Adherence and Infectivity of Green Fluorescent Protein-Labeled *Pseudomonas plecoglossicida* to Ayu *Plecoglossus altivelis*

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**ABSTRACT**—We report green fluorescent protein (GFP) as a useful reporter molecule to track the occurrence of *Pseudomonas plecoglossicida* in ayu *Plecoglossus altivelis*. The gfp gene was placed in a shuttle vector pME4510 and then three kinds of GFP vectors, pSKL01, pSKT03 and pSKN04 were constructed, in which lac, tac, and *neophosphotransferase II* (*npt2*) promoters drove the expression of GFP, respectively. A promoterless GFP vector was designated as pSKP02. *P. plecoglossicida* carrying pSKT03 gave the highest expression of GFP. In addition, pSKT03 was relatively stable under non-selective pressure and there was no significant changes in the *in vitro* growth and pathogenicity of the cells carrying this plasmid. The cells of fluorescent *P. plecoglossicida* attaching to the body surface of ayu were easily detected under a fluorescence microscope. It was revealed that they adhered predominantly to the microscopic injuries in the skin and fins.

**Key words:** *Pseudomonas plecoglossicida*, green fluorescent protein, *Plecoglossus altivelis*, adherence, ayu

*Pseudomonas plecoglossicida* is a pathogen causing bacterial hemorrhagic ascites (BHA) in ayu (*Plecoglossus altivelis*) in Japan (Wakabayashi et al., 1996; Nishimori *et al.*, 2000). Sukenda and Wakabayashi (1999, 2000) described the kinetics of this bacterium in the experimentally infected fish based on the microbiological culture and real-time quantitative polymerase chain reaction. To study the infection process of this disease, it was apparent that many aspects could be investigated more easily if a visible marker could be introduced into the bacterial cells.

Recently, the gfp gene encoding green fluorescent protein (GFP) from a jellyfish (*Aequorea victoria*) was shown to be expressed in both eukaryotic and prokaryotic cells (Chalfie *et al.*, 1994). GFP is a monomeric 238 amino-acid protein that emits green light when excited with blue light. The usefulness of GFP in tagging a pathogenic bacterium of fish has been illustrated by Ling *et al.* (2000). *Edwardsiella tarda* was tagged with *Escherichia coli* vector (pGFPuv) and the infection kinetic of the bacterium was studied *in vitro* and *in vivo*. However, the current lack of constructs for diverse groups of bacteria make these systems inapplicable for other bacteria as the pGFPuv could not be expressed in *P. plecoglossicida*.

Previous experiments show that immersion challenge is considered as a suitable challenge model in reproducing BHA which resembles a natural infection (Sukenda and Wakabayashi, 1999; 2000). Moreover, the skin is likely the main portal of entry for *P. plecoglossicida*, since the bacterium was found in early stages post-infection (Sukenda and Wakabayashi, 2000). Many authors have noted that injuries are among the main sites of bacterial adherence on the skin and fins of fish (Schneider and Nicholson, 1980; Crouse-Eisnor *et al.*, 1985; Ventura and Grizzle, 1987; Speare and Mirstamali, 1992). However, the precise mechanism

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of their adherence to the body surface of fish is as yet unknown. We used GFP as a molecular marker for P. plecoglossicida to study the adherence sites of P. plecoglossicida in the body surface of ayu. Its adherence was also compared with non-pathogenic Escherichia coli as well as microspheres.

Materials and Methods

Bacterial strain, plasmids, media and culture conditions

The bacterial strains and plasmids used in this study are listed in Table 1. E. coli JM109 and P. plecoglossicida FPC 951T (= ATCC 700383T) carrying GFP constructs were grown in LB medium containing gentamicin (15 µg/mL) at 37°C and 25°C, respectively.

DNA cloning and polymerase chain reaction (PCR)

Plasmid extraction, restriction enzyme, PCR products purification, agarose gel electrophoresis, gel-isolated DNA fragment purification, dephosphorylation and ligation were carried out using standard methods (Sambrook et al., 1989), and following the appropriate manufacturer's instructions. Restriction and other enzymes were purchased from Takara Co., Ltd., Japan. Oligonucleotide primers were synthesized by Life Technologies Inc., USA. PCR amplification was achieved using Gene AmpPCR System 2400, Perkin-Elmer. The amplification was performed in 50 µL containing 1.5 mM MgCl₂, 0.5 mM dNTPs (adenine, guanine, cytosine, and thymine), 20 pM of each primer, 2.5 units of Taq DNA polymerase and 100 ng of template DNA. A total of 30 cycles were run as follows: denaturation at 94°C for 30 s, annealing at appropriate temperature for 1 min and extension at 72°C for 1.5 min, followed by 1 cycle at 72°C for 5 min.

Construction of shuttle vectors expressing GFP

Four GFP vectors were constructed in this study. A broad-host range plasmid of pME4510 served as a shuttle cloning vector and gfp cDNA was isolated from pGFPuv. The first plasmid construct was pSKL01 in which lac promoter was used to drive expression of GFP. The promoter and gfp gene were isolated by PCR at an annealing temperature of 58°C from pGFPuv, using two primers with the following sequences: 5'-GGCCGAAATTCTTAAATGCGAGCTG-3' and 5'-AGTTGGGAATTCTTATTGT-3'. Both of primers contained an EcoRI restriction site (underlined). The 988 bp of PCR product was then ligated into pME4510 linearized with EcoRI. The second plasmid construct was a promoterless, pSKP02. It was constructed by restriction of pME4510 with EcoRI and HindIII and ligated into 750 bp fragment encoding the promoterless GFP isolated from pGFPuv by cutting with EcoRI and HindIII. From this plasmid, we constructed two other vectors, namely pSKT03 and pSKN04 in which tac and npt2 promoter drove the expression of GFP, respectively. The pSKP02 was linearized with HindIII and XbaI. PCR product carrying the tac promoter fragment was amplified from pMBB66EH at an annealing temperature of 58°C using primers 5'-ATAAAGCTTCTGGCAAATATTCTGAAA-3' and 5' CCTCTCTAGAATTGTTATCCGCTCACA-3'. The npt2 promoter was amplified from Tn5-Mob at an annealing temperature of 60°C, using primers 5'-CTCTCTAGAAGGTAGCTTGGAGG-3' and 5'-CTCTCTAGACAGCCATTGACATCCTTGGCCG-3'. The first and second promoters of both pairs contained a HindIII site and a XbaI site (underlined). The PCR products were cut with HindIII and XbaI and ligated to the vector.

Cells transformations

All plasmid vectors were transformed into E. coli JM109 using a standard heat-shock transformation (Sambrook et al., 1989) and were electroporated into P. plecoglossicida using method as described by Bloemberg et al. (1997).

GFP fluorescence

The expression of GFP-expressing cells was

<table>
<thead>
<tr>
<th>Strain</th>
<th>Description</th>
<th>Reference/source</th>
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<tbody>
<tr>
<td>E. coli JM109</td>
<td>competent cell</td>
<td>Takara</td>
</tr>
<tr>
<td>P. plecoglossicida FPC 951T</td>
<td>wild type</td>
<td>Nishimori et al. (2000)</td>
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<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Description</th>
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<tr>
<td>pGFPuv</td>
<td>AmpR, Plac, gfp</td>
<td>Clontech</td>
</tr>
<tr>
<td>pMME66EH</td>
<td>Plac, AmpR</td>
<td>ATCC 37620</td>
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<tr>
<td>pSUP5011</td>
<td>Tn5-Mob</td>
<td>Simon (1984)</td>
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<td>pSKN04</td>
<td>GmR, Pnpt2, gfp</td>
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assessed by eye using a UV transillumination (Vilber Lourmat, France) and fluorescence microscope (Olympus SZX-12, Japan) fitted with GFP filter set to examine colonies, and an Olympus BX-60 microscope to examine cells. The amount of fluorescence emitted by cultures was also assayed using a F-2000 fluorescence spectrophotometer (Hitachi, Japan) set to excite the cells at 395 nm and detect emission at 509 nm. Culture was grown in LB medium with the antibiotic overnight and diluted to $A_{600}$ of 1.4 in the fresh LB medium. The cells were washed twice and diluted in the same volume of PBS, and the fluorescence was measured immediately.

**Growth of cells and plasmid stability**

*P. plecoglossicida* carrying the appropriate GFP-plasmid constructs were grown overnight in the presence of antibiotic LB broth. The cells were washed twice and diluted 1:1000 in LB broth without antibiotics, and growth of the cells was determined hourly at $A_{600}$. For plasmid stability assay, sequential propagation under non-selective conditions were performed by inoculating with 1:1000 (v/v) of the previous culture daily. The bacterial culture was sampled daily to assess plasmid survival by comparing duplicate colony counts on selective and non-selective plates.

**Infectivity of fluorescent *P. plecoglossicida* to ayu**

Two groups of fish were immersed for 15 min in $10^7$ CFU/mL of fluorescent *P. plecoglossicida* (pSKT03) and *P. plecoglossicida* (wild type), respectively. A control group was immersed in PBS. Kidney samples from dead fish were inoculated on LB plate with or without antibiotic. Pieces of tissue were immediately mounted in Tissue-Tek OTC Compound (Sakura, Japan), frozen in liquid nitrogen and then cut into 7 μM sections on a cryostat (Sakura, Japan).

**Adherence of fluorescent *P. plecoglossicida* to ayu**

Fluorescent *P. plecoglossicida* FPC 951 (pSKT03), fluorescent *E. coli* JM109 (pSKL01), and one-micron fluoresbrite® bright blue (BB) carboxylated microspheres (Polyscience, Inc., USA) were used. The bacterial cells and microspheres were suspended in PBS at $10^7$ CFU/mL and $10^6$ particles/mL, respectively. A mixture of bacteria and microspheres was made by adding an equal volume of the microspheres suspension to the bacterial suspension.

Seven different immersion experiments were performed as follows. In the experiments nos. 1, 2 and 3, fish were immersed in the fluorescent microsphere, fluorescent *E. coli*, and fluorescent *P. plecoglossicida* suspensions, respectively for 15 min and then dipped in 1 L of a 0.05% trypan blue (TB) solution for 10 min. In the experiments nos. 4, 5 and 6, fish were immersed in the mixture of fluorescent microspheres and fluorescent *E. coli*, the mixture of fluorescent microspheres and fluorescent *P. plecoglossicida*, and the mixture of fluorescent microspheres and *P. plecoglossicida* (wild type), respectively. In the experiment no. 7, fish were immersed in fluorescent *P. plecoglossicida* suspension only. The fish were killed by a lethal dose of anesthetic (2-phenoxethanol) and fixed in a 4% paraformaldehyde for 1 d. The skin of the body surface was peeled off, and the fins were cut off from individuals. These tissues were mounted on glass slides with slow fade light antifade kit (Molecular probes, USA). All specimens were observed in detail under a fluorescence microscope.

**Results**

**Construction of plasmids**

We successfully developed GFP vectors for use in *P. plecoglossicida* (Fig. 1). The GFP vector, pSKL01 (Fig. 1A), was constructed in which lac promoter drove the expression of GFP. When pSKL01 was introduced into *P. plecoglossicida*, this resulted in weak green fluorescence of GFP which could still be observed under a fluorescence microscope (Fig. 2A). Thus, to intensify fluorescence, other constructs were prepared in which other promoters were used to express GFP. First, we constructed the promoterless GFP vector, designated pSKP02 (Fig. 1D), by cloning promoterless GFP fragments from pGFPuv to the pME4510. When pSKP02 was introduced into *P. plecoglossicida*, no fluorescence could be observed (Fig. 2D). However, when tac promoter was cloned in the upstream of gfp gene, yielding pSKT03, and introduced into *P. plecoglossicida*, the resulting fluorescence of the transformants was stronger (Figs. 1B and 2B). Identical approach was used in which pSKNO4 with npt2 promoter was used to drive GFP expression (Figs. 1C and 2C).

**GFP expression**

Expression of GFP in *P. plecoglossicida* was demonstrated by fluorescence microscopy of individual cells with all the constructs, as shown in Fig. 2. When fluorescence was measured by fluorometry, the emission was as follows: 4, 14, 5, and 1 fluorescent units, for *P. plecoglossicida* carrying pSKL01, pSKT03, pSKN04, and wild type, respectively. The emission of 980, 53, 826, and 2 fluorescent units were for *E. coli* carrying pSKL01, pSKT03, pSKN04, and wild type, respectively. All GFP constructs lead much more fluorescence in *E. coli* than *P. plecoglossicida*.

**Plasmid stability**

Colonies grown on the present antibiotic media gave uniform fluorescence, but those grown on media not containing antibiotic showed fluorescent and non-fluorescent colonies that indicated plasmid loss. The stability of the GFP-plasmid constructs were investigated...
during sequential propagation in the absence of antibiotic selection, and show that after subculturing and growth for 4 successive days, the cells, ranging from 25% (pSKL01) up to 70% (pSKT03) and 72% (pSKN04), still carried their respective plasmid (Fig. 3). However, the relative highly instability of plasmid pSKL01 under nonselective condition suggests that this plasmid may not be suitable for long-term experiments without antibiotic pressure.

**Plasmid burden**

Such differences in the growth rate occurred because of the metabolic load on the cells caused by the need to replicate the plasmid, and to express its gene, the growth of the cells in liquid LB medium, with and without GFP plasmid, was assayed. The results show that under laboratory conditions, the growth rate of *P. plecoglossicida* with or without the GFP expressing plasmid were identical (Fig. 4), indicating that there is no significant burden to the cells carrying these plasmids.

**Infectivity and adherence assays**

No mortalities or clinical signs of disease were observed in negative control fish. At day 10 post infection, cumulative mortalities in 10 experimentally infected fish reached 90% (pSKT03) and 80% (wild type). All diseased fish showed clinical signs and gross pathological features consistent with those described in fish with BHA (Wakabayashi et al., 1996). Fluorescent *P. plecoglossicida* (pSKT03) was also observed in the kid-
Adherence of GFP-labeled *P. plecoglossicida*

2. Fluorescence micrographs of *P. plecoglossicida* carrying GFP constructs. (A) pSKL01 (lac promoter), (B) pSKT03 (taq promoter), (C) pSKN04 (npI2 promoter), and (D) pSKP02 (promoterless).

5. Cryostate section of kidney of ayu infected with fluorescent *P. plecoglossicida* showing numerous fluorescent bacteria.

6. Fluorescent microspheres adhering to the body surface of ayu: (A) light microscopy, (B) fluorescence microscopy. Necrotic tissues stained with trypan blue (arrow head) and fluorescent microspheres (arrow) are shown.

7. Fluorescent microspheres and fluorescent *P. plecoglossicida* adhering to the body surface of ayu: (A) fluorescent microspheres, (B) fluorescent *P. plecoglossicida*. 8. Fluorescent microspheres and fluorescent *E. coli* adhering to the body surface of ayu: (A) fluorescent microspheres, (B) fluorescent *E. coli*. Scale bars for 2A–2D and 5 are 10 μm, and 6A–8B are 30 μm.
Fig. 3. Stability of plasmids in *P. plecoglossicida* under non-selective conditions.

**Discussion**

Green fluorescent protein cDNA has been expressed in a variety of both gram-positive and gram-negative bacteria (Lewis and Errington, 1996; Valdivia *et al.*, 1996; Bloemberg *et al.*, 1997). In the present study, we described the construction of new GFP-containing plasmids for use in *P. plecoglossicida*. Plasmids pSKL01, pSKT03 and pSKN04 carry a bright mutant of GFP (Crameri *et al.*, 1996), which allowed easy detection with a standard fluorescence microscope. In the presence of the lac promoter, high expression of GFP were seen in *E. coli*, whereas a much weaker of induction was seen in *P. plecoglossicida*. This is consistent with previous observations that the lac promoter is expressed weakly in pseudomonads (Matthysse *et al.*, 1996; Rist and Kertesz, 1998). When another construct was made in which the nptII promoter was used to express GFP in pSKN04, intenser fluorescence was seen in *P. plecoglossicida*. The expression of GFP in *P. plecoglossicida* was the highest when taq promoter was used to drive GFP expression. This suggested that *P. plecoglossicida* carrying pSKT03 could be used for *in vivo* experiments using fish models. In addition, pSKT03 was relatively stable under non-selective pressure and there was no significant burden to the cell carrying this plasmid.

The advantages of using GFP as a marker include ease of detection, no requirement for exogenous substrate or energy source, and no processing of the cells (Chalfie *et al.*, 1994; Kremer *et al.*, 1995). Histological study confirmed the presence of fluorescent *P. plecoglossicida* (pSKT03) in fish infected with this strain. The conventional method using indirect antibody labeling needs longer processing times for bacterial detection. Using the GFP markers, bacteria in cryostate sections were directly visualized without any immunological treatment. Moreover, by tracking GFP-expressing bacteria in animal models, the distribution of bacteria in tissue and transmission pathway could be clarified (Ling *et al.*, 2000).

In addition to the new research possibilities by using GFP as a reporter for *P. plecoglossicida*, we demonstrated the useful technical application of the plasmids to visualize the bacteria in the body surface of infected fish. As shown for the adherence assay in this study, the adherence of fluorescent *P. plecoglossicida* as well as *E. coli* on the body surface of ayu could be easily detected using fluorescence microscopy. The present study revealed that bacteria adhered predominantly in the area where microspheres were found. Kiryu and Wakabayashi (1999) demonstrated that microspheres adhered predominantly to the surface of microscopic injuries that were stained with trypan blue. However, since the fluorescence of bacteria disappeared when stained with trypan blue, we used fluorescent micro-
spheres indirectly to specify microscopic injuries. Since the fluorescent *P. plecoglossicida* as well as *E. coli* were almost always located in the area where the fluorescent microspheres were also present, we concluded that the injuries were the main site of bacterial adherence to the skin and fins of fish. However, pathogenic bacteria need to survive at adhesion sites and to escape from the host defense system. These processes seem to decide the fate of *P. plecoglossicida* and *E. coli* in the microscopic injuries. Further studies on the mechanism of invasion of the adhered bacteria into the host tissues are needed.

Several new variants of the GFP have recently become commercially available. pBFP2 (Clontech) is a mutant expressing blue fluorescence. Another recently available one is a red-shift excitations variant (pEGFP). The availability of these variants will significantly extend the usefulness of GFPs in bacteria and will enable us to compare the ability of different strain and species to infect fish in co-infection experiments. For example, the expression of the *gfp* gene in *P. plecoglossicida* and blue-fluorescent variant in other fish pathogenic bacteria such as *Flavobacterium psychrophilum*, which also causes another serious disease of ayu, would allow a comparison of the infection kinetics of the two different species in co-infection experiments using ayu as a model animal. Moreover, with the help of these, it should be possible to visualize gene regulation events, in bacteria, in cell culture and in the host, to identify specific adaptation mechanisms and improve understanding of host-pathogen interactions that make persistent infection possible (Josenhans et al., 1998).

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