INTRODUCTION

Edwardsielllosis is a serious disease problem in many cultured freshwater and seawater fishes worldwide (Plumb 1999, Muroga 2001). The causative agent of the disease, Edwardsiella tarda, has been isolated from a wide variety of animals, including mammals and humans (Sakazaki 2001). In Japan, E. tarda has been known to be an important pathogen, particularly in Japanese eel Anguilla japonica, Japanese flounder Paralichthys olivaceus and red sea bream Pagrus major cultures, since the 1970s. Although persistent studies have been performed on the disease and the agent, there is no efficacious control method for this chronic disease due to poor understanding of the infection mechanism in addition to lack of effective chemotherapy. A number of virulence factors of E. tarda have been reported, including the ability to invade epithelial cells (Janda et al. 1993), to resist phagocyte-mediated killing (Ainsworth & Chen 1990, Iida & Wakabayashi 1993), and to produce toxins such as dermatotoxins (Ullah & Arai 1983) and haemolysins (Hirono et al. 1997, Strauss et al. 1997). Among these virulence factors, the ability to resist phagocyte-mediated killing by failing to trigger the respiratory burst gives it a niche advantage in the host (Srinivasa Rao et al. 2001). In addition, we have recently reported that the anti-apoptotic effect of its infection in murine
macrophages is important for efficient intracellular growth of bacteria in the macrophages (Okuda et al. 2006). However, only scant information is available on the pathogenicity of *E. tarda*.

Many gram-negative pathogenic bacteria use a conserved protein secretion machinery termed ‘Type III secretion system’ (TTSS) to transport virulence factors and cause disease. The TTSS apparatus consists of approximately 20 to 25 proteins. The unique feature of the TTSS is a needle-like structure through which particular proteins (termed ‘effectors’) are injected into host cells. Effectors play an important role in the pathogenic relationship between host and bacterium. Injection of effectors into host cells occurs through pores (termed ‘translocons’) formed in the host cell membrane by the TTSS using Type III-secreted proteins (Ghosh 2004). The TTSS encoded within *Salmonella* pathogenicity island 2 (SPI2) is essential for its intracellular accumulation in macrophages. SseB, SseC, and SseD proteins secreted from the TTSS encoded within SPI2 are reported to function as translocons (Nikolaus et al. 2001). Using TnphoA transposon tagging and the proteomics approach, a TTSS was found in *Edwardsiella tarda* that was similar to that encoded within SPI2 of *Salmonella* (Srinivasa Rao et al. 2003, Tan et al. 2005). Three TTSS proteins of *E. tarda*, homologous to SseBCD were identified and these proteins were designated as EseB, EseC, and EseD (Srinivasa Rao et al. 2004, Tan et al. 2005). We reported that, similar to *Salmonella*, the TTSS encoded within *E. tarda* is required for its intracellular accumulation in murine macrophages (Okuda et al. 2006). This suggests that the function of the TTSS of *E. tarda* is to deliver effectors into macrophages using EseBCD proteins as translocons in order to survive in the host cells.

Here we investigate the role of the EseBCD proteins in intracellular replication of the bacterium in murine macrophages by investigating the requirements for the secretion of these proteins.

**MATERIALS AND METHODS**

**Bacterial strains, bacterial growth, and media.** The wild-type strain, *Edwardsiella tarda* FK1051, was isolated from a diseased Japanese flounder, *Paralichthys olivaceus*. The Type III mutant mET1229 is a TTSS-deficient mutant of *E. tarda* FK1051, i.e. the *esaV* gene in FK1051 was knocked out, as described previously (Okuda et al. 2006).

Insertion mutation was carried out in the *eseB*, *eseC*, and *eseD* genes in the wild-type strain as described previously (Tan et al. 2005) with a slight modification using the suicide vector plasmid, pRE112 (Okuda et al. 2001). Internal fragments of each gene were amplified by PCR using the primers listed in Table 1, i.e. a 236 bp fragment from position 89 to 324 in the *eseB* gene (accession number: AY643478), a 501 bp fragment from position 224 to 724 in the *eseC* gene (AY643478), and a 237 bp fragment from position 103 to 339 in the *eseD* gene (AY643478) were amplified using *ΔeseB*-1 and *ΔeseB*-2 primers, *ΔeseC*-1 and *ΔeseC*-2 primers, and *ΔeseD*-1 and *ΔeseD*-2 primers, respectively (Table 1). The primers were designed where the amplification has a SacI site and stop codons at the 5’ end and a SacI site at the 3’ end. The PCR amplification was performed using Taq polymerase (Ex Taq, Takara) at 30 cycles: 30 s denaturation at 95°C, 1 min annealing at 60°C, an extension step at 72°C for 1.5 min, and then a final extension step at 72°C for 5 min. The amplified fragments were digested using SacI and cloned into SacI-cleaved pRE112. The plasmids (designated as pΔeseB, pΔeseC, and pΔeseD) were first constructed using an *Escherichia coli* SY327λpir background and *E. coli* SM10λpir was transformed with them. These plasmids were mobilized into *Edwardsiella tarda* FK1051 using conjugation. Transconjugants were selected on Trypto-Soya agar (TSA, Nissui) supplemented with chloramphenicol (30 µg ml⁻¹) and colistin (10 µg ml⁻¹). Insertional inactivation then occurred, resulting in the integration of the cloned suicide vector

<table>
<thead>
<tr>
<th>Designation</th>
<th>Sequence</th>
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<tbody>
<tr>
<td>ΔeseB-1</td>
<td>5’-GAG AGA GCT CTG ACT GAC TGA GCA TCA TGA CCG AC-3’</td>
</tr>
<tr>
<td>ΔeseB-2</td>
<td>3’-AGA GGA GCT CCG TGA TGT TGT GAT-3’</td>
</tr>
<tr>
<td>ΔeseB-3</td>
<td>5’-GAG AGG ATC CAT GAC TGT CAA TAC AGA CTA C-3’</td>
</tr>
<tr>
<td>ΔeseB-4</td>
<td>5’-AGA GGC CCG CCG GCG GAT ATT CTG GGC GAT GGA T-3’</td>
</tr>
<tr>
<td>ΔeseC-1</td>
<td>5’-GAG AGA GCT CTG ACT GAC TGA AGC AGG ATC ACG ATC CTA AG-3’</td>
</tr>
<tr>
<td>ΔeseC-2</td>
<td>5’-AGA GGA GCT CGC ATT GTC CTT ATG CAG-3’</td>
</tr>
<tr>
<td>ΔeseC-3</td>
<td>5’-AGA GAC ATT CAT GGT GCG CCG CCA AGC GGC A-3’</td>
</tr>
<tr>
<td>ΔeseC-4</td>
<td>5’-AGA GGC CCG CCG GGC GAT ATT GGC GGC GAT GTT-3’</td>
</tr>
<tr>
<td>ΔeseD-1</td>
<td>5’-AGA GGC CCG CCG GGC GAT ATT GGC GGC GAT GAT-3’</td>
</tr>
<tr>
<td>ΔeseD-2</td>
<td>5’-AGA GGC CCG CCG TTA ACG CCG TGT CTA-3’</td>
</tr>
<tr>
<td>ΔeseD-3</td>
<td>5’-AGA GGC CCG CCG GGC GAT ATT GGC GGC GAT GGC GAT-3’</td>
</tr>
<tr>
<td>ΔeseD-4</td>
<td>5’-AGA GGC CCG CCG GGC GAT ATT GGC GGC GAT GGC GAT A-3’</td>
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into the *eseB*, *eseC*, and *eseD* genes in the wild-type genome as described previously (Tan et al. 2005). Single strains retaining the mutant *eseB*, *eseC*, or *eseD* gene were selected as the EseB, EseC, and EseD mutants, respectively.

FK1051 was grown in Tripto-Soy broth (TSB, Eiken) at 30°C. The Tyee III mutant mET1229 was grown in TSB supplemented with kanamycin (50 µg ml⁻¹), EseB, EseC, and EseD mutants were grown in TSB supplemented with chloramphenicol (30 µg ml⁻¹). Formalin-killed FK1051 was prepared as described previously (Clemens et al. 2004).

The influence of the incubation temperature on EseB, EseC, and EseD secretion was determined as follows: the wild-type strain and mET1229 were grown in TSB medium at 20, 25, 30, and 37°C to an absorbance of light with a wavelength of 600 nm (A600) = 0.9.

The influence of the pH in the culture media on Ese-BCD secretion was determined as described previously (Beuzon et al. 1999). The wild-type strain and mET1229 were grown in pH 7.4 TSB medium at 25°C until an A600 = 0.6, and then the cells were inoculated into fresh TSB media at pH 5.5, 6.5, 7.4, or 8.5 and incubated at 25°C for 24 h.

**J774 cells.** J774 cells, a murine macrophage-like cell line derived from BALB/c mice, were obtained from the American Type Culture Collection. J774 cells were maintained in RPMI1640 (Sigma) supplemented with 10% FCS.

**Infection of J774 cells with *Edwardsiella tarda* strains.** A 24-well tissue culture plate was inoculated with 7.5 × 10⁴ J774 cells per well and incubated overnight at 37°C and under 5% CO₂. The cells were infected with either the wild-type *E. tarda* strain or the mET1229, EseB, EseC, and EseD mutants at a multiplicity of infection (moi) of 1 for 30 min. Pre-warmed tissue culture medium containing 200 µg ml⁻¹ gentamicin was then added. After 1.5 h incubation, the medium was removed, the cells were washed twice with PBS and new tissue culture medium was added. The J774 cells were incubated for 0, 5, 12, or 22 h at 37°C and under 5% CO₂. At the end of each time period, the cells were washed twice with PBS, lysed with 1% Triton X-100 and cultured on plates to estimate the colony forming units (CFU) in the macrophages.

**Preparation of secreted proteins.** Secreted proteins in the culture supernatant were isolated using TCA precipitation. TCA was added to the culture supernatant to 10% and incubated on ice for 15 min. After centrifugation at 15,000 × g for 15 min, the precipitated proteins were washed 3 times with acetone and dried. The dried proteins were dissolved in PBS and the concentration was determined using the Bradford assay (BioRad).

**Protein analysis.** Secreted protein profiles were analyzed using sodium dodecylsulfate-polyacrylamide gel electrophoresis (SDS-PAGE) according to the method of Laemmli (1970). A 5 µg sample was applied to 12.5% gel. Protein bands were visualized using Coomassie Brilliant Blue R-250 (CBB; Wako) or Silver Stain Kit II (Wako).

**Amino acid sequencing.** The proteins separated by SDS-PAGE were transferred to a polyvinylidene fluoride (PVDF) membrane (Millipore). After staining the PVDF membrane with CBB, protein bands in the membrane were excised. The amino acid sequence was determined using a G1005A Protein Sequencing System (Hewlett-Packard).

**Labeling of acidified phagosomes with LysoTracker red DND-99.** LysoTracker red DND-99 (Molecular Probes), an acidotropic dye, consists of a weak base conjugated to a red fluorophore and is used as a marker of phagosomal acidification and maturation (Toyooka et al. 2005). J774 cells were pretreated with LysoTracker (25 nM) in RPMI1640 for 2 h before infection. The cells were then infected with the wild-type strain or mET1229 at an moi of 20 for 30 min and pre-warmed tissue culture medium containing 200 µg ml⁻¹ gentamicin and 25 nM LysoTracker was added. After incubation for 1.5 h, the medium was removed, washed twice with PBS, and tissue culture medium containing LysoTracker (25 nM) was added. The J774 cells were incubated for 1 h at 37°C and <5% CO₂. At the end of time period, the cells were washed twice with PBS, fixed in 4% paraformaldehyde for 20 min, washed again with PBS, and treated with NH₄Cl. Then the cells were made permeable with 0.2% Triton X-100. Cover slips covered with the cells were incubated in anti-*Edwardsiella tarda* rabbit serum (1:640) for 1 h at 37°C and <5% CO₂, washed with Tris-buffered saline Tween (TBS-T), and incubated with Alexa Fluor 488 chicken anti-rabbit IgG antibody (Molecular Probes) (1:100). The cover slips were mounted on glass slides and phagosomes containing bacteria were examined using a confocal laser scanning microscope system (LSM 510, Carl Zeiss). The percentage of acidic phagosomes were calculated as the ratio of the green bacteria to the total number of bacteria (total ca. 50 to 60 bacteria). The statistical analysis was performed using the 2-tailed t-test.

**RESULTS AND DISCUSSION**

**Identification of proteins secreted via the TTSS of *Edwardsiella tarda***

The supernatant proteins of the wild-type strain and the Type III mutant (mET1229) are shown in Fig. 1A. The major 3 bands at the positions 55, 25,
and 22 kDa were detected in the wild-type, but not in the mET1229 samples. The N-terminal amino acid sequence (10 residues) of the 25 kDa protein was determined (Fig. 1B); however, the 55 and 22 kDa sequences were not determined because they have an N-terminal block. A BLAST search shows that the 25 kDa protein is similar to the predicted N-terminal sequence of EseB, and has a sequence similarity to Salmonella SseB, that functions as a translocon (Srinivasa Rao et al. 2004). In Salmonella, a set of substrate proteins consisting of SseB, SseC, and SseD are secreted by the TTSS and these proteins are predominantly assembled into complexes on the bacterial cell surface to function as the translocon for the translocation of effector proteins such as SspH1 and SspH2 (Nikolaus et al. 2001). Since the EseB, EseC and EseD proteins of Edwardsiella tarda are previously reported to be the homologues of Salmonella SseB, SseC and SseD (Srinivasa Rao et al. 2004), we considered the 55 and 22 kDa proteins to be EseC and EseD, respectively. To confirm that the 25, 55 and 22 kDa proteins were EseB, EseC, and EseD, respectively, we constructed knockout mutants in the genes coding for EseB, EseC, and EseD in E. tarda (Fig. 2). As expected, the knockouts of EseB, EseC, and EseD resulted in the disappearance of the 25, 55, and 22 kDa proteins, respectively. However, the knockout of the gene coding for EseB caused the disappearance of the 55 kDa protein (EseC) as well as the 25 kDa protein (EseB) and the knockout of the gene coding for EseD resulted in the disappearance of the 55 kDa protein (EseC) as well as the 22 kDa protein (EseD). This suggests that the knockout of the eseB, eseC, or eseD gene influences the secretion of the mutual genes. In Salmonella, similar observations are reported where the sseC mutant causes down-regulation of the expression of SseD and the sseD mutant results in the down-regulation of SseC (Chakravortty et al. 2005). However, we cannot rule out the possibility that disappearance of the secretion of the 55 kDa protein (EseC) may also be due to polar effects on the 25 kDa protein (EseB) since the eseC gene is located downstream of the eseB gene (Tan et al. 2005), and in another report, the deletion mutant of AeseB in a different strain did not affect the secretion of EseC and EseD (Zheng et al. 2007). On the other hand, the eseC gene is located upstream of the eseD gene (Tan et al. 2005); therefore, the observation of reduced secretion of the 55 kDa protein (EseC) may not be due to polar effects on the 22 kDa protein (EseD), but may be similar to the observation reported previously, where, in Salmonella, the sseD mutant results in the down-regulation of SseC (Chakravortty et al. 2005).
Requirement of EseB, EseC, and EseD proteins for intracellular replication of *Edwardsiella tarda*

The average number of viable wild-type cells replicating in J774 cells increased 5 to 22 h after infection, while the EseB, EseC, and EseD mutants decreased in a time-dependent manner. Most of the mutants entering J774 cells were no longer viable 22 h after infection (Fig. 3). This suggests that the EseB, EseC, and EseD proteins are essential for intracellular survival of *Edwardsiella tarda* in murine macrophages. This was similar to observations in *Salmonella*, i.e. the growth of strains carrying mutations in *sseB* or *sseC* within murine macrophages was severely decreased (Hensel et al. 1998). Tan et al. (2005) reported that intracellular growth of the *eseB*, *eseC*, and *eseD* insertional mutants in fish phagocytes decreased moderately, compared to
the wild-type strain. However, we found that 5 h after infection, intracellular growth of the eseB, eseC, and eseD mutants in murine macrophages decreased considerably, compared to the wild-type strain. This difference between the observations of Tan et al. (2005) and our data may be due to differences between fish phagocytes and murine macrophages.

**Effect of culture conditions on the secretion of the EseBCD proteins**

We investigated whether secretion of the EseBCD proteins was influenced by culturing bacterial cells in media at different pHs. We show in Fig. 4 that EseBCD proteins were secreted when the wild-type strain was cultured in media at neutral (pH 7.4) or alkaline (pH 8.5) pH, while the secretion of these proteins was faint when the wild-type strain was cultured in media at an acidic pH (pH 5.5). This was different for the Sse-BCD proteins found in Salmonella (Nikolaus et al. 2001), where the SseBCD proteins were most secreted when bacterial cells were cultured in media at acidic pH (pH 5.0). This suggests that after ingestion by macrophages Edwardsiella tarda prevents acidification of the phagosomes in order to allow secretion of the EseBCD proteins through the TTSS.

Srinivasa Rao et al. (2004) previously reported that the EseB and EseD protein production was markedly reduced at 37°C incubation but was observed by 2-dimensional gel electrophoresis (2-DE) followed by silver staining. Similarly in our study, the EseBCD proteins were secreted at highest levels when the wild-type strain was incubated at 25 to 30°C, while secretion of these proteins was markedly reduced at 37°C (Fig. 5A), as detected by SDS-PAGE followed by silver staining (Fig. 5B), which coincides with the observation by Srinivasa Rao et al. (2004). This temperature dependency of translocon protein secretion will be of importance in clarifying the infection mechanism of Edwardsiella tarda in cultured fishes, which are usually reared in water with temperatures < 30°C. Further investigations are required to reveal the relationship between fish virulence, or resistance of fish macrophages, and secretion of the translocon proteins.
pH of phagosomes after *Edwardsiella tarda* infection

*J*774* cells infected with the formalin-killed wild-type strain, wild-type strain or mET1229 mutant were stained using the LysoTracker (Fig. 6). The majority of phagosomes in the *J*774 cells phagocytosing the formalin-killed wild-type strain or mET1229 colocalized with red acidic particles 1 h after infection. In contrast, phagosomes with the wild-type strain did not colocalize with acidic particles (Fig. 6). The differences between the formalin-killed wild-type and wild-type strains and between the wild-type strain and mET1229 were significant (p < 0.002 and p < 0.001, respectively).

This suggests that unidentified effector proteins through the TTSS may be secreted from the wild-type strain to prevent fusion of acidic particles with the phagosomes, resulting in the neutralization of the phagosomes. We speculate that neutralization of phagosomes is a prerequisite for the efficient secretion of the EseBCD translocon proteins that are essential for the successive secretion of unidentified effector proteins. Further investigation in identifying these unidentified effector proteins may provide important insights into the mechanism initiating intracellular replication of *Edwardsiella tarda* in macrophages.

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**LITERATURE CITED**


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