Using Integrated Bioinformatics Strategy to Identify Critical Factors for the Structural Integrity of Salmonella T3SS

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Abstract

Type-III secretion system of Gram-negative bacteria is the major molecular machine responsible for the infection of host cells and the in-host survival of the bacteria. The T3SS is composed of three structural components: basal body, needle, and export apparatus. The needle is an extracellular protein complex that recognizes host cells and transports bacterial effector proteins into the host cells. The basal body forms a channel across the bacterial membranes and also provides structural support to the needle. The export apparatus selects effector proteins and initiates the transportation of these proteins. Since these three structural components are formed by specific proteins, abolishing the interaction of these proteins will disrupt the structural integrity of one or more structural components of T3SS, and eventually affect the infection and/or virulence of the bacteria. In this study, we analyzed the sequential, structural, and interactomic features of Salmonella T3SS structural proteins. We found that these structural proteins have abundant short and/or long disordered regions that overlap with other structured/functional regions. We identified critical interaction patterns and hub proteins SipB, SpaO, and SpaS, in the interactome of T3SS structural proteins. We also predicted novel binding motifs for six T3SS structural proteins of which the interaction partners are unknown. These results are expected to shed light on future studies in the fields of T3SS structural integrity and drug discovery.

Keywords: Protein intrinsic disorder, MoRF, Binding motif

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Introduction

Type III Secretion System (T3SS) is a needle-like appendage found on the surfaces of many gram-negative bacteria, such as Salmonella, Escherichia coli (E.coli), Vibrio, Yersinia, and Chlamydia. T3SS facilitates these bacteria to invade host cells and to cause various diseases, including: typhoid fever, food poisoning, diarrhea, plague, and many others[1,2]. A T3SS apparatus measures 60–80 nm in length and 8 nm in diameter with an interior lumen of ~3 nm in diameter. The major function of T3SS is to export specific proteins (effector proteins) from bacterial cytoplasm to the extracellular to invade and to manipulate host cells. For this reason, T3SS is also called injectisome or injectosome. A single bacterium normally expresses one but may have more T3SSs[3,4]. The bacterial genome may contain different clusters of genes that express different types of T3SS. Salmonella contains two clusters of T3SS genes, which are Salmonella Pathogenicity Island 1 (SPI-1) that is responsible for host invasion, and Salmonella Pathogenicity...
Island 2 (SPI-2) that is critical for the survival of the bacterium inside host cell[3]. Two systems are similar to each other, but also present remarkable differences[9]. *Enterohaemorrhagic E. coli* also has two clusters of T3SS genes[73]. However, the functional differences between them are not completely discovered.

T3SSs from various bacterial species share very similar structures. A T3SS appendage contains three structural parts: transmembrane basal body, extracellular needle, and cytoplasmic export apparatus[9]. The structures of the basal body and needle are similar to that of the base and hook of flagellar filament[10], indicating their common origin of evolution. The basal body is integrated with two layers of membrane of bacterium and is thus composed of three portions: a ring integrated with the outer membrane of the bacterium (OM ring), a ring integrated into the inner member of the bacterium (IM ring), and a periplasmic rod connecting both rings. The needle is connected to the basal body and extends to the extracellular part of the bacterium. The needle is a syringe-like structure composed of needle filament, tip, and translocon. The needle filament is composed of many proteins of the same type, which enclose an interior lumen with a diameter of 3 nm to facilitate the secretion of effector proteins[11-15]. The tip of T3SS needle is able to trigger the secretion of effector proteins[13] upon being activated by specific environmental factors, such as contact with specific host cells, temperature, pH value, osmolarity, and many others[16]. The export apparatus is a dynamic complex of multiple proteins connected to the IM ring of T3SS[17,19]. The function of the export apparatus is to control the secretion of effector proteins. After being secreted into host cells, these effector proteins are able to manipulate the host cells in multiple ways[20-23]. More specific examples of host cell manipulation include: inducing the host cell to engulf the bacterium; inducing apoptosis[24] by the interaction between *Shigella flexneri* effector IpA and caspase 1 of host cell[25]; and entering the nucleus of host cell and then activating the expression of genes that are beneficial for bacterial infection as shown in the case of *Xanthomonas* effector TAL[50]. Deletion of proteins in the export apparatus significantly influenced the secretion of other proteins[17,19,27-30] and hence may alter the virulence and ability of affection of the bacteria.

Clearly, T3SS is critical for the pathogenicity of Gram-negative bacteria. Recent studies found that defects of T3SS significantly reduced the ability of infection of bacteria[31-35]. Therefore, disrupting the assembly of T3SS eliminated the virulence, but didn’t kill the bacteria. These observations suggested a new alternative strategy for drug development by focusing on anti-virulence and anti-infective targets[37-40]. This alternative strategy does not add selection pressure towards drug resistance and hence may indicate a solution to the fast developed drug resistance of traditional antibiotics that are normally either bactericidal or bacteriostatic[41], and provide additional clinical treatment of bacterial infection. For these reasons, T3SS has become a very important target for drug development[37]. Several small molecules have been reported to disrupt the assembly and function of T3SS in various model systems[42-44].

To facilitate mechanistic studies on the structural integrity of T3SS and to facilitate the discovery of new drugs that target T3SS structural proteins, thorough understanding on the three-dimensional structures and on the interaction partners of T3SS structural proteins becomes a prerequisite. The T3SS system is normally composed of tens of proteins, which are categorized into two groups: structural proteins that are responsible for the assembly of the basal body and needle of T3SS; and translocator proteins that facilitate the translocation of other bacterial proteins from bacterial cytoplasm to the outside. Many proteins that compose of T3SS are multi-domain proteins. 3D-dimensional structures for some of them have been identified[44-46]. However, not all the T3SS proteins are structured. Intrinsically disordered regions have been found in several individual T3SS proteins, such as *Salmonella* SipD[47], *Ecoli* EspD[48], *Yersinia* LcrV[49], and *Pseudomonas* PopB[50]. These disordered regions are frequently involved in protein-protein interactions[47,48,50-52]. Nonetheless, the overall abundance and functional importance of intrinsic disorder in the T3SS proteins still remains unclear.

In our previous analysis on the distribution of protein intrinsic disorder among over 3,500 species, bacterial proteomes were estimated to have about 20-24% of intrinsic disorder each[53]. In many pathogens, disordered regions are correlated to the virulence[54-56]. Therefore, it becomes interesting to see how much intrinsic disorder is in the T3SS system and what functions it has. In this study, we characterized various sequential, structural, and functional features of the structural proteins of *Salmonella* T3SS, identified their interaction patterns, and predicted novel binding motifs. The results of the study can be used for further studies of identifying new targets and developing new strategies to disrupt the assembly of T3SS and therefore to reduce the infection and virulence of bacteria.

**Method**

**T3SS structural proteins**

Since the proteins that make up T3SS are highly conserved across many different species of Gram-negative bacteria on their sequences, structures, and functions[61-63], we chose T3SS structural proteins of *Salmonella typhimurium* SPI-1 in this study. The number of T3SS structural proteins in *Salmonella* was estimated to be around 20 in previous studies[64]. Among which, several proteins may have general-purpose functions. Therefore, we selected nineteen proteins as listed in Table 1. These nineteen proteins may be split into three groups: (1) Basal body structural proteins: PrgH, PrgK, InvG, InvH, and PrgJ; (2) Needle structural proteins: PrgI, SipD, SipB, and SipC; and (3) Export apparatus structural proteins: InvC, InvI, OrgB, SpaO, InvA, InvE, SpaP, SpaR, SpaQ, and SpaS. The amino acid sequences of these proteins were extracted from UniProt. The 3D structures of these proteins or of segments of these proteins were extracted from PDB.

**Protein-protein interaction network**

Both STRING[64] and DIP[65] databases were used to search for interaction partners of the T3SS structural proteins. STRING is one of the most well maintained and most prevailing databases for protein-protein interactions. We only selected interactions that had been experimentally validated for our further analysis. DIP is another well designed database for experimentally identified protein-protein interactions. The information from DIP was used to supplement the results of STRING.
Table 1: List of Salmonella T3SS structural proteins.

| Structural parts | Protein Name | Length | Function | UniProt ID | PDB ID | PDB Sequence | PDB Coverage |
|------------------|--------------|--------|----------|------------|---------|--------------|-------------|
| Basal body       | PrgH*        | 392    | IM ring  | P41783     | 4G2S; 4G11 | 11-119; 170-392 | 84.7%       |
|                  | PrgK*        | 252    | IM ring  | P41786     | 4W4M; 4OYC | 19-92; 96-200  | 73.8%       |
|                  | InvG         | 562    | OM ring  | P35672     | 4G08    | 22-178       |             |
|                  | InvH         | 147    | OM ring  | P0CL43     | ---     | ---          |             |
|                  | PrgJ         | 101    | Inner rod| P41785     | ---     | ---          |             |
| Needle           | PrgI         | 80     | Filament | P41784     | 3ZQE    | 1-80         | 100%        |
|                  | SipD         | 343    | Tip      | Q56026     | 3NZZ    | 38-110; 132-342 | 82.8%       |
|                  | SipB**       | 593    | Translocon| Q56019    | 3TUL    | 83-115; 127-170; 182-226 | 20.6%     |
|                  | SipC         | 409    | Translocon| P0CL47    | ---     | ---          |             |
| Export apparatus | InvC         | 431    | ATPase   | P0A1B9     | ---     | ---          |             |
|                  | InvI (SpaM)  | 147    | Central stalk | P40612    | ---     | ---          |             |
|                  | OrgB         | 226    | Peripheral stalk | E1WAB4    | ---     | ---          |             |
|                  | SpaO         | 303    | C-ring homolog | P40699    | 4YXA    | 145-213; 232-297 | 44.6%       |
|                  | InvA**       | 685    | Gate     | P0A113     | 2X49    | 357-685      | 47.9%       |
|                  | InvE         | 372    | Gate-keeper | P35671    | ---     | ---          |             |
|                  | SpaR**       | 263    | IM partner| P40701    | ---     | ---          |             |
|                  | SpaQ**       | 86     | IM partner| P0A1L7    | ---     | ---          |             |
|                  | SpaS**       | 356    | ---      | P40702    | 3C01    | 239-345      | 30.0%       |

N.B. (*) Proteins with single trans-membrane segment. (**) Proteins with multiple trans-membrane segments. When proteins have multiple PDB entries, the X-ray structure was selected by firstly the highest sequence coverage and then the highest resolution. The PDB ID column may have more than one PDB IDs when the protein has multiple PDB entries for different segments. A single PDB ID may contain multiple segments of the same protein as shown by the numbers in the PDB Sequence column. In this case, the missing residues between segments were not crystallized and are thus ordered. The values in the PDB Coverage column present the fractions of amino acids included in the PDB structures.

Disorder prediction

PONDR-VLXT[66] and PONDR-FIT[67] were used to predict the structural flexibility of proteins. PONDR-VLXT is very powerful in identifying hydrophobic clusters inside intrinsically disordered regions. PONDR-FIT is one of most accurate predictors for disorder prediction and for providing biologically relevant information[68]. The combination of PONDR-VLXT and PONDR-FIT was used by us to analyze the correlation between protein intrinsic disorder and function in many projects, such as reprogramming factors of pluripotent stem cells[69], structural flexibility and mechanisms for methionine oxidation[70], evolution of P53[71], PTEN interactome[72], virulence factors[73], yeast mitosis factors[74], DBC1[75], Emerin[76], etc. Both of these two predictors take amino acid sequence as input. The output is a list of per-residue scores for all the residues in the sequence. Residues with scores higher than 0.5 are assigned as disordered residues, while residues with scores lower than 0.5 are interpreted as structured residues. Consecutive disordered residues form an Intrinsically Disordered Region (IDR). When all the residues in a sequence are disordered, the entire protein is predicted to be In Intrinsically Disordered Protein (IDP).

Secondary structure prediction

NetSurfP[77] was used to predict the secondary structure of protein sequences. NetSurfP is an ensemble predictor that provides not only the high-quality predictions, but also the reliability of each prediction. In addition to secondary structure, NetSurfP also provides prediction of accessible surface area.

Functional motifs/domains

Pfam[78] was used to search for functional domains of the T3SS structural proteins. We also used MoRF-II[79,80] and ANCHOR[81] to predict potential binding motifs inside each of the T3SS structural proteins. MoRF-II was designed to identify short motifs that locate inside intrinsically disordered regions and transform their conformations from coil to helix upon binding to partners. The binding motifs identified by MoRF-II very frequently overlap with dips in the disorder profile made from disorder scores predicted by PONDR-VLXT. ANCHOR is able to predict highly hydrophobic segments inside disordered regions based on the calculation of interaction energy. These two predictors are able to identify binding motifs of different preferences.

Results

As shown in Table 1, ten out of nineteen T3SS structural proteins don’t have PDB structures. In the rest nine proteins that have PDB structures, four have structures for at least 70% of each of the sequences, and the other five only have structures for less than 50% of each of the full length sequences. Therefore, most of the T3SS structural proteins or protein regions still do not have
experimentally validated structures. To check whether or not those structure-unknown proteins and/or regions are IDPs or IDRs, PONDR-FIT was used to predict all nineteen proteins. The per-residue predictions for all the proteins were presented in Figure 1(a). Curves below dashed lines represent structured regions, of which the 3D structures should be able to obtain through experimental methods. Curves above dashed lines denote IDRs that don’t have rigid 3D structures under physiological conditions. When analyzing the results, the predictions of structure-known proteins and regions were used as a control set to compare with their corresponding experimentally observed structured regions to examine the prediction accuracy of the predictor. Clearly, all the regions that have PDB X-ray structures have been predicted to be structured except ~AA160–240 of SipB and ~AA40–110 of SipD. These two regions possess structures of helical bundle as shown in Figure 1(b). The amino acid sequences corresponding to these two regions are highly charged and hydrophilic. Nonetheless, segments of these two sequences form amphiphilic helices as demonstrated in the figure by the colors on different sides of these helices. These amphiphilic helices use their hydrophobic sides to interact with each other and to form helical bundles. After taking this factor into consideration, it is clear that the results of PONDR-FIT prediction match to the experimental results very well.

Figure 1a
Disorder prediction of nineteen Salmonella T3SS structural proteins. The proteins were organized in three groups: proteins in basal body (upper left panel), proteins in needle (lower left panel), and proteins in export apparatus (right panel). The name of each protein was labeled in the corresponding plot. In all the plots, x-axis is the index of amino acid along the sequence, and y-axis shows the predicted per-residue disorder score generated from PONDR-FIT. The curve in each plot is the disorder profile made from PONDR-FIT predictions for all residues in each protein. Dashed line in the middle of each plot indicates the boundary between disordered (y ≥ 0.5) and structured (y < 0.5) residues. Horizontal bars are categorized by their colors: blue - regions that have PDB structure; gray - Pfam domains; dark green – transmembrane segments; dark red – coiled coils. (b) PDB structures of two needle structural proteins: SipD (PDB id: 3NZZ) and SipB (PDB id: 3TUL). The region from Ala132 to Gln342 of SipD was colored by its secondary structures. The other region of SipD (Gly36-Ser110) was colored by the types of amino acids (white - hydrophobic; red – negatively charged; blue – positively charged; and green - polar). SipB was also colored by the types of amino acids in the same way. In both structures, discontinued regions indicate missing residues in the structure. (c) Combined analysis of disorder predictions from both PONDR-FIT (gray) and PONDR-VLXT (black) for InvE, InvG, and SipC. This plot is amended from (a) and therefore all the other annotations are the same as those in (a).

Disorder prediction also identified other structured regions that don’t have experimentally observed structures. These predicted structured regions can be classified into four groups: (1) Regions with multiple transmembrane segments, such as ~AA20-320 of InvA, ~AA10-210 of SpaP, ~AA10-80 of SpaQ, ~AA10-260 of SpaR, ~AA20-200 of SpaS, and ~AA320-420 of SipB. It has been well realized that solving the structure of transmembrane proteins is really challenging; (2) Regions that are shorter than 50–60 residues, e.g., ~AA180-224 of SipC. Many other proteins also have such short structure-prone regions. These short
structure-prone regions may not have strong-enough hydrophobic interactions to maintain rigid 3D structures\cite{50}; (3) Regions overlapped with Pfam domains, including: ~AA180–320 and ~AA360–420 of InvG, ~AA20–80 and ~AA140–340 of InvC, ~AA40–210 of InvE, and SipC. Further analysis on disorder prediction of both PONDR-FIT and PONDR-VLXT as shown in Figure 1(c) indicated that each of these Pfam domains is a combination of short structural-prone region(s) and long IDR(s). For this reason, although the functional roles of these regions have been characterized, the structures of these domains are still not defined; (4) regions with uncharacterized features and functions, such as ~AA10–AA100 of PrgJ, AA220–360 of InvE, ~AA20–140 of SpaO, and ~AA60–320 of OrgB.

The predicted protein intrinsic disorder is not negligible in the T3SS structural proteins, with 21.2% of disordered residues by PONDR-FIT predictor and 27.3% of disordered residues by PONDR-VLXT predictor. Almost all the proteins have disordered residues at N- and/or C-termini. Disordered regions were also found in the middle of or throughout sequences. InvH and InvI are both disorder-dominant proteins. Although both of them have structure-prone regions, these regions are short and have considerable levels of flexibility as indicated by the values of their disordered scores. Therefore, these two proteins may not form rigid structures under physiological conditions. Another five proteins (SipB, SipC, SipD, InvE, and OrgB) have long IDRs that have at least 30 consecutive disordered residues. In which, the IDRs of SipB and SpaS connect other structured/functional domains, and the IDRs in the rest three proteins form entire N- or C-terminal disordered domains. In addition to long IDRs, short IDRs can be observed in all other proteins, such as inside functional Pfam domains (~AA420–460 of InvG), linking different structured domains (~AA210–230 of SpaO), separating transmembrane segments (~AA100–130 of SpaR), and inside structured domains (~AA220–230 of PrgH).

Figure 2 presents analysis on the abundance of intrinsic disorder in these proteins. Each protein in this figure is described by three bars from left to right representing: length of the protein, length of the longest IDR in that protein, and fraction of disordered residues in that protein. The first quantity shows the dimension of the protein. The other two quantities describe different aspects of disordered content: length of longest IDR shows the dimension of a consecutive segment of disordered residues; fraction of disordered residues shows the total amount of disordered residues in the protein. These two quantities can be combined to characterize the distribution of disordered residues in a protein. For example, PrgJ has 16.3% of disordered residues and its longest IDR has 12 residues. By taking into consideration that PrgJ has 80 residues in total, it can be concluded that almost all of the disordered residues in PrgJ are in the longest disordered region. Another example is SipC, which contains 57.5% of disordered residues and has 90 residues in the longest IDR. Since this protein has 409 residues, it can be expected that this protein may have multiple long IDRs. When measuring the overall abundance of protein intrinsic disorder, seven proteins (InvH, SipD, SipB, SipC, InvI, InvE, and SpaP) out of nineteen contain IDRs longer than 30AA, these seven proteins also have more than 25% of disordered residues in their sequences. In which, SipC has the highest fraction of disordered residue of 57.5%. InvI, InvH, and SipD have more than 40% of disordered residues. In terms of the length of IDR, SipD has the longest IDR of 126 residues. SipB, SipD, and InvE have long IDRs that are near or over 90 residues. In addition, SpaP, InvI, and InvH have long IDRs that have at least 30 disordered residues.

Figure 2: Distribution of predicted protein intrinsic disorder in nineteen structural proteins of Salmonella T3SS. X-axis shows 19 proteins listed in Table 1. Two y-axes were used in the figure. The first y-axis is on the left and shows the number of amino acids in each protein (protein length, dark cyan) or the longest IDR (dark pink). The second y-axis is on the right and presents the fraction of predicted disordered residues in each protein (dark yellow). The long-dashed line corresponds to the 1st Y-axis and equals to 30AA. The short dashed line matches to 25% on the 2nd Y-axis.

Figure 3 shows protein-protein interaction networks of 19 T3SS structural proteins. Out of these 19 proteins, three proteins (SipB, SpaO, and SpaS) have interactions with proteins that do not belong to T3SS, ten (PrgJ, PrgH, PrgJ, PrgK, InvC, InvG, InvA, SpaP, SpaQ, and SpaR) are bound by one or more T3SS structural proteins, the other six (SipC, SipD, InvH, InvE, InvI, and OrgB) do not have any experimentally validated interaction partners. In more details, all of the three proteins in the first group (SipB, SpaO, and SpaS) have multiple interaction partners. Both SipB and SpaS are transmembrane proteins with at least two transmembrane segments (Figure 1(a)). Meanwhile, both of them contain coiled coil(s) (Figure 1(a)). In the second group of which each of the ten proteins interacts with other T3SS proteins, three of them (PrgJ, InvG, and SpaR) may interact each with itself. These three proteins either have PDB structures (PrgJ, and N-terminal part of InvG), or were predicted to be structured (C-terminal of InvG, and SpaR). In addition, nine proteins (PrgJ, InvG, PrgH, PrgJ, PrgK, InvA, SpaP, SpaQ, and SpaR) in the second group interact with SpaS. InvC, another protein in the second group, interacts with SpaO. SpaR also interacts with SpaP and SpaQ. Both SpaP and SpaQ have another common interacting protein FlG, which is not a T3SS structural protein. In the third group that contains six non-interactive proteins, SipC and SipD are needle structural proteins, InvH is basal body structural protein located in the OM ring, the other three proteins (InvE, InvI, and OrgB) are components of export apparatus. In terms of functions, InvH and OrgB don’t have clear functional annotations, and the other four proteins may have PDB structure, Pfam domain, coiled coil, and/or transmembrane segment.
The predicted binding motifs from both MoRF-II and ANCHOR tire sequence or the N-ter of the Pfam domain. The locations of tured domain. Predicted binding motifs are in the N-ter of the en- second half of the N-ter IDR and the first half of the C-ter struc -ure-prone domain, with an identified Pfam domain covering the -tured domain. InvE has an N-ter IDR followed by a C-ter struct -ure. In the export apparatus proteins, four (InvA, InvI, OrgB, and InvA in the export apparatus (right panel)). In all these plots, y-axis shows the index of amino acid along the protein sequence, y-axis presents per-residue disorder scores predicted from both PONDR-FIT and PONDR-VLXT. Curves in gray are disor-der profiles from PONDR-FIT prediction, while curves in black are dis-order profiles from PONDR-VLXT prediction. The dashed lines in the middle are the boundary of disordered (y>=0.5) and structured (y<0.5) residues. Horizontal bars represent regions of specific interests (from top to bottom): dark green – binding motifs predicted by MoRF-II; red – binding motifs predicted by MoRF-II; dark cyan – helical segments predicted by NetSurfP; pink – beta-strands predicted by NetSurfP; gray – Pfam domains; dark red – coiled coils; dark yellow – transmembrane segments; blue – regions with PDB structures.

**Discussions**

T3SS is the major molecular machine responsible for bacterial infection and virulence of Gram-negative bacteria. The T3SS is composed of about 20 proteins by forming three structural sections: basal body, needle, and the export apparatus. To further explore the functional roles of the T3SS structural proteins, we predicted potential binding motifs of these proteins using both MoRF-II and ANCHOR predictors. Eight out of nineteen T3SS structural proteins were found to have predicted binding motifs as shown in Figure 4. InvH is one of the structural proteins in basal body. This protein is predict-ed to be highly flexible with several very short structure-prone segments. It doesn’t have any known structural or functional domains. Both of MoRF-II and ANCHOR identified a binding motif located at ~AA60 and ~AA80, respectively. Secondary structure analysis by NetSurfP showed that these two segments are helices. SipB, SipC, and SipD are needle structural proteins. SipB has multiple predicted binding motifs on/near both ends of its structure-known domain in the N-terminal half of the entire sequence. Since the structured domain is composed of coiled coils, the predicted binding motifs also intersperse on the linkers of those coiled coil segments. SipC has a ~60AA N-ter-inal disordered region, a ~200AA structure-prone domain in the middle, and a ~120AA disordered region at the C-terminal. All the predicted binding motifs locate in the C-terminal region. SipD has a ~40AA N-terminal disordered region, followed by another ~60AA helical bundle, and another ~200AA structured domain at the C-terminal. All the predicted binding motifs are in the N-terminal disordered region and/or the ends of helices in the helical bundle. In the export apparatus proteins, four (InvA, InvE, InvI, and OrgB) have predicted binding motifs. InvA has eight transmembrane segments in the N-terminal half and another structured domain in the C-terminal half. A predicted MoRF motif is right between the transmembrane domain and structured domain. InvE has an N-ter IDR followed by a C-ter structure-prone domain, with an identified Pfam domain covering the second half of the N-ter IDR and the first half of the C-ter struct-ured domain. Predicted binding motifs are in the N-ter of the entire sequence or the N-ter of the Pfam domain. The locations of the predicted binding motifs from both MoRF-II and ANCHOR are consistent to each other. InvI is a short but almost fully disordered protein, with the entire sequence being annotated as a Pfam functional domain. This domain has two coiled coils in the middle. Multiple binding motifs were predicted throughout the entire sequence. OrgB is another unannotated protein. It is com-posed of ~50AA N-ter IDR and a ~180AA C-ter structure-prone domain. A MoRF motif was identified in the N-ter IDR. In brief, T3SS structural proteins have multiple short binding motifs, which can be used to regulate the interaction between them and other proteins. More interestingly, all the six proteins that don’t have interaction partners in protein-protein interaction databases (Figure 3) were predicted to have binding motifs.
needle is responsible for host detection and transport of infection and virulence factors. The basal body constructs a channel across bacterial membranes and provides structural support to the needle. The export apparatus facilitate the selection and transport initiation of various effector proteins. Each of the structural sections is formed by multiple proteins, which are also called T3SS structural proteins. Clearly, interrupting the interaction of structural proteins inside T3SS or manipulating the interaction between T3SS structural proteins and other bacterial proteins may disrupt the structural integrity of T3SS, and thus affect the affection and virulence of the bacteria. Such a strategy provides an alternative way for drug development that is not related to bactericidal or bacteriostatic and hence poses less selective pressure of developing drug resistance on the bacteria. Studies on the structural biology of the T3SS proteins are critical for understanding the structural integrity of T3SS and the manipulation of the T3SS structures.

Not all of the T3SS structural proteins have experimentally observed structures. By analyzing the results of disorder prediction from both PONDOR-FIT and PONDOR-VLXT predictors, we found that the T3SS structural proteins have significant amount of protein intrinsic disorder. The disordered residues stay in both N- and C-termini of each protein, connect various structured and/or functional domains, and form large disordered functional domains that may have more than one hundred residues. These disordered regions may contain various structural motifs, such as coiled coil, helix, and beta-strand, and are critical for intra- and intermolecular interaction[51].

By analyzing the protein-protein interaction networks of all the T3SS structural proteins, we identified several hub proteins in the networks and specific patterns of interaction. Among nineteen T3SS structural proteins, three (SipB, SpaO, and SpaS) have multiple non-T3SS interaction partners. SpaO and SpaS can both interact with FlIG, a non-T3SS protein. SpaP, SpaQ, SpaR, and SpaS also form a closed loop in the protein-protein interaction network. Prgl, InvG, and SpaR are each able to form multimer with itself. Therefore, these results provide a new strategy of selecting critical target for regulating the interactions and assembly of T3SS.

Six out of nineteen T3SS structural proteins don’t have experimentally observed interaction partners. We applied MoRF-II and ANCHOR predictors and identified multiple binding motifs in these six proteins, as well as in another two proteins. The binding motifs are in terminal regions of proteins, in the linker region of structured domains, at the edge of structured domains, or inside long disordered regions. The discovery of these binding motifs provides critical information for the experimental validation of the interaction partners and interaction patterns of these proteins.

Clearly, integrated analysis by combining sequential, structural, interactomic analysis of the T3SS structural proteins revealed the abundance of protein intrinsic disorder in this system, identified specific patterns of protein-protein interaction, and discovered novel binding motifs in multiple T3SS structural proteins. The results are expected to facilitate further studies on the manipulation of inter-molecular interaction, disruption of structural integrity of T3SS, and selection of drug target.

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