



Atomic Structure of IgID Demonstrates Its Role as a Component of the Baseplate Complex of the *Francisella* Type VI Secretion System

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ABSTRACT Francisella tularensis, a Tier 1 select agent of bioterrorism, contains a type VI secretion system (T6SS) encoded within the Francisella pathogenicity island (FPI), which is critical for its pathogenesis. Among the 18 proteins encoded by FPI is IqID, which is essential to Francisella's intracellular growth and virulence, but neither its location within T6SS nor its functional role has been established. Here, we present the cryoEM structure of IgID from Francisella novicida and show that the Francisella IgID forms a homotrimer that is structurally homologous to the T6SS baseplate protein TssK in Escherichia coli. Each IqID monomer consists of an N-terminal β -sandwich domain, a 4-helix bundle domain, and a flexible C-terminal domain. While the overall folds of IgID and TssK are similar, the two structures differ in three aspects: the relative orientation between their β -sandwich and the 4-helix bundle domains; two insertion loops present in TssK's β -sandwich domain; and, consequently, a lack of subunit-subunit interaction between insertion loops in the IgID trimer. Phylogenetic analysis indicates that IgID is genetically remote from the TssK orthologs in other T6SSs. While the other components of the Francisella baseplate are unknown, we conducted pulldown assays showing IgIJ interacts with IgID and IgIH, pointing to a model wherein IgID, IgIH, and IgIJ form the baseplate of the Francisella T6SS. Alanine substitution mutagenesis further established that IgID's hydrophobic pocket in the N-terminal β -sandwich domain interacts with two loops of IgIJ, reminiscent of the TssK-TssG interaction. These results form a framework for understanding the hitherto unexplored Francisella T6SS baseplate.

IMPORTANCE Francisella tularensis is a facultatively intracellular Gram-negative bacterium that causes the serious and potentially fatal zoonotic illness, tularemia. Because of its extraordinarily high infectivity and mortality to humans, especially when inhaled, F. tularensis is considered a potential bioterrorism agent and is classified as a Tier 1 select agent. The type VI secretion system (T6SS) encoded within the Francisella pathogenicity island (FPI) is critical to its pathogenesis, but its baseplate components are largely unknown. Here, we report the cryoEM structure of IgID from Francisella novicida and demonstrate its role as a component of the baseplate complex of the Francisella T6SS. We further show that IgID interacts with IgIJ and IgIH, and propose a model in which these proteins interact to form the Francisella T6SS baseplate. Elucidation of the structure and composition of the Francisella baseplate should facilitate the design of strategies to prevent and treat infections caused by F. tularensis.

KEYWORDS type VI secretion system, baseplate complex, cryo-electron microscopy, *Francisella*, IgID, intracellular pathogen, contractile injection system

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rancisella tularensis is a facultatively intracellular Gram-negative bacterium that causes the serious and potentially fatal zoonotic illness, tularemia (1, 2). As a zoonosis, its animal reservoirs include rabbits and aquatic rodents with biting insects, such as ticks and deer flies, acting as vectors, though possibly nonessential to its enzootic persistence (3). Humans acquire the disease via bites from infected insects or by contact with infected animals or materials from the infected animals (2). Because of its extraordinarily high infectivity (inhalation of as few as 10 bacteria can cause a serious infection (4, 5)) and mortality, especially when inhaled, *F. tularensis* is considered a potential bioterrorism agent and is classified as a Tier 1 select agent (6).

Francisella causes disease by replicating intracellularly within cells of the infected animals and its intracellular life cycle has been well studied (7–9). After uptake by a process termed "looping phagocytosis" (10), the bacterium initially resides within a phagosome that acquires only limited late endosomal/lysosomal markers (7, 8). Subsequently, the bacterium permeabilizes its phagosome and replicates extensively within the cytosol (7, 8). A cluster of genes encoded on an island called the Francisella pathogenicity island (FPI) is essential to the capacity of the bacterium to disrupt its phagosome and replicate intracellularly (11, 12). It is thought that the FPI encodes an atypical type 6 secretion system (T6SS) (13). T6SS resembles an inside-out contractile phage and injects toxins and effector molecules into target prokaryotic or eukaryotic cells (14). The canonical T6SS consists of a membrane complex that anchors the system to the bacterial membrane, a baseplate complex, and a metastable contractile sheath that, upon contraction, drives an inner tube tipped with a spike and effector proteins across the bacterial inner and outer membranes and into the target cell (15).

Because the level of homology with other T6SSs is low and some of the components of other T6SSs have not been identified in the FPI, the Francisella T6SS has been classified as a separate system (T6SSii) (16). F. tularensis has three major subspecies of clinical relevance to humans: F. tularensis subsp. tularensis (type A, found only in North America), F. tularensis subsp. holarctica (type B, found both in North America and in Europe), and F. tularensis subsp. mediasiatica (found primarily in central Asia) (17). A closely related species, F. novicida (Fn), is often used as a model to study the Francisella T6SS because of several features: it is of low virulence; it has the same intracellular life cycle (9); its genome has 98% identity at the nucleotide level with F. tularensis subsp. tularensis and subsp. holarctica (18); and it is genetically more tractable because it has a single FPI rather than two as in F. tularensis. While the proteins encoded on the Francisella T6SS show little sequence homology with T6SSs of other bacteria, the structures that have been determined to date show strong structural homology with components of other T6SSs. For example, we determined the structure of the Francisella T6SS sheath by cryoEM, showing that it was composed of IgIA/IgIB heterodimers that form hexagonal rings, which stack via an interlaced array of beta-strands (19). The sheaths of the canonical T6SS (20) and R-pyocin (21) have been shown to have structurally homologous sheath structures with corresponding interlacing β -strand meshwork. We have shown (22) that the Francisella central spike complex consists of trimeric PdpA and VgrG that has both structural similarities to, and differences from, the central spike of the canonical T6SS and other contractile injection systems with known structures.

The precontraction T6SS baseplate encloses the central spike and serves as a scaffold upon which the sheath and tube are assembled. While the atomic structure of the baseplates of canonical T6SS (23), R-pyocin (24), and T4 phage (25) have been determined, neither the composition nor the structure of the *Francisella* T6SS baseplate is known. The FPI encoded protein IgID is essential for phagosome escape and intracellular growth (26–28), and virulence in animals (29–31), but its function and role in the *Francisella* T6SS have not been established. Here, we report the purification of IgID from Fn and the determination of its cryoEM structure at 3.0 Å resolution. We show that IgID forms homotrimers with structural homology to TssK of the canonical T6SS. We also demonstrate that IgID interacts with IgIJ and IgIH and suggest how they serve as components of the *Francisella* T6SS baseplate.

RESULTS

Purification and cryoEM structure of Francisella novicida IgID. The *igID* gene was deleted from Fn and expressed under the control of the *F. tularensis* live vaccine strain bacterioferritin promoter on a plasmid with a 3× FLAG tag. The results showed that the complemented strain was fully functional in phagosomal escape (Fig. S1). T6SS was induced by growing the bacteria in high KCl as described previously (19, 22). Analysis of the purified material by Coomassie blue staining (Fig. S2A) showed a single protein band at about 47 kDa that corresponded to the IgID immunoreactive band. In the final step of purification, IgID eluted on Sephacryl S200 gel filtration in the same position as sweet potato amylase (MW 200 kDa), indicating that it forms a multimeric complex (Fig. S2A). Western immunoblotting confirmed that the protein staining band corresponded with a single band of immunoreactive material (Fig. S2B).

To obtain the structure of IqID, we recorded 2880 cryoEM images. We carried out a single-particle analysis (Fig. S2C to F and Fig. S3A) and generated a 3.0 Å-resolution reconstruction of sufficient quality (Fig. S3B) for the model building of the major parts of the protein. The structure was in a homotrimeric state with dimensions of 86 Å \times 82 Å containing three IgID molecules (Fig. 1B), consistent with size exclusion chromatography. Among the 398 residues of the full-length protein, residues M1-Y268 were resolved in our map. Each subunit of the IgID trimer contained an N-terminal β sandwich (residues M1 to P146) and a middle 4-helix bundle domain (residues Y159 to Y268) (Fig. 1A). The N-terminal β sandwich started with a long α -helix followed by a β -sheet consisting of three anti-parallel β-strands. Between residues M1 to Y268, the cryoEM density corresponding to two connection loops (E52 to G57; D89 to D93) was visible but of insufficient quality to support atomic modeling (Fig. 1C). The C-terminal domain was not visible in our 3 Å map with C3 symmetry indicating the flexibility relative to the N-terminal part. After applying symmetry expansion and local classification, we obtained a map with extra density corresponding to the C-terminal domain (Fig. 1D and Fig. S2G). The extra densities were at a lower resolution and not symmetrical among the three IgID protomers, suggesting flexibility of this region.

The trimeric structure of IgID resolves extensive intersubunit contacts (Fig. S4A), involving more than 30 residues in each protomer (Fig. S4B to D). Interactions were primarily through N-terminal helices forming a three-helix bundle and middle 4-helix bundle domain, further enhanced by domain-swapped-like interactions of the N-terminal loop with the adjacent protomer's β sandwich. The interactions for trimerization were largely hydrophobic, with some salt bridges and hydrogen bonds involved.

IgID is homologous to E. coli TssK with similarities and differences. Our results demonstrate that the Francisella IgID is structurally homologous to the canonical T6SS baseplate protein TssK (Fig. 2A and B). While the sequence identity and similarity between IgID and TssK (E. coli) are only 18.8% and 33.9%, respectively (Fig. S4F), the secondary structures (Fig. S4F) and the 3D structures of the two proteins are similar (Fig. 2A and B). Both IgID and TssK are homotrimers, containing an N-terminal β sandwich, middle 4-helix bundle domain, and C-terminal domain. By superimposing the IgID protomer structure onto TssK (Fig. 2C), they are automatically aligned by the 4-helix bundle domain based on maximum matching. It is apparent that the 4-helix bundle domain aligns well, while the N-terminal β sandwiches do not match even though both are composed of N-terminal helices and β strands. When we intentionally align the two structures by N-terminal β sandwiches, the two 4-helix bundle domains shift away from each other (Fig. 2D). These comparisons show that the relative orientation between their β -sandwich and the 4-helix bundle domains is different even though the overall folds of IgID and TssK are similar. Looking closely at the alignment of N-terminal β sandwiches, we found two loops (loop1: A108 to R124, loop2: E134 to E139) protrude in TssK compared to IgID (Fig. 2D). In the IgID trimer, the three protomers separate as three individual lobes (Fig. 2E). While in TssK trimer, a ring-like structure is formed mediated by intersubunit loop interactions mainly by hydrogen bonds (Fig. 2F and Fig. S4E). In the published TssK structures with a C-terminal domain, either by crystallography or cryoEM (23, 32), not all the C-terminal domains of three protomers can

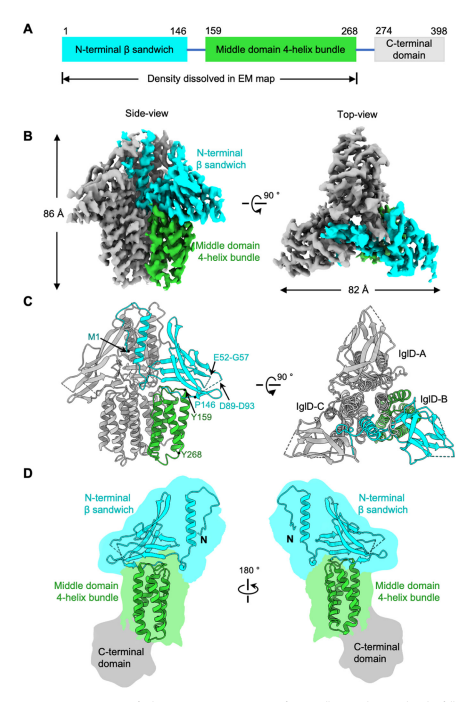


FIG 1 CryoEM structure of IgID. (A) Domain organization of *Francisella novicida* (Fn) IgID. The full-length protein consists of an N-terminal β sandwich (cyan), middle domain 4-helix bundle (green), and C-terminal domain (light gray). The C-terminal domain was not modeled. (B) Two different views of the cryoEM density map of IgID homotrimer. One protomer is represented with domains shown in the same color scheme as in A, the other two protomers are colored gray. (C) Two different views of IgID homotrimer structure. The structure is colored as in B. The three protomers are labeled as IgID-A, IgID-B, and IgID-C. (D) IgID monomer structure is shown in two different views. Domains are colored as in A. The N termini of the protein are labeled. Cartoon illustrations of full-length IgID monomer with C-terminal domain are shown based on the low-resolution map.

be traced, consistent with our assumption that the C-terminal domain was flexible. The C-terminal domain of TssK was shown to interact with the cytoplasmic domains of membrane complex (TssL and TssM) in T6SS. The flexibility of the C-terminal domain may provide a flexible link to maintain the anchorage of the baseplate to the membrane complex.

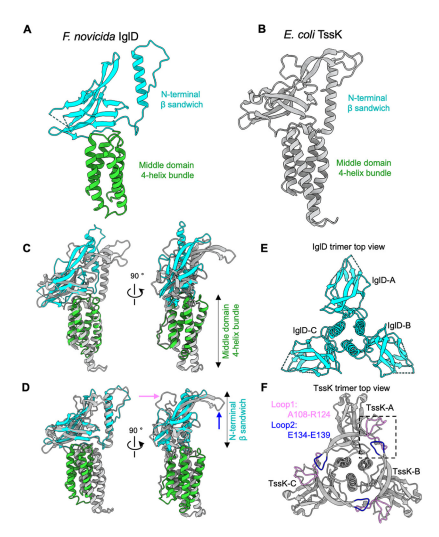


FIG 2 Structure comparison of IgID and TssK. (A) Structure of Fn IgID monomer. Domains are colored as in Fig. 1A. (B) Structure of EAEC TssK monomer (PDB accession no. 5MWN; Chain A). (C) Superposition of Fn IgID (color) and EAEC TssK (gray) monomers by 4-helix bundle domain. (D) Superposition of Fn IgID (color) and EAEC TssK (gray) monomers by N-terminal β-sandwich domain. The two arrows point to two insertion loops in TssK compared with IgID. The magenta one points to loop1 (A108-R124), the blue one points to loop2 (E134-E139). (E) IgID trimer top view, 4-helix bundle domain was omitted for clarity. The three protomers are labeled as IgID-A, IgID-B, and IgID-C. (F) TssK trimer top view, 4-helix bundle domain was omitted for clarity. The three protomers are labeled as TssK-A, TssK-B, and TssK-C. Loop1 in each protomer is colored magenta, and loop2 in each protomer is colored blue. The boxed region shows loop1 from TssK-A and loop2 from TssK-B.

The DeepMind "AlphaFold2" artificial intelligence program has been employed to predict protein folding (33). After we obtained the structure of IgID, we sought to compare the model predicted by this program with the experimentally derived structure. We ran AlphaFold2 on google Colab (34) requesting the prediction of IgID as a monomer (Fig. S5A). The predicted structure consisted of an N-terminal β sandwich, middle 4-helix bundle domain, and C-terminal domain as with the experimentally derived structure. The superimposition of the two structures showed that they were not the same (Fig. S5B). We then divided the structure into 3 rigid domains and superimposed them individually. The N-terminal β sandwich, aside from the N-terminal helix, aligns very well with a root mean square deviation (RMSD) of 1.9 Å on 96 residues (Fig. S5C). For the 4-helix bundle domain of the two structures (Fig. S5D), the RMSD was 1.2 Å on 110 residues. Comparison of the experimental and predicted protomer structures revealed that the structures of each globular domain are highly similar even though the overall architectures are not the same. The C-terminal domain predicted by

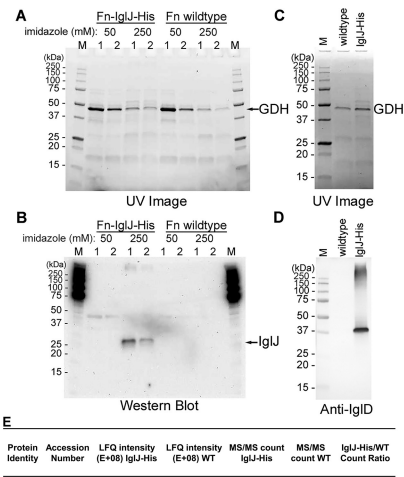
AlphaFold2 (Fig. S5E) is an assembly of two β sheets, one with five antiparallel β strands and the other with three. We then used AlphaFold2 to predict IgID as a homotrimer (Fig. S5F), which we had already established experimentally. The predicted trimer structure was unsatisfactory mainly because it lost the N-terminal helix trimerization and the domain-swapped interaction between protomers (Fig. S5F).

Identification of Francisella baseplate proteins by affinity pulldown and mass spectrometry. We set out to identify other Francisella baseplate components by affinity pulldown and mass spectrometry. In the canonical T6SS, the baseplate consists of 4 proteins, TssE, TssF, TssG, and TssK, that interact to form a wedge complex with stoichiometry TssK₆-TssG₁-TssF₂-TssE₁ and 6 wedge complexes assemble to form the baseplate (23). While our structural study demonstrated that IgID is the TssK ortholog, the other components of the Francisella T6SS baseplate have not been identified. We hypothesized that the Francisella protein IqIJ corresponds to the TssG of canonical T6SS based on its small size, absence of another assigned role in the T6SS, and the fact that it is essential for T6SS assembly, phagosome escape, and virulence (35). To test this hypothesis, we replaced the chromosomal iglJ gene of Fn with a gene encoding a C-terminal His₁₈ epitope tag (IgIJ-His) and showed that the strain was capable of T6SSmediated IgIC secretion and intracellular growth (Fig. S6). We then induced the T6SS in recombinant and wild-type (WT) control Fn by growing the bacteria in TSBC with 5% KCI. The bacteria were pelleted, washed to remove free amines, cross-linked with dithiobis succinimidyl propionate (DSP), lysed by sonication in RIPA buffer, and the IgIJ-His and its associated proteins enriched by Ni-NTA agarose affinity chromatography. While the main protein pulled down by the Ni-NTA affinity resin with or without His-tagging of IgIJ was found by mass spectrometry to be glutamate dehydrogenase (GDH) (Fig. 3A), Western immunoblotting confirmed enrichment of IgIJ (Fig. 3B) and demonstrated specific pulldown of IgID (Fig. 3C and D), i.e., IgIJ and IgID immunoreactive bands were present in the pulldown samples of the IgIJ His-tagged strain, but not that of the nontagged parental strain. The similar band intensities of GDH (Fig. 3A) confirmed comparable loading of the affinity resin with the IgIJ His-tagged and nontagged samples.

To identify additional proteins interacting with IgIJ, we acetone-precipitated the IgIJ-His and wild-type samples, digested them with trypsin, and identified and quantified the peptides of the constituent proteins by liquid chromatography with tandem mass spectrometry (LC-MS/MS)-based proteomics analysis. The relative specific enrichment of the proteins was determined by calculating the MS/MS spectral count ratios of proteins identified in the IgIJ-His sample versus the wild-type sample (Fig. 3E). Of the 5 proteins with the highest spectral count ratios, 4 were FPI encoded proteins (IgIJ, IgID, IgIH, and IgIA). Of the five proteins, IgIJ, IgID, and IgIH had the highest absolute spectral counts and the highest label-free quantification (LFQ) intensities in the IgIJ His-tagged sample, suggesting these three proteins interact as components of the *Francisella* baseplate wedge complex.

To test our hypothesis that IgIH is a component of the baseplate wedge complex, we replaced the chromosomal *igIH* gene of Fn with a gene encoding a C-terminal TwinStrep-His₁₈ dual epitope tag (IgIH-TS-His), induced the T6SS in the recombinant and wild-type control Fn by growing the bacteria in TSBC with 5% KCl, and performed DSP cross-linking and affinity pulldown using Ni-NTA agarose as described above. Western immunoblotting demonstrated that IgID was pulled down in the IgIH-TS-His expressing strain but not in the wild-type strain (Fig. 4A). IgIJ was not probed in the IgIH pulldown immunoblot because IgIJ was not epitope-tagged in the strain used and antibody to IgIJ was not available to us. However, mass spectrometry-based proteomic analysis of the samples demonstrated that IgIJ was pulled down in the IgIH-TS-His expressing strain but not in the wild-type strain (Fig. 4B). Together, these results are consistent with the hypothesis that IgID, IgIJ, and IgIH interact as T6SS baseplate proteins.

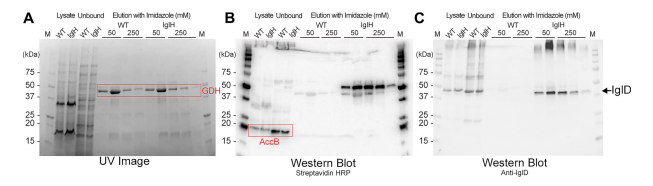
Interrogation of IgID and IgIJ interactions by alanine-substitution mutagenesis. Next, we explored how IgID and IgIJ interact and what are the structural homologs of IgIJ and IgIH in canonical T6SS. While BLAST searches of IgIJ and IgIH yield no significant sequence similarity with any non-Francisella proteins, their size and predicted



Protein Identity	Accession Number	LFQ intensity (E+08) IgIJ-His	LFQ intensity (E+08) WT	MS/MS count IgIJ-His	MS/MS count WT	IgIJ-His/WT Count Ratio
lglJ	A0Q7H9	88.2	0.7	597	16	37.3
UvrA	A0Q5P3	3.5	1.0	103	3	34.3
IgID	A0Q7I2	40.7	0.1	650	19	34.2
lglH	A0Q7H6	23.5	0.1	381	12	31.8
IgIA	A0Q7I5	7.0	0.4	198	8	24.8
GDH	A0Q832	75.1	207.7	1954	1073	1.8

FIG 3 IgIJ-His pulls down IgID. Wild-type (WT) Fn or Fn expressing IgIJ-His were grown in high KCI, cross-linked with DSP, lysed, and IgIJ-associated proteins enriched by nickel-agarose affinity chromatography. (A) SDS-PAGE gel showing proteins eluted in successive fractions with 50 mM and 250 mM imidazole visualized by stain-free UV imaging. (B) Western immunoblot using anti-His epitope antibody identifies IgIJ-His in the Fn IgIJ-His sample and not in the WT sample. (C and D) Samples eluted with 250 mM imidazole were further analyzed by SDS-PAGE with stain-free UV imaging of proteins (C) and Western immunoblotting using polyclonal antibodies against IgID (D). (E) MS/MS analysis of IgIJ-His and WT pulldown samples.

secondary structures suggest that IgIJ and IgIH correspond to baseplate proteins TssG and TssF, respectively, of canonical T6SS. In the cryoEM structure of the enteroaggregative *Escherichia coli* (EAEC) T6SS baseplate, TssG has two loops ("feet") extending from its C-terminal domain and each loop has three conserved hydrophobic residues in a triangular array, with the hydrophobic side chains of each loop projecting into a hydrophobic pocket formed by the N-terminal domains of a TssK trimer (36). Cherrak et al. (37) also identified a conserved sequence in the N-terminal domain of canonical TssK of EAEC corresponding to residues forming the hydrophobic pocket of the TssK trimer and further described each of the two TssG C-terminal feet as having a conserved pattern of three "LG repeats", wherein "LG" refers to a hydrophobic residue followed by a residue with either a small (e.g., glycine, serine) or basic (e.g., lysine, arginine) side chain. Sequence alignment of TssGs of other T6SSi revealed deviations from the "LG"



		LFQ intensity (E+08)			3)	MS/MS count			lglH/WT		
		<u>lgIH</u>		<u>WT</u>		<u>lgIH</u>		<u>WT</u>		Count Ratio	
Protein	Accession	50	250	50	250	50	250		250	50	250
Identity	Number	mM	mM	mM	mM	mM	mM	50 mM	mM	mM	mM
IgIH	A0Q7H6	4834	1846	0	0	481	625	0	0	∞	∞
IgID	A0Q712	245	83	0.05	0	76	115	2	0	38	∞
IgIB	A0Q7I4	90	35	0.23	0	70	91	4	0	17.5	∞
lglJ	A0Q7H9	86	23	0	0	33	43	0	0	∞	∞
IgIA	A0Q7I5	20	9	0.04	0	17	27	3	0	5.7	∞
GDH	A0Q832	20508	3679	9444	220	2122	1494	1585	431	1.3	3.4

FIG 4 Epitope-tagged IgIH pulls down IgID. Wild-type (WT) Fn or Fn expressing IgIH with TwinStrep and His-tags were grown in high KCl, cross-linked with DSP, lysed, and IgIH-associated proteins enriched by nickel-agarose affinity chromatography. (A) SDS-PAGE gel showing proteins in the lysate, affinity column pass-through (unbound), and eluted in successive fractions with 50 mM and 250 mM imidazole visualized by stain-free UV imaging with glutamate dehydrogenase (GDH) as the most abundant protein eluted by imidazole in both samples. Comparable amounts of GDH demonstrated comparable loading of the affinity resins. (B) Western immunoblot using Streptavidin-peroxidase identifies elution of TwinStreptagged IgIH in the IgIH lanes but not the WT lanes. The endogenous biotinylated protein, AccB (56), was identified at comparable levels in WT and IgIH lanes of lysate and unbound samples, again showing comparable loading of the affinity resins. (C) Anti-IgID immunostaining identifies pulldown of IgID in the Fn IgIH sample but not in the WT sample. (D) MS/MS analysis of acetone precipitated pellets of the 50 mM and 250 mM eluates of the IgIH and WT samples.

repeat motif, such as the presence of acidic residues, rather than small or basic residues, in some "LG" repeats of TssG of Klebsiella pneumoniae, Vibrio cholerae, and Yersinia pseudotuberculosis (36). Whereas the basic side chains of EAEC TssG "LG" repeat residues form salt bridges with acidic residues above the N-terminal domain (NTD) hydrophobic pocket of EAEC TssK (36), the acidic residues in the "LG" repeat of these "outliers" may also form salt bridges with basic residues above the NTD of their corresponding TssKs. It is noteworthy that the N terminus of IqID is largely consistent with the NTD consensus sequence of canonical T6SSi TssK (Fig. 5A), including the pivotal residues (W8, L14, L19) that contribute to the hydrophobic pocket (Fig. 5B and Fig. S7) interacting with LG repeats of the TssG feet. We have also identified conserved sequences within predicted loop structures near the C terminus of IgIJ (i.e., the putative "feet") similar to the TssG "LG repeat" motif described by Cherrak et al. (37) that could interact with the hydrophobic pocket in the NTD of IgID, matching the TssK-TssG interactions of canonical T6SS (Fig. 6A). As shown in Fig. 7A, Francisella IgID is phylogenetically closer to the T6SSiii TssKs of Flavobacteria and Bacteroidales, whose NTD has conserved residues similar to those of canonical TssK and Francisella IgID (Fig. 5A). Alignment of canonical TssG sequences (T6SSi) with those of *Flavobacteria* and Bacteroidales TssG (T6SSiii) revealed that, whereas the last "LG" repeat of canonical TssG was typically about 40 residues from the C terminus, the last "LG" repeat of T6SSiii TssG was at the very C terminus, with the hydrophobic residue being the last residue (Fig. 6A). We identified similar "LG" repeat sequences in loop regions of the C terminus

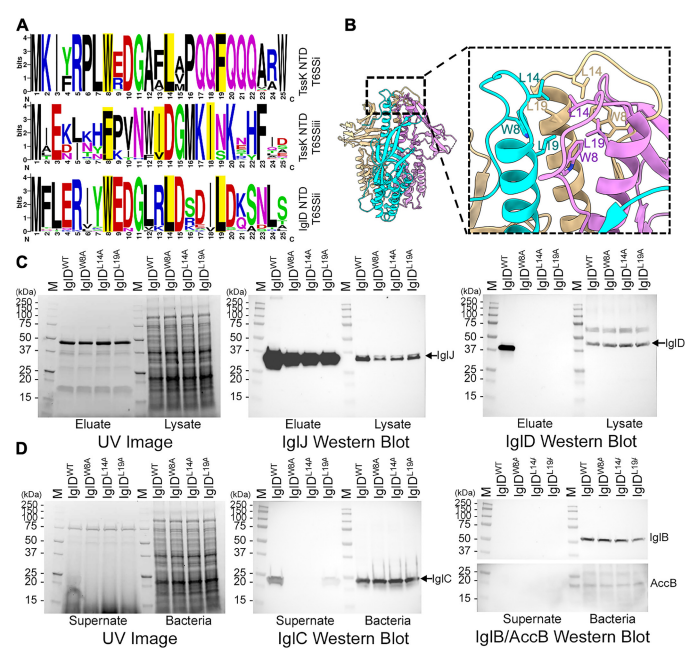


FIG 5 IgID N-terminal domain residues are essential for interaction with IgIJ and IgIC secretion. (A) Comparison of consensus N-terminal domain sequences of canonical T6SSⁱⁱ TssK, T6SSⁱⁱⁱ TssK, and Francisella (T6SSⁱⁱ) IgID. Conserved residues contributing to the hydrophobic pocket of T6SSⁱ and T6SSⁱⁱⁱ TssK and T6SSⁱⁱⁱ IgID are highlighted. (B) Key residues (W8, L14, L19) consistent with TssK-NTD in the IgID structure are shown in stick presentation. Individual protomers of the IgID homotrimer are colored cyan, magenta, and tan. (C) Alanine substitution of IgID NTD residues W8A, L14A, or L19A abolishes the IgIJ pulldown of IgID. Stain-free UV image shows comparable lane loading. IgIJ-FLAG Western blot shows the capture of IgIJ in parental and Ala-substitution strains. IgID Western blot shows pulldown of IgID only in the parental strain. (D) IgID NTD Ala substitution mutants are defective for IgIC secretion. Stain-free UV image shows comparable loading of lanes. IgIC Western blot shows secretion of IgIC into culture supernatant fluid only by the parental strain and comparable levels of IgIC in the Iysates of all strains. IgIB/AccB Western blot shows the absence of leakage of cytosolic proteins IgIB or AccB into the culture supernatant fluid.

of *Francisella* IgIJ, with the final hydrophobic residue of the last "LG" repeat usually being the last residue of the sequence (Fig. 6A). While the first putative foot of IgIJ is entirely within a loop region, the first "LG" repeat (V242, E243) of the second foot is predicted to be the last two residues of β -strand (Fig. S8F). However, the remainder of the second foot (residues 244 to 257) is predicted to be a loop (Fig. S8F), that would allow the hydrophobic side chains of the "LG" positions to form a triangular array for insertion into the IgID NTD hydrophobic pocket. Residue 243 (highly conserved

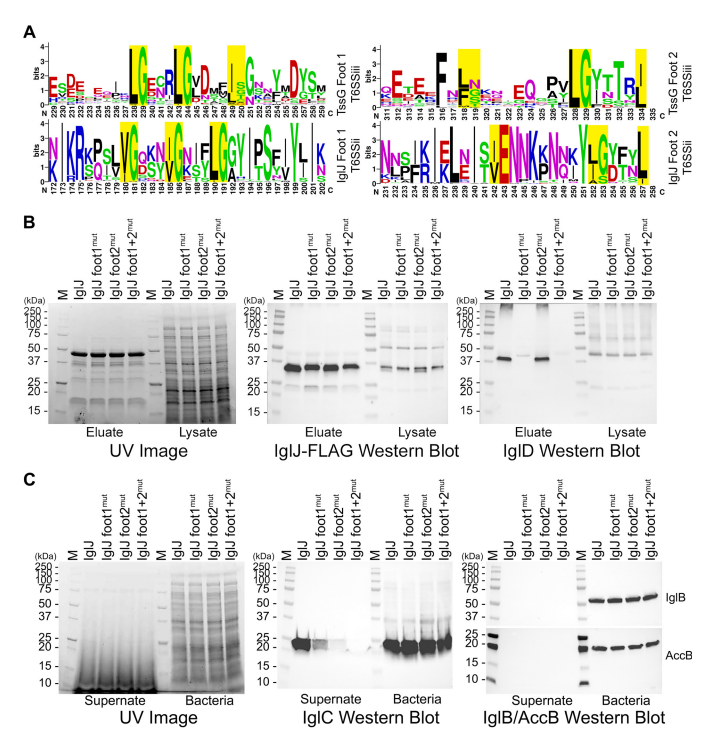


FIG 6 Alanine substitutions in the C terminus of IgIJ disrupt pulldown of IgID and decrease IgIC secretion. (A) Comparison of consensus sequences of putative T6SSiii TssG and T6SSiii IgIJ foot1 and foot2. Putative LG repeat positions of T6SSiii TssG feet (A, top lines) and corresponding sequences conserved in Francisella species (A, bottom lines) are highlighted. (B) Alanine substitution of the three hydrophobic residues of IgIJ foot1 (I180, V185, L190) abolishes IgIJ pulldown of IgID. Stain-free UV image shows comparable lane loading. IgIJ-FLAG Western blot shows the capture of IgIJ in parental and Ala-substitution strains. IgID Western blot shows pulldown of IgID in the parental strain and the strain with IgIJ foot2 substitutions (V242, E243, L257), but not the strains with alanine substitutions in IgI foot1 or both foot1 and foot2. (C) Impact of alanine substitutions (V242, E243, L257), but not the strains with alanine substitutions in IgIC secretion. Stain-free UV image shows comparable loading of lanes. IgIC Western blot shows that alanine substitution of foot1 markedly decreases IgIC secretion, substitution in foot2 even more dramatically decreases IgIC secretion, and substitution in both feet abolishes IgIC secretion. Western blot of the bacterial lysates shows comparable levels of IgIC in all of the strains. IgIB/AccB Western blot shows the absence of leakage of cytosolic proteins IgIB or AccB into the culture supernatant fluid.

Structure and Function of Francisella IgID

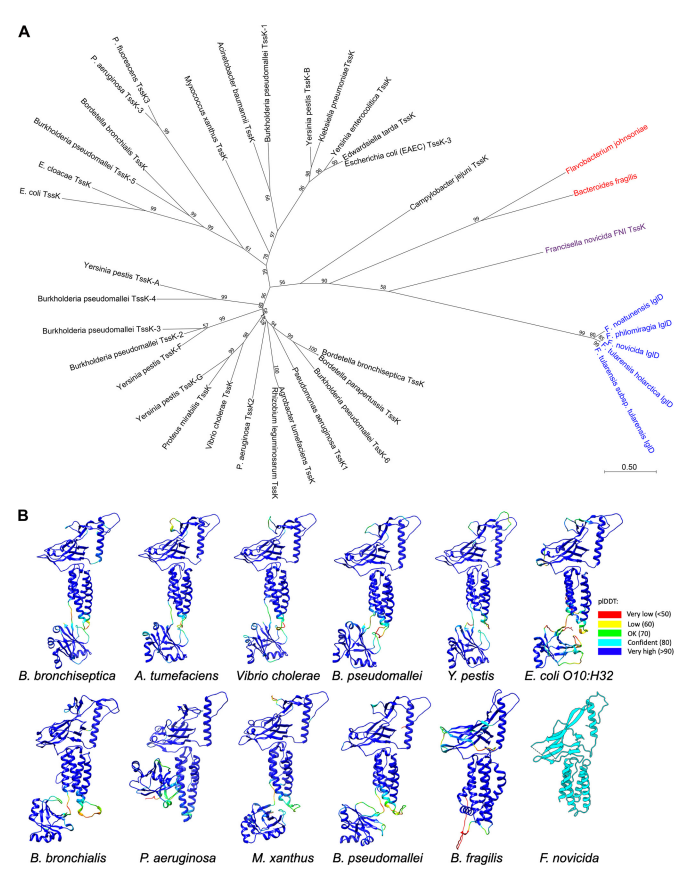


FIG 7 Phylogenetic relationships of IgID and TssK as inferred by using the Maximum Likelihood method and JTT matrix-based model (37). (A) T6SSⁱ are shown in black; T6SSⁱⁱ (*Francisella*) are shown in blue, and T6SSⁱⁱⁱ are shown in red. The tree with the highest log likelihood (-29351.96) is shown. The (Continued on next page)

glutamate) is acidic (Fig. 6A), rather than basic, but its sidechain may form a salt bridge with conserved basic residue 13 (arginine or lysine) above the hydrophobic pocket of the IgID NTD (Fig. 5A and Fig. S7). All the alignments and predictions indicate that IgIJ functions as the TssG homolog of EAEC T6SS to interact with IgID NTD hydrophobic pocket by two feet.

To test the hypothesis that IgIJ interacts with IgID NTD by two feet, we carried out a pulldown assay with alanine substitutions on IgIJ and IgID. In EAEC, mutation of any of the highly conserved residues that contribute to the TssK NTD hydrophobic pocket (W8, L14, F19) leads to a loss of TssG-TssK interaction and T6SS function (37). Francisella IgID has highly conserved NTD residues in the same positions (Fig. 5A and B), and we have found that replacing any one of these three residues with alanine (W8A, L14A, or L19A) abolishes the capacity of IgIJ to pulldown IgID (Fig. 5C) and abolishes T6SS IgIC secretion (Fig. 5D). In the case of EAEC, deletion of either foot1 or foot2 of TssG also blocks TssK pulldown and T6SS biological function (37). However, we found that deletion of either the putative foot2 region of IgIJ or the deletion of both putative feet resulted in undetectable IgIJ expression, as well as loss of IgIC secretion (unpublished data). Therefore, we examined the impact of alanine substitutions in putative IgIJ foot1 (I180A, V185A, L190A) and foot 2 (V242A, E243A, L252A, L257A). In foot2 we substituted alanine both for the hydrophobic residue and the highly conserved E243 to eliminate the potential salt bridge interaction. We found that the 4 alanine substitutions in foot2 did not block the IqIJ pulldown of IqID (Fig. 6B) and that the three alanine substitutions in foot1 or the combined foot1 and foot2 alanine substitutions were sufficient to block IgIJ pulldown of IgID (Fig. 6B). We observed marked impairment of IgIC secretion by F. novicida with the 3 alanine substitutions in IgIJ foot1, even greater impairment of IgIC secretion with the 4 alanine substitutions of foot2, and the abolition of IgIC secretion with combined substitutions in both foot1 and foot2 (Fig. 6C). The intracellular growth by these alanine substitution mutants within macrophage-like THP-1 cells paralleled their capacity for IgIC secretion (Fig. S9). Specifically, we observed severe impairment of intracellular growth with any of the three alanine substitutions in the IgID NTD, marked impairment in intracellular growth with the 3 IgIJ foot1 substitutions, even more marked impairment of growth with the foot2 alanine substitutions, and the combined foot1 and foot2 substitutions were as impaired in growth as a complete ially deletion (Fig. S9). These data indicate that the putative IgIJ foot1 "LG" repeats are necessary and sufficient to enable IgIJ-IgID interaction under the stringent conditions of the pulldown assay, which includes high salt (0.3 M NaCl) and 1% Triton X-100, both used to prevent nonspecific pulldown. Under the conditions of the pulldown assay, foot1 may play a stronger role in interacting with IgID because its loop is fixed within two regions of secondary structure, whereas the extreme C-terminal foot2 loop is inherently more flexible and disordered. On the other hand, within the intact bacterium (i.e., under physiological conditions and in the presence of accessory and chaperone proteins), the combination of hydrophobic and salt bridge interactions of foot2 may make it more important in interacting with the IgID NTD. While alanine substitutions in both putative feet of IgIJ were required for complete disruption of T6SS IgIC secretion, alanine substitutions in any of the three key conserved hydrophobic residues contributing to the IgID NTD hydrophobic pocket were sufficient to disrupt both pulldown and IgIC secretion because alteration of the IgID NTD hydrophobic pocket disrupts interactions with both IgIJ feet. These data support IgIJ and IgID interacting directly by two feet and NTD hydrophobic pocket similar to TssG and TssK interactions, which are essential for T6SS secretion, indicating IgIJ is a homolog to TssG.

FIG 7 Legend (Continued)

percentage of trees in which the associated taxa clustered together is shown next to the branches. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the JTT model, and then selecting the topology with a superior log likelihood value. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. This analysis involved 38 amino acid sequences. Evolutionary analyses were conducted in MEGA11 (39). (B) AlphaFold2 prediction results for TssK/IgID from representative species in (A). The source organism is labeled below each structure. The AlphaFold2 prediction models are colored according to pLDDT confidence. The last one is the Fn IgID structure solved in this study.

Phylogenetic analysis shows that IgID is genetically remote from the TssK orthologs in other T6SSs. In canonical T6SS, TssK is situated at the interface between the baseplate and the membrane complex of the T6SS. Its N-terminal domain interacts with the TssG of the baseplate and its flexible C-terminal domain interacts with the membrane complex (23). As such, TssK functions as an adaptor between the baseplate and membrane complex (23). Whereas the T6SS baseplate proteins TssF, TssG, and TssE have structural orthologs in the baseplates of extracellular contractile injection systems (e.g., myophage, R pyocins, and insecticidal antifeeding prophages [Afps]), TssK and IgID have no corresponding structural counterparts in extracellular contractile injection systems. This is likely attributable to the fact that contractile phages, R pyocins, and Afps have no structure corresponding to a membrane complex.

T6SSs have been divided into 4 classes, with the Francisella T6SS being in a class by itself (T6SSi), whereas canonical T6SSs of Gammaproteobacteria – e.g., Vibrio cholerae and Pseudomonas aeruginosa – belong to T6SSi; Flavobacterium johnsoniae and Bacteroides fragilis belong to T6SSiii, and the Candidatus Amoebophilus asiaticus Afp-like system belongs to T6SSiv and is phylogenetically closer to R pyocins and myophage than to other T6SSs (38). Phylogenetic analysis with MEGA11 (39) using IgID sequences of Francisella and TssK sequences of other bacteria confirmed that IgID of F. tularensis is phylogenetically remote from other TssK proteins (Fig. 7A). The AlphaFold2 predictions for TssKs from each main branch of the phylogenetical tree show that TssKs from different species exhibit highly similar folding architecture as Francisella IgID whose structure was determined in this study (Fig. 7B), demonstrating again that IqID is a structural homolog to TssK. Fn also has a second T6SS-like gene cluster called the F. novicida island (FNI), which encodes a hypothetical TssK-like protein (40). Phylogenetic analysis places the FNI TssK in between the T6SSⁱⁱ FPI IgID and the T6SSⁱⁱⁱ TssKs of *F. johnsoniae* and B. fragilis (Fig. 7A), suggesting that it may have been acquired by horizontal transfer. The FNI TssK and FPI IgID proteins have sequence homology only at their C-terminal domains (IgID residues 319 to 377) and no significant sequence homology in the regions for which we determined the structure. We did not identify any peptides attributable to the FNI tssK gene product in our mass spectrometry studies, and it is not known whether the FNI tssK gene is transcribed or functions as part of a T6SS.

DISCUSSION

In this study, we show that *Francisella* IgID forms homotrimers that are structural orthologs of baseplate protein TssK of canonical T6SS. We show by cross-linking affinity pull-down studies that IgID interacts with IgIJ and IgIH, and we propose that these proteins function as orthologs of canonical TssG and TssF in the *Francisella* T6SS baseplate (Fig. 8). We used AlphaFold2 to predict the IgIH and IgIJ structures based on their sequences (Fig. S8). The structure of IgIH predicted (Fig. S8E) by AlphaFold2 aligned well with the published structure of TssF (RMSD = 6.2 Å, TM-align (41) score of 0.55). On the other hand, the AlphaFold2 predicted structure of IgIJ (Fig. S8B) was very dissimilar to the published structure of TssG while a model of IgIJ predicted by I-tasser using TssG as a template (Fig. S8C) showed high similarity with TssG. Additional structural studies will be required to confirm our hypothesis that IgJ and IgIH are the *Francisella* orthologs of TssG and TssF.

Each wedge of the canonical T6SS baseplate also contains a fourth protein, TssE, a sheath initiator protein that interacts via a β -strand with the handshake domain of the sheath subunits closest to the baseplate. The *Francisella* TssE ortholog has not yet been identified and the *Francisella* genome lacks any protein with significant homology with canonical TssE.

Elucidation of the structure and composition of the *Francisella* baseplate will be useful in devising strategies to prevent and treat infections caused by *F. tularensis*. In this regard, it is noteworthy that Cherrak et al. (37) developed a biomimetic cyclic peptide that blocks the interaction of TssG feet with the highly conserved N-terminal domain hydrophobic pocket of TssK, thus blocking T6SS assembly and virulence. Because we showed that the *Francisella* N-terminal domain has both sequence and structural homology with canonical TssK and that mutation of any of the conserved hydrophobic

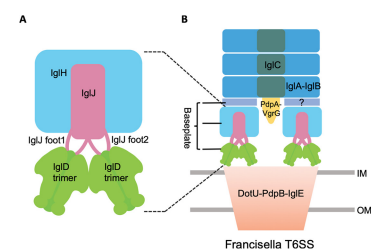


FIG 8 Schematic illustration of Fn T6SS baseplate assembly. (A) IgID trimer (colored green), IgIJ (magenta), and IgIH (light blue) are components of the Fn T6SS baseplate. IgIJ interacts with two IgID homotrimers via foot1 and foot2. (B) The baseplate docks to the DotU-PdpB-IgIE membrane complex (orange) and further recruits the tail complex (dark blue and dark green). The question mark within the baseplate indicates the unknown component(s). In particular, the *Francisella* sheath initiator protein, i.e., the TssE ortholog, has not been identified. OM, outer membrane; IM, inner membrane.

residues of the NTD leads to a loss of IgIJ-IgID interaction and a loss of T6SS secretion, a similar strategy will likely be effective for *Francisella*.

MATERIALS AND METHODS

Purification of IgID. Fn expressing N-terminally 3× FLAG-tagged IgID was grown in trypticase soy broth containing 0.2% L-cysteine (TSBC), 20 µg/mL kanamycin, and 5% KCl, rotating at 150 rpm at 37°C to an optical density (OD) of approximately 2. The bacteria were pelleted by centrifugation at 5000 \times gfor 1 h and resuspended in 100 mL of 50 mM Tris-HCl, pH 8.0, 1 mM EDTA, 1 mM N-ethylmaleimide (NEM), 1 mM phenylmethylsulfonyl fluoride (PMSF), supplemented with 100 μ g/mL lysozyme and EDTAfree protease inhibitor (Calbiochem, 1:1000) and benzonase (VWR, 1:10,000). The sample was sonicated with a probe tip sonicator (Cell Disruptor model W-375, Heat Systems Ultrasonics, Plainview, NY) while stirring in an ice bath, clarified by centrifugation at 44,400 \times g for 23 min, and applied to a Q-Sepharose column (XK-26, 13 cm bed height, GE Healthcare). The column was washed with 50 mM Tris-HCl, eluted with a 0 to 1 M NaCl gradient, and fractions containing FLAG-immunoreactive material were identified by sandwich ELISA in which the ELISA plate (high bind, polystyrene 96-well ELISA plate, Costar, Corning Inc.) was coated for 90 min with rat anti-FLAG monoclonal antibody (Novus Biologicals, 10 μ g/mL in PBS), blocked with 1% BSA, incubated with fractions from the column (diluted 1:100 in 50 mM Tris-HCl, pH 7.5, containing 0.15 M NaCl [TBS], 0.1% BSA), washed, incubated with mouse anti-FLAG-peroxidase conjugate (MilliporeSigma, 1:2000), washed, and peroxidase developed with TMB substrate (Thermo Fisher Scientific) according to the manufacturer's directions. FLAG-immunoreactive fractions were pooled, applied to an anti-FLAG agarose (0.5 mL, Sigma) column, and washed sequentially with (i) 50 mM Tris-HCl containing 0.5 M NaCl, (ii) 50 mM Tris-HCl, pH 7.5, containing 10% glycerol, 10 mM MgCl₂ and 10 mM ATP, and (iii) TBS. The resin was eluted with 10 mL of 3× FLAG peptide (0.1 mg/mL) in TBS. The eluate was concentrated to 1 mL with a 100,000 MW cutoff spin concentrator (Millipore) and applied to a Sephacryl S200HR gel filtration column that was pre-equilibrated with TBS. UV absorbance at 280 nm of the eluate from the column was monitored with a UV monitor (2238 LKB Uvicord SII). FLAG-immunoreactivity in gel filtration fractions was evaluated by a direct ELISA by diluting aliquots from the column 100-fold with 0.05 M NaHCO₃, pH 9.6, and adding 0.1 mL/well of a high bind, polystyrene 96-well ELISA plates (Costar, Corning Inc.). After 90 min at room temperature, the wells were blocked with 1% BSA in TBS, washed three times with TBS, and incubated for 60 min at room temperature with horseradish peroxidase (HRP)-conjugated mouse anti-FLAG monoclonal antibody (clone M2, MilliporeSigma) diluted 1:2000 with TBS-1% BSA. The wells were washed three times with TBS, and peroxidase activity was developed with 0.1 mL per well of tetramethylbenzidine (TMB) substrate (Thermo Fisher Scientific) according to the manufacturer's directions. The reaction was stopped by adding 0.1 mL/well of 2 M H,SO₄, and the absorbance at 450 nm was measured with a microplate reader (iMark, Bio-Rad). The peak fractions containing FLAG-immunoreactive material were pooled and concentrated to 1 mL with a 100,000 MW cutoff spin concentrator (MilliporeSigma).

CryoEM sample preparation and image collection. For cryoEM sample preparation, 3 μ L of purified IgID protein was applied to glow-discharged Quantifoil R 1.2/1.3 gold grids. The grids were blotted with filter paper and then flash-frozen in liquid ethane using FEI Vitrobot Mark IV (Thermo Fisher Scientific). An FEI TF20 cryoEM instrument was used to optimize the freezing conditions. The best grids were obtained with 60 s

glow discharge using air and with the Vitrobot sample chamber set at 8°C temperature, 100% humidity, 6 s blotting time. and 1 blotting force.

CryoEM grids were loaded to a Titan Krios electron microscope (Thermo Fisher Scientific) equipped with a Gatan imaging filter (GIF) Quantum LS and a Gatan K3 Summit direct electron detector. The microscope was operated at 300 kV with the GIF energy-filtering slit width set at 20 eV at superresolution mode, yielding a pixel size of 1.1 Å on the sample level. Movies were recorded with SerialEM at a nominal magnification of $81,000\times$. Defocus was set to -1.8 to $-2.6~\mu m$. The total exposure time for each movie was set to 2 s, fractionated equally into 40 frames, giving a total dosage of \sim 50 e-/Å2 per movie. A total of 2,880 movies were collected for image processing.

Single particle cryoEM reconstruction and model building. The workflow of single-particle analysis is summarized in Fig. S3. Frames in each movie were aligned for drift correction with the GPU-accelerated program MotionCor2 (42), generating two averaged micrographs, one with dose weighting (for particles extraction and final reconstructions) and the other without (used for manually screening, particle picking and defocus determination). The averaged micrographs have a calibrated pixel size of 1.1 $\mbox{\normalfont\AA}$ on the specimen scale. Defocus values of micrographs were determined by CTFFIND4 (43). The following image processing was done using RELION 3.1 (44, 45). A total of 7,479,951 particles were picked initially and extracted with a box size of 180 pixels. After 2D classification, 3,701,614 particles were selected. Some of the 2D classes showed strong features of a 3-fold symmetry (Fig. S2D). Therefore, the C3 symmetry was used in the following 3D classifications and 3D refinements until the symmetry expansion. The selected particles were subjected to 3D classification and 3D refinement generating a construction with a resolution of 3.9 Å. Then skip aligning 3D classification was applied. Three good classes totaling 1,138,462 particles were selected and combined, followed by 3D refinement, then re-extracted from dose-weighted micrographs with a box size of 300 pixels. Another round of 3D local refinement was performed with the re-extracted particles, yielding a final 3.0 Å-resolution reconstruction which was used for model building. Symmetry expansion followed by skip align class3D and local refinement without any symmetry (C1) was applied, resulting in a 3.8 Å-resolution reconstruction with a faint density corresponding to the C-terminal domain. We used a Gaussian filter (width 3) in UCSF Chimera to generate a low-resolution reconstruction to identify it (Fig. S2G). We also performed de novo C1 reconstruction and obtained a 4.0 Å reconstruction. As expected, this resolution was worse than that (3.8 Å) of our previous C1 reconstruction. The resolution of the cryoEM map was estimated based on the gold-standard Fourier shell correlation criterion, FSC = 0.143. Data collection and processing statistics are given in Table S1.

The atomic model of IgID was built and refined manually in COOT (46). The model was further refined using Phenix (47) in real space with secondary structure, Ramachandran, and rotamer restraints. Refinement statistics of the models are summarized in Table S1. AlphaFold2 predictions were run on google Colab. Figures and movies were generated using UCSF Chimera (48) and ChimeraX (49).

IgIJ-His pulldown, western immunoblotting, and mass spectrometry. Parental Fn U112 and Fn in which the chromosomal iglJ gene was replaced with a gene encoding a C-terminal His₁₈ epitope tag were each grown in one L of TSBC with 5% KCl to an optical density of 2. The His-tagged and parental bacteria were processed identically in parallel. The bacteria were pelleted by centrifugation ($5000 \times q$ for 90 min) and washed three times by centrifugation with 50 mM sodium phosphate, pH 7.5, with 5% KCl to remove residual free amines. The washed pellets were resuspended in 35 mL of 50 mM sodium phosphate buffer, 5% KCl, and then freshly prepared dithiobissuccinimidyl propionate (DSP, Lomant's Reagent, Thermo Fisher Scientific) in DMSO was added to achieve a final concentration of 1% DMSO. The bacteria were cross-linked for 45 min while turning end-over-end at room temperature. The reaction was stopped by the addition of 1 M Tris HCl, pH 7.5, to achieve a final concentration of 50 mM. The bacteria were pelleted by centrifugation at $48,000 \times g$ for 30 min and resuspended in 10 mM Tris HCl, pH 8.0, 1% Triton X-100, 0.1% sodium deoxycholate, 0.1% sodium dodecyl sulfate, 0.3 M NaCl, 1 mM PMSF, 1 mM NEM, and EDTA-free protease inhibitor cocktail (1:100, Calbiochem) and benzonase (1:1000). The samples were sonicated with a probe tip sonicator and insoluble material pelleted by centrifugation at $25,000 \times g$ for 60 min at 4°C. The clarified lysates were adjusted to contain 10 mM imidazole and rotated overnight with 0.25 mL Ni-NTA agarose that had been equilibrated with the same buffer (Qiagen). The Ni-NTA resins were washed with 10 mM Tris-HCl, 0.3 M NaCl, 1% Triton X-100, 0.1% deoxycholate, and 0.1% SDS containing (RIPA buffer) 10 mM imidazole, and eluted sequentially with 20 mM, 50 mM, and 250 mM imidazole in the same buffer (8 column volumes of each, i.e., 2 mL collected in 1 mL fractions). IgID immunoreactive material and IgIJ-His were detected by Western immunoblotting in the 250 mM eluates of the IgIJ-His sample but not in the parental sample. Protein in the 250 mM eluates was acetone precipitated by adding 4 volumes of acetone, storing the samples overnight at -20°C, and centrifuging at $10,000 \times g$ for 10 min at 4°C. The pellets were washed twice with 4:1 acetone:water at 4°C, air dried for 30 min at room temperature, and stored at -20°C. Further processing was conducted by the UCLA Proteome Research Center. The pellet was resuspended in 8 M urea, 100 mM Tris-HCl, pH 8.5; reduced with 5 mM tris(2carboxyethyl)phosphine (TCEP); alkylated with 10 mM iodoacetamide, and digested with sequencing-grade trypsin. The peptide mixture was desalted, fractionated on-line using C18 reverse phased chromatography, and analyzed using tandem mass spectrometry on a Q-Exactive mass spectrometer (Thermo Fisher Scientific). Data analysis was performed using IP2 (Integrated Proteomics Applications) against an Fn U112 database (taxid no. 401614) and filtered using a decoy-database estimated false discovery rate of less than 0.01.

IgIH-TwinStrep-His pulldown and western immunoblotting. Parental Fn U112 and Fn in which the chromosomal *igIH* gene was replaced with a gene encoding a C-terminal TwinStrep-His₁₈ dual epitope tag were each grown in 250 mL of TSBC with 5% KCl to an optical density of 2. The epitope-tagged and parental bacteria were processed identically in parallel. The bacteria were pelleted and washed to remove free amines as described above; resuspended in 10 mL of 50 mM sodium phosphate buffer, 5% KCl; cross-linked for 45 min with 1 mM DSP; pelleted; lysed in RIPA buffer, and affinity enriched with Ni-

NTA agarose (0.2 mL resin per sample) as described above. IgID immunoreactive material and IgIH-TS-His were detected by Western immunoblotting in the 50 mM and 250 mM eluates of the IgIH-TS-His sample but not in the nontagged parental sample.

Gene deletion and epitope tagging. The 1 to 1.5 kb upstream and downstream neighbor regions flanking the *igID* or *igIJ* genes were amplified from genomic DNA isolated from Fn U112. The upstream and downstream fragments were joined with overlap extension PCR and cloned into pMP590 for use to generate gene deletion mutants by allelic exchange (19, 50). For the generation of an Fn strain expressing *igIJ*-his in the chromosome, the *igIJ* gene was amplified with a short linker encoding GSGSGSGSGSG and an expression tag of 18 histidine residues at the 3'-end, and the upstream, *igIJ*-his and downstream fragments were joined with overlap extension PCR and cloned into pMP590. Alanine substitutions of igID and igIJ were generated using overlapping extension PCR. For episomal expression, *igID* was amplified from Fn genomic DNA with a FLAG expression tag at the 5'-end and cloned into pFNLTP6 (51) in which the *gro*E promoter was replaced by the *bfr* promoter of *F. tularensis* subsp. *holarctica* live vaccine strain. All plasmid constructs were confirmed by nucleotide sequencing.

T6SS-mediated IgIC secretion. Fn strains were inoculated in TSBC containing 5% KCl at A550 of 0.05 and grown at 37° C, 240 rpm for 14 h. Bacteria were pelleted by centrifugation at 11,000 rpm for 10 min. The supernatant fluid was collected, diluted 32-fold in 50 mM bicarbonate buffer, pH 9.6, and added to a 96-well high binding assay plate (Corning, Kennebunk, ME). After a 2-h incubation, the plate was washed with PBS and blocked with 1% BSA in TBS before incubating with rabbit anti-IgIC or IgID primary antibody (1:1000) for 90 min and subsequently HRP-conjugated goat anti-rabbit IgIG(H+L) (1:2000) for 1 h at room temperature. The enzymatic reaction was developed by the addition of TMB substrate and stopped with 2 M $_{\rm L2}$ SO $_{\rm 4}$ solution according to the manufacturer's directions (Pierce TMB, Thermo Fisher). The absorbance at 450 nm and 570 nm was read using an iMark Microplate Reader (Bio-Rad).

Intracellular growth measurement. Human THP-1 monocytic cells (ATCC TIB-202) were differentiated with PMA for 3 days and infected with Fn strains at a multiplicity of infection ratio of 2:1 (bacteria: cell) for 2 h. The infected monolayer was treated with a culture medium containing 10 μ g/mL gentamicin for 30 min to kill extracellular bacteria, washed with HBSS, and the last wash was replaced with DMEM containing 10% FBS and 0.1 μ g/mL gentamicin. At 1 and 21 h postinfection, the cells were lysed with 1% saponin in PBS for 5 min at 1 h and 21 h postinfection, serially diluted in PBS, and the diluted lysates plated on GCII chocolate agar to determine the number of bacteria in the monolayer.

Phagosome integrity assessment. Phagosomal integrity was assessed by a modification of the method of Checroun et al. (52) as previously described (53). THP-1 monocytic cells were differentiated with PMA on coverslips and infected with Fn for 5 h. The infected monolayer was treated with 4% paraformaldehyde in PBS for 1 min followed by 4% paraformaldehyde in 25 mM HEPES-KOH, pH 7.0, 125 mM potassium acetate, 2.5 mM magnesium acetate (KHM) with 15% sucrose, and 50 μg/mL digitonin for 1 min. The monolayer was washed twice with KHM buffer with 15% sucrose and incubated with rabbit anti-F. tularensis or chicken anti-Fn antibody in KHM buffer with 15% sucrose and 0.1% BSA for 15 min at 37°C. The monolayer was washed twice with KHM buffer with 15% sucrose and fixed for 30 min in 4% paraformaldehyde in 0.075 M sodium phosphate. The monolayer was then washed twice with PBS and permeabilized with 1% Triton X-100 in TBS for 30 min. Intracellular Fn was subsequently stained with appropriate primary and secondary antibodies. Cell and bacterial DNA were stained with DAPI. The coverslips were mounted with Prolong gold antifade medium (Life Sciences) and imaged with an Eclipse TE2000 (Nikon) inverted fluorescence microscope equipped with a SPOT camera and software.

In silico analyses. For phylogenetic tree generation, TssK sequences and IgID sequences of representative bacteria of the various T6SS subtypes (corresponding to the species included in our previously published phylogenetic tree (54)) were downloaded from Uniprot and a phylogenetic tree was generated using the Maximum Likelihood Method by MEGA11 (39).

Consensus sequences of the TssK and IgID NTD were prepared by downloading TssK and IgID sequences from BLAST and aligning the sequences with the MUSCLE multisequence alignment program (https://www.ebi.ac.uk/Tools/msa/muscle/). Outliers were eliminated and consensus sequences were displayed by WebLogo (55).

Data availability. The cryoEM density map has been deposited in the Electron Microscopy Data Bank under accession codes EMD-27656. The atomic coordinate has been deposited in the Protein Data Bank under accession code 8DOL.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, JPG file, 0.3 MB.

FIG S2, JPG file, 0.6 MB.

FIG S3, JPG file, 0.8 MB.

FIG S4, JPG file, 2 MB.

FIG S5, JPG file, 0.8 MB.

FIG S6, JPG file, 0.1 MB.

FIG S7, JPG file, 0.4 MB.

FIG S8, JPG file, 1.5 MB.

FIG S9, JPG file, 0.2 MB.

TABLE S1, PDF file, 0.04 MB.

ACKNOWLEDGMENTS

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M.A.H. and Z.H.Z. conceived and supervised the project. D.L.C. and B.-Y.L. prepared IgID sample; X.L. and X.Y. made cryoEM grids, performed cryoEM imaging, and data processing; X.L. built the atomic models and illustrated the structures; D.L.C. and B.-Y.L. prepared bacterial strains, performed biochemistry analysis, secretion, phagosomal escape and intracellular growth assays. X.L., D.L.C., B.-Y.L., Z.H.Z., and M.A.H. interpreted the data and wrote the manuscript; and all authors reviewed and approved the paper.

We declare no competing interests.

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