Natural competence for transformation is a developmental program that allows certain bacteria to take up free extracellular DNA from the environment and integrate this DNA into their genome. Thereby, natural transformation acts as mode of horizontal gene transfer and impacts bacterial evolution. The number of genes induced upon competence induction varies significantly between organisms. However, all of the naturally competent bacteria possess competence genes that encode so-called DNA-uptake machineries. Some components of these multi-protein complexes resemble subunits of type IV pili and type II secretion systems. However, knowledge on the mechanistic aspects of such DNA-uptake complexes is still very limited. Here, we discuss some new findings regarding the DNA-uptake machinery of the naturally transformable human pathogen *Vibrio cholerae*.

**Cellular Localization of the DNA-Uptake Machinery within Naturally Competent *V. cholerae* Cells**

Horizontal gene transfer (HGT) is a major driving force of bacterial evolution. The rapid exchange of genetic information mediated by HGT enables bacteria to adapt to new environmental niches, to spread harmful traits such as antibiotic resistance cassettes or pathogenicity islands and to maintain genome integrity. HGT by means of conjugation and transduction relies on cell-cell contacts and/or mobile genetic elements, whereas natural transformation consists of the uptake of naked DNA from the environment. The ability to acquire exogenous DNA is called natural competence, a physiological state that is of transient nature for most bacterial species. The development of natural competence is often a highly regulated process, which frequently requires environmental cues such as nutrient availability or species-specific competence pheromones. Initiation of natural competence results from the expression of so-called competence genes, which in part encode specialized proteins that constitute the so-called DNA-uptake machinery. Contrary to the different regulatory networks controlling the onset of natural competence, the components of the DNA-uptake machinery are often conserved in Gram-positive and Gram-negative bacteria. Moreover, many of these proteins share homology with type IV pili (Tfp) and type II secretion systems (T2SS). Thus, it is tempting to speculate that DNA uptake by naturally competent bacteria occurs via nearly universal machineries. However, DNA-uptake complexes, especially their composition and cellular localization, have so far been poorly characterized. Moreover, most of the existing information is based on data that were acquired from the naturally competent Gram-positive bacterium *Bacillus subtilis* and, more recently,
also from *Streptococcus pneumoniae*.\(^{15,16}\) However, the situation is different and less well studied for Gram-negative bacteria, where external DNA must be first translocated across the outer membrane (and the periplasmic space).

In our recent study, we aimed to gain insight into the process of DNA uptake and to visualize the DNA-uptake machinery in the human pathogen *Vibrio cholerae* using a cell-biology based approach.\(^{17}\) In this Gram-negative bacterium, natural competence is induced during growth on chitinous surfaces,\(^{7,18}\) which the bacterium encounters in its natural reservoir.\(^{19}\) Substantial information concerning the regulatory circuit of natural competence and transformation of this organism has been gathered in less than a decade (first reviewed by Seitz and Blokesch\(^ {7}\) and later also by Sun et al.\(^ {20}\)). However, the DNA-uptake machinery of *V. cholerae* has never been investigated and has remained a mystery until recently.\(^ {17,21,22}\)

As a first step, Seitz and Blokesch identified the minimum competence regulon of *V. cholerae* based on previously unpublished and published expression data (\(^ {18,23}\) and Blokesch, unpublished), locus organization, homology to other naturally competent bacteria and conservation in other naturally transformable *Vibrio* species.\(^ {17}\) A minimal (most likely still incomplete) competence set of 19 genes was identified with the majority of genes encoding proteins with homology to biogenesis or structural components of Tfp.

To verify the importance of these candidate genes with respect to DNA uptake and natural transformation, each gene was deleted from the parental *V. cholerae* O1 El Tor strain, and the mutant was assessed for its transformability. Notably, pilD was excluded in this study as the deletion mutant displayed a strong growth defect (both in liquid culture and on solidified agar plates). Furthermore, the gene coding for the single-strand binding protein (ssb) could not be deleted.\(^ {17}\) The latter finding is in agreement with recent studies that report the essentiality of ssb in *V. cholerae*.\(^ {24-26}\) All of the other mutants were impaired for transformation even though low numbers of transformants were consistently obtained for strains lacking components of the Tfp portion of the DNA-uptake machinery. Such residual transformability was never observed for those mutants devoid of the competence genes *comEA*, *comEC*, and *comF* as well as the *recA* mutant, which served as a control in this assay.\(^ {17}\) The encoded competence proteins were further categorized (using a recently developed whole-cell duplex PCR DNA-uptake assay\(^ {27}\)) as required for transport across the outer membrane or inner membrane. All of the Tfp-related proteins and ComEA were required for DNA-uptake across the outer membrane, whereas ComEC, ComF, and RecA were necessary only for inner membrane DNA translocation or recombination.\(^ {17}\) Moreover, the results demonstrated that all of the steps involving Tfp-related components or ComEA function upstream of the inner-membrane channel ComEC, whereas the cytoplasmic protein RecA was confirmed as acting downstream of ComEC (similar to DprA; Seitz and Blokesch, data not shown). Finally, as the *comEC* and *comF* mutant of *V. cholerae* consistently displayed the same phenotype with respect to the absence of transformability and the accumulation of transforming DNA within the periplasmic space, it was hypothesized that the two proteins ComEC and ComF work in concert in mediating the translocation of the incoming DNA across the inner membrane (Fig. 1).\(^ {17}\)

As a second step, Seitz and Blokesch imaged a variety of competence-related proteins (Fig. 2).\(^ {17}\) First, the authors used immunofluorescence labeling against an affinity tag that was genetically added to the major pilin subunit PilA. Using this approach, the competence-induced Tfp became evident. Notably, this structure extended well beyond the outer membrane, thereby contradicting the previously hypothesized pseudopilus model for *V. cholerae*.\(^ {18}\) For the majority of piliated cells (95%) only one pilus structure was detectable per cell (Fig. 2). In the absence of each of the Tfp-related components encoded by the minimum competence gene set, the cells appeared non-piliated. However, a pilus structure was still observable in the mutants lacking either a Tfp-unrelated competence gene (*comEA*, *comEC*, *comF*, and *recA*; all non-transformable) or the potential retraction ATPase (PilT).\(^ {17}\) Interestingly, the latter mutant population contained a significantly higher fraction of cells with two or more Tfps.

Next, the authors sought to also localize other competence proteins that were produced as translational fusions (with fluorescent proteins) and replaced the native proteins. All of the competence-induced cells displayed several distinct foci for the secretin PilQ as well as for the putative traffic ATPase PilB (Fig. 2), whereas no localization pattern was obvious for the retraction ATPase PilT.\(^ {17}\)

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**Figure 1.** The ComEC and ComF proteins might drive DNA import into the cytoplasm. Based on the similar phenotypes of both the *comEC* and *comF* mutants, it was suggested that ComF works in concert with ComEC in the translocation of the DNA across the inner membrane.
However, how can such a discrepancy between a single pilus and several outer membrane secretins (PilQ) and elongation ATPases (PilB) be explained? To address this question, Seitz and Blokesch demonstrated that the Tfp consistently co-localized with one of the PilQ foci. Moreover, the PilB ATPase foci were dynamic within the cell (which was not the case for a PilB variant with a mutated Walker B motif) and sometimes aggregated in close proximity of the PilQ protein complexes. The authors therefore hypothesized that each cell might contain several pre-assembled Tfp complexes that might solely lack the major pilin subunit and that pilus elongation – one at a time – occurs upon stimulation of pilin polymerization through PilB (Fig. 2). Such a model would suggest that a higher percentage of cells contain more than one pilus during PilB overproduction, which was exactly what the authors described.17

**Putative Roles of other Competence Tfp-Associated Proteins**

Another important finding of this study was that a gene cluster of five genes (e.g., VC0857 to VC0861) encoding hypothetical proteins or putative type IV pilins contributed significantly to efficient transformation.17 Interestingly, a minor pilin (ComP) was recently shown to directly interact with the species-specific DNA of Neisseria28 thereby answering one of the many open questions in the field of natural transformation, namely “which, if any, protein acts as a receptor for transforming DNA”29 in Neisseria. However, the comP mutants of N. gonorrhoeae, although impaired for natural transformation, are properly piliated and exert a normal Tfp function.30 We recently showed that in contrast to Neisseria and Haemophilus influenzae, V. cholerae does not differentiate between species-specific and species-non-specific DNA at the level of the DNA-uptake process,27 thus excluding a role of the VC0857-0861 operon-encoded proteins in species-specific DNA recognition. Notably, no pili were observed in any of the five mutant strains, consistent with the absence of transforming DNA in the periplasm.17 A putative function of these proteins could therefore be to activate the Tfp assembly machinery, which could potentially work in a similar manner as recently demonstrated for three
minor pseudopilins involved in T2SS assembly in *Klebsiella oxytoca*.

Another competence gene that was part of this study but not further investigated was VC1612. In previous studies, the VC1612 gene was slightly upregulated when *V. cholerae* was grown on chitin surfaces and was inducible by the transformation regulatory protein TfoX. The VC1612 gene product was initially annotated as a putative fimbrial biogenesis and twitching motility protein, putative. Furthermore, BLAST comparisons with PilF of *P. aeruginosa* indicate that the VC1612 protein of *V. cholerae* (suggested annotation: PilF) might act as a pilotin, thereby assisting the PilQ secretin in its assembly.

### The Conserved Operon Structure of *pilMNOPQ*

While comparing the competence/Tfp-related genes of different organisms, it became obvious that there were many variations with respect to the organization of the genes (e.g., whether they clustered together in operons or not). For example, whereas the major pilin-encoding gene *pilA* and the gene coding for the elongation ATPase PilB exist in a single operon in *V. cholerae* (supplementary table of ref. 17), this is not the case for many other Tfp-containing organisms (for an example, see the genetic map of the Tfp genes of *Mycococcus xanthus*). Interestingly, the *pilMNOPQ* operon (Fig. 4) is conserved in many proteobacteria, including for example *P. aeruginosa* strain PAO1, *Pseudomonas stutzeri* A1501, *Escherichia coli* K12, *Legionella pneumophila* strain Paris, *Legionella longbeachae* strain NSW150, and *M. xanthus* (according to the GenoList database; see also review).

What is so special about these five genes that warrants such a conserved operon structure? Indeed, many studies have recently addressed potential protein-protein interactions between these five proteins, and all of these studies suggested that the encoded proteins form a complex that connects the inner and outer membrane (including the outer membrane secretin PilQ). Interestingly, Friedrich et al. recently demonstrated that this protein complex is assembled in a sequential outside-in pathway, which therefore occurs opposite to the gene succession (PilQ-P-O-N-M). It is tempting to speculate that the genetic organization aims to avoid the toxic effects of the gene products that are encoded toward the end of the operon. Thus, the conserved gene sequence would ensure that a given stoichiometry is preserved and that the assembly is only initiated after all of the proteins are synthesized (e.g., full-length mRNA is available in the cell). Notably, the enhanced expression of PilQ and *pilP* in trans results in severe toxicity in both *V. cholerae* and *E. coli*.

In summary, we conclude that we are only beginning to understand the basic components of the DNA-uptake machinery of *V. cholerae* and its functionality.
Moreover, the composition and mechanistic aspects of type IV pili and the DNA-uptake machineries are generally not well understood, even though many hypothetical models of these multiprotein complexes exist. Notably, many of those models have not changed dramatically over the last decade (compare, for example, refs. 47 and 48) but by now are much better supported by experimental data.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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Figure 4. The conserved pilMNOQ operon encoding components of the Tfp-part of the DNA-uptake machinery. For details, see text.
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