Ecological implications of gene regulation by TfoX and TfoY among diverse Vibrio species

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Summary
Bacteria of the genus Vibrio are common members of aquatic environments where they compete with other prokaryotes and defend themselves against grazing predators. A macromolecular protein complex called the type VI secretion system (T6SS) is used for both purposes. Previous research showed that the sole T6SS of the human pathogen V. cholerae is induced by extracellular (chitin) or intracellular (low c-di-GMP levels) cues and that these cues lead to distinctive signalling pathways for which the proteins TfoX and TfoY serve as master regulators. In this study, we tested whether the TfoX- and TfoY-mediated regulation of T6SS, concomitantly with natural competence or motility, was conserved in non-cholera Vibrio species, and if so, how these regulators affected the production of individual T6SSs in double-armed vibrios. We show that, alongside representative competence genes, TfoX regulates at least one T6SS in all tested Vibrio species. TfoY, on the other hand, fostered motility in all vibrios but had a more versatile T6SS response in that it did not foster T6SS-mediated killing in all tested vibrios. Collectively, our data provide evidence that the TfoX- and TfoY-mediated signalling pathways are mostly conserved in diverse Vibrio species and important for signal-specific T6SS induction.

Introduction
Secretion systems that enable the export of macromolecules across membranes are often associated with pathogenesis in Gram-negative bacteria. However, these machineries and their substrates also play important roles in the interactions between bacteria and other microorganisms in their natural environment (Costa et al., 2015; Green and Mecsas, 2016). An important secretion system, the type VI secretion system (T6SS), was first described in Vibrio cholerae and is a syringe-like killing device involved in virulence and interbacterial warfare (Pukatzki et al., 2006; Ho et al., 2014; Russell et al., 2014). T6SSs are present in ~25% of sequenced Gram-negative bacteria, including pathogenic and nonpathogenic species, and are defined by a set of 13 core components (Das and Chaudhuri, 2003; Bingle et al., 2008; Boyer et al., 2009). These components build a macromolecular complex that shares both structural and functional homology with contractile bacteriophage tails, such as in the phage T4 (Leiman et al., 2009; Basler, 2015; Taylor et al., 2018). Briefly, a tail-like structure, which is composed of hemolysin coregulated protein (Hcp) hexamers and sharpened by spike-like tip proteins (VgrG and PAAR), is surrounded by a contractile sheath structure and polymerizes onto a membrane-spanning multiprotein complex. When the sheath contracts, the effector-decorated tip complex is propelled outwards together with the inner tube to puncture the membrane of a neighbouring target cell and inject toxic effector proteins (reviewed by Zoued et al., 2014; Cianfanelli et al., 2016). Each T6SS encodes its own sets of variable effector proteins that determine the specificity and the biological function of the system, as they can impact both eukaryotic host cells and bacterial competitors. To prevent self-intoxication and to kin-discriminate siblings, bacteria produce effector-specific neutralizing immunity proteins, which are frequently encoded adjacent to the effector gene in a bicistronic unit (Durand et al., 2014; Alcoforado Diniz et al., 2015). The structural components of the T6SS are mostly encoded within larger gene clusters, which are variable with respect to genetic content and organization. The genes coding for the secreted core components (Hcp, VgrG, PAAR) are often found in smaller auxiliary clusters together with the genes that encode the effectors-immunity pairs (Boyer et al., 2009). Many bacterial genomes encode more than one T6SS, which likely serve different needs (Boyer et al., 2009). Accordingly, bacteria employ a wide variety of regulatory mechanisms, from common cues to specific regulatory
cascade, to ensure appropriate expression of T6SS genes (Miyata et al., 2013), which likely vary within species when multiple T6SS systems are present. Studying the various T6SS systems and their regulation is, therefore, important as such knowledge sheds light onto the evolution of bacterial species in response to new niches and their interaction with environmental hosts.

*V. cholerae* O1 El Tor strains, which are responsible for the current seventh pandemic of cholera (further referred to as pandemic strains), harbour a single T6SS. In contrast, many non-cholera *Vibrio* species encode several T6SSs, such as clinical isolates of *V. parahaemolyticus* and environmental isolates of *V. vulnificus*, both of which carry two T6SSs (Yu et al., 2012; Church et al., 2016). The genome of the squid symbiont *V. fischeri*, on the other hand, codes for one T6SS even though a recent study by Speare et al. identified a secondary T6SS in several *V. fischeri* isolates that fostered niche domination within the squid’s light organ (Speare et al., 2018).

The T6SS of pandemic *V. cholerae* is silent under standard laboratory conditions (Pukatzki et al., 2006). In contrast to pandemic strains, the O37 serogroup strains V52 and ATCC25872 are toxigenic but nonpandemic and their T6SS is constitutively activated, as is the case for V52 and ATCC25872 are toxigenic but nonpandemic and contrast to pandemic strains, the O37 serogroup strains are silent under standard laboratory conditions (Pukatzki et al., 2006). The genome of the squid symbiont *V. fischeri*, on the other hand, codes for one T6SS even though a recent study by Speare et al. identified a secondary T6SS in several *V. fischeri* isolates that fostered niche domination within the squid’s light organ (Speare et al., 2018).

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Interestingly, *V. cholerae* and most other members of the genus *Vibrio* (vibrios) contain a TfoX homologue named TfoY (Pollack-Berti et al., 2010; Metzger et al., 2016). However, despite the homology to TfoX and its common annotation as a competence regulator, TfoY is neither essential nor sufficient for competence induction in *V. cholerae* (Metzger et al., 2016). TfoY also induces the T6SS of *V. cholerae*, though the TfoX and TfoY regulons are non-overlapping, with TfoX co-inducing competence and TfoY orchestrating a defensive reaction. This defensive reaction includes enhanced motility and the co-production of the T6SS with additional extracellular enzymes, some of which are known to intoxicate amoebal predators (Metzger et al., 2016; Van der Henst et al., 2018). Moreover, TfoX is naturally produced upon growth on chitinous surfaces (Meibom et al., 2005), while TfoY production is inhibited by c-di-GMP, an secondary messenger in bacteria, such that it is only present at low intracellular c-di-GMP levels (Inuzuka et al., 2016; Metzger et al., 2016).

Given the presence of TfoX and TfoY in many vibrios, we aimed to understand their role in non-cholera species. We show that the T6SS activation is highly conserved across vibrios, with the majority of tested species concomitantly inducing natural competence and motility when TfoX and TfoY, respectively, are produced. Interestingly, we observed different scenarios when comparing those *Vibrio* species that encode more than one T6SS, with TfoX and TfoY not fulfilling the same inducing role for all T6SSs. We, therefore, conclude that some vibrios specifically induce one or the other T6SS under different environmental conditions driven by either TfoX or TfoY, while other species combine forces to simultaneously induce both T6SSs.

**Results and discussion**

**Conservation of T6SS production by TfoX and TfoY in pandemic isolates of *V. cholerae***

Since its discovery in a nonpandemic strain of *V. cholerae*, it has been reported that the T6SS of pandemic strains is silent under standard laboratory conditions (Pukatzki et al., 2006). We recently showed that the two regulatory proteins TfoX and TfoY foster T6SS production and, accordingly, interbacterial competition in pandemic *V. cholerae* (Borgeaud et al., 2015; Metzger et al., 2016). The previous study that addressed TfoY, however, was based on a single strain of *V. cholerae*, namely strain A1552 (Yildiz and Schoolnik, 1998). This human isolate originated from foodborne transmission on an airplane returning to the US from Peru/South America (Blokesch, 2012a) and belongs to the West-African South American (WASA) lineage of the currently ongoing seventh cholera pandemic (Domman et al., 2017; Matthey et al., 2018).
are frequently reported, this study was designed to first confirm the generality of TfoX- and TfoY-dependent T6SS activation in pandemic strains of *V. cholerae*. To do this, we tested five isolates from three different countries (Bangladesh, Peru and Bahrain) for T6SS-dependent killing of *Escherichia coli*. Notably, we used a derivative of the first sequenced isolates of *V. cholerae*, pandemic strain N16961 (Heidelberg *et al.*, 2000), in which the frameshift mutation in the quorum-sensing regulator gene hapR was repaired (Kühn *et al.*, 2014). As demonstrated in Supporting Information Fig. S1, all strains behaved similarly to A1552 upon TfoX or TfoY production. We, therefore, suggest that the dual control over T6SS by these two regulators is conserved among pandemic *V. cholerae*.

**Functionality of TfoX and TfoY homologues from other *Vibrio* species in *V. cholerae***

Pollack-Berti *et al.* showed that TfoX and TfoY existed in all fully sequenced *Vibrio* species (Pollack-Berti *et al.*, 2010), so we wondered whether these homologues would act in a comparable manner in non-cholera vibrios despite the fact that these *Vibrio* species are often adapted to different environmental niches (Le Roux and Blokesch, 2018) and their homologous proteins might, therefore, serve different, adaptation-specific functions. We, therefore, compared the protein sequences of TfoX and TfoY from four different *Vibrio* species, *V. cholerae*, *V. parahaemolyticus*, *V. alginolyticus* and *V. fischeri*, with the additional species chosen because previous studies had already described the genetic organization of their T6SS (Salomon *et al.*, 2013; 2014; 2015; Speare *et al.*, 2018). Moreover, these additional species have isolates that contain two T6SSs. We, therefore, considered the possibility that TfoX and TfoY, which are produced under vastly different conditions in *V. cholerae* (Metzger and Blokesch, 2016), might be specialized for the induction of a single T6SS in those organisms that harbour several systems. By aligning the protein sequences, we observed a high level of protein conservation for TfoX and TfoY, with a sequence identity of 68%/66%, 67%/66% and 59%/67% when the homologous proteins of *V. parahaemolyticus* (TfoX<sub>vp</sub>/TfoY<sub>vp</sub>), *V. alginolyticus* (TfoX<sub>vaa</sub>/TfoY<sub>vaa</sub>) and *V. fischeri* (TfoX<sub>vf</sub>/TfoY<sub>vf</sub>) were compared to the respective homologues of *V. cholerae* (Fig. 1A). Given these identity values, we concluded that the TfoX proteins of *V. parahaemolyticus* and *V. alginolyticus* were more closely related to the homologous protein of *V. cholerae*, whereas TfoX from *V. fischeri* diverged the most, which is in accordance with the phylogenetic distances between the species (Fig. 1A). Notably, TfoY remained equally conserved despite the phylogenetic divergence among the species.

Given the high level of protein conservation, we next tested whether the protein homologues from the non-cholera vibrios could function in *V. cholerae*. For this purpose, we cloned the respective genes preceded by an arabinose-inducible promoter (P<sub>BAD</sub>) onto a site-specific transposon (mini-Tn7) together with the gene that encodes the arabinose-responsive regulatory protein AraC (Supporting Information Table S1). Next, we grew the genetically engineered strains and the parental WT strain in the presence of arabinose (as inducer), isolated their RNA and then quantified the transcript levels of representative T6SS-encoding genes. As shown in Fig. 1B, heterologous production of the diverse TfoX proteins resulted in increased T6SS transcript levels compared to the uninduced conditions, though the level of induction was slightly lower than the production of *V. cholerae*’s own TfoX, especially for the most diverging protein variant from *V. fischeri* (TfoX<sub>vf</sub>). We confirmed these data at the protein level through the detection of the inner tube protein of the T6SS, Hcp (Fig. 1C). Consistent with the production of Hcp and the increase of the T6SS transcript level, these strains were able to kill *E. coli* in an interbacterial competition assay, with the least predatory behaviour observed for the most divergent TfoX<sub>vf</sub>-carrying variant (Fig. 1D).

TfoX of *V. cholerae* was first identified as the master regulator of natural competence for transformation (Meibom *et al.*, 2005; Metzger and Blokesch, 2016). The TfoX-induced competence regulon includes those genes that encode the structural components of the DNA-uptake machinery, which primarily consists of a central pilus structure (encoded by pil genes) (Seitz and Blokesch, 2013) and the DNA binding protein ComEA that is essential for reeling DNA across the outer membrane and into the periplasmic space (Seitz and Blokesch, 2014; Seitz *et al.*, 2014; Matthey and Blokesch, 2016). We, therefore, tested whether the TfoX variants from the other *Vibrio* species would likewise foster competence-gene expression, which was indeed the case (Fig. 1E). Compared to the other variants, the more diverse TfoX<sub>vf</sub> did not induce the same high transcript levels of comEA, which, accordingly, also led to lower transformation frequencies (Fig. 1F). Notably, comEA expression in *V. cholerae* is regulated by an intermediate transcription factor, QstR, which is dependent on the production of both TfoX and HapR (Lo Scudato and Blokesch, 2013; Jaskólska *et al.*, 2018). HapR is the master regulator of the QS system, meaning that comEA expression is dependent on a functional QS system and on high cell density. As *V. fischeri* - but not the other tested *Vibrio* species - lacks a QstR homologue, we suggest that TfoX<sub>vf</sub> induces comEA at lower levels, as it is adapted to the more diverse QS circuit of *V. fischeri* and the absence of QstR (Milton, 2006; Verma and Miyashiro, 2013).

To assess the functionality of the TfoY homologues in *V. cholerae* we first tested the ability of the TfoY variants to foster natural transformation, given that these proteins are frequently annotated as ‘DNA transformation protein
V. induced and uninduced conditions, as shown on the material. Transformation frequencies reexpression. The indicated strains were grown under inducible conditions, and the genomic DNA of A1552-lacZ-Kan served as the transforming tfoY

J. Motility was scored on soft agar with and without arabinose as an inducer. The motility phenotype was quanti

F. Natural transformation using genomic DNA is maintained in a

D, I. Interspecies killing assay with

C, H. Detection of Hcp protein produced by TfoX- and TfoY-expressing cells. Cells were grown in the absence or the presence of the inducer (arabinose) as indicated below the image. Detection of tfoX- and tfoY-derived Hcp was confirmed using Western blots. Identical residues are highlighted in shades of blue (above threshold of 65%).

B–J. Chromosomally-located tfoX and tfoY from chola (V. cholerae) and non-chola Vibrio species (V. parahaemolyticus, V. alginolyticus and V. fischeri) were expressed under the control of the arabinose-inducible PBAD promoter in V. cholerae. The colour code (TfoX-induced, green; TfoY-induced, blue) is used throughout the graphs. The WT strain lacking an inducible copy of tfoX or tfoY served as the negative control. The fold change (tfoX- or tfoY-induced over parental WT strain) of relative gene expression of (B, G) T6SS genes or (E) representative competence genes is shown.

C, H. Detection of Hcp protein produced by TfoX- and TfoY-expressing cells. Cells were grown in the absence or the presence of the inducer (arabinose) as indicated below the image. Detection of $\alpha_{Hcp}$ served as a loading control.

D, I. Interspecies killing assay with E. coli as prey. V. cholerae harbouring different inducible versions of (D) tfoX or (I) tfoY were co-cultured with the prey on LB agar plates supplemented with arabinose. The survival of the prey was determined on selective LB agar plates and is depicted as CFU per ml.

F. Natural transformation using genomic DNA is maintained in a V. cholerae strain expressing tfoX from non-cholera vibrios but is nonfunctional upon tfoY expression. The indicated strains were grown under inducible conditions, and the genomic DNA of A1552-lacZ-Kan served as the transforming material. Transformation frequencies reflect the number of transformants divided by the total number of CFUs. < dl, below detection limit.

J. Motility was scored on soft agar with and without arabinose as an inducer. The motility phenotype was quantified as the ratio between the induced and uninduced conditions, as shown on the Y-axis. Abbreviations: V.c., V. cholerae A1552; V.p., V. parahaemolyticus RIMD2210633; V.a., V. alginolyticus 12G01; V.f., V. fischeri ES114. Bar plots represent the average of at least three independent biological replicates (± SD). Statistical significance is indicated (*p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001).

Fig. 1. TfoX and TfoY proteins from diverse Vibrio species are functional in V. cholerae.

A. Phylogenetic tree and protein sequence conservation of TfoX and TfoY across Vibrio species. The phylogenetic tree was generated on phylot.biobyte.de, a phylogenetic tree generator, based on NCBI taxonomy. The protein sequences of TfoX and TfoY were aligned using ClustalOmega, and subsequently modified with Jalview. Identical residues are highlighted in shades of blue (above threshold of 65%).

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TfoX’ or ‘TfoX-family DNA transformation protein’ due to their similarity to TfoX. However, no transformants were detected upon the production of any of these proteins, indicating their inability to induce natural competence in *V. cholerae* (Fig. 1F). Conversely and consistent with the high sequence similarity of the proteins from the four organisms, all TfoY variants functioned at the same level as the *V. cholerae* TfoY to increase T6SS transcript levels (Fig. 1G), to induce Hcp protein production (Fig. 1H), and, consequently, to foster interbacterial killing of *E. coli* (Fig. 1I). In addition, the four TfoY variants were able to induce motility as a T6SS-independent phenotype, as visualized on swarming agar plates and quantified in Fig. 1J. Taken together, these data indicate that TfoX and TfoY are conserved proteins among these four *Vibrio* species and that the TfoY variants can fully replace the *V. cholerae* innate protein, while the TfoX variants are functionally replaceable at high levels for TfoX_\textsubscript{up}/TfoX_\textsubscript{Vs} and intermediate levels for TfoX_\textsubscript{Vf}.

**TfoX but not TfoY induces *V. fischeri’s* primary T6SS**

While the complementation assay in *V. cholerae* described above provided a first insight into the conservation of TfoX and TfoY from diverse vibrios by indicating their similarity to those of *V. cholerae*, the experiments did not tell us whether these proteins would be involved in similar phenotypes as in *V. cholerae* in their native hosts. We, therefore, genetically engineered the different *Vibrio* species to place inducible copies of their own tfoX and tfoY genes onto a mini-Tn7 transposon that was then integrated into their own genomes (Supporting Information Table S1). Next, we analysed the different phenotypes under uninduced and induced conditions. For *V. fischeri* strain ES114 (Supporting Information Table S1), we showed that TfoX production led to a high induction of T6SS transcripts, while TfoY only partially induced a subset of T6SS genes (Fig. 2A). Consistent with these expression data, we witnessed the T6SS-mediated killing of *E. coli* only upon TfoX induction, while TfoY induction did not change the number of surviving co-cultured prey bacteria (Fig. 2B).

Pollack-Berti *et al.* showed that TfoX was required for natural competence for transformation in *V. fischeri* (Pollack-Berti *et al.*, 2010), as is the case for *V. cholerae* (Meibom *et al.*, 2005). We, therefore, measured relative transcript levels of representative competence genes in this organism and confirmed their induction upon TfoX production (Fig. 2C). Surprisingly, while we only observed a mild induction of comEA relative to the levels of induction seen in *V. cholerae* (Fig. 1), we witnessed an almost 10-fold upregulation of litR, which encodes the main QS regulator LitR (homologue of *V. cholerae’s* HapR). This phenotype differs from what is known about competence regulation in *V. cholerae* in which case hapR transcript levels are left unchanged upon TfoX induction (Meibom *et al.*, 2005; Metzger and Blokesch, 2016). We, therefore, speculate that an increase in LitR levels can compensate for the lack of the intermediate regulator QstR in *V. fischeri*. Indeed, the QstR protein of *V. cholerae* is required for competence-mediated DNA uptake in two ways: (i) through direct and indirect induction of certain competence genes, such as comEA, as described above; and (ii) due to its ability to downregulate the dns gene (Lo Scrudato and Blokesch, 2013; Jaskólska *et al.*, 2018) encoding for the extracellular nuclease Dns, which degrades transforming material outside the cell and within the periplasmic space (Blokesch and Schonlik, 2008; Seitz and Blokesch, 2014). Notably, the HapR protein of *V. cholerae* also silences dns expression, but only partially in the absence of QstR (Lo Scrudato and Blokesch, 2013). It is, therefore, likely that the increased production of LitR upon TfoX production in *V. fischeri* compensates for the absence of QstR to ensure sufficient repression of dns, which, ultimately, would allow DNA uptake to occur. It is not clear, however, why the comEA transcripts were not induced to higher levels, though we recently demonstrated for *V. cholerae* that the comEA gene requires an additional but so far unidentified regulatory input apart from TfoX and QstR (Jaskólska *et al.*, 2018). Indeed, artificial QstR production was sufficient to induce the T6SS genes in this organism at comparable levels as the upstream regulatory protein TfoX, even in the absence of HapR, while the comEA transcript levels were only partially increased compared to a noninduced WT strain. We, therefore, hypothesize that this input signal might be conserved in *V. fischeri*, though missing under the tested conditions. This would lead to a high overall competence-gene induction upon TfoX production but lower comEA transcripts as compared to *V. cholerae* (Fig. 1E).

Lastly, we determined that the motility of *V. fischeri* changed upon TfoX production (Fig. 2D), even though the difference between the TfoY-induced and TfoY-uninduced state was not as pronounced as for *V. cholerae* (Fig. 1J). We argue that this might be caused by a basal production level of TfoY in this organism that does not rely on the artificial induction. A higher basal TfoY level would result in a higher basal motility, which would, therefore, appear as a less pronounced induction upon additional TfoY production. There was also a decrease in motility observable upon TfoX production (Fig. 1D), which is consistent with the idea of a basal TfoY level, as the expression data showed that TfoX significantly lowered the tfoY transcript levels (Fig. 2A). To see whether this is a conserved regulatory phenomenon, we tested whether TfoX production also lowered tfoY levels in *V. cholerae*, which was indeed the case with an average repression of 4.85 (± 0.5)-fold (based on three independent biological experiments; p = 0.0001). For *V. fischeri* TfoY production lowered tfoX transcript levels (Fig. 2A), which was likewise the case for *V. cholerae* though to a
lower extend (1.45 ± 0.1-fold; p = 0.025), suggesting that both proteins work independently of each other and are mutually exclusive. This finding contradicts a previous study on *V. fischeri* in which the authors concluded that TfoY was required for efficient TfoX-mediated transformation (Pollack-Berti *et al.*, 2010). Notably, both studies used different conditions (e.g. inducible TfoX/TfoY expression in this study versus a transposon-inserted mutant of *tfoY* (Pollack-Berti *et al.*, 2010)) and further studies are, therefore, required to conclusively show if TfoY is indeed involved in natural transformation in *V. fischeri* or not, as we suggest based on the herein-described data.

**TfoX and TfoY have a different impact on T6SS induction in V. alginolyticus**

While *V. cholerae* strains and the examined *V. fischeri* strain ES114 contain only a single T6SS, several other *Vibrio* species contain more than one. Hence, we asked whether, and if so how, TfoX and TfoY affect the production of different T6SSs within the same organism. To address this question, we first tested the contribution of these regulators to the two T6SSs (T6SS1 and T6SS2) of *V. alginolyticus* in an *E. coli*-killing assay using wildtype (WT) *V. alginolyticus* or its *hcp1* (referred to as ΔT6SS1 throughout the text), *hcp2* (referred to as ΔT6SS2 throughout the text) or double mutant as a predator. Consistent with a previous report by Salomon *et al.* (2015), WT *V. alginolyticus* did not show interbacterial predation at 37 °C on standard LB medium (e.g. without additional salt; Fig. 3A). Upon production of TfoX or TfoY, however, predation by the WT was significantly increased. This effect was less pronounced but reproducible in the strain that only carried the complete T6SS2 gene set (ΔT6SS1), while a strain that only carried a complete T6SS1 gene set (ΔT6SS2) was not able to kill the *E. coli* cells, indicating that T6SS1 is
Fig. 3. TfoY is a universal T6SS inducer in V. alginolyticus, while TfoX is specific for T6SS2. A–C, E, F. Interspecies killing assay of V. alginolyticus with E. coli as prey. V. alginolyticus strains were co-cultured with E. coli prey for 4 h at (A, E) 37°C or (B, C, F) 30°C on plain LB agar plates (C) or plates supplemented with arabinose (A, B, E, F, to induce tfoX or tfoY, as indicated below the graph). The recovery of the prey is indicated as CFU/ml on the Y-axis.

D. TfoX induces competence genes in V. alginolyticus. The relative expression of representative competence genes was scored in WT and tfoX-induced strains as indicated on the Y-axis.

G. Motility ratio between uninduced and induced V. alginolyticus strains is indicated. Details as in Fig. 2D. Abbreviations: WT, V. alginolyticus strain 12G01; ΔT6SS1, 12G01Δhcp1; ΔT6SS2, 12G01Δhcp2; ΔT6SS1ΔT6SS2, 12G01Δhcp1Δhcp2; ΔtfoX, 12G01ΔtfoX; ΔtfoY, 12G01ΔtfoY; ΔtfoXΔtfoY, 12G01ΔtfoXΔtfoY; ΔluxR, 12G01ΔluxR. Bar plots represent the average of at least three independent biological replicates (± SD). Statistical significance is indicated (*p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001).

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Notably, the genes occurred, indicating the protein TfoX-mediated induction of representative competence tent bacteria (Seitz in ratcheting the DNA into the periplasmic space in competence genes transcripts and bacterial motility upon other regulatory protein levels and a high motility are linked throughout the genus Vibrio, even in organisms that contain lateral flagella in addition to the polar flagellum (McCarter, 2001).

TfoX and TfoY are both dedicated to one specific T6SS in V. parahaemolyticus

Lastly, we tested the contribution of TfoX and TfoY to T6SS regulation in V. parahaemolyticus. As with V. alginolyticus, this bacterium also contains two T6SSs, T6SS1 and T6SS2, which were previously shown to be differentially regulated by HapR in V. cholerae (homologues are LuxR in V. alginolyticus and OpaR in V. parahaemolyticus) (Lo Scrudato and Blokesch, 2012; Metzger and Blokesch, 2016; Jaskólska et al., 2018), which we assumed to be a conserved feature in other Vibrio species (see data below).

In addition to T6SS activation, we also tested the level of competence genes transcripts and bacterial motility upon TfoX and TfoY induction. As shown in Fig. 3D, a significant TfoX-mediated induction of representative competence genes occurred, indicating the protein’s conserved action. Notably, the comEA transcript induction levels again turned out to be unexpectedly low compared to the induction levels in V. cholerae (Fig. 1E). ComEA plays a primary role in ratcheting the DNA into the periplasmic space in competent bacteria (Seitz et al., 2014), and we, therefore, expect that these low transcript levels would lead to insufficient amounts of the ComEA proteins and defective DNA uptake. As comEA is co-regulated by the TfoX- and QS pathways in V. cholerae (Suckow et al., 2011; Lo Scrudato and Blokesch, 2012), comparable to the T6SS genes (Borgeaud et al., 2015; Metzger et al., 2016), we wondered whether QS was impaired in the tested V. alginolyticus strain. As demonstrated in Fig. 3E and F, TfoX-mediated T6SS induction worked efficiently in the WT but was impaired in a luxR-minus strain (the hapR homologue in this organism; Supporting Information Table S1). These data indicate that QS is functional in the WT strain and support the notion that TfoX and LuxR co-regulate the T6SS2, consistent with earlier data that showed a requirement of LuxR for basal T6SS2 activity in V. alginolyticus (Yang et al., 2018). TfoY-mediated T6SS activation, however, occurred independently of LuxR (Fig. 3E and F) consistent with what is known for V. cholerae (Metzger et al., 2016). Based on these data, we conclude that the additional but so far unknown regulatory input required for comEA expression aforementioned (Jaskólska et al., 2018) might also be a prerequisite in V. alginolyticus and that this pathway and not the QS network overall might be non-functional in either the strain used in this study or under the tested conditions or both. The motility-inducing phenotype of TfoY, on the other hand, was very pronounced in V. alginolyticus (Fig. 3G), strengthening the idea that TfoY protein levels and a high motility are linked throughout the genus Vibrio, even in organisms that contain lateral flagella in addition to the polar flagellum (McCarter, 2001).

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**Fig. 4.** TfoX and TfoY each induce a dedicated T6SS in *V. parahaemolyticus.*

A. E. Interspecies killing of *E. coli* by *V. parahaemolyticus*. *V. parahaemolyticus* strains (WT, T6SS1-, T6SS2-, T6SS1/T6SS2-minus or opaR-minus) were co-cultured with *E. coli* prey for 4 h at 30°C on LB agar plates supplemented with arabinose to induce tfoX or tfoY (as indicated below the graph). The recovery of the prey is indicated on the Y-axis. The recovery of the prey is indicated on the Y-axis.

B. Basal T6SS activity is independent of TfoX and TfoY. WT *V. parahaemolyticus* or its tfoX-, tfoY- or tfoX/tfoY-minus variants were tested for their ability to reduce the number of *E. coli* prey cells. The T6SS double mutant (∆T6SS1 ∆T6SS2) served as a control.

C. Relative expression of representative competence genes in WT or tfoX-induced bacteria.

D. Quantification of TfoX- or TfoY-induced motility phenotypes. Details as in Fig. 2D.

E. TfoX- and TfoY-dependent phenotypes in vibrios.

Abbreviations: WT, *V. parahaemolyticus* POR1 (RIMD2210633 derivative); ∆T6SS1, POR1Δhcp1; ∆T6SS2, POR1Δhcp2; ∆T6SS1ΔT6SS2, POR1Δhcp1Δhcp2. Bar plots represent the average of at least three independent biological replicates (± SD). Statistical significance is indicated (*p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001).

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caused this basal T6SS2 activity (Fig. 4B). Interestingly, a recent study showed T6SS2-dependent killing of a *V. parahaemolyticus* strain that lacked a newly identified effector immunity pair (RhsP and RhsPi) by its parental WT strain, even in the absence of TfoX or TfoY induction (Jiang et al., 2018). These data, therefore, suggest that the basal killing activity that we observed against *E. coli* as a prey might be more enhanced against nonimmune siblings or, alternatively, that the experimental conditions were different to the current work and that under such conditions the T6SS2 is highly active. Lastly, we cannot exclude the possibility that strain differences or the domestication of certain strains has caused the difference in basal T6SS activity. Notably, the focus of the current study was on the contribution of TfoX or TfoY to T6SS activity, which we unambiguously demonstrate to enhance T6SS activity in *V. parahaemolyticus*.

Examining other TfoX- and TfoY-mediated phenotypes confirmed the increase of competence gene transcripts and enhanced motility respectively (Fig. 4C and D). However, *comEA* transcript levels were again unexpectedly low even though the induction levels of *qstR* were comparable with what we observed for *V. cholerae* (Fig. 1E). While these data were similar to the above-described observations for *V. alginolyticus* (Fig. 3D), which suggested a *comEA*-specific expression defect, we nonetheless tested whether the *V. parahaemolyticus* strain used in this study was QS proficient or QS deficient. Indeed, previous studies in different *Vibrio* species described gain-of-function mutations in *luxO*, encoding a repressor of HapR/LitR/OpaR synthesis (Gode-Potratz and McCarter, 2011; Kimbrough and Stabb, 2015; Stutzmann and Blokesch, 2016) and clinical *V. parahaemolyticus* isolates with such *luxO* mutations were previously described to be locked in a pathogenic state (Kernell Burke et al., 2015). To exclude that this has happened in the strain we were working with, we first sequenced *luxO* and *opaR* and confirmed that both genes were mutation free. Additionally, we deleted *opaR* from the WT strain and compared this strain to its TfoX- or TfoY-inducible derivatives (Supporting Information Table S1) for an *E. coli*-killing assay. As shown in Fig. 4E, the TfoY-induced prey killing was maintained upon deletion of *opaR*, while the TfoX-mediated T6SS activity was abrogated. Since OpaR is produced at high cell density, its deletion blocks QS-dependent signalling. We, therefore, conclude that the WT strain has a functional OpaR and that TfoX-mediated T6SS induction in *V. parahaemolyticus* requires coregulation by OpaR, similar to what we have shown for *V. cholerae* (Borgeaud et al., 2015; Metzger et al., 2016) and for *V. alginolyticus* (Fig. 3E and F). These data also support the idea that the low expression levels seem specific for *comEA* and not based on a general mutation within the QS cascade. The low level of the *comEA* transcripts also explains why we were unable to naturally transform *V. parahaemolyticus* even after TfoX induction. Interestingly, Chimalapati et al. recently reported rare transformants for competence-induced *V. parahaemolyticus* with frequencies between $1 \times 10^{-8}$ and $2.7 \times 10^{-6}$ (Chimalapati et al., 2018), which corresponds to 0.01% to 1% of what is commonly observed for *V. cholerae* (see Fig. 1 and (Lo Scrudato and Blokesch, 2012) despite the ~1000-fold molar excess of the selective resistance marker that was used as transforming DNA in this study. It should be noted that these experiments were based on the heterologous expression of the *V. cholerae* tfoX homologue from a multicopy plasmid, which makes a direct comparison with the current study on the bacterium’s indigenous TfoX protein impossible.

Our data suggest that TfoX induces the T6SS2 in *V. parahaemolyticus*. As previous studies showed that the chitin sensors ChiS, TfoS and the downstream-regulated small RNA TfoR, which is required for tfoX mRNA translation, are highly conserved in diverse *Vibrio* species (Li and Roseman, 2004; Yamamoto et al., 2011; 2014), we suggest that TfoX is exclusively produced upon growth on chitinous surfaces, as first demonstrated for *V. cholerae* (Meibom et al., 2004; 2005). The production of TfoY is less understood. In a previous study on *V. cholerae*, we showed that lowered c-di-GMP levels led to the production of TfoY under standard laboratory conditions (e.g. in LB medium) and we and others suggested that this c-di-GMP-dependent control occurred at the translational level (Inuzuka et al., 2016; Metzger et al., 2016). Interestingly, Gode-Potratz et al. demonstrated that growth on surfaces lowered the levels of the secondary messenger c-di-GMP in *V. parahaemolyticus* (Gode-Potratz et al., 2011). Consistently, Salomon et al. established that surface sensing correlated with Hcp1 production in this organism (Salomon et al., 2013). Based on these data, we speculated that surface sensing, TfoY production and T6SS1 induction might be linked. To address this hypothesis and especially the link between low c-di-GMP levels and TfoY production, we genetically engineered *V. parahaemolyticus* to carry a translational fusion between its indigenous tfoY gene and the gene that encodes super-folder GFP (sfGFP). In addition, we included a transcriptional mCherry-encoding reporter gene behind this construct, which, ultimately, allowed us to monitor the transcriptional and translational control of tfoY. Next, we incorporated either an empty mini-Tn7 transposon (Tn) or transposons that carried arabinose-inducible genes coding for a c-di-GMP-producing diguanylate cyclase (Tn-vdcA) or for a phosphodiesterase (Tn-cdpA) into this strain (Supporting Information Table S1). These strains as well as the transposon-deficient parental strain and the WT without any fluorescent protein-encoding genes were then grown in the
presence of the inducer arabinose followed by a western blot analysis to visualize the TfoY-sfGFP fusion protein or the transcriptional reporter protein mCherry. Using this approach, we observed that TfoY was produced solely at low c-di-GMP levels (Fig. 4F). Importantly, based on the comparable levels of mCherry in all reporter strains, we concluded that the tfoY transcript levels did not significantly change in response to increased or decreased c-di-GMP levels (Fig. 4F). This finding contradicts a recent study in V. cholerae that concluded that TfoY was induced at both low and high intracellular concentrations of c-di-GMP and that this regulation occurred at the translational and transcriptional levels, the latter due to activation by the c-di-GMP-dependent transcription factor VpsR (Pursley et al., 2018). We, therefore, constructed a similar dual translational-transcriptional reporter strain in V. cholerae (Supporting Information Table S1) and tested this strain under normal, increased or decreased c-di-GMP levels. Using this approach, we were able to detect an increase in the TfoY protein under low c-di-GMP levels compared to normal conditions, while an increase in intracellular c-di-GMP did not result in higher TfoY protein levels (Supporting Information Fig. S2), consistent with our previous findings (Metzger et al., 2016). Importantly, we also did not observe a change in the transcriptional reporter under high c-di-GMP conditions (Supporting Information Fig. S2), despite normal VpsR function in the pandemic V. cholerae strain (A1552) used in this study (Yildiz et al., 2001), which confirmed the data presented above for V. parahaemolyticus. An explanation for this discrepancy might be that we engineered the construct at the native locus of tfoY within the V. parahaemolyticus or V. cholerae genomes, while Pursley et al. were unsuccessful in detecting tagged TfoY variants using a similar approach. They, therefore, decided to construct a TfoY-GFP translational-fusion-encoding reporter gene on a plasmid, which could change the expression pattern compared to the gene’s native locus. In addition, these authors performed their experiments in a Vibrio polysaccharide mutant as parental strain (∆vpsL), which is deficient in biofilm formation (Pursley et al., 2018), while our strain maintained a normal ability for this process.

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Collectively, we conclude that the lowered c-di-GMP levels observed in surface-sensing *V. parahaemolyticus* (Gode-Potratz et al., 2011) might trigger TfoY induction, which induces the T6SS1 and subsequently initiates motility, though the exact mechanism for the latter remains to be discovered. The data provided here for *V. parahaemolyticus*, therefore, support our previous suggestion that TfoY triggers a defensive escape reaction that might allow *Vibrio* species to defend themselves against bacterial competitors and eukaryotic predators. Interestingly, a recent study that profiled the gene expression of *V. parahaemolyticus* reported increased tfoY expression (VP1028; falsely annotated as tfoX and, therefore, discussed as competence regulator in this study) within infected infant rabbits when compared to *in vitro* conditions (Livny et al., 2014), suggesting that TfoY might also play a role *in vivo* and, therefore, in pathogenesis.

**Conclusion**

In this study, we investigated the ecological implications upon TfoX and TfoY production in diverse *Vibrio* species. We showed a conserved pattern of competence and motility induction by TfoX and TfoY, respectively, and demonstrated that the link between the TfoX and TfoY regulatory proteins and T6SSs is highly conserved among cholera and non-cholera *Vibrio* species (Fig. 5), highlighting the general importance of these regulators. Notably, TfoX induced at least one T6SS in every tested species concomitantly with an interbacterial killing phenotype. Hence, we suggest that chitin-mediated TfoX production (Meibom et al., 2005; Metzger and Blokesch, 2016) and the resulting coupling between T6SS-facilitated neighbour killing and competence-mediated DNA uptake (Borgeaud et al., 2015) might be an ancient phenotype maintained in most *Vibrio* species. Our study also suggests that TfoX and TfoY are mutually exclusive (Fig. 2A and data presented above). Indeed, TfoX-mediated tfoY repression seems like a prerequisite for proper colonization of chitinous surfaces, as TfoY-induced motility would otherwise counteract this process. Thus, even upon contact with the surface and the reported decrease in c-di-GMP levels (Gode-Potratz et al., 2011), tfoY translation would not occur on chitin due to the absence/low levels of the template mRNA. In contrast, when vibrios sense non-chitinous surfaces, resulting in the absence of the chemoattracting chitin-degradation product chitobiose (Keyhani and Roseman, 1999), TfoY would be produced. Interestingly, the effects of TfoY turned out to be more versatile compared to TfoX (Fig. 5). While the c-di-GMP-dependent inhibition of TfoY translation seemed conserved, at least in the two tested *Vibrio* species (Fig. 4E and Supporting Information Fig. S2), the regulators seem to have diverged in the different organisms. Indeed, TfoY induces both T6SSs in *V. alginolyticus* and one T6SS in *V. cholerae* and *V. parahaemolyticus* (out of two T6SSs present in the latter bacterium) but this regulator does not impact the activity of the T6SS of *V. fischeri* under the tested conditions. However, several isolates of *V. fischeri*, other than the squid isolate ES114, contain a secondary T6SS. This T6SS is located on a strain-specific genomic island on the small chromosome and was recently shown to contribute to competitor elimination within the host’s light organ (Speare et al., 2018). While we were unable to genetically engineer the T6SS2-containing fish symbiont MJ11 strain for technical reasons, it is tempting to speculate that this secondary T6SS of *V. fischeri* might be controlled by the TfoY protein. In line with this idea is the fact that *V. fischeri*’s secondary T6SS is broadly conserved among vibrios and is similar to the TfoY-inducible T6SS1 of *V. alginolyticus* and *V. parahaemolyticus* (Speare et al., 2018), which we confirmed by BLAST analyses of *V. fischeri*’s T6SS2 genes VFMJ11_A0804 to VFMJ11_A0809. Notably, Speare et al. proposed the possibility that this secondary T6SS2 was present in the common ancestor of *V. fischeri* and related *Vibrio* species but was eventually lost in niche-adapted strains for which interbacterial killing was no longer advantageous. We extend this hypothesis and speculate that niche-adapted vibrios might no longer require a TfoY-mediated defensive response against either competing bacteria or grazing predators. Future studies are required to address these interesting evolutionary hypotheses.

**Experimental procedures**

**Bacterial strains, plasmids and growth conditions**

The *V. cholerae*, *V. alginolyticus*, *V. parahaemolyticus* and *V. fischeri* strains and plasmids used in this study are listed in Supporting Information Table S1. *E. coli* strains DH5α (Yanisch-Perron et al., 1985), TOP10 (Invitrogen), SM10::pir (Simon et al., 1983), S17-1::pir (Simon et al., 1983) and MFDpir (Ferrieres et al., 2010) were used for cloning purposes and/or served as donor in bacterial mating experiments.

Unless otherwise stated, the *V. cholerae*, *V. alginolyticus*, *V. parahaemolyticus* and *E. coli* strains were grown aerobically in Lysogeny broth (LB; 10 g/l of tryptone, 5 g/l of yeast extract, 10 g/l of sodium chloride; Carl Roth) or on LB agar plates at 30°C or 37°C. *V. fischeri* strains were cultured aerobically in LB salt (LBS) medium (10 g/l of tryptone, 5 g/l of yeast extract, 20 g/l of sodium chloride, 20 mM Tris–HCl [pH 7.5], 0.2% glycerol; adapted from (Dunlap, 1989; Graf et al., 1994)) or LBS agar plates at 28°C. LB/LBS motility plates had less agar (0.3%) than standard LB/LBS agar plates (1.5%). The following compounds were added if required at the given concentrations: arabinose (0.02% or 0.2%), diaminopimelic acid (DAP; 0.3 mM) or the antibiotics...
the based genetic engineering technique was used to replace our group (De Souza Silva and Blokesch, 2010; Blokesch, TransFLP gene disruption method previously described by Kan, the resistance gene (below), derived from pGP704-Sac28, was used for genetic A1552 chromosome. Plasmid pGP704-Sac-Kan (see digestion was performed on both the PCR product and the plasmid pGP704-Sac28 (Meibom et al., 2004). Marine broth 2216 (BD Difco™ 2216) was used to isolate single colonies of V. alginolyticus and V. parahaemolyticus after E. coli counter-selection.

Genetic engineering of strains and plasmids

DNA manipulations were performed according to standard molecular biology-based protocols (Sambrook et al., 1982). Enzymes were purchased from the listed companies and were used as recommended by the manufacturer: Pwo polymerase (Roche), Taq polymerase (Promega), restriction enzymes (New England Biolabs). Following initial screening by PCR (using bacterial cells as templates), genetically engineered strains and plasmids were verified by Sanger sequencing (Microsynth, Switzerland).

V. cholerae strains were genetically modified using both a gene-disruption method based on the counter-selectable plasmid pGP704-Sac28 (Meibom et al., 2004) or the TransFLP gene disruption method previously described by our group (De Souza Silva and Blokesch, 2010; Blokesch, 2012b; Borgeaud and Blokesch, 2013). This transformation-based genetic engineering technique was used to replace the tfoY gene by a translational and transcriptional tfoY-mCherry::gfp fusion at the natural locus on the V. cholerae A1552 chromosome. Plasmid pGP704-Sac-Kan (see below), derived from pGP704-Sac28, was used for genetic modifications of V. parahaemolyticus and V. alginolyticus.

To construct the counter-selectable plasmid pGP704-Sac-Kan, the aph gene was amplified from a kanamycin-resistance gene (aph)-carrying transposon using primers with overhanging SspI and Bsal restriction sites. A restriction digestion was performed on both the PCR product and the vector pGP704-Sac28, which were subsequently ligated, to replace the bla gene of pGP704-Sac28 with the aph gene, resulting in plasmid pGP704-Sac-Kan (Supporting Information Table S1).

All pBAD-derived plasmids harbouring tfoX-strep or tfoY-strep genes from different Vibrio species were constructed in the following way: The genes were amplified with Strep-tagII-encoding primers using gDNA of the respective parental Vibrio strain as a template. The restriction-enzyme-digested PCR product was subsequently cloned into plasmid pBAD/MycHisA (Supporting Information Table S1). The resulting plasmids served as templates for fragments containing araC, the arabinose-inducible promoter Pbad and the respective tfoX or tfoY genes, which were subcloned into the mini-Tn7-containing delivery plasmids (Supporting Information Table S1). The plasmid pGP704-Tn-CmR is a derivative of pGP704-mTn7-minus-SacI (Nielsen et al., 2006), where the aacC1 gene within the mini-Tn7 was replaced by the cat gene. To this end, the PCR-amplified cat gene (including its promoter) of pBR-FRT-Cat-FRT2 (Metzger et al., 2016) was cloned into pGP704-mTn7-minus-SacI, which was digested using restriction enzymes SbfI and EcoRV. For the insertion of this mini-Tn7 transposon into the Vibrio spp. chromosomes, a tri-parental mating strategy was employed (Bao et al., 1991). The donor plasmids are indicated in Supporting Information Table S1.

Natural transformation assay

Natural transformation assays in liquid were performed with minor modifications to the previous protocol (Lo Scrudato and Blokesch, 2012). This assay is chitin-independent and uses strains carrying an arabinose-inducible copy of tfoX-strep or tfoY-strep (both from V. cholerae, V. alginolyticus, V. parahaemolyticus or V. fischeri) on the chromosome. V. cholerae strains were pre-grown overnight in LB medium and further grown in the presence of the arabinose inducer up to an optical density at 600 nm (OD600) of 1.0. At this point, 0.5 ml of the cultures were supplemented with 1 μg of genomic DNA. For all natural transformation assays performed in V. cholerae, the genomic DNA of strain A1552-lacZ-Kan (Marvig and Blokesch, 2010) served as transforming material. Cells were further incubated under shaking conditions at 30°C for 4 h. Serial dilutions were spotted on LB to count the total number of cells and on kanamycin-containing LB agar plates to select the transformants. Transformation frequencies were calculated as the number of colony-forming units (CFUs) of the transformants divided by the total number of CFUs. Averages of at least three biologically independent experiments are provided. For statistical analyses, the data were log-transformed (Keene, 1995) and significant differences were determined by the two-tailed Student’s t-test. When no transformants were recovered, the value was set to the detection limit to allow for statistical analysis.

Gene expression analysis by quantitative reverse transcription PCR

Quantitative reverse transcription PCR (qRT-PCR)-based transcript scoring in V. cholerae, V. alginolyticus, V. parahaemolyticus and V. fischeri was performed following a previously established protocol (Lo Scrudato and
Blokesch, 2012). Strains with and without an arabinose-inducible copy of specified genes (tfoX-strep or tfoY-strep) were grown for 6 h in 2.5 ml LB or LBS supplemented with 0.2% arabinose. Cultures (2 ml) were processed for RNA isolation and subsequent cDNA synthesis. Relative gene expression values were normalized against the gyrA transcript levels. Fold changes were determined using the relative expression values of the induced strains divided by the values of the parental WT strain (as specified for each inducible construct in the figures). Averages of at least three biologically independent experiments (± SD) are provided with statistical analyses (two-tailed Student’s t-test) being done on the log-transformed data (Keene, 1995).

Interbacterial killing assay using E. coli as prey

The E. coli killing assay was performed following a previously established protocol with minor adaptions (Borgeaud et al., 2015). The E. coli prey cells and the given predator cells were mixed at a ratio of 1:10 and spotted onto membrane filters on pre-warmed LB and/or LBS agar plates (± 0.2% ara). After 4 h of incubation at 28°C (predator: V. fischeri) or 30°C and 37°C (as indicated for the predators V. cholerae, V. alginolyticus and V. parahaemolyticus), bacteria were resuspended and serial dilutions were spotted onto antibiotic-containing LB agar plates to enumerate the CFUs (shown as CFU/ml). Arabinose-uptake-deficient E. coli TOP10 and its derivative TOP10-TnKan served as prey. Significant differences were determined by a two-tailed Student’s t-test on log-transformed data of at least three biological replicates. If no prey cells were recovered, the value was set to the detection limit to allow calculation of the average of the three independent biological experiments and statistical analysis.

Motility assay

The motility of the Vibrio species was assessed by spotting 2 μl of the respective overnight culture onto freshly prepared LB and/or LBS motility agar plates (containing 0.3% agar) with or without 0.2% arabinose. Following the incubation at 30°C for 6 h for V. cholerae, V. parahaemolyticus and V. alginolyticus, or at room temperature for 7 h for V. fischeri, the diameters of the bacterial swimming were determined. The motility induction was calculated by dividing the swimming diameter of induced versus uninduced strains. The averages of at least three independent experiments (± SD) are provided. For statistical analyses, a two-tailed Student’s t-test was performed.

SDS-PAGE and western blotting

Cell lysates were prepared as described previously (Metzger et al., 2016). In brief, after cultivation with or without arabinose for 3 or 6 h, bacterial cell pellets were resuspended in Laemmli buffer, adjusting for the total number of bacteria according to the OD_{600} values. Proteins were separated by sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis and western blotted as described (Lo Scudatto and Blokesch, 2012). Primary antibodies against Hcp (Eurogentec) (Metzger et al., 2016), GFP (Roche, Switzerland) and mCherry (BioVision, USA distributed via LubioScience, Switzerland) were used at a 1:5000 dilutions, and E. coli Sigma70 (BioLegend, USA distributed via Brunschwig, Switzerland) was used at a 1:10 000 dilution. Goat anti-rabbit horseradish peroxidase (HRP) and goat anti-mouse HRP (both diluted 1:20 000; Sigma-Aldrich, Switzerland) served as secondary antibodies. LumiLightPLUS western blotting substrate (Roche, Switzerland) was used as an HRP substrate and the signals were detected using a ChemiDoc XRS+ station (BioRad).

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Author contributions

Conception, design and analysis: L.C.M, N.M., and M.B.; performed research: L.C.M, N.M., C.S., E.J.C. and M.B.; wrote the manuscript: L.C.M. and M.B. with input from N.M.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s website:

Table S1. Bacterial strains and plasmids used in this study.

Fig. S1. TfoX- and TfoY-induced T6SS production is conserved in pandemic V. cholerae strains. Interspecies killing assay between diverse V. cholerae strains and E. coli as prey. The V. cholerae O1 El Tor strains tested are as follows: A1552, N16961rep (with repaired frameshift mutation in hapR), C6709, E7946 and P27459 as indicated below the graph. Co-culturing with E. coli occurred on LB agar plates supplemented with arabinose to induce tfoX or tfoY where indicated. The parental strains without inducible copies of the regulatory genes served as a control. Prey recovery is indicated as CFU/ml on the Y-axis. Bar plots represent the average of three independent biological replicates (± SD). Statistical significance is indicated (*p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001).

Fig. S2. TfoY production is translationally but not transcriptionally controlled by c-di-GMP in V. cholerae. V. cholerae reporter strains carrying a gene encoding for a translational fusion between TfoY and mCherry (tfoY-mCherry) with or without a downstream transcriptional reporter gene (gfp) at the gene’s native chromosomal locus were genetically manipulated to insert inducible copies vdcA or cdpA into their genome (inside a mini-Tn7 transposon). Cells were then grown under inducible conditions to increase or decrease intracellular c-di-GMP concentrations, as shown by the arrows above the images. Detection of TfoY-mCherry and GFP occurred through western blotting. The reporter strain lacking the transposon as well as the parental WT served as controls. Detection of σ70 served as a loading control.