Pathogenicity and virulence regulation of *Vibrio cholerae* at the interface of host-gut microbiome interactions

Ansel Hsiao\(^a\) and Jun Zhu\(^b\)

\(^a\)Department of Microbiology & Plant Pathology, University of California Riverside, Riverside, CA, USA; \(^b\)Department of Microbiology, Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA, USA

**ABSTRACT**

The Gram-negative bacterium *Vibrio cholerae* is responsible for the severe diarrheal pandemic disease cholera, representing a major global public health concern. This pathogen transitions from aquatic reservoirs into epidemics in human populations, and has evolved numerous mechanisms to sense this transition in order to appropriately regulate its gene expression for infection. At the intersection of pathogen and host in the gastrointestinal tract lies the community of native gut microbes, the gut microbiome. It is increasingly clear that the diversity of species and biochemical activities within the gut microbiome represents a driver of infection outcome, through their ability to manipulate the signals used by *V. cholerae* to regulate virulence and fitness *in vivo*. A better mechanistic understanding of how commensal microbial action interacts with *V. cholerae* pathogenesis may lead to novel prophylactic and therapeutic interventions for cholera. Here, we review a subset of this burgeoning field of research.

**Introduction**

*Vibrio cholerae* is a Gram-negative bacterium responsible for the severe human diarrheal disease cholera. Cholera is characterized by voluminous watery diarrheas and vomiting, which may rapidly lead to hypovolemic shock, acidosis, and death, with an untreated case-fatality rate as high as 50%. Seven recorded major cholera pandemics have been recorded since 1871, though the disease likely has an ancient association with humans. Although the development of oral rehydration has reduced the treated case fatality rate substantially, cholera continues to impose an enormous global health burden. One contributor to this is the high morbidity of cholera; though patients are likely to survive infection due to rehydration therapy, the debilitating diarrhea characteristic of cholera continues for days even if the pathogen is cleared with antibiotics. Cholera is widely distributed, and represents a threat to public health in many parts of Asia, Africa, and Latin America. Each year sees 1.3–4.0 million cases of cholera, and 21,000–143,000 deaths worldwide, as estimated by the World Health Organization (WHO) [1]. Therefore, infection with *V. cholerae* remains an important health and economic concern [2–5]; the Roadmap to 2030 from the WHO Global Task Force on Cholera Control (GTFCC) envisages a plan for a 90% reduction in deaths from cholera [6].

Although *V. cholerae* is serologically diverse with more than 206 serogroups reported, only O1 and O139 serogroups have been known to cause epidemic cholera [2]. The first six cholera pandemics were caused by classical biotype *V. cholerae*, whereas the El Tor biotype is responsible for the current, seventh pandemic. These two *V. cholerae* biotypes differ considerably; El Tor strains generally cause a milder form of cholera than that caused by classical strains and apparently evolved as better survivors in the aquatic environment [7]. Currently, El Tor strains are predominant everywhere that *V. cholerae* O1 can be found [2].

Between epidemics, *V. cholerae* natively resides in aquatic environments such as freshwater lakes and rivers, where these bacteria interact with various surfaces in the form of biofilms, which form an important survival mechanism [8]. From these aquatic reservoirs, *V. cholerae* spreads to populations of the natural host, humans, through contamination of water and food. Upon human colonization, virulence is due primarily to the production of cholera toxin (CT), which alters host cell signal transduction pathways and leads to cell damage and diarrhea, and the toxin coregulated pilus (TCP), which is critical for colonization of the intestinal...
The profuse watery diarrhea of cholera then disseminates the pathogen back into the environment, where the cycle of fecal contamination can continue, leading to epidemic spread in human populations.

*V. cholerae* is able to respond to environmental signals in the transition from the aquatic environment into the gastrointestinal tract, and regulate virulence genes coordinately to allow for colonization, survival, virulence, and subsequent broad dissemination. The environmental conditions of the gut are tightly bound to the activities of the native microbial community of the gastrointestinal tract, the gut microbiome. Due to the rapidly expanding field of research, this review will focus on just a subset of recent work on how *V. cholerae* adapts to *in vivo* environments by responding to intestinal signals produced and modulated by the host and by commensal microbes.

**Regulatory networks to coordinately activate virulence genes during infection**

A body of work on the ecology of cholera has demonstrated that *V. cholerae* is an autochthonous aquatic organism, occupying brackish water and estuarine environments when not associated with the human host [10,11], underlined by the presence of multiple adaptive strategies for persistence in aquatic reservoirs. Outside of the human host, *V. cholerae* can be found complexed with marine organisms such as crustacean zooplankton [12,13]. *V. cholerae* have been found as biofilms on zooplankton host exoskeleton, as well as in the gut of these marine organisms. Planktonic *V. cholerae* can also respond to low temperatures or nutrient limitation by phenotypic transition to a viable but non-culturable (VBNC) state [14] characterized by reduced cell size and minimal metabolic activity. These environmental persistence strategies have major consequences for the transition into the host gastrointestinal tract. Biofilm-associated *V. cholerae* is more acid-resistant than planktonic cells, which aids in the transition through the low-pH environment of the stomach and ultimately into the distal small intestine, the preferred site of human colonization [15]. There, *V. cholerae* also needs to exit biofilm structures in order to disseminate into the gut mucosa. This process is aided by gut-localized environmental factors such as bile [16].

As they enter the host, *V. cholerae* cells are exposed to a series of changes, such as temperature, osmolarity, oxygen concentration, and exposure of antimicrobial agents (e.g. bile salts). Given the dramatically different environmental conditions between the aquatic reservoir and the host mucosa, it is unsurprising that *V. cholerae* have evolved numerous regulatory mechanisms designed to tailor the production of factors permitting optimal host colonization.

The ability of *V. cholerae* to colonize and cause disease in hosts requires production of a number of virulence factors during infection. The two major virulence determinants of *V. cholerae* are encoded by two separate genetic elements. Cholera toxin, which causes the diarrhea characteristic of cholera, is encoded by *ctxAB* genes on the lysogenic CTXφ bacteriophage [17]. *V. cholerae* also produces toxin-coregulated pilus (TCP), which are required for intestinal colonization both in animal models and in human volunteers [18,19]. TCP is thought to be a polymer of the main structural subunit, TcpA, and serves as the receptor for the CTXφ bacteriophage [17,20]. The genes required for TCP synthesis, including tcpA as well as accessory colonization factor (acf) genes and the genes encoding the virulence transcriptional activators ToxT and TcpP, are located on a 40-kb *Vibrio* pathogenicity island [21]. Coordinate expression of *V. cholerae* virulence genes results from the activity of a cascading system of regulatory factors [22].

The primary direct transcriptional activator of *V. cholerae* virulence genes, including *ctxAB* and tcpA, is ToxT, a member of the AraC/XylS-family of transcriptional regulators [23], members of which often bind effector molecules and/or oligomerize to affect transcriptional regulation at target promoters [24]. Unsaturated fatty acids can directly bind to and inhibit the activity of ToxT [25–27]. To activate transcription, ToxT recognizes and binds to a degenerate 13-bp DNA sequence, the “toxbox” [28]. FadR, the master regulator of fatty acid metabolism, modulates ToxT activity at both transcriptional and posttranslational levels [29], suggesting that the expression of genes involved in fatty acid biosynthesis and virulence are intertwined.

A complex regulatory pathway controls the expression of ToxT (Figure 1). The ToxR protein was identified as the first positive regulator of *V. cholerae* virulence genes; the genes involved in the transcriptional cascade resulting in toxT expression and consequent virulence gene activation is thus often referred to as the “ToxR regulon” [30]. Acting in conjunction with TcpP (see below), ToxR activates the transcription of toxT [31–33]. ToxR directly regulates the transcription of many genes [34], including *ompU* and *ompT*, which encode the major porins of *V. cholerae* [35,36]. ToxR is a bitopic membrane protein containing a cytoplasmic DNA-binding domain, a single transmembrane domain, and a periplasmic domain. ToxR activity requires the presence of another inner membrane...
protein, ToxS; deletion of toxS negatively impacts ToxR transcriptional activity [37], suggesting that ToxS serves as an effector of ToxR function by influencing the stability and/or the dimerization of ToxR [23,38,39]. Recent work has also shown that ToxR-ToxR protein–protein interactions are significantly increased in response to ToxR operators and the co-activator ToxS [40].

To activate the expression of toxT, ToxR acts with a second transcription activator, TcpP, which is also membrane-localized and contains a cytoplasmic winged helix-turn-helix (w-HTH) domain [32]. TcpP, like ToxR, requires the presence of a membrane-bound effector protein, TcpH, which interacts with TcpP [41]. TcpP is degraded by a protease in the absence of TcpH, and during conditions unfavorable for virulence gene activation [42,43]. TcpP binds to the toxT promoter just upstream of the −35 element and is a direct toxT activator [44]. Overexpression of TcpP alone activates toxT expression [45,46], but binding of ToxR to the upstream of TcpP-binding site is required for TcpP-mediated expression of toxT at endogenous expression levels [47,48].

Two activators encoded by unlinked genes, AphA and AphB, regulate transcription of tcpPH. AphA is a dimer with an N-terminal winged-helix DNA binding domain that is structurally similar to those of MarR family transcriptional regulators [49]. AphA cannot activate transcription of tcpPH alone, but requires interaction with the LysR-type regulator AphB that binds downstream of AphA binding site [50]. This interaction is thought to stabilize AphB binding to its recognition site and result in activation of the tcpPH promoter. In addition, AphB enhances the expression of toxR [51]. Expression of aphA is also controlled by a quorum-sensing system [50,52–54], discussed in detail below. This process means that virulence gene expression declines at high cell density and is thought to contribute to the self-limiting nature of V. cholerae infections.

**Gastrointestinal signals modulate virulence**

Pathogens that cause diseases of complex animal hosts require clever strategies for survival and multiplication during the dynamic conditions found during infection. Often pathogens take advantage of host-specific signals to modulate their gene expression in order to adapt the new environments. V. cholerae encounters a variety of unique host-specific signals, including bile, differences in osmolarity, oxygen availability, changes in pH as it travels from the aquatic reservoir to the stomach and into the intestines. Ample research has demonstrated that this bacterium has evolved to rapidly respond to these signals to promote survival and proliferation in the gut, and below we review a subset of these in vivo signals.

**Bile**

Bile is a digestive secretion primarily involved in emulsifying and solubilizing dietary lipids to aid absorption, and is composed of bile acids, cholesterol, phospholipids, and IgA [55]. Bile acids, the predominant

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**Figure 1.**

*V. cholerae* virulence regulatory networks. Major transcriptional regulators and their corresponding signals are shown. In particular, the master regulator ToxT activates virulence genes which products are involved in synthesis of the key virulence determinants TCP and CT. The expression of toxT is regulated by TcpP and ToxR. QS: quorum sensing. ←: activation; ↓: repression.
component of bile, are synthesized in the liver from cholesterol as primary bile acids, often conjugated to amino acids such as taurine and glycine. Bile is stored in the gall bladder, to be secreted into the small intestine with the intake of food, where due to the local pH they are often found as primary bile salts. In the intestines, bile acids/salts mediates digestive processes, and are modified by the action of gut microbes into several secondary bile molecules [56–58]. Up to 95% of bile acids are reabsorbed from the distal ileum, to be passed back to the liver via portal circulation to be re-conjugated to amino acids and re-secreted in a process called enterohepatic circulation [59,60].

Bile can destabilize membranes and disrupt bacterial cellular homeostasis via its detergent-like properties. As a result, the large quantity of bile secreted by liver every day represents a challenge for invading pathogens [61]. V. cholerae is highly resistant to bile through the action of efflux pumps and by outer membrane porins selectively restricting the influx of bile salts [62–65]. The compartment-specific cycling of bile acids means that these molecules can serve as a convenient spatiotemporal cue for gene regulation in microbes adapted to the small intestine. Bile has been shown to promote V. cholerae motility, which is required for efficient colonization [66,67]. Most prominently, V. cholerae is able to use a set of largely primary bile salts (taurocholate, glycocholate) to activate virulence gene expression [68]. This activation is mediated through the transmembrane transcription factor TcpP. Various genetic and biochemical analyses indicate that in the absence of bile salts, one of the two cysteine residues in the periplasmic domain, Cys^{218}, forms an inhibitory intramolecular disulfide bond with the other cysteine residue Cys^{207}. Taurocholate promotes the formation of C^{207}–C^{207} intermolecular disulfide bond formation of TcpP and dimerization, and thus increasing the activity TcpP. Further investigation [69] showed that bile salts inhibit the reductase activity of DsbA, a conserved oxidoreductase in the bacterial periplasm that participates in protein folding by introducing disulfide bonds into proteins [70]; indeed, DsbA induces TcpP dimerization in the presence of primary bile salts such as taurocholate. Calcium has been shown to enhance virulence by promoting bile salt-induced TcpP–TcpP interaction [71]. Bile salts can also prevent ToxR proteolysis and promote ToxR-ToxR protein interaction and ToxRS complex formation [40,72]. One bile salt, taurocholate, promotes V. cholerae dispersal from biofilm structure [16], in addition to activating virulence genes, suggesting that if V. cholerae is ingested as a biofilm to protect against reduced stomach pH, it has coopted the host-derived bile salt signal to detach from the biofilm and go on to activate virulence. Bile is a highly complex mixture and its components have been reported to have different effects on V. cholerae infection; as mentioned above, unsaturated fatty acids that are found in bile bind to ToxT and inhibit its transcriptional activity [25]. Crude bile has been found to decrease cholera toxin production, and fatty acids with bile can repress other aspects of V. cholerae virulence through ToxT [25,73].

**Anaerobiosis**

Given the presumed low oxygen concentration of the gut [74], it is unsurprising that anaerobiosis serves as one of the host environmental factors that modulate virulence factor production [75]. V. cholerae encounters low oxygen concentrations in the upper intestine, and transcriptional examinations of in vivo grown bacteria have confirmed the expression of metabolic genes responsible for anaerobic energy metabolism [76]. Anaerobic respiration of trimethylamine N-oxide (TMAO) enhance cholera toxin production, and TMAO induces more severe symptoms in the infant mouse model [77]. Oxygen availability has been shown to modulate virulence gene expression in a number of gastrointestinal pathogens, such as Shigella [78], Salmonella [79,80], and enterohaemorrhagic E. coli [81]. Under anaerobic conditions, tcpP expression increases, and work has shown that this effect acts through AphB [82]; anaerobiosis enhances oligomerization and activity of AphB [83]. Specifically, one key AphB cysteine residue (Cys^{335}) is oxidized under aerobic conditions. Under low oxygen conditions, AphB Cys^{335} is reduced, which promotes oligomerization and subsequently enhances AphB activity. Intriguingly, during the transition from oxygen-rich to oxygen-poor environments, the rate of reduction of AphB is slow. Another redox-sensing regulator, OhrR, whose cysteine reduction is faster than that of AphB, is needed to jump-start virulence gene expression [84]. Anaerobiosis also promotes ToxR-TcpP interaction, which is important for virulence gene induction [85]. Other global regulatory systems for anaerobic metabolism, such as the ArcA/ArcB two-component system, may also be involved in modulating expression of V. cholerae virulence genes [86], though the exact mechanism of this regulation is unclear.

**Catabolite sensing and nucleotide-derived second messengers**

In recent years, it has become more and more apparent that nucleotide-derived small molecules that accumulate intracellularly are important for bacterial environmental adaptation. These so-called second messengers often have global regulatory effects across bacterial species [87]. In V. cholerae, these small molecules control important physiological functions such as virulence and biofilm...
formation. Among them, the well-studied cyclic adenosine monophosphate (cAMP) is primarily associated with the regulation of carbon utilization. The cAMP regulation is mediated through the cyclic AMP (cAMP) receptor protein (CRP), which binds DNA in response to the intracellular availability of cAMP. CRP-cAMP negatively regulates virulence genes, including cholera toxin genes, TCP pilin gene tcpA, and tcpP [88,89]. Recently, the Camilli group used chromatin immunoprecipitation coupled with DNA sequencing (ChiP-seq) to map the distribution of CRP binding sites across the V. cholerae genome [90] and found that CRP-regulated genes substantially overlaps the ToxR regulon, and that CRP also controls additional virulence factors not regulated by ToxR, such as production of RTX toxin. Another well-studied molecule is guanosine pentaphosphate ((p)pppGpp), which is a primary signal in the bacterial stringent response [91]. In V. cholerae, ((p)pppGpp) signaling pathways regulates antibiotic tolerance [92]. In recent years, the importance of cyclic dinucleotide second messengers has become widely appreciated. Bis (3′,5′)-cyclic dimeric guanosine monophosphate (c-di-GMP) has been found in all major bacterial phyla. C-di-GMP is produced by diguanylate cyclases and degraded by phosphodiesterases [93]. The V. cholerae genome contains 42 genes encoding putative diguanylate cyclases/phosphodiesterases [94]. In V. cholerae, c-di-GMP enhances biofilm matrix production and represses motility as well as virulence gene expression [95–101], suggesting that c-di-GMP may be regulated in response to a variety of environmental signals. Additionally, the most recently identified bacterial cyclic dinucleotide signaling molecule, the hybrid 3′, 3′-cyclic GMP-AMP (cGAMP), is synthesized by the enzyme DncV, whose expression is regulated by the master virulence regulator ToxT [102]. cGAMP binds to the effector called CapV, a phospholipase, and regulates virulence, chemotaxis, and fatty acid metabolic genes [103].

**ROS/RNS**

One of the major stresses in the intestine that V. cholerae must overcome are those generated by exposure to reactive radical species. Reactive compounds, including reactive oxygen species (ROS) and reactive nitrogen species (RNS), are produced in the gut during V. cholerae infection [104–106]. Several proteins have been identified in V. cholerae ROS resistance, including catalases (KatG and KatB), peroxiredoxin (PrxA), organic hydroperoxide resistance protein (OhrA), a redox-regulated chaperone (Hsp33), and a DNA-binding protein from starved cells (DPS) [107–107–110]. ROS resistance in V. cholerae is tightly regulated by a variety of mechanisms. OxyR is required to activate catalase genes and dps, and is modulated by another OxyR homolog, OxyR2 [107,109,111]. Quorum sensing systems [54], PhoB/PhoR two-component systems [112], and the virulence regulator AphB also play important roles in the V. cholerae oxidative stress response [113]. Interestingly, ROS may also enhance the V. cholerae mutation rate in vivo, which results in increased catalase production and increased biofilm formation, leading to colonization advantages in ROS-rich intestines [114]. Adaptive responses in many bacterial pathogens are induced by nitric oxide and RNS. For example, in an inflamed gut, NO is converted to nitrate, which is used by enteric pathogens, such as S. Typhimurium and E. coli, as an anaerobic respiration substrate, leading to pathogen expansion at the expense of the native gut microbiota [115–118]. V. cholerae lacks a nitrite reductase, but nitrate reduction in V. cholerae can occur at alkaline pH during hypoxic growth, whereas in acidic conditions, accumulation of NO2 from NO3 simultaneously limits growth while preserving viability [119]. It has also been shown that a NO-activated transcriptional regulator, NorR, positively regulates hmpA, which encodes a flavohemoglobin that is critical for V. cholerae RNS resistance in vitro and in vivo [120,121].

**Temporal gene expression during V. cholerae infection**

Facultative bacterial pathogens such as V. cholerae that transit from environmental reservoirs to host populations and back are rarely in a static environmental context. Thus, these bacteria must continuously adapt, re-tooling their transcriptional and translational repertoires in order to suit the varied environments they encounter. For example, toxins, host-specific adherence/attachment and invasion factors, loosely classified under the designation “virulence genes,” are activated in an infection-specific manner, whereas the expression of some genes important for environmental survival are down-regulated during infection in order to evade host defense mechanisms (Figure 2).

**In vivo induced genes**

Very often virulence genes are only highly expressed in vivo; in vitro conditions developed to induce virulence gene expression may thus seem artificial. For instance, the laboratory conditions required for activation of TCP and CT in the classical biotype (30°C, pH 6.5, and low osmolarity) are obviously different from the conditions in the small intestine [122]. In the El Tor biotype, one artificial condition (AKI) has been defined that induces expression of CT and TCP
In an elegant study [124], Camilli and colleagues used a recombination-based in vivo expression technology (RIVET) to study in vivo timing of gene induction at the single-cell level. They found that while choler toxin gene is induced monophasically in the small intestine, the expression of tcpA, which encodes the major subunit of TCP, is induced biphasically in two temporally and spatially separable events during V. cholerae infection. Intriguingly, ToxR, TcpP, and ToxT are all required to activate virulence genes in vitro but only ToxT is fully required in vivo. Later RNA-Seq analysis identified that transcripts elevated in infected rabbits and mice relative to laboratory media are regulated not only by previously identified regulators, but also by genes and small RNAs previously not linked to virulence [125], suggesting that virulence activation in vivo is more complex than previously thought.

**In vivo repressed genes**

Pathogenic bacteria must contend with an in vivo environment that is under the protection of host mechanisms capable of rapidly identifying and eliminating foreign microorganisms. Successful pathogens often have mechanisms to survive the recognition by components of the host immune system. Using a differential fluorescence-activation approach, Hsiao et al. reported [126] that among the genes that are repressed by V. cholerae during infection are those encoding for the biogenesis of a Type IV mannose-sensitive hemagglutinin (MSHA), a structure that is produced in vitro and is important for V. cholerae biofilm formation [127], but not during early colonization of the infant mouse model. This is an important process for V. cholerae during infection, as host-secreted IgA nonspecifically binds to V. cholerae cells in an MSHA-dependent manner. Bacteria bound by secretory immunoglobulin in the gut can become entrapped in the mucus layer of the intestine, excluding them from the epithelium and leading to clearance by bulk flow [128–130]. Interestingly, it has been shown that the transcription factor responsible for the negative regulation of msh gene transcription is ToxT, the same transcription factor responsible for direct virulence gene induction [131]. Thus, this pathogen very
efficiently combines its responses to host stimuli in order to simultaneously upregulated factors necessary for colonization (ctx and tcp genes), while using these same factors to transcriptionally repress the biogenesis of anti-colonization factors such as MSHA. Recently, another study used a different approach to identify in vivo-repressed genes in V. cholerae and found that clcA, encoding an H+Cl− transporter, is repressed during infection [132]. While ClcA facilitates survival under low pH (e.g., the stomach), its activity becomes detrimental under the alkaline conditions found in the small intestine; indeed, constitutive expression of clcA reduces colonization fitness.

Late induced genes

Schild and colleagues employed a modified the tnpR RIVET system described above [124,133] to screen for genes only induced late in infection [134]. The library of promoters can be culled at any given point during infection by the oral administration of kanamycin to infected animals, which kills those cells that have already expressed tnpR and resolved the neo-sacB cassette. Interestingly, many of the genes found to have been induced in later stages of infection are involved in bacterial metabolism, indicating that V. cholerae initiates a transcriptional program to prepare for life outside the host. Using a host-to-environment transition assay, these genes are shown to be important for V. cholerae to persist within cholera stool and/or aquatic environments. In addition, biogenesis genes required for the environmental adhesin, MSHA pili is also upregulated, further suggesting that V. cholerae are transcriptionally committing to life outside of the host during exit.

Quorum sensing regulates V. cholerae transmission and dissemination

Quorum sensing (QS) refers to the phenomenon in which bacteria produce and exchange chemical signals to monitor population density [135–138]. Many Gram-positive and Gram-negative bacteria use quorum sensing to control a variety of physiological functions. V. cholerae QS regulatory systems are highly complex. At low cell densities, the components of the QS pathway act as kinases to phosphorylate LuxO, which in turn activates the transcription of small RNAs (qrr1-qrr4) that destabilize mRNA of hapR, encoding the QS master regulator [139] and activate aphA, encoding the virulence regulator. At high cell densities, two sets of autoinducers, CAI-1 ((S)-3-hydroxytridecan-4-one) and AI-2 ((2S,4S)-2-methyl-2,3,3,4-tetrahydroxytetrahydrofuran borate) [140–142], bind to cognate sensors on the bacterial surface and induce conformational changes in the sensors [143], which results in dephosphorylation of LuxO. Thus, expression of the Qrr sRNAs is repressed by AI-2 and CAI-1. In addition, two other receptor proteins, CqsS and VpsS, have been reported to channel information through LuxO [144]. More recently, Recently, Bassler’s group has discovered another QS system [145] that does not require LuxO or the Qrr sRNAs, but rather relies on another autoinducer, DPO (3,5-dimethylpyrazin-2-ol), which is synthesized by the threonine dehydrogenase Tdh. DPO is sensed by VqmA, a LuxR-type transcriptional regulator [146], which induces the transcription of the VqmR sRNA. VqmR inhibits transcription of multiple target, including vpsT [97,147] that encodes a key activator of biofilm formation, as well as the virulence regulator gene aphA [148,149].

V. cholerae employs QS systems to temporally control virulence during infection. It has been shown that QS represses virulence gene expression and biofilm formation while activating production of extracellular proteases, suggesting the importance of QS in entering and exiting the host [15,54,150,151]. Surface-attached V. cholerae (as in biofilms) may be the major entry route for V. cholerae infection as simple filtration using used sari cloths reduces cholera significantly [152]. The biofilm structure may be critical during entry into the host in order to protect against acid shock in the stomach [15]. After reaching the intestine, dispersal of individual cells from the biofilm leads V. cholerae to transition away from a high cell density state, leading to de-repression of virulence. However, the reduction in cell density alone may not be sufficient to completely inhibit HapR-mediated repression of virulence gene expression. By penetrating mucus barrier, V. cholerae utilize flagellar regulatory systems to further repress hapR [153]. Later in the infection, the number of V. cholerae in the intestine increases, and quorum sensing again represses CT and TCP production and activates production of proteases, which serve as detachases [154]. Detachment from the epithelium could permit individual cells to establish new infection foci in the intestine or to exit the host. This mucosal escape response is also mediated by the stationary phase alternative sigma factor RpoS [155]. In addition to these quorum sensing molecules, it has been shown that other small molecule metabolites, such as indole and cyclo(Phe-Pro) (cFP) that accumulate to high concentrations at the stationary phase, inhibit virulence through a global regulator LeuO [156,157]. The QS regulon also consists of a number of additional genes involved in chitin-induced natural competence, stress responses, and hemolysin production, phage
production, among others [113,158–161], suggesting of the importance of QS mechanisms to adaptation to diverse environmental niches [136].

Interestingly, a large percentage of natural V. cholerae isolates are QS-deficient [162–164], implying a selective advantage of QS mutants in nature. It has been reported that these QS mutants actively cheat through signaling others to produce QS-dependent “public goods”[165], suggesting that social cheating may drive QS deficiency emergence within V. cholerae natural populations. The loss of QS regulation may also improve virulence gene expression in the gut.

The role of the gut microbiome in V. cholerae pathogenesis

The gut microbiome is a highly diverse assemblage of microorganisms representing all domains of life, though dominated by the eubacteria. While many human body sites host resident or transiently present microbes, the gastrointestinal tract represents the densest site of continuous microbial colonization; it is thought that bacterial cells in the gut outnumber human somatic cells, and the genetic diversity of the assembled genomes of gut resident species far eclipses that of humans [166,167]. The gut microbiome has been implicated in numerous host phenotypes, and research has recently begun to focus on the molecular interactions of this commensal microbial community and enteric pathogens such as V. cholerae (Figure 2).

Impact of cholera on the gut microbiome

The characteristic “rice water stool” produced during acute cholera is dominated by vibrios [168], and was an early clue as to the etiology of the disease in humans. Culture-dependent and culture-independent efforts have since defined at a much greater taxonomic resolution the effects of cholera on the bacteria within human gut microbiome. The fulminant diarrhea caused by cholera has predictably deleterious effects on the abundance and diversity of the gut microbial population. Culturing studies have shown a multi-log decrease in cultivable non-Vibrio bacteria during cholera in stool during acute disease in adults compared to convalescent populations [168]. More recently, metagenomic techniques, focusing largely on high-throughput sequencing of PCR amplicons of the 16S ribosomal small subunit gene in fecal specimens, have been used to probe for gut microbiome changes at high taxonomic resolution. Hsiao et al. closely tracked adult cholera patients in Bangladesh from presentation at clinic to 3 months of convalescence, and showed that the microbiome became dramatically less diverse during diarrhea, becoming dominated by mostly streptococcal species [169]. Existing gut microbes were detected at very low abundance, but over the course of recovery from diarrhea expanded to re-populate the gut in a successional process similar to the initial colonization of the gut during infancy and childhood. This drop in diversity is paralleled in findings in children with cholera [170]. Subsequent 16S amplicon sequencing-based studies in malnourished children with cholera showed an increase in Enterobacteriaceae, Veillonellaceae, and Streptococcaceae during infection [171]. A transient but dramatic dysbiosis in microbial structure has since been reported not only for cholera, but diarrhea of multiple etiologies [172,173] and in severe malnutrition [174]. These patterns in fecal microbiome complexity and membership during dysbiosis are paralleled in the small intestine, where E. coli, Streptococci, and aerobic lactobacilli were found in abundance in the duodenum and jejunum by culturing [175].

Gut microbiome structure as a driver of colonization resistance

A role for commensal microbes in V. cholerae infection outcome has been long recognized. Freter et al. showed in the 1950s that guinea pigs and mice whose commensal flora had been depleted by antibiotics were susceptible to V. cholerae colonization, while untreated animals were highly resistant [176]. This is in contrast to germfree mice, which support very high levels of V. cholerae colonization in both the distal small intestine and distal gut compartments [177]. Limitations in the ability to define, culture, and establish in vivo complex mixtures of microbes in the gut has until recently prevented work at the species resolution on the role of the human, as opposed to the rodent, gut microbiome, which differ in both species richness and composition [178].

Several recent studies have used a variety of molecular and animal model approaches to examine the role of human microbiome structure in susceptibility to infection. Hsiao et al. established in germfree mice defined communities of cultured human gut commensal bacteria closely modeled on normal healthy human gut communities [169]. The defined model healthy human microbiome were found to be highly resistant to V. cholerae, and one gut microbe commonly found in healthy human populations, Blautia (formerly Ruminococcus) obeum drove a large proportion of this colonization resistance phenotype. Specific exclusion of this species in communities established in germfree mice yielded significantly higher V. cholerae colonization, and direct competition of V. cholerae and B. obeum led to a 2-log decrease in pathogen load compared to germfree conditions. This suggested that human microbiomes could be a susceptibility factor for V. cholerae colonization, and
that these effects may be highly specific to certain microbiome members; engineered communities with and without *B. obeum* in the above study had negligible differences in overall phylogenetic diversity, and the closely related *Blaustia torques* showed no effects on *V. cholerae*. Midani et al. examined complete human fecal microbiomes of household contacts of cholera patients in Bangladesh that did or did not subsequently develop symptomatic infection to identify microbial correlates of infection susceptibility in full human microbiomes [179]. Using a machine learning approach, they identified several species associated with household contacts that remained uninfected, including *Blaustia, Ruminococcus,* and *Prevotella* species, while *Streptococcus, Prevotella,* and *Blaustia* species were higher in infected contacts. Differences in alpha diversity were not associated with subsequent infection. That species of the same genus were associated with both infected and uninfected outcomes further highlights the likely specificity of species and their biochemical functions in determining the outcome of exposure to *V. cholerae*.

Work by Alavi et al. has experimentally confirmed the role of differences in human gut microbiomes in *V. cholerae* colonization outcome [180]. These studies established defined model and complete human fecal microbiomes in germfree mice and suckling animals with their native murine microbiomes cleared using antibiotics. High-*Streptococcus* microbiomes modeled after diarrhea- and malnutrition-disrupted gut communities were highly susceptible to *V. cholerae* infection. Surprisingly, complete fecal microbiomes from healthy US donors transplanted into suckling animals yielded ~30-fold differences in subsequent *V. cholerae* colonization. These findings suggest that commensal microbe-dependent disease susceptibility differences are not strictly dichotomous between “normal” vs “diseased” microbiome states, and that interpersonal differences in microbiome structure can drive infection outcomes. Furthermore, this suggests that disruption of the microbiome by other infectious diarrheas or malnutrition may also be a risk factor for cholera.

*Commensal microbial bile metabolism modulates V. cholerae virulence*

Similarly to *V. cholerae*, commensal gut microbes adapted to the small intestine have evolved mechanisms to deal with the bacteriostatic qualities of bile. Microbial bile salt hydrolase (*bsh*) enzymes can remove the conjugated amino acid of primary bile salts, and a series of microbial enzymes can dehydroxylate the 7th position of sterol backbone of bile (7α-dehydroxylation). Microbial *bsh* enzymes are broadly distributed within the genomes of human gut-associated species and play an important role in microbial bile tolerance and bile recirculation. Deconjugation of amino acids attached to primary bile salts such as taurocholate and glycocholate reduces the hydrophilicity of these molecules, leading to a drop in their detergent-like effects [185,186]. Though many gut commensal microbes are refractory to genetic manipulation, mutations in *bsh* genes in *Lactobacillus amylovorus* and *L. plantarum* have been associated with sensitivity to bile and bile salts [186,187]. That microbial metabolism of bile salts is a major player in the composition of the bile acid pool has been demonstrated in germfree animal systems; in germfree mice, essentially all small intestinal bile are conjugated primary bile salts (taurocholate and tauromuricholate) [188]. Bioinformatic studies have identified several broad *bsh* phylotypes within commensal microorganisms, which have different predicted bile salt substrates and activity [58]. The presence of specific

*B. obeum* has been shown to upregulate production of an AI-2 molecule in response to *V. cholerae* infection when colonized in germfree mice [169]. Expression of the *B. obeum* AI-2 synthase LuxS in an AI-2-null *E. coli* was sufficient to restrict *V. cholerae* colonization. This signaling was independent of the canonical AI-2 sensor LuxP, as *luxP* deletion did not rescue *V. cholerae* colonization in the presence of *B. obeum*. This suggests that AI-2 molecules produced by different members of the gut microbiome may differentially integrate into the virulence regulatory cascade. LuxS homologs are widely distributed in the genomes of gut commensals [182], though it is uncertain if there is substantial structural diversity in the resulting autoinducer molecules. Two structures of AI-2 have been elucidated, the AI-2 of *Vibrio* being a furanosyl borate diester, in contrast to a non-borated molecule synthesized by *Salmonella* [140,183]. Cross-feeding experiments demonstrate that *V. cholerae* can respond to AI-2 from *E. coli* [184]. That LuxP seems to be dispensable in *V. cholerae* regulatory responses to *B. obeum* AI-2 suggests that there may be further structural diversity in this QS molecule as made by different gut commensals.

**Commensal microbial quorum sensing in V. cholerae infection**

The complex and tightly coordinated virulence regulatory cascade of *V. cholerae* can be modulated by autoinducer molecules from various sources. One mechanism of interaction between commensal microbes and *V. cholerae* pathogenesis is the production of cross-species autoinducers. *V. cholerae* responds to a set of species-specific (CAI-1) and potentially cross-species (AI-2, DPO, ethanolamine) signals in gene regulation [144,149,181]. The human commensal
sets of bacteria in the small intestine can thus lead to differential activity against components of the bile acid pool, with consequent effects on the chemical environment of the small bowel.

Recent work has shown that bsh activity by *V. cholerae*-restricting microbes such as *B. obeum* strongly contributes to the outcome of *V. cholerae* infection [180]. Using an *in vitro* screen for activity against bile acids, Alavi et al. showed that *B. obeum* encodes for a bsh level with high activity against taurocholate, a key activator of *V. cholerae* virulence in mice and humans [68]. Overall levels of bsh activity were higher in organisms characteristic of healthy human gut microbiomes *in vitro* and bsh genes were more abundant in sequenced metagenomic datasets of fecal samples from healthy Bangladeshi adults compared to dysbiotic, *V. cholerae*-susceptible microbiomes. Genes encoding for these enzymes were found in lower abundance in post-diarrhea fecal microbiomes, and taurocholate deconjugating activity was less frequently found in isolates of species characteristic of these dysbiotic guts communities. *B. obeum* bsh activity has been directly correlated to the ability of complete complex human fecal microbiomes to resist *V. cholerae* infection in suckling mice; communities with *B. obeum* led to lower tcpA expression during infection of suckling animals [180]. This was independent of AI-2 production, as *B. obeum* reduced tcpA activation by intestinal tissues even when boiled to remove AI-2, and constitutive expression of *B. obeum* bsh was able to significantly restrict *V. cholerae* colonization. This suggests that microbiome dysbiosis can lead to increased *V. cholerae* susceptibility via a loss of bsh activity, and represents a recurrent and individual-specific window of vulnerability to infection in human populations in cholera endemic areas.

**Driving disease resistance using the microbiome**

Several proof-of-principle studies have demonstrated the effectiveness of engineering probiotic strains to modulate *V. cholerae* colonization *in vivo*. Duan et al. used a probiotic *E. coli* Nissle strain engineered to produce CAI-1 via constitutive *cqsA* expression, and showed that introduction of this strain into suckling mice reduced susceptibility to subsequent *V. cholerae* challenge in a QS-dependent manner [189]. Recently, Mao et al. reported the effectiveness of an engineered *Lactococcus lactis* that can serve as a sensor of *V. cholerae* infection via detection of *V. cholerae*-specific autoinducers and consequent production of an enzymatic indicator. This strain was also able to promote colonization resistance via decreases in intestinal pH [190]. Co-opting the metabolic activities of commensal microbes has also been shown to reduce *V. cholerae* susceptibility. Expression of other autoinducers produced by microbiome members such as AI-2 (via luxS expression) [169] and manipulation of bile acid pools (via bsh expression) [180] has demonstrated effectiveness in mouse infection models, using *E. coli* as an expression platform. Engineered *V. cholerae* may also have “probiotic” effects on subsequent infection; recent studies have shown that a non-virulent candidate live vaccine strain rapidly outcompeted fully virulent *V. cholerae* in infant rabbits [191]. However, it is unclear whether the use of existing probiotic species for introduction of specific microbial functions will have durable efficacy as a prophylactic compared to introduction of fully gut-adapted species with multiple levels of activity; commercial probiotic strains have demonstrated low effective colonization in germfree animals [192] and *V. cholerae* are rapidly excluded by normal commensal microbes in animal models in the absence of successful colonization leading to toxin production and diarrhea [169]. The identification of human commensal microbes with efficacy against *V. cholerae* virulence, for example *B. obeum*, the source organism for the luxS and bsh mechanisms described above, may allow for more stable prophylactic effects on the host due to these organisms being better adapted to colonization of the human gastrointestinal tract.

**Inter-bacterial competition and V. cholerae virulence**

*V. cholerae* has evolved numerous mechanisms as part of infection that are able to deplete the native microflora of the small intestine. The profuse watery diarrhea of cholera is one obvious element of this process, as this rapidly drops the levels of commensal microbes in both rice water stool and the small intestine [168,169,175]. Recent work has identified other biochemical pathways by which this human pathogen can influence gut commensal levels. The Type-six secretion system (T6SS) is a molecular complex that enables *V. cholerae* to kill other Gram-negative cells and modulate host cell behavior via contact-dependent cell-cell secretion of effector proteins [193,194]. These systems have been identified in several Gram-negative species associated with the GI tract [195,196]. Several lines of evidence suggest that this mechanism of microbial competition plays an important role in *V. cholerae* pathogenesis. Genetic deletion of the T6SS leads to a colonization defect or loss of virulence in several animal models, including infant rabbits [197], *Drosophila* [198] and suckling mice [180,199]. *V. cholerae* uses the T6SS to directly and indirectly target competitor commensal microbes.
This interbacterial competition can be via direct killing; *V. cholerae* is able to use T6SS to kill murine commensal *E. coli* isolates during colonization of sucking mice [199]. Interestingly, this killing leads to an increased upregulation of virulence gene expression over and above that seen during the *in vitro* to *in vivo* transition. This has also been demonstrated in a *Drosophila* model of infection, where T6SS interactions with the commensal *Acetobacter pasteurianus* led to increased disease symptoms including host death [198]. T6SS can also play a role in displacing competitive commensal organisms as demonstrated in *V. cholerae* colonization of zebrafish [200], where T6SS mediated increased gut motility to provide a competitive advantage to the pathogen over resident bacteria.

The T6SS may also be regulated by processes related to commensal metabolism in the gut. Several studies have also shown a link between Q5 and T6SS regulation; the key Q5 regulator HapR has been shown to directly initiate T6SS gene expression [201] and indirectly via the transcription factor QstR [202]. Q5-dependent T6SS regulation has been shown to be conserved across pandemic and non-pandemic strains of *V. cholerae* [203]. The presence of numerous interspecies signaling molecules in the gut may thus also be associated with upregulation of competitive cell-cell killing mechanisms during infection. The layer of highly glycosylated mucins that form the intestinal epithelial mucus barrier is also sensed by *V. cholerae* to de-repress expression of T6SS related genes [204]. T6SS gene expression can also be regulated by specific bile acids; deoxycholic acid, a metabolic product of cholic acid formed by 7α-dehydroxylation by microbial action, has been shown to inhibit T6SS assembly [204]. These processes may intersect with the action of the gut microbiome. Commensal microbes are found in large numbers in mucus [205,206]; several microbes, for example *Akkermansia muciniphila* [207], are known to be highly adapted to use human mucus components as nutritional sources, while the role of microbial enzymes in bile metabolism has been heavily studied. Processing by commensal microorganisms of mucin and bile may thus play important but as yet un-elucidated roles in T6SS regulation.

**Perspectives and future studies**

As experimental and bioinformatic techniques for probing microbe-microbe and microbe-host interactions during *V. cholerae* infections advance, we can expect new discoveries in how the gut microbiome shapes the *in vivo* signals that drive *V. cholerae* infection. While oral rehydration with antibiotic treatment is highly effective at reducing mortality due to cholera, cholera remains a formidable global health challenge demanding of new therapeutic or prophylactic approaches. While the development of antibiotic resistance in *V. cholerae* is of growing concern [208], therapeutic or prophylactic approaches using the microbiome that target colonization, virulence gene expression, or nutrient competition is unlikely to be as strong a driver for the development of resistance mechanisms. Therapies leading to reductions in mortality also largely do not reduce the high morbidity of cholera. It is increasingly clear that the variability of the microbiome, from individual to individual, and within individuals in response to environmental stimuli, determines a personalized infection outcome, depending on the presence and activity of key commensal microbes. Thus, manipulation of microbial colonization, if understood mechanistically and rationally designed, may offer durable prophylactic or therapeutic efficacy against cholera. Probiotics that can durably establish in the context of complex human fecal gut communities, as opposed to simplified animal model system, critical for such approaches; the identification of commensals such as *B. obeum* with anti-virulence activity that are native to the human gastrointestinal tract and able to establish colonization across different human gut microbiomes is thus of increasing significance [169,180].

Another avenue of research of key importance to the field is the development of new and tractable experimental models to identify candidate microbes mediating infection resistance and to test targeted microbiome manipulations. Mice without native murine microbes, whether germfree [169,180,192] or antibiotic-cleared [199], can serve as hosts for complete or defined model human fecal gut microbiomes in the context of *V. cholerae* colonization and infection. These systems allow researchers to insert, remove, or mix microbes and assemblages to determine their effects on pathogenesis and *in vivo* pathogen fitness. While *V. cholerae* behavior differs in germfree adult mice compared to suckling animals by lower CT expression [169], this experimental model allowed for the targeted establishment of human gut bacteria to evaluate their effects on colonization, and extended studies of *in vivo* metabolism and cell-cell interactions that are difficult to accomplish with shorter-term models such as suckling mice or suckling rabbits.

Consistency of effects on colonization resistance will be of key importance in identifying candidate next-generation probiotics. Combinatorial approaches, where microbial subsets are randomized to determine whether specific microbes play active roles *in vivo* regardless of other colonizing microbes, have also been used to identify contributors to host immunity and *V. cholerae* colonization resistance in the
context of quite different background microbiomes [180,209], a necessary characteristic in a broadly efficacious next-generation probiotic.

The role of diet and nutrition, both of the host and of V. cholerae during colonization, may also play an important role in future work. Microbiome structure is strongly driven by host diet [210–213]. However, the role of dietary components, and what nutrient sources are most important for rapid V. cholerae proliferation in vivo, have not been extensively studied. The role of commensal microbes in setting the stage for early infection through contribution to the gut nutrient landscape is also largely unknown. Prebiotic approaches, which use targeted nutritional interventions to drive specific effects on gut microbiomes, may also be a fruitful area of research for targeted microbiome modification as applied to V. cholerae colonization resistance.

The expansion in multi-omics approaches, microbial ecological analysis, and experimental animal models of V. cholerae infection and in vivo fitness have revealed numerous fascinating aspects of the biology of this important human pathogen and how it is impacted by the variable membership and biochemistry of the human gut microbiome. Future advances will capitalize on the translational potential of these findings, as pathogenesis starts to incorporate new understandings of the microbiome to produce prophylactic and therapeutic interventions for cholera.

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