A Mutant with Aberrant Extracellular LcrV-YscF Interactions Fails To Form Pores and Translocate Yop Effector Proteins but Retains the Ability To Trigger Yop Secretion in Response to Host Cell Contact

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The plasmid-encoded type three secretion system (TTSS) of Yersinia spp. is responsible for the delivery of effector proteins into cells of the innate immune system, where these effectors disrupt the target cells’ activity. Successful translocation of effectors into mammalian cells requires Yersinia to both insert a translocon into the host cell membrane and sense contact with host cells. To probe the events necessary for translocation, we investigated protein-protein interactions among TTSS components of the needle-translocon complex using a chemical cross-linking-based approach. We detected extracellular protein complexes containing YscF, LcrV, and YopD that were dependent upon needle formation. The formation of these complexes was evaluated in a secretion-competent but translocation-defective mutant, the YscFD28AD46A strain (expressing YscF with the mutations D28A and D46A). We found that one of the YscF and most of the LcrV and YopD cross-linked complexes were nearly absent in this mutant. Furthermore, the YscFD28AD46A strain did not support YopB insertion into mammalian membranes, supporting the idea that the LcrV tip complex is required for YopB insertion and translocon formation. However, the YscFD28AD46A strain did secrete Yops in the presence of host cells, indicating that a translocation-competent tip complex is not required to sense contact with host cells to trigger Yop secretion. In conclusion, in the absence of cross-linkable LcrV-YscF interactions, translocon insertion is abolished, but Yersinia still retains the ability to sense cell contact.

Type three secretion systems (TTSS) are employed by a number of Gram-negative pathogens to transfer effector proteins from the bacterial cytosol across the plasma membrane of eukaryotic cells, a process called translocation (1). These systems are essential virulence mechanisms for pathogens because they enable the pathogen to interfere with host defenses and thereby establish a replication niche inside a host (2,3). In the pathogenic Yersinia spp., the plasmid-encoded TTSS translocates a set of five to six effector proteins, called Yops, which antagonize the functions of innate immune cells during animal infection (4–7). The Yersinia TTSS resembles a syringe-like apparatus with three distinct parts: the base, which spans both the inner and outer membranes; the needle, which protrudes from the base and forms a hollow tube that is a YscF protein polymer; and the tip complex, which rests at the distal end of the needle (8). The tip complex has been visualized by electron microscopy (EM) (9–11) and appears to be a homopentamer of LcrV (11, 12). This observation has been corroborated by the modeling of an LcrV pentamer onto the tip of the homologous Shigella polymer (9) and by oligomerization studies of LcrV and its Pseudomonas aeruginosa homologue, PcrV, which show that when these proteins are purified, they form a pentameric ring structure (13). The base and needle are sufficient for secretion of Yops into the extracellular environment although the regulation of secretion is altered in the absence of LcrV (14–16). Translocation of effectors across host cell membranes requires LcrV (17–19). In addition to its location at the needle tip, LcrV is secreted and is also found in the bacterial cytosol, where it plays a regulatory role in Yop secretion (14–16, 20,21). For translocation of Yops to occur, Yersinia must insert two proteins, YopB and YopD, into the membranes of targeted cells (1, 22). YopD, but not YopB, has been found in purified needle preparations from Yersinia enterocolitica; however, its location on or within these needles has not be visualized by EM (10, 11). While it has been speculated that the YopD in these complexes is due to secreted but insoluble YopD which copurifies with needles (11, 12), recent work has also shown that YopD copurifies with LcrV-capped needles (10). YopB and YopD form a hetero-oligomeric pore in mammalian plasma membranes, termed the translocon, and are critical for the formation of a pore through which effector proteins are thought to travel (1,22). Purified YopB and YopD can insert into lipid bilayers (23, 24), most likely because of their hydrophobic nature. However, by themselves, they do not form a functional translocon that permits entry of effector proteins into cells (23, 25). Extracellular LcrV is essential for the assembly and insertion of the translocon (11, 26) and has been implicated in determining the channel size of the translocation pore (12, 25, 27). In the absence of LcrV, YopD, but not YopB, still inserts into membranes (12). This suggests an active role for LcrV in the insertion of YopB. Combined, these observations lead to a model in which LcrV interacts with YopB and/or YopD and guides them to form a functional translocon, permitting the translocation of effector Yops. It is currently unclear, however, if the role of LcrV in YopB insertion and/or translocon assembly requires LcrV associ-
Plasmids

table 1 strains and plasmids used in this study

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Plasmids

table 2 primers used in this study

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translocation-defective mutant (30), failed to form YscF-lcrV cross-linked complexes. Also it failed to insert YopB into host cell plasma membranes yet still sensed host cell contact, as indicated by Yop secretion. These results support a model in which an LcrV tip complex is critical for translocon insertion but is not required for Y. pseudotuberculosis to detect the presence of host cells and initiate secretion of Yops.

**MATERIALS AND METHODS**

**Strains and bacterial culture conditions.** Bacterial strains are listed in Table 1. Y. pseudotuberculosis cells were cultured in Luria broth (L-broth) at 26°C overnight with aeration. Unless otherwise indicated, Y. pseudotuberculosis cultures were diluted 1:40 into secretion medium (2X yeast extract-tryptone [YT] medium supplemented with 20 mM Na-oxalate and 20 mM MgCl₂) and incubated at 26°C for 1.5 h with aeration and then at 37°C for 1.5 h with aeration to induce synthesis of the TTSS. Isopropyl β-D-1-thiogalactopyranoside (IPTG; 30 μM) was added at the shift to 37°C to induce expression of yscF when necessary. To monitor the sensitivity of the wild type (WT), the ΔyopN strain, and a strain expressing YscF with the mutations D28A and D46A (YscFD28AD46A) to a range of calcium concentrations, bacteria were grown in 2X YT medium supplemented with 5 mM EGTA and 20 mM MgCl₂, and the medium was supplemented with various concentrations of CaCl₂ as indicated in the figures. To monitor whether Y. pseudotuberculosis secreted Yops in RPMI medium, Y. pseudotuberculosis cells were grown in 2X YT medium plus 5 mM CaCl₂ for 2 h at 26°C and shifted to growth at 37°C, and then the bacteria were gently spun down and resuspended in RPMI medium and grown for 1 h at 37°C. M9 minimal medium supplemented with 0.4% glucose and a 1% solution of alanine, aspartate, cysteine, glutamate, glycine, histidine, isoleucine, leucine, methionine, phenylalanine, proline, serine, threonine, tryptophan, tyrosine, and valine was used in cross-linking experiments as indicated in the figure legends. When cells were grown in M9 medium, the time for secretion induction was increased from 1.5 h to 3 h to compensate for the reduced growth rate.

The pTRC99A-yscF plasmid (29) was used as a template for site-directed mutagenesis to generate the plasmid pTRC99A-yscFD28AD46A. Primers used are listed in Table 2. The lcrV gene and the yscF genes were deleted in the IP2666 Δ5 (IP2666 ΔyopHEMO1) and IP2666 Δ5N (IP2666 ΔypHEMO1N) strain backgrounds by allelic exchange to create IP2666 Δ5F (IP2666 ΔyopHEMO1 ΔyopF), Δ5SF (IP2666 ΔyopHEMO1 ΔyopF), Δ5V (IP2666 ΔyopHEMO1 ΔlcrV), and Δ5NV (IP2666 ΔyopHEMO1N ΔlcrV) strains. pCD442-lcrVKO or pCD442-yscFKO (29) was introduced into Y. pseudotuberculosis as described previously (34).

**Cross-linking of Y. pseudotuberculosis protein.** For cross-linking experiments, Y. pseudotuberculosis cells were washed and resuspended in warm phosphate-buffered saline (PBS) at an optical density at 600 nm (OD600) of 0.5, except where it is indicated in the text that Y. pseudotuberculosis cells were not washed before exposure to cross-linking agents. Fifty microliters of the Y. pseudotuberculosis suspension was subjected to cross-linking with 1 mM bis(sulfosuccinimidyl) suberate (BS3), 1 mM ethylene glycol bis(sulfosuccinimidyl)succinate (sulfo-EGS), or 1 mM 3,3'-dithio-bis(sulfosuccinimidylpropionate) (DTSSP) dissolved in water. Cross-linking reactions were allowed to proceed for 30 min at 37°C. After the cross-linking step, cells were pelleted and resuspended in 2X sample buffer (100 mM Tris, pH 6.8, 4% SDS, 30% glycerol, 1% bromophenol blue). In addition, 4% β-mercaptoethanol (BME) was added to the sample buf-
fer unless DTSSP was used. Proteins were separated by SDS-PAGE, transferred to polyvinylidene difluoride (PVDF) membranes, and visualized by Western blot analysis. Western blotting was carried out using anti-rabbit antibodies to YopB (1:10,000) (gift from J. Bliska, Stony Brook), YopD (1:10,000), LcrV (1:10,000), or YscF (1:15,000). Both Cy3 (1:2500)-conjugated and horseradish peroxidase (HRP; 1:10,000)-conjugated secondary anti-rabbit antibodies (Invitrogen) were used for visualization.

Immunoprecipitation. Y. pseudotuberculosis cells were grown in 2 × YT medium for 1.5 h at 26°C and then shifted to 37°C for an additional 1.5 h with aeration. Cells were treated with cross-linking reagents as described above. Bacteria were then pelleted and resuspended in 100 μl of boiling buffer (50 mM Tris-Cl, pH 7.5, 1 mM EDTA, 1% SDS) and heated for 5 min at 95°C. Lysates were centrifuged to remove insoluble proteins, and supernatants were added to 900 μl of ice cold TNET buffer (50 mM Tris-Cl, pH 7.5, 150 mM NaCl, 5 mM EDTA). SigmaFast protease inhibitor cocktail (1×) was added to each reaction mixture. Fifty microliters of a 1:1 solution of protein A beads in TNET buffer was incubated with 2 μl of anti-LcrV serum and then added to the buffered lysate. Mixtures were allowed to incubate at 4°C for ~18 h, and then the beads were removed, washed in TNET buffer, and eluted by the addition of 2× sample buffer with 4% BME. Samples were heated to 95°C, and proteins were separated by SDS-PAGE and visualized by Western blotting. In some cases, blots were stripped in acid stripping buffer (25 mM glycine, 1% SDS, pH 2.0) for 30 min at room temperature and then reprobed with another antibody.

Secretion of Yops into culture supernatants. Secretion of Yops into the culture supernatant was performed as described previously (29). The secreted proteins were separated by SDS-PAGE and visualized by Coomassie staining. For determination of YopE secretion into tissue culture supernatants of infected cells, HEP-2 cells were infected as described below in RPMI 1640 medium lacking fetal bovine serum (FBS). Following infection, supernatants were removed, clarified by centrifugation, and precipitated with trichloroacetic acid (TCA) as described previously (29). Secreted proteins were separated by SDS-PAGE, and YopE protein was visualized by Western blot analysis with anti-YopE antibody (1:10,000).

Indirect IF microscopy. Y. pseudotuberculosis was grown in Yop secretion medium at 37°C for 90 min, fixed, and analyzed by immunofluorescence (IF) as described previously (37). Micrographs were taken with a Nikon inverted TE2000-U microscope with a Photometrics charge-coupled-device (CCD) camera at a magnification of ×60 using MetaVue software (Molecular Devices, Sunnyvale, CA), DAPI (4′,6′-diamidino-2-phenylindole) and Alexa Fluor 594 were visualized using Nikon UV-2E/C and G-2E/C filters, respectively. Images were pseudocolored and merged using NIS-Elements software.

Tissue culture. HEP-2 cells were maintained in RPMI 1640 medium (Cellgro) with 5% FBS at 37°C and 5% CO₂. For plasma membrane isolation experiments, cells were grown to confluence in T-75 flasks (approximately 1 × 10⁶ cells). For all other tissue culture experiments, HEP-2 cells were seeded into either 6- or 96-well tissue culture treated plates at 6 or 1.5 × 10⁵ cells per well, respectively, 24 h prior to the experiment.

The following procedure for HEP-2 cell infections was used for all tissue culture experiments. Y. pseudotuberculosis cells were gently washed in PBS and resuspended in 37°C RPMI medium with 5% FBS supplemented with 30 μM IPTG. This medium was then used to replace culture medium on HEP-2 cells. Infection was initiated by centrifuging the bacteria onto the cells at 290 × g for 5 min at room temperature (RT). Plates were incubated at 37°C in 5% CO₂ for the remainder of the experiment.

Cell-rounder assay. Cell-rounder experiments were performed as described previously (38). The infection proceeded for 60 min before imaging. Cells were examined on a Nikon Eclipse TE2000-U microscope (Melville, NY).

Plasma membrane isolation and cross-linking. Y. pseudotuberculosis cells were grown in 2 × YT medium at 26°C for 2 h, 30 μM IPTG was then added, and the bacteria were shifted to 37°C for 2 h. A total of 5 × 10⁹ Y. pseudotuberculosis cells were gently washed in warm PBS and resuspended in 5 ml of RPMI medium containing 30 μM IPTG. HEP-2 cells were grown to confluence of approximately 1 × 10⁵ cells in a T-75 flask and washed gently with warm PBS. Five milliliters of RPMI medium containing Y. pseudotuberculosis and IPTG was added to the confluent flask, resulting in a multiplicity of infection (MOI) of ~50:1. Flasks were incubated without centrifugation for 1 h at 37°C in 5% CO₂, after which time flasks were placed on ice for 5 min. The medium was removed from the flasks, the monolayer was gently washed with cold PBS, and the cells were removed by scraping into 5 ml of cold PBS. Cells were centrifuged at 290 × g for 5 min, resuspended in ice-cold swelling buffer (1 mM EGTA, 1 mM MgCl₂, and 1 mM dithiothreitol [DTT]), and allowed to sit on ice for 5 min. After swaying, cells were again pelleted at 290 × g for 5 min and resuspended in 1 ml of sucrose buffer (250 mM sucrose, 3 mM imidazole, pH 7.4) and 1× SigmaFast protease inhibitor cocktail. The cells were transferred to a Dounce homogenizer and stroked gently 10 times with the loose pestle. Cells lysates were then transferred to a 15 ml conical tube and centrifuged at 900 × g for 5 min to pellet insoluble material and bacteria. Supernatants were collected, and pellets were washed with 1 ml of sucrose buffer and centrifuged at 900 × g for 10 min. Supernatants were combined, transferred to a 2-ml Eppendorf tube, and centrifuged at 14,000 × g to remove any remaining insoluble material. For cross-linked samples, a 250 mM stock solution of 1,5-difluoro-2,4-dinitrobenzene (DFDNB) in dimethylformamide (DMF) was prepared and added to the lysate at a final concentration of 500 μM. Samples were incubated on ice for 2 h before being quenched with the addition of 25 mM Tris, pH 8. Both cross-linked and nontreated samples were centrifuged at 125,000 × g to pellet membrane vesicles. Pellets were resuspended in 2× sample buffer with 4% BME and separated by SDS-PAGE. Proteins were visualized by Western blotting.

RESULTS

HMW complexes of LcrV and YopD, but not YopB, can be detected in the presence of Y. pseudotuberculosis needles. To probe for protein-protein interactions occurring between YscF, LcrV, YopB, and YopD, we established a chemical cross-linking assay using amine-to-amine cross-linkers. In order to eliminate complications from potential cross-links between effector Yops and TTSS components, a Y. pseudotuberculosis strain that lacks five effector Yops, the YopHEMOJ strain (Δ5) was used. Y. pseudotuberculosis cells grown under Yop-secreting conditions were washed gently in PBS and exposed to cross-linkers. Cells were lysed, and YscF, LcrV, YopD, or YopB was analyzed by Western blotting for cross-linked complexes. Sulfot-EGS, a 16.1-A, non-membrane-permeable NH₃-NH₃ cross-linker, generated reproducible cross-links containing YscF, LcrV, or YopD (Fig. 1A to C). High-molecular-weight (HMW) species were detected that correspond to multimers of LcrV or LcrV with other proteins (Fig. 1B). One particular LcrV-containing band corresponds to the molecular weight of an LcrV-YscF cross-link (Fig. 1A). A band of similar molecular weight was also recognized by the YscF antibody (Fig. 1B), supporting the idea that this band represents an association between surface-localized LcrV and YscF proteins. The LcrV complexes were detected only when YscF was present (Fig. 1A). This indicates that formation of these complexes requires the presence of needles and suggests that that these complexes are both extracellular and associated with Y. pseudotuberculosis. In addition, HMW cross-linked protein complexes containing YopD were also detected when Y. pseudotuberculosis was grown under secretion-inducing conditions (Fig. 1C). These multimers were dependent on both YscF and LcrV, indicating that external YopD complexes may require both needle formation and tip complex formation in order to form. We were unable to ob-
serve HMW structures containing YopB with this or other crosslinkers (Fig. 1D and data not shown), suggesting either that YopB does not associate on the surface of Y. pseudotuberculosis under Yop-secreting conditions or that YopB cannot be cross-linked by these cross-linkers. In summary, the sulfo-EGS cross-linker provides us with a tool to investigate whether the putative YscF-LcrV and/or the HMW complexes of LcrV and YopD are formed when TTSS proteins with various mutations are expressed.

To confirm that the 46-kDa band was a cross-linked species consisting of LcrV and YscF (Fig. 1A and B), we exposed Y. pseudotuberculosis to a membrane-impermeable, amine-to-amine, cleavable cross-linker, DTSSP (12 Å). A 46-kDa band was detected by Western blotting with DTSSP (see Fig. 1A; also data not shown). Cross-linked LcrV and YscF were immunoprecipitated, the cross-links were cleaved with BME, and the immunoprecipitates were probed by Western blotting (Fig. 2A and B). We found that YscF was detected in immunoprecipitates of cross-linked LcrV and, likewise, that LcrV was detected in the YscF immunoprecipitates, confirming that LcrV and YscF were directly cross-linked to each other (Fig. 2A and B).

We next investigated whether LcrV associates with YopD or YopB under Yop-secreting conditions. After chemical cross-linking with sulfo-EGS, LcrV was immunoprecipitated and analyzed by Western blotting. The LcrV-YscF and one other HMW LcrV-containing complex were immunoprecipitated (Fig. 2C, frame i), consistent with HMW complexes observed in whole bacterial lysates. Our inability to detect the highest HMW complex after immunoprecipitation seen in whole-cell lysates (Fig. 1A) may be because the LcrV antibody is unable to immunoprecipitate the complex, or because its low abundance and HMW make it difficult to visualize by Western blotting. To determine if YopD or YopB was present in the immunoprecipitated complexes, the blot was reprobed with antiserum recognizing YopD or YopB. Neither YopD nor YopB (Fig. 2C, frames ii and iii) was detected in these complexes, suggesting that the HMW complex consists of either multimers of LcrV or LcrV associated with other secreted proteins. Given the observations that LcrV forms HMW complexes only when needles are generated, that LcrV cross-links with YscF, and that a pentamer of LcrV is found on the needle tip (11, 12), it is plausible that these HMW LcrV complexes are cross-linked tip complexes.

The translocation-defective YscFD28AD46A strain secretes Yops in a calcium-regulated manner. Having established that LcrV cross-linked to YscF and that secreted LcrV and YopD form HMW complexes in association with Y. pseudotuberculosis, we investigated whether a translocation-defective mutant of Yersinia had altered tip complex formation or altered YopD complex formation by studying the cross-linking patterns of these proteins. A yscF mutant, the yscFD28AD46A strain, which was originally isolated and characterized by Torruellas et al. in Y. pestis (30), was unable to translocate Yops into mammalian cells. This yscF mutant in Y. pestis was also unresponsive to calcium since Yops were secreted into supernatants at calcium concentrations up to 7 mM (30). However, it formed typical needle polymers as measured by BS2 cross-linking. To investigate the phenotypes of this mutation in Y. pseudotuberculosis, we expressed yscFD28AD46A in a ΔyopN strain (29). Surprisingly, when expressed in Y. pseudotuberculosis, the yscFD28AD46A-expressing strain was still sensitive to calcium (Fig. 3A) as the yscFD28AD46A mutant strain secreted WT levels of Yops when it was grown in a calcium-free medium (Fig. 3A). When calcium was added back to the medium, both the WT and the yscFD28AD46A-expressing strains stopped secreting detectable levels of Yops in the presence of 2.5 mM CaCl2, whereas the ΔyopN mutant continued to secrete Yops in the presence of 5.0 mM CaCl2.

![Image](http://jb.asm.org/)

**FIG 2** YscF and LcrV can be immunoprecipitated (IP) after cross-linking. (A and B) A WT, ΔyscF (ΔF), or ΔlcrV (ΔV) Y. pseudotuberculosis strain was exposed to DTSSP and then lysed. Lysates were split and immunoprecipitated with LcrV or YscF antibodies. Cross-links within the immunoprecipitates were cleaved with BME. Proteins were separated by SDS-PAGE and analyzed by Western blotting with anti-LcrV (A) or anti-YscF (B). (C) LcrV was immunoprecipitated from lysates of Y. pseudotuberculosis exposed to sulfo-EGS with anti-LcrV antibody. Proteins were eluted from protein A beads by boiling in sample buffer. Proteins were analyzed by Western blotting with anti-LcrV (i), anti-YopD (ii), and anti-YopB (iii) antibodies. Each experiment was repeated twice, and representative blots are shown. Asterisks indicate incompletely stripped LcrV. ΔS, YopN ΔyopF, ΔSN, YopN ΔHMOJ ΔlcrV; ΔSF, YopN ΔHMOJ ΔyscF; α, anti.
and 7.5 mM CaCl₂ (Fig. 3A and data not shown). To investigate if Y. pseudotuberculosis expressing YscFD28AD46A formed YscF polymers, Y. pseudotuberculosis was grown in the absence of calcium and exposed to the cross-linker BS3 (Fig. 3B). No differences in the molecular weights of cross-linked complexes were observed between the WT YscF and YscFD28AD46A strains. Interestingly, however, we consistently observed variation in the intensity of specific bands (Fig. 3B, asterisks), suggesting that there may be differences in the polymers formed by this mutant. We also verified that YscF localized to the surface of Y. pseudotuberculosis cells by immunofluorescence (IF) microscopy. In both /H9004 yscF strains expressing WT YscF and expressing YscFD28AD46A, punctate staining was observed on nonpermeabilized Y. pseudotuberculosis cells, consistent with extracellular localization of YscF polymers (Fig. 3C) (37, 38). We next assessed the ability of this mutant to support translocation of YopE using an HEp-2 cell-rounding assay, which detects the activity of the effector YopE on the actin cytoskeleton of translocated cells (39). Notably, the yscFD28AD46A-expressing strain was incapable of causing HEp cell rounding, suggesting that it is unable to translocate YopE (Fig. 3D). Combined, these observations indicate that the YscFD28AD46A mutant in Y. pseudotuberculosis is calcium regulated and generates needles but cannot support Yop translocation into mammalian cells.

Translocation-defective YscFD28AD46A needles associate aberrantly with LcrV. To further probe why needles carrying the YscFD28AD46A mutation fail to translocate Yops, we used IF and chemical cross-linking to investigate whether LcrV was found in association with needles. Both the nonpermeabilized WT yscF- and YscFD28AD46A-expressing strains, pYscFD28_46A, pTRC99A-yscFD28AD46A, were grown under secretion-inducing conditions for 1.5 h and then fixed and mounted onto coverslips. Cells were labeled with anti-YscF antibody and visualized with Alexa Fluor-594-conjugated anti-rabbit secondary (red). Bacteria were counterstained with DAPI (blue). Frame i, ΔyscF(pTRC99A-yscF); frame ii, ΔyscF(pTRC99A-yscFD28AD46A); frame iii, ΔyscF(pTRC99A). Each experiment was repeated a minimum of two times. Representative experiments are shown for each.

FIG 3 Characterization of the YscFD28AD46A needle in Y. pseudotuberculosis. (A) The WT IP2666, ΔyopN, or ΔyscF strain carrying pTRC99A (pTRC), pTRC99A-yscF (pYscF), or pTRC99A-yscFD28AD46A (pYscFD28_46A) was grown in 2X YT medium plus 5 mM EGTA and 20 mM MgCl₂ supplemented with the indicated amounts of CaCl₂ for 2 h at 26°C. Expression of YscF from pTRC99A was induced with the addition of IPTG when bacteria were shifted to 37°C. Supernatants from cultures grown for 2 h at 37°C were precipitated with TCA. Secreted proteins were separated by SDS-PAGE and visualized by Coomassie blue. (B) Bacteria were grown under secretion-inducing conditions for 90 min and then exposed to 1 mM BS3 or water for 30 min. Whole cells were solubilized in sample buffer and subjected to Western blot analysis with antibody against YscF. Asterisks indicate HMW bands that vary in intensity between the WT YscF- and YscFD28AD46A-expressing strains. pYscFD28_46A, pTRC99A-yscFD28AD46A. (C) Y. pseudotuberculosis cells were grown under secretion-inducing conditions for 1.5 h and then fixed and mounted onto coverslips. Cells were labeled with anti-YscF antibody and visualized with Alexa Fluor-594-conjugated anti-rabbit secondary (red). Bacteria were counterstained with DAPI (blue). Frame i, ΔyscF(pTRC99A-yscF); frame ii, ΔyscF(pTRC99A-yscFD28AD46A); frame iii, ΔyscF(pTRC99A). (D) HEp-2 cells were seeded into 96-well plates and then infected with Y. pseudotuberculosis at an MOI of 50:1. Images were taken after 1 h of incubation at 37°C. Frame i, ΔyscF(pTRC99A-yscF); frame ii, ΔyscF(pTRC99A-yscFD28AD46A); frame iii, ΔyscF(pTRC99A). Each experiment was repeated a minimum of two times. Representative experiments are shown for each.
revealed several notable differences between WT yscF-expressing and yscFD28AD46A-expressing Y. pseudotuberculosis. When yscFD28AD46A-expressing cells were exposed to sulfo-EGS, there was a stark reduction in the amount of the LcrV-YscF cross-linked product and in the HMW complexes containing LcrV compared to levels in Y. pseudotuberculosis expressing WT YscF (Fig. 4B to E). Furthermore, while YopD was secreted (Fig. 3A), we detected almost no HMW complexes containing YopD associated with yscFD28AD46A-expressing bacteria (Fig. 4D). These observations, combined with the IF results, indicate that although LcrV is secreted and detected on the surface of the mutant, the YscFD28AD46A needles have an altered association with LcrV. Furthermore, this altered association interferes with the formation of the LcrV-YscF cross-link and with HMW forms of LcrV and YopD with associated Y. pseudotuberculosis.

To test if the conformation of the LcrV cross-linked complexes had been only modestly altered such that sulfo-EGS was no longer able to cross-link LcrV to YscFD28AD46A and if another cross-linker might still reveal an association, we examined whether LcrV-YscFD28AD46A cross-links were detectable using DTSSP. The WT yscF-expressing strain had a strong LcrV-YscF cross-link with DTSSP. However, no LcrV-YscF cross-linked product was observed in the strain expressing yscFD28AD46A (Fig. 5A). This indicates that the interaction between LcrV and YscFD28AD46A polymers is altered enough so that it cannot be detected by two different cross-linkers. Curiously, the HMW LcrV complexes that were detected with sulfo-EGS (Fig. 1) were not readily detected with DTSSP (Fig. 5), indicating that these complexes cannot be detected with a shorter cross-link arm. However, a different cross-linked product of approximately 55 kDa was observed (Fig. 5A), which could be an LcrV dimer or LcrV cross-linked with another protein.

We found that the LcrV-YscF cross-linked product was not detected with either sulfo-EGS or DTSSP, but LcrV was detected...
in a punctate pattern on the surface of *Y. pseudotuberculosis* cells by IF. Therefore, we investigated whether our manipulations prior to cross-linking dislodged a tip complex that may have a weaker association with the YscFD28AD46A needle and be washed away prior to exposure to a cross-linker. To address this possibility, *Y. pseudotuberculosis* was grown in a minimal medium in the absence of the amino acids that quench sulfo-EGS, i.e., lysine, glutamine, arginine, and asparagine, and then *Y. pseudotuberculosis* was exposed to sulfo-EGS without a prior washing step. Again, under these conditions, an LcrV-YscF cross-link was detected in the strain carrying a WT YscF allele but was not detected in the strain carrying the yscFD28AD46A allele (Fig. 5B). In summary, these results suggest that the YscFD28AD46A needles still associate with LcrV but that the conformation of the needle-tip complex is sufficiently different to change the availability of the lysines in YscF and/or LcrV for cross-linking with two different reagents. Furthermore, the YscFD28AD46A needles are unable to support the formation of the HMW complexes containing LcrV and YopD.

**YscFD28AD46A needles target YopB and YopD aberrantly to the plasma membrane of mammalian cells, phenocopying a ΔlcrV mutant.** LcrV is critical for YopB and YopD to form productive pores in mammalian cell membranes (12). However, how LcrV facilitates the insertion of the translocon is unknown. The observation that YscFD28AD46A needles have an altered association with LcrV, coupled with the observation that this mutant fails to translocate Yops, suggested that the association of LcrV with YscF may be crucial for insertion of YopB and/or YopD into host cell membranes. To test whether the yscFD28AD46A strain could target YopB and YopD to the plasma membrane of infected cells, plasma membranes were purified from infected cells, and the amount of YopD, YopB, and LcrV associated with the membrane was measured by Western blotting (Fig. 6). For these assays, a strain of *Y. pseudotuberculosis* that lacks the five effectors and the regulatory protein YopN (ΔSN strain) was used because it was previously reported to increase the detectable amount of membrane-associated pore-forming proteins in sheep red blood cells (SRBCs) (12, 23). This phenomenon was also observed for our *Y. pseudotuberculosis* ΔSN strain after infection of HEp-2 cells (unpublished observation). YopB, YopD, and LcrV copurified with plasma membranes of HEp-2 cells infected with the ΔSNF strain expressing WT *yscF* (Fig. 6A). In contrast, plasma membranes from HEp-2 cells infected with the yscFD28AD46A-expressing strain did not contain YopB, and the amount of YopD and LcrV associated with these membranes was reduced (Fig. 6A). The reduction of YopD and the absence of YopB in plasma membranes from cells infected with YscFD28AD46A strain represented the same phenotype as that seen with strains lacking LcrV (Fig. 6B) (12). In summary, these results suggest that the altered association of LcrV with YscFD28AD46A needles leads to a failure to insert YopB into plasma membranes and defective translocon formation, which accounts for the translocation-negative phenotype observed in this strain.

**Plasma membrane-associated LcrV oligomerizes despite aberrant interactions with YscFD28AD46A needles.** Since LcrV was associated with plasma membranes from HEp-2 cells infected with either *yscF* or yscFD28AD46A-expressing *Y. pseudotuberculosis* (Fig. 6), we asked whether LcrV was oligomerized when it associated with membranes. To address this question, the plasma membranes of HEp-2 cells infected with *Y. pseudotuberculosis* harboring either the WT *yscF* allele or the mutant yscFD28AD46A allele were collected following exposure to DFDNB, a 3-Å amine-to-amine cross-linker. After the membranes were concentrated, the presence of HMW complexes of LcrV was assessed by Western blotting (Fig. 5). DFDNB was used because only very weak cross-linked products were detected with sulfo-EGS (data not shown). At least four HMW LcrV-containing complexes were observed in both strains (Fig. 7, asterisks). Interestingly, the LcrV-YscF cross-link was not detected with plasma membrane-associated LcrV. This suggests that this pool of LcrV is no longer associated with the needle or that DFDNB cannot cross-link LcrV with YscF or that the LcrV-YscF cross-link band is obscured by a comigrating band at 46 kDa. In summary, these results indicate that LcrV can form HMW complexes in association with mammalian membranes despite having abnormal association with needles when it is secreted from a YscFD28AD46A strain. These HMW LcrV complexes, however, are not sufficient to support YopB insertion into the membrane, but rather an association of LcrV with the YscF polymer appears necessary for proper insertion of YopB.

**Yop secretion in response to cell contact is triggered in the yscFD28AD46A mutant.** It is currently unknown what signal initiates effector secretion from *Y. pseudotuberculosis* in the presence of host cells. Possibilities include that the tip complex may directly sense host contact or may interact with the YopB/YopD pore, leading to a conformational shift that is propagated through the needle (1, 9, 30). Since LcrV has an aberrant association with polymers of YscFD28AD46A based on cross-linking results (Fig. 4B), the study of this YscF mutant may indicate whether a normal tip complex, as defined by the association of YscF and LcrV detected by chemical cross-linking, is needed to trigger Yop secretion in the

![FIG 6](http://jb.asm.org/) Association of LcrV, YopB, and YopD with HEp-2 plasma membranes after infection with the WT YscF or YscFD28AD46A strain. HEp-2 cells were infected with *Y. pseudotuberculosis* strains at an MOI of 50:1. After 1 h, the plasma membranes were collected, and the proteins were separated by SDS-PAGE. Proteins were visualized by Western blot analysis with anti-YopB, anti-LcrV, anti-YopD, and anti-Na/K ATPase antibodies. (A) Strains lacking *yopHEMOJN yscF* (ΔSNF) and expressing pYscF, pYscFD28AD46A, or pTRC99A were induced in 30 μM IPTG at 37°C. (B) Western blot analysis of the *ΔyopHEMOJN* (ΔSN), ΔyopHEMOJN ΔlcrV (ΔSNV), and ΔyopHEMOJN ΔyscF (ΔSNF) strains. The experiments were performed twice, and a representative blot is shown.
presence of host cells. To determine if the yscFD28AD46A strain initiated Yop secretion after exposure to mammalian cells, HEp-2 cells were infected with *Y. pseudotuberculosis* expressing either WT yscF or yscFD28AD46A. After 1 h, the proteins secreted into the tissue culture supernatants were collected, and the presence of YopE was detected by Western blot analysis. In the absence of HEp-2 cells, YopE was detected only in the medium containing constitutively secreting HEp-2 cells, YopE was detected only in the medium containing calcium plus 5 mM CaCl₂ for 2 h at 26°C and then grown at 37°C for 2 h. Strains carrying pTRC99A or plasmid derivatives were exposed to IPTG to induce expression of YscF. (A) HEp-2 cells were infected at an MOI of 50:1 (+). Equivalent numbers of bacteria were added to wells lacking HEp-2 cells (−). After a 1-h incubation at 37°C, supernatants were collected, and proteins were analyzed by TCA precipitation and separated by SDS-PAGE. Proteins were visualized by Western blot analysis with anti-YopE antibody. (B) After growth at 37°C, bacteria were gently spun and grown for an additional hour in RPMI medium. Supernatants were collected, and proteins were precipitated with TCA, separated by SDS-PAGE, and visualized by Coomassie blue dye. Each experiment was repeated at least twice, and a representative blot is shown.

**DISCUSSION**

The translocation of effectors across the membrane of target cells during infection is an essential step in *Yersinia* pathogenesis; however, the mechanisms that drive this process have remained poorly understood (8). One particularly enigmatic question has been whether the tip complex is involved in host cell sensing and acts as the trigger for effector release from the bacteria. A second question is whether contact between the tip complex and the needle is required to form a pore. Continuous contact between the two would support the idea that a conduit is formed through which translocating effectors traverse. In order to answer these questions, assessment of the tip complex is necessary. Previous elegant methods to analyze the tip complex have involved using EM reconstructions of purified needle complexes (11, 13). However, this assay is technically challenging and requires specialized equipment, so it is not amenable to a high-throughput approach necessary to measure the effects of various mutations and/or conditions. In addition, purification of needle complexes may disrupt weak interactions or fail to reveal transient interactions that may occur among components. Therefore, we developed a simple biochemical cross-linking assay to evaluate the state of the LcrV, YscF, YopB, and YopD. With this assay we demonstrated that we could observe cross-links of LcrV to YscF. Furthermore, we found that LcrV formed HMW complexes that are dependent upon YscF and are consistent with multimers of LcrV, suggesting that these HMW LcrV complexes represent the tip complex. In addition, the observation that no YopB-YscF cross-linked complexes were detected indicates that not all TTSS substrates passing through the needle are cross-linked with YscF. This suggests that the interaction between YscF and LcrV is specific and does not arise during its transit through the needle.

Interestingly, HMW YopD-containing complexes formed in an YscF- and LcrV-dependent manner, indicating that they are extracellular but associated with *Yersinia*. Veenendaal et al. demonstrated that both IpaB and IpaC (orthologues of YopB and YopD, respectively) are localized to the tips of purified *Shigella flexneri* needles in an IpaD-dependent manner (IpaD is an orthologue of LcrV in *Shigella*) (40). In contrast, YopB and YopD have not been observed on the *Yersinia* needle tip. While YopD, but not YopB, has been detected in preparations of purified needle complexes, this copurification may be due to insoluble YopD (11, 12). We were unable to demonstrate a direct interaction between LcrV and YopD even though the YopD HMW complexes were dependent on LcrV. Therefore, LcrV may transiently interact with YopD to facilitate formation of these complexes, and we may have been unable to catch these transient interactions in our cross-linking studies. Alternatively, our inability to detect HMW YopD-con-
taining complexes in the ΔlcrV strain may be due to the fact that less YopD is secreted in the absence of LcrV (41), and, hence, its concentration may not have been sufficient to form multimers. Finally, these YopD HMW complexes may arise from secreted, insoluble forms of YopD that may associate with the bacteria. Since purified YopD and LcrV interact with each other (42), we favor the idea that LcrV directly interacts with YopD extracellularly on Y. pseudotuberculosis cells but in a manner that our cross-linker was unable to capture.

No YopB-associated proteins were detected with the cross-linker used in this study. The lack of cross-linked YopB complexes on the bacterial surface is consistent with the observation that YopB is not associated with purified needle (11). Furthermore, this result suggests that any transient YopB interactions with the needle complex cannot be cross-linked by the amine-to-amine cross-linker used here and that not all putative type III secretion substrates can be cross-linked with YscF and/or LcrV. If YopB is not associated with needle-tip complexes, then YopB acts differently than IpaB from Shigella, which localizes at the tip prior to host cell contact (43). In fact, IpaB is proposed to be a member of the tip complex itself with four monomers of IpaD and one monomer of IpaB (43). However, it is not the case in Yersinia, where LcrV exists as a homopentamer both by modeling predictions (9) and experimental evaluation (12, 13). On the other hand, since LcrV is involved in the insertion of YopB into membranes (12), it seems plausible that an interaction between YopB and LcrV occurs but that this interaction is unstable and transient, that it occurs only upon contact with the plasma membrane, and/or that it could not be detected with sulfo-EGS. The YscFD28AD46A mutant may be defective in interacting with YopB for several reasons. The polymer itself may be defective in harboring and/or inserting YopB into the membrane, or YopB may require an YscF-LcrV interaction prior to insertion into the membrane. For example, the YscF-LcrV interaction may be critical for a particular YopB interaction, which, in turn, may be necessary for YopB placement in the membrane.

The existence of mutations in yscF that lead to a calcium-blind phenotype in Yersinia and of mutations in MxiH, the Shigella ortholog of YscF, that lead to Congo red insensitivity in Shigella, suggests that the needle is involved in sensing the environment (29, 30, 44). One such mutant in Y. pestis, the yscFD28AD46A strain, was unable to sense calcium, leading to constitutive secretion of IpaC to the tip upon induction of secretion (47). Roehrich et al. demonstrated that a small deletion mutation in the C terminus of IpaB (ipaBΔ3) leads to constitutive secretion and recruitment of IpaC into the membrane (43). In Y. pestis, we observed that the YscFD28AD46A mutant fails to interact with the tip complex in a manner that can be detected by cross-linking with sulfo-EGS. If sensing cell contact occurs by the tip complex or by YopB, then we would predict that this mutant may be unable to secrete Yops in response to cells. Astonishingly, we found that this mutant readily secretes Yops in response to cell contact, indicating that neither a normal tip complex nor YopB insertion into the plasma membrane is required for sensing host cells. The LcrV at the YscFD28AD46A polymer tip may still retain the ability to sense cell contact and propagate this message, even though it no longer retains the ability to facilitate YopB insertion. Alternatively, the mechanism for sensing cell-contact may be LcrV independent in Yersinia.

One critical function of LcrV is to mediate organization of the bacterial membrane. Deane et al. (9) proposed a model whereby the needle polymer undergoes a conformational shift upon induction of secretion. In support of this model, they showed that mutations that interfere with regulatory mechanisms governing secretion lie in regions of MxiH required for monomer-monomer interactions (9). It has been hypothesized that the trigger for a conformational change in the needle polymer comes from a conformational change that occurs in the IpaB protein, a component of the Shigella tip complex, presumably in response to cell contact (47).
able to form an interaction with the tip complex and form HMW LcrV complexes detectable by chemical cross-linking still retains the ability to sense contact with host cells. However, it cannot insert YopB into the membrane and generate a functional translocon. Therefore, a normal tip complex appears to be critical for pore formation. Additional work will elucidate the mechanism by which Yersinia senses contact with host cells. However, whereas IpAB is required for sensing cells in Shigella, insertion of the Yersinia ortholog, YopB, is not required for sensing of cells by Yersinia, suggesting that the cell-sensing mechanisms used by these two bacteria may differ.

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