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Analysis of cell envelope proteins

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► **To cite this version:**

Mickaël Desvaux, Michel Hébraud. Analysis of cell envelope proteins. Handbook of Listeria monocytogenes, Dongyou Liu ; CRC Press, Taylor and Francis Group, pp.359-393, 2008, Chapter 12. hal-02910867

HAL Id: hal-02910867

<https://hal.inrae.fr/hal-02910867>

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12 Analysis of Cell Envelope Proteins

Mickaël Desvaux and Michel Hébraud

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12.1 INTRODUCTION

From a morphological point of view, the most fundamental dichotomy within prokaryotes (the term “prokaryotes” is used here in its primary etymological sense—that is, single-celled organisms without nuclei as opposed to eukaryotes, without any further phylogenetic considerations¹) is between those bound by a single biological membrane (monoderm prokaryotes)—that is, the cytoplasmic membrane, and those bound by two concentric but topologically different membranes (diderm prokaryotes)—that is, the inner membrane (cytoplasmic membrane) and the asymmetric outer membrane.² In accordance with holistic and teleonomic concepts, organisms are far more than mere collections of genes,^{3,4} and such difference in membrane organization, and thus cell compartmentation, is not trivial but has profound phylogenetic, structural, metabolic, and physiological implications. Based on the most recent advances in biological evolution and megaclassification of organisms,^{5–7} monoderm prokaryotes are regrouped under the term Monodermata (also called Unibacteria), which essentially includes Archaea together with Posibacteria (formerly called Gram-positive bacteria).

It is worth stressing that the term “Gram-positive bacteria” is terminologically ambiguous, especially for researchers interested in aspects related to bacterial cell envelope (e.g., protein secretion or surface proteins).⁸ From its origin, a positive or negative result given by Gram staining method

indicates whether or not bacteria retain the stain respectively. Later on, the difference in staining was related to profound divergence in structural organization of the cell envelope, briefly: (1) a cytoplasmic membrane surrounded by a thick cell wall in Gram-positive bacteria, and (2) a cytoplasmic membrane surrounded by a thin cell wall beneath the outer membrane in Gram-negative bacteria. Molecular analyses further revealed that, contrary to Gram-negative bacteria, Gram-positive bacteria correspond to a phylogenetically coherent grouping of prokaryotes within the domain Bacteria with phylum BXIII Firmicutes (low G+C mole percent) and phylum BXIV Actinobacteria (high G+C mole percent).^{9,10} However, from Gram staining to cell envelope organization to taxonomic grouping, each step represents some approximations, which often result in misleading or incoherent statements in the literature. For example, some members of Firmicutes and Actinobacteria phyla do not retain Gram stain because of (1) the absence of a cell wall (e.g., bacteria from the genus *Mycoplasma*), (2) a too thin cell wall (e.g., some members of the genus *Clostridium*), or (3) the presence of a waxy outer sheath preventing penetration of the stain (e.g., species from the genus *Mycobacterium*).

Inversely, some bacteria not taxonomically related to Gram-positive bacteria retain the Gram stain (e.g., some members of the phylum BIV Deinococcus-Thermus). More confusingly, some bacteria clearly possessing a Gram-negative-like cell envelope architecture are in fact phylogenetically related to the taxonomic group of Gram-positive bacteria (e.g., *Thermotoga maritima* currently classified in phylum BII Thermotogae,¹¹ or *Fusobacterium nucleatum* belonging to phylum BXXI Fusobacteria).^{12,13} Some other phyla regroup bacteria exhibiting both cell envelope structures (Gram-negative-like or Gram-positive-like cell envelope)—for example, BVI Chloroflexi or BVII Thermomicrobia.¹⁴ Even in some deep branches of the phylum Firmicutes, some bacteria clearly exhibit Gram-negative cell envelope ultrastructure (e.g., in genus *Desulfotomaculum*, *Selenomonas*, *Syntrophomonas*, or *Coprothermobacter*).² Therefore, it appears in numerous cases that the term “Gram-positive bacteria” cannot describe at once a particular Gram staining result, cell envelope organization, and taxonomic group; thus, when employing this term it is extremely important to specify what it refers to. Because of fewer terminological ambiguities, the terms “Monodermata” or “monoderm bacteria” will be preferred to describe prokaryotic cells surrounded by a single biological membrane but without any further phylogenetic considerations. For the purpose of the present review, the term “Gram-positive bacteria” will be used to describe bacteria with a cell envelope composed of (1) a cytoplasmic membrane, and (2) a cell-wall composed at least of peptidoglycan.

Listeria species are monoderm bacteria possessing a thick cell wall retaining Gram stain and belonging to phylum Firmicutes, class Bacilli, order Bacillales, and family Listeriaceae,⁹ and as such are Gram-positive bacteria in all meaning of the term. *L. monocytogenes* is undoubtedly the species that has attracted most attention, considering its frequent occurrence in food coupled with a high mortality rate.¹⁵ Still, the genus *Listeria* comprises six species: (1) two pathogenic ones (*L. monocytogenes*, a human pathogen, and *L. ivanovii*, a ruminant pathogen), and (2) four non-pathogenic relatives (*L. innocua*, *L. seeligeri*, *L. welshimeri*, and *L. grayi*).^{16,17} Only two completed *L. monocytogenes* genome sequences are currently available—*L. monocytogenes* 1/2a EGD-e and 4b F2365^{18,19}—but several other strains are being unassembled¹⁸ or sequenced (<http://www.ncbi.nlm.nih.gov/genomes/lproks.cgi>). Among other species, *L. innocua* CLIP11262¹⁹ and *L. welshimeri* SLCC5334²⁰ are the only genomes available, but *L. ivanovii* PAM55, *L. seeligeri* SLCC3954, and *L. grayi* CLIP12515 are currently being sequenced.¹⁷ Since the genomes of *L. monocytogenes* 1/2a F6854 and 4b H7858 are unfinished, some genes cannot be properly identified; also, final assembly of these genomic sequences may reveal homologues at a later date. Because no clear conclusion can be drawn from genomic analysis of unfinished genomes,²¹ this review will only focus on completed genome sequences of *L. monocytogenes* strains.

12.2 PROTEIN SECRETION SYSTEMS

Within the cell envelope, *Listeria* species can exhibit a large variety of proteins; some of them can even interact with the cell surroundings and thus constitute the surfaceome (i.e., the subset of pro-

tein exposed on the bacterial cell surface). It is worth reminding that, on one hand, cell wall is not an impermeable barrier and cell envelope proteins can interact with the environment without ever having a domain that leaves the confine of the cell wall⁸ and that the extracellular milieu can penetrate the cell wall, so proteins do not necessarily need to poke out into the environment.²² On the other hand, protein localization into the cell envelope is no guarantee that it is cell surface exposed *stricto sensu* as proteins can be masked by overlying components such as capsule polymer, for example.⁸ Nevertheless, for the purpose of the present review, cell surface proteins will refer to gene products that are attached to the cell wall and/or cytoplasmic membrane and interacting with the external side, whereas cell envelope proteins will refer to all gene products present within the cell wall and/or the cytoplasmic membrane.

While cell surface proteins are systematically cell envelope proteins, the opposite is not necessarily true (e.g., proteins attached to the cytoplasmic membrane but interacting only with the cytoplasm). Still, all cell surface proteins (and most cell envelope proteins) must be first translocated to the cytoplasmic membrane via a protein secretion system before attaching to membrane or cell wall components and thus remaining in contact with the external side. Concerning the functions of cell envelope proteins, they are extremely diverse, ranging from transporters and enzymes involved in various metabolic pathways (such as carbohydrates, proteins, nucleotides, or lipids), signal transductions, adhesion and colonization determinants, to virulence factors. It is worth stressing that among cell surface proteins, some so-called moonlighting proteins can be present.²³ Such proteins are multifunctional in the sense that they conduct enzymatic and/or nonenzymatic activities, sometimes taking part in widely divergent pathways, especially when present at different subcellular locations. For example, enolase, a cytoplasmic protein normally involved in glycolytic pathways, was found on the listerial cell surface, which can bind to human plasminogen.²⁴

In Didermata (corresponding to Gram-negative bacteria, also called Negibacteria),^{5,25} six major protein secretion systems (numbered from Type I to Type VI, i.e., T1SS to T6SS) are currently recognized and are restricted to these microorganisms.^{26–29} In fact, protein secretion systems are categorized primarily by translocation mechanisms across the outermost lipid bilayer, which corresponds to the outer membrane in diderm bacteria but to the cytoplasmic membrane in monoderm prokaryotes. To date in monoderm bacteria, six systems are described as allowing protein secretion^{30–33}—that is, protein transport from inside to outside cell cytoplasm—namely, (1) the Sec pathway (secretion, TC #3.A.5; TC#: transport classification number),³⁴ (2) the Tat pathway (twin-arginine translocation, TC #2.A.64), (3) the FEA (flagella export apparatus, TC #3.A.6.1), (4) the FPE (fimbriin-protein exporter, TC #3.A.14), (5) the holins (hole-formers, TC#1.E.), and (6) the Wss (WXG100 secretion system, proteins with WXG motif of ~100 residues). To be complete, the MscL family (large conductance mechanosensitive ion channel, TC #1.A.22) and the putative Tad (tight adherence) apparatus could also be added to the list,^{35,36} even though experimental evidence is not currently available in monoderm bacteria. Once translocated by one of these systems, a protein can remain associated to the cell envelope, be released into the extracellular milieu, or be translocated into a host cell.

As depicted in Figure 12.1, identification of protein secretion systems in *Listeria* involved screening of genome coding sequences (CDS) against various databases as well as bibliographic analyses. From there, Sec, Tat, FPE, FEA, holins, and Wss were identified in *L. monocytogenes*³⁷ (Figure 12.2). While some components of these secretion systems have been experimentally investigated, in *Listeria*, protein translocation per se has never been ascertained in any of them yet.

12.2.1 SEC SYSTEM

The presence, remarkable conservation, and essential nature of the Sec translocon in all living cells have given rise to the notion of a general secretory pathway (GSP) but also led to confusing statements in the literature.³⁸ As illustrated in Figure 12.3, all components of Sec translocon are encoded in *L. monocytogenes*. In addition to the SecYEG protein conducting channel, the signal recognition

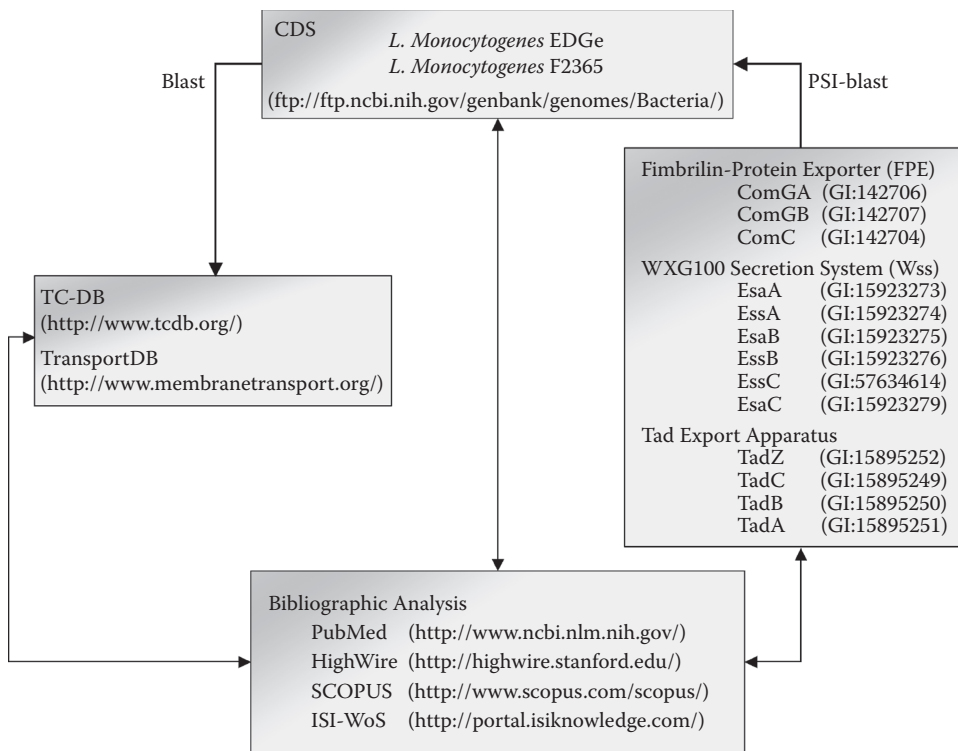


FIGURE 12.1 Genomic identification of protein secretion systems in *Listeria* species.³⁷ Prior to bioinformatic analysis, complete genome, coding sequences (CDS), and original annotation data sets were downloaded from GenBank. Each CDS was screened for the capacity to encode a component of a protein secretion system following BLAST against TCDB¹⁴⁶ and TransportDB.²⁴⁸ These analyses revealed the presence of Sec components and partners as well as FEA subunits, Tat components and holins. MscL and ABC transporter truly implicated in protein secretion could not be identified. The identification of FPE was based on PSI-BLAST searches using GenBank amino acid sequences of ComGA, ComGB, and ComC from *B. subtilis* as queries. Similarly, Wss was identified using EsaA, EssA, EsaB, EssB, EssC, and EsaC from *S. aureus* as amino-acid sequence queries. Using protein sequences of *Clostridium acetobutylicum* as queries,³³ Tad system components could not be identified. Overall, bibliographic analyses were also performed from various databases.

particle (SRP) and the SRP receptor are ubiquitous and essential in all domains of life.³⁹ In *E. coli*, SRP interacts with nascent signal peptide for cotranslational translocation and specific integration of inner membrane proteins, whereas the targeting factor and chaperone SecB interacts with the mature part of the protein and allows post-translational translocation via Sec.⁴⁰ As in all Gram-positive bacteria,⁴¹ SecB and CsaA (analogous to SecB in *B. subtilis*³⁰) are absent from *L. monocytogenes*. In *E. coli*, three auxiliary proteins (SecD, SecF, and YajC) form a transmembrane complex loosely associated with SecYEG and increase the overall efficiency of protein translocation through the cytoplasmic membrane.⁴²

Contrary to SecDF-YajC, the cytosolic ATPase SecA is essential to Sec-dependent translocation in bacteria as it provides the driving force for stepwise export of the protein.⁴³ A SecA paralogue (i.e., SecA2) has been identified in several Gram-positive bacteria including *L. monocytogenes*.⁴⁴ Contrary to *Streptococcus gordonii*, for example,⁴⁵ presence of SecA2 in *L. monocytogenes* is not accompanied by duplication of SecY. While SecA2 is not essential and its relationship with SRP/Sec is unknown, it clearly allows the secretion of a subset of proteins in *L. monocytogenes* (e.g., Iap,⁴⁴ NamA,⁴⁶ and FbpA⁴⁷). Interestingly, the membrane protein FbpA lacks a putative N-terminal signal peptide. As in *B. subtilis*,³⁰ two paralogues of YidC could be identified in *L. monocytogenes*: SpoIIIJ and YqjG.³⁷ In *E. coli*, the polytopic membrane protein YidC is necessary for cotranslational

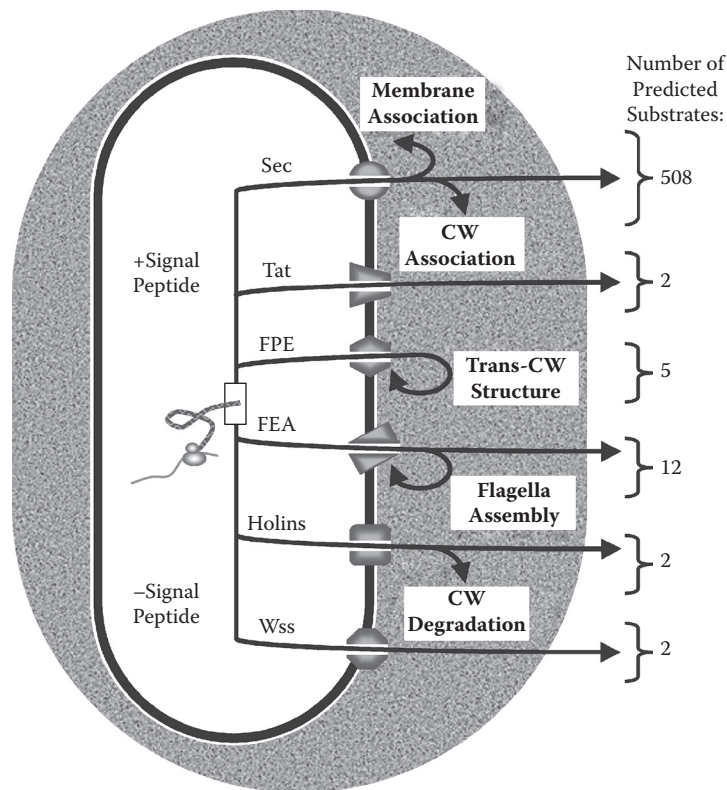


FIGURE 12.2 Schematic overview of protein secretion pathways in *L. monocytogenes* EGD-e.37 Proteins to be translocated can exhibit (+) or not (–) an N-terminal signal peptide (with the exception of Sec pathway, which can translocate proteins with or without signal peptide by alternative mechanisms). Proteins translocated via the Sec pathway remain membrane associated or cell wall associated, are released into the extracellular milieu, or would even be injected into an eukaryotic host cell. Proteins exported via Tat would most certainly be cell surfaced or released into the extracellular milieu. FPE would be involved in the formation of transcell-wall structures. FEA is involved in flagella assembly. Proteins exported by holins seem secreted into the extracellular milieu or involved in cell wall degradation. WXG100 proteins would be secreted into the extracellular milieu. The number of translocated proteins by each pathway is given from most recent estimations. CW, cell wall; Sec, secretion; FPE, fimbriin-protein exporter; Tat, twin-arginine translocation; FEA, flagella export apparatus; Wss, WXG100 (proteins with WXG motif of ~100 amino acyl residues) secretion system.

insertion of all integral membrane proteins (IMPs).⁴⁸ YidC is a versatile pathway since it can be Sec-, SecA-, and/or SecB independent. In *B. subtilis*, studies have showed that SpoIIIJ and YqjG play a role in the folding of several secreted proteins and can work independently to insert integral membrane proteins.⁴⁹

Signal peptide of translocated preprotein is cleaved off by a membrane-bound signal peptidase (SPase). Different classes of N-terminal signal peptide are recognized and are cleaved by different types of SPases. Signal peptides of proteins targeted to Sec are of two classes: class 1 and class 2. Class 2 signal peptides are present in lipoproteins and are cleaved off by SPase II (for further details, see section 12.3.1.2). As depicted in Figure 12.3, precursor proteins exhibiting a class 1 signal peptide meet different fates; that is, they can (1) insert in cytoplasmic membrane and thus become integral membrane proteins (for further details, see section 12.3.1.1), (2) remain attached covalently or noncovalently to cell wall components (for further details, see section 12.3.2), (3) be released into the extracellular milieu, or (4) be injected into a eukaryotic host cell via pore formed by Sec-secreted listeriolysin O in a process called cytolysin-mediated translocation (CMT).^{50,51} It is worth noting that CMT has never been as yet reported in *Listeria*.

cell wall requires sortases. Proteins emerging from the Sec apparatus and exhibiting an LPXTG-like motif C-terminally located (for further details, see section 12.3.2.1) are recognized by membrane-associated sortase.⁵⁵ Transpeptidase sortase attacks the TG bond of the LPXTG-like motif, capturing cleaved polypeptide as a thioester-linked acyl enzyme at its active site cystein residue.⁵⁶ Subsequently, this complex is resolved by the nucleophilic attack of the amino group of the cross-bridge within lipid II precursor. Based on phylogenetic analyses, sortases are now classified into four classes, designated A, B, C, and D.⁵⁷ In *L. monocytogenes*, two sortases are present (SrtA and SrtB; Figure 12.3).

As observed in other Gram-positive bacteria, sortase of class A (also called SrtA subfamily) in *L. monocytogenes* is encoded only once in the genome, resembles a Type II membrane protein, and is necessary for the anchoring of the majority of LPXTG-containing proteins.⁵⁸ Sortase of class B (SrtB subfamily) recognizes a particular type of sorting signal (i.e., an NXZTN motif), which suggests a lower stringency of the recognition motif of SrtB compared to SrtA.⁵⁹ Captivatingly, from investigations in *Streptococcus pyogenes* and *Staphylococcus aureus*, glycosylated LPXTGase, an enzyme that cleaves the C-terminal LPXTG motif, is the first enzyme found that is produced by nonribosomal peptide (NRP) synthesis.^{60,61} It is known that NRP synthesis (and similarly related polyketide synthesis) occurs in Bacilli class, where NRPs are assembled in the cytoplasm by large megaproteins called NRP synthetases consisting of a series of active modules carrying out catalysis and modification of the tethered growing peptide chain.⁶² However, investigations in *S. aureus* suggest that enzymes responsible for cell wall assembly may also be involved in the construction of LPXTGase.⁶¹ Finally, it cannot be excluded that such a nonribosomally synthesized enzyme be also present and involved in LPXTG-like protein anchoring in *L. monocytogenes*.³⁷

Substrates of the Sec system are generally considered as exhibiting an N-terminal signal peptide composed of three domains: (1) The N-domain contains positively charged amino terminus, (2) the H-domain is a hydrophobic core region, and (3) the C-domain contains the cleavage site.⁶³ It must be emphasized, however, that it is not the case for all proteins (e.g., some SecA2-dependent and/or YidC-dependent proteins). Still, the presence of an N-terminal signal peptide indicates a protein is targeted to membrane. Despite lack of amino acid sequence similarity, signal peptides can be detected with good accuracy by various documented and publicly available applications (Table 12.1). The first methods developed were SigCleave and SPScan, which were implementations of a simple weight matrix approach.⁶⁴ While SigCleave is part of the EMBOSS suite and also available by an interface on the World Wide Web, SPScan is only available as part of the GCG suite and thus requires ability to work under Unix-like environment. Comparing the two programs, SPScan has clearly better predictive performance in terms of secretory protein and cleavage site recognition, especially for prokaryotic proteins.⁶⁵ Nearly a decade later, SignalP, a promising method based on a

TABLE 12.1
Bioinformatic Resources for Prediction of Bacterial N-Terminal Signal Peptides

Application	Method	Websserver	Ref.
SigCleave	Position weight matrix	http://bioweb.pasteur.fr/seqanal/interfaces/sigcleave.html	64
SPScan	Position weight matrix	none	64
SignalP	Neural network	http://www.cbs.dtu.dk/services/SignalP/	67
	Hidden Markov model		
PrediSi	Position weight matrix	http://www.predisi.de/	69
SOSUisignal	Global physicochemical analysis	http://bp.nuap.nagoya-u.ac.jp/sosui/sosuisignal/	70
Phobius	Hidden Markov model	http://phobius.binf.ku.dk/	71
PSORTb	Support vector machine	http://www.psорт.org/psортb/	74
	Hidden Markov model		
SPdb	BLAST	http://proline.bic.nus.edu.sg/spdb/	76

neural network, was released⁶⁶ and has undoubtedly become the most popular method for predicting N-terminal signal peptide. Since the first available version 1.1, SignalP has been substantially improved up to the latest version 3.0.⁶⁷

While version 1.1 is definitively out of date, both versions 2.0 and 3.0 use either a neural network (NN) or HMM. When comparing SignalP v2.0-NN, -HMM, and SPScan, it appears that (1) SPScan predicts correctly more proteins as secreted than SignalP v2.0-NN or -HMM; (2) SignalP v2.0-NN and -HMM are superior in predicting the correct cleavage site; (3) SignalP v2.0-NN lags behind SPScan and SignalP v2.0-HMM in classifying correctly the proteins, the latter providing the best prediction; and (4) SignalP v2.0-NN is the best for predicting of the correct cleavage site.⁶⁵ In other words, these methods are complementary in predicting an N-terminal signal peptide. The main improvement in SignalP v3.0 is increased accuracy in prediction of signal peptidase cleavage sites.⁶⁷ In comparative analyses, SignalP3.0 performs significantly better than other machine learning and HMM methods. Despite performance improvement in the latest SignalP v3.0, however, it appears that SignalP v2.0-NN remains the best signal prediction program.⁶⁸

A position weight matrix approach was improved by a frequency correction, which takes into consideration the amino acid bias (i.e., PrediSi).⁶⁹ SOSUisignal is a global structure analysis based on physicochemical features of the three signal peptide domains—N-, H-, and C-domains—and discriminates between cleavable and anchoring signal sequences.⁷⁰ Since a signal peptide contains a hydrophobic H-domain, there is a risk of erroneously identifying a transmembrane α -helix as a signal peptide or, conversely, classifying a protein with a signal peptide H-domain region as an IMP. In order to discriminate between the two, a combined TM topology and signal peptide predictor has been developed: Phobius.⁷¹ Phobius significantly reduces false classifications of signal peptides compared to SignalP. Another machine learning approach used for prediction of signal peptides is support vector machine (SVM), which can predict signal peptides with great accuracy.⁷² Such an implementation of an SVM combined with an HMM is part of PSORTb,⁷³ now applicable to both Gram-positive and Gram-negative bacteria.⁷⁴ Finally, SPdb, a repository of experimentally determined and computationally predicted signal peptides, is also accessible via BLAST (basic local alignment search tool) search.^{75,76}

It can be stressed again that these analyses only predict the presence of signal peptide, meaning that the protein is targeted to the cytoplasmic membrane. However, it does not necessarily mean the protein is translocated across the cytoplasmic membrane via Sec or released into the extracellular milieu. Indeed, proteins translocated via Tat or FPE also possess N-terminal signal peptides with additional features, which are not identified by the previous tools (Table 12.1). Thus, final prediction of a protein translocated via Sec requires additional inspections (see sections 12.2.2 and 12.2.3). Concerning proteins translocated by the Sec system and possessing a signal peptide, they can (1) be released into the extracellular medium or injected into a host cell, (2) remain associated to the cell wall by covalent or noncovalent interactions, or (3) remain associated to the cytoplasmic membrane by transmembrane domains (including H-domain of uncleaved signal peptide) or be lipoproteins (see section 12.3). Thus, final localization prediction of Sec substrates requires a combination of tools for prediction of function, motifs, and TMDs. It is also recommended to combine these results with those from tools dedicated to prediction of protein subcellular localization in Gram-positive bacteria (Table 12.2).

NNPSL was the first tool developed for such prediction and is based on an NN.⁷⁷ SubLoc,⁷⁸ PSORTb,⁷³ CELLO⁷⁹ (recently extended to prediction in Gram-positive bacteria⁸⁰), and LOctree⁸¹ are basically SVM. These tools have their own advantages and weaknesses,⁸² and some of them, like PSORTb, combine a variety of individual predictors. Proteome Analyst is a novel type of machine-learning classifier that involves several steps in the prediction process, such as BLAST search against Swiss-Prot database and naïve Bayesian classifiers.⁸³ From the most recent studies on performance of prediction tools, PSORTb and Proteome Analyst achieve the highest overall precision.⁸⁴ Gpos-PLoc, another type of ensemble classifier, was recently developed where several basic

TABLE 12.2
Bioinformatic Resources for Prediction of Subcellular Localization of Proteins in Gram-Positive Bacteria

Application	Method	Webserver	Ref.
NNPSL	Neural network	http://www.doe-mbi.ucla.edu/~astrid/astrid.html	77
SubLoc	Support vector machine	http://www.bioinfo.tsinghua.edu.cn/SubLoc/	78
PSORTb	Support vector machine Ensemble classifier	http://www.psорт.org/psортb/	73
CELLO	Support vector machine	http://cello.life.nctu.edu.tw/	80
LOCtree	Support vector machine	http://cubic.bioc.columbia.edu/services/loctree/	81
Proteome Analyst	Ensemble classifier	http://pa.cs.ualberta.ca:8080/pa/	83
Gpos-PLoc	Ensemble classifier	http://202.120.37.186/bioinf/Gpos/	85
DBSubLoc	BLAST	http://www.bioinfo.tsinghua.edu.cn/~guotao/intro.html	86
PSORTdb	BLAST	http://db.psорт.org/	87
PA-GOSUB	BLAST	http://www.cs.ualberta.ca/~bioinfo/PA/GOSUB/	88
Augur	Ensemble classifier	http://bioinfo.mikrobio.med.uni-giessen.de/augur/	80

classifiers were fused and optimized for predicting subcellular localization of Gram-positive bacterial proteins.⁸⁵ Finally, several databases (derived from previously described prediction tools) are available following BLAST search (DBSubLoc,⁸⁶ PSORTdb,⁸⁷ and PA-GOSUB⁸⁸). Augur is another database especially dedicated to protein localization on cell surface of Gram-positive bacteria.⁸⁹ Once again, final prediction of secreted proteins (and localization) should combine results from these various bioinformatic tools.⁸⁴

Using SignalP v2.0 to predict signal peptide region and TopPred v2.0 to exclude other transmembrane domains, 86 proteins were predicted as secreted into the extracellular medium from genomic analysis of *L. monocytogenes* EGD-e.¹⁹ In *L. monocytogenes* F2365, 420 proteins were predicted with a putative N-terminal signal peptide, including 2 with a YSIRK motif.¹⁸ Performing extensive genomic analyses, which combined results from SignalP v2.0, SigCleave, SOSUI, PSORT, and TMPinGS, the number of proteins bearing an N-terminal signal peptide was estimated at 525 in *L. monocytogenes* EGD-e, including 255 IMPs and 270 exported proteins where 121 would be released into the extracellular milieu.⁹⁰ All 14 virulence factors characterized so far in *L. monocytogenes* are most likely translocated via the Sec translocon.³⁷ Among the 121 proteins originally predicted as secreted via Sec and released into the extracellular milieu, a closer look revealed that four prepilins—that is, ComGC (Lmo1345), ComGD (Lmo1344), ComGE (Lmo1343), and ComGG (Lmo1341)—should be removed from the output since they would form trans-cell-wall structure following translocation via FPE.³⁷ Proteomic analysis of supernatant from liquid culture of *L. monocytogenes* EGD-e allowed the identification of 54 out of 117 proteins predicted as extracellular, including virulence factors Hly, PlcA, and PlcB.⁹⁰

12.2.2 TAT PATHWAY

The term twin-arginine translocation (Tat) was coined from the systematic presence of RR motif in signal peptide of proteins translocated via this secretion system.⁹¹ The [ST]RRXFLK motif straddles the N-domain and H-domain of N-terminal signal peptide.⁹² Contrary to the Sec translocon, the main feature of this pathway is its ability to translocate proteins in a folded state. General knowledge on the precise succession and mechanistic events leading to protein secretion via this pathway remains rudimentary.⁹³ The generally accepted translocation model was first proposed by Mori and Cline,⁹⁴ where Tat translocation follows a cycle in which TatBC functions in the specific recognition of the substrate and TatA functions as the pore-forming unit. An alternative model proposes

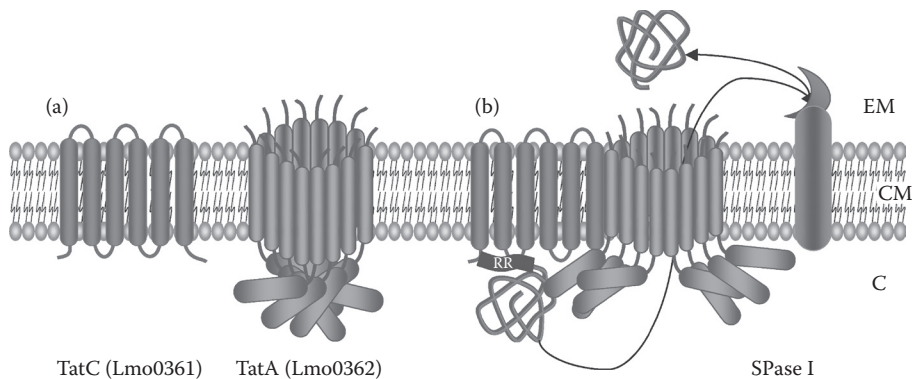


FIGURE 12.4 Tat translocon in *L. monocytogenes*.³⁷ During Tat secretion, the general model proposes a cyclical assembly of components. (A) In resting state, Tat machinery components are separately present in the cytoplasmic membrane (i.e., TatB and TatA). (B) Once Tat substrate protein precursor binds to the TatC in an energy-independent step, this complex associates with TatA in a step driven by transmembrane proton electrochemical gradient. This association would persist until completion of protein transport across the membrane driven by proton motive force. Tat signal peptide is subsequently cleaved by SPase I and Tat machinery components disassembled, as depicted in (A). RR, twin-arginine motif in Tat signal peptide; EM, extracellular milieu; CM, cytoplasmic membrane; C, cytoplasm.

that membrane integration could precede Tat-dependent translocation and the membrane targeting process may require ATP-dependent N-terminal unfolding-steps energy.⁹⁵

Still, components of Tat translocon differ in number between Gram-negative and Gram-positive bacteria.³⁰ The most baffling difference is the absence of TatB from all Gram-positive bacteria sequenced so far, although it is an essential components of Tat translocon in *E. coli*, which is used as a paradigm.⁹⁶ As in most Gram-positive bacteria, Tat translocon in *L. monocytogenes* is encoded in one locus and is composed of only two proteins, TatA and TatC³⁷ (Figure 12.4). TatC is a large IMP generally considered as the primary site for signal-peptide recognition.⁹⁷ TatA is a membrane protein that oligomerizes to form a protein-conducting channel where the number of subunits would adjust in function of the Tat substrate size.⁹⁸ In TatA, a cytoplasmic lid region acts as a gate and would open following association of TatC–substrate complex with TatA, then inducing conformation change and protein translocation driven by proton motive force. Translocated protein is finally released after cleavage by SPase I.⁹⁶ A Tat translocon does not seem to be systematically present in *L. monocytogenes* as no component could be identified in *L. monocytogenes* F2365. The Tat system has never been experimentally investigated in *Listeria*; thus, its expression, functionality, involvement of one or three SPases I, or proteins secreted via this pathway remain unknown.³⁷

Three tools are currently available to discern Tat substrates (Table 12.3) and TATFIND was the first program especially devoted to such identification.⁹⁹ In its original available version, TATFIND v1.2, prediction was based on two criteria: (1) presence of conserved Tat motif ZRRZZZ within the first 35 amino acid residues, where Z represents a defined set of permitted residues; and (2) presence

TABLE 12.3
Bioinformatic Resources for Prediction of Tat Signal Peptides

Application	Method	Webserver	Ref.
TATFIND	Physicochemical analysis and regular expression	http://signalfind.org/tatfind.html	100
TatP	Neural network and regular expression	http://www.cbs.dtu.dk/services/TatP/	101
TATPred	Naïve Bayesian network	http://www.jenner.ac.uk/logP/JennerlogPcalc.htm	102

of an uncharged stretch of at least 13 amino acids downstream of the twin arginine. In the latest version, TATFIND v1.4, search for a single charged residue in positions +2 and +5 relative to the RR was included.¹⁰⁰ TatP v1.0 incorporates signal peptide and cleavage site prediction based on a combination of two artificial neural networks followed by a postfiltering of the output based on regular expression RR[FGAVML][LITMVF].¹⁰¹ Compared to TATFIND v1.2, TatP generates far fewer false positive but slightly more false negative predictions. TATPred is the latest algorithm based on naïve Bayesian network developed for prediction of Tat substrates.¹⁰² Compared to TatP, TATPred appears as the most robust and reliable predictor with higher sensitivity of prediction.

According to TATFIND search, only two Tat substrates could be identified in *L. monocytogenes* EGD-e.⁹⁹ One of these putative Tat substrates, however, is also present in *L. monocytogenes* F2365, where the Tat system is not encoded.³⁷ These substrates have never been reported as present in the extracellular milieu of *L. monocytogenes*. While it has been long assumed that the RR motif was highly specific and conserved in Tat substrates, it must be stressed that substitutions of one arginine, or in some cases both arginines, by lysine¹⁰³ or that natural proteins harboring very distantly related RR motifs¹⁰⁴ could still permit targeting and translocation via the Tat pathway.¹⁰⁵ This indicates that Tat system specificity is more flexible than originally thought and thus presence of Tat substrate cannot be systematically identified by bioinformatic analysis.

12.2.3 FPE

Components of fimbriin-protein exporter (FPE) of Gram-positive bacteria are homologous to proteins required for secretion of substrate proteins in Gram-negative bacteria, namely, some ATPase and IMP components of the Type II protein secretion system (T2SS), Type 4 piliation system (Tfp), and Type IV protein secretion system (T4SS), as well as archaeal flagella.¹⁰⁶ As in all Gram-positive bacteria where it has been reported so far,^{33,107,108} components of FPE in *L. monocytogenes* are encoded in a *comG* operon, except for *ComC* located elsewhere on the chromosome. Protein exporters of the FPE family consist of two constituents—ComGA and ComGB—that would function together in an ATP-hydrolysis-dependent export of proteins across the cytoplasmic membrane^{109,110} (Figure 12.5). ComGA is an ATPase localized to the cytoplasmic side of the membrane that could participate in modeling of pilus-like structure.¹⁰⁹ As a homologue to PilC of Tfp and PulF of T2SS,¹⁰⁹ ComGB is an IMP having three putative TMDs that could play the role of a protein-conducting channel.¹¹¹ ComC is a Type 4 prepilin peptidase involved in cleavage of N-terminal signal peptide of class 3;¹¹² this signal peptidase belongs to the aspartic acid protease family.¹¹³

While ComC is required for maturation, translocation, and assembly of prepilins, an initial translocation event across the cytoplasmic membrane has not been clearly elucidated. As prepilin signal peptide is cleaved at the cytoplasmic side between the N- and H-domains, prepilins are certainly not translocated by the Sec or Tat pathways and the hypothesis of ComGAB involvement is favored. However, YidC contribution cannot be excluded³⁰; prepilins were originally thought to insert spontaneously in the membrane bilayer but with the current knowledge of membrane protein insertion this hypothesis should not be privileged (see section 12.3). Four Type 4 prepilins are encoded in *comG* locus by the *comGC*, *comGD*, *comGE*, and *comGG* genes¹¹⁰; ComGF is presumably an IMP. Once matured and translocated, pilins form a trans-cell-wall macromolecular complex where monomers are covalently linked by disulphide bonds.¹¹⁴ Since this structure is involved in bacterial competence and does not form a proper Type 4 pilus, it was named competence pseudopilus.

In *B. subtilis*, Type 4 prepilins exhibit N-terminal signal peptides with a conserved motif [KR]G▼F[TSI][LTY][VLIP][EA] located between the N- and H-domains where ▼ indicates the predicted cleavage site.¹¹⁰ In *Listeria*, the motif is slightly different—that is, [NPRS][GA]▼F[TS]L[VLIP][EF]—and is found in five putative prepilins (i.e., ComGC, ComGD, ComGE, ComGF, and ComGG).³⁷ In *B. subtilis*, the highly conserved phenylalanine at position +1 is aminomethylated by ComC, which thus appears bifunctional as it is also involved in prepilin processing.³⁰ Using ScanProsite syntax,¹¹⁵ search for consensus motif [GA]F[TS]LX[EF] located between the N- and

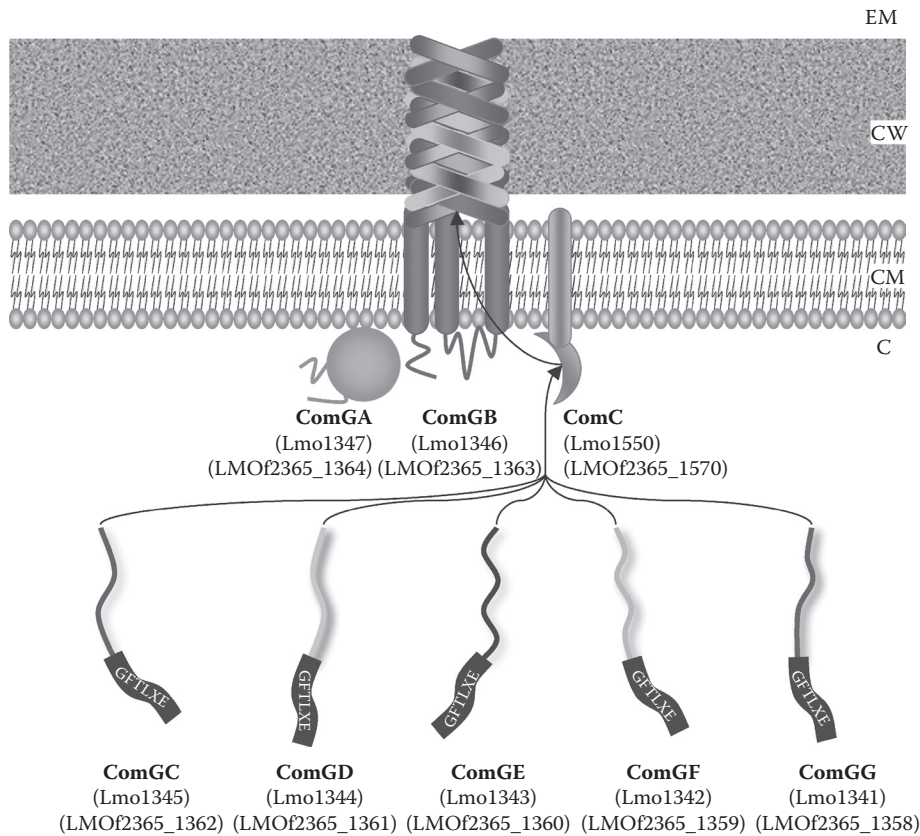


FIGURE 12.5 FPE in *L. monocytogenes*.³⁷ The prepilins initially float in the CM; initial insertion into the membrane is certainly Sec or Tat independent but the involvement of ComGAB remains to be ascertained. After processing by ComC signal peptidase, ComGA and ComGB would be involved in assembly of pseudo-pilins to form a trans-cell-wall pilus-like structure. GFTLXE, conserved [GA]F[TS]LX[EF] motif in Type 4 prepilin signal peptide from *Listeria*; CW, cell wall; EM, extracellular milieu; CM, cytoplasmic membrane; C, cytoplasm.

H-domains of predicted signal peptide can thus be performed in order to identify putative FPE substrates in *Listeria*. The FPE system has never been experimentally investigated in *Listeria*; thus, its expression, functionality, and involvement in bacterial competence remain to be established.¹⁰⁸

12.2.4 FEA

L. monocytogenes produces up to six peritrichous flagella, which are down-regulated at 37°C, although variation from one strain to another was reported.^{116,117} Regulation of listerial flagella is not entirely understood and appears rather complex since at least five regulators involved in its expression have been identified so far: FlaR,¹¹⁸ PrfA,¹¹⁹ DegU,¹²⁰ MogR,¹²¹ and GmaR (Lmo0688 also called WcaA).¹²² Interestingly, the antirepressor GmaR is bifunctional since it also functions as a glycosyltransferase for flagellin FlaA¹²² and glycosylation with β -O-linked N-acetylglucosamine was indeed established for FlaA.¹²³ This investigation constituted the first description of β -O-GlcNac post-translational modification on a prokaryotic protein, though flagella glycosylation is not essential for motility in *L. monocytogenes*.¹²⁴ As motility mediators, flagella are important in colonization of abiotic surfaces and host cell invasion but do not function as adhesins.^{124,125}

Interestingly, FlaA was also demonstrated as exhibiting a peptidoglycan-hydrolyzing activity that might play a role during flagella assembly.^{126,127} Indeed, some flagellar components are assembled

on the bacterial cell surface where local digestion of cell wall sacculus might be required—namely, for (1) the rod proteins (i.e., FlgB, FlgC, FliE, and FlgG), (2) the hook/junction proteins (i.e., FliK, FlgD, FlgE, FlgK, and FlgL), and (3) the filaments proteins FlaA and FliD. As in Gram-negative bacteria,^{128–132} these proteins lack a cleavable N-terminal signal peptide and are presumably translocated by the flagella export apparatus (FEA) composed of FlhA, FlhB, FliH, FliL, FliP, FliO, and FliH. In *Listeria*, all flagella components are encoded in a single flagellar–motility–chemotaxis cluster of 41 genes,³⁷ where FEA and its potential substrates could be identified by homology search. In Didermata, T3SS refers to a secretion system where translocation apparatus is homologous to injectisomes (T3aSS) and flagella (T3bSS),^{51,133} both of which are involved in secretion of extracellular proteins.^{134,135} As already stressed, however, this terminology is restricted to Gram-negative bacteria. In monoderm bacteria, involvement of FEA in secretion of extracellular protein has only been suggested in *Bacillus thuringiensis*.¹³⁶

12.2.5 HOLINS

Holins (hole-formers) are small membrane proteins of phage origin that essentially control endolysin function in a process leading to bacterial apoptosis.^{137–139} A current model for the holin-endolysin system proposes that holins accumulate in the cytoplasmic membrane, whereas endolysins accumulate in the cytoplasm^{140,141} (Figure 12.6). At a programmed time, holins oligomerized to form pores in the cytoplasmic membrane, allowing release of endolysins into the extracytoplasmic space leading to cell lysis following cell wall degradation and membrane disruption. Homo-oligomeric pore complexes formed by holins would provide a passive but specific translocation system.¹⁴² Generally, holin and its specific endolysin are genetically encoded in tandem. Some holin genes possess a dual start motif, which results in the expression of two distinct proteins with dramatically opposed function since one would promote autolysis (holin) and the other would inhibit it (antiholin).¹⁴³ Such regulation can also occur between proteins encoded at different loci (e.g., *lrgAB/cidAB* operons in *Staphylococcus aureus*).^{144,145} Holins are an extremely diverse group of proteins with 23 distinct families recognized in TC-DB (transport classification database),¹⁴⁶ although they can be grouped into three classes based on membrane topologies.¹⁴⁷ Class 1 holins exhibit three helical TMDs, whereas class 2 holins have two TMDs. Besides classes 1 and 2, which cover most holins, a third class was identified on the basis of T protein of phage T4 where only a single TMD is present.¹⁴⁸

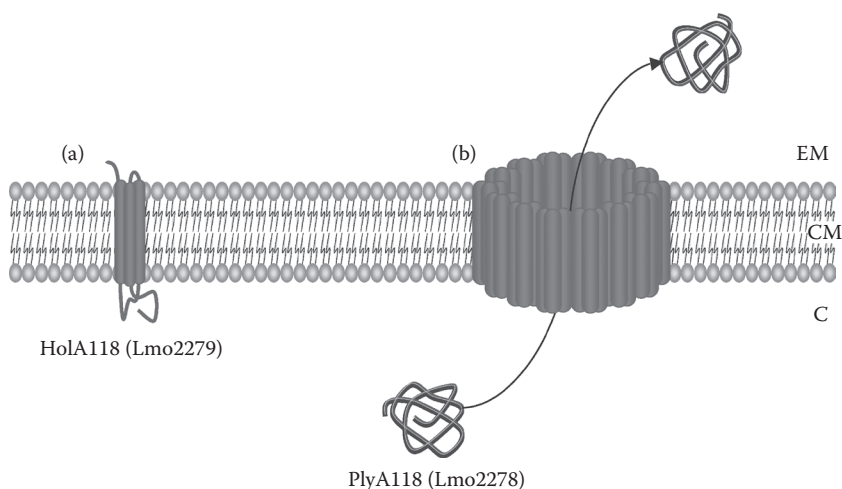


FIGURE 12.6 ϕ A118 holin-endolysin system in *L. monocytogenes* EGD-e.³⁷ (A) HolA118 is a class 1 holin (i.e., with 3 TMDs). (B) HolA118 oligomerizes in the CM to form a pore allowing translocation and activation of endolysin Ply118. EM, extracellular milieu; CM, cytoplasmic membrane; C, cytoplasm.

The number of holins encoded in *Listeria* varies between strains and species³⁷; only one holin could be identified in *L. monocytogenes* F2365, whereas five holins were found encoded in non-pathogenic strain *L. innocua* CLIP11262. However, only three distinct families of holins were identified in *L. monocytogenes* (i.e., as belonging to bacteriophage 118, TcdE, and bacteriophage 11 families). Holins of ϕ A118 family (HolA118) were first identified and investigated in *L. monocytogenes* although a homologue is at least also encoded in *L. innocua*.^{37,149} HolA118 is not encoded by all *L. monocytogenes* species as it is absent from *L. monocytogenes* F2365. Native holin HolA118 is a 93-amino-acid-long protein belonging to class 1, but its encoding gene is subjected to dual translational initiation, which leads to a second 83-amino-acid-long protein called HolA118(83) acting as an antiholin.¹⁵⁰ Gene encoding phage lysin of ϕ A118 (PlyA118) systematically clusters with gene encoding HolA118.^{18,151} The endolysin PlyA118 is an L-alanoyl-D-glutamate peptidase hydrolyzing the cross-linking bridges of cell wall peptidoglycan and thus responsible for bacterial lysis in a programmed cell death.

Holins belonging to TcdE family are encoded in all sequenced *Listeria* but as ϕ 11 holins they have never been experimentally investigated. TcdE holin was investigated in *Clostridium difficile*, where toxigenic strains produce two large toxins, TcdA and TcdB, of major importance in bacterial virulence, which would be translocated across the cytoplasmic membrane by TcdE.^{152,153} While in *C. difficile* all these genes are encoded within a pathogenicity locus, no genes coding for toxins or virulence factors could be identified in *Listeria*.³⁷ However, a putative autolysin lacking a signal peptide was systematically present (i.e., genes encoding Lmo0129 and LMOF2365_0147 in *L. monocytogenes* EGD-e and F2365, respectively). Although this particular holin family has never been investigated per se in *Listeria*, proteomic analysis in *L. monocytogenes* EGD-e disclosed the presence of Lmo0129 in supernatant of bacterial cultures, suggesting this secretion pathway is active in this species.⁹⁰ In *Listeria*, ϕ 11 holins were only identified in unassembled genome of *L. monocytogenes* F6854 and nonpathogenic *L. innocua* CLIP11262, where they systematically clustered with genes encoding amidases presumably involved in cell wall degradation.³⁷

12.2.6 Wss

Wss stands for proteins with WXG motif of ~100 residues (WXG100) secretion system.¹⁵⁴ WXG100 is a new superfamily of proteins around 100 amino acids long, possessing a coil-coil domain and bearing a conserved WXG motif located in the middle region. First identified members of this superfamily were paralogues ESAT-6 (early secreted antigen target of 6 kDa) and CFP-10 (culture filtrate protein 10) from *Mycobacterium tuberculosis*. While ESAT-6 and CFP-10 are specific and experimentally investigated proteins, WXG100 (PF06013) is an established and generic terminology more appropriate to describe protein members of this family, especially those that have not been experimentally investigated yet. No generic terminology for the different components of Wss apparatus has been established yet. Presence of a novel protein secretion system was clearly suggested by bioinformatic analysis.¹⁵⁴ In *B. subtilis*, genes encoding WXG100 proteins appeared to cluster systematically with *yukab*, which are predicted to encode membrane bound ATPases with FtsK/SpoIIIE domains. Similar genetic organization was observed in some *Corynebacterium*, *Mycobacterium*, *Streptomyces*, *Bacillus*, *Clostridium*, *Listeria*, and *Staphylococcus* species.^{31,154} YukAB homologues appear encoded as single or two CDS. To date, Wss seems phylogenetically restricted to Gram-positive bacteria and has been experimentally investigated only in *M. tuberculosis*, *M. smegmatis*, and *S. aureus*.

In *Mycobacterium*, two WXG100 proteins are secreted: ESAT-6 and CFP-10.¹⁵⁵ Recently, a C-terminal signal sequence required for secretion via Wss was unraveled in CFP-10.¹⁵⁶ *Mycobacterium* Wss apparatus was named Snm (secretion in mycobacteria) and is composed at least of^{155,157}: (1) Snm1, Snm2, and Snm6 containing NTP-binding motifs (where Snm1 and Snm2 are homologous to YukAB); (2) Snm4, which is an IMP; (3) Snm5 and Snm7 with uncharacterized functions,

and Snm8 (i.e., a membrane anchored serine protease). Snm permits translocation of ESAT-6 and CFP-10 as well as their heterodimerization.^{158,159} In *S. aureus*, the Wss was named Ess (ESAT-6 secretion system) and is encoded in a locus composed of eight CDS including two WXG100 paralogues—EsxA (Ess extracellular protein A) and EsxB—as well as:¹⁶⁰ (1) EssC (Ess protein C) homologous to YukAB; (2) EssA, EssB, and EsaA (ESAT-6 secretion accessory protein A), which are IMPs; and (3) EsaB and EsaC, which predict cytoplasmic chaperones. Compared to *B. subtilis*, where a putative Wss was primarily uncovered, EssB and EsaB appear homologous to YukC and YukD, respectively. In *S. aureus*, no homologue to Snm4, Snm5, Snm6, Snm7, or Snm8 was found, whereas in mycobacteria, no homologue to EssA, YukC, EsaA, and YukD could be identified. In both *M. tuberculosis* and *S. aureus*,^{160,161} Wss is important and critical for bacterial pathogenicity, though the function of WXG100 proteins in virulence remains obscure.¹⁶²

Syntenly is highly conserved between Wss encoding loci of *S. aureus* and *L. monocytogenes*.¹⁶⁰ However, compared to *Mycobacterium* species or *S. aureus*, only a single copy of Wss locus is present in each sequenced *Listeria* genome.³⁷ Following homology with *S. aureus* and *Mycobacterium*, Wss in *L. monocytogenes* is represented in Figure 12.7. From one report,¹⁶³ it seems that WXG100 protein is not required for virulence of *L. monocytogenes*. Still, protein expression, system functionality, and involvement in bacterial virulence of Wss remain to be established.

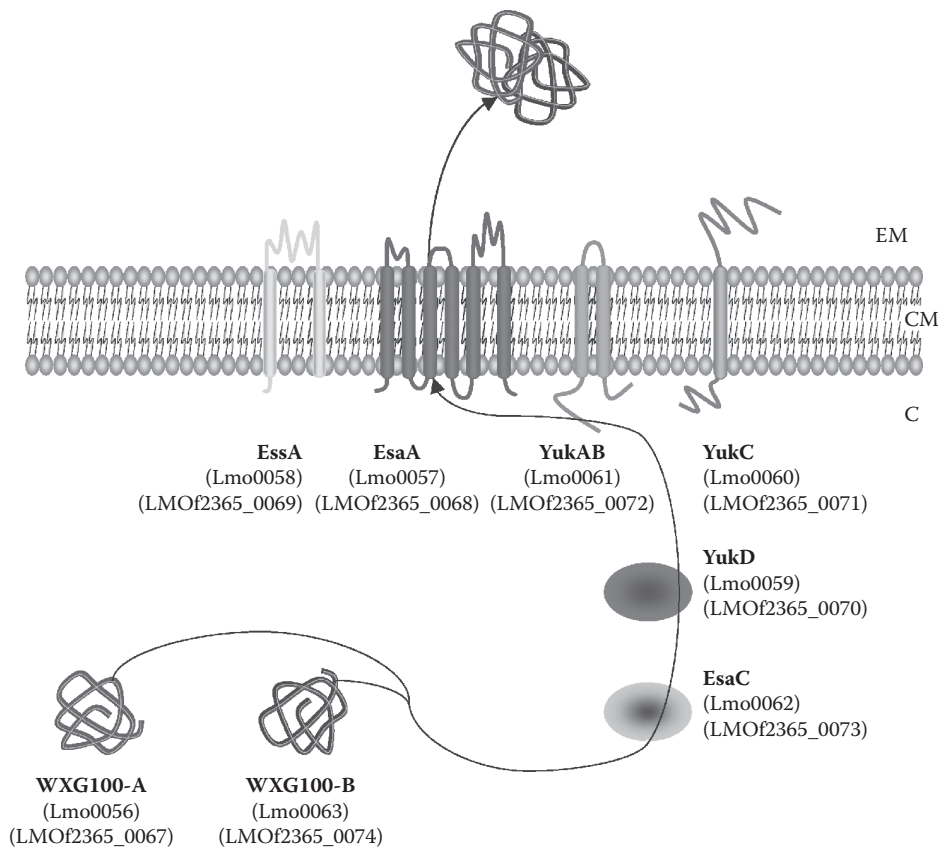


FIGURE 12.7 Wss in *L. monocytogenes*.³⁷ WXG100 proteins would interact with putative cytoplasmic chaperones YukD and EsaC before being translocated by the Wss apparatus constituted of EssA, EsaA, YukC, and YukAB; in the course of translocation the two WXG100 proteins would finally form a heterodimer. EM, extracellular milieu; CM, cytoplasmic membrane; C, cytoplasm.

12.3 CELL ENVELOPE-ASSOCIATED PROTEINS

Cell envelope of Gram-positive bacteria is primarily composed of a single biological membrane (i.e., the cytoplasmic membrane) and a cell wall made up of peptidoglycan (which in turn consists of linear polysaccharide chains cross-linked by short peptides).¹⁶⁴ Besides peptidoglycan, the rigid cell-wall of Gram-positive bacteria contains large amounts of wall-associated polymers, also called “secondary” cell wall polymers (SCWPs), which can be classified into three distinct groups: (1) teichoic acids (i.e., polyol phosphate polymers, including lipoteichoic acids), (2) teichuronic acids, and (3) other neutral or acidic polysaccharides that cannot be assigned to the two former groups (e.g., lipoglycans).^{55,165,166} The SCWPs, present in various proportions, are either covalently linked to the peptidoglycan backbone (i.e., teichoic acids) or tethered to a lipid anchor moiety. Except for teichoic and teichuronic acids, the structure and biosynthesis of other SCWPs are largely unknown. It must be stressed that in almost all phylogenetic branches of Archaea and Bacteria, the cell envelope is also constituted of a proteinaceous S-layer (regular crystalline surface layer), which forms the outermost cell-wall layer.¹⁶⁷ The S-layer entirely coats the bacterial cell surface and is composed of (glyco)proteins, which bind by noncovalent interactions to cell wall components and are arrayed in a two-dimensional lattice.¹⁶⁷ S-layer, however, is not present in all Gram-positive bacteria as it is absent from all members of *Listeria* genus. Within the cell envelope, proteins can associate with cytoplasmic membrane or cell wall components.⁸

12.3.1 MEMBRANE-ASSOCIATED PROTEINS

Membrane-associated proteins include membrane integrated proteins as well as peripheral membrane proteins. Being different from membrane integrated proteins, peripheral membrane proteins do not possess membrane spanning domains. Membrane integrated proteins are anchored within the lipid bilayer and thus systematically exhibit hydrophobic transmembrane domains (TMDs), which are normally α -helices for proteins found in the cytoplasmic membrane. Peripheral membrane proteins include (1) lipoproteins, (2) subunits of membrane-associated complexes, and (3) proteins interacting with membrane components due to electrostatic and/or hydrophobic/steric properties.¹⁶⁸ Following recommendations of the Gene Ontology (GO) Consortium for describing location of cellular components (one of the three organizing principles of GO with biological process and molecular function),¹⁶⁹ two classes of membrane-related location are distinguished (Figure 12.8). First, intrinsic to plasma membrane (GO:0031226) refers to proteins with covalently attached moiety embedded in the cytoplasmic membrane, which splits into (1) integral to plasma membrane (GO:0005887) corresponding to membrane integrated proteins, where some part of the peptide sequence spans all or part of the cytoplasmic membrane; and (2) anchored to plasma membrane (GO:0046658) corresponding to proteins tethered to the cytoplasmic membrane by a nonpolypeptidic covalently attached anchor: lipoproteins. Second, extrinsic to plasma membrane (GO:0019897) refers to proteins neither anchored by covalent bonds to any moiety nor directly embedded in the cytoplasmic membrane; some of these proteins can be (1) primarily present in the cytoplasm (GO:0005737) but interact with membrane components, or (2) subcomponents localized within protein complex (GO:0043234).

12.3.1.1 Integral Membrane Proteins

As already mentioned, all bacterial IMPs are presumably inserted into the cytoplasmic membrane via YidC homologues^{48,170,171} (i.e., SpoIIIJ and YqjG in Gram-positive bacteria³⁰; see section 12.2.1). The Sec-independent function of YidC homologues is conserved and essential for bacterial cell growth as it works like a membrane protein insertase.¹⁷² YidC plays a major role in the folding step of transmembrane-spanning domains but the exact mechanism of functioning is not fully understood.¹⁷³ YidC would facilitate the insertion of membrane proteins by providing a special amphiphilic surface, which would overcome the repulsion of the hydrophobic protein segments by polar head groups. In addition, polar residues seem to be protected against the hydrocarbon core of

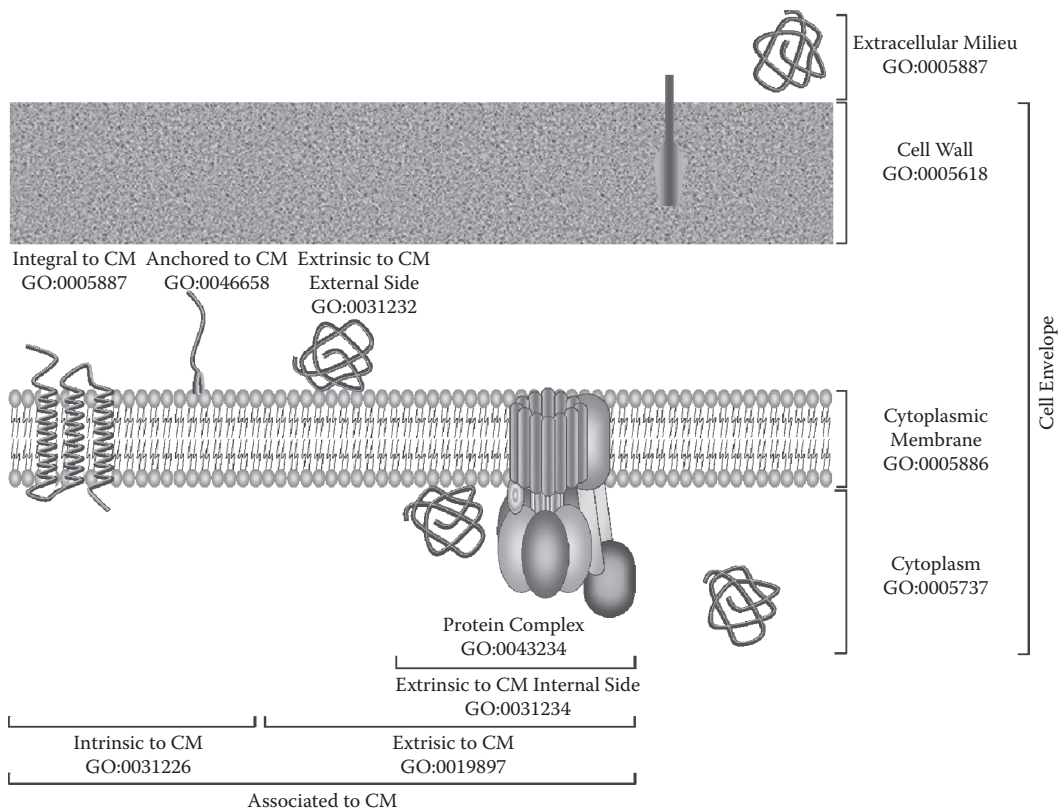


FIGURE 12.8 Description of protein localization following GO in Gram-positive bacteria.¹⁷⁹ In monoderm bacteria four subcellular compartments can be distinguished: (1) cytoplasm, (2) cytoplasmic membrane, (3) cell wall, and (4) extracellular milieu. A membrane-associated protein can be intrinsic or extrinsic. Proteins intrinsic to CM are either integral to membrane (i.e., integral membrane proteins) or anchored to CM, essentially lipoproteins with the restriction of lipoproteins having TMDs. Proteins extrinsic to CM can be located on the external or internal side of the CM (i.e., in exoplasmic or cytoplasmic compartment, respectively). They can interact more or less temporarily with membrane components or be part of membrane protein complex (e.g., F_1F_0 ATP synthase δ subunit) as indicated on the schema. A protein can also have multiple final localization. Importantly, because cell wall of Gram-positive bacteria is permeable, extracellular milieu penetrates it. CM, cytoplasmic membrane; GO, gene ontology.

the membrane by YidC. In the case of Sec-dependent translocation, the protein would be stabilized and then folded by contact with YidC after leaving the Sec YEG channel. It is quite possible that the transmembrane segments could fold and interact with each other even within SecYEG-YidC machinery. It has been suggest that YidC functions as an assembly site for hydrophobic domains, so it may be necessary for its attaching to the individual subunits of multisubunit membrane complex.¹⁷⁴ It is worth noting that, in *E. coli*, targeting, translocation, and insertion of IMPs are considered cotranslational and thus SRP dependent.¹⁷⁵

Translocation of polypeptide chain is promoted by signal peptides and interrupted by another type of topogenic element called stop-transfer sequence^{176,177}; both types of topogenic sequences act as α -helical transmembrane domains. Multiple uncleaved signal peptides can be found all along the amino acid sequences; when located N-terminally, they can be cleaved or not. These types of topogenic elements have a $C_{out}-N_{in}$ topology and when uncleaved are also called signal-anchors or Type II signals.¹⁷⁸ Similarly, one or more stop-transfer sequences with an $N_{out}-C_{in}$ topology can be present in polypeptide chain and are also called Type II signals. Single-spanning membrane proteins are discriminated on the basis of Type I or Type II signal (Figure 12.9). Polytopic membrane

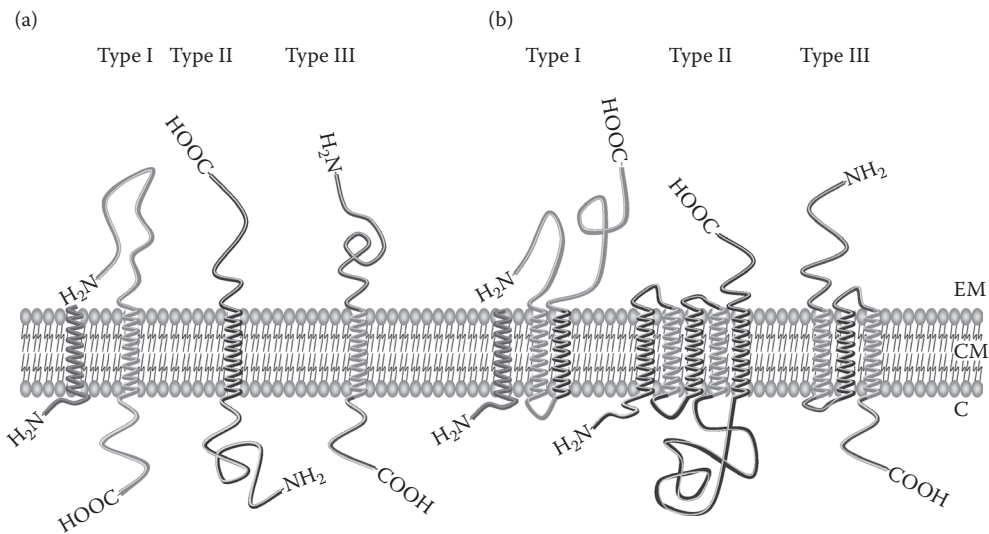


FIGURE 12.9 Classification and topology of IMP in cytoplasmic membrane.¹⁷⁹ (A) Three types of single-spanning membrane proteins can be discriminated: (1) Type I proteins possess a cleavable N-terminal signal peptide and thus have a Type I signal or stop-transfer sequence with N_{out}-C_{in} topology; (2) Type II proteins have a Type II signal or signal-anchor sequence with a C_{out}-N_{in} topology, which can correspond to an uncleavable N-terminal signal peptide; and (3) Type III proteins have reverse signal-anchor sequence (i.e., with a N_{out}-C_{in} topology) and are sometimes described as Type I proteins without a cleavable signal peptide since the reverse signal-anchor sequence is a Type I signal. (B) Three types of multispanning-membrane proteins (i.e., with a number of TMDs higher than 1) can be distinguished based on whether the most N-terminal TMD is either (1) cleaved by a SPase (i.e., Type I); (2) spans the membrane with an N_{out}-C_{in} orientation (i.e., Type II); or (3) have a C_{out}-N_{in} orientation. Various numbers of TMDs are present in multispanning-membrane proteins where alternates Type I and II signals. EM, extracellular milieu; CM, cytoplasmic membrane; C, cytoplasm.

proteins are built up of a series of Types I and II modules that initiate and halt the translocation of the polypeptide chain. Such IMPs are classified on the basis of the orientation of the most N-terminal TMD spanning the lipid bilayer¹⁷⁹ (Figure 12.9).

Numerous tools are available to predict IMPs and their topology. Table 12.4 is an attempt to review all of them. These tools are based on various approaches, such as (1) statistical analyses (e.g., Tmpred or TMSTAT); (2) hydrophobicity analyses (e.g., SOSUI or TopPred); (3) NNs (e.g., PREDTMR or PHDhtm); (4) SVMs (e.g., SVMtm); or (5) HMMs (e.g., HMMTOP or THUMBUP). Some of them—for example, ConPred or TUPS—combine results of several models. Readers are invited to study related publications listed in Table 12.4 in order to get further insight into the methods used. The total number of IMPs encoded in *L. monocytogenes* genomes is estimated to be 1204 and 733 in *L. monocytogenes* EGD-e and F2365, respectively.¹⁸ Virulence factors ActA and SvpA are cell surface exposed IMPs of *L. monocytogenes* exhibiting a hydrophobic tail (i.e., a carboxyl terminal region containing a hydrophobic domain followed by positively charged residues).^{180,181} Following genomic analysis, a total of 11 surface proteins with hydrophobic tails have been predicted in *L. monocytogenes* EGD-e.¹⁹ Contrary to what is sometimes assumed, cell surface IMPs should not be restricted only to proteins with a hydrophobic tail¹⁸²; indeed, depending on the number and organization of Type I and Type II signals in IMPs, final protein topology can result in the cell surface exposure of functional domains located not only N- or C-terminally but also in loops. With FbpA as an example,⁴⁷ it can be noticed that from experimental investigations, some proteins appear located within the cytoplasmic membrane despite the absence of predicted signal peptide and TMD.

TABLE 12.4
Bioinformatic Resources for Prediction of TMDs

Application	Method	Webserver	Ref.
TMpred	Statistical analysis	http://www.ch.embnet.org/software/TMPRED_form.html	249
TopPred	Hydrophobicity analysis	http://bioweb.pasteur.fr/seqanal/interfaces/toppred.html	250
PHDhtm	Neural network	http://npsa-pbil.ibcp.fr/cgi-bin/npsa_automat.pl?page=/NPSA/npsa_htm.html	251
DAS	Hydrophobicity analysis	http://www.sbc.su.se/~miklos/DAS/	252
TMAP	Statistical analysis	http://bioweb.pasteur.fr/seqanal/interfaces/tmap.html	253
TSEG	Hydrophobicity analysis	http://www.genome.jp/SIT/tsegdir/tseg_exe.html	254
TMHMM	Hidden Markov model	http://www.cbs.dtu.dk/services/TMHMM-2.0/	255
SOSUI	Hydrophobicity analysis	http://bp.nuap.nagoya-u.ac.jp/sosui/	256
PREDTMR	Neural network	http://athina.biol.uoa.gr/PRED-TMR2/	257
kPROT	Statistical analysis	http://bioinfo.weizmann.ac.il/kPROT/	258
TMSTAT	Statistical analysis	http://bioinfo.mbb.yale.edu/tmstat/	259
HMMTOP	Hidden Markov model	http://www.enzim.hu/hmmtop/	260
TMFinder	Hydrophobicity analysis	http://www.bioinformatics-canada.org/TM/	261
DAS-TMfilter	Hydrophobicity analysis	http://www.enzim.hu/DAS/DAS.html	262
SPLIT	Statistical analysis	http://split.pmfst.hr/split/	263
THUMBUP	Hidden Markov model	http://sparks.informatics.iupui.edu/Softwares-Services_files/thumbup.htm	264
UMDHMMTMHP	Hidden Markov model	http://sparks.informatics.iupui.edu/Softwares-Services_files/umdhmm.htm	264
TUPS	Combination	http://sparks.informatics.iupui.edu/Softwares-Services_files/tups.htm	264
BPROMPT	Bayesian belief network	http://www.jenner.ac.uk/BPROMPT/	265
SVMtm	Support vector machine	http://ccb.imb.uq.edu.au/svmtm/svmtm_predictor.shtml	266
ConPred	Combination	http://bioinfo.si.hirosaki-u.ac.jp/~ConPred2/	267
HMMTM	Hidden Markov model	http://biophysics.biol.uoa.gr/HMM-TM/	268
MINNOU	Hydrophobicity analysis	http://minnou.cchmc.org/	269
MEMSAT	Statistical analysis	http://bioinf.cs.ucl.ac.uk/psipred/	270

12.3.1.2 Lipoproteins

In monoderm bacteria, lipoproteins are attached to the outer surface of the cytoplasmic membrane via a covalently bound lipid anchor.³² Systematically, these proteins are first translocated in a Sec-dependent manner and thus possess N-terminal signal sequences. Such signal peptides, however, belong to class 2 as it exhibits a conserved lipobox motif in the C-domain.³⁰ It can be noticed, however, that in *E. coli* YidC plays an important role in targeting and translocation of some lipoproteins.¹⁸³ Lipobox includes invariably a cysteine residue located just after the cleavage site of signal peptide. After translocation of the prolipoprotein, a common post-translational modification involves a prolipoprotein diacylglyceryl transferase (Lgt), which adds an N-acyl diglyceride group from a glycerophospholipid to the SH-group of the lipobox cysteine.¹⁸⁴ This thioether linkage allows protein anchoring to the membrane thanks to the insertion of the diacylglyceryl group into the lipid bilayer. Subsequently, SPase II (also called Lsp for lipoprotein signal peptidase) cleaves off the signal peptide and the cysteine becomes the N-terminal residue.¹⁸⁵ In contrast to *E. coli*, however, lipidation by Lgt (Lmo2482) in *Listeria* is neither essential for bacterial growth nor a prerequisite for activity of Lsp.¹⁸⁶ Once signal peptide is cleaved off, the amino-terminal cysteine residue is usually acylated at the free amino group by a phospholipid/apolipoprotein transacylase (Lnt), resulting

TABLE 12.5
Bioinformatic Resources for Prediction of Lipoproteins

Application	Method	Webserver	Ref.
ScanProsite	Profile (PS51257) search Pattern (G+LPP) search	http://www.expasy.org/tools/scanprosite/	189, 190
DOLOP	Pattern search	http://www.mrc-lmb.cam.ac.uk/genomes/dolop/	191
LipoP	Hidden Markov model	http://www.cbs.dtu.dk/services/LipoP/	192
SPELip	Neural network	http://gpcr.biocomp.unibo.it/predictors/	193
LipPred	Naïve Bayesian network	http://www.jenner.ac.uk/LipPred/	194

in protein anchoring to membrane long chain fatty acid.¹⁸⁷ This additional post-translational modification step, however, does not seem to be conserved in all bacteria,¹⁸⁴ as an *Int* gene is apparently lacking from all sequenced members of Firmicutes phylum, and *Listeria* species is no exception.¹⁸⁸ It can be further noticed that some lipoproteins are IMPs integrated to the cytoplasmic membrane by TMDs in a YidC-dependent manner.¹⁷³

As summarized in Table 12.5, several resources can be applied for genomic identification of lipoproteins. In PROSITE,¹⁸⁹ lipobox motif was previously referred to as PS00013 and defined by the regular expression {DERK}(6)-[LIVMFWSTAG](2)-[LIVMFYSTAGCQ]-[AGS]-C, where {DERK}(6) means that none of the four amino acids are allowed in the first six positions relative to the cleavage site. The pattern had two additional rules: (1) The cysteine must be between positions 15 and 35, and (2) at least one positively charged residue (K or R) must be within one of the first seven N-terminal residues. This pattern (i.e., a qualitative motif description based on a regular expression-like syntax) is now replaced by a profile (i.e., a quantitative motif description based on the generalized profile syntax), referred to as PS51257 and defined as prokaryotic membrane lipoprotein lipid attachment site profile. Originally, lipobox search using PS00013 was known to generate a significant proportion of false-positives, which prompted the need to improve the syntax of this regular expression. Thus, a refined pattern named G+LPP (for Gram-positive lipoprotein) and using PROSITE syntax—that is, [MV]-X(0,13)-[RK]-{DERKQ}(6,20)-[LIVMFESTAG]-[LVIAM][IVMSTAFG][AG]-C—was developed.¹⁹⁰ This pattern appears more specific for the identification of Gram-positive bacterial lipobox and allows greater discrimination against false-positives compared with PS00013. Thus, lipobox can be predicted by scanning polypeptidic sequences for the presence of PS51257 profile or G+LPP pattern using ScanProsite.¹¹⁵

However, correct sequence assignment as putative lipoprotein also requests that the lipobox is localized within an N-terminal Sec-dependent signal peptide where it covers H- and C-domains; signal peptide can be predicted following analysis with previously described tools (Table 12.1). DOLOP compiles similar criteria by scanning query sequences for presence of (1) a lipobox within the first 40 residues from the N-terminus with the consensus as [LVI][ASTV][ASG][C], (2) positively charged amino acid in n-domain of signal peptide, and (3) at least 7–22 residues between the predicted lipobox and the charged residue.¹⁹¹ LipoP is based on an HMM and discriminates among lipoprotein signal peptides (cleaved by SPase II), other signal peptides (cleaved by SPase I), n-terminal membrane helices, and cytoplasmic protein following attribution of scores.¹⁹² Despite having been primarily developed for Gram-negative bacteria, LipoP can also efficiently identify lipoproteins in Gram-positive bacteria. The only feeble point may be that when handling lipoproteins with transmembrane regions, the HMM misses, in some cases, the lipoprotein signal peptide. SPELip is an NN-based method for prediction of signal peptide and integrating a regular expression search based on PROSITE pattern.¹⁹³ LipPred is based on test against a naïve Bayesian network allowing the identification of lipoprotein with a calculated index for prediction confidence.¹⁹⁴ When compared to other available methods, LipPred can be considered as the most accurate for detection of lipoprotein signal sequence and SPase II cleavage site. Finally, lipoprotein-associated domains

can be searched using HMMs from Pfam or Tigrfam: namely, in Firmicutes, PF00938, PF01347, PF01540, PF02030, PF03180, PF03202, PF03260, PF03304, PF03305, PF03330, PF03640, PF04200, PF05481, TIGR00363, TIGR00413, TIGR01742, TIGR01533, and TIGR02898. Proteins identified following this approach should, however, be considered with great care and ideally confirmed by methods listed previously for lipoprotein modification/processing motif.

Following search for presence of PS00013 pattern and signal peptide using SignalP v2.0, 68 lipoproteins were originally identified in the genome of *L. monocytogenes* EGD-e.¹⁹ Pfam and Tigrfam searches allowed the identification of 70 lipoproteins in *L. monocytogenes* F2365, whereas the number of lipoproteins was estimated at 63 in *L. monocytogenes* EGD-e.¹⁸ Using the computational pipeline Augur,⁸⁹ where lipoproteins are identified on the basis of G+LPP pattern match, 65 lipoproteins were identified in *L. monocytogenes* EGD-e.¹⁸ Generating a new HMM from 26 verified lipoproteins by proteomic analysis in *L. monocytogenes* EGD-e, the number of lipoproteins was reestimated down to 62 in *L. monocytogenes* EGD-e and 56 in *L. monocytogenes* F2365.¹⁸⁶ Despite discrepancy in predicted number, lipoproteins constitute the largest family of putative surface proteins in *Listeria* (68 out of 133 originally predicted in *L. monocytogenes* EGD-e).¹⁹ These lipoproteins are putatively involved in various metabolic pathways (e.g., as substrate-binding components of ABC transport systems); remarkably, no biological function could be assigned for a large proportion. In spite of their predominance as surface proteins of Gram-positive bacteria, very few lipoproteins have been biochemically characterized.¹⁹⁵ In *Listeria*, only five have been more specifically investigated and thus confirmed, at least partially, in term of biological function: (1) TcsA (Lmo1388), a CD4⁺ T cell-stimulating antigen presented by major histocompatibility complex class II molecules¹⁹⁶; (2) GbuC (Lmo1016), a glycine betaine binding-protein part of an ABC transport system¹⁹⁷; (3) the substrate binding protein OpuCC (Lmo1426) part of an ABC L-carnitine transport¹⁹⁸; (4) OppA (Lmo2196), another ABC substrate binding protein mediating the transport of oligopeptides¹⁹⁹; and (5) the virulence factor LpeA (Lmo1847) involved in bacterial entry into eukaryotic infected cells.²⁰⁰

12.3.1.3 Extrinsic Membrane Proteins

No bioinformatic tool is currently available to identify such proteins, which are most of the time primarily predicted as extracellular or cytoplasmic depending on their presence on the external or internal side of the cytoplasmic membrane (Figure 12.8). Thus, their identification requires a deep understanding of bacterial physiology and excellent general literature survey. Some of these proteins, which are not intrinsic to the cytoplasmic membrane, can be subunits of membrane protein complexes such as F₁F₀ATP synthetase (GO:0045260), fumarate reductase complex (GO:0045284), or ABC (ATP binding cassette) transporters (GO: 0043190). Some other extrinsic membrane proteins can interact more or less temporarily with other membrane components, including other membrane-associated proteins. For example, in the SRP-dependent pathway, ribosomal proteins interact with Sec translocon in the course of cotranslational translocation,²⁰¹ or in two-component systems, response regulators interact with membrane bound sensors.²⁰² It should also be noticed that some cytoplasmic proteins can associate with the lipid bilayer by weak interactions and by no means be functionally associated with membrane components. To date, the number of extrinsic membrane proteins has never been estimated in *L. monocytogenes*.

12.3.2 CELL WALL-ASSOCIATED PROTEINS

Cell wall-associated proteins are either covalently linked to peptidoglycan when possessing a C-terminal LPXTG motif or noncovalently linked to cell wall components by a cell wall-binding domain (CBD).⁸ In Gram-positive bacteria, six CWBDs are currently characterized: CWBD of Type 1 (CWBD1), CWBD of Type 2 (CWBD2), Lysin motif domain (LysM), GW modules, S-layer homology domain (SLHD), and WXL domain (WXL).^{8,203} In *Listeria*, however, only proteins with

LPXTG, LysM, GW, and WXL motifs have been identified so far.^{182,203,204} In *L. monocytogenes*, proteins exhibiting such motifs systematically possess Sec-dependent N-terminal signal peptide. These motifs can be found using RPS-BLAST (reverse position-specific BLAST)²⁰⁵ or HMM²⁰⁶ from different databases—namely, InterPRO,²⁰⁷ Pfam,²⁰⁸ SMART,²⁰⁹ TIGRfam,²¹⁰ and SuperFamily.²¹¹

12.3.2.1 LPXTG Motif

LPXTG motif (IPR001899, PF00746, TIGR01167) is found in proteins covalently attached to cell wall by sortases⁵⁵ (see section 12.2.1). This motif consists of a pattern varying around LPXTG, a hydrophobic domain, and a positively charged tail.²¹² Cross-bridging of the protein to cell wall by sortase would occur in four steps.^{213,214} Following translocation across the Sec apparatus, membrane-associated sortase recognizes the LPXTG motif and cleaves it before linking proteins to cell wall precursor lipid II.²¹⁵ The proteins thus linked to lipid II are further incorporated into the cell wall by transglycosylation and transpeptidation reactions that generate peptidoglycan. Forty-one proteins with LPXTG motif substrates of StrA have been identified in *L. monocytogenes* EGD-e,¹⁹ whereas two proteins with an NXZTN motif are recognized by StrB.^{181,216} In *L. monocytogenes* F2365, a total of 44 LPXTG-like proteins have been identified.¹⁸ StrA is required for bacterial virulence as among its protein substrates several virulence factors have been identified, such as InlA.^{59,217} Compared to StrA, StrB plays a minor role both in terms of number of proteins anchored to cell wall and involvement in bacterial virulence, although virulence factor SvpA is substrate to StrB.²¹⁶

12.3.2.2 Noncovalently Attached Cell Wall Proteins

Even though most cell-associated proteins contain an LPXTG motif in *Listeria*, several proteins bear domains involved in noncovalent attachment to the components of cell wall. Three motifs are clearly established as involved in noncovalent cell wall attachment in *Listeria*: LysM, GW, and WXL.^{182,203,204} Besides these known attachment domains, other proteins found in the cell wall are retained by putative CWBDs, alternative or unknown mechanisms.

12.3.2.2.1 LysM

LysM (IPR002482, PF01476, SM00257, SSF54106) is a motif about 40 residues long and composed of three α -helices with a general peptidoglycan binding function.²¹⁸ It is found in a variety of enzymes mostly involved in bacterial cell wall degradation. When present, this motif is often repeated several times in the protein sequence. Interestingly, proteins bearing this motif can attach to surface of Gram-positive bacteria other than the ones that synthesized it.²¹⁸ In *L. monocytogenes*, several proteins bear LysM domains; among others, P60 (also called Iap or CwhA) can be cited as it is also considered as a virulence factor.²¹⁹ It is worth stressing here that contrary to previous assumption,^{19,182} NlpC/P60 domain should not be considered as the motif involved in cell wall binding *stricto sensu* (see section 12.3.2.2.4). In *L. monocytogenes* EGD-e and F2365 genomes, six LysM proteins, including P60 orthologue, were identified following bioinformatic analysis (Desvaux and Hébraud, unpublished data).

12.3.2.2.2 GW

GW (SSF82057) module was originally identified in *L. monocytogenes* within internalin InlB.²²⁰ This module is about 80 amino acids, contains a highly conserved dipeptide Gly-Trp, and interacts with lipoteichoic acids allowing cell surface attachment.¹⁸² GW modules are found in multicopy, as in InlB, where three copies are present in the C-terminal region, or in Ami, where eight modules are present. It also appears that the higher the number of GW modules is, the stronger is the attachment to the bacterial cell wall^{221,222}; proteins exhibiting only one GW module would not be surface attached at all. It is interesting to note that GW modules are related to Src homology-3 (SH3) clan (CL0010) and more specifically prokaryotic SH3 of Type 3, or SH3b (IPR013247, PF08239), but are unlikely to act as functional mimics of SH3 domains since proline-binding sites are blocked or

destroyed in GW domains.²²³ In *L. monocytogenes* EGD-e, nine GW proteins were identified, most of which (including Ami) exhibit an amidase domain.^{19,204} This indicates that this class of protein would be mainly involved in cell wall degradation in *L. monocytogenes*, although Ami is also considered as a virulence factor involved in bacterial adhesion to infected cells.^{182,224}

12.3.2.2.3 WXL

Following genomic analysis of *Enterococcus faecalis*, a novel C-terminal cell wall binding motif was uncovered and named WXL domain.²⁰³ This conserved domain is characterized by a first WXL motif and an additional YXXX[LIV]TWXLXXXP motif found further downstream; the two WXL domains are separated by between 66 and 247 residues. WXL domain was found in CDS of several genomes of low G+C Gram-positive bacteria, including *L. monocytogenes* where four proteins bearing such domain were identified.²⁰³ In *E. faecalis*, it was demonstrated that WXL domain is a determinant of bacterial subcellular protein. Indeed, its presence conferred cell surface display of the protein, whereas specific deletions into the domains prevented its display. Moreover, neither proteins nor carbohydrates were necessary for cell wall attachment but peptidoglycan was a binding ligand for WXL domain. As LysM, WXL can attach to cell wall of a variety of Gram-positive bacteria. From genome-wide analysis of Gram-positive bacteria,²²⁵ it appeared that genes encoding WXL proteins seem to cluster and that some N-terminal regions of these proteins are involved in utilization of plant complex polysaccharides. It was also suggested that some WXL proteins might mediate interactions between different bacteria species.²⁰³ In *Listeria*, physiological function of such proteins awaits to be established and their presence on bacterial cell surface remains to be demonstrated.

12.3.2.2.4 Other Noncovalently Attached Cell Wall Proteins

ChW motif (IPR006637, SM00728, PF07538) stands for clostridial hydrophobic domain with a conserved W residue and was first uncovered from bioinformatic analysis of *Clostridium acetobutylicum* genome.²²⁶ As GW, ChW contains a highly conserved Gly-Trp dipeptide motif and was suggested to be involved in cell surface attachment.³³ This repetitive domain can be found several times along the protein sequence (up to nine copies). ChW proteins were speculated to be part of a molecular complex on bacterial cell surface dedicated to degradation of polymer and surface adhesion.²²⁶ One putative serine protease bearing three copies of ChW motif is encoded in genome of *L. monocytogenes* F2365 (Desvaux and Hébraud, unpublished data); its expression, secretion, cell surface display, and function remain to be established. Similarly, function of SH3b is as yet unknown, but *Staphylococcus simulans* lysostaphin contains such a domain in its C-terminal region.²²⁷ Since this region mediates protein binding to bacterial cell wall, SH3b may have this function.

It is important to note that despite the absence of cell wall binding motifs in some enzymes involved in cell wall degradation, such proteins have affinity for cell wall components via their enzymatic active site. Thus, secreted proteins with cell wall degradation domains, such as NlpC/P600 (IPR000064, PF00877)²²⁸ or N-acetylmuramoyl-L-alanine (IPR002508, PF01520), can be localized in cell wall. However, such domains should not be considered as cell wall binding motifs per se since primary function of these enzymes is to cleave cell wall components following when they find a new cleavage site or are released into the extracellular milieu. In *L. monocytogenes*, these enzymes are involved in numerous cellular processes.¹²⁶

It is now well known that many metabolic enzymes can be surface localized in Gram-positive bacteria.²²⁹ Such proteins lack N-terminal signal peptide and are supposedly secreted by pathways alternative to the known ones. SecretomeP is a bioinformatic tool dedicated to the prediction of such proteins.²³⁰ However, instead of secretion through the cytoplasmic membrane, these proteins could be released from the bacterial cell following autolysis and then attached to cell surface of nonlysed cells. In *Streptococcus pneumoniae*, release of cytoplasmic proteins is triggered by competent cells and originates from lysis of noncompetent cells.²³¹ This tightly controlled phenomenon was named allolysis and involves several cell wall hydrolases. Once released and associated with bacterial cell surface, such proteins generally moonlight.²³ For example, in *S. pneumoniae*, the glycolytic enzyme

glyceraldehyde 3-phosphate dehydrogenase genolase exhibits plasmin(ogen)-binding activity on the bacterial cell surface and thus significantly enhances bacterial virulence.²³² In *L. monocytogenes*, several proteins primarily predicted as cytoplasmic were also identified in the cell wall fraction, including enolase, which was demonstrated to bind human plasminogen.²⁴ Functions of other cytoplasmic proteins found at this subcellular location (e.g., chaperone DnaK, elongation factor TU, or glyceraldehyde-3-phosphate dehydrogenase) remain to be elucidated as well as protein motifs involved in cell wall attachment.

12.4 CONCLUSIONS AND PERSPECTIVES

The proteomic technologies are certainly the most powerful and appropriate to provide global and accurate information on the expression, structure, and function of proteins. Since the first description of protein extraction and separation using two-dimensional gel electrophoresis (2-DE) in 1975,^{233–235} many advances (use of new detergents, immobilized pH gradient, new apparatus for IsoElectroFocalization [first dimension], and SDS-PAGE [second dimension]) have been brought to improve protein solubilization and resolution as well as reproducibility and implementation of the techniques. Over the past two decades, 2-DE progressively became the classical method of choice to separate and compare complex mixtures of proteins and was mainly applied for soluble intracellular proteins. In the field of bacteriology, an increasing number of investigations using comparative proteomic approaches was devoted to the characterization of adaptive responses—namely, to various physicochemical stresses or to the effect of a gene mutation. At this time, however, protein identification was difficult and time consuming as it essentially involved Edman degradation and sequence alignment from short and on limited numbers of amino acid sequences. Consequently, these early studies generally remained quite descriptive and phenomenological.

However, two occurrences gave a considerable impetus to proteomic analyses: (1) the availability of ever growing amounts of genomic sequence data, and (2) important advances in mass spectrometry technology for ionization and detection of large molecules such as peptides and proteins. Indeed, data obtained with mass spectrometry analysis—namely, peptide mass fingerprinting or fragmentation—could from then be matched against databases of all known gene products and thus greatly facilitated protein identification. The use of 2-DE and MS tools to separate and identify proteins is now widespread in all domains of life science. One of the consequences of this remarkable progress was the possibility to establish 2-DE databases containing several hundreds of identified protein spots available on proteome reference maps. Thus, the first bacterial 2-DE database was established on *E. coli* cytosoluble proteins separated into different pH gradients and regularly brought up to date.^{236,237} Similar but generally more limited 2-DE databases are available for other bacterial species, including *L. monocytogenes* EGD-e²³⁸ (<http://www.clermont.inra.fr/proteome>).

Another consequence was the possibility and need to investigate further the different cell compartments and thus following cell fractionation to explore thoroughly the different subcellular proteomes that, in Gram-positive bacteria, include (1) cytoplasmic proteins, (2) membrane associated proteins, (3) cell-wall associated proteins, and (4) proteins secreted in the extracellular milieu. While extracting and separating cytoplasmic and extracellular proteins could be achieved rather easily and efficiently, classical 2-DE procedures failed to give a good overview of proteins present in the cell envelope—that is, proteins associated to cytoplasmic membrane or cell wall. Beyond the well-known limitations of 2-DE gel-based technology (the inability to separate or to reveal low-abundance proteins, high molecular mass and extreme *pI* [isoelectric point] proteins), other limitations appeared much more problematic for the cell surface subproteomes due to the intrinsic properties of cell envelope associated proteins. Indeed, multitransmembrane proteins are generally highly hydrophobic and are either almost impossible to solubilize during the extraction procedure or not recovered in the second dimension of 2-DE due to self-aggregation and irreversible precipitation in IEF. On the other hand, proteins noncovalently or *a fortiori* covalently attached to the cell wall peptidoglycan are very difficult to extract and require specific and laborious treatments not directly

compatible with classical 2-DE separation. Altogether, this considerably hampers proteomic analysis in classical 2-DE gel-based technology.

Different strategies are now developed to tackle the difficulties to analyze these cell envelope subproteomes. These strategies can associate different protocols of protein extraction with different techniques of separation and mass spectrometry.²³⁹ Several studies have attempted to extract membrane associated proteins of Gram-positive bacteria by combining protocols described for Gram-negative or eukaryotes organisms.²⁴⁰ Thus, the extraction procedures could include enzymatic treatment, fractionation of broken cell by centrifugation, use of chemical agents such as zwitterionic detergents for solubilization of hydrophobic proteins,²⁴¹ solvents for delipidation,²⁴² or protein extraction and separation.²⁴³ For example, such a combinational approach has been used to efficiently characterize by 2-DE the cell-wall and membrane-associated subproteomes of the Gram-positive bacterium *Staphylococcus xylosus*.²⁴⁴

A different protocol originally developed for *Bacillus cereus*²⁴⁵ has been applied for the global extraction of *L. monocytogenes* cell-surface proteins combining the protein solubilization by SDS with a classical SDS-PAGE separation.²⁴⁶ The 1-DE and 2-DE separations were both used to characterize the cell wall subproteome of *L. monocytogenes*.²⁴ In this case, proteins were extracted by the sequential action of two salts at high concentration and their identification was performed by N-terminal sequencing and peptide mass fingerprint obtained with matrix assisted laser desorption ionization time of flight mass spectrometer (MALDI-TOF MS). It is interesting to note that among the 55 identified proteins, only 27 possessed a peptide signal, including 4 proteins with cell wall-binding motifs (2 GW proteins and 2 LysM proteins), 20 lipoproteins, and 3 proteins with no predictable surface association motif. The 28 remaining proteins without peptide signal were primarily predicted with cytoplasmic functions and nothing could explain how they managed to cross the cytoplasmic membrane or how they associated with bacterial cell wall. Such unusual localization of cytoplasmic proteins leads to the suggestion that they could moonlight on the bacterial cell surface, although no experimental evidence could back up such a hypothesis.

More recent strategies and technologies consist in analyzing peptide mixture obtained by tryptic digestion of cell envelope protein samples issued from stringent protocols extraction (e.g., combining cell mechanic broken) fractionation by centrifugation, and treatment with high concentration of SDS at 100°C, then at 80°C.^{58,247} An alternative approach consists in “shaving” the bacterial surface with a specific protease (such as trypsin) to cleave surface-exposed proteins.²³⁹ After lyophilization to remove SDS, the peptide hydrolysate is then separated by two-dimensional liquid chromatography coupled to tandem mass spectrometer (2-D LC MS/MS). This separation technique, termed “shotgun proteomic” or multidimensional protein identification technology (MudPIT), uses a two-dimensional liquid chromatography to separate a tryptic peptide mixture where a strong cation exchange is applied in the first dimension and a reverse phase is applied in the second dimension. The separated peptides are subjected online to analysis by fragmentation (MS/MS) in an electrospray ionization MS. Peptide fragmentation spectrum is further used to identify the original protein via query against databases.

Besides a significant gain of time, the use of 2-D LC MS/MS overcomes all limitations of gel-based 2-DE previously cited. Even hydrophobic proteins can be identified thanks to amino acid cleavage sites accessible to tryptic digestion in exposed regions of the protein. These new approaches allowed extraction of proteins following treatments that are not always compatible with classical 2-DE. Consequently, the set of proteins identified with high number of transmembrane spanning regions or LPXTG motif (i.e., covalently anchored to cell wall) has been significantly enlarged. The other development of 2-D LC MS/MS concerns the possibility to perform quantitative proteomics for comparative analysis of samples pretreated with amino acid tags or labels such as the ICATTM (isotope-coded affinity tags), iTRAQTM, or SILAC technologies.²³⁹

The years to come will undoubtedly see the development and improvement of these new exploring methods of subproteomes. Everyone will have the possibility to map protein expression and to compare several biological samples with high throughput, sensibility, and resolution. In spite of

this very attractive progress and considering at least its complementarity, classical 2-DE technique remains irreplaceable. Indeed and contrary to LC MS/MS approaches, gel-based 2-DE allows one to separate simultaneously several hundreds of proteins at once and to visualize shifts due to post-translational modifications. The implementation of two or more complementary proteomic strategies would be one of the keys to generate valuable information on the role of cell envelope proteins in pathogenic processes, bacterial communication, sensing of and exchange with its environment, motility, and adhesion on and colonization of biotic or abiotic surfaces.

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