Extracytoplasmic-Stress-Responsive Pathways Modulate Type III Secretion in *Yersinia pseudotuberculosis*

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Three signal transduction pathways, the two-component systems CpxRA and BaeSR and the alternative sigma factor $\sigma^E$, respond to extracytoplasmic stress that facilitates bacterial adaptation to changing environments. At least the CpxRA and $\sigma^E$ pathways control the production of protein-folding and degradation factors that counter the effects of protein misfolding in the periplasm. This function also influences the biogenesis of multicomponent extracellular appendages that span the bacterial envelope, such as various forms of pili. Herein, we investigated whether any of these regulatory pathways in the enteropathogen *Yersinia pseudotuberculosis* affect the functionality of the Ysc-Yop type III secretion system. This is a multicomponent molecular syringe spanning the bacterial envelope used to inject effector proteins directly into eukaryotic cells. Disruption of individual components revealed that the Cpx and $\sigma^E$ pathways are important for *Y. pseudotuberculosis* type III secretion of Yops (*Yersinia* outer proteins). In particular, a loss of CpxA, a sensor kinase, reduced levels of structural Ysc (*Yersinia* secretion) components in bacterial membranes, suggesting that these mutant bacteria are less able to assemble a functional secretion apparatus. Moreover, these bacteria were no longer capable of localizing Yops into the eukaryotic cell interior. In addition, a cpxA lrQ double mutant engineered to overproduce and secrete Yops was still impaired in intoxicating cells. Thus, the Cpx pathway might mediate multiple influences on bacterium-target cell contact that modulate *Yersinia* type III secretion-dependent host cell cytotoxicity.

Type III secretion (T3S) is an extensively studied virulence mechanism used by many gram-negative bacterial pathogens of both animals and plants. Resembling a syringe composed of ~25 protein components that span the bacterial envelope (52, 56), T3S systems (T3SSs) efficiently secrete and deliver toxic effectors into target cells during intimate contact with a host (27, 30). This enables bacteria to evade host immune responses and facilitate often lethal infections.

Pathogenic bacteria occupy a diverse array of infection niches inside a host. Each niche boasts a unique combination of environmental signals that serve as a detailed grid reference, allowing individual bacteria to monitor their precise location (4, 9). However, bacterial infections are dynamic processes, so bacteria must constantly recognize and respond to different sets of environmental cues depending on the stage of infection. This active spatial and temporal environmental fingerprint is interpreted by multiple and overlapping control factors that coordinate regulation of bacterial virulence gene expression, ensuring that each virulence factor is expressed only when needed (4, 9).

The bacterial cell envelope is a critical data-processing center. It contains multiple relay pathways that transmit external signals into the bacterial cytoplasm. These activate subsets of DNA binding proteins that cooperate with RNA polymerase to specifically stimulate gene transcription (4, 9). At least three of these pathways respond to extracytoplasmic stress (ECS): the two-component systems CpxRA and BaeSR and the alternative sigma factor $\sigma^E$ (RpoE) (1, 22, 79). Growth in adverse environments triggers the CpxRA- and $\sigma^E$-responsive elements to induce the production of protein-folding and degradation factors designed to counter protein misfolding in the bacterial periplasm. It is less clear what is induced by the BaeSR system, although an *Escherichia coli* transcriptome analysis indicated a regulon that includes genes related to carbohydrate and multidrug transporters, chemotaxis responses, and flagellum biosynthesis (69). Another group of proteins, collectively functioning in the phase shock response, may also aid in tolerance to periplasmic stress (61). Interestingly, these pathways are now also linked to bacterial pathogenesis (16, 79, 85). They ensure the accurate assembly of multicomponent virulence determinants spanning the bacterial envelope, such as P pilis (41, 45), type IV bundle-forming pilis (68), and curli (47).

Human-pathogenic *Yersinia* spp. all harbor a large virulence plasmid encoding the Ysc-Yop T3SS (11). Use of this system to translocate several toxic effectors into the target cell interior enables *Yersinia* to resist phagocytosis and proliferate extracellularly (24, 91). Yop synthesis and secretion are controlled by temperature and calcium in laboratory media or target cell contact (76, 84). While the virulence plasmid encodes several built-in regulatory elements (55), only a few additional chromosomally encoded regulatory factors have been described: YmoA (12) and its regulated proteolysis via the ClpXP and Lon proteases (44), adenylyl cyclase (Cya) and the cyclic AMP receptor (Crp) (75), the SmpB-SsrA translational control system (70), DNA adenylation methylase overproduction (48), and the tsaA allele of unknown function (17). Although the
mechanism by which most of these components modulate T3S remains obscure, it illustrates how the inherent complexity of T3SSs requires fine-tuning by the activities of multiple regulatory factors. We propose that Ysc-Yop biogenesis in the bacterial envelope also engages one or more of the ECS response pathways. Herein, we describe the systematic disruption of components T3SSs requires fine-tuning by the activities of multiple regulatory factors. We propose that Ysc-Yop biogenesis in the bacterial envelope also engages one or more of the ECS response pathways. Herein, we describe the systematic disruption of components T3SSs requires fine-tuning by the activities of multiple regulatory factors.

**MATERIALS AND METHODS**

**Bacterial strains, plasmids, and growth conditions.** Bacterial strains and plasmids are listed in Table 1. Bacteria were routinely cultivated in Luria-Bertani (LB) agar or broth at either 26°C (Y. pseudotuberculosis) or 37°C (E. coli) with aeration. To examine T3S, we used Y. pseudotuberculosis YPIII/pIB102 (serotype III), in which pIB102 is the virulence plasmid encoding the Ysc-Yop T3SS. This plasmid is a derivative of pIB1 (7, 29), distinguished only by a kanamycin resistance cartridge inserted into the virulence cassette. Kanamycin resistance was conferred from the mutagenesis vector pJF006. This fragment was cloned via XbaI/KpnI restriction and subcloned into pBBR1MCS-3 (51) to create plasmid pKEC010. Gene expression is under the control of IPTG (isopropyl-B-D-thiogalactopyranoside).

**TABLE 1. Bacterial strains and plasmids used in this study**

<table>
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<tr>
<th>Strains or plasmid</th>
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<td><strong>E. coli</strong></td>
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<tr>
<td>S17-1</td>
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<td><strong>Y. pseudotuberculosis</strong></td>
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<tr>
<td>YPIII/pIB102</td>
<td>yadA::Tn5 Km' (wild type)</td>
<td>8</td>
</tr>
<tr>
<td>YPIII/pIB102 [cpxA in-frame deletion of codons 41 to 449, Km']</td>
<td>This study</td>
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<td>YPIII/pIB102 [cpxA in-frame deletion of codons 41 to 449, Km']</td>
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<td><strong>Plasmids</strong></td>
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<td>pC4-TOPO</td>
<td>TA cloning vector, Km'</td>
<td>Invitrogen</td>
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<tr>
<td>pMD4</td>
<td>Suicide plasmid carrying sacBR, Cm'</td>
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<td>pJE006</td>
<td>925-bp Xhol/XbaI PCR fragment of cpxA(41–449) in pMD4, Cm'</td>
<td>This study</td>
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<td>485-bp Xhol/XbaI PCR fragment of cpxR(11–193) in pMD4, Cm'</td>
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<td>Expression vector, Cm'</td>
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<td>pKEC021</td>
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<td>pET22b(+)</td>
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<td>pMF581</td>
<td>1,381-bp NdeI/EagI PCR fragment of cpxA in pET22b(+), Cb'</td>
<td>This study</td>
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mM CaCl2 (medium with Ca²⁺). Induction of secretion followed after a growth shift from 26°C to 37°C. Where required, antibiotics were added at the final concentrations of 100 μg per ml carbenicillin (Cb), 50 μg per ml kanamycin (Km), 20 μg per ml spectinomycin (Sp), and 25 μg per ml chloramphenicol (Cm).

**Plasmid construction.** Amplified DNA fragments used for constructing individual in-frame deletions were generated by overlap PCR (38). The primer combinations used to create each mutation are listed in Table 2 and were synthesized by either DNA Technology A/S (Aarhus, Denmark) or TAG Copenhagen A/S (Copenhagen, Denmark). All amplified fragments were first cloned into pCR4-TOPO TA (Invitrogen AB, Stockholm, Sweden) and then sequenced commercially (MWG Biotech AG, Ebersberg, Germany). Confirmed fragments were then subcloned into the Xhol/XbaI-digested suicide mutagenesis vector pDM4 (58). Conjugal mating experiments with Y. pseudotuberculosis and selection for the appropriate allelic exchange events used conventional methodology with one exception: standard LB agar with appropriate antibiotics was used to selectively cultivate Yersinia after conjugation (28, 58).

**Full-length cpxA and cpxR were amplified by PCR using the primer combinations listed in Table 2. The cpxA fragment was cloned into pET22b (Novagen, Madison, WI) using NdeI/EagI restriction to develop pMF581. The cpxR fragment went through a series of cloning events: it was first cloned into pET22b using NdeI/Xhol-compatible ends to give pKED010 and was then excised using Xbal/Xhol restriction and subcloned into pBRR1MCS-3 (51) to create plasmid pMF687 before a final round of cleavage from pMF687 to yield an XbaI/KpnI fragment that was subcloned into the low-copy-number expression vector pMMB208 (62) to give pKED021. In addition, cpxR(D51A), encoding an alanine substitution of the phosphorylatable aspartate residue at position 51, was amplified from the mutagenesis vector pF006. This fragment was cloned via XbaI/KpnI restriction into pMMB208, yielding pF015. Gene expression is under the control of IPTG (isopropyl-B-D-thiogalactopyranoside).
Table 2. Oligonucleotides used in this study

<table>
<thead>
<tr>
<th>Purpose</th>
<th>Oligonucleotide pair(s) (sequence)*</th>
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<tr>
<td>Mutagenesis (plasmid)</td>
<td>ΔbaeR (pMF682) pbaeSa (5'-GTC ATG GTG GTA AGT GAC AGC AAC ATC CAT GAA GAG-3') and pbaeSb (5'-TAT TAA CAC TAA CAT GCA AGT G-3') pbaeSc (5'-TGG ATG GTG GTA AGT GAC AGC AAC ATC CAT GAA GAG-3') and pbaeSd (5'-TCT AGT GTC ATG GAC AAC ATC CAT GAA GAG-3')</td>
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<td>Complementation</td>
<td>ΔbaeR (pKEC009) pbaeRa (5'-TGG ATG GTG GTA AGT GAC AGC AAC ATC CAT GAA GAG-3') and plcrFrev (5'-TCT GCA CAC CTT CCA GAT ATC AGC GGG TTC TTA TAC AG-3')</td>
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<tr>
<td>Mutagenesis (plasmid)</td>
<td>ΔapcB (pMF586) ppcxB (5'-AGC TTC GAG AAC GAA CAT TGA CCG CAT C-3') and ppcxA (5'-AGC TTC GAG AAC GAA CAT TGA CCG CAT C-3')</td>
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<td>ROLE OF CpxRA IN YERSINIA TYPE III SECRETION</td>
<td>pcppR (pJF006) pcppR (5'-TAA CTA CAC TAA CAT GAA GAG-3') and pcppR (5'-TAA CTA CAC TAA CAT GAA GAG-3')</td>
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* The nucleotide sequences in italics identify the incorporated XhoI, XbaI, NdeI, KpnI, or EagI restriction sites used for cloning of the amplified DNA fragments. The underlined sequence indicates the complementary overlap between respective primers in the overlap PCRs.
and grown for approximately 1.5 h to an optical density at 600 nm of 0.4 to 0.6. RNA was immediately stabilized by the addition of the RNAProtect bacterial reagent (QIAGEN Nordic, West Sussex, United Kingdom) and then extracted using the Nucleospin RNA II method (Macherey-Nagel, Düren, Germany) including an on-column DNase treatment. To generate cDNA, 0.2 μg of total RNA was reversed transcribed with the ImProm-II reverse transcription (RT) system (Promega, Madison, WI). To confirm the absence of contaminating DNA in the RNA preparations, a duplicate reaction lacking the RT enzyme was always performed in parallel. Transcripts derived from genes of the T3S and belonging to ECS regulons were detected by PCR with the specific internal primer combinations listed in Table 2. The conditions for DNA amplification in a 20-μl reaction mixture included 0.5 μl template and the Easy-A high-fidelity PCR cloning enzyme (Stratagene). Fractionated DNA fragments were visualized by using a Fluor-S Multimager (Bio-Rad), and the density of each fragment was analyzed with Quantity One software (version 4.2.3; Bio-Rad Laboratories, Sundbyberg, Sweden). Cycling conditions used for cDNA amplification were an initial denaturation step at 94°C for 3 min; denaturation at 94°C for 30 s, annealing at 50°C for 30 s, and extension at 72°C for 30 s (5 cycles); and then denaturation at 94°C for 30 s, annealing at 55°C for 30 s, and extension at 72°C for 30 s (25 cycles), before a final extension step at 72°C for 5 min in a GeneAmp PCR system.

Analysis of Yop synthesis and secretion. Routine analysis of Yop synthesis and secretion was performed by using established methods (25, 26, 28, 71). Protein fractions separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and subjected to immunoblotting were identified with rabbit polyclonal antiserum raised against a total pool of secreted Yops or the nonsecreted Y. pestis V47-2. Whole-membrane preparations.

Identification of Yersinia ECS-responsive pathways. T3S requires the assembly of ~25 components, which together span the bacterial envelope. Coordinated assembly of these building blocks might require periplasmic quality control systems to deal with misfolded or misassembled proteins. Known quality control systems of the bacterial envelope that respond to ECS include a pair of two-component phosphorelay systems, the sensor kinases BaeS and CpxA and their respective cognate response regulators BaeR and CpxR, and the alternate sigma factor RpoE (1, 22, 79). To investigate if the biogenesis of the Ysc-Yop T3S of Y. pseudotuberculosis engages one or more of the ECS-responsive pathways, we used the genome sequence generated from Y. pseudotuberculosis IP32953 serotype I (10) to locate the baeSR operon (10), baeR, cpxR, cpxA, and rpoE genes. The operons and their genetic neighborhood are shown in Fig. 1A. The gene arrangement surrounding the three operons bears some similarity to that found for E. coli K-12, particularly with those genes flanking the baeSR operon (6). However, alleles unique to Yersinia with no known function disrupt this gene order. This is most evident for the genes in the neighborhood of the cpxR4 operon (Fig. 1A). Despite these differences, identity to the equivalent proteins of E. coli is high for BaeS (64% amino acid identity to BaeS of E. coli), BaeR (73%), CpxA (80%), CpxR (89%), and RpoE (92%).

Using RT-PCR, we first analyzed the expression of these genes in Yersinia cells grown at low (26°C) and high (37°C) temperatures in BHI medium that is restrictive (plus Ca²⁺) or permissive (minus Ca²⁺) for Yop synthesis and secretion. Although not mimicking ysc- or yop-inducible expression, Yersinia clearly reacted to growth under these conditions. Without exception, Ca²⁺ depletion elevated the three ECS regulons at low growth temperatures (Fig. 1B). However, this same medium restricted gene expression within these regulons at high growth temperatures. Moreover, baeS, cpxR, and rpoE were maximally expressed at elevated temperatures in the presence of Ca²⁺. Intriguingly, therefore, it appears that under conditions where T3S gene expression is maximal, ECS responsiveness is actually reduced. We interpret this seemingly inverse relationship to indicate cross talk between T3S and the ECS-responsive pathways in Yersinia. However, it might not be as straightforward as ECS-responsive pathways overseeing the T3SS assembly process.

Functional Ysc-Yop type III secretion requires an intact Cpx pathway. As a consequence of these findings, we generated in-frame, near-full-length deletions of baeS, baeR, cpxA, cpxR, and rpoE in Y. pseudotuberculosis YPIII/pIB102 serotype III (Fig. 1A). We confirmed by RT-PCR that none of these deletions generated a polar effect on the transcription of genes flanking each deletion (Fig. 1C and data not shown). The null mutants were given the following designations: YPIII40/pIB102 (∆baeS), YPIII29/pIB102 (∆bace), YPIII08/pIB102 (∆cpxR), YPIII07/pIB102 (∆cpxA), and YPIII09/pIB102 (∆rpoE).

Yersiniae induce the secretion of Yops through the Ysc-Yop system when grown at elevated temperatures in the absence of Ca²⁺ ions. We used these growth conditions to examine Yop secretion by mutant bacteria deficient in a particular ECS-responsive pathway. Mutants no longer producing BaeS, BaeR, or CpxR still maintained the production and secretion of Yops, occurring only in the absence of Ca²⁺ in a pattern akin to that of wild-type bacteria (Fig. 2A). In contrast, the ΔcpxA null mutant displayed a marked reduction in Yop production and secretion. This phenotype is due to the loss of the CpxA sensor kinase, because functional secretion could be restored in this mutant when ectopically producing a wild-type copy of CpxA (Fig. 2A). It should be noted that the ΔcpxA null mutant, but not the ΔcpxR equivalent, has a slight defect in growth during routine culturing (data not shown). This minor defect is not alone expected to cause the loss of T3S, because even wild-type Y. pseudotuberculosis cells cease to grow when producing and secreting Yops.

We also observed that the ΔrpoE null mutant was defective for Ysc-Yop T3S (data not shown). However, this mutant was apparently unstable and/or acquired second-site suppressor mutations that rapidly restored wild-type-like levels of Yop production and secretion. These isolates have not been further characterized. Given that rpoE is an essential gene in many...
bacteria, we mutated rseA and rseB, giving rise to strains YPIII28/pIB102 (ΔrseA) and YPIII42/pIB102 (ΔrseB), respectively (Fig. 1). These genes in E. coli encode proteins that modulate the σE response; RseA possesses anti-σE activity, and RseB exerts its negative effect by binding to RseA (19, 59). Their function in Yersinia would appear to be similar to that observed in E. coli, since amino acid identity (63% for RseA and 62% for RseB) is high and the gene arrangement within the operon is essentially conserved (Fig. 1). Interrogation of these mutants revealed that a loss of rseA consistently resulted in a modest elevation of Yop synthesis and secretion levels, with small amounts visible even under Yop-restrictive conditions (Fig. 2A). This is consistent with the loss of rpoE negatively affecting bacterial Ysc-Yop T3S (data not shown); the absence of rseA would be expected to activate the σE-dependent ECS response (2, 49). Thus, both the Cpx and σE ECS-responsive pathways affect Yersinia Ysc-Yop T3S.

The Cpx pathway affects type III secretion system biogenesis. Functional CpxA is necessary for the full production and secretion of Yops into laboratory media. We wondered...
whether this secretion defect was due to an inability to assemble fully functional secretion complexes. T3SSs are thought to be assembled in an ordered fashion whereby an appropriate incorporation of a preceding component serves as the platform for the subsequent assembly of additional components (50, 88). We therefore inspected whether the loss of CpxA induced a general assembly defect or if just one or at most a few specific components of the Ysc-Yop T3SS were affected. To begin to address this, we purified total membrane preparations of *Y. pseudotuberculosis* grown under both T3S-permissive and -restrictive conditions. We analyzed protein content by immunoblot using available rabbit antibodies raised against the core components YscU (54), YscJ (95), and YscP (23, 46); the needle components YscF and LcrV (37, 46, 63); and the secreted effector YopH (74). To control for membrane isolation, we also utilized antibodies that recognized the outer membrane protein TolC (89) and the inner membrane protein FtsH (90) in the blotting procedure. Interestingly, only levels of needle-associated YscJ, YscF, and LcrV and the effector YopH were reduced in the absence of CpxA (Fig. 3). YscU and YscP levels remained consistent regardless of the strain analyzed. Based on these findings, we suspect that the Cpx pathway might exert an activity at a specific step beyond the integration of YscU and YscP into the growing structure. This point would represent a Cpx-dependent T3S assembly checkpoint. Alternatively, it could indicate that the assembly of YscU and YscP does not involve the periplasm, thereby avoiding any involvement of the Cpx pathway.

**Analysis of T3S gene transcription.** It is also possible that CpxA may influence the activity of a regulator that in turn affects transcription from promoters lying upstream of specific

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**FIG. 2.** Analysis of Yop synthesis and secretion from *Y. pseudotuberculosis* strains grown either with (+) or without (−) Ca²⁺. Total Yops associated with bacteria or secreted free into the growth medium (expression fraction) (top) or contained solely within the cleared culture medium (secreted fraction) (bottom) were separated by SDS-PAGE and identified by immunoblot analysis using polyclonal rabbit antisera recognizing all secreted Yops or the cytoplasmic YerA chaperone. Where indicated (+), IPTG was added at a final concentration of 0.4 mM upon a temperature shift. (A) Wild-type strain YPIII/pB102, baeS null mutant strain YPIII40/pB102, baeR null mutant strain YPIII29/pB102, cpxA null mutant strain YPIII08/pB102, cpxA null mutant strain YPIII07/pB102, complemented ΔcpxA/cpxA⁺ strain YPIII07/pB102, pMF581, recA null mutant strain YPIII28/pB102, and recB null mutant strain YPIII42/pB102 are shown. (B) lcrQ::Sp-Sm mutant strain YPIII/pB26, lcrQ::Sp-Sm ΔcpxA double mutant strain YPIII07/pB26, and complemented lcrQ::Sp-Sm ΔcpxA/cpxA⁺ strain YPIII07/pB26, pMF581 are shown. (C) Noncomplemented ΔcpxR vector alone (YPIII08/pB102, pMMPB208), complemented ΔcpxR with nonphosphorylatable CpxR(D51A) (YPIII08/pB102, pJF015), and complemented ΔcpxR/cpxR⁺ (YPIII08/pB102, pKEC021) are shown. Molecular masses shown in parentheses are deduced from the primary sequence.

**FIG. 3.** Analysis of total membrane fractions of *Yersinia*. Bacteria were grown at 37°C in BHI medium either with (+) or without (−) Ca²⁺. The bacteria were washed and harvested by centrifugation, and the re-suspended pellets were sonicated to recover soluble material. The clarified lysate was subjected to ultracentrifugation to pellet total membrane material. Proteins contained within this material were fractionated by SDS-PAGE and immunoblotted using antibodies recognizing YerA (89) and the inner membrane protein FtsH (90) in the blotting procedure. Interestingly, only levels of needle-associated YscJ, YscF, and LcrV and the effector YopH were reduced in the absence of CpxA (Fig. 3). YscU and YscP levels remained consistent regardless of the strain analyzed. Based on these findings, we suspect that the Cpx pathway might exert an activity at a specific step beyond the integration of YscU and YscP into the growing structure. This point would represent a Cpx-dependent T3S assembly checkpoint. Alternatively, it could indicate that the assembly of YscU and YscP does not involve the periplasm, thereby avoiding any involvement of the Cpx pathway.

**Analysis of T3S gene transcription.** It is also possible that CpxA may influence the activity of a regulator that in turn affects transcription from promoters lying upstream of specific
T3S genes. However, one of the difficulties with pinpointing the precise function of CpxA during T3S is that the Ysc-Yop system is regulated by a phenomenon known as feedback inhibition (11). If secretion is compromised, the negative regulatory element LcrQ (YscM1 and YscM2 in *Yersinia enterocolitica*) remains trapped inside the bacteria, where it still exerts its repressive effect, so that both Yops production and secretion remain low (76). To analyze *ysc-yop* transcription in the *Y. pseudotuberculosis* wild type, the ΔcpxR null mutant, and the ΔcpxA null mutant, we employed the RT-PCR technique in which the template was cDNA derived from mRNA isolated from exponentially grown bacteria cultivated under T3S-permissive conditions (BHI broth depleted of Ca^{2+} ions). PCRs used specific primer pairs recognizing the genes encoding late-secreted products encompassing effectors (*yopE*, *yopH*, and *yopK*) and translocators (*yopD*), cytoplasmic chaperones (*yerA* and *lerG*), structural proteins (*yscF* and *yscN*), and regulatory proteins (*yopN* and *lerF*). To control for the responsiveness of the Cpx regulon, four confirmed CpxRA-regulated genes, at least in *E. coli*, were included in the analysis: *cpxP* (encoding a negative regulatory element of the Cpx signaling cascade) (13, 80), *dsbA* (disulfide bond catalyst) (14, 77), *ppiA* (peptidyl-prolyl cis-trans isomerase) (77), and *degP* (serine-threonine protease) (15, 77). Like their counterparts in *E. coli*, we confirmed that the expression of these four genes was modulated by the functional status of the Cpx pathway: it was reduced in the absence of CpxR but elevated in the absence of CpxA (Fig. 4). When T3SS gene transcription was analyzed, *yopE*, *yopH*, *yopK*, and *yopD* were the only ones displaying reduced transcription in the ΔcpxA null mutant (Fig. 4). This low expression could essentially be complemented by providing back a wild-type copy of inducible *cpxA* in trans. Significantly, these genes all encoded products that are among the last to be secreted. Generally, the transcription of all other tested T3SS genes was apparently unaffected by the status of the Cpx pathway.

To examine if reduced *yopE*, *yopH*, *yopK*, and *yopD* transcription was due to the effect of feedback inhibition in a ΔcpxA null mutant, we short-circuited this phenomenon by introducing a mutated *lerQ* allele into the ΔcpxA null mutant background (76). Significantly, in this ΔcpxA *lerQ*::Sp-Sm double mutant, transcription from these four gene promoters under T3S-permissive conditions was still notably lower (Fig. 4). This is evidence that the reduced transcription seen in CpxA-defective bacteria is more likely due to a direct effect of an altered Cpx pathway and not necessarily due to feedback inhibition caused by the loss of a competent T3SS. This suggests some action of the Cpx pathway on control regions of a specific subset of T3S genes. Moreover, it is interesting that the transcription of all other T3S genes assayed in the ΔcpxA null mutant was not visibly altered (Fig. 4). In particular, this occurred for *yscF* even though a significant loss of YscF production was visualized (Fig. 3). It is possible, therefore, that CpxA might also influence the levels of another subset of T3S components by an undisclosed posttranscriptional mechanism.

**Disruption of the intrinsic negative regulatory loop restores T3S.** It was a surprise to find that the reduced transcription seen in the ΔcpxA null mutant was also observed in an isogenic double mutant lacking the negative repressive element LcrQ. Since a loss of LcrQ elevates Yop synthesis and secretion even by bacteria grown under conditions that are not normally T3S permissive, we utilized the ΔcpxA *lerQ*::Sp-Sm double mutant to analyze levels of secreted Yops. Significantly, secretion levels were restored in the ΔcpxA *lerQ*::Sp-Sm double mutant similarly to the ΔcpxA parental strain (Fig. 2B). These observations...
are in accordance with the increased presence of the YscF, YscI, and LcrV structural components, all found to be reduced in the single mutant, in the bacterial envelope (Fig. 3). This suggests that the secretion defect observed in the ∆cpxA null mutant can be suppressed in bacteria lacking the negative regulatory loop that in turn permits consistent Yop synthesis. Presumably, these higher protein production levels increase the odds of assembling at least a few T3SSs that are functional, even when CpxA is lacking. An understanding of how this occurs, particularly when the transcription of a subset of T3S genes remains low, requires further investigation.

Removal of CpxA inhibits Yop effector translocation into eukaryotic cells. T3SSs are designed to translocate antihost effectors into infected eukaryotic cells. We wanted to determine if the reduced secretion of Yops into laboratory medium by the ∆cpxA null mutant also equated to a deficiency in Yop translocation. We performed a HeLa cell cytotoxicity assay with the various Y. pseudotuberculosis strains. This well-established assay qualitatively scores YopE cytotoxicity by the monitoring of cell monolayer morphology (83). YopE specifically induces cellular cytotoxicity via its GTPase-activating activity towards small G proteins involved in regulating actin-nucleating dynamics (5, 92). Despite secreting low levels of Yops in vitro, the ∆cpxA null mutant failed to induce any cytotoxicity towards target cells (Fig. 5). This was distinct from the complemented mutant, which induced a degree of cellular cytotoxicity similar to those seen for wild-type and ∆cpxA null bacteria. Significantly, the ∆hse5, ∆hseR, ∆hse4, and ∆hseB null mutants could still translocate the YopE cytotoxin normally (data not shown).

We considered that the ∆cpxA null mutant did not translocate YopE because it was poorly secreted (see Fig. 2A). This proved not to be the case, however, because a ∆lerQ cpxA double mutant, which secretes Yops abundantly (Fig. 2B), also poorly translocated YopE. This was due solely to a defect in CpxA, since ∆cpxA lerQ::Sp-Sm ectopically expressing CpxA efficiently restored full cytotoxicity (Fig. 5). Hence, it appears that CpxA may have at least two roles, one required to ensure the efficiency of Ysc-Yop T3S of proteins originating from inside bacteria and another that could modulate the timely translocation of this secreted protein cargo into the eukaryotic cell interior.

Cpx responsiveness is inhibitory to T3S. Our data imply that CpxA can function independently from the cognate response regulator CpxR. Perhaps CpxA can cross talk with another unidentified response regulator(s). At least in E. coli, however, no experimental support for this exists (94). A more plausible explanation concerns the levels of accumulated phosphorylated CpxR (CpxR-P). Normally, the phosphatase activity of CpxA keeps the level of CpxR-P low (81). However, in the absence of CpxA, CpxR can be phosphorylated by low-molecular-weight phosphodonors such as acetyl phosphate (13, 15). In this context, the presence of high levels of CpxR-P may attenuate Y. pseudotuberculosis T3S. To test this, we examined the T3S profile of the ∆cpxR null mutant expressing an IPTG-inducible wild-type copy of cpxR in trans. We compared this profile to that obtained from the parental strain and two control strains in which the ∆cpxR null mutant harbored only the empty vector or a derivative producing a mutant form, CpxR(D51A), which is unable to be phosphorylated. After IPTG induction, elevated levels of both wild-type CpxR and CpxR(D51A) notably reduced both the production and secretion of Yops even under T3S-permissive growth conditions (Fig. 2C). Furthermore, the overexpression of CpxR, but not the CpxR(D51A) variant, prevented the translocation of the YopE cytotoxin into HeLa cell monolayers (data not shown) (Fig. 5I). Because excessive copies of CpxR are toxic to bacteria (20), we also performed this analysis in the absence of IPTG induction. Significantly, low-level production of the CpxR(D51A) nonphosphorylated variant maintained a normal T3S profile, but similarly produced wild-type CpxR still had a negative impact on Yop secretion and synthesis (Fig. 2C).

While the latter strain does exhibit a moderate growth defect during routine culturing (data not shown), it is comparable to the ∆cpxA null mutant that is also thought to produce high levels of phosphorylated CpxR (13, 15). Thus, a tantalizing interpretation of these data is that it is the uncontrolled activation of the Cpx response leading to high levels of CpxR-P,
and not its removal, that compromises the pathogenicity of Y. pseudotuberculosis.

**DISCUSSION**

In this study, we have established that the Cpx two-component signal transduction pathway is important for T3S by Y. pseudotuberculosis, implicating it as an important regulatory circuit in *Yersinia* pathogenicity. This further underscores previous studies that described a role for CpxRA in the phase-variable expression and/or assembly of several adhesion organelles in pathogenic strains of *E. coli* (21, 34, 41, 45, 47, 68, 72) and for the cellular attachment, invasion, and intracellular survival of *Salmonella enterica* serovar Typhimurium (40). Interestingly, the latter effect might be mediated by influences on T3S by this bacterium (65), which has also been observed for *Shigella sonnei* (60, 66, 67).

It was anticipated that the Cpx pathway would serve a quality control function, overseeing the assembly of working T3SSs. This is based on members of the CpxR-P regulon functioning in the periplasm to assist in protein folding and degradation. These include peptidyl prolyl cis-trans isomerases (PPIases) (32), the disulfide bond catalyst DsbA (96), and the serine protease and chaperone DegP (HtrA) (73). In particular, DsbA is already known to help fold the T3S secretins in *Yersinia pestis* (43), *Shigella flexneri* (93), and *S. enterica* serovar Typhimurium (57) and for T3S gene expression in *Pseudomonas aeruginosa* (31). However, we are unable to demonstrate a role for DsbA in *Y. pseudotuberculosis* T3S (K. E. Carlsson, unpublished data). This is also the case for the various periplasmic PPIases (M. S. Francis, unpublished data). On the other hand, *Y. pseudotuberculosis* lacking DegP is affected in T3S, but this mutant is highly unstable (Carlsson, unpublished). Interestingly, DegP is responsible for degrading misfolded proteins in the periplasm during P-pilus assembly in *E. coli* (42, 45). Perhaps DegP assumes a similar role during T3S biogenesis in *Yersinia*. Nevertheless, our data herein indicate that Cpx pathway activation, resulting in accumulated CpxR-P, actually negatively impacts *Yersinia* T3S. These observations create a conundrum: why would an ECS-sensing pathway controlling the production of useful protein-folding and degradation factors be deleterious to the assembly of a complex multicomponent T3SS spanning the bacterial envelope? We will endeavor to address this intriguing connection in the future, initially by focusing on the effect of CpxR-P accumulation in *Yersinia* cytoplasm. For example, it would be beneficial to examine the effect of CpxR-P on T3S when the signal transduction cascade is still intact. This might be possible by overexpressing *nlpE*, encoding an outer membrane lipoprotein from *E. coli* (87).

It is interesting that the ΔcpxA null mutant exhibits two distinct regulatory effects. In a subset of genes (such as *yopE*, *yopH*, *yopK*, and *yopD*) that encode late-secreted substrates, both transcription and translation are reduced. Apparently, this effect is not a consequence of feedback inhibition (a functional T3SS is required for the expression of genes encoding secretion substrates) (11) because this same phenotype is still observed in an isogenic double mutant lacking the LcrQ-dependent negative regulatory loop that relieves feedback inhibition. However, another phenotype was observed for *yscF*, encoding the needle component, where transcription remained normal even though protein levels were significantly reduced. This indicates that at least YscF could also be controlled posttranscriptionally. The mechanism(s) underpinning these two distinct regulatory pathways remains unclear, although an undisclosed CpxA-dependent posttranscriptional regulatory pathway is an implicated feature of T3S control in *S. sonnei* (60).

In all likelihood, elevated CpxR-P levels negatively regulate T3S by *Yersinia*. In *E. coli*, CpxR-P is thought to recognize a putative consensus sequence, 5′-GTAAA(N)₅GTAAA-3′, located in target gene promoters (20). However, scanning the promoters of *Yersinia* T3S genes failed to reveal any obvious CpxR-P recognition motifs. Therefore, maybe CpxR-P does not directly engage T3S gene promoters. In *E. coli*, the CpxR-P regulon boasts over 100 different members. In this sense, CpxR-P might directly control the production of another factor(s) associated with the regulation of T3S genes. Apart from the above-mentioned periplasmic quality control factors (DsbA, DegP, and the PPIases), the CpxR-P regulon also affects ribosome assembly, the production of tRNAs, mRNA processing, small regulatory RNAs, RNA binding proteins, and chaperones (20). Thus, there is a plethora of factors that could potentially differentially influence individual components of the Ysc-Yop T3SS. To find out why gene subsets are regulated differently, determining their regulatory mechanisms and determining how they converge together to coordinate T3S are our future goals. It is also worth considering that the partial overlap between the Cpx and RpoE (σ²E) ECS-responsive pathways can mean that the loss of CpxA may alter the expression of σ²E-regulated genes (22). This might also have an impact on T3S in *Y. pseudotuberculosis*, given that σ²E is probably required.

The alternative sigma factor σ²E is proposed to modulate a core regulon responsible for the maintenance of lipopolysaccharide and porin biosynthesis conserved across a number of bacteria. In addition, another variable regulon apparently performs pathogenesis-related functions necessitated by the unique environmental niches inhabited by individual bacteria (82). The finding that ΔrpoE and ΔrseA null mutants were affected in T3S supports a pathogenicity role for σ²E in *Y. pseudotuberculosis*. In contrast, a loss of RseB was not seen to alter T3S. This is not surprising, judging from studies of *E. coli* (19). The ΔrseB null mutant still probably possesses RseA activity, suggesting that RseA has the ability to modulate σ²E-dependent effects on T3S independently of RseB. Given that rpoE is an essential gene in several bacteria (18, 35), which was reflected in the instability of our mutant described here, the ΔrseA null mutant is important since it will provide opportunities to dig deeper into the connections between σ²E and T3S. In this sense, it is also significant that survival of our ΔrpoE mutant likely occurs through the acquisition of second-site suppressor mutations (18). Interestingly, these mutations restore T3S, suggesting that they position in genes whose products also modulate T3S.

We were unable to assign a function for the BaeSR two-component phosphorelay system in *Yersinia* T3S in vitro or in the intoxication of target cells. This system has not yet been implicated in bacterial pathogenesis (79, 85), but a genome-wide analysis revealed that the overproduction of BaeR affected the expression of *E. coli* gene clusters associated mainly
with chemotaxis, flagellar biosynthesis, maltose transport, and multidrug efflux (69). Furthermore, genetic studies of \textit{E. coli} and \textit{S. enterica} indicate that BaeSR is important for resistance to antimicrobial compounds (3, 36, 39, 64) and chemical-induced ECS (78). It remains to be seen what physiological functions the BaeSR pathway modulates in \textit{Yersinia} and whether this impacts bacterial survival in the environment or during a host infection.

Finally, by removing the LcrQ negative regulatory element, we could create an isogenic \textit{Y. pseudotuberculosis} \(\Delta\text{cpxA}\) null mutant to consistently produce and secrete Yops. However, this strain was still significantly impaired in its ability to intoxicate infected cell monolayers with Yops, a defect that was complementable by the ectopic expression of CpxA. This indicates one of two possibilities: either T3S occurs differently in the presence of cells, such that the \(\Delta\text{cpxA}\) null mutant has a different effect in vivo, or the Cpx pathway affects additional processes required for T3S-dependent cytotoxicity. It is possible that the CpxR-P regulon of \textit{Yersinia} includes other loci that enable eukaryotic cell contact. In fact, \textit{Yersinia} strains lacking CpxA do associate poorly with cells, whereas contact by CpxR-defective bacteria is enhanced (our unpublished data). Furthermore, the modulation of cell contact in these strains is dependent on levels of the predominant \textit{Yersinia} adhesin, invasin (our unpublished data). These collective findings provide additional support for CpxRA responsiveness being an important elicitor of bacterial pathogenicity.

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