Yops (pTTSS) or with a pathogenicity plasmid coding for a defective translocation apparatus (pYV5.15) with MOI 20 for 2 or 4 h to determine GILZ mRNA expression by real-time RT-PCR. Means + SD of 2 independent experiments. Significant differences compared to pYV' are indicated by asterisks (p<0.05). (B) Immunoblot of GILZ protein expression induced by strains with single yop deletions. Representative immunoblot and quantification means + SEM of 4 independent experiments normalized to untreated. Significant differences compared to ΔyopT are indicated by asterisks (p<0.05). (C) Infection with a ΔyopT deletion strain and derivative strains complemented with an additional plasmid encoding wildtype (pyopT) or protease deficient (pyopT<sub>C139A</sub>) YopT. Representative immunoblot analysis of GILZ protein expression in the cytosolic fraction and of RhoA in cytosolic and membrane fraction (left) and quantification means + SD of GILZ expression of two representative experiments normalized to untreated (right).
GILZ protein expression induced by strains translocating only one Yop affecting Rho GTPases. Cells were infected with Yersinia translocating YopE, YopE, YopT or YopO. Infections with pYY, ΔyopT and a strain translocating only enzymatically inactive YopE-Ova were used as controls.

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Expression in Epithelial Cells

Clostridium Difficile Toxin B Induces Massive GILZ Expression in Epithelial Cells

The cleavage of the membrane-bound forms of Rho GTPases by YopT leads to their release from membranes, thus abrogating downstream effector contact. We investigated whether other toxins which affect Rho GTPases such as C. difficile toxin B (TcdB 10463) or C. limosum C3 transferase might also induce GILZ transcription and expression. Toxin B inhibits the small Rho GTPases RhoA, Rac, Cdc42, RhoG and TC10 by glucosylation. The activity of toxin B was indicated by rounding of HeLa cells 2 h after treatment (data not shown). In addition, glucosylation of Rac1 was demonstrated by immunobLOTS using the antibody mAb102 which only recognizes non-glucosylated Rac1 (Figure 3A). Moreover, toxin B induced RhoB expression as has been previously reported [26] (Figure 3A). C3 transferase exclusively inactivates RhoA - B and -C by ADP-ribosylation. To enable sufficient uptake, the uptake mechanism of the binary C. botulinum C2 toxin was exploited: HeLa cells were treated with a recombinant C2IN-C3lim fusion toxin that is comprised of the catalytic domain of C. limosum C3 transferase and the adaptor domain of the C2I component. This allows efficient cellular internalization via the activated C2IHa transport component [27]. The toxic activity of C2IN-C3lim fusion toxin was monitored and validated by induction of typical morphological changes of the cells and by sequential ADP-ribosylation of cell lysates after toxin treatment (Figure S2). GILZ protein expression was analyzed by immunobLOTS. Toxin B rapidly induced GILZ expression (Figure 3A) while C3 transferase induced GILZ expression only marginally (Figure 3B). In similar, treatment of Mono Mac 6 cells with toxin B or DEX led to increased GILZ expression six hours after stimulation (data not shown).

Recent reports pointed out that several mouse GILZ (Tsc22d3) and probably also human GILZ isoforms exist [28,29]. So far there are two mouse isoforms annotated in NCBI Protein: Tsc22d3 isoform 1 (NP_001070832.1, 201 aa, 22 kDa) named also long GILZ (L-GILZ) and a shorter isoform 2 (NP_034416, 137 aa, 15 kDa), the prototypic GILZ [28]. Notably, Bruscoli et al. found L-GILZ to appear at a molecular weight of ~ 28 kDa. In a recent screening for host factors interacting with herpex simplex protein, human orthologues of these GILZ isoforms have been found: human GILZ isoform 1 (NP_932714.1, 200 aa, 22 kDa) and isoform 2 (NP_004080, 134 aa, 15 kDa) [30]. While the long GILZ isoforms show some differences in the sequence N-terminal of the Tsc22 box, the short isoforms differ only by two amino acids.

Therefore, we addressed the question whether the human orthologue of L-GILZ is also expressed in HeLa cells besides 15 kDa GILZ examined so far. To provide additional evidence, that observed immunoreactive bands really represent GILZ isoforms, cells were transfected with siRNA specific for human GILZ 3’UTR and thus repressing all transcripts of human GILZ. Analysis of HeLa cell lysates showed two dominant bands which were recognized by our polyclonal antibody preparation and which were repressed by the pan-GILZ siRNA: We found a strong and constitutively expressed GILZ protein that migrated at slightly below 30 kDa (human Tsc22d3 isoform 1/L-GILZ) while treatment of cells with DEX or toxin B only markedly induced expression of the 15 kDa GILZ protein (human Tsc22d3 isoform 2; Figure 3C). Our data thus parallel the results of Soundararajan et al. who found L-GILZ expressed constitutively in various murine tissues while treatment with the corticoid aldosterone induced GILZ isoform 1 expression [29]. In the presented study we therefore continued focusing on 15 kDa GILZ expression that was also induced by YopT protease activity.

GILZ mRNA expression was analyzed using primers that bound to the GILZ 3’-UTR throughout the study. With these pan-GILZ primers also GILZ mRNA expression was also detectable already 2 h after toxin B treatment (Figure 3D).

Elevated GILZ mRNA Levels Result from Transcriptional Activation of the GILZ Promoter

To elucidate, whether increased GILZ mRNA levels were the result of enhanced transcription, HeLa cells were transfected with a luciferase reporter gene preceded by 2088 bases of the GILZ promoter region (p2088-Luc), as such constructs have proven sufficient to assay GILZ transcription in previous studies [11]. Measurements of luciferase activity in lysates of cells transfected with p2088-luc were in line with detection of GILZ mRNA and GILZ protein levels. Significantly increased GILZ promoter activity was detected only in cells infected with pYY or pITSS yopE53-yopT or treated with toxin B or DEX (as compared to uninfected and untreated cells, Figure 3E). Thus, GILZ induction under these conditions results at least partially from enhanced GILZ mRNA de novo synthesis. Notably, toxin B showed comparable potency to activate the GILZ promoter as glucocorticoids.

Involvement of Rho GTPase – MAP Kinase Signaling in GILZ induction

Further studies of GILZ induction were focused mainly on toxin B because of its stronger trans-activation of the GILZ promoter. The observation of GILZ induction by YopT and toxin B but not by YopE or YopO and only marginally by C3 toxin indicated that inactivation of Rho family GTPases may not be solely sufficient for potent GILZ induction. Consequently, inhibition of the effector protein Rho associated kinase kinase (ROCK), an important transducer of RhoA signaling as well as treatment of cells with lovastatin, a general saponylation blocker of Rho family GTPases were largely ineffective in generating GILZ expression (data not shown). To investigate whether modulation of Rho GTPase expression levels may affect the transcriptional activity of the GILZ promoter, first HeLa cells were transfected with different expression vectors leading to overexpression of RhoA, Rac1, CDC42 or RhoB. Transient transfection led to overexpression of these proteins compared to transfection with empty control vector (Figure 4A, upper panel). Interestingly, overexpression of RhoA or RhoB specifically and significantly repressed basal GILZ promoter activity, as opposed to expression of Rac1 or CDC42 (Figure 4A, lower panel). This result clearly indicates that distinct Rho GTPases modulate transcriptional regulation of GILZ. In a second experiment HeLa cells were transfected for 48 h with either siRNA specific for RhoA or RhoB or both and immunoblots were performed (Figure 4B). RhoB expression was hardly detectable but inhibition of RhoA expression led to upregulation of RhoB expression. Neither inhibition of RhoA alone nor of RhoA and RhoB in combination led to upregulation of GILZ expression, indicating that expression levels of RhoA and RhoB might be...
Figure 3. GILZ is expressed upon stimulation with C3 exotoxin or toxin B. (A) HeLa cells were incubated with toxin B (50 ng/ml) for indicated time spans and immunoblots were performed from whole cell lysates using anti-Rac1 mAb102 recognizing only non-glucosylated Rac-1, and an anti-Rac1 mAb antibody recognizing total Rac1 as well as antibodies recognizing RhoB, actin or GILZ. (B) Immunoblot of GILZ and actin expression upon stimulation of HeLa cells with C2IN-C3lim (100 ng/mL) + C2IIa (200 ng/mL) for indicated time spans. (C) To explore the expression of additional GILZ isoforms, HeLa cells were transfected with 7.5 nM siRNA specific for GILZ or control siRNA for 48 h and subsequently either left untreated or stimulated with C. difficile toxin B (50 ng/ml) or 100 μM DEX for 4 h. Arrows mark three GILZ isoforms which were inhibited by the used GILZ siRNA. Note that only isoform 1 was induced by the used stimuli. (D) Cells were stimulated with toxin B for 2 or 4 h to determine GILZ mRNA expression by real-time RT-PCR. Mean + SD of 2 independent experiments normalized to untreated. (E) To assay transcriptional activity of the GILZ promoter cells were transfected with a luciferase reporter under control of a 2088 bp GILZ promoter and co-transfected with pCMV-β-gal (for standardization) 24 h before infection with a Y. enterocolitica pYV' and various mutant strains or treatment with DEX or toxin B. Means + SD of 4 independent experiments normalized to untreated. Significant differences compared to untreated are indicated by asterisks (p<0.05).

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involved in modulating transcriptional activity but are not sufficient for regulation of GILZ expression. As toxin B has been reported to elicit ERK and p38 phosphorylation in treated cells [31,32] we examined the possible contribution of MAPK activation in toxin B mediated GILZ induction. We detected moderate p38 and ERK phosphorylation peaking as early as 10 min after toxin B treatment which subsequently vanished (Figure 4C).

To test whether MAPK activation contributed to GILZ induction, we analyzed the ability of toxin B to induce GILZ expression in the presence of inhibitors of MAPK phosphorylation and treated cells with SB 202190 (target: p38) or PD 98059 (ERK) 2 h prior to toxin B stimulation and GILZ protein was detected by immunoblot analysis at 6 h or 24 h after stimulation. In conclusion, induction of GILZ expression by bacterial toxins did not only require the inhibition of Rho GTPases. We speculate that several activities of these toxins are necessary, such as both inhibition of distinct RhoGTPases and activation of MAP kinases. Toxin B Transactivates the GILZ Promoter via a Canonical E-box

To test which promoter elements might be of relevance for toxin B induced GILZ expression HeLa cells were co-transfected with the p2088 GILZ promoter luciferase reporter and pHM6 based plasmid for overexpression of the indicated Rho GTPases. Means + SD of 3 independent experiments normalized to untreated. Significant differences compared to control vector transfection are indicated by asterisks (p < 0.05). In a control experiment, HeLa cells were transfected in the same setting and cell lysates were used for immunoblots to detect RhoA, RhoB, Cdc42 and Rac1 expression. (B) HeLa cells were transfected for 48 h with indicated concentrations of siRNA. Immunoblots were performed from cell lysates for RhoA and RhoB and from cytosolic extracts for GILZ. (C) Toxin B treatment leads to fast and transient MAPK phosphorylation. After treatment of HeLa cells with toxin B for the indicated time spans, levels of phosphorylated as well as total ERK and p38 were assayed by immunoblot. (C) Toxin B induced GILZ expression is mediated by both ERK and p38 MAPK. Cells were pretreated with MAPK phosphorylation inhibitors SB 202190 (p38) or PD 98059 (ERK) 2 h prior to toxin B stimulation and GILZ protein was detected by immunoblot analysis at 6 h or 24 h after stimulation. doi:10.1371/journal.pone.0040730.g004

Figure 4. Role of Rho GTPases and MAP kinases for GILZ expression. (A) Overexpression of RhoA or RhoB lowers basal GILZ levels. HeLa cells were co-transfected with the p2088 GILZ promoter luciferase reporter and pHM6 based plasmid for overexpression of the indicated Rho GTPases. Means + SD of 3 independent experiments normalized to untreated. Significant differences compared to control vector transfection are indicated by asterisks (p < 0.05). In a control experiment, HeLa cells were transfected in the same setting and cell lysates were used for immunoblots to detect RhoA, RhoB, Cdc42 and Rac1 expression. (B) HeLa cells were transfected for 48 h with indicated concentrations of siRNA. Immunoblots were performed from cell lysates for RhoA and RhoB and from cytosolic extracts for GILZ. (C) Toxin B treatment leads to fast and transient MAPK phosphorylation. After treatment of HeLa cells with toxin B for the indicated time spans, levels of phosphorylated as well as total ERK and p38 were assayed by immunoblot. (C) Toxin B induced GILZ expression is mediated by both ERK and p38 MAPK. Cells were pretreated with MAPK phosphorylation inhibitors SB 202190 (p38) or PD 98059 (ERK) 2 h prior to toxin B stimulation and GILZ protein was detected by immunoblot analysis at 6 h or 24 h after stimulation. doi:10.1371/journal.pone.0040730.g004

Toxicity B Transactivates the GILZ Promoter via a Canonical E-box

To test which promoter elements might be of relevance for toxin B induced GILZ expression HeLa cells were transfected with GILZ promoter-luciferase constructs containing promoter fragments of different length (ranging from -416 bp to -2088 bp upstream of the transcription initiation site, Figure 5A). As reported previously [11,33] removal of the glucocorticoid responsive elements (GRE)
3, 4 and 5 (-1526 construct) abrogated DEX mediated promoter activation. In the case of toxin B stimulation however, transfection with even the most truncated GILZ-promoter fragment (-416) proved sufficient for trans-activation (Figure 5B).

Conserved transcription factor (TF) binding sites in this sequence comprise a FHRE (FHRE1), two Oct1 (Oct1a and Oct1b) and one GRE element (GRE1), a canonical E-box (myc1) and a non-canonical E-box (myc2). To single out the promoter motif required for GILZ induction by toxin B we introduced into the GILZ-1940 promoter region point mutations disrupting consensus sequences of identified trans-activating sites (Figure 6A). Measurement of luciferase activity in HeLa cells transfected with this derivative of the p1940 GILZ promoter luciferase construct revealed decreased luciferase activity of all mutants except for myc2 compared to p1940 (Figure 6B). This data indicates that FHRE1, Oct1, GRE elements and the canonical E-box myc1 positively affect basal activity of the GILZ promoter while the non-canonical E-box myc2 enables repression of basal GILZ promoter activity. While FHRE1, Oct1 and myc2 mutated promoters retained their responsiveness to toxin B- and DEX-treatment, relative increase of GILZ promoter activity (compared to unstimulated cells transfected with the same construct) by DEX was only influenced by mutation of GRE1+2. In contrast, GILZ promoter trans-activation induced by toxin B was only affected by mutation of the canonical E-box myc1 (Figure 6C).

**Figure 5. Differential activation of truncated GILZ promoter elements by toxin B or DEX.** HeLa cells were transiently transfected with pCMV-β-gal for 24 h and co-transfected (A) with p2088-Luc or luciferase reporters fused to shortened promoter regions, containing the indicated TF binding sites. GRE: glucocorticoid responsive element, FHRE: forkhead responsive elements, c-myc: c-myc binding site (E-box), Oct: octamer binding site (B) Transfected cells were treated with 100 μM DEX or 50 ng/ml toxin B for 6 h. Subsequently luciferase assays were performed. Results are expressed as fold induction compared to unstimulated (none) cells and represent the mean ± SEM of three experiments performed in triplicates \( (p<0.05) \). doi:10.1371/journal.pone.0040730.g005

**Toxin B Treatment and pYV+ Infection Results in TF Binding to the GILZ Promoter Canonical E-box**

To identify proteins which may bind to the canonical E-box (myc1), electrophoretic mobility shift assays (EMSA) were performed using a GILZ promoter dsDNA fragment (GILZ-63/37) comprising the canonical E-box and two partially overlapping putative Ceb binding elements (CRE) (Figure 7A). Stimulation of HeLa cells with toxin B as well as infection with _Y. enterocolitica_ pYV+ - but not _DyopT_- increased protein complex binding to GILZ-63/37 as soon as 30 minutes after stimulation with toxin B (Figure 7B and C). Competition experiments revealed that binding of TF protein could be abrogated by addition of excess unlabeled GILZ-63/37 DNA probe, CRE I mut or CRE II mut probe (Figure 7D). In contrast, an excess of cold probe with mutated myc1 E-box (E-box mut) was not able to compete with GILZ-63/37. We therefore conclude that the E-box but not the flanking CRE elements are crucial for increased binding of transcription factors to this GILZ DNA region after treatment of cells with toxin B.

**Toxin B Treatment Results in GILZ Promoter Activation by USF1 and USF2**

The CANNTG E-box sequence has been characterized as binding motif of the large basic/helix-loop-helix/leucin zipper (b/HLH/Z) TF family [34]. The human c-Myc oncoprotein is probably the most prominent member of this family, but co-incubation with antibody directed against c-Myc did not change
Figure 6. Importance of specific cis-elements for toxin B induced GILZ promoter trans-activation. (A) Recognition sequences of cis-elements which were mutated. Mutated base pairs are highlighted using bold letters. (B) HeLa cells were transfected with p1940-Luc and different mutated derivatives for 24 h and subsequently stimulated with DEX or toxin B for 6 h. Data are shown as relative light units (RLU) standardized to β-Gal activity and protein concentration or (C) as fold induction after DEX or toxin B stimulation compared to untreated conditions of each individual expression vector. Means ± SEM of four independent experiments are shown. Asterisks indicate significant differences between DEX or toxin B stimulation compared to uninfected (p < 0.05).

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gel-shift behavior of the $GILZ^-63/-37$ probe as was the case with antibodies specific for Creb and p65 that have been included as negative controls (Figure 7E).

In HeLa cells, the b/HLH/Z TF upstream stimulatory factor (USF) has been described to bind the E-box motif with high affinity and to activate transcription of downstream genes [35].

Addition of anti-USF-1 or anti-USF-2 antibody to EMSA samples specifically abrogated or supershifted DNA-protein complex formation, respectively (Figure 7E). This data shows that toxin B induced $GILZ^\zeta$ promoter activation is associated with increased binding of USF-1 and USF-2 to the canonical E-box.

Inhibition of both USF-1 and USF-2 protein expression by transfection with USF-1 and USF-2 siRNA abrogated basal as well as toxin B induced $GILZ^\zeta$ promoter activation.
Discussion

Upon interaction with epithelial cells, Y. enterocolitica yopV* inhibits gene expression induced by other bacterial factors such as Inv, YadA and YopB very efficiently [3,4,5,6]. Only a few genes are upregulated in epithelial cells infected with Y. enterocolitica yopV* [6]. As demonstrated here, infection of HeLa cells with Y. enterocolitica results in activation of the GILZ promoter followed by enhanced gene transcription and GILZ protein expression.

Previous studies showed that several GILZ isoforms exist in mice as well as in humans [28,29,30]. Research has been focused on the 15 kDa isoform 2 so far, but it has become clear that other isoforms may be differentially expressed and may display different downstream activities. Thus, these studies also analyzed the expression of the larger isoform 1 (L-GILZ). In various murine tissues and cultured kidney epithelial cells, L-GILZ was found to be constitutively expressed, while GILZ expression was induced by aldosterone treatment [29]. In primary mouse myoblast cultures however, DEX treatment induced both L-GILZ and GILZ expression [28].

We transfected HeLa cells with siRNA matching the 3’UTR of GILZ to prove that detected protein bands represent indeed GILZ protein. By using polyclonal GILZ antibody, we detected strong constitutive expression of L-GILZ at almost 30 kDa. In fact, the L-GILZ transcript encodes only a 22 kDa protein, when translated from the canonical translation start codon. However, it has been demonstrated in mouse cells that translation of this GILZ isoform starts earlier at a non-canonical CUG codon, thus resulting in a larger protein [28]. Comparison of mouse (NM_001077364.1) and human (NM_198057.2) GILZ mRNA transcripts revealed that the same non-canonical start codon preceded by a Kozak consensus sequence is also present in human GILZ mRNA and our western blot analysis showed that human L-GILZ is also predominantly translated as a larger protein in HeLa cells, while a faint band indicates that low amounts of the canonical translation product also exist.

Additional stimulation or infection of HeLa cells only induced expression of 15 kDa GILZ isoform 2, further referred to as GILZ in this study. Unfortunately, there is no uniform nomenclature of GILZ isoforms. We have adopted the numbering of GILZ isoforms that is used by NCBI Protein, but the nomenclature of other databases (e.g. UniProt) or publications may be completely different.

Infection with different Yersinia mutants which are not able to secrete or translocate Yops demonstrates that translocation of Yops into the cytosol HeLa cells is necessary for GILZ induction. In addition we provide evidence that YopT but not YopE or YopO which are also known to inhibit Rho GTPases is required to induce GILZ expression.

Infection of HeLa cells with Yop deletion mutants revealed that GILZ expression was abrogated upon infection with a Y. enterocolitica yopT deletion mutant and complementation with yopT restored GILZ expression. In contrast, deletion of yopP, yopE, yopM or yopH did not affect GILZ expression, implying that YopT might be crucial for GILZ expression.

Infection with a mutant translocating only YopT into HeLa cells clearly induced GILZ expression indicating that no other Yops are required to induce GILZ expression. Consistently mutants translocating YopE or YopO were not able to induce GILZ expression.

For catalytic protease activity of YopT the amino acids C139, H258 and D274 are crucial [38]. In line with this finding infection experiments with a yopT deletion mutant complemented with yopTC139A encoding protease deficient YopT demonstrated that...
protease activity of YopT was required for increased GILZ expression.

YopT is a cysteine protease which cleaves RhoA, Rac, and Cdc42 [21,38,39] directly upstream of the C-terminal isoprenylated cysteine. Which Rho GTPases are affected by YopT inside the cells is still unclear. It has been reported that translocated YopT inserts into the plasma membrane. There it cleaves RhoA but not Rac or Cdc42 removing membrane bound RhoA as well as depleting the cytoplasmic RhoA - GDI-1 pool. This finally results in accumulation of RhoAn the cytosol [40]. Other publications however indicate that upon *Listeria pseudotuberculosis* infection YopT cleaves RhoA as well as Rac and Cdc42 from nascent phagosomes and affects RhoG [41,42].

Since YopT is able to induce GILZ expression, we tested whether other bacterial toxins inhibiting Rho GTPases could also trigger GILZ expression. In line with this hypothesis *C. difficile* toxin B which leads to UDP-glucosylation of Rho GTPases [43] and inactivates RhoA, Rac, Cdc42, RhoG and TC10 or indirectly activates RhoB [44], strongly induced GILZ expression two, four and six h after start of treatment. After 24 h GILZ expression faded, possibly because of toxin B degradation and because of the activation of RhoB expression that in turn may inhibit GILZ expression. In contrast, C3 transferase which affects RhoA, -B, and -C by ADP-ribosylation only marginally induced GILZ expression. The low impact of C3 transferase on GILZ expression might suggest that inactivation of RhoA, RhoB or RhoC is not sufficient to trigger significant *GILZ* expression. Similarly, YopE does not induce GILZ expression. YopE possesses GTPase activating activity that has been shown to reduce RhoA, Rac1, and Cdc42 activity in *vitro* [45] but seems to act preferentially on Rac1 and RhoG in *vivo* [42,46]. Evidence that modulation of Rho GTPases may affect GILZ expression was provided by data showing that overexpression of RhoA or RhoB partially inhibited transcriptional activity of the *GILZ* promoter. However, inhibition of either RhoA, RhoB or both using siRNA did not upregulate GILZ expression. On the one hand these data argue that inhibition of those Rho GTPases may play a minor or no role for GILZ expression. On the other hand they also emphasize that we are far away from understanding all the consequences of enzymatic modifications of GTPases induced by pathogenic bacteria.

Searching for additional stimuli, we focused on the strong toxin B induced GILZ expression. Besides inhibiting Rho GTPases, toxin B is also activating pro-inflammatory signaling. Earlier reports [31,32,47] pointed out that treatment of cells with toxin B resulted in MAP kinase activation that is independent of Rho GTPase UDP-glucosylation. Since inhibition of MAP kinases inhibited GILZ expression partially, our results suggest at least a minor contribution of p38 as well as ERK activity in enhancing toxin B induced GILZ expression. However, it is very unlikely that IL-10 acts as an effector for indirect toxin B induced GILZ expression, because pre-treatment of toxin B stimulated HeLa cells with 100 to 200 ng/ml neutralizing anti-human IL-10 antibody [JES3-29F1, BD Pharmingen] had no impact on toxin B induced GILZ expression (data not shown).

Taken together, rather than suggesting a linear induction pathway, expression of (inducible isosform 2) GILZ appears to be regulated by diverse signals induced by some pathogenicity factors.

To shed some light on the downstream events of GILZ induction by YopT and toxin B we examined activation of the GILZ promoter. Previous studies of GILZ promoter activation revealed two principal mechanisms so far: IL-2 deprivation leads to binding of FoxO3 to the forkhead responsive elements (FIRE) and transcriptional activation. Deletion of the three distal glucocorticoid responsive elements of the GILZ promoter abrogates DEX mediated transcriptional activation of GILZ, demonstrating a crucial role of GRE elements for DEX mediated GILZ expression [11,33]. Our data show that similar to DEX, toxins modifying GTPases such as *C. difficile* toxin B and YopT lead to trans-activation of the GILZ promoter. Further studies using shorter GILZ promoter elements demonstrated that in contrast to DEX, a truncated promoter fragment starting at −416 is sufficient for toxin B-induced GILZ promoter trans-activation, however.

Mutations affecting cis-elements in this region revealed that only mutation of a canonical E-box (myc1) abrogates toxin B induced *GILZ* promoter activity. This further differentiates *GILZ* induction by toxin B from that promoted by IL-2 deprivation or glucocorticoids and defines a novel activation mechanism of the *GILZ* promoter. In line with this finding the binding of transcription factors to this canonical E-box increased after stimulation with toxin B or infection with *Y. enterocolitica* pYV*+,* but not with a ΔyopT deletion mutant. This strongly supports the idea that toxin B and YopT use a common downstream pathway to induce GILZ expression.

Previous studies reported interaction of the ubiquitously expressed upstream stimulatory factors (USF)-1 and -2 with canonical E-box regulatory elements (CANNTG) [34]. Consequently, binding of USF-1 and USF-2 as key elements in toxin B-mediated gene expression in response to bacterial virulence factor action.

In addition, we tested the hypothesis whether YopT induced GILZ expression may contribute to *Tessina* mediated inhibition of NF-kB activity. By overexpression of GILZ we could demonstrate that GILZ is able to inhibit NF-kB activation in epithelial cells. Our data are in line with an earlier report showing that GILZ inhibits NF-kB activation in T cells [14]. Having shown that YopT induces GILZ and that GILZ inhibits NF-kB activation, we postulated that YopT might also be involved in *Tessina*-mediated inhibition of NF-kB activation. However, while infection with *T. enterocolitica* pTTSS-yopE55+yopT translocating only YopT into HeLa cells led to GILZ expression, inhibition of NF-kB activity was not achieved (data not shown). It has been published earlier that *T. pseudotuberculosis* YopT moderately inhibited NFκB activation [25]. We could reproduce this finding, however in our hands only overexpression of YopT into HeLa cells resulted in some inhibition of NF-kB activity. Moreover an association between YopT induced GILZ expression and the slight reduction of NF-kB...
activity could not be shown. While the difference in the effectiveness of YopT translated while Yersinia infection is most likely due to the use of different strains, the pathways of YopT mediated NFκB inhibition in either case are probably the same. As discussed by Viboud et al. [25] Yop GTPases promote IkB degradation by JNK activation [50] or by the PKC-ERK pathway [51]. Removal of Yop GTPases from their membrane anchor by YopT therefore should inhibit these ways of NFκB activation and account for the decreased NFκB activity resulting from YopT action.

We also investigated a possible role of GILZ in the induction of apoptosis by toxin B which has been shown to trigger apoptosis by caspase dependent as well as independent pathways. However, knock-down of GILZ did not significantly change the ratio of apoptotic cells.

Therefore, the physiological relevance of GILZ in bacterial infections stays elusive. Several studies highlighted putative physiological effects of GILZ in mice which might potentially modulate immune responses against Yersinia. On the one hand, transgenic mouse overexpressing GILZ show augmentation of Th1 cytokine production as well as induction of Th1 response, leading to protection of mice in a colitis model [54]. On the other hand LPS, an important cofactor triggering inflammatory response in bacterial infection dampened GILZ expression in cultured alveolar macrophages as well as in lung tissues [55] indicating that bacterial factors not only upregulate but also downregulate GILZ expression. The respective contributions of the opposing effects of different bacterial components thus may influence the magnitude and the role of GILZ expression in bacterial infections. For instance, in case of C. difficile infection it was shown that the C. difficile toxin A and the here characterized GILZ inducer toxin B induce a strong inflammatory response [56] and are responsible for the acute inflammation in the gut characterized by pseudo-membrane formation [57]. However, besides the toxins A and B several other components of C. difficile like surface layer proteins [55] or flagella also induce inflammation [56]. Interestingly the pro-inflammatory response and intestinal secretion in the gut induced by C. difficile toxin A can be reduced by exogenous DEX treatment and is also down-regulated by endogenous glucocorticoid production [58]. From this data, one may speculate that both toxin A as well as glucocorticoid induced GILZ might be of physiological relevance during C. difficile infection in limiting the pro-inflammatory response and intestinal secretion. The study of GILZ knockout mice may help to verify this hypothesis.

GILZ knockout mice lacking all four isofoms were recently described and it was demonstrated that GILZ expression results in infertility of male mice but had no impact on inflammatory responses at all [59]. Thus for instance, no difference in the pro-inflammatory response after simultaneous treatment with DEX and LPS was observed between bone marrow derived macrophages of GILZ knockout or wildtype control mice indicating that GILZ has rather no impact on inflammatory responses which is in line with our data. Nevertheless, these mice might prove useful to elucidate the role of GILZ in mouse models of Yersinia enterocolitica or Clostridium difficile infection.

Taken together we could clearly demonstrate that - in addition to previously shown trigger mechanisms - some bacterial toxins such as YopT and toxin B induce GILZ expression. In contrast to dexamethasone treatment and IL-2 deprivation, toxin B and Yersinia YopT mediated regulation of GILZ expression depends on binding of USF-1 and USF-2 to the GILZ promoter. These results open a new branch of investigation in the field of host pathogen interactions.

Materials and Methods

Bacterial Strains and Growth Conditions

All bacteria were grown in Luria-Bertani broth (LB). For infection with Y. enterocolitica strains or E. coli HB101 plv1914 (E. coli plv1914) [60], overnight cultures grown at 27°C or 37°C, respectively, were diluted to an OD₆₀₀ of 0.2 in LB and incubated for 3 h at 37°C. All bacterial strains used in this study are listed in Table 1. The strains ΔyopT yopO⁻ or ΔyopT yopO⁻ ΔrpoA were generated by electroporation of the plasmids pIM279 or pISO1, respectively, into the strain ΔyopT [61].

Generation of Yersinia Mutants

To generate the pTTSS yopO strain secreting only YopO but no other Yops, the plasmid pBM-yopO was electroporated into WA-314 pTTSS and positive colonies were screened by selection with chloramphenicol. To yield the plasmid pBM-yopO the pYV plasmid of wild type Y. enterocolitica strain WA-314 was used as a template to amplify the yopO gene in two fragments, thereby eliminating the internal BamHI site using the primers 5'-GGA TCT ATG AAA ATC ATG GGA ACT ATG TCA CC-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' for fragment 1 and 5'-GTC TGAG TCA CAT CTA TCC CCG-3' and 5'-ATC GTG CAA CCT TCA CTC GAG AGG ATT CAT CAG GAC GAC ACA CAC-3' for fragment 2. Both fragments were used as templates in a PCR yielding the unified yopO gene, using the primers 5'-GGA TCT ATG AAA ATC ATG GGA ACT ATG TCA CC-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' as primers and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' as a template to amplify the yopO gene in two fragments, thereby eliminating the internal BamHI site using the primers 5'-GGA TCT ATG AAA ATC ATG GGA ACT ATG TCA CC-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' for fragment 1 and 5'-GTC TGAG TCA CAT CTA TCC CCG-3' and 5'-ATC GTG CAA CCT TCA CTC GAG AGG ATT CAT CAG GAC GAC ACA CAC-3' for fragment 2. Both fragments were used as templates in a PCR yielding the unified yopO gene, using the primers 5'-GGA TCT ATG AAA ATC ATG GGA ACT ATG TCA CC-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' as primers and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' as a template.

DNA Constructs

The GILZ (p2088) comprising the nucleotide sequence from -2088 to +20 flanked by KpnI and HindIII sites was inserted into pGL3 basic (Promega, Madison, WI) resulting in p2088-Luc [11]. The generation of the constructs p1940, p1526, p854, p416, GRE 1+2 mut and FHRE 1+3 mut has been described elsewhere [11,33]. Point mutations were introduced into the plasmid p1940-Luc by the Quick change PCR based strategy (Stratagene, Cedar Creek, USA) to generate the additional constructs using the following phosphorylated primers: c-myc 1- mut, 5'-GAC GCA GGC GGC TCC TCC-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3'. The yopO gene together with the yopO and yopO promoter region was amplified using the primers 5'-GGA TCT ATG AAA ATC ATG GGA ACT ATG TCA CC-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' and 5'-GTC TCT GGT CCA GAT GCT TCT GAA TCC TCT GCA GTG AAG GTT CGA CG-3' as a template.

In order to generate the expression plasmid pHM6-GILZ-HA, the coding sequence of human GILZ was amplified from cDNA of dexamethasone stimulated HeLa cells using the primers 5'-GCA GCC AAG CCT TAT GAA CAC CGA AAT GTA TCA G-3' (fw) and 5'-GGA CAC GTC CCT TAC GTA CAC CGC AGA ACC-3' (rev) and ligated into pCRH Blunt TOPO (Invitrogen). The
**Table 1.** Bacterial strains used in this study.

| Designation | Genotype or Description | Reference or source |
|-------------|-------------------------|---------------------|
| pYV+        | Y. enterocolitica serotype O8; clinical isolate; WA-314 pYV08+ | [68]          |
| pYV         | Serotype O8, virulence plasmid cured derivative of Y. enterocolitica WA-314 (WA-C) | [68]          |
| WA-314 pKD46 | WA-314 pKD46          | [69]          |
| ΔyopE       | WA-C pYVΔyopE 17–203  | [69]          |
| ΔyopH       | WA-C pYVΔyopH17–455   | [69]          |
| ΔyopM       | WA-C pYVΔyopM 17–351   | [69]          |
| ΔyopP       | WA-C pYVΔyopP 17–272   | [69]          |
| ΔyopT       | WA-C pYVΔyopT          | [61]          |
| pYVΔyopT    | WA-C harbouring pYVΔyopT and pBC18R pYV yopT, sycT (pIM279) | [20,61], this study |
| pYVΔyopTΔC139A | WA-C harbouring pYVΔyopT and pBC18R pYV yopT ΔC139A, sycT (pISO1) | [61,70], this study |
| pYV505      | Tn5 insertional inactivation of ycd | [18]          |
| pTTSS       | WA-C with plasmid pTTSS encoding the type three secretion system and YadA but lacking YopE, YopH, YopM, YopO, YopT and YopP | [71]          |
| pTTSS yopE ΔCA3, ova | WA-C pTTSS with plasmid pME53-Ova247–355 | [72]          |
| pTTSS yopE yopP | WA-C pTTSS with plasmid pME53-yopT | [72]          |
| pTTSS yopE | WAC pTTSS with plasmid pBM53-yopE | [72]          |
| pTTSS yopE yopP | WAC pTTSS with plasmid pBM53-yopP | [72]          |
| E.coli plv 1914 | E. coli HB101 strain expressing Y. enterocolitica Inv | [66]          |
| pCE138-G1   | HindIII - SaI fragment of yopE 138-gfp in pACYC184; HindIII - yopE 138 - BamHI - gfp - SaI | [73]          |
| pTTSS yopO  | WA-C pTTSS with plasmid pBM-yopO | [74], this study |

All PCR amplificates have been checked by sequencing prior to subcloning, pNFkB-Luc was purchased from Stratagene (Amsterdam, Netherlands) and pCMV-ß-galactosidase (pCMV-ß-gal) from Clontech (Palo Alto, CA), pUHD-USF1mutBR (USF-1 DN) was a kind gift from M. Eilers [64].

**Cell Culture and Infection**

HeLa cervical epithelial cells (ATCC CCL-2.1) were grown in RPMI 1640 (Biochrom KG, Berlin, Germany) supplemented with 10% fetal bovine serum (Sigma Chemical, St. Louis, MO), 2 mM L-glutamine (Biochrom KG; Berlin, Germany), penicillin (100 U/ml), and streptomycin (100 µg/ml) (Biochrom KG; Berlin, Germany) in a humidified 5% CO₂ atmosphere at 37°C.

For infection, cells were washed with PBS and incubated in antibiotic-free medium. Cells were infected with a MOI of 20 or as indicated and incubated at 37°C. After 1 h gentamicin (100 µg/ml) was added to kill extracellular bacteria. At indicated time points after infection, cells were lysed to perform subsequent analyses.

**Transfection of siRNA**

siRNA was purchased from Qiagen, Hilden, Germany and comprised the following sequences (sense strand); siUSF1, 5’-GACC CCA ACC AGU GGU GCU ATTtT-3, siUSF2, UAG CCA CAC UGG UUG GGU CdtTdT-3’ [65], siC 5’-UUC UCC GAA CGU GUC AGC UdTdT-3’ (AllStars negative control siRNA), siGILZ 5’-CAC CGU GGU GAA GAC CGU G dTdT-3’. siRNA transfection was done using HiPerfect (Qiagen,Hilden, Germany) transfection reagent according to the manufacturers fast-forward protocol, adding first siRNA mixture and then cells (6×10⁴ HeLa

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*HindIII/Pmel GILZ fragment of pCRII Blunt-GLZ was then ligated into the expression vector pHM6 (Roche Diagnostics Mannheim, Germany)*.

The plasmids pHM6-YopT or pHM6-YopTΔC139A were generated by PCR using the primers 5’-GGT ACC ATG AGT ATT CAC GGA CAC-3’ (fwd) and 5’-GGC GCC GCC GGT TGT AAA CCT CCT TGG-3’ (rev) and as templates pIM279 or pISO1, respectively. After ligation into pCRII Blunt TOPO, the amplicons were cut with HindIII or NotI and ligated into pHM6 resulting in pHM6-YopT and pHM6-YopTΔC139A.

For constitutive expression of Rho GTPases in mammalian cells we generated the following constructs: RhoA: pHM6-RhoA225N-HA was generated by amplification of RhoA225N inserted into pGEX-2T [62] using the primers 5’-ACT CAT AAG CCT TGG TGC CAT CCG GAA GAA ACT GGT G-3’ (fwd) and 5’-ACA GTT TTC ACA AGA CAA GCC AAC CAC CAG-3’ (rev). After ligation into pCRII Blunt TOPO, RhoA225N was donated as a HindIII/EcoRI fragment to pHM6. Cdc42/RhoB/TC10: Coding sequences contained in pRK5 [63] have been amplified with 5’-GGT ACC ATG AGT CAT AAG TGT GTT GTT-3’ (fwd) and 5’-GGC GCC GCC GGT CAT AGC AGC ACA CAC CT-3’ (rev) (Cdc42), 5’-AAC TGG CAT GTT GCC CAT CCG C-3’ (fwd) and 5’-GAA TTC TTC TCA TAG CAC CTT GCA GCA GTT GAT GCA G-3’ (rev) (RhoB) or 5’-AAC TGG CAT GCC CCG AGC CCG CCG C-3’ (fwd) and 5’-GGG GCC GCC GGT CAC GTA ATT AAA CAA CAC-3’ (rev) (TC10). After ligation into pCR2.1 TOPO, insertions were donated as HindIII/EcoRI (RhoB) or HindIII/XhoI (TC10) fragments to pHM6. Rac1: pCDNA3.3-HA-Rac1 was a kind gift from Klaudia Giehl.

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cells in the 24-well format for luciferase experiments, 5·10^5 HeLa cells in the 6-well format for western blot analysis. Minimal siRNA concentrations yielding extensive knockdown were applied to avoid off-target effects. End concentrations were 100 nM for siUSF1/2 and 10 nM for siGILZ. SiRNA transfection was performed 72 h prior to further treatment when using siUSF1/2, or 48 h for GILZ siRNA.

**Transient DNA Transfection**

For DNA transfection, 5·10^4 HeLa cells were seeded in 24-well plates and transfected on the following day in Opti-MEM (Invitrogen; Karlsruhe, Germany) with DNA constructs using ExGen transfection reagent (Fermentas, Ontario, Canada) according to the manufacturer's instructions. After 4 h, the transfection medium was replaced by the normal culture medium and cells were incubated for another day at 37°C. For luciferase assays cells were transfected with 125 ng of the respective luciferase reporter construct and co-transfected with 125 ng of pCMV-β-gal to assay transfection efficiency. For expression of Rho GTPases, GILZ, USF-1 DN or YopT, cells were transfected with 250 ng of the respective expression plasmids.

**Stimulants/Inhibitors**

Clostridium difficile toxin TcdB10463 was kindly provided by Dr. I. Just (Institute of Toxicology, Hannover Medical School, Hannover, Germany). For delivery of C3 transfease into cells, the C2IN-C3lim fusion toxin was used together with the activated transport component C2IHa from the C. botulinum C2 toxin [27]. Dexamethasone was purchased from Sigma Chemical (St. Louis, MO).

MAPK inhibitors SB 201290 and PD 98059 were purchased from Calbiochem (E. Merck, Darmstadt, Germany). Both inhibitors were used at final concentrations of 10 μM. Inhibitor treatment started 1 h before toxin B treatment and was continued until cells lysis. Cell viability was confirmed by microscopic observation.

**Quantitative RT-PCR Analysis**

Total RNA of infected HeLa cells in 6-well plates was extracted using the RNeasy Mini Kit (Qiagen, Hilden, Germany). 2 μg of RNA were reverse transcribed as described earlier [66]. Real-time PCR was performed on a LightCycler (Roche Diagnostics, Mannheim, Germany), using LC-FastStart DNA Master SYBR Green Kit (Roche Diagnostics; Mannheim, Germany), using LC-FastStart DNA Master SYBR Green Kit (Roche Diagnostics; Mannheim, Germany). *GILZ* mRNA expression was quantified using the primers 5'-GGG CAG AGC CAC TAA ACT TG-3' and 5'-GCC TTC ACG AAA CAG AGG AG-3'. As a standard an ampiclon generated with these primers was used. The Real-time PCR was carried out with an annealing temperature of 68-58°C for 45 cycles. *GILZ* mRNA expression levels were normalized to 18S rRNA expression as secondary control.

**Electrophoretic Mobility Shift Assays (EMSA)**

GILZ Induction by Bacterial Toxins

To evaluate luciferase activity, the Luciferase Reporter Gene Assay, high sensitivity (Roche Diagnostics, Mannheim, Germany) was used. Cells were washed twice with PBS and lysed with luciferase buffer according to the manufacturer’s instructions. Lysates were centrifuged and supernatants removed for determination of β-galactosidase activity. Luciferase activity was standardized on β-galactosidase activity and protein concentration (relative light units).

**Sequential ADP-ribosylation of Rho Proteins**

The ADP-ribosylation status of Rho was determined by sequential ADP-ribosylation. To this end, the cell lysates were incubated for 20 min at 37°C with biotin-labelled NAD+ and C2IN-C3lim (100 ng/mL). The proteins were separated by SDS-PAGE, blotted onto nitrocellulose and the biotin-labelled, i.e. ADP-ribosylated Rho was detected with streptavidin-peroxidase by Western blotting.

**Antibodies and Immunoblotting**

Antibodies used for EMSAs and immunoblotting were purchased from Santa Cruz Biotechnology, Santa Cruz, CA (rabbit anti-human Myc (N-262), rabbit anti-human USF-1 (H86), rabbit anti-human USF-2 (C20), mouse anti-RhoA (26C6)), from Cyto-skeleton Inc, Chicago, USA (mouse anti-Rac1 mAb, mouse anti-CDC42 mAb), from BD Bioscience, Heidelberg, Germany (mouse anti-Rac1 clone 102) and from Cell Signaling Technology, Boston, MA (rabbit anti-RhoB). Polyclonal antibodies directed against GILZ were prepared by immunizing rabbits (Biosen, Berlin, Germany) with a GST-GILZ fusion protein. Subsequently, serum was affinity purified using recombinant GST-GILZ protein. To detect GILZ 4 h or as indicated after infection/stimulation by immunoblotting, cells were lysed by three freeze/thaw cycles in PBS containing complete protease inhibitor cocktail (Roche Diagnostics, Mannheim, Germany) plus DTT (400 μM). Cell fragments were spun down by centrifugation at 15,000 g for 10 min at 4°C. In assays the membrane fraction was analysed the pellet was dissolved in a buffer containing 150 mM NaCl, 20 mM Tris-HCl, pH 7.6, 1% Triton X and protease inhibitor cocktail. Protein concentration was determined by the Bradford method using Bio-Rad protein assay (Bio-Rad, Hercules, CA) and up to 120 μg of protein were loaded on a 15% SDS gel. Separated proteins were transferred electrophoretically to an Immobilon-P PVDF membrane (Millipore, Bedford, MA). The membrane was blocked with 50% soy milk. For detection of GILZ expression the polyclonal rabbit anti-GILZ antibody was used followed by incubation with a goat anti-rabbit IgG (H+L) as a secondary antibody and a horseradish peroxidase-conjugated (HRP) donkey anti-goat IgG (H+L) antibody (Jackson Immuno Research Laboratories, West Grove, PA) as a tertiary antibody. Immunoblots were stripped and actin expression was determined by using a mouse anti-actin antibody (Sigma-Aldrich, St. Louis, MO) and HRP labeled rabbit anti-mouse IgG antibody (DAKO, Glostrup, Denmark). For determination of USF-1 and USF-2 expression as secondary antibody a HRP labeled goat-antirabbit IgG antibody (Santa Cruz Biotechnology, Santa Cruz, CA) was used. The detection was carried out using the chemiluminescence detection kit (ECL) from Amersham Biosciences (Uppsala, Sweden).

**Luciferase Reporter Assay**

To evaluate luciferase activity, the Luciferase Reporter Gene Assay, high sensitivity (Roche Diagnostics, Mannheim, Germany) was used. Cells were washed twice with PBS and lysed with luciferase buffer according to the manufacturer’s instructions. Lysates were centrifuged and supernatants removed for determination of protein content, β-galactosidase and luciferase activity. Luciferase activity was standardized on β-galactosidase activity and protein concentration (relative light units).
nondenaturating polyacrylamide gel using 0.5 x TBE (25 mM Tris-HCl, 25 mM boric acid, 0.5 mM EDTA) as running buffer. Gels were transferred to Whatman 3M paper and dried under vacuum. Protein binding was assessed via autoradiography.

Measurement of Apoptosis
Apoptosis was measured as the fraction of cells with hypoploid nuclei [67]. Briefly, cells were harvested 24 h or 48 h after treatment, washed and resuspended in hypotonic lysis buffer containing propidium iodide. After 10 min incubation on ice, one volume of PBS was added and cells were analyzed on a FACScalibur Cytometer (BD Biosciences; Heidelberg, Germany).

Statistics
If not stated otherwise, data shown in the figures are from representative experiments. Comparable results were obtained if not otherwise stated in a total of at least two experiments. Differences between mean values were analyzed using either one-way ANOVA or the Student’s t test. P<0.05 was considered statistically significant.

Supporting Information
Figure S1 Overexpression of GILZ or YopT inhibits NF-kB promoter activity. HeLa cells were transfected with pNF-kB-Luc luciferase reporter and pCMV-β-gal for 24 h and (A) co-transfected with either pHM6 or pHM6-GILZ. Subsequently, HeLa cells were infected with E. coli pInv1914 expressing invasin (MOI 100) for 6 h and luciferase assays were performed. Means ± SD of two independent experiments are shown. (B) HeLa cells were transfected with pNF-BB Luc and pCMV-β-gal and co-transfected with pHM6, pHM6-YopT+ or pHM6-YopT,C139A for 24 h. Subsequently HeLa cells were left untreated or infected with E. coli pInv1914 expressing invasin (MOI 100) for 6 h and luciferase assays were performed. Results of A and B are expressed as fold induction compared to uninfected cells treated with empty vector control and represent the mean ± SEM of a representative experiment performed in quadruplicates. Asterisks indicate a significant difference between NF-kB activation of cells transfected with pHM6 compared to cells transfected with pHM6-YopT+ (p<0.05). (C, D) HeLa cells were transfected with siGILZ and cultured for 24 h and subsequently transfected with pHM6 or pHM6-YopT for additional 24 h. Cell lysates were harvested to determine GILZ and β-actin expression by immunoblot. NF-kB driven luciferase activity was assayed described for B. Means ± SEM of three independent experiments. (TIF)

Figure S2 HeLa cell intoxication by C3 toxin Rho ADP-ribosylation. HeLa cells were incubated at 37°C with C2IN-C3lim (100 ng/mL) + C2IIa (200 ng/mL). After 2, 4 and 6 h pictures were taken to demonstrate the C3-induced change in cell morphology (A) and the cells were lysed. B. The percentages of cells showing “C3-morphology” were calculated from the pictures. Values are given as mean ± S.D. (n = 3); ** p<0.005. C. The ADP-ribosylation status of Rho from the cells was determined by sequential ADP-ribosylation. To this end, the cell lysates were incubated for 20 min at 37°C with biotin-labelled NAD+ and C2IN-C3lim (100 ng/mL). The proteins were separated by SDS-PAGE, blotted onto nitrocellulose and the biotin-labelled, i.e. ADP-ribosylated Rho was detected with streptavidin-peroxidase by Western blotting. The ADP-ribosylated Rho is shown. Comparable amounts of blotted lyase proteins were confirmed by Ponceau S-staining (not shown). (Note: In this experimental setting unlabeled Rho ADP-ribosylation in the intact cells competes with biotin-labelled ADP-ribosylation after lysis. A strong signal therefore means that Rho was not ADP-ribosylated in the intact cells, a weak signal indicates ADP-ribosylation of Rho by the toxin in the intact cells prior to lysis). (TIF)

Figure S3 Impact of GILZ on toxin B triggered apoptosis. HeLa cells were transfected with siGILZ for 48 h (A) and subsequently stimulated with Toxin B for additional 24 h or 48 h. Apoptotic cells were detected by Nicoletti assay. Results are expressed as mean ± SEM of three independent experiments (B). (TIF)

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Author Contributions
Conceived and designed the experiments: MK IBA HB EB. Performed the experiments: MK DG TG PG BM SB SL. Analyzed the data: EB MK DG TG PG SB HB. Contributed reagents/materials/analysis tools: SM BL SL. Performed the experiments: MK DG TG PG BM SB SL. Analyzed the data: EB MK DG TG PG SB HB. Wrote the paper: MK EB. Substantial corrections and discussion of manuscript: BL MLAL MP IS HB RZ IBA.

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