

Molecular and genetic characterization of the TonB2-cluster TtpC protein in pathogenic vibrios

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Abstract TtpC is a fourth required protein in the TonB2 energy transduction system in *Vibrio anguillarum*. TtpC is necessary for iron transport mediated by the TonB2 system and is highly conserved in all pathogenic vibrio species studied to date as well as several marine organisms. We show here that the TtpC proteins from selected pathogenic vibrio species can function with the TonB2 system of *V. anguillarum* to allow iron transport mediated by a chimeric TonB2 system where the native ExbB2, ExbD2 and TonB2 function with an episomally expressed TtpC *in trans* from a different species. The discovery that inter-species complementation occurs can be used to identify the functional regions of the TtpC proteins and will lead to an investigation of the mechanism of interaction between the TtpC protein and other members of the TonB2 system.

Keywords TtpC · Vibrio · Iron transport

Abbreviations

2,3-DHBA 2,3-Dihydroxybenzoic acid
IPTG Isopropyl-beta-D-thiogalactopyranoside

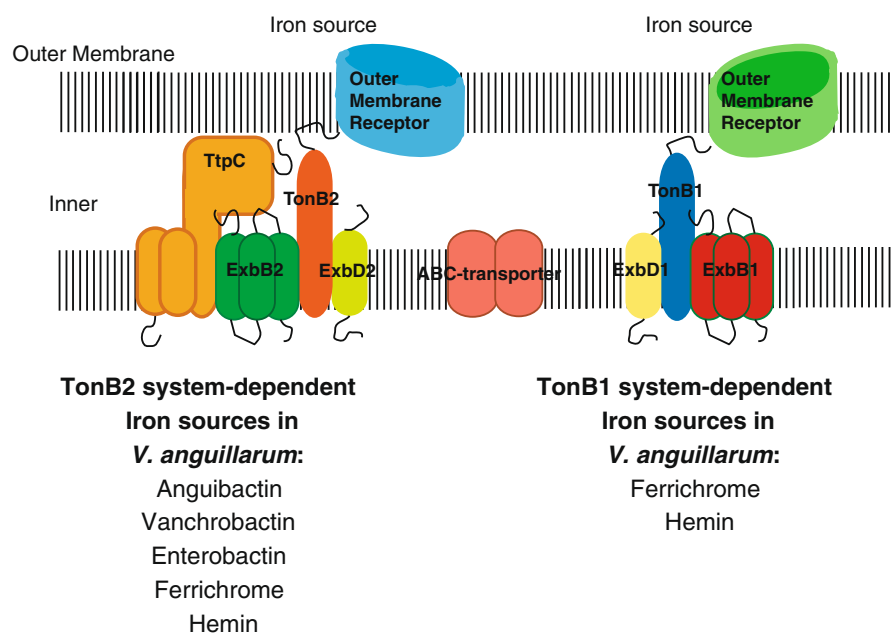
Introduction

Pathogenic bacteria possess virulence factors that allow them to invade a vertebrate host and establish colonization of the host while evading the innate and adaptive immune systems. One of these virulence factors is the ability to compete with the host cells for iron. Iron is an essential element for nearly all species and vertebrate hosts have developed mechanisms to sequester free iron in efforts to deter the growth of pathogenic bacteria. In biological fluids, iron is always bound in a complex with iron-binding proteins such as transferrin, lactoferrin or as part of the oxygen-carrying heme group in red blood cells. Thus, bacterial pathogens scavenging for iron must also possess iron-binding proteins and mechanisms to internalize bound iron in order to establish an infection. Some species express outer-membrane receptors specific for iron-binding host proteins like transferrin or heme. Alternatively, bacteria can synthesize siderophores that are used to scavenge iron from eukaryotic iron-binding proteins. Siderophores are low molecular weight compounds that have an extremely high affinity for iron, in some cases more so than host iron-binding proteins, and are secreted by the bacterium to scavenge iron from the surrounding environment. In Gram-negative species, the iron-siderophore complexes are bound on the surface of the outer membrane by receptor proteins. Most bacterial species express several outer-membrane receptors, each one specific for a different ferric iron-siderophore complex.

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Depending on the outer membrane transport protein and inner membrane permeases being used, the ferric iron or the iron-siderophore complex is internalized into the periplasmic space and then transported through the inner membrane to the cytosol where it can be used by the cell as a co-factor in oxidation–reduction reactions and as a transcriptional co-repressor with the master-regulator Fur (Ernst et al. 1978). The energy required to power all of these steps of iron uptake is derived from the proton motive force in the inner membrane of the Gram-negative cell. This energy is harvested and transduced to the iron-siderophore complex-binding proteins in the outer membrane via a complex consisting of a TonB-family protein, the transducer, and related energy harvesting proteins, ExbB and ExbD (Brinkman and Larsen 2007). ExbB is necessary to stabilize TonB (Fischer et al. 1989; Skare and Postle 1991; Postle and Kadner 2003). The TonB protein interacts with a “TonB box” region on the periplasmic domain of the outer membrane receptor (Crosa et al. 2004). Much of the elucidation of energy transduction between the cellular membranes of Gram-negative bacteria has been determined in *Escherichia coli*. *E. coli* has one classical TonB system whereas the pathogenic vibrio species possess at least two, and in some species, three TonB systems that appear to be specific for the transport of various iron sources (Fig. 1) (Stork et al. 2004).

Fig. 1 Two TonB systems in *V. anguillarum* allow the bacterium to utilize different iron sources



Requirement of a fourth protein

The presence of TonB along with the accessory proteins ExbB and ExbD is conserved across many Gram-negative species. In *V. anguillarum* however, a fourth protein was discovered to be necessary for ferric anguibactin transport mediated by the TonB2 system (Stork et al. 2007). This protein, TtpC is encoded in the *tonB2* gene cluster of *V. anguillarum*.

Using sucrose-density gradients, it was shown that TtpC is located in the inner membrane. Through the use of bioassays to test iron-source utilization, it was shown that TtpC is necessary for ferric-anguibactin transport mediated by the TonB2 system in the fish pathogen *V. anguillarum* (Table 1) (Stork et al. 2007). In addition, the TtpC homologue in the human pathogen *V. cholerae* is essential for enterobactin transport mediated by the *V. cholerae* TonB2 system. Although *V. anguillarum* has two TonB energy transduction systems, the TtpC protein is specific for the TonB2 system and not necessary for iron transport mediated by the TonB1 system in *V. anguillarum* (Table 1).

Homology of TtpC proteins in several pathogenic vibrio species

Based on the data presented in Table 1 showing that TtpC is necessary in the TonB2 system of both

Table 1 TtpC is necessary for iron transport mediated by the TonB2 system in *V. anguillarum* and *V. cholerae* (Stork et al. 2007)

Strains	Iron sources ^c				
	Ferric ammonium citrate	Anguibactin	Enterobactin	Ferrichrome	Heme
<i>V. anguillarum</i>					
775 Wild Type	+	+	+	+	+
775 <i>tonB1</i> [−] , <i>ttpC</i> [−]	+	−	−	−	−
775 <i>tonB1</i> [−] , <i>ttpC</i> [−] / <i>pexbB2</i> , <i>exbD2</i> , <i>tonB2</i> _(va) ^d	+	−	−	−	−
775 <i>tonB1</i> [−] , <i>ttpC</i> [−] / <i>pttpC</i> , <i>exbB2</i> , <i>exbD2</i> , <i>tonB2</i> _(va)	+	+	+	+	+
775 <i>ttpC</i> [−]	+	−	−	+	+
<i>V. cholerae</i>					
CA401 Wild Type	+	ND ^b	+	+	+
CA401 <i>exbB2</i> [−] / <i>i</i>	+	ND	−	+	+
CA401 <i>exbB2</i> [−] / <i>pexbB2</i> , <i>exbD2</i> , <i>tonB2</i> _(va)	+	ND	−	+	+
CA401 <i>exbB2</i> [−] / <i>pttpC</i> , <i>exbB2</i> , <i>exbD2</i> , <i>tonB2</i> _(va)	+	ND	+	+	+
CA401 <i>ttpC</i> [−]	+	ND	−	+	+

^a + or − indicates growth or lack of growth around the specified iron source

^b ND, Not determined

^c 5 µl of each iron source was spotted on the surface of the plates in the following concentrations; Ferric ammonium citrate: 500 µg/ml; enterobactin: 1.0 mg/ml; anguibactin: 1.0 mg/ml; ferrichrome: 1.0 mg/ml; hemin: 20 µg/ml

^d Mutations in chromosomally encoded genes were complemented by genes expressed from the low copy-number expression vector pACYC177

V. anguillarum and *V. cholerae* and that the TonB2 system from *V. anguillarum* can support iron transport of TonB2-specific iron sources in *V. cholerae*, we asked the question: How similar are the TtpC proteins to one another?

To compare the sequences we used the T-coffee server to perform a multiple sequence alignment of the amino acid sequences of the TtpC protein from five pathogenic vibrio species. The Clustal W formatted alignment and the score are shown in Fig. 2. The multiple sequence alignment produced an overall score of 66 (Poirot et al. 2003). Individual alignments of the *V. anguillarum* TtpC sequence to the TtpC sequence of other pathogenic vibrios by BLASTp reveal amino acid sequence similarity between 73 and 80% (Altschul et al. 1990).

Interspecies complementation

After determining the very high similarity between the amino acid sequence of the TtpC proteins in the pathogenic vibrio species, we asked: Are the TtpC proteins from *V. cholerae* and *V. vulnificus* similar enough to the TtpC protein from *V. anguillarum* that

they would be able to complement a chromosomal deletion in the *V. anguillarum* *ttpC*? We observed that *V. cholerae* *ttpC* can complement a Δ *ttpC* mutation in *V. anguillarum* for the uptake of ferric-anguibactin, but *V. vulnificus* *ttpC* cannot. We also observed that the *V. anguillarum* *ttpC* was the only complementing gene that restored growth on all tested iron sources (Table 2). The lack of growth of the *V. anguillarum* Δ *ttpC*, Δ *angA*/*pttpC*_{vc} strain around enterobactin as an iron source indicates that while the TtpC proteins from the different species are highly similar, slight differences may impede apparent involvement with certain outer membrane receptors in *V. anguillarum*.

Requirement of TtpC dependent on TonB2

We observed that TtpC is necessary for growth of *V. anguillarum* on iron sources that utilize only the TonB2 system for transport through the outer membrane. Next, we set out to determine if TtpC was necessary for transport of iron sources that utilize both TonB systems in *V. anguillarum*. We performed bioassays using ferrichrome as an iron source capable

Alginolyticus	--mmkwslswalistaalmpytf-----ssdallqkaqgenrqqqshnvaresgfkqt
Parahaemolyticus	--mikkwslswalistaamphttf-----asdtllqkaqgenrqqqthnasresgfkkt
Anguillarum	--mvfslslcl-----tllafnsf-----astelsvnakvenkqqqqhniereagfkqt
Cholerae	mtlrktllalalslpfafvsvan-----asnslvqgaqtqekaqqqhnqreagfvqt
Vulnificus	---mkgfktiaaallaslfmvssahaepkslnellknvksesiveskenkaregeflad
	:. :. . : * :. * . . * ** *
Alginolyticus	eqdlgaiknklvaeraalqaeadslsvtfgeneaqlaaleklrletgslgelfgvvrqn
Parahaemolyticus	egelgaiknklvaeraalqaeadtllsvtfseenaqlaaleklrletgslgelfgvvrqn
Anguillarum	eqqlrrakaeaqrnrdlqaqadtlstqfsnennrlarleeqlrletgslgelfgvvrqn
Cholerae	aqelgaakaellaernrlqkeadqlssqfsndentlarleetrletgslgemfgvvrqn
Vulnificus	rnkqaellkqakaqлкаetalgeqlkatfddndkkltelteqlrqrsgtlgemfgvvrqy
	:. : * : . : * . * : : * : * * . : * : * : * : * : *
Alginolyticus	akeleselkssvtgvdanayqkdidaivaakslptltqlqamwrsmeeqikasgemanvs
Parahaemolyticus	akeleselkhavtgvdhsyqkdieaivaakslptlqlqamwrsmeeqvtasgelakvs
Anguillarum	aa-lsselessitavdrhqtysrvgeivaakslpsmpqlgtlwhsyvgeiqiasgelasrvq
Cholerae	akelqselqsvtgvpebrahqqsidvvaaktlpsmaqlrglwqamseeirasgqvktte
Vulnificus	agefkgflfnasqnavqfpqrdalltklaeskelpsteelesfwhvtlqqiivsgtsttp
	* :. . : . : * : : : * * : * : * : : : * : .
Alginolyticus	ftllngegreqtvsgrvlgsmaillddtgyvkwngqrgdavnlylrqpesgpt--an----ti
Parahaemolyticus	ftlldgegkeqtvnvrqlgamaillddtgyvkwngqrgdavnlylrqpengpt--as----af
Anguillarum	iaifngdqgtqnvaaayrigtf-1vtdeggyvkwngkdqaataylkkpddngpi--fs---gl
Cholerae	iqwlngqgetqtpalrqlsglgliseqgyvkwdnarqqalsyqqllpsdfpt--fshirrtl
Vulnificus	atvvyyeggeavrdvtlvgefnaiadgkyvtyvptgtkfeelsrgpranirrhvagfvs
	: * : * : * : : : * : : : *
Alginolyticus	ssgdidalvidpsrgilleqlansptladrlnaggvvgkiilglilaigllialvrgaslm
Parahaemolyticus	tngeidslvidpsrgilleqlansptkladrnaggvvgkiilvllaigllialvrgslf
Anguillarum	ntgeieslvdpdpsrgilleqlaltptlaerlqmggavvniilvllglliafyrgrfalv
Cholerae	vdgdvvtmvdpsrgvllleqlaltptfsqrlqaggvignvilvllgvglialyrgaila
Vulnificus	tsssyepflfidpsrgvllslvgsptvkeridggvgvyilamgalgalialfclyrllr
	. . . : * * : * . * : * : * : * : * : * : *
Alginolyticus	isrqkimkqlktpaqpgnnplgrvlavvyqdkdhksvealelrllaeavvdeqthlekglsm
Parahaemolyticus	iarhkiakqlknptkaednplgrvlsvyqndkhksvealelrllaeavvdeqthlekglsm
Anguillarum	mtbhkikaqlrldvqkagnplgrvlsvynkeqrsvealelrllaeavvdeqthlekglsm
Cholerae	tlrqkikaqlknpeqpnnplgrilavynkeqqrsvalelrllaeavvdeqnhletglsm
Vulnificus	viggkmrkqaksdavignplgeviqayqdhkdnledleakldeilrnapiakierfivs
	* : * : . * * : * : * : * : * : * : *
Alginolyticus	lkllaalaplmlglgtvtgmietfqvitqfgngdpkvmaggismalvtvtlglvlsamp11
Parahaemolyticus	lkllaalaplmlglgtvtgmietfqvitqfgngdpkvmaggismalvtvtlglvlsamp11
Anguillarum	lkllaalaplmlglgtvtgmietfqvitqfgngdpkvmaggismalvtvtlglvlsai11
Cholerae	lkllaalaplmlglgtvtgmietfqvitqfgngdpkvmaggismalvtvtveglaai11
Vulnificus	iklfasvapillglgtvmgmietfqgaitlfgtdpklmaggisealvttmlglvvaip11
	: * : * : * : * * * * * * * * * * * * * * * * * : * : * * *
Alginolyticus	lahnvllssqaenirsilekqgiglvaeqaerdmppnnkshsntlaenaa
Parahaemolyticus	lahnvllssqaesirnilekqgiglvaeqaerdmppnnkngrnntiaenaa
Anguillarum	lahnillssqaenirsilekqgiglvaaqaedeqcas-----mrnaa
Cholerae	lahnilsaqaearnilekqgiglvaaqaerdc--gaavtqsrveqaa
Vulnificus	ffytltvhskgrrrlvtleeqsagfiaryqekhltae-----r
	: : : : : * * : * * * *

Table 2 *V. cholerae* ttpC complements Δ ttpC mutation in *V. anguillarum* for growth around 2,3-DHBA

Strains ^c	Iron sources ^b				
	Ferric ammonium citrate	Enterobactin	2,3-DHBA	Vanchrobacin	Hemin
<i>V. anguillarum</i>					
775/pMMB208	+ ^a	+	+	+	+
775 Δ ttpC, Δ angA	+	–	–	–	+
775 Δ ttpC, Δ angA/pttpCva	+	+	+	+	+
775 Δ ttpC, Δ angA/pttpCvc	+	–	+	–	+
775 Δ ttpC, Δ angA/pttpCvv	+	–	–	–	+

^a + or – indicates growth or lack of growth around the specified iron source

^b 1 μ l of each iron source was spotted on the surface of the plates in the following concentrations; ferric ammonium citrate: 10 mg/ml; enterobactin: 1.0 mg/ml; 2,3-DHBA: 3.0 mg/ml; vanchrobacin: 1.0 mg/ml; hemin: 20 μ g/ml

^c Mutations in chromosomally encoded genes were complemented by genes expressed from the low copy-number expression vector pMMB208. Expression was induced with 1 mM IPTG

Table 3 TtpC is necessary for usage of iron sources when transport is mediated by the TonB2 system

Strains ^c	Iron sources ^b			
	TonB1- and TonB2-mediated transport		TonB2-mediated transport	
	Ferric ammonium citrate	Ferrichrome	Enterobactin	2,3-DHBA
<i>V. anguillarum</i>				
775/pMMB208	+ ^a	+	+	+
775 Δ tonB1, Δ ttpC, Δ angA	+	–	–	–
775 Δ tonB1, Δ ttpC, Δ angA/pttpCva	+	+	+	+
775 Δ tonB1, Δ ttpC, Δ angA/pttpCvc	+	+	–	+
775 Δ tonB1, Δ ttpC, Δ angA/pttpCvv	+	–	–	–

^a + or – indicates growth or lack of growth around the specified iron source

^b 1 μ l of each iron source was spotted on the surface of the plates in the following concentrations; ferric ammonium citrate: 10 mg/ml; enterobactin: 1.0 mg/ml; 2,3-DHBA: 3.0 mg/ml; ferrichrome: 1.0 mg/ml

^c Mutations in chromosomally encoded genes were complemented by genes expressed from the low copy-number expression vector pMMB208. Expression was induced with 1 mM IPTG

(López et al. 2008). This strain still had a wild type TtpC protein so ttpC was also deleted in the tonB1, tonB2 double mutant and complemented with tonB from *E. coli* on a the low copy-number vector pMMB208 (Morales et al. 1991) to determine if the TonB from *E. coli* needed a functional TtpC protein to energize transport of ferric-anguibactin in *V. anguillarum*. Bioassays with these strains were performed and are presented in Table 4. The TonB protein from *E. coli* does not need a functional TtpC protein to energize the transport of ferric-anguibactin. From this, we may conclude that while the function of TonB1va or TonBec do not require TtpC, the TtpC protein is required for TonB2 function in *V. anguillarum* (Tables 3 and 4).

Lack of anguibactin uptake or lack of anguibactin production?

When 2,3-DHBA is spotted on the surface of the *V. anguillarum* bioassay plates as an iron source, the Δ ttpC, Δ angA mutant strains complemented with *V. anguillarum* and *V. cholerae* ttpCs are able to grow but when 2,3-DHBA is supplied to the Δ ttpC, Δ angA mutant strain complemented with the *V. vulnificus* ttpC, no growth is observed (Table 2). Because of this observation, we asked the question: Is the lack of growth seen in the mutant complemented with *V. vulnificus* ttpC due to a lack of iron transport mediated by the TonB2 system or is it due to that particular strain not being able to produce anguibactin

Table 4 In *V. anguillarum*, the TonB protein from *E. coli* does not require TtpC to mediate transport of the endogenous siderophore anguibactin

Strains ^c	Iron sources ^b		
	Ferric ammonium citrate	Enterobactin	Anguibactin
<i>V. anguillarum</i>			
775/pMMB208	+	+	+
CSL68: $\Delta tonB1, \Delta tonB2$ /pMMB208	+	–	–
CSL64: $\Delta tonB1, \Delta tonB2$ /p $ton8_{E. coli}$	+	–	+
775 $\Delta ttpC$ /pMMB208	+	–	–
775 $\Delta tonB1, \Delta tonB2, \Delta ttpC$ /pMMB208	+	–	–
775 $\Delta tonB1, \Delta tonB2, \Delta ttpC$ /p $tonB_{E. coli}$	+	–	+

^a + or – indicates growth or lack of growth around the specified iron source

^b 1 μ l of each iron source was spotted on the surface of the plates in the following concentrations; ferric ammonium citrate: 10 mg/ml; enterobactin: 1.0 mg/ml; anguibactin: 1.0 mg/ml

^c Mutations in chromosomally encoded genes were complemented by genes expressed from the low copy-number expression vector pMMB208. Expression was induced with 1 mM IPTG

Table 5 Growth around anguibactin-producing *V. anguillarum* strains

Indicator strains	Anguibactin-producing strains ^b					
	775 Wild type	$\Delta ttpC$	$\Delta ttpC, \Delta angA$ /p $tttC_{va}$	$\Delta ttpC, \Delta angA$ /p $tttC_{vc}$	$\Delta ttpC, \Delta angA$ /p $tttC_{vv}$	Ferric ammonium citrate
CC9-8	– ^a	–	–	–	–	+
CC9-16	+	+	–	–	–	+
CC9-8 + DHBA	–	–	–	–	–	+
CC9-16 + DHBA	+	+	+	+	+	+

^a + or – indicates growth or lack of growth around the specified iron source

^b 3 μ l of each anguibactin-producing strain was spotted on the surface the plate

from the 2,3-DHBA? To answer this question, we used the anguibactin-indicator strains *V. anguillarum* CC9-8 (Tolmasky et al. 1988), which cannot produce or transport anguibactin and *V. anguillarum* CC9-16 which cannot produce anguibactin but is proficient in transport. 3 μ l of overnight, IPTG-induced culture of the complemented $\Delta ttpC$, $\Delta angA$ mutants grown in CM9 minimal media were used as the iron sources in the bioassay (Table 5). When 2,3-DHBA was added to the bioassay plate, CC9-16 (Tolmasky et al. 1988), was able to grow around all of the supernatants tested indicating that all strains were producing anguibactin when grown on 2,3-DHBA.

Future directions

The genetic complementations of the $\Delta ttpC$ mutation in *V. anguillarum* show that the *V. vulnificus* TtpC is

unable to complement the $\Delta ttpC$ mutant to restore growth around the TonB2-specific iron sources. These results illuminate a potential mechanism by which TtpC could be working. Previous data from M. Stork shows that protein–protein interactions occur between TtpC and TonB2 in *V. anguillarum* (Stork et al. 2007). Based on these observations, we postulate that the interactions between the TonB2 protein and the *V. vulnificus* TtpC in the *V. anguillarum* $\Delta ttpC$, $\Delta angA$ mutant strain are either not occurring all together or a necessary interaction between the two proteins, or possibly, with the accessory proteins ExbB2 and ExbD2, is unstable and does not allow for energy transduction by the TonB2 system.

Based on the results from Tables 2, 3 and 4, the TtpC protein may be essential for interactions between the TonB2 system and the outer-membrane receptor. This theory is supported by the observation

that the *V. cholerae* TtpC is able to complement growth around anguibactin as an iron source but is unable to complement growth around vanchrobactin or enterobactin as iron sources. We note that these potential interactions could be very specific due to the high similarity between the *V. anguillarum* and *V. cholerae* TtpC protein sequences.

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