A novel cell wall-anchored peptidoglycan hydrolase (autolysin), IspC, essential for *Listeria monocytogenes* virulence: genetic and proteomic analysis

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We have recently concluded that a *Listeria monocytogenes* 86 kDa immunogenic surface protein, IspC, is a cell wall-anchored peptidoglycan hydrolase (autolysin), capable of degrading the cell wall peptidoglycan of the bacterium itself. To determine if this enzyme has any biological functions and/or plays a role in virulence, we in-frame-deleted the *ispC* gene from the *L. monocytogenes* chromosome. This ΔispC mutant exhibited complete abrogation of expression of IspC and displayed no defects in *in vitro* growth, colony and microscopic morphologies, or biochemical characteristics. Lack of IspC led to attenuated virulence in mice, evidenced by a significant reduction in bacterial counts in livers and brains and no mortality compared with the wild-type. Furthermore, the data from assays using various eukaryotic cells for adhesion, invasion, actin tail formation, plaque formation and intracellular growth indicated that the mutant was severely attenuated in virulence in a cell culture model in a cell type-dependent manner. The findings that (i) the mutant was impaired for adhesion to certain eukaryotic cells, and (ii) both purified IspC and its C-terminal cell wall-binding domain were capable of binding sheep choroid plexus (SCP) epithelial cells and Vero cells, supported the role of IspC as an adhesin in virulence. The ΔispC mutant exhibited a marked defect in adhesion to and invasion of SCP cells but not human brain microvascular endothelial cells (HBMEC), suggesting that IspC is necessary for crossing the blood–cerebrospinal fluid barrier. Proteomic and immunological analysis showed a reduced surface expression of some known or putative virulence factors (e.g. ActA, InlC2 and a flagellin homologue, FlaA) due to IspC deficiency. Altogether, this study demonstrates that IspC, expressed as a minor autolysin *in vitro*, is not important for cell division or separation but is essential for full virulence of *L. monocytogenes in vivo*.

**INTRODUCTION**

Autolysins are peptidoglycan hydrolases that are capable of cleaving the covalent bonds in the cell wall peptidoglycan (murein) of bacteria from which they are produced (Shockman & Holtje, 1994). Several autolysins from *Listeria monocytogenes*, a Gram-positive intracellular bacterium that infects humans leading to the development of a severe food-borne illness, including p60, P45, Ami, MurA and Auto, have been identified and characterized (Popowska, 2004). Bacterial autolysins are involved or implicated in pathogenesis (Berry *et al.*, 1989; Cabanes *et al.*, 2004; Canvin *et al.*, 1995; Lenz *et al.*, 2003; Lock *et al.*, 1992; Mani *et al.*, 1994; Milohanic *et al.*, 2001) in addition to their biological functions, involving cell wall expansion, cell division, cell separation, chemotaxis, biofilm formation, genetic competence, protein secretion, antibiotic-induced lysis, sporulation, and formation of flagella (Smith *et al.*, 2000). Autolysin-deficient mutants, including an *LytA* mutant of *Streptococcus pneumoniae* (Berry & Paton, 2000), an *AtlE* mutant of *Staphylococcus epidermidis* (Rupp *et al.*, 1999), and Ami, *Auto*, p60 and *MurA* mutants of *L. monocytogenes* (Cabanes *et al.*, 2004; Lenz *et al.*, 2003; Milohanic *et al.*, 2001; Pilgrim *et al.*, 2003) are less virulent in animal models than their parental wild-type strains. The mechanisms that underlie the involvement of autolysins in virulence are not fully understood and need to be investigated further. The autolytic activity and the cell wall-binding domain (CWBD) of bacterial autolysins have been shown to be involved in virulence. LytA of *Strep. pneumoniae* and MurA and p60 of *L. monocytogenes* are thought to mediate the release of cytoplasmic toxins or...
proinflammatory deceased cell wall components resulting in subsequent tissue injuries (Jedrzejas, 2001; Lenz et al., 2003; Lock et al., 1992; Tuomanen, 2000). Restoration of virulence in an L. monocytogenes p60-defective mutant strain requires expression of the full-length autolysin p60 with an intact catalytic domain (Lenz et al., 2003). The CWBDs of autolysins, made up of repeated glycine-tryptophan (GW) dipeptide modules, contribute to bacterial virulence by promoting adherence to eukaryotic cells, as shown for Ami of L. monocytogenes (Milohanic et al., 2001), and to extracellular matrix proteins, as shown for Aas of Staphylococcus saprophyticus (Hell et al., 1998) and AtlC of Staphylococcus caprae (Allignet et al., 2002).

We have recently demonstrated the peptidoglycan hydrolase (autolysin) activity of IspC, an L. monocytogenes (serotype 4b) immunogenic surface protein of 774 aa with a calculated molecular mass of 85.87 kDa and a theoretical pI of 9.4 (Wang & Lin, 2007; Yu et al., 2007). IspC is similar in modular domain structure to Auto and Ami of L. monocytogenes and contains an N-terminal catalytic domain (amino acids 24–197) and a C-terminal CWBD (amino acids 198–774) consisting of seven GW modules (Wang et al., 2007; Wang & Lin, 2007). Several lines of experimental evidence accumulated in our recent studies (Wang & Lin, 2007; Yu et al., 2007) have suggested that IspC is potentially involved in pathogenesis. This evidence includes the following: (i) antibody to IspC is present in rabbits infected with live L. monocytogenes but not in animals receiving heat-killed bacteria, suggesting that this protein is specifically induced or upregulated in vivo during infection; (ii) IspC is localized on the cell surface; and (iii) the protein possesses autolytic activity. Both Ami and Auto of L. monocytogenes are involved in pathogenesis, but they act via different mechanisms (Cabanes et al., 2004; Milohanic et al., 2001). Ami contributes to bacterial adhesion to eukaryotic cells, whereas Auto is not involved in adhesion but contributes to the entry of L. monocytogenes into eukaryotic cells. This suggests the necessity of examining the role of the newly identified L. monocytogenes autolysin IspC in pathogenesis.

Here we conducted an analysis of the L. monocytogenes ispC gene and its product using genetic and proteomic approaches in an attempt to define the biological function of this cell-wall-anchored peptidoglycan hydrolase and its role in virulence. Data obtained from this investigation indicate that IspC is a novel factor that contributes to the virulence of L. monocytogenes through mechanisms involving the adhesive properties of the C-terminal CWBD, and that it regulates the surface display of other virulence factors, presumably by the autolytic activity of its N-terminal catalytic domain.

METHODS

Bacterial culture and plasmid vector. The L. monocytogenes serotype 4b strain LI0521 (wild-type) and its ispC in-frame deletion mutant strain (ΔispC) were grown in brain heart infusion (BHI) broth or on BHI agar plates. Escherichia coli DH5α was used in cloning experiments and cultured in Luria–Bertani (LB) medium.

Purification of recombinant proteins. The recombinant IspC and its C-terminal CWBD (amino acids 198–774) fused C-terminally to a GFP variant optimized for brighter fluorescence when excited by UV light (GFPuv–CBD1) were expressed in E. coli and purified to electrophoretic homogeneity by using Ni-NTA agarose affinity chromatography followed by SP Sepharose cation-exchange chromatography (for IspC only), as described previously (Wang et al., 2007; Wang & Lin, 2007).

Construction of ispC in-frame deletion mutant. A DNA fragment composed of the 453 bp sequence upstream of the ispC ORF and the first three codons was derived by PCR from the L. monocytogenes (serotype 4b strain LI0521) genomic DNA with the primer pair P517 (5'-ATCGAGTCAGGAAATCATAAAGGCTCCTCA-3') and P496 (5'-TAAAAGCTGAATCTTATTACCTATCGTG-3') and P518 (5'-AAGGTCGATTGATGAAAGCTAC-3') and SalI site underlined). A DNA fragment containing the 405 bp sequence downstream of the ispC ORF and the last eight C-terminal codons was similarly obtained by PCR with the primer pair P495 (5'-ATGATAAAATCGAGCTTTTACAAACGTTAAATAGAC-3') and P518 (5'-AAAAGTCGATTGAATGAAAGCTAC-3') and SalI site underlined. The two PCR fragments were spliced together as described previously (Wang & Lin, 2007) using primers P495 and P496, resulting in an internal deletion of 764 aa of the coding sequence within ispC. The fused DNA fragment was inserted into the SacI and SalI sites of pAUL-A (kindly provided by Professor T. Chakraborty, Institute of Medical Microbiology, Justus-Liebig University, Germany) to create pAUL-AΔispC. Following electrophoration of the recombinant plasmid into competent E. coli DH5α, bacteria were screened for targeted gene deletion as described elsewhere (Schafferkordt et al., 1998). A ΔispC mutant was confirmed by PCR analysis of the genomic DNA using a primer pair internal to the deletion region [P304 (5'-GATGTTAAAACTGTCCTTATTAC-3') and P285 (5'-TTAGCTCCTCGGATCCTAAATAGAC-3')] and a primer pair external to the deletion region [P481 (5'-CAGCCAAGAAGATTGTCGCTAAA-3')] and P518], by sequencing the PCR products.

Confirmation of the ΔispC mutant at the protein level. Surface expression of IspC was examined by immunofluorescence microscopy analysis of live bacteria (the ΔispC mutant and the wild-type) probed with rabbit antiserum raised against purified recombinant IspC (RzispC), as described previously (Wang & Lin, 2007). IspC in the total protein extracts (see below) from the mutant and wild-type strains was also analysed by Western blotting using RzispC. The bacteriolytic activity of the wild-type and the ΔispC mutant was assessed by analysis of the cell surface protein extracts (see below) equivalent to 1 ml of culture with an OD620 of 1.0 in a 12 % renaturing SDS-PAGE gel containing 0.2 % (w/v) autoclaved Micrococcus lysodeikticus ATCC 4698 (Sigma), as described previously (Wang & Lin, 2007).

Passage of L. monocytogenes in mice. The ΔispC mutant and wild-type strains were passaged in vivo in mice to eliminate the possibility that in vitro growth may reduce the bacterial virulence. Both strains (100 μl, 2 × 109 cells ml−1) in PBS, pH 7.2, were intravenously inoculated into BALB/c mice. Each strain recovered from livers of infected mice at day 3 post-infection was grown to OD620 equivalent to 1 ml of culture with an OD620 of 1.0 in a 12 % renaturing SDS-PAGE gel containing 0.2 % (w/v) autoclaved Micrococcus lysodeikticus ATCC 4698 (Sigma), as described previously (Wang & Lin, 2007).

Phenotypic analysis. Comparative phenotypic analysis of the ΔispC mutant and the wild-type with respect to their in vivo growth, and biochemical and morphological characteristics was performed. The
growth curve was established by subculturing each strain at the same starting concentration from the overnight culture in triplicates in 50 ml BHI broth (pH 7.2), BHI broth (pH 7.2) containing 0.3 M NaCl (osmotic stress), and BHI broth (pH 4.5; acidic stress). Samples of cultures were taken at various time points for 48 h to measure OD_{620}. The catechol test, Christie–Atkins–Munch-Petersen (CAMP) test (Staphylococcus aureus and Rhodococcus equi), H$_2$S test, oxidase test, nitrate reduction test and motility test (25 and 35 °C) were conducted according to established methods (MacFaddin, 2000). Carbohydrate utilization under fermentative and oxidative conditions was tested by using API 50 CH strips (bioMérieux) using CHB/E medium according to the manufacturer’s instructions. The colony morphology of the mutant was examined by culturing bacteria on tryptic soy blood agar (TSBA) plates. The microscopic morphology of bacterial cells was examined at various phases of growth (i.e. early exponential, mid-exponential, late-exponential and stationary phases) using phase-contrast light microscopy, transmisision electron microscopy (TEM) and cross-sectional TEM.

**Extraction of total bacterial proteins.** The ΔispC mutant and wild-type strains at early exponential growth phase, equivalent to 5 ml of culture with an OD$_{620}$ of 1.0, were treated with chloramphenicol (20 μg ml$^{-1}$) to inhibit protein synthesis and washed with ice-cold PBS containing protease inhibitor cocktail (Roche) and 20 μg chloramphenicol ml$^{-1}$ (PBS-CC). Bacteria were lysed in 500 μl PBS-CC in a FastPrep Blue tube on a FastPrep apparatus FP120 (Qiagen) with five bursts of 45 s on setting 6.5. The cell lysates were boiled for 10 min after addition of 500 μl 2 × SDS-PAGE sample buffer [0.1 M Tris/HCl, pH 6.8, 40 % (v/v) glycerol, 20 % (v/v) β-mercaptoethanol, 4 % (w/v) SDS and 0.02 % (w/v) bromophenol blue] and centrifuged at 14,100 × g to collect total bacterial proteins (the supernatant) for SDS-PAGE analysis.

**Extraction of bacterial surface proteins.** Samples of the ΔispC mutant and wild-type strains equivalent to 50 ml of culture with an OD$_{620}$ of 0.5 were collected at various growth phases (early exponential, mid-exponential, late-exponential and stationary) and washed as described above. Cell pellets were resuspended in 1 ml 2 × SDS-PAGE sample buffer and boiled for 10 min. The supernatants containing surface-extracted proteins were collected by centrifugation at 14,100 × g for 5 min and stored at −20 °C until use.

**Analysis of cell surface proteins by MS and Western blotting.** Bacterial surface proteins (150 μl) from the ΔispC mutant and wild-type strains were separated by SDS-PAGE using a 4 % stacking gel and a 12 % resolving gel on a Protein II xi system (Bio-Rad) and stained with Coomassie brilliant blue. Resolved protein bands of interest were excised from the wild-type samples and subjected to in-gel tryptic digestion for liquid chromatography–tandem MS (LC–MS–MS) analysis, as described elsewhere (Vasilescu et al., 2005). The MS–MS data were then analysed against the protein database using the Mascot search engine. Western blot analysis of surface proteins was performed as described previously (Wang & Lin, 2007), with rabbit anti-InLA, anti-InLB, anti-ActA and anti-InIC2 antibodies.

**In vivo virulence assay.** Six- to eight-week-old female BALB/c mice were used in in vivo infection experiments with the ΔispC mutant and the wild-type strain. For each bacterial strain, four groups of six mice each were intravenously inoculated with 5 × 10$^8$ bacteria in 100 μl PBS. Brains, livers and spleens were aseptically removed from one particular group (n=6) at 6, 24, 48 and 72 h post-infection, homogenized in 5 ml 1% buffered peptone water (BPW) using a stomacher, and spread on BHI agar plates with 0.1 ml of homogenates at 10-fold serial dilutions. Bacteria were enumerated after incubation at 37 °C for 24 h.

**Cell lines and cell culture.** The cell lines Caco-2 (human colon carcinoma enterocyte-like epithelial cell), Hep-G2 (human hepatocellular carcinoma cell), Vero (African green monkey kidney cell), L132 (human embryonic lung fibroblast), Hela (human cervical epithelial cell), HBMEC (human brain microvascular endothelial cell), SCP (sheep choroid plexus epithelial cell), L2 (murine fibroblast cell) and J774 (murine macrophage) were grown in MEM complete medium (1 × minimal essential medium supplemented with 10% heat-inactivated fetal bovine serum (FBS) or 5% horse serum for SCP, 0.1 mM non-essential amino acids, 2 mM L-glutamine and 0.15% sodium bicarbonate). Sodium pyruvate was added at 1 mM into the MEM complete medium for Hep-G2. HBMEC (Cell Systems) was cultured on gelatin-coated surface in Cell Systems complete (CS-C) medium according to the manufacturer’s instructions. All eukaryotic cells were grown at 37 °C and 5% CO$_2$ in a humidified incubator.

**Adhesion, invasion and intracellular growth assays.** The ability of bacteria with respect to adhesion to-, invasion of-, and intracellular growth within eukaryotic cells was assessed basically as described elsewhere (Bergmann et al., 2002; Rowan et al., 2000), with some minor modifications. Briefly, 2 × 10$^5$ cells (1 × 10$^5$ HBMEC cells) were seeded in 24-well tissue culture plates (VWR), grown for 48 h, and infected with 0.5 ml bacterial suspension (ΔispC mutant or wild-type strain) in MEM complete medium (for HBMEC, 25 mM HEPES and 1 mM sodium pyruvate was added to the medium) for 1 h at the desired m.o.i. for each cell line: m.o.i. 50 for Hela and L132, m.o.i. 100 for Vero, m.o.i. 20 for Caco-2, Hep-G2, SCP and HBMEC, and 2 × 10$^5$ bacteria for J774. For the adhesion assay, eukaryotic cells were washed five times with 3 ml PBS [Dulbecco’s Modified Eagle Medium (DMEM; high glucose) for HBMEC] and lysed by 1 ml 1% (v/v) Triton X-100. Viable bacteria were enumerated by plating 0.1 ml lysates at 10-fold serial dilutions on BHI agar plates. For the invasion assay, infected eukaryotic cells were washed and incubated in 1 ml medium containing 100 μg gentamicin ml$^{-1}$ for 1.5 h. After washing, surviving intracellular bacteria were quantified as described above. For intracellular growth assays, infected eukaryotic cells were washed and incubated in medium containing 100 μg gentamicin ml$^{-1}$. Surviving bacteria were quantified as above at 2, 4, 6 and 8 h after addition of gentamicin. All cellular adhesion, invasion and growth assays were performed in triplicate.

**Actin tail formation.** J774 cells (2 × 10$^5$ per well) were seeded into two-well Lab-Tek chamber slides (Fisher) and grown to ~90% confluence. After washing with DMEM minus FBS (37 °C), cells were infected with 10$^5$ bacteria (mutant or wild type) in 1 ml of the same medium for 1 h, and then incubated in complete MEM containing 100 μg gentamicin ml$^{-1}$ for an additional 3 or 6 h. After washing with PBS, cells were fixed in 4% (v/v) paraformaldehyde in PBS for 12 min, permeabilized in 0.2% (v/v) Triton X-100 in PBS for 8 min, and blocked in 3% (w/v) BSA in PBS for 30 min. Bacteria were stained with rabbit antiserum to ActA for 1 h followed by incubation with Alexa Fluor 647 goat anti-rabbit secondary antibody (Invitrogen) for 1 h. Actin tails were stained with Alexa Fluor 488 Phalloidin (Invitrogen) for 30 min. Nuclei were stained with 4’,6-diamidino-2-phenylindole (DAPI; Sigma) at 10 μM in PBS for 3 min. Slides were mounted with Vectashield mounting medium (Vector Laboratories) and examined on an Olympus BX51 fluorescence microscope (Olympus Canada).

**Immunofluorescence and immunogold TEM.** Immunofluorescence and immunogold TEM procedures (Wang & Lin, 2007) were used to detect the in situ surface expression of IspC or ActA in live bacteria of the ΔispC mutant and wild-type strains.

**Plaque assay.** Murine L2 cells (3 × 10$^5$ per well) were seeded into six-well plates (VWR) and grown to confluence. After washing with DMEM minus FBS (37 °C), cells were infected with 10$^5$ bacteria.
(ΔispC mutant or wild-type strain) in 1 ml of the same medium for 1 h. After washing with complete MEM medium (37 °C) containing 10 μg gentamicin ml⁻¹, 3 ml agar overlay (42 °C) containing 1 × DMEM, 10 % FBS, 10 μg gentamicin ml⁻¹ and 0.7 % agarose was added to each well, and the plates were incubated for 4 days at 37 °C. Plaques were visualized on a Zeiss Axiovert-10 inverted microscope equipped with a Nikon digital camera at a total magnification of 45 and their relative sizes (diameters) estimated using ImageJ version 1.37a (National Institutes of Health).

**RESULTS**

**Construction and phenotypic characterization of an ispC in-frame deletion mutant**

To elucidate the biological function of the 86 kDa autolysin IspC (Wang & Lin, 2007) and its potential role in virulence, an IspC-deficient *L. monocytogenes* strain was successfully created from the parental strain *L. monocytogenes* serotype 4b via homologous recombination using the pUAL-A-based integration–excision technique. This mutant, which has a chromosomal in-frame deletion of 2292 bp within the 2325 bp *ispC* ORF (Fig. 1a), designated ΔispC, was confirmed by PCR with two sets of primer pairs (Fig. 1b) followed by DNA sequencing. With primers P304 and P285, which target the deleted region, no PCR product was obtained from the ΔispC mutant chromosomal DNA, in contrast to a product derived from the wild-type strain that corresponded to its calculated size of 452 bp. With primers P481 and P518, which anneal outside the deleted region, a PCR product close to the expected size of 993 bp was amplified from the mutant chromosomal DNA and, as expected, was smaller than that (~3200 bp) derived from the wild-type. DNA sequencing of the PCR product derived from the ΔispC mutant with primers P481 and P518 also confirmed a target deletion of 2292 bp within the *ispC* ORF.

The ΔispC mutant exhibited the abrogation of IspC expression in *in vitro* culture, as evidenced by the data from three separate experiments. (i) Western blot analysis of total bacterial proteins revealed that a protein band of ~86 kDa from the wild-type recognized by rabbit anti-IspC polyclonal antibody (RzispC) was not present in the mutant (Fig. 1c). (ii) Renaturing SDS-PAGE analysis of the surface protein extract for peptidoglycan hydrodase activity with respect to *M. lysodeikticus* cell wall showed that a bacteriolytic band of ~86 kDa associated with the wild type was not found in the mutant (Fig. 1d). (iii) Immunofluorescence microscopy with RzispC showed no fluorescent staining on live mutant bacteria but detected IspC on the surface of wild-type bacteria (Fig. 1e).

Interestingly, surface peptidoglycan hydrodase activity profiles from the wild-type and the ΔispC mutant showed that IspC is a minor autolysin in *in vitro* culture.

The ΔispC mutant was indistinguishable from the wild-type strain with respect to the following: colony morphology on TSBA plates, bacterial morphology at various growth phases in BHI broth at 37 °C as revealed by phase-contrast microscopy (Fig. 1f), TEM (data not shown) and sectional TEM (Fig. 1g), motility at 25 and 35 °C, haemolysis on sheep blood agar plates, carbohydrate utilization on API 50 CH strips, growth curve at 37 °C in BHI broth or growth in BHI under acidic (pH 4.5) or osmotic (0.3 M NaCl) stress, and a number of biochemical tests (i.e. catalase test, CAMP test (Staph. aureus and *R. equi*), H₂S test, oxidas e test and nitrate reduction test).

**IspC is required for virulence *in vivo***

To evaluate whether the *ispC* gene contributes to virulence, BALB/c mice were infected intravenously with the ΔispC mutant and the wild-type strain. The deficiency of IspC in bacteria significantly (*P*<0.01) reduced the bacterial load in the target organs (by approximately fivefold in brain and approximately sevenfold in liver) at 48 h post-infection with the exception of spleen, while the bacterial counts in these organs were similar between the mutant and the wild-type at 6 and 24 h (Fig. 2). The requirement of IspC for virulence was further demonstrated by the fact that the mutant did not cause any mortality at 72 h, in contrast to the death of two out of six mice infected with the wild-type strain. This result prompted us to further investigate the role of IspC in virulence at the various stages of the infectious cycle, using cultured eukaryotic cells.

**Involvement of IspC in bacterial adhesion in a cell type-dependent manner**

In earlier studies, GW modules of several autolysins, including Ami of *L. monocytogenes* (Milohanic et al., 2001), Aas of *Staph. saprophyticus* (Hell et al., 1998) and AtlC of *Staph. caprae* (Allignet et al., 2002), were shown to be involved in bacterial adhesion. To determine if the surface autolysin IspC, which contains seven GW modules in its C-terminal CWBD (Wang & Lin, 2007) promotes the adherence of bacteria to eukaryotic cells, the ΔispC mutant and the wild-type were assayed for binding to several types of eukaryotic cells (Fig. 3). The primary attachment of the ΔispC mutant to Hep-G2, Vero and SCP was reduced by
approximately two- to threefold \((P<0.01)\) in comparison to that of the wild-type strain, while both strains were similar in adhesion to Caco-2, L132, Hela, HBMEC and mouse macrophage J774. This indicates that IspC plays a role in mediating bacterial adhesion to normally non-phagocytic cells in a cell type-dependent manner.

**IspC is required for invasion in a cell type-dependent manner**

To determine whether IspC contributes to the invading capability of *L. monocytogenes*, internalization of the \(\DeltaispC\) mutant and the wild-type into various normally non-phagocytic eukaryotic cells was investigated (Fig. 4). The mutant exhibited a reduction in internalization into Caco-2, Hep-G2, Vero, L132 and SCP by two- to threefold \((P<0.01)\) relative to the wild-type, while it entered Hela and HBMEC as efficiently as the wild-type. Thus, IspC is involved in invasion of normally non-phagocytic cells by *L. monocytogenes* in a cell type-dependent manner.

**Direct binding of recombinant IspC and its C-terminal CWBD to the surface of SCP and Vero cells**

The observation that the \(\DeltaispC\) mutant possessed weaker adhesive and invasive characteristics with respect to certain eukaryotic cell types prompted us to further study the adhesive properties of purified IspC and its C-terminal CWBD (amino acids 198–774) fused to GFPuv (GFPuv–CBD1) with respect to SCP and Vero cells; in these studies the mutant exhibited an impaired adhesion and invasion ability. Fluorescence microscopy showed that both the purified recombinant IspC and GFPuv–CBD1 bound to
SCP and Vero (Fig. 5). Stronger fluorescent signals were observed for binding of these proteins to SCP than to Vero. Fixation and fixation followed by permeabilization of these eukaryotic cells prior to fluorescence staining resulted in similar fluorescence images of cells (data not shown), indicating that the binding event occurred on the cell surface. Binding of IspC or its C-terminal CWBD to the cell surfaces of SCP and Vero was specific, as no fluorescence staining on the cell surfaces was observed when cells were probed with rabbit preimmune serum or purified GFPuv. Under the same assay conditions, almost no fluorescence was observed on the surface of HBMEC probed with purified IspC or GFPuv–CBD1 (data not shown).

Deletion of ispC impairs the display of surface proteins

To determine whether the chromosomal deletion of ispC would alter the display of bacterial surface proteins, including known virulence factors, as an alternative mechanism leading to the reduced adhesive and invasive capacity of the mutant, surface proteins from the ΔispC mutant and the wild-type at various growth phases were analysed by SDS-PAGE followed by MS (Fig. 6a) or by Western blotting (Fig. 6b) using specific antibodies. Two protein bands of ~90 and ~30 kDa present in the wild-type were undetectable in the mutant by SDS-PAGE, and were subsequently identified by MS as ActA and a flagellin protein (a FlaA homologue), respectively. Western blot analysis using rabbit antiserum to ActA showed that expression of ActA was not abolished in the mutant but was dramatically reduced at all growth phases, in contrast to the wild-type (Fig. 6b). The decrease in the amount of ActA was also observed with Western blot analysis of total cell lysates of the mutant (data not shown). Reduced surface display of ActA in the mutant was further demonstrated by immunofluorescence microscopy (Fig. 6c, top and middle panels) and immunogold TEM (Fig. 6c, bottom panel) probed with rabbit anti-ActA antibody. Western blot analysis using rabbit anti-InlC2 antibody revealed a significant reduction in InlC2 expression in the mutant, although this protein was not visible on the SDS-PAGE gel. Three surface proteins having a similar expression level in the mutant and the wild-type were identified to be InlB precursor and p60, by MS, and InlA, by Western blot analysis using rabbit anti-InlA antibody. The absence of an alteration in InlB expression was further confirmed by Western blot analysis using rabbit anti-InlB antibody.

IspC is involved in actin tail formation in the early stages of intracellular infection

The observation that the surface display of a few proteins, including ActA, was strikingly reduced in the ΔispC mutant prompted us to further investigate whether the abrogation of IspC expression affects the intracellular motility of
bacteria by examining, in comparison with the wild-type, the formation of actin tails in the intracellular niche after infection with the mutant of mouse macrophage J774 cells. The mutation led to formation of much shorter and weaker actin tails at the early infection stage (i.e. 3 h) than the wild-type, while at the late infection stage (i.e. 6 h), formation of the actin tails was similar in the mutant and the wild-type (Fig. 7).

IspC contributes to cell-to-cell spread

A contribution of IspC to cell-to-cell spread of bacteria may be expected, as the formation of actin tails was impaired in early stages of infection with the ΔispC mutant. This was macroscopically confirmed by a plaque assay of a murine L2 fibroblast monolayer infected with the mutant as compared with the wild-type. The relative plaque size

**Fig. 3.** Quantitative analysis of adhesion of the ΔispC mutant to various eukaryotic cell lines in comparison with the wild-type (WT). The mutant and wild-type strains were used to infect a particular cell line at a predetermined m.o.i. (see Methods) for 1 h and the bacteria associated with the eukaryotic cells were quantified by plating the cell lysates after washing away free bacteria. The quantity of mutant bacteria is calculated relative to that of the wild type (set as 100%) and presented as the mean ± SD (n=3). *Statistically significant difference (P<0.05).

**Fig. 4.** Quantitative analysis of entry of the ΔispC mutant into various eukaryotic cell lines. The mutant and wild-type (WT) strains were used to infect a particular cell line in triplicate at a predetermined m.o.i. (see Methods) for 1 h followed by further incubation with 100 μg gentamicin ml⁻¹ for 1.5 h. The intracellular bacteria were determined as in Fig. 3. The quantity of mutant bacteria is calculated relative to that of the wild type (set as 100%) and presented as the mean ± SD (n=3). *Statistically significant difference (P<0.05).
(expressed in arbitrary units; a.u.) formed by the mutant strain (175.5 ± 18.444 a.u., n=22) was significantly smaller (P<0.01) than that of the wild-type (206.375 ± 25.595 a.u., n=24) (Fig. 8), indicating the reduced capacity for bacterial cell-to-cell spread of the ΔispC strain.

IspC contributes to bacterial intracellular growth at the later infection stage

Autolysins have dual effects on bacterial growth and survival (favouring growth and causing bacterial death). The effect of IspC on intracellular survival and growth was assessed in two representative cell lines (Vero and J774) following infection with the ΔispC mutant and the wild-type over an 8 h time-course. The mutant showed an approximately twofold reduction in growth within both Vero and J774 cells compared with the wild-type (Fig. 9, Vero, P<0.01; J774, P<0.05) at 8 h post-infection, whereas the mutant was similar in growth rate to the wild-type before this time point, indicating that IspC extends the existence of intracellular bacteria and promotes bacterial intracellular growth at the later infection stage.

DISCUSSION

In this work, we showed evidence that a cell wall-anchored peptidoglycan hydrolase (autolysin), IspC (Wang & Lin, 2007), recognized also as the target of the humoral immune response to listerial infection (Yu et al., 2007) and expressed as a minor autolysin *in vitro* (this study), is not important for cell division or separation during *in vitro* growth but is required for full virulence of *L. monocytogenes*. By in-frame-deleting the *ispC* gene, we have determined the effect of IspC deficiency on phenotypic characteristics of *L. monocytogenes* (i.e. *in vitro* growth, colony and cell morphologies, and biochemical properties), virulence of the bacterium in mouse and eukaryotic cell models of infection, and display of other surface proteins.
The findings from these analyses and from the analysis of the binding to eukaryotic cells of the purified IspC and of its C-terminal CWBD (amino acids 198–774) fused to GFP (Wang & Lin, 2007) have shed new light on our understanding of the molecular mechanisms by which a surface peptidoglycan hydrolase contributes to bacterial pathogenesis.

The in vivo study demonstrated a marked attenuation of virulence of the ΔispC mutant for mice. Similarly, in vivo studies with animal models (mice, rats and guinea pigs) of infection have shown that mutant strains defective in the synthesis of autolysins, including AtlE of Staph. epidermidis (Rupp et al., 2001), p60, Ami and Auto of L. monocytogenes (Cabanes et al., 2004; Milohanic et al., 2001; Pilgrim et al., 2003), and LytA of Strep. pneumoniae (Berry & Paton, 2000), are less virulent than the wild-type strains. Colonization of liver and brain by L. monocytogenes was much more IspC-dependent than was that of spleen. The observed effect on virulence, as reflected by the impaired ability to colonize the target organs (liver and brain) of the ΔispC mutant, is unlikely to be attributable to the growth
rate, colony or cell morphology, or biochemical characteristics of the mutant strain, because in vitro the ΔispC mutant is similar in these aspects to the wild-type strain. These findings suggest that IspC does not function in cell division or separation during in vitro growth, for which other autolysins, p60 and MurA (NamA), have been shown to be necessary in L. monocytogenes (Carroll et al., 2003; Machata et al., 2005; Pilgrim et al., 2003). Therefore, IspC, deficiency of which is responsible for the observed attenuated virulence in mice, is involved in the establishment of L. monocytogenes infection in vivo.

This study has further characterized the ΔispC mutant in detail using a cell culture infection model employing various eukaryotic cell types that the bacterium normally encounters during in vivo infection, leading to a better understanding of the principal roles of IspC in pathogenesis. The results obtained from these experiments demonstrate that IspC significantly facilitates the infectious process of L. monocytogenes at multiple steps that are known to require several key virulence factors (Vazquez-Boland et al., 2001), such as internalins (InlA and InlB), responsible for bacterial entry into host cells, phospholipases (PlcA and PlcB) and listeriolysin O (LLO), for escape from the phagosomes, and the actin polymerization (actin tail)-promoting protein ActA, for intracellular movement and cell-to-cell spread. The impaired capability of the ΔispC mutant to adhere to Hep-G2, Vero and SCP cells but not to other eukaryotic cells (Caco-2, L132, Hela, HBMEC and J774) suggests that IspC functions in pathogenesis as an adhesin, mediating the attachment of the bacterium to certain eukaryotic cells, which presumably express unidentifed receptor(s) specific for IspC. Localization of IspC on the cell surface (Wang & Lin, 2007) meets the requirement for this protein to act as an adhesin. The adhesin nature of IspC is supported by the observation that purified IspC was capable of binding to SCP and Vero via its C-terminal CWBD, made of seven GW modules. The adhesive properties of other autolysins, including Aas of Staph. saprophyticus (Hell et al., 1998), AtlC of Staph. caprae (Allignet et al., 2002), AtlE of Staph. epidermidis (Heilmann et al., 1997) and Ami of L. monocytogenes (Milohanic et al., 2001), have been shown. Comparison of the adherence of the ΔispC mutant to various eukaryotic cells with the invasion of these cells by the mutant points to some interesting facts: (i) the mutant, which has an impaired ability to adhere to the cell lines Hep-G2, Vero and SCP, is less invasive to these cells; and (ii) the mutant strain, which has no reduction in its ability to adhere to Hela and HBMEC, is fully capable of invading these cells. One exception to this is that although the ΔispC mutant was capable of adhering to Caco-2 and L132, it showed impaired ability to invade these cells. These results indicate that IspC is necessary for entry of L. monocytogenes into

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**Fig. 7.** Detection of actin tail formation in mouse macrophage J774 cells following infection with the ΔispC mutant and the wild-type (WT). Actin tail formation promoted by ActA from the mutant and wild-type strains was examined at the early (3 h) and later (6 h) stages of J774 infection. The actin tails (green), bacteria (red) and nuclei (blue) were stained with Alexa Fluor 488 Phalloidin and rabbit anti-ActA antibody, followed by interaction with Alexa Fluor 647 goat anti-rabbit antibody and DAPI, respectively. The images were visualized with a fluorescence microscope equipped with a digital video camera. The difference in actin tail formation between the mutant and the wild-type is indicated with arrows.
specific types of eukaryotic cells, although this may not be always dependent on IspC-mediated adhesion. Surprisingly, the autolysin Auto of *L. monocytogenes*, similar in domain organization to two other *L. monocytogenes* autolysins, Ami and IspC, with an affinity for certain eukaryotic cells (Milohanic et al., 2001; Wang & Lin, 2007; this study), is not required for adhesion to, but is required for entry into, eukaryotic cells (Cabanes et al., 2004). It appears that the importance of a particular autolysin in *L. monocytogenes* pathogenesis is dependent on the cell type (epithelial cells, fibroblasts, hepatocytes, endothelial cells and macrophages) that the bacterium encounters during *in vivo* infection.

It was interesting to observe that adhesion to and invasion of SCP epithelial cells but not HBMEC by *L. monocytogenes* was dependent on the product of *ispC*. This is a novel finding with implications for the role that IspC plays in *L. monocytogenes* infection of the brain. There are two brain barriers: the cerebral capillary endothelium, as the barrier between the blood and the brain parenchyma; and the choroid plexus epithelium, as the barrier between blood and the cerebrospinal fluid (CSF) (Tuomanen, 1996). It has been shown that efficient invasion of HBMEC by *L. monocytogenes* depends on InlB (Greiffenberg et al., 1998). We have first used SCP cells to demonstrate that *L. monocytogenes* invades these cells in an IspC-dependent manner. This suggests that when cultured human epithelial cells from the choroid plexus are not available, SCP cells are a good cell culture model for *in vitro* study of adhesion to and entry into *L. monocytogenes* to breach the blood–brain barrier, causing

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**Fig. 8.** Box plot comparison of the sizes of plaques formed from murine L2 cells infected with the ΔispC mutant and the wild-type (WT). At least 20 plaques were randomly selected for each strain for determination of their relative sizes (diameter in arbitrary units; a.u.) by analysing the plaque images with ImageJ software. The lower boundary of the box, the line within the box, and the upper boundary of the box correspond to the lower quartile (the 25th percentile), the median and the upper quartile (the 75th percentile), respectively. Error bars above and below the box mark the 90th and 10th percentiles. Solid circles represent the unusual observations (outliers) greater than the 90th percentile or less than the 10th percentile.

**Fig. 9.** Quantitative analysis of the intracellular growth of the ΔispC mutant in Vero and J774 cells. The intracellular growth assay (see Methods) was performed to assess bacterial intracellular growth. Mutant and wild-type (WT) bacteria were used to infect two representative cell lines (Vero and J774) at a predetermined m.o.i. for 1 h followed by further incubation with 100 μg gentamicin ml⁻¹. The numbers of intracellular viable bacteria were determined 2, 4, 6 and 8 h after addition of gentamicin as in Fig. 3, and are presented as the mean ± SD (n=3). *Statistically significant difference (P<0.05).
meningitis or encephalitis. The entry of the bacterium into the epithelial cells of the choroid plexus is mediated by IspC via interaction with an unknown receptor; when within the epithelial cells, the bacterium undergoes cell-to-cell spread or enters into the CSF to undergo an extracellular phase prior to causing brain infection (encephalitis). Demonstration of the presence of L. monocytogenes in CSF (Brouwer et al., 2006, Schluter et al., 1996) supports the notion of L. monocytogenes undergoing an extracellular phase. Alternatively, L. monocytogenes could invade the microvascular endothelial cells through the interaction of InlB with a specific, as yet unidentified receptor, leading to encephalitis. The important but different functions of both IspC and InlB in brain pathogenesis caused by L. monocytogenes may be explained by the facts that both proteins contain the C-terminal CWBD made up of repeated GW modules, with variation in number and amino acid sequence for each protein (Braun et al., 1997; Wang & Lin, 2007); this domain was shown to be responsible for interaction of IspC with SCP epithelial cells (this study), and presumably accounted for binding of InlB to HBMEC.

Formation of shorter and weaker actin tails during early infection of J774 cells with the ΔispC mutant and of smaller sizes of plaques following infection of L2 fibroblasts with the mutant, and significant reduction in growth of the mutant within Vero and J744 cells suggest that efficient intracellular movement, cell-to-cell spread, and intracellular survival of the bacterium necessarily depend on the expression of the ispC gene. We observed a marked reduction in the amount of the surface protein ActA due to the absence of IspC during in vitro growth. This seems to suggest that IspC regulates the surface display of ActA, presumably through its autolytic activity, and thus promotes intracellular movement and cell-to-cell spread. However, the expression of ActA on the bacterial surface within the infected cells (J774) did not seem to be affected by lack of IspC expression in the ΔispC mutant (our unpublished data). Deletion of the p60 gene from L. monocytogenes has been shown to affect the polarization of ActA on the bacterial surface, leading to loss of actin-based motility (Pilgrim et al., 2003). It may be speculated that the deletion of ispC alters the polarization of ActA on the surface of bacteria during the intracellular phase of growth, resulting in shorter and weaker actin tails in early stages of infection.

Using proteomic and immunological analysis, we have assessed the effect of IspC deficiency on the display of surface proteins. The reduced surface expression in the ΔispC mutant of ActA, a putative LPXTG motif-containing internalin, InIC2, and a flagellin-like protein homologous to the 30 kDa flagellin FlaA with peptidoglycan hydrolase activity in L. monocytogenes EGD-e (Popowska & Markiewicz, 2004) suggests that IspC may also be indirectly involved in pathogenesis, because these surface proteins were demonstrated to be, or were implicated as, virulence factors. In addition to the role of ActA in promoting the actin-based intra- and intercellular movement of L. monocytogenes, this protein has been shown to mediate the attachment to and entry of the bacterium into eukaryotic cells (Alvarez-Dominguez et al., 1997). Although the role of InlC2 in pathogenesis remains undefined, this protein may be required for bacterial survival under osmotic and/or stationary-phase stress (Dramsi et al., 1997; Kazmierczak et al., 2003). The FlaA protein, a structural component of flagella, is capable of facilitating the initial association of the bacterium with and effective invasion of epithelial cells (Dons et al., 2004), and enhances L. monocytogenes infectivity after ingestion (O’Neil & Marquis, 2006). Thus, the attenuated virulence of the ΔispC mutant observed with mouse and cell culture infection models may be partly due to the reduced surface expression or display of other known or putative virulence factors. It is unlikely that the C-terminal region of IspC, which is made up of seven GW modules and functions to anchor the protein to the cell wall (Wang & Lin, 2007), is responsible for maintaining the proper display of other surface proteins. We propose that the autolytic activity conferred by the N-terminal catalytic domain of IspC (Wang & Lin, 2007) breaks the chemical bonds within cell wall peptidoglycan to alter the surface properties (e.g. structure, charge and surface modification), as has been shown for p60 of L. monocytogenes (Pilgrim et al., 2003) and Atl of Staph. aureus (Takahashi et al., 2002), providing the cell wall architecture necessary for the proper polarization or display of virulence factors such as ActA, thereby allowing their function. This is also supported by our unpublished observation that the mutant strain was harder to disrupt by physical (ultrasonication) or enzymic (lysozyme digestion) methods than the wild-type. The inspection of proteins that exhibited a reduced surface expression (ActA, InlC2 and a FlaA homologue) and those that were unaffected (InlB, InlA and p60) appears to indicate that the influence on the display of surface proteins of IspC is independent of a particular surface-targeting mechanism, because various surface-targeting mechanisms (Bierne & Cossart, 2007; Cabanes et al., 2002) such as the LPXTG motif in InlA and InlC2, a C-terminal hydrophobic domain in ActA, a C-terminal domain consisting of repeated GW modules in InlB, and the LysM domain in p60, are employed in these proteins. The indirect role of IspC in virulence by promoting the surface display of other virulence factors is in contrast to the findings that the autolysins Auto and p60 of L. monocytogenes do not alter the expression of other major virulence factors such as InlA, InlB, ActA and LLO (Cabanes et al., 2004; Pilgrim et al., 2003). Thus, the present study is the first to demonstrate a dual role for a minor L. monocytogenes autolysin in virulence.

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REFERENCES


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