INTRODUCTION

Type III secretion systems (T3SSs) produced by assorted Gram-negative bacteria consist of ~20 different proteins that sequentially assemble into a syringe-like device that spans four layers of the cell envelope (the inner membrane, the periplasm, the peptidoglycan layer, and the outer membrane). Once fully assembled, bacteria utilize this structure to inject effector proteins into the eukaryotic host cells. Several of these proteins form two well-characterized, and independently assembled oligomeric ring structures—the outer membrane secretin ring and the inner membrane ring (Blocker et al., 2001; Burghout et al., 2004; Diepold et al., 2010; Kimbrough & Miller, 2000, 2002; Koster et al., 1997; Kubori et al., 2000; Ogino et al., 2006). Additionally, an inner membrane export apparatus and cytosolic sorting complex boast ~10 well-conserved proteins common to all flagella- and non-flagella T3SSs (Butan et al., 2019; Diepold et al., 2011; Fabiani et al., 2017; Johnson et al., 2019; Lara-Tejero et al., 2011; Wagner et al., 2010; Zhang et al., 2017). Their role
may involve substrate recognition, sorting, and energizing for substrate export. A needle appendage spans the membrane rings and extends beyond the bacterial surface (Broz et al., 2007; Marlovits et al., 2006; Wood et al., 2008). These components are listed with universal nomenclature in Supplementary Table S1.

The needle appendage is comprised of "early" substrates that represent the first set of components secreted by a functional T3SS (Zhang et al., 2017), and is the conduit through which unfolded substrates are exported to the outside (Dohlich et al., 2014; Radics et al., 2014). A second set composed of "middle" substrates are secreted and either assemble at the needle tip or sit distal to the needle tip to initiate pore formation in the eukaryotic cell membrane (Matei et al., 2011; Nauth et al., 2018; Park et al., 2018; Russo et al., 2019). Pore formation is required for the correct delivery into the eukaryotic cell interior of a third set of secreted components—the "late" effector proteins. Hence, substrate secretion might be temporally coordinated according to their individual function; "early" secreted substrates possess functions required first, 'middle' substrates function next, while the "late" secreted substrates are the translocated effectors (Dewoody et al., 2013; Diepold & Wagner, 2014; Osborne & Coombes, 2011).

Three clinically relevant pathogenic Yersinia species have served as models to study type III secretion, namely Y. pestis, Y. pseudotuberculosis, and Y. enterocolitica. All three species possess the ability to colonize and establish a disease state in humans. This ability is in part due to all harboring a common virulence plasmid encodes a highly homologous Ysc (Yersinia secretion)—Yop (Yersinia outer protein) T3SS (Cornelis et al., 1998). The function of the Ysc-Yop T3SS is to target the cytosol of eukaryotic cells a small but potent effector arsenal. Consequently, Yersinia can replicate extracellularly because the internalized effectors modulate phagocytic, apoptotic, and proinflammatory cellular signaling mechanisms (Grabowski et al., 2017; Pha & Navarro, 2016; Philip et al., 2016).

In Yersinia Ysc-Yop T3S, there is experimental support for a hierarchical substrate secretion and effector targeting model (Mahdavi et al., 2014; Thorslund et al., 2011). For example, Diepold and colleagues identified that a tripartite complex of YscX-YscY-ScTV is involved in the control of "early" substrate export (Diepold et al., 2012). ScTV is a core component of all T3SSs and is a relatively fixed inner membrane component of the export apparatus (Diepold et al., 2015). On the other hand, the components YscX and YscY are unique to the Ysc-Yop T3SS phylogenetic clade that is also produced by bacteria including species of Pseudomonas, Aeromonas, Photobacterium, and Vibrio (Gazi et al., 2012; Gurung et al., 2018). YscX is believed to be a T3S substrate chaperoned in the cytoplasm by YscY (Day & Plano, 2000), and both are essential for T3S activity (Day & Plano, 2000; Iriarte & Cornelis, 1999). One or more functions of native YscX and YscY appears to be evolutionary optimized to contribute unique and essential information for the physical assembly of the Ysc-Yop T3SS by pathogenic Yersinia sp. because related proteins failed to restore T3SS function in Yersinia mutants lacking yscX and/or yscY. This was despite a reciprocal YscX–YscY and a YscX–YscY–ScTV interactions between member proteins from different bacteria (Gurung et al., 2018).

The initial discovery of YscX and its important role in Ysc-Yop T3S activity occurred over 20 years ago (Day & Plano, 2000; Iriarte & Cornelis, 1999). Nonetheless, new functional insight has rarely occurred since these seminal publications. This present study addresses this critical knowledge gap by building on the previous findings of Diepold and colleagues (Diepold et al., 2012). In this regard, we hypothesize that YscX secretion is important for T3S activity. Indeed, we demonstrate herein that YscX is a true T3S substrate containing a bona fide secretion recognition signal located at the N-terminus. Moreover, disruption of this N-terminal secretor domain creates a general defect in Ysc-Yop T3S, and an effect of this is improper localization of the needle components SctI and ScTF to the bacterial periphery. Altogether, our study provides new insights into the hierarchal targeting and export of early substrates during assembly of the Ysc-Yop T3SS in Y. pseudotuberculosis. These findings can be applicable to other bacteria such as species within the Pseudomonas, Aeromonas, Photobacterium, and Vibrio genera that are producing a system belonging to the Ysc-T3SS clade.

## RESULTS

### 2.1 The N-terminal segment of YscX is an effective T3S export signal

YscX is an apparent early Ysc-Yop T3SS substrate (Day & Plano, 2000; Diepold et al., 2012). However, the secretion signal that mediates its secretion has not been identified. Since T3S export signals of substrates are typically located at the N-terminus (Löwer & Schneider, 2009; McDermott et al., 2011; Wang et al., 2013), we examined if the N-terminal residues of YscX could function as an independent secretion signal to promote the T3S of a β-lactamase reporter. Capitalizing on a reporter system established for studies of the SctB N-terminus (Amer et al., 2011), a series of translational fusions between the 5' end of yscX (including the native Shine-Dalgarno (SD) sequence) and a promoterless bla allele lacking a secretion signal were generated. Plasmids were maintained in trans, and expression of each fusion was controlled by an IPTG-inducible promoter. We assayed by Western blotting total protein fractions recovered from pelleted bacteria as well as protein secreted to the extracellular medium that was collected from trichloroacetic acid-precipitated bacteria-cleared culture supernatants. Sequences of yscX encoding all (FULL) or the first 50, 25, 20, 15, 10 and 5 amino acids were all sufficient to produce generous levels of β-lactamase, a portion of which was secreted (Figure 1a). This secretion was dependent on the presence of a functional T3SS, because an isogenic mutant lacking ScTV, an integral T3SS component, failed to secrete these fusions (Figure 1a). Critically, the secretion efficiency peaked at YscX25-Bla (circa 25%) and
descended to YscX10-Bla (circa 3%) (Figure 1b). Additionally, YscX5-Bla fusion secretion occurred at an efficiency of <1%, and visualization required overexposure of the immunoblot image (Figure S1a). On the other hand, the smallest yscX fusion of 1 amino acids did not visibly promote secretion of the reporter (Figures 1 and S1a). Moreover, we also noted poor steady-state levels of accumulated YscX1-Bla fusion compared to all others (Figure 1a). However, this reduction in accumulation could not be explained by a simple increase in protein turnover, for this smaller fusion was stable (Figure S1b). On the other hand, longer fusions beginning with YscX20-Bla were increasingly less stable, and this instability was most dramatic for the largest YscX50-Bla and YscXFULL-Bla fusions (Figure S1b). This instability would explain the poor secretion efficiency of these larger fusions (Figure 1). Taken altogether, a secretion signal of YscX must minimally consist of 10 or more N-terminal residues. Furthermore, translation efficiency and product stability are two factors that visibly influence the levels of detectable Bla fusions.

2.2 | Deletions within the YscX export signal disrupt its secretion

Given that the N-terminus of YscX is an effective T3S export signal, we wanted to address what impact deleting this region has on YscX synthesis and secretion. Three in-frame deletion mutations in the 5-prime terminus of the yscX allele were introduced in cis on the virulence plasmid to generate mutant bacteria encoding the variants YscXΔ3-7, YscXΔ8-12, and YscXΔ13-22. To visualize YscX by Western blot, a rabbit polyclonal anti-YscX antiserum was raised against the predicted immunogenic peptide (NH2- ) CLHRAQDYRRELDTL(- CONH2) encompassing the residues 70 to 83 of YscX. Its utility in Western blotting was first checked in protein samples derived from Yersinia strains that either lacked YscX or expressed different YscX variants (Figure S2). With this antibody, we first analyzed the susceptibility of each YscX variant to endogenous proteases as an indicator of their ability to form native tertiary folds. Generally, low levels of detectable
accumulated wild type YscX remained similar over time, suggesting it to be stable as long as the cognate T3S chaperone YscY was always present (Figure S3). This corroborates earlier findings of the stabilizing effect of YscY on pre-secretory pools of YscX (Day & Plano, 2000). Critically, there were no obvious differences in stability of accumulated YscXΔ3-7, YscXΔ8-12, and YscXΔ13-22 (Figure S3).

Next, we compared the three YscX mutants to parent bacteria for the ability to secrete YscX during growth in T3S-restrictive (BHI broth plus Ca²⁺ ions) and T3S-permissive conditions (BHI broth minus Ca²⁺ ions). Interestingly, we could only detect the steady-state accumulation of native YscX and very low amounts of YscXΔ3-7 associated with the bacterial pellet (Figure 2a, upper panel). Native YscX and YscXΔ3-7 were secreted free into the culture medium under

**Figure 2** Analysis of T3S activity by *Y. pseudotuberculosis* producing YscX variants devoid of N-terminal sequence. Overnight cultures of *Y. pseudotuberculosis* that either harbored an intact lcrQ allele (a and c; LcrQ⁺) or lacked the lcrQ allele (b and d; LcrQ⁻) were sub-cultured into BHI medium in the presence (+) or absence (−) of calcium ions at 26°C for 1 h and at 37°C for 3 h. Protein in the bacterial pellet (Synthesis) and either protein recovered from trichloroacetic acid-precipitated bacterial-free culture supernatant (Secretion—YscX) or protein directly sampled supernatant (Secretion—Yops) were collected and solubilized in sample buffer. Cross-linking of surface localized SctF (E) was performed by growing *Yersinia* strains in non-permissive T3S media (plus Ca²⁺). Where indicated (+), the membrane-impermeable chemical cross-linker BS³ was added to the bacteria, subsequently quenched with Tris–HCl and bacteria pellets were solubilized in sample buffer. Protein samples were then fractionated by 15% acrylamide (for YscX and SctF) or 12% acrylamide (for Yops) SDS–PAGE, wet-blotted onto PDVF membrane and then detected using rabbit polyclonal anti-YscX, anti-SctE, anti-SctB, anti-YopE or immune-absorbed monospecific anti-SctF antibodies. Panels A and C: Parental (YscXwt), YPIII/pIB102; ΔsctU, YPIII/pIB75; ΔyscX, YPIII/pIB880; ΔyscY, YPIII/pIB890; yscXΔ3-7, YPIII/pIB88002; yscXΔ8-12, YPIII/pIB88003; yscXΔ13-22, YPIII/pIB88004. Lanes B and D: ΔlcrQ null mutant (YscXwt), YPIII/pIB26; ΔsctU, ΔlcrQ, YPIII/pIB75-26; ΔyscX, ΔlcrQ, YPIII/pIB880-26; ΔyscY, ΔlcrQ, YPIII/pIB890-26; yscXΔ3-7, ΔlcrQ, YPIII/pIB88002-26; yscXΔ8-12, ΔlcrQ, YPIII/pIB88003-26, yscXΔ13-22, ΔlcrQ, YPIII/pIB88004-26. Approximate molecular mass values shown in parentheses were deduced from primary amino acid sequences. The arrow indicates a faint protein band representing YscXΔ3-7 that is only clearly visible with prolonged exposure times. Panel E: Parental (YscXwt), YPIII/pIB102; ΔsctU, YPIII/pIB75; ΔyscX, YPIII/pIB880; ΔsctF null mutant, YPIII/pIB202; yscXΔ3-7, YPIII/pIB88002; yscXΔ8-12, YPIII/pIB88003, yscXΔ13-22, YPIII/pIB88004. The predicted molecular mass of monomeric SctF is given in parenthesis, while approximate sizes of protein molecular weight standards are given to the left.
these assay conditions (Figure 2a, lower panel and Figure S2a, lower panel). In the ΔsctU null mutant control strain that is T3SS-defective, and in the ΔyscY null mutant control strain that is devoid of T3S chaperone activity toward YscX (and is also T3S-defective), production and secretion of YscX were not detectable (Figure 2a).

A dramatic reduction in steady-state levels of YscXΔ3-7, YscXΔ8-12, and YscXΔ13-22 made an analysis of their secretion in isogenic strains inconclusive. A deletion of lcrQ—encoding for a negative regulatory element of Yops synthesis—into the YscXΔ3-7, YscXΔ8-12, and YscXΔ13-22 producing bacteria circumvented this. LcrQ (genetic equivalent YscM1 and YscM2 in Y. enterocolitica) is an anti-activator critical to feedback inhibit the Ysc- Yop T3SS. Removal of lcrQ leads to de-repression and constitutive production of Yops synthesis even in T3SS-defective bacteria (Li et al., 2014b; Petersson et al., 1996; Rimpiläinen et al., 1992). Accordingly, analysis of the lcrQ mutant for YscX synthesis and secretion displayed a typically de-regulated phenotype, constitutively producing YscX during growth in both non-inductive (BHI plus Ca²⁺ ions) and inductive (BHI minus Ca²⁺ ions) growth conditions, while still only secreting YscX in inductive growth conditions (Figure 2b). Additionally, we could now easily detect YscX synthesis in all strains except for the ΔyscX and ΔyscY null mutants, yet only parental Yersinia as well as the YscXΔ3-7 producing strain secreted YscX free into the surrounding medium (Figure 2b). Employing a deletion of lcrQ to de-repress Yop synthesis improved the detection of accumulated bacterial-associated pools of YscX variants and confirmed a lack of secretion of YscXΔ8-12 and YscXΔ13-22, but not of YscXΔ3-7. Hence, stably produced YscX is a bona fide T3S substrate of Y. pseudotuberculosis, and this secretion
FIGURE 3 Type III secretion activity from bacteria expressing YscX chimeras with a reciprocally exchanged N-terminal secretion signal. Expression and secretion of chimeric substrates with a reciprocally exchanged N-terminal secretion signal. Overnight cultures were subcultured into BHI broth lacking calcium and then grown at 26°C for 1 h and then at 37°C for 3 h. Expression and secretion of YscX variants (a) were determined in bacterial pellet (a—upper panel) and TCA precipitated proteins freely released into culture supernatants (a—lower panel) following growth of Yersinia in secretion-permissive conditions (−Ca²⁺). Protein samples were fractionated by a 12% acrylamide SDS–PAGE and then transferred onto a PVDF membrane support for immune-detection with rabbit polyclonal anti-YscX antiserum. Yops (b) associated with total fraction (proteins within intact bacteria and secreted into the culture medium, b—upper panel) and or with supernatant (secreted free to the extracellular medium, b—lower panel) were analyzed following growth of Yersinia in BHI medium in the presence (+) or absence (−) of calcium ions. Protein samples were fractionated by a 12% acrylamide SDS–PAGE and then transferred onto a PVDF membrane for immune-detection with rabbit polyclonal antisera to SctE, SctB and YopE. Surface localized SctF were cross-linked (c) following growth in non-permissive T3S media (plus Ca²⁺). The membrane-impermeable chemical cross-linker BS² was added where indicated (+) and subsequently quenched with Tris–HCl. Bacteria pellets derived protein samples were then fractionated by 15% acrylamide SDS–PAGE, wet-transferred to PVDF and detected with immune-absorbed monospecific anti-SctF antiserum. Synthesis fractions (upper panels) represent bacterial pellet. Secretion fractions (lower panels) signify proteins freely released into the culture supernant. Panel a and b: Parental (YscXΔ9), YPIII/pIB1102; ΔyscX null mutant, YPIII/pIB8800; ΔyscY null mutant, YPIII/pIB8800; ΔsctU null mutant, YPIII/pIB75; SctBΔ2-15-YscX, YPIII/pIB88001; YopEΔ2-15-YscX, YPIII/pIB88007; SctFΔ2-15-YscX, YPIII/pIB88006; LcrQΔ2-15-YscX, YPIII/pIB88005; SctUΔ2-15-YscX, YPIII/pIB88008; YscUΔ2-15-YscX, YPIII/pIB88009; YscXΔ2-15-SctB, YPIII/pIB62520; YscΔ2-15-YopE, YPIII/pIB581; YscXΔ2-15-SctF, YPIII/pIB20202; YscXΔ2-15-LcrQ, YPIII/pIB2602. Approximate molecular mass of YscX is shown in parenthesis and was deduced from primary amino acid sequence. Panel c: Parental (YscXΔ9), YPIII/pIB1102; ΔsctU, YPIII/pIB75; ΔyscX, YPIII/pIB8800; ΔsctU null mutant, SctBΔ2-15-YscX, YPIII/pIB88001; SctFΔ2-15-YscX, YPIII/pIB88006; YscUΔ2-15-SctF, YPIII/pIB20202, YscXΔ2-15-SctB, YPIII/pIB62520. The predicted molecular mass of monomeric SctF is given in parenthesis, while approximate sizes of protein molecular weight standards are given to the left.

2.3 The ability to secrete Yops correlates with YscX secretion

YscX plays a role in promoting T3S (Day & Plano, 2000; Diepold et al., 2012; Iriarte & Cornelis, 1999). We wondered whether YscX secretion is important for this process. Using the strains producing a non-secreted form of YscX i.e.: the variants YscXΔ8-12 or YscXΔ13-22 allowed us to investigate this point. We used the membrane-impermeable cross-linker BS² to examine whether the early substrate—the needle component SctF—could polymerize at the surface of YscXΔ8-12, YscXΔ13-22, and YscXΔ13-22 producing bacteria. We detected monomeric SctF located in the bacterial cytoplasm, and therefore protected from the membrane impermeable BS² cross-linker, in samples derived from these bacteria as well as in parental bacteria and the T3SS-defective full-length yscX and sctU deletion mutants (Figure 2e). Only in parental bacteria and YscXΔ8-12 producing bacteria, the cross-linking agent BS² cross-linked secreted SctF, indicative of higher order structures representative of the T3S needle (Figure 2e). We visualized no cross-linked surface-located SctF in the T3SS-defective full-length yscX and sctU deletion mutants or in bacteria producing YscXΔ8-12 or YscXΔ13-22 (Figure 2e). The sctF null mutant served as a specificity control to demonstrate that the monospecific anti-SctF antibodies did not cross-react with any other Yersinia protein (Figure 2e). Altogether, these data demonstrate that a failure to export YscXΔ8-12 or YscXΔ13-22 coincided with an inability of these bacteria to assemble a complete T3SS as measured by the absence of a surface appendage composed of polymerized SctF.

We anticipated that strains lacking surface polymerized SctF had abolished T3SS activity. To examine this point, samples derived from YscXΔ8-12, YscXΔ13-22, and YscXΔ13-22 producing bacteria were subject to Western blotting to probe for the middle substrates, SctB and SctE, and the late substrate YopE (commonly termed as Yops). As already discussed, Y. pseudotuberculosis lacking yscX, yscY, or sctU are feedback-inhibited for Yops production (Figure 2c) (Bröms et al., 2005; Lavander et al., 2002), but the introduction into these strains of an lcrQ deletion to avoid the production of this repressor element restored partial Yops synthesis in both T3S-restrictive (high Ca²⁺) and T3S-permissive growth media (low Ca²⁺) (Figure 2d). Not surprisingly, however, these bacteria could not secrete Yops regardless of whether LcrQ was present (Figure 2c) or absent (Figure 2d). Significantly, this LcrQ-dependent Yop synthesis and secretion profile was mirrored in the mutant bacteria producing YscXΔ8-12 or YscXΔ13-22(Figure 2c, d). In contrast, LcrQ+ bacteria producing native YscX or the stable and secreted YscXΔ8-7 variant still actively synthesized Yops in a controlled low Ca²⁺-dependent manner (Figure 2c), and this control was lost upon removal of lcrQ (Figure 2d). Critically, these bacteria maintain tight control of this secretion process because it occurred only during growth in T3S-permissive conditions.

These Yop profiles corroborated with the low calcium growth characteristics of these bacteria. Consistent with being repressed for Yop synthesis, the full-length mutations of sctU, yscY, and yscX all resulted in LcrQ−-bacteria exhibiting a calcium independent (Cl) growth phenotype, being able to grow at 37°C regardless of the calcium concentration in the growth medium (Figure S4). Similarly, a Cl growth phenotype was also observed for LcrQ−-bacteria producing either YscXΔ8-12 or YscXΔ13-22 that were also unable to synthesize Yops (Figure S4). In contrast, both LcrQ+ parent bacteria and LcrQ−-mutant bacteria producing the stable and secreted YscXΔ8-7 variant required calcium for growth at 37°C (Figure S4). Moreover, deletion of the lcrQ allele from all strains prompted a temperature sensitive (TS) growth phenotype (Figure S4). This
is consistent with the lcrQ deletion effect being epistatic over any mutations that compromise T3SS assembly and that culminate in bacteria constitutively producing Yops regardless of Ca\(^{2+}\) concentration.

Taken together, bacteria that produce and secrete YscX\(_{2-15}\) still maintain Ysc-Yop T3SS assembly, Yops secretion, and control, whereas these features are lost in bacteria that can produce but not secrete YscX\(_{2-15}\) and YscX\(_{2-15}\). Hence, the ability to secrete any T3SS substrate (early, middle, and late) correlates with YscX secretion. The data suggest also that the N-terminal YscX secretor domain is involved in dual functions, having roles in both YscX secretion and T3SS assembly. Moreover, the first 3 to 7 residues of the YscX N-terminus are dispensable for both YscX secretion and T3SS assembly and activity.

### 2.4 T3SS assembly requires an untainted YscX N-terminus

Our N-terminal deletion analysis identified a requirement for the YscX secretor domain in Ysc-Yop T3SS assembly and activity, which corroborates earlier observations that YscX is needed for needle assembly and to orchestrate secretion of ‘middle’ translocator and ‘late’ effector substrates (Diepold et al., 2012). To investigate the possibility to uncouple YscX secretion from T3SS assembly, we took advantage of the fact that the reciprocity of secretion signals among the Ysc-Yop T3SS substrates is well-known (Amer et al., 2011; Sorg et al., 2007). We created a small series of in cis-encoded YscX chimeric variants that have had the native N-terminal codons 2 to 15 exchanged with a variety of functional secretion signals from the early substrates SctF, the chimera was termed (SctF\(_{2-15}\)-YscX) and LcrQ (LcrQ\(_{2-15}\)-YscX), the middle substrate SctB (SctB\(_{2-15}\)-YscX) and the late substrate YopE (YopE\(_{2-15}\)-YscX). Additionally, we generated two chimeric YscX substrates that contain codons 2 to 15 derived from the genetically analogous proteins PscX from *Pseudomonas aeruginosa* which is considered a non-secreted substrate (PscX\(_{2-15}\)-YscX) (Bröms et al., 2005; Gurung et al., 2018; Yang et al., 2007) and SctX from *Photorhabdus luminescens* (SctX\(_{2-15}\)-YscX) (Duchaud et al., 2003; Gurung et al., 2018). As controls, we generated reciprocal chimeric substrates in SctB, YopE, SctF, and LcrQ by substituting the 2–15 N-terminal codons with equivalent generated reciprocal chimeric substrates in SctB, YopE, SctF, and LcrQ respectively. These control strains produce recombinant SctB, YopE, SctF, and LcrQ but allow detection of native YscX.

We grew bacteria in BHI broth in T3S permissive (minus Ca\(^{2+}\)) and T3S restrictive (plus Ca\(^{2+}\)) growth conditions (Figure 3a, upper panel). Unsurprisingly, bacteria producing YscX\(_{2-15}\)-SctF could not support Yops production and secretion (Figure 3b). In parallel, we also looked for SctF secretion and its ability to polymerize at the bacterial surface to form the needle appendage—the final step in the assembly of an active Ysc-Yop T3SS. In our assay, the non-membrane permeable chemical cross-linker BS\(^{3}\) supported SctF polymerization. Given the phenocopy existing among all six N-terminal YscX chimaera-producing bacteria, we examined SctF polymerization only in the representative bacteria producing SctB\(_{2-15}\)-YscX or SctF\(_{2-15}\)-YscX. Although we detected non-membrane SctF located in the bacterial cytoplasm (and protected from the membrane impermeable BS\(^{3}\) cross-linker) in bacteria producing SctB\(_{2-15}\)-YscX or SctF\(_{2-15}\)-YscX cross-linked surface-located SctF was absent (Figure 3c). Hence, inserting a heterologous N-terminal signal sequence was not sufficient to maintain YscX function.

To rule out the possibility that defective SctF surface polymerization and T3SS activity was actually due to the failed secretion ability of the YscX chimera, we analyzed YscX secretion uncoupled from YscX-dependent T3SS assembly. This analysis was performed using our reporter assay, where we again translationally fused DNA segments encoding the first 25 residues of SctF\(_{2-15}\)-YscX, LcrQ\(_{2-15}\)-YscX, SctB\(_{2-15}\)-YscX, YopE\(_{2-15}\)-YscX, PscX\(_{2-15}\)-YscX, and SctX\(_{2-15}\)-YscX (including the native SD sequence) to the *bla* allele whose expression is controlled by an IPTG-inducible promoter. Except for two variants—SctF\(_{2-15}\)-YscX and PscX\(_{2-15}\)-YscX—all others produced recombinant β-lactamase in sufficient quantities to allow secretion by parental bacteria to be detected (Figure S5). Once again, deletion of sctU encoding a component of the
T3SS completely abrogated this secretion (Figure S5). Levels of the SctF_3-7-YscX and PscX_3-7-YscX products were simply too low to determine if they could be secreted. Thus, with two exceptions it is apparent that the heterologous secretion signals are active and maintain the ability to support the T3S of recombinant substrates, such as chimeric YscX.

Hence, YscX chimeras that differ from native YscX only at the N-terminus, by virtue of an exchanged type III recognition and secretion signal, abolished any aspect of T3SS export, including SctF surface polymerization (see Figure 3c), the ability to assemble a competent T3SS able to secrete middle and late substrates such as SctB, SctE, and YopE (see Figure 3b). Accordingly, we conclude that some rudimentary aspects of T3S must rely upon non-redundant peptide sequence in the extreme YscX N-terminus. Given the complexity of T3SS biogenesis and function, this aspect is expected to involve protein interactions with this region of YscX.

2.5 | Interplay between YscX and the T3S chaperone YscY

YscY stabilizes YscX, so the YscY-YscX protein complex is obviously important for T3SS activity (Day & Plano, 2000; Iriarte & Cornelis, 1999). Moreover, T3SS chaperones often engage with their cognate substrates through a chaperone binding domain that locates immediately adjacent the N-terminal secretion signal (Francis, 2010). Thus, we next investigated if YscX mutants defective in T3SS assembly and function have lost an ability to interact with YscY. In our hands, the yeast two-hybrid system has proven useful in demonstrating a YscX–YscY protein–protein interaction (Bröms et al., 2005). We, therefore, generated GAL4 activation domain (AD) fusions to yscX allelic variants in the vector pGADT7. These were co-transformed along with the pGBK7 derivative containing yscY fused to the GAL4 DNA binding domain (DBD) into the Saccharomyces cerevisiae reporter strain AH109 (James et al., 1996). We confirmed the stable production in yeast of all recombinant variants (Figure S6). Next, we judged a protein–protein interaction according to the ability of the host yeast strain to grow on minimal agar medium lacking either adenine or histidine. A GAL4 AD fusion to wild type YscX and the three GAL4 AD fusions to YscX_Δ3-7, YscX_Δ8-12, and YscX_Δ13-22 could all still bind to GAL4 DBD fused to native YscY binding—as measured by yeast growth—although the degree of YscX_Δ8-12-YscY interaction was comparatively inferior (Table 1 and Figure S7). Regardless, this degree of interaction with chaperone was sufficient to stabilize pre-secretory pools of all three YscX variants (Figure S3). Given that YscX stability depends on YscY (Day & Plano, 2000; Gurung et al., 2018; Iriarte & Cornelis, 1999), these data reflect that the YscX–YscY complex remains intact in the YscX mutants of Yersinia.

2.6 | A core element of Ysc-Yop T3SS assembly, SctV, remains a target of defective YscX

Based upon a previous report (Diepold et al., 2012), a tripartite complex forms between YscX–YscY and the integral inner membrane protein SctV, a core component of all T3SSs. This tripartite complex may contribute to establish a competent T3SS by coordinating early substrate secretion (Diepold et al., 2012). Hence, we wondered whether manipulation of the YscX N-terminus violates YscX–YscY binding to SctV. A yeast three-hybrid assay was established to detect this because both YscX and YscY are required simultaneously for complexing with SctV. A soluble variant of SctV containing only amino acid residues 322 through to 704 was cloned into the plasmid pGADT7 creating a fusion to the C-terminus of the GAL4 activation domain. We also cloned a full-length parental yscY allele into the MCSII site in the plasmid pBRIDGE creating a fusion with the GAL4 DNA binding domain. In this vector, the yscY allelic variants were then cloned under the control of a methionine repressible promoter (MCSII). Tripartite interactions were determined followed by growth on histidine lacking agar media in the presence or absence of methionine or by β-galactosidase measurements after growth in equivalent liquid media replete with histidine. We observed a specific interaction with SctV only in the presence of both YscY and YscX proteins (Table 2 and Figure S8). Every modified YscX variants could still engage with SctV so long as YscY was also present (Table 2 and Figure S8). As determined by interactions in yeast, vivid defects in Ysc-Yop T3SS induced by manipulation of the YscX N-terminus is not due to an inability of these mutants to form a tripartite complex with YscY and SctV. Hence, YscX function must involve another critical molecular target.

2.7 | Manipulation of YscX N-terminus affects localization of Sctl and SctF

Disruption of the YscX secretory domain caused a general secretion defect. We wanted to understand why this defect occurs. The
assembly of T3S injectisome is hierarchal, such that the deployment of cell-proximal components occurs before the more distal segments. We wondered if the N-terminus of YscX is required for an aspect of this injectisome assembly process. We first performed Western immunoblotting against different components of T3SS contained within standardized total protein samples (a mix of protein associated with bacteria and the culture supernatant) derived from Y. pseudotuberculosis associated with bacteria and the culture supernatant) derived from Western immunoblotting against different components of T3SS assembly checkpoint control. This assay allowed a quick appraisal for altered production of any structural T3SS component in the yscX, lcrQ mutant backgrounds. This necessitated the production of specific antibodies against purified components of the membrane spanning rings (SctC, SctD, and SctJ), the inner membrane export apparatus (SctV and SctU), and the cytoplasmic components (SctK, SctN, and SctO) (Table S1). In the parental strain, bands were detected representing a protein with an apparent molecular mass of ~60 kDa with anti-SctC antibody, ~42 kDa with anti-SctD antibody, ~25 kDa with anti-SctJ antibody, ~70 kDa with anti-SctV antibody, ~25 kDa with anti-SctK, ~45 kDa with anti-SctN antibody, and ~19 kDa with anti-SctO antibody (Figure 4a), which is consistent with the observations of others (Blaylock et al., 2006; Gauthier & Finlay, 2003; Koster et al., 1997; Li et al., 2014b; Mukerjea & Ghosh, 2013; Silva-Herzog et al., 2008; Soto et al., 2017). All these bands were absent in the Y. pseudotuberculosis YPIII strain lacking the T3SS encoded virulence plasmid, which confirms the antibody specificity (Figure 4a). Importantly, we could not identify any dramatic deviations between these expression profiles and those derived from the N-terminal YscX mutants (Figure 4a).

Since basal body component production in YscX mutants is not affected, we next investigated if these structures are competent for secretion of “early” substrates such as the SctI adapter/washer and SctF needle filament (Torres-Vargas et al., 2019; Zilkenat et al., 2016). The parental and YscXΔ3–7 strains produced and secreted comparable amount of SctI and SctF consistent with a functional T3SS (Figure 4b). While secretion of SctF was calcium-dependent, it was interesting to note SctI secretion irrespective of calcium concentration as reported in a previous study (Figure 4b) (Wood et al., 2008). We also observed reduced accumulation of Sctl and SctF in mutants lacking YscX and YscY compared to parental Yersinia and the mutant producing YscXΔ3–7 (Figure 4b). The lower accumulation of Sctl and SctF was consistently mimicked by Yersinia strains producing YscXΔ8–12 and YscXΔ13–22. Moreover, this pool of Sctl and SctF was

### TABLE 2 Protein–protein interactions in the yeast three-hybrid assay

| Yeast three-hybrid constructs | Derivative of pBridge | Derivative of pGADT7 | Interaction |
|-----------------------------|----------------------|---------------------|------------|
| Plasmid | Binding domain (MCSI) | MCSI | Activation domain | HIS3 | β-Galactosidase activity (LacZ) [fold induction*]| |
| pBRIDGE | none | none | pGADT7 | – | 0.95 ± 0.15 |
| pBRIDGE | none | none | pJMG043 (SctV322–704) | – | 0.79 ± 0.12 |
| pMF442 | YscY* | none | pGADT7 | – | 0.808 ± 0.17 |
| pMF442 | YscY* | none | pJMG043 (SctV322–704) | – | 0.836 ± 0.13 |
| pJMG070 | YscY* | YscX* | pGADT7 | – | 0.742 ± 0.13 |
| pJMG070 | YscY* | YscX* | pJMG043 (SctV322–704) | +++ | 234.8 ± 29.39 [281] |
| pJMG071 | YscY* | SctF2–15 YscX* | pJMG043 (SctV322–704) | +++ | 237 ± 37.58 [283] |
| pJMG072 | YscY* | LcrQ2–15 YscX* | pJMG043 (SctV322–704) | +++ | 196.6 ± 45.53 [235] |
| pJMG073 | YscY* | SctB2–15 YscX* | pJMG043 (SctV322–704) | +++ | 172.8 ± 5.35 [207] |
| pJMG074 | YscY* | YopE2–15 YscX* | pJMG043 (SctV322–704) | +++ | 128.8 ± 9.07 [154] |
| pJMG075 | YscY* | PscX2–15 YscX* | pJMG043 (SctV322–704) | +++ | 136.2 ± 9.15 [163] |
| pJMG076 | YscY* | SctX2–15 YscX* | pJMG043 (SctV322–704) | +++ | 162.2 ± 22.75 [194] |
| pJMG077 | YscY* | YscXΔ3–7* | pJMG043 (SctV322–704) | +++ | 131.8 ± 9.92 [158] |
| pJMG078 | YscY* | YscXΔ18–12* | pJMG043 (SctV322–704) | +++ | 149.4 ± 23.86 [179] |
| pJMG079 | YscY* | YscXΔ13–22* | pJMG043 (SctV322–704) | +++ | 184.6 ± 38.00 [221] |

Notes: lacZ is a reporter gene in S. cerevisiae Y190 and HIS3 is a reporter gene in S. cerevisiae HF7c. Activation of these genes imply an interaction between interactive protein partners fused to the binding domain (MCSI on pBRIDGE) and the activation domain on pGADT7. Expression of the gene cloned into multiple cloning site II (MCSII) is expressed in the absence of methionine. β-Galactosidase activity is recorded in Miller units and is the mean of at least five independent measurements from liquid cultures of logarithmic yeast cells. HIS3* represents strong growth (+++) to no growth (--) on minimal medium devoid of histidine and methionine, recorded after day 4. The standard error of the mean was calculated using Excel software (Microsoft Office suite).

*The relative increase in β-Galactosidase activity brought about by the methionine-dependent expression of the yscX allele from MCSII. Baseline β-Galactosidase activity is taken from yeast harboring the plasmids pMF442 and pJMG043.
not secreted (Figure 4b) by the non-functional yscX mutants, despite all strains producing equivalent levels of other core structural T3SS components (Figure 4a). This corroborated with the non-secreting control strain lacking SctU (Figure 4b). Hence, a general secretion defect is associated with a specific defect in SctI and SctF synthesis and secretion.

As this might indicate a disordered assembly of the T3SS where one or more individual T3S components exhibit asymmetric distribution in different sub-cellular compartments. To investigate this, sub-cellular localization of different Ysc components was analyzed in fractions derived from the lcrQ mutant with intact YscX (parental background) or producing the variants YscX_∆3−7, YscX_∆8−12, YscX_∆13−22. Analyzed sub-cellular fractions representing the cytoplasm, inner membrane, periplasm and outer membrane fractions were interrogated by Western blotting using our battery of specific antibodies to individual Ysc/Sct proteins. We first confirmed fraction purity using rabbit antibodies to proteins known to be enriched in the outer membrane (OmpA) (Confer & Ayalew, 2013), periplasm (SurA) (Mas et al., 2019), inner membrane (FtsH) (Bittner et al., 2017), and the cytoplasm (H-NS) (Grainger, 2016) (Figure S9). Next, we observed an enrichment of SctC in the outer membrane fractions, SctJ lipoprotein in the periplasmic fraction, and an integral inner membrane protein SctV in the inner membrane fractions (Figure 5a), which corroborated earlier studies (Gauthier & Finlay, 2003; Silva-Herzog et al., 2008). We also detected SctD predominately in the inner membrane fraction (Figure 5a), consistent with it being a component of the inner ring structure (Hu et al., 2017; Lountos et al., 2012).

We also observed low levels of all these proteins in the cytoplasmic fraction confirming consistent expression profiles across all strains (Figure 5a). In addition, we observed the cytosolic SctN, SctO, and SctK proteins enriched in the inner membrane fraction (Figure 5a). This is not surprising given their association with integral membrane proteins during their dynamic involvement in active T3SS assembly as shown previously for Salmonella (Lara-Tejero et al., 2011). Critically, there was no difference in localization of all these tested components between the parental and the N-terminal YscX mutants. We interpret these data to indicate that assembly of the bacterial envelope spanning “basal body” structure occurs independently of YscX.

**Figure 4** YscX influence on the production of Ysc structural components. *Y. pseudotuberculosis* from overnight cultures were sub-cultured into BHI medium in the presence (+) or absence (−) of calcium ions at 26°C for 1 h and at 37°C for 3 h. Protein samples were fractionated by SDS–PAGE in the range of 12 to 18% acrylamide and then transferred onto a PDVF membrane support for immune-detection with rabbit polyclonal antiserum to SctC, SctD, SctJ, SctV, SctN, SctO, SctK and DnaJ (a) as well as to SctI and SctF (b). Synthesis fractions represent bacterial protein (protein within intact bacteria and secreted in the culture medium). Secretion fractions represent protein freely released into the culture supernatant. Strains: ΔlcrQ null mutant (YscXwt – parental), YPIII/pIB26; ΔsctU, ΔlcrQ, YPIII/pIB75-26; ΔyscX, ΔlcrQ, YPIII/pIB880-26; ΔyscY, ΔlcrQ, YPIII/pIB890-26; yscX_∆3−7, ΔlcrQ, YPIII/pIB88002-26; yscX_∆8−12, ΔlcrQ, YPIII/pIB88003-26; yscX_∆13−22, ΔlcrQ, YPIII/pIB88004-26; YPIII (PC) is cured of the virulence plasmid. Arrows indicate specific protein band of interest and asterisks identify unknown non-specific cross-reacting proteins. Approximate molecular masses are shown in parenthesis and were deduced from primary amino acid sequence.
function. This corroborates earlier findings where SctQ, which itself needs many of the T3SS components for its assembly, was shown to localize perfectly fine in a yscX null mutant (Diepold et al., 2010).

We then assessed if the N-terminal YscX deletions affected subcellular localization of native SctI and SctF. Strikingly, only parental Y. pseudotuberculosis and the YscXΔ3–7 producing strain could export detectable levels of SctI to the periplasm (Figure 5b). This was despite all strains being capable of targeting SctI for secretion as judged by its widespread presence in the inner membrane fractions (Figure 5b). The absence of SctI in the periplasm of the ΔyscX null mutant and bacteria producing YscXΔ8–12 corroborates observed defects in SctI assembly. These observations suggest that the N-terminal region of YscX is critical for the earliest steps in T3SS export, including the biogenesis of the Yersinia T3SS needle complex composed of at least SctI and SctF.

2.8 | Localization of YscX in an active T3SS

Our observations implied that YscX orchestrates ordered export and/or assembly of the early substrates SctI and SctF. We wondered...
if defective YscX influenced this role in bacteria containing an intact T3SS. To address this, we first generated the mutant forms of YscX appended with a FLAG™ epitope at the N-terminus. In the low copy number pMMB208 vector, we cloned the alleles under control of an IPTG inductor promoter, before introducing them into parental *Y. pseudotuberculosis*. When induced ectopically with 0.4 mM IPTG, we detected in the synthesis fraction comparable levels of all FLAG-YscX variants (Figure 6a, upper panel). This production did not interfere with the ability of these bacteria to secrete SctB. However, a slight reduction in percent SctE and YopE secretion by bacteria expressing FLAG-YscXΔ8-12 and FLAG-YscXΔ13-22 was observed (Figure 6b, lower panel and 6C). Critically, this coincided with a loss of secretion of both FLAG-YscXΔ8-12 and FLAG-YscXΔ13-22 by a fully functional T3SS (Figure 6a, lower panel). In contrast, both FLAG-YscXwt and FLAG-YscXΔ3-7 were efficiently secreted (Figure 6a, lower panel). This corroborated our earlier in cis observation (see Figure 2a). Using these same secretion-permissive growth conditions, we performed a sub-cellular fractionation on this strain collection. The ectopically produced FLAG-tagged proteins enriched in the inner membrane fraction, and with a minor amount detected in the cytoplasm fraction (Figure 6d). Inner membrane enrichment of the FLAG-YscXΔ8-12 and FLAG-YscXΔ13-22 occurred even though a fully functional T3SS failed to secrete these two proteins (Figure 6a, lower panel). Hence, all three YscX N-terminal variants target the inner membrane, and although not confirmed, presumably meant that all could co-locate to the T3S base, suggesting an interaction with SctV and YscY. However, only FLAG-YscXΔ3-7 was capable of secretion. Lack of FLAG-YscXΔ8-12 and FLAG-YscXΔ13-22 secretion could not be due to interference with Sctl and SctF assembly because the export of middle and late T3S substrates continued, albeit at reduced efficiency.

3 DISCUSSION

Based on the findings in this study, we propose a model that demonstrates the essential need for YscX to establish Ysc-Yop T3SS activity (Figure 7). A consequence of the general secretion defect caused by non-secreted YscX mutants was an inability to correctly localize Sctl and SctF and, therefore, a failure to assemble the needle. Hence, the coordinated assembly of the Sctl adapter/washer followed by the SctF needle filament indicates that YscX secretion is a key element in mediating spatiotemporal control of early Sctl and SctF substrates during Ysc-Yop T3SS assembly.

Our data do not disclose how YscX imparts this control but remains consistent with the suggestion that it is involved in recruitment and sorting of Sctl and SctF for prioritized secretion (Wagner et al., 2018). However, the loss of control is not necessarily due to a lack of YscX secretion per se. Alternatively, YscXΔ8-12 and YscXΔ13-22 may form non-productive interactions with the sorting platform, either because the binding is too tight or because it blocks access points for other substrates. This would result in a general secretion blockage of any substrate, including YscXΔ8-12 or YscXΔ13-22 themselves. To investigate these possibilities, we attempted an immunoprecipitation experiment to detect a direct interaction between YscX with either Sctl or SctF, which was unsuccessful. However, this negative result is not definitive because the proposed interactions would be transient and initially require formation of the tripartite YscX–YscY–SctV complex, which is an established element of YscX function (Diepold et al., 2011, 2012, 2015).

The fate of secreted YscX remains an enigma. We initially considered that secreted YscX was a minor component of the *Yersinia* T3SS needle following assembly of the basal body, given the role of YscX in Sctl and SctF needle biogenesis. However, preliminary results from probing purified needle fractions by Western blot with anti-YscX antisera did not support this hypothesis. Moreover, subcellular localization experiments were also plagued by detection sensitivity issues in immunoblot assays using the anti-YscX antisera. Hence, follow-up efforts to address this enigma must first uncover novel YscX variants that uncouple its roles in needle assembly and general secretion from its own secretion. Then, using these genetic tools combined with a functional fluorescent tagging system, it would be possible to visualize YscX export and targeting at the single cell level with live cell microscopy in real time.

Incidentally, the observation that YscX secretion is a prerequisite for needle biogenesis suggests that YscX may represent a new substrate secretion class secreted prior to the 'early' needle substrates, Sctl and SctF. This notion is corroborated by the YscX production and export profile in the SctU(N263A) cleavage site mutant background (Sorg et al., 2007). In this strain, there is some selective enrichment of YscX secretion among the known early-secreted substrates. This prompted us to examine if YscX influenced the SctU auto-cleavage event, a known mechanism of hierarchal secretion control (Bjornfot et al., 2009; Frost et al., 2012; Ho et al., 2017). However, we could demonstrate that SctU auto-cleavage was unaffected by the absence of functional YscX (Figure S10). Thus, despite evidence suggesting that YscX forms a new "pre-early" substrate secretion class, the mechanism orchestrating this secretion order is unclear. Elucidating this mechanism could be difficult because the whole secretion process is unlikely to be linear, even if one would manipulate a population of bacteria to synchronize their secretion.

In this study, we identified YscX as an indispensable component of Ysc-Yop T3SS biogenesis, regulation, and function. Reflecting this essentiality, the YscX amino acid sequence is predominately invariant among 115 fully sequenced strains of *Y. pestis*, *Y. enterocolitica*, and *Y. pseudotuberculosis*. The only exception is observed in a group of 7 sequenced *Y. enterocolitica* strains, where this form of YscX differs by a single Glu to Asp amino acid substitution present at position 28 (BLASTP version 2.10.1+ with the search performed on May 27, 2020) (Altschul et al., 1997, 2005). Hence, YscX function cannot tolerate sequence variation, and this reflects our own genetic complementation studies with both YscX (and YscY) family members that revealed total specificity of YscX (and YscY) for *Y. pseudotuberculosis* T3SS activity (Gurung et al., 2018).

Given the strict coupling of YscX function to the cognate YscY T3S chaperone (Francis, 2010; Pallen et al., 2003), studies aimed
to pry apart YscX-YscY interdependence will shed light on a potential new class of T3S chaperones. Furthermore, interactions between YscX, YscY, and SctV are sequential with the formation of YscX-YscY binding needed for interactions with SctV to occur (Diepold et al., 2012). This indicates that SctV engagement would need conformational changes in YscX and YscY. Advances in this

FiguRe 6  Fate of N-terminal FLAG™-tagged YscX variants produced in Yersinia with an intact T3SS. Parental bacteria (YPIII/pIB102) were conjugated with vector alone (pMMB208) or vector harboring N-terminal FLAG-tagged yscX alleles (FLAG-YscX). Following an overnight culture, Y. pseudotuberculosis strains were sub-cultured in BHI broth lacking calcium at 26°C for 1 h and then shifted to 37°C for 3 h. Where indicated, IPTG at a concentration of 0.4 mM was added during the temperature shift. Protein samples from bacterial pellet (Synthesis) and trichloroacetic acid-precipitated bacterial supernatant (Secretion—for YscX variants) or directly sampled supernatant (secretion—Yops) were fractionated by acrylamide SDS–PAGE for recombinant YscX separation (a) or for Yops separation (b). Samples were immunoblotted with mouse anti-FLAG monoclonal antibody (a) or a cocktail of rabbit polyclonal anti-SctE, anti-SctB and anti-YopE antibodies (b). At least three independent experiments were used to quantify relative SctE, SctB and YopE synthesis and secretion values ± standard errors of the means using Image Lab 6.0.1 (Bio-Rad) (c). Percent total secretion values were calculated as the ratio between the amount of secreted and total protein. Sub-cellular fractions representing the cytoplasm, inner membrane, periplasm and outer membrane were prepared from parental bacteria harboring N-terminal FLAG-tagged yscX alleles (FLAG-YscX variants) to determine the localization of recombinant YscX variants (d). Lysate refers to the non-fractionated total cell extract. Samples were analyzed by Western blotting using the mouse anti-FLAG monoclonal antibody. Strains: parent (YscX^+), YPIII/pIB102; parent/empty vector, YPIII/pIB102, pMMB208; ΔyscX null mutant, YPIII/pIB880; ΔyscX/empty vector, YPIII/pIB880, pMMB208; parent/FLAG-YscX^+, YPIII/pIB102, pJMG242; parent/FLAG-YscXΔ3-7, YPIII/pIB102, pJMG349; parent/FLAG-YscXΔ8-12, YPIII/pIB102, pJMG350; parent/FLAG-YscXΔ13-22, YPIII/pIB102, pJMG352. Approximate molecular mass values shown in parentheses were deduced from primary amino acid sequences of YscX-YscY binding needed for interactions with SctV to occur (Diepold et al., 2012). This indicates that SctV engagement would need conformational changes in YscX and YscY. Advances in this
process would catalyze opportunities to dissect how this tripartite complex may recognize and prepare the needle components for export, a process that must also involve YscX release from the complex and subsequent export from the bacteria. To this end, the non-secreted mutant forms of YscX described in this study will be helpful in dissecting tripartite complex function in the context of Ysc-Yop T3SS activity, and will also contribute valuable knowledge to elucidate the YscX secretion enigma.

The nature of T3SS injectisomes is generally quite diverse. Therefore, a relevant question concerns how broadly applicable YscX is to all such T3SSs. It is clear from our previous work that YscX is unique to the Ysc-T3SS clade that is produced by certain bacterial species within the Yersinia, Pseudomonas, Aeromonas, Photobacterium, and Vibrio genera (Gurung et al., 2018). YscX–YscY heterodimer formation is conserved in this clade, as is an YscX–YscY complex formed with SctV (Gurung et al., 2018), which together mediates export and/or assembly of early T3S substrates (Diepold et al., 2012). In the non-Ysc-T3SS clades that all lack YscX and YscY homologs, the equivalent SctV protein does not appear to possess any additional domains that could accommodate the functions of YscX. This is corroborated by the fact that SctV from Y. pseudotuberculosis and S. enterica–Typhimurium are not interchangeable (Ginocchio & Galán, 1995). Thus, it is utterly intriguing that the Ysc-T3SS clade has evolved a unique system in YscX–YscY–SctV for coordinating early substrate secretion. Several structural studies have proposed a conserved region in the cytoplasmic domain of SctV that is thought to recognize a chaperone-substrate complex and prepare substrates for secretion (Erhardt et al., 2017; Kuhlen et al., 2021; Majewski et al., 2020; Matthews-Palmer et al., 2021; Xing et al., 2018; Yuan et al., 2021). Hence, SctV has a conserved universal role in all T3SSs, although bacteria producing the Ysc-T3SS clade seems to have customized this function in a way that relies on the YscX–YscY complex. What was the force driving the evolution of this YscX–YscY–SctV connection remains a key question. Nevertheless, in light of a newly discovered accessory protein secreted by the Salmonella SPI-1 T3SS that controls needle filament assembly (Kato et al., 2018), or a novel chaperone-mediated secretion switching from early to middle substrates in the Salmonella SPI-2 T3SS (Takaya et al., 2019), it is feasible that non-Ysc T3SSs do possess alternative mechanisms serving the same purpose as does YscX–YscY function. Presumably, specific variations have evolved among different T3SS families given the evolutionary divergence of T3SS that has occurred with respect to different host cell tropisms (Abby & Rocha, 2012).

As a tool to report on YscX secretion, YscX-Bla fusion proteins were engineered based on the experiences from a previous study by this laboratory (Amer et al., 2011). The method is convenient, but it is necessary to keep in mind a couple of caveats. Although the native β-lactamase reporter protein is stable (Amer et al., 2011), the measurable levels of accumulated β-lactamase reporter protein can be influenced by at least two factors. The first factor is the efficiency

![FIGURE 7 Schematic representation of YscX N-terminus mediated control of the Ysc-Yop T3SS. The N-terminal region of YscX (magenta circle) harbors an independent secretion signal that promotes its T3S-dependent secretion. Manipulation of this N-terminal region (blue circle) either by targeted deletion or by defined domain swapping with equivalent secretion signals from other T3S substrates completely disrupts T3SS activity. However, all N-terminal YscX variants maintain an equivalent bipartite interaction with YscY and/or a tripartite interaction with YscY and SctV. Moreover, they are recruited to the inner membrane base in an unbiased manner. The N-terminal region of YscX harbors an additional and a specific recognition signal critical for export and/or assembly of early substrates SctI and SctF. Inset—colored shapes used to depict YscX variants, YscY, SctV, SctI and SctF. IM—inner membrane, OM—outer membrane.](image-url)
of translation of the appended sequence at the N-terminus of the reporter protein. This is illustrated herein with the YscX1-Bla and YscX2-Bl a fusions. Both generated stable product (see Figure S1b), but the amount of the YscX1-Bl a product accumulated is much lower than for the YscX2-Bl a (see Figure 1a). This is a clear indication that the yscX sequence encoding the first 5 codons translates more efficiently than does the shorter sequences of 1 codon. The second factor is the change in overall folding of recombinant Bl a fusion proteins that can lead to increased susceptibility to digestion by intracellular proteases. This is exemplified by a comparison of the longer fusions YscX15-Bl a and YscX20-Bl a. Both accumulate equivalent steady-state levels, suggesting similar translation efficiencies (see Figure 1a). However, the larger YscX20-Bl a fusion is noticeably less stable, and stability deteriorates further with the appending of longer yscX sequences to the N-terminus of Bl a (see Figure S1b). It is important to be kept in mind these caveats since they would influence any translational reporter system. Nevertheless, in our hands not only did the Ysc-Bl a translational reporter fusions identify the first ten codons as a minimal N-terminal secretion signal for YscX, but they also indicated that elements of yscX translation control exist within the N-terminus region. This important observation will be followed up in future work, because mechanisms of translation control are under-appreciated in the T3SS research community. Furthermore, we are also keen to address why appending a tag at the YscX C-terminus has a dramatic negative impact on the integrity of the recombinant YscX. We infer from this work that the C-terminus of YscX is intolerant to genetic manipulation. Realistically, this effect can be understood only after a structure of YscX has been resolved.

Finally, one other serendipitous finding from this work concerned the loss of secretion control in the reciprocal YscX2-15-LcrQ chimera. It was evident that in non-secretion-permissive conditions bacteria producing and secreting a variant of LcrQ that contained residues 2 to 15 derived from YscX secreted at least SctB, SctE and YopE. This is a bold phenotype and mimicking loss-of-function lcrQ mutants (Li et al., 2014a; Rimpiläinen et al., 1992; Stainier et al., 1997; Wulff-Strobel et al., 2002). There is little understanding of the molecular mechanism of LcrQ secretion control even in light of new findings linking LcrQ with intracellular proteases. This is exemplified by a comparison of the longer fusions YscX15-Bl a and YscX20-Bl a. Both accumulate equivalent steady-state levels, suggesting similar translation efficiencies (see Figure 1a). However, the larger YscX20-Bl a fusion is noticeably less stable, and stability deteriorates further with the appending of longer yscX sequences to the N-terminus of Bl a (see Figure S1b). It is important to be kept in mind these caveats since they would influence any translational reporter system. Nevertheless, in our hands not only did the Ysc-Bl a translational reporter fusions identify the first ten codons as a minimal N-terminal secretion signal for YscX, but they also indicated that elements of yscX translation control exist within the N-terminus region. This important observation will be followed up in future work, because mechanisms of translation control are under-appreciated in the T3SS research community. Furthermore, we are also keen to address why appending a tag at the YscX C-terminus has a dramatic negative impact on the integrity of the recombinant YscX. We infer from this work that the C-terminus of YscX is intolerant to genetic manipulation. Realistically, this effect can be understood only after a structure of YscX has been resolved.

4 | EXPERIMENTAL PROCEDURES

4.1 | Bacterial strains and growth conditions

The bacterial strains we used in this study are listed in Table S2. Bacteria were routinely cultivated in Luria Bertani (LB) agar or broth at either 26°C (Y. pseudotuberculosis) or 37°C (E. coli) with aeration. Antibiotics were added when required at the final concentrations of Cb: 100 μg per ml (carbenicillin—Cb), 50 μg per ml (kanamycin—Km) and 25 μg per ml (chloramphenicol—Cm). Analysis of T3SS by Y. pseudotuberculosis occurred at 37°C in Brain heart infusion (BHI) broth. Media containing Ca2+ ions was the non-inducing condition (BHI supplemented with 2.5 mM CaCl2), while media devoid of Ca2+ ions was the inducing condition (BHI supplemented with 20 mM MgCl2 and 5 mM Ethylene glycol-bis-[β-aminoethyl ether]-N,N,N’,N”-tetraacetic acid).

4.2 | PCR amplification and sequence analysis

Amplified DNA fragments obtained by PCR used the appropriate oligonucleotide combination listed as online supplementary information (Table S3). Sigma-Aldrich Co (Dorset, England) synthesized these custom oligonucleotides. Amplified fragments were confirmed to be mutation free by being first cloned using the InStAclone PCR cloning kit (Thermo Fisher Scientific, Inc.) and then sequenced by Eurofins MWG Operon AG (Ebersberg, Germany).

4.3 | Site-directed mutagenesis and allelic exchange by homologous recombination

For the construction of in-frame yscX deletions and chimeric variants harboring exchanged N-terminal secretor domains, mutagenized DNA fragments were generated by overlap PCR (Horton & Pease, 1991). DNA fragments were cloned into the XhoI-BamHI digested suicide meganuclease vector, pDM4 (Milton et al., 1996). These plasmids transferred into E. coli S17-1,pir served as the donors in conjugal matings with the appropriate Y. pseudotuberculosis recipients. Selection based on Cm resistance enabled the isolation of conjugates having an initial single homologous recombination event that lead to integration of the entire meganuclease plasmid. This was followed by conventional sucrose sensitivity measures (Milton et al., 1996) to enhance selection for appropriate secondary homologous recombination events that facilitated exchange of wild type alleles with mutated alleles and concomitant meganuclease plasmid excision. Verification of generated in cis mutations on the virulence plasmid utilized a combination of diagnostic PCR and sequence analysis prior to use.

4.4 | Generation of epitope-tagged protein expression constructs

yscX incorporating either a His6 N-terminal or C-terminal epitope tag was PCR amplified and ligated into the PsiI and BamHI sites within the pMMB208 expression vector. In addition, yscX and N-terminal deletions yscX variants with a 5-prime FLAG™ epitope were PCR amplified and cloned into BamHI-EcoRI digested pMMB208. Similarly, yscY incorporating either a FLAG™
N-terminal or C-terminal epitope tag was PCR amplified and ligated into PsI and BamHI restricted pMMB208. This placed the epitope-tagged alleles under an IPTG inducible promoter. The constructs in E. coli S17-1::pir were mobilized into the appropriate Yersinia background.

4.5  Detection of type III substrate synthesis and secretion

Induction of type III substrate synthesis and secretion from Y. pseudotuberculosis was essentially performed as previously described (Amer et al., 2011). Sampled protein content was either directly from 2 ml bacterial cultures or from 10 ml culture volumes first precipitated with trichloroacetic acid. Sampling direct from the bacterial suspension, which contains a mix of protein associated with bacteria and protein released into the supernatant, assessed total protein content. Sampling of the cleared supernatant provided an assessment of the secreted protein levels. Fractionation of collected protein samples was performed by SDS-PAGE. To detect individual proteins, SDS–PAGE fractionated protein was wet transferred onto PVDF membranes and subjected to immunoblotting. Chemiluminescent detection utilized the Thermo Scientific Pierce ECL 2 Western Blotting Substrate.

4.6  Antibodies and antibody production

Primary rabbit polyclonal antiserum against the secreted SctE, SctB, and YopE were a gift from Hans Wolf-Watz (Umeå University, Sweden). Primary rabbit polyclonal antiserum against the synthetic peptide (NH₂-CLHRAQDYRRELDTL-(CONH₂) incorporated the residues 70 to 83 of YscX was generated by AgriSera AB (Vännäs, Sweden). Antibodies to various other Ysc antibodies were generated by Agrisera AB and were raised against the purified proteins ZZ-SctO, His-SctJ, His-SctK, His-SctN, His-SctC (residues 26 to 242), SctI and SctD (residues 143 to 419). Mikael Lindberg at the Protein Expertise Platform, Umeå University, expressed and purified all proteins. Mouse monoclonal anti-Penta-His antibody and mouse monoclonal anti-Flag M2 antibody were purchased from Qiagen GmbH (#34669; Hilden, Germany) and Sigma (#F3165; St Louis, Missouri, USA) respectively. Secondary antiseras was anti-rabbit or anti-mouse antibody conjugated to horseradish peroxidase (GE Healthcare, Buckinghamshire, United Kingdom).

4.7  Yeast plasmid construction, transformation, and n-hybrid assays

The yscXΔ35-7 yscXΔ87-12 and yscXΔ132-22 alleles were amplified by PCR using primers described in Table S3 and translationally fused to the GAL4 activation domain in the yeast two-hybrid vector pGADT7 (Clontech Laboratories, Palo Alto, CA) (Table S2). The generation of a construct expressing the wild type yscX allele in pGADT7, and a construct expressing the native yscY allele fused to the GAL4 DNA binding domain in pGBK7 have been described previously (Bröms et al., 2005; Francis et al., 2001). Transformation of the S. cerevisiae reporter strain AH109 was performed as described earlier (Francis et al., 2000). Protein interactions from multiple independent transformations were determined by measuring the activation of the ADE2 reporter gene activation and the HIS3 reporter gene during growth on tryptophan and leucine minus SD synthetic minimal medium also lacking either adenine or histidine, respectively. The latter also required the addition of 4 mM 3-aminotriazole in the growth media to overcome any risk of false positives (James et al., 1996).

To confirm the stable production of the bacterial proteins in yeast, we adapted a previously described protocol (Francis et al., 2000). Overnight cultures were grown in minimal selection medium. A volume of 0.1 was used to inoculate 50 ml of fresh non-selective YPD medium. Cultures were then grown at 30°C until mid-log phase (OD₆₀₀ between 0.4 and 0.6), chilled quickly, and then harvested by centrifugation at 1000 RCF for 5 min at 4°C. Pellets were washed in equal volume ice-cold H₂O, then immediately frozen in a dry ice-ethanol bath. Cracking buffer [8 M urea, 5% (v/v) SDS, 40 mM Tris–HCl, pH 6.8, 0.1 mM EDTA, 0.4% (w/v) bromophenol blue] was prewarmed to 60°C, and a final concentration of 1% (v/v) β-mercaptoethanol. Complete protease inhibitor cocktail (Boehringer Mannheim; used at a concentration recommended by the manufacturer) and 174.2 μg/ml phenylmethylsulphonyl fluoride (PMSF) were added immediately before use. Subsequently, PMSF was added every additional ~7 min. Yeast was rapidly thawed at 60°C in 300 μl of cracking buffer and transferred directly into a 0.8 volume of glass beads contained in 2 ml reaction tubes. Samples were heated at 70°C for 10 min before a vigorous 1 min vortex. Cell debris and unbroken cells were pelleted at 4°C for 5 min at 14,000 rpm. Supernatants were collected into fresh ice-cold 2 ml reaction tubes containing PMSF. Following denaturation at 95°C for 5 min, samples were immediately fractionated by SDS–PAGE and protein content visualized by immunoblot. Proteins fused to the GAL4 DNA binding domain were detected with mouse monoclonal anti-GAL4 DNA BD antibody (#630403, Clontech Laboratories Inc). Proteins fused to the GAL4 activation domain were detected with mouse monoclonal anti-HA antibody (clone 12CA5) (Roche Diagnostics Corp.).

Yeast three-hybrid assays used pGADT7 and the pBridge vector (Clontech Laboratories). This method is assayed for specific protein–protein interactions in the context of a third protein. From the pBridge vector, YscY was constitutive expressed as a Gal4 DNA BD fusion protein through the alcohol dehydrogenase promoter (MCIS cloning site). Additionally, variants of YscX were set up to be conditionally expressed from the MET25 promoter (cloning site MCSII). Yeast three-hybrid assays used pGADT7 and the pBridge vector (Clontech Laboratories). This method is assayed for specific protein–protein interactions in the context of a third protein. From the pBridge vector, YscY was constitutive expressed as a Gal4 DNA BD fusion protein through the alcohol dehydrogenase promoter (MCIS cloning site). Additionally, variants of YscX were set up to be conditionally expressed from the MET25 promoter (cloning site MCSII). Yeast three-hybrid assays used pGADT7 and the pBridge vector (Clontech Laboratories). This method is assayed for specific protein–protein interactions in the context of a third protein. From the pBridge vector, YscY was constitutive expressed as a Gal4 DNA BD fusion protein through the alcohol dehydrogenase promoter (MCIS cloning site). Additionally, variants of YscX were set up to be conditionally expressed from the MET25 promoter (cloning site MCSII).
both *S. cerevisiae* Y190 and HF7c strains to permit dual reporter gene measurements via β-galactosidase assays (lacZ reporter) and growth on histidine dropout media (HIS3 reporter), respectively.

### 4.8 Construction and analysis of YscX translationally fused to signalless β-lactamase

Various length translational fusions linking the 5-prime region of yscX, including the predicted Shine-Dalgarno (SD) sequence, to truncated *bla* were then generated in pAAA010 that contains a secretion signal less *bla* reporter under IPTG inducible control. This was achieved by *BamHI*-KpnI cloning in two ways. Larger DNA fragments (>75 base pairs) were first amplified by PCR with the appropriate primer pairings listed in Table S3 and using lysed YP111/pIB102 as a source of template DNA. Smaller DNA fragments (<45 base pairs) were formed by the annealing of two complementary oligonucleotides prior to DNA ligation to the vector (Table S3). Analysis of recombinant β-Lactamase synthesis and secretion followed the trichloracetic acid-precipitation procedure for large-scale Yop synthesis and secretion analysis. After Western blot of fractionated protein, fusion proteins were detected with a primary mouse monoclonal anti-β-Lactamase antibody (MA1-20370, Thermo Scientific) followed by incubation with α-rabbit antiserum conjugated with horse radish peroxidase (GE Healthcare) and Pierce ECL 2 Western Blotting Substrate.

### 4.9 Intrabacterial protein stability

Intrabacterial protein stability in the presence of intracellular proteases was assessed after growth in BHI (Brain Heart Infusion) medium supplemented with 2.5 mM CaCl₂ (BHI + Ca²⁺) as described previously (Feldman et al., 2002). Fractionation of collected protein samples was by SDS–PAGE. For specific detection, protein was transferred onto a PVDF membrane. Membrane bound native YscX was detected with rabbit polyclonal YscX antiserum and recombinant YscX-Bla fusions with a rabbit polyclonal β-lactamase antiserum (#AB3738, Millipore), followed by horseradish peroxidase conjugated anti-rabbit antibody prior to chemiluminescent detection with the Pierce ECL 2 Western Blotting Substrate.

### 4.10 SctF surface localization and chemical cross-linking

Overnight cultures from *Yersinia* strains were grown with shaking at 26°C in 2 ml of BHI broth supplemented with 2.5 mM CaCl₂. Subsequently, 0.1 volumes of bacterial suspension were subcultured into 3 ml fresh media and incubated for 3 h at 37°C. After each culture was standardized by A₆₀₀, 1 ml volumes were harvested by centrifugation at 8000 g for 5 min at 4°C. Each bacterial pellet was gently suspended in 1 ml of cold 20 mM HEPES, 2.5 mM CaCl₂ (pH 8). Bacterial surface proteins were cross-linked for 30 min at RT with the non-cleavable, membrane-impermeable, amine-reactive cross-linker Pierce BS³ (Thermo Scientific) at a final concentration of 5 mM. Cross-linking reactions were quenched for 15 min by addition of Tris–HCl (pH 8.0) to a final concentration of 20 mM. Cell fractions were collected by centrifugation at 12,200 g for 5 min at 4°C. Bacterial pellets were then suspended in 100 µl of 1X sample buffer and analyzed by 12% acrylamide SDS–PAGE and immunoblotting with rabbit anti-SctF polyclonal antiserum (a gift from Hans Wolf-Watz) that underwent several rounds of immunoabsorption with purified SctF to enhance its monospecificity.

### 4.11 Cell fractionation and Ysc component localization

Bacterial cell fractionation of *Y. pseudotuberculosis* was performed as described previously with some modifications (Filip et al., 1973; Obi et al., 2011; Vesto et al., 2018). Following induction of T3S, cells from 2 ml of bacterial culture were normalized to the amount of bacterial cells at an optical density of 600 nm. Pelleted cells were obtained by centrifugation at 6000 g for 10 min at 4°C, washed once with 1 ml of PBS and resuspended in 1 ml of ice-cold osmotic shock buffer (30 mM Tris–HCl, pH 8, 20% (w/v) sucrose). After addition of EDTA to a final concentration of 2.5 mM and 15 min gentle agitation at 4°C, cells were pelleted at 3000 g for 12 min, resuspended in 300 µl of ice-cold 5 mM MgSO₄, and kept at ice for 20 min. The supernatant collected after centrifugation at 14,000 g for 10 min represented the periplasmic fraction. The pellet was then resuspended in 1 ml of lysis buffer (100 mM Tris–HCl, pH 8.0, 10 mM EDTA) and lysed on ice by sonication. Intact bacteria and unbroken cells were removed by centrifugation at 14,000 g for 5 min at 4°C. The supernatant was subjected to ultracentrifugation at 185,000 g for 35 min at 4°C to recover cytoplasmic proteins (supernatant). The pellet (total membrane fraction) was then washed with 500 µl of 20 mM potassium phosphate buffer, pH 7.0 at 208,000 g for 12 min. To separate inner and outer membrane proteins, the pellet was resuspended in 250 µl of 0.5% sarcosyl in 20 mM potassium phosphate buffer, pH 7.0 and incubated at room temperature for 30 min with gentle agitation. This mixture was centrifuged at 20,800 g for 45 min at 4°C. The soluble fraction contained the inner membrane proteins. The pellet enriched in outer membrane proteins was resuspended in 250 µl of 30 mM Tris–HCl, pH 8.0. All the sub-cellular fractions were standardized to the same concentration as measured by BCA protein assay kit (Pierce). Following standardization, 200 µl of each cellular fraction was mixed with 50 µl of 4 X SDS sample buffer and analyzed by SDS/PAGE (12%–18%) and immunoblotting. To confirm the integrity of cellular fractionation, rabbit polyclonal antibodies raised against H-NS, FtsH, SurA, and OmpA were used as the cytoplasmic, inner membrane, periplasmic, and outer membrane marker respectively (Obi et al., 2011; Sal-Man et al., 2013).
4.12  |  Kinetics of SctU autoproteolysis

The construction of pML13 (sctU appended with a 3-prime FLAG™ and cloned under the tac promoter of pMMB66EH) has been described previously (Lavander et al., 2002). The plasmid pML13 was conjugated into recipient Yersinia strains that either contained YscX variants or lacked YscX. Cultures of Yersinia were grown in BHI—Ca2+ at 26°C for 1 h. Following expression of SctU-FLAG™ with 0.4 mM IPTG, cultures were grown for an additional 1 h at 37°C. At time point 0, chloramphenicol was added to a final concentration of 50 μg/ml to the bacterial culture to inhibit de novo protein synthesis. Thereafter, bacterial cultures were collected at indicated time points and normalized to the number of bacterial cells at an optical density of 600 nm. Protein samples associated with bacterial pellets were then suspended in 1 X SDS sample buffer to be analyzed by SDS–PAGE and immunoblotted with mouse monoclonal anti-Flag M2 antibody.

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CONFLICT OF INTEREST

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

AUTHOR CONTRIBUTIONS

Conceived and designed the experiments: JMG, AAA, AD, MSF. Provided critical reagents: SC. Performed the experiments: JMG, AAA, AD. Analyzed the data: JMG, AAA, SC, AD, MSF. Wrote the paper: JMG, MSF. All authors read and approved the paper.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

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