Immunoresponse of Coho salmon immunized with a gene expression library from *Piscirickettsia salmonis*

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**ABSTRACT**

We have used the expression library immunization technology to study the protection of Coho salmon *Oncorhynchus kisutch* to the infection with *Piscirickettsia salmonis*. Purified DNA from this bacterium was sonicated and the fragments were cloned in the expression vector pCMV-Bios. Two libraries were obtained containing 22,000 and 28,000 colonies and corresponding to approximately 8 and 10 times the genome of the pathogen, respectively. On average, the size of the inserts ranged between 300 and 1,000 bp.

The plasmid DNA isolated from one of these libraries was purified and 20 µg were injected intramuscularly into 60 fish followed by a second dose of 10 µg applied 40 days later. As control, fish were injected with the same amount of DNA of the vector pCMV-Bios without insert. The titer of IgM anti-*P. salmonis* of vaccinated fish, evaluated 60 days post-injection, was significantly higher than that of the control group injected with the vector alone. Moreover, this response was specific against *P. salmonis* antigens, since no cross reaction was detected with *Renibacterium salmoninarum* and *Yersinia ruckeri*.

The vaccinated and control fish were challenged 60 days after the second dose of DNA with $2.5 \times 10^7$ *P. salmonis* corresponding to 7.5 times the LD$_{50}$. At 30 days post-challenge, 100% mortality was obtained with the control fish while 20% of the vaccinated animals survived. All surviving fish exhibited a lower bacterial load in the kidney than control fish.

The expression library was also tested in Balb/c mice and it was found that the humoral immune response was specific to *P. salmonis* and it was dependent on the amount of DNA injected.

**Key terms:** Piscirickettsia salmonis, genetic vaccine, expression library immunization.

**INTRODUCTION**

*Piscirickettsia salmonis* is the etiologic agent of Salmonid Rickettsial Septicemia (SRS) a disease responsible for extensive mortalities in the Chilean salmon industry. This agent was identified for the first time in 1990 by Fryer (Fryer et al., 1990). The disease occurs mainly in coho salmon (*Oncorhynchus kisutch*). It was found later that this pathogen affects other farmed salmon species (Gaggero, et al., 1995) including salmonids from other countries besides Chile (Brocklebank et al., 1993; Rodger and Drinan, 1993; Olsen et al., 1997).

The pathogen is a Gram-negative obligate intracellular bacterium found in cytoplasmic vacuoles of infected cells. The organism is pleomorphic, predominantly coccoid in shape and ranging 0.5 to 1.0 µm in diameter (Fryer et al., 1992). The pathogen was initially isolated from the kidney of a diseased adult coho salmon and successfully cultured in a cell line derived from Chinook salmon (*Oncorhynchus tshawytscha*) embryos, known as CHSE-214 cells (Fryer et al., 1990).

Presently, the SRS has been partially controlled by means of the use of antibiotics. These are not fully effective and have problems derived from their toxicity and generation of resistance. A more convenient procedure to prevent the infection is the use of vaccines. However, no effective vaccines are currently available against *P. salmonis*. This pathogen grows very slowly, taking 15 to 20 days to develop a full cytopathic effect making it difficult to produce large amounts of cells and to
separate them from host material, as required for an effective purify bacterin formulation.

Nucleic acid vaccines represent a new approach to the control of infectious agents (Leong et al., 1997). These novel experimental vaccines are both easy to construct and to produce, however the problem still remains of discovering what gene or genes to include in the genetic immunization vector to obtain an efficacious vaccine. This is particularly a problem with pathogens with larger genomes such as bacteria and parasites. One approach that circumvents this problem is the expression library immunization technology developed by Barry and coworkers (Barry et al., 1995). The procedure involves making an expression library from the whole genome of the pathogen in appropriate genetic immunization vectors. This expression library is used in successive immunizations and challenges to eventually strip down the genome into protecting genes (Johnston and Barry, 1997). These concepts were experimentally demonstrated when Barry and collaborators (Barry et al., 1995) showed that immunization of mice with expression libraries of Mycoplasma pulmonis induced protection against a challenge with the pathogen.

Here we report that immunization of Coho salmon with an expression library of P. salmonis induced antibody immunity and partial protection to a challenge with the pathogen. We have shown previously that DNA constructs containing P. salmonis coding (Tischler et al., 2002) regions elicit immunoresponses in mice (Wilhelm et al., 2002).

Growth and purification of P. salmonis

P. salmonis LF-89 was continuously propagated in the Chinook salmon embryo cell line CHSE-214 (CRL 1681 ATCC) (Lannan et al., 1984) using antibiotic-free minimal essential medium (MEM) plus 10% bovine serum as described elsewhere (Fryer et al., 1990, Jamett et al., 2001). The titer of P. salmonis was determined using a plaque formation assay (Lannan et al., 1991). Briefly, serial dilutions of P. salmonis were plated on fresh monolayers of CHSE-214 cells in 24-well plates and incubated at 16°C for 24 h. After the attachment of P. salmonis to the cells, the culture was overlaided with 0.7% methylcellulose in MEM. Plates were incubated at 16°C for 8 days and the number of plaques determined under phase microscopy. P. salmonis was purified using Percoll density gradient centrifugation (Jamett et al., 2001). The final pellet of purified bacterium was suspended in a solution containing 50 mM Tris-HCl, (pH 7.5), 200 mM NaCl, and the concentration of protein was measured (Smith et al., 1985). To determine the number of bacterial cells/ml, serial dilutions in the Tris-saline solution were counted under phase microscopy in a Neubauer chamber. In this procedure, 1 mg of protein corresponds to approximately 3.8 x 10^9 P. salmonis cells.

Preparation of genomic DNA from P. salmonis

Genomic P. salmonis DNA was prepared and purified as described (Wilhelm et al., 2003). Briefly, purified P. salmonis cells were first treated with DNAse I (Ogarkova and Balayeva, 1984) following the protocols of Hallberg (1994) and Binder (1995). The cells were suspended in 1 ml of DNAse I buffer (5 mM MgCl₂, 0.1 mM CaCl₂, 4 mM 2-mercaptoethanol, 0.1 mM p-methylsulphonylfluoride, 200 mM NaCl, 200 mM KCl and 50 mM Tris HCl (pH 7.4), treated with 20 µl of a 10 mg/ml DNAse I solution and incubated at 25°C for 60 minutes. Then, the cells were pelleted by centrifugation, resuspended in 0.5 ml of

MATERIALS AND METHODS

Bacterial strains

The bacterial strains used were: P. salmonis LF-89 (ATCC VR 1361), Escherichia coli HB101, ATCC 8739, Renibacterium salmoninarum ATCC 33209 and Yersinia ruckeri ATCC 29473.
PBS with 0.1 ml of 0.1 M EDTA, pelleted again by centrifugation and resuspended in cell lysis buffer (0.75 M sacarose, 400 mM NaCl, 20 mM EDTA, 0.2% SDS, 1 mg of proteinase K and 50 mM Tris HCl (pH 9.0)). The suspension was incubated at 58°C for 1 hour with gently stirring. The resulting DNA solution was extracted once with phenol saturated with Tris-HCl (pH 8.0) and twice with chloroform: isoamyl alcohol (24:1) and precipated for 30 min with 0.4 volumes of 5 M ammonium acetate and 2.5 volumes of absolute ethanol at -20°C. The pellet was washed with cold 70% ethanol and dissolved in 1 mM EDTA, 10 mM Tris HCl (pH 8.0).

Preparation of P. salmonis expression libraries

Purified P. salmonis DNA was sonicated at 4°C in aliquots of 15 mg each to an approximate size of 1,000 to 2,000 base pairs, as determined by agarose gel electrophoresis. Subsequently, the extremes of the DNA fragments were repaired using DNA polymerase I Klenow fragment and ligated overnight at 14-16°C to blunt-ended linearized plasmid pCMV-Bios (Figure 1) in a 10:1 fragment: vector molar relationship using T4 DNA ligase at a concentration of 20 U/µl. The ligation mixture was used to transform competent E. coli HB101 cells. For immunization, plasmid DNA pooled from 4 preparations from a total of 22,000 colonies was lyophilized and suspended in 0.9% of NaCl to a final DNA concentration of 0.1 mg/ml.

Immunization of Balb/c mice

Balb/c mice 6-8 weeks old were immunized by injection in the anterior tibia muscle with 5, 20 and 50 µg respectively of vector or library DNA dissolved in 0.9% NaCl, 25% sacarose and 50 ppm of tricaine metasulphonate.

Immunization of salmon

Coho salmon of approximately 20 g were treated with 30 ppm of tricaine metanesulphonate for 10 min and injected with 20 µg of vector or library DNA in 200 µl of sterile 0.9% NaCl, 25% sacarose and 50 ppm of tricaine metasulphonate split in four sites (50 µl each) at the dorsal anterior muscle. After 41 days, fish were immunized with an additional aliquot of 10 µg of DNA dissolved in 100 µl of the above solution, using the above procedure. All fish were maintained at an average temperature of 12°C, in alternating dark and light periods of 12 hr each.

The presence of P. salmonis-specific IgM antibodies in the serum of salmon immunized with DNA was determined by ELISA. Ninety six well microtiter plates were coated with a 3 µg/ml solution of Aulacomya ater bioadhesive (Burzio et al., 1996), incubated 30 min at room temperature, washed and dried for 30 min at 37°C. The wells of the coated plate were activated with 100 µl of a suspension containing 10 µg/ml of P. salmonis bacterial extract in PBS (pH 7.2) and incubated for 60 min at 37°C. Subsequently, the plates were incubated with a solution of 0.5% Tween 20 and 5% non-fat milk for 60 min at 37°C and dried for 60 min at 37°C.

The sera to be analyzed were serially diluted, incubated in each well for 3 hr at
room temperature. Then, the dilutions were discarded and the wells were washed 3 times with 0.02% Tween 20 in PBS. Subsequently, 100 µl of a solution of anti salmon IgM monoclonal antibody was added to each well and incubated 2 hr at room temperature. After washing 3 times with 0.02% Tween 20 in PBS, 100 µl of a peroxidase-conjugated anti mice IgG monoclonal antibody were added to each well, incubated 30 min at room temperature in the dark, developed with TMB and hydrogen peroxide and read at 450 nm.

Determination of the LD50

A total of 56 coho salmon of approximately 150 g each, were divided in 7 groups of 8 salmon each. Each group was inoculated by intraperitoneal injection with 200 µl of a suspension, containing increasing amounts of *Piscirickettsia salmonis* LF-89 in PBS containing 200 mM NaCl. The fate of the fish was followed for 8 weeks and the accumulative mortality was recorded for each group of fish treated with the different dose of pathogen.

Challenge of fish immunized with DNA formulations

Coho salmon immunized with the DNA expression library as well as with pCMV-Bios vector alone (control) were anesthetized and challenge by intraperitoneal injection of 200 µl of a suspension containing 200 mM NaCl containing *P. salmonis* (7.5 times the LD50). The challenge experiment was carried out for 30 days post-challenge and the mortality was verified daily. The presence of *P. salmonis* in dead fish was measured by immunofluorescence in kidney and brain tissues. The level of anti *P. salmonis* IgM was measured in selected animals by ELISA.

ELISA and Immunofluorescence

Polystyrene microtiter plates (PolySorp) were coated with the adhesive polyphenolic protein purified from the mussel *Aulacomya ater* (Buzio et al., 1996). Briefly, 100 µl of a 3 µg ml⁻¹ solution of the adhesive in 0.1% acetic acid was applied to each well and incubated for 30 min at room temperature. The acidic solution was discarded and the plate was air dried at 37°C for 1 h. To bind a given antigen to the coated plate, 100 µl of antigen solution in PBS containing between 0.75 and 12 µg ml⁻¹ of protein was applied to each well and incubated for 2 h at 37°C. After discarding the solution, the plate was washed once with distilled water and then blocked with a solution containing 2% casein and 2% sucrose in PBS for 2 h at room temperature. The blocking solution was discarded, and the plate was air dried for 2 h at 37°C and stored at 4°C in a sealed plastic envelope with desiccant. Further ELISA procedures and color development was carried out as described before (Aguayo et al., 2002).

To prepare the tissue extract, about 200 mg of salmon kidney was transferred to a plastic bag together with 2 volumes (w/v) of a solution containing 0.3 M sucrose, 0.15 M NaCl, 5 mM MgCl₂, 1 mM CaCl₂, and 10 mM sodium phosphate buffer (pH 7.5). The bag was sealed to avoid contamination and the kidney was macerated with a spatula. The macerate was transferred to a sterile Eppendorf tube and centrifuged at 2000 x g for 5 min.

To detect the presence of *P. salmonis* in fish tissues, samples of kidney and brain were subjected to indirect immunofluorescence using monoclonal antibody 7G4 as described before (Jamett et al., 2001).

RESULTS AND DISCUSSION

Culture of *P. salmonis*

*P. salmonis* is a strict intracellular bacteria which grows only in cultured cells such as salmon CHSE-214 cells. Figure 2 shows an electron microscopy picture of the CHSE-214 cells infected with *P. salmonis*. This picture depicts some of the characteristics of the intracellular growth of the bacteria such as their localization inside vacuoles of the host cells, the
presence of their double membrane with an ondulant external membrane and their appearance as electron-dense bodies in the bacterial cytoplasm. Cultured CHSE-214 cells infected with *P. salmonis* are considered ready for harvesting when nearly 100% of the cells showed cytopathic effects. At this stage the percentage of viable *P. salmonis* cells is always higher than 95%, the number of CHSE-214 cells is about 100,000 cells/cm² and the number of *P. salmonis* is about 1x10³ bacteria/cell.

**Purification of *P. salmonis***

Due to the large difference in genome size between bacteria and salmon, nuclei contamination of bacterial cells is a significant problem in obtaining highly purified bacterial DNA devoid of salmon DNA. One can calculate that one nuclei of a salmon cell contains 6,000 more DNA that one *P. salmonis* cell. Our first approach was the use of density gradient purification in sacarose. However, this procedure yielded bacteria contaminated with salmon DNA detected by amplification of the salmon growth hormone gene by PCR (results not shown). Because of this, we decided to use DNase I which could degrade salmon nuclear DNA and leave intact the pathogen DNA protected by the bacterial double membrane. This procedure was previously used by Ogarkova and Balayeva to obtain *Rickettsia prowazekii* DNA free of host chicken DNA (Ogarkova and Balayeva, 1984). Also, Binder (1995) used DNAse I to purify mitochondrial DNA and RNA from contaminating chromosomal DNA.

DNase I is very efficient, in the presence of Mg++ and traces of Ca++, in degrading single and double stranded DNA. Once completed the digestion, the enzyme can be completely inhibited by a mixture of 10 mM EDTA and 0.2 mM EGTA. The purity and integrity of the high MW bacterial DNA obtained by this procedure was demonstrated by a combination of gel electrophoresis to verify its size, amplification of *P. salmonis* 16S rRNA gene and by amplification (or its absence) of the salmon growth hormone gene (results not shown).

**Construction of *P. salmonis* expression libraries***

The *P. salmonis* genomic library was obtained employing a shotgun cloning procedure (Hengen, P., 1997) using DNA fragments obtained by sonication (Figure 3). Two libraries were obtained, each from 15 µg of *P. salmonis* DNA, resulting in 22,242 and 28,365 colonies respectively. Considering this number of colonies and assuming DNA fragments of 1,000 base pairs and a bacterial genome size of 1 x 10⁶ it is possible to calculate that they are equivalent to 22.2 and 28.4 bacterial genomes. Correcting for orientation in the vector (1/2) and reading frame (1/3), these libraries represented approximately 3.7 and 4.7 bacterial genomes, respectively.

The presence of DNA inserts in the plasmids from the DNA library was demonstrated by double digestion of
plasmid DNA from 12 randomly selected transformant colonies with the endonucleases EcoRI and HindIII. The results of Figure 4 show that all of them have inserts. Plasmid DNA from colonies 2, 4, 5, 9, 10, 11 and 12 gave 2 fragments, one of 900 to 1,100 base pairs and another of 450 to 600 base pairs. DNA from colonies 3 and 6 yielded only one fragment of about 550 to 600 base pairs and colony 8 yielded 2 fragments of approximately 200 to 300 base pairs. The presence of multiple fragments is probably the result of internal EcoRI or HindIII sites in the cloned DNA. The analysis was extended to DNA from 30 bacterial colonies with very similar results. These experiments indicate that about 80% of the plasmid library contains DNA inserts of the expected size.

**Immunization of Coho salmon with the expression library and challenge with P. salmonis.**

Fish were vaccinated as described in Materials and Methods. Sixty days after vaccination fish were challenged with one dose of $2.5 \times 10^7$ *P. salmonis* which corresponded to 7.5 times the determined LD50. One week after the challenge, the fish showed a slower activity quite different to non-challenged animals. About day 10 after challenge fish started to die. After 20 days, 100% of vector vaccinated control fish and 80% of the DNA vaccinated were dead. This percentage was unchanged in the following weeks (Figure 5). The vaccinated
fish that survived recovered their appetite and vitality with a behavior identical to the unchallenged control fish. All dead fish were analyzed macroscopically and found to show all the signs described for SRS. These were anemia as evidenced by white gills; high amounts of mucous liquid in the abdominal cavity, and evident hemolysis and destruction of organs such as kidneys, spleen, liver, brain and heart. Direct immunofluorescence with anti-\textit{P. salmonis} monoclonal antibody 7G4 (Jamett et al., 2001) revealed the presence of a very high amount of bacteria in the kidney. The kidney is the target organ to detect infectious agents in salmonids due to its

![Figure 5](image-url)

**Figure 5.** Accumulated mortality of vaccinated and non-vaccinated Coho salmon after a challenge with \textit{P. salmonis}. □: control fish; ●: vaccinated fish.

![Figure 6](image-url)

**Figure 6.** Detection of \textit{P. salmonis} in salmon kidney tissues by direct immunofluorescence. Detection was carried out using monoclonal antibody 7GH conjugated to FITC. A, B, C and D: kidney tissue of dead fish after challenge; E, F, G and H: kidney tissue of non-vaccinated non-challenged fish; I, J, K and L: kidney tissue of vaccinated fish that survived the challenge.
hemaetopoyetic function in fish. The large amount of bacteria is in agreement with the high capacity of infection and replication of this bacteria in this host. This result is shown in Figure 6 where four representative fish are analyzed by immunofluorescence.

The vaccinated fish that survived the challenge were sacrificed and analyzed. These fish did not show the anatomopathologic signs present in non-surviving fish. All organs were found healthy and normal. In addition, analysis by immunofluorescence showed significantly less *P. salmonis* compared to the amount found in dead fish. It appears therefore that there is a direct correlation between the organ damage and the amount of bacteria detected. We believe this lower damage and lower bacterial load are a consequence of the immunization with the library DNA. However, it is also possible that the higher resistance of these fish may be due to intrinsic genetic factors of a subgroup of fish. The somewhat low percentage of protection may be due to several reasons. One could the very high amount of challenge bacteria used in this first attempt (2.5 x 10^7) which is significantly higher that the amount involved in a natural infection. Another explanation could be the very high complexity of the library used which would imply a very low concentration of the hypothetical effective genes.

**Evaluation of immunoresponse of vaccinated Coho salmon**

The humoral response of vaccinated and non vaccinated control fish was evaluated by ELISA after the first and second dosis of vaccine. The results are shown in Figure 7. At 40 days, fish vaccinated with 20 µg of library DNA showed a level of *P. salmonis* IgM higher than control fish. At 100 days after vaccination, which included a second

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**Figure 7.** Detection of anti-*P. salmonis* IgM in salmon vaccinated with DNA from the library.

1st DNA: response at 40 days after the first immunization with 20 µg of DNA. 2nd DNA: response at 60 days after a second immunization with 10 µg of DNA.
Figure 8. Specificity of the humoral response of salmon vaccinated with the *P. salmonis* DNA library.

Figure 9. Dose-dependent anti *P. salmonis* humoral response of mice vaccinated with the *P. salmonis* DNA library.
dosis of 10 µg of library DNA at day 60, the level of IgM did not increase significantly. This humoral response appears specific for *P. salmonis* since sera of fish vaccinated with the library did not react significantly with antigens of *Renibacterium salmoninarum* or *Yersinia ruckeri*, which are common pathogens of salmonids (Figure 8).

**Evaluation of immunoresponse in mice**

We were also interested in determining if the library DNA was able to induce a humoral response in mice. With this objective Balb c mice were injected with 5, 20, 50 and 250 µg of library DNA in one dosis. The level of antibodies was measured by ELISA using purified *P. salmonis* as antigen. The results are shown in Figure 9. They show that the response is dose-dependent with a very clear effect at the 250 µg dosis. This is quite similar to reports by others (Davis et al., 1993). This humoral response detected in immunized mice is specific for *P. salmonis*. No response was detected in ELISA assays using *Renibacterium salmoninarum* or *Yersinia ruckeri* as antigens (Figure 10).

**CONCLUSIONS**

The results described in this paper indicate that an experimental vaccine composed of a wide collection of expressable *P. salmonis* coding regions is able to produce a measurable level of protection in salmon to a challenge with the pathogen. This protection has been demonstrated by a decreased level of mortality, decreased bacterial load and the presence of specific anti *P. salmonis* IgM showed by the vaccinated fish.

The experimental approach described here can be continued further. It may now be possible to deconvolute this genomic expression library by successive immunizations and challenge with pools.

**Figure 10.** Specificity of the anti *P. salmonis* humoral response of mice vaccinated with the *P. salmonis* DNA library.
of decreasing complexity until the DNA clones responsible for protection can be identified and isolated.

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