In vitro activation of rainbow trout macrophages stimulates inhibition of Renibacterium salmoninarum growth concomitant with augmented generation of respiratory burst products

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ABSTRACT: Rainbow trout head kidney macrophages pretreated for 48 h with a supernatant which contained macrophage activating factor (MAF) obtained from Concanavalin A/phorbol myristate acetate stimulated rainbow trout leucocytes express an enhanced ability to inhibit Renibacterium salmoninarum growth in vitro compared with untreated controls. Enhanced 'killing' capacity of these macrophages was detectable after 24 h exposure to R. salmoninarum in vitro. Addition of N⁶-methyl-L-arginine in vitro did not diminish bacterial growth inhibition by MAF-activated macrophages, suggesting that nitric oxide radicals were not involved in this event. However, MAF-treated macrophages expressed enhanced respiratory burst activity compared with control macrophages after a 3 d exposure to R. salmoninarum in vitro. In addition, it was demonstrated that R. salmoninarum was susceptible to hydrogen peroxide (H₂O₂) in vitro using a cell-free assay. Introduction of catalase to the in vitro macrophage assay abrogated the 'killing' activity, suggesting that H₂O₂ release from activated macrophages was involved in the inhibition of R. salmoninarum growth. The results are discussed in relation to other intracellular pathogens and the possible importance of cell-mediated immunity for protection against bacterial kidney disease.

KEY WORDS: Renibacterium salmoninarum · Macrophage activation · Respiratory burst · Growth inhibition

INTRODUCTION

Bacterial kidney disease (BKD), induced by the Gram-positive bacillus Renibacterium salmoninarum, is a significant infection of cultured and feral salmonids (Evelyn 1993). The disease is typically slowly progressive and systemic, characterised by renal and muscular granulomata, abdominal ascites, internal haemorrhage and anaemia. The bacillus is able to survive within cells of the mononuclear phagocyte system (Young & Chapman 1978), a scenario reminiscent of several bacterial (Dannenberg 1991, Baldwin et al. 1993) and parasitic infections in mammals (Liew & Cox 1991). This intracellular strategy may afford protection from extracellular mechanisms of host defence, including antibody binding and complement fixation. Indeed, this may account for the lack of protection conferred by many R. salmoninarum antigen preparations although they stimulate elevated agglutinin levels in salmonids (Baudin Laurencin et al. 1977, Evelyn 1993). Furthermore, opsonization of R. salmoninarum with specific antisera and complement significantly enhances their survival and growth following phagocytosis by normal rainbow trout macrophages (Bandin et al. 1995). Access to intracellular sites by pathogens often in-
volves mimicry of recognition molecules such as selectins and integrins (Sandros & Tuomanen 1993) or fixation of complement to induce phagocytosis via complement receptors (Kaufmann 1993). In this way cellular defences may be subverted, since phagocytosis by these ligands does not trigger bactericidal pathways such as the release of toxic reactive oxygen radicals, and once intracellular, pathogens may further thwart host defences by preventing phagolysosomal fusion or by entering the cytosol (Baldwin et al. 1993). It is presumed that R. salmoninarum must employ similar tactics. The bacterium is known to activate the alternative complement pathway with subsequent covalent binding of C3b, and this enhances phagocytosis by trout macrophages in vitro (Rose & Levine 1992).

In mammals, intracellular pathogens such as Ehrlichia spp. and Leishmania spp. can be eliminated through experimental upregulation of cell-mediated immune responses (Liew et al. 1990, Park & Rikihisa 1992). Particular cytokines appear to play pivotal roles as paracrine or autocrine modulators of macrophage function in resistance to various intracellular pathogens and include interferon (IFN)-γ, tumor necrosis factor (TNF)-α and interleukins (IL) 1 and 6 (Titus et al. 1991, Nauciel & Espinasse-Maes 1992, Denis & Ghadirian 1994). Typically, these cytokines upregulate phagocyte cytoidal pathways, including the release of toxic reactive oxygen and nitrogen species (Ding et al. 1988, Thomson 1991). In fish, it is known that salmonid phagocyte functions can be upregulated experimentally in vitro using supernatants from antigen or mitogen stimulated leucocytes, which are thus deemed to contain a macrophage activating factor (MAF) (Graham & Secombes 1988, Marsden et al. 1994). MAF treatment of macrophages results in elevated bactericidal activity against Aeromonas salmonicida, probably as a result of elevated exposure to toxic H2O2 species (Sharp & Secombes 1993). However, evidence is growing to support the notion that nitric oxide synthase is present in teleosts (Schober et al. 1993, Hardie et al. 1994b, Schoor & Plumb 1994, Wang et al. 1995) and thus nitrogen radicals may also have a protective role to play in disease elimination in these animals.

Salmonid macrophages do show a weak capacity to inhibit Renibacterium salmoninarum growth in vitro when cultured together for 5 or more days, although the mechanism for this elimination is not understood (Bandin et al. 1992). Therefore, the aim of this study was to ascertain whether the cellular immune response in salmonids may potentially afford protection to R. salmoninarum through upregulation of phagocyte functions by cytokines. To this end, MAF-containing supernatants were examined for their ability to upregulate rainbow trout macrophage-mediated inhibition of R. salmoninarum growth in vitro. This was demonstrated after a 7 d culture period with bacteria, known to be sufficient time to allow some inhibition by normal macrophages (Bandin et al. 1992), and after a 1 d co-culture period, previously shown to be insufficient time for growth inhibition. The potential mechanisms of this phenomenon were also investigated.

**MATERIALS AND METHODS**

**Fish.** Rainbow trout Oncorhynchus mykiss weighing 300 to 500 g were obtained from College Mill trout farm, Almond Bank, Perthshire, UK. Fish were acclimated to 14°C in aerated, dechlorinated mains water and fed twice daily on commercial trout pellets (EWOS) for at least 2 wk before use.

**Preparation of head kidney macrophage monolayers.** Phagocyte-enriched head kidney leucocyte suspensions were obtained as previously described (Secombes 1990). Briefly, head kidney leucocyte suspensions were prepared in heparinized Leibovitz medium (L15, Gibco) and separated on a 34%/51% discontinuous Percoll density gradient. The phagocyte-enriched fraction was collected from the interface, washed twice in L15 and resuspended to 2 x 10^7 viable cells ml^-1 in L15 containing 0.1% foetal calf serum (FCS, Gibco). Aliquots (100 μl) of this suspension were dispensed into wells of a 96 well microtitre plate (Nunc). After 3 h at 18°C, non-adherent cells were washed off with L15 medium. The remaining macrophage monolayers were incubated with 100 μl L15 plus 5% FCS and washed again with this medium before use. All solutions were antibiotic-free.

**Production of MAF supernatants.** A MAF supernatant was produced using the method of Graham & Secombes (1988), with the exception that antibiotics were excluded from this protocol to avoid effects on subsequent bactericidal assays (see below). Briefly, head kidney leucocytes obtained from a rainbow trout were stimulated with mitogens (10 μg Concanavalin A ml^-1 plus 5 μg phorbol myristate acetate ml^-1, Sigma) for 3 h. After thorough washing, treated cells were cultured in L15 plus 10% FCS for a further 48 h before the supernatant was harvested.

The presence of MAF activity in the supernatant was confirmed by its ability to augment respiratory burst activity of test macrophages in vitro as assessed by the ferricytochrome c reduction technique (see below). Optimal stimulatory effects were observed at a 1:4 dilution of the supernatant (data not shown), and subsequent macrophage stimulations used this concentration of MAF.

**Bacteria.** A strain of Renibacterium salmoninarum (MT426) originally isolated from a clinical outbreak of
BKD in rainbow trout in Scotland was obtained from the SOAEFD Marine Laboratory culture collection. This isolate expressed characteristics associated with virulence (Bruno 1988), including an autoaggregating capacity and the presence of the surface-associated p57 protein as ascertained by SDS-PAGE (data not shown). Bacteria were cultured on Mueller Hinton Agar (Difco) supplemented with 0.1% L-cysteine hydrochloride for 7 d at 15°C. Bacteria were washed off the agar with L15 medium, washed twice, counted and then serially diluted in L15 plus 5% FCS or phenol-red-free Hanks’ balanced salt solution (HBSS, Gibco) for bactericidal or cell free assays, respectively (see below). Before bactericidal assays, FCS was screened to confirm that there were no deleterious effects on R. salmoninarum growth in vitro.

Co-culture assays. These assays are based upon procedures previously described (Graham et al. 1988, Bandín et al. 1992). Macrophage monolayers from 3 fish per experiment, either untreated or pretreated for 48 h with MAF, were washed twice with L15 before the addition of 50 μL L15 plus 5% FCS. Macrophage monolayers which had been washed and lysed with 50 μL cold sterile distilled water served as controls. A volume of 50 μL of R. salmoninarum suspended in L15 plus 5% FCS was then added to macrophage monolayers in triplicate to give concentrations ranging from 0.25 to 2 X 10^7 bacteria well^-1. In addition, various micromolar concentrations of N^6-methyl-L-arginine (L-NMMA, Sigma) were added to some wells to inhibit putative nitric oxide synthase activity (Wang et al. 1995). Furthermore, 0.3 mg catalase ml^-1 (Sigma) was introduced in some wells to degrade hydrogen peroxide (H_2O_2) produced during the in vitro assay. Throughout, reagents were diluted in appropriate media and added to give a constant final volume of 200 μL well^-1 across all treatments.

Microtitre plates were then centrifuged for 5 min at 150 x g, to aid contact between bacteria and macrophages. After a period of 1 or 7 d at 18°C, infected cultures were again centrifuged at 150 x g, the supernatant aspirated, and 50 μL of cold sterile distilled water added, which effectively lysed the macrophages, releasing any intracellular bacteria. A volume of 100 μL of L15 containing 5% FCS was added to all wells, and plates which had only been exposed to macrophages for 1 d were given a 6 d ‘grow up’ period so that the relationship between macrophage exposure time and bacterial level could be assessed at a fixed and therefore comparable time point. In addition, because of the lower starting concentrations of bacteria used and the slow in vitro doubling time of R. salmoninarum, the 6 d grow up period was required to permit the bacteria to reach concentrations within the detection range of the assay. Thus, after a total of 7 d in vitro for all cultures, bacterial viability was quantified by the addition of 10 μL 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT, 5 mg ml^-1; Sigma) per well. The optical density (600 nm) of each well was assessed after exactly 15 min using a multiScan spectrophotometer (MDC). In some instances (the catalase co-culture experiment) data were expressed as a percentage of the values obtained using untreated macrophages.

Cell-free generation of oxygen radicals. To determine the susceptibility of R. salmoninarum to oxygen radical species a photochemical radical-generating system was employed (Karczewski et al. 1991). In this system R. salmoninarum were exposed to levels of oxygen radicals, such as superoxide anion (approx. 7 nmol well^-1), consistent with quantities that might be generated from MAF-activated macrophages in vitro (Jang et al. 1994). Briefly, washed R. salmoninarum were serially diluted in phenol-red-free HBSS containing 12.5 mM methionine and 0.1 mM riboflavin. Aliquots (100 μL) were then dispensed into wells to give a range of 1.6 x 10^4 to 10^6 bacteria well^-1. Catalase (0.3 mg ml^-1) or superoxide dismutase (SOD, Sigma, 300 IU ml^-1) was also introduced to some wells to degrade any generated hydrogen peroxide or superoxide radicals, respectively. The riboflavin was then photoreduced by a 5 min exposure to a 21 W light source with a Kodak no. 47 blue filter (360 to 520 nm) at a range of 7 cm, the methionine serving as a hydrogen donor. Following this, the plates were incubated for 5 h at room temperature in natural light to support further radical generation, then centrifuged at 150 x g for 5 min and the supernatant carefully removed. Finally, 100 μL of L15 plus 5% FCS was added per well to support R. salmoninarum growth for the next 7 d at 18°C. Bacterial growth was assessed by MTT reduction as outlined above.

Respiratory burst activity. Macrophage monolayers, some of which had been pretreated for 48 h with MAF, were washed twice with L15 before the addition of R. salmoninarum at concentrations ranging from 2.5 x 10^6 to 3.2 x 10^7 bacteria in 100 μL L15 plus 5% FCS. The microtitre plates were then centrifuged for 5 min at 150 x g to aid contact between bacteria and macrophages. After 3 d at 18°C, infected cultures were centrifuged and the supernatant aspirated. The plates were carefully washed twice with phenol-red-free HBSS, using centrifugation and aspiration steps. To each well, 100 μL HBSS containing ferricytochrome c (2 mg ml^-1, Sigma) and phorbol myristate acetate (1 μg ml^-1) was then added to stimulate the respiratory burst. As a control for specificity, 300 IU SOD ml^-1 was added to some wells. The optical density of each well was measured at 550 nm after 30 min using a multiScan spectrophotometer, with the relevant SOD control act-
ing as the blank. Optical densities were converted to nmol O₂ produced, as described by Pick (1986), i.e. by multiplying by a factor of 15.87.

Statistics. Data were analysed using 1- and 2-way analysis of variance (ANOVA), and Student's t-test.

RESULTS

The results show that macrophages pretreated with a MAF supernatant for 48 h express a markedly enhanced (p < 0.001) ability to inhibit in vitro growth of *Renibacterium salmoninarum* across all bacterial concentrations tested, compared with untreated counterparts, when incubated with the bacteria for 7 d (Fig. 1). Untreated macrophages did not show this activity compared with lysed macrophage controls. Indeed, bacterial growth was significantly higher (p < 0.001) in the intact macrophage group compared with the lysed control. The effectiveness of MAF pretreatment of macrophages was clear even after only 24 h of contact between cells and bacteria (Fig. 2), as determined overall by 2-way ANOVA (p < 0.01). Indeed, with the exception of the highest bacterial concentration, the other bacterial concentrations showed typically that MAF pretreatment of macrophages resulted in a significant (p < 0.05) reduction in bacterial growth, although clearly not to the same degree as that seen after a 7 d co-culture period. The inclusion of L-NMMA in vitro did not diminish the effect of MAF-activated...
HBSS only
riboflavin + methionine
riboflavin + methionine + SOD
riboflavin + methionine + catalase
riboflavin + methionine + SOD + catalase

![Graph showing bacterial growth](image)

Fig. 4. Effect of a 5 h exposure to chemically generated oxygen radicals on the in vitro growth of varying concentrations of *Renibacterium salmoninarum*. In some wells superoxide dismutase (SOD, 300 IU ml⁻¹) and/or catalase (0.3 mg ml⁻¹) were introduced to sequester O₂⁻ or H₂O₂ respectively. Growth was assessed following a 7 d 'grow up', as measured by MTT reduction (optical density 620 nm). Data are means ±SE of triplicate values.

Utilising a photochemical generation method, it was demonstrated that *Renibacterium salmoninarum* was susceptible to oxygen radical species. When bacteria were exposed to an activated riboflavin and methionine solution, a significant (p < 0.001) reduction in growth occurred across all bacterial concentrations tested compared with bacteria in HBSS alone (Fig. 4). This result was attributable to radical products released, as neither methionine nor riboflavin solutions showed individual effects in vitro (data not shown).

Furthermore, when SOD and catalase were introduced jointly to break down O₂⁻ and H₂O₂, respectively, from this system, bacterial growth remained at levels that were not significantly different to the HBSS control. To elucidate whether one or both radical species were responsible for killing, each enzyme was added individually. The addition of SOD alone did not prevent significant (p < 0.001) growth inhibition from occurring. In contrast, catalase inclusion in vitro again retained bacterial growth at levels not significantly different to the HBSS control.

Examination of the respiratory burst activity of macrophages which had been exposed to *Renibacterium salmoninarum* for 3 d showed that there was a significant (p < 0.001) difference between control cells and MAF-treated cells across all bacterial concentrations tested (Fig. 5). Notably, the respiratory burst of untreated cells became exhausted using at least 2 doubling dilutions of bacteria before MAF-treated cells. Interestingly, the proportional difference in respiratory burst activity between MAF-treated and untreated cells increased in magnitude with increasing bacterial numbers (Fig. 5).

As greater respiratory burst activity in MAF-treated macrophages would potentially result in elevated H₂O₂ release in vitro, catalase was introduced into in vitro co-culture assays to determine if H₂O₂ was involved in MAF-mediated growth inhibition. The results of this experiment (Fig. 6) further demonstrate that MAF can mediate significant (p < 0.001) *Renibacterium salmoninarum* growth inhibition by macrophages, although the effect was not as dramatic as in Fig 1 due to the use of a different batch of MAF super-
showed that untreated trout macrophages could
acquire `killing' activity when co-cultured for 5 or more
days with *R. salmoninarum*. This apparent disparity
may be resolved when experimental variables such as
the level of macrophage activation *in vivo* prior to
tissue culture or whether macrophages became nutri-
tionally deprived *in vitro*, a factor shown to stimulate
spontaneous activation in mammalian phagocytes
(Walker et al. 1991), are considered. Nevertheless,
cytokine-activated macrophages are clearly able to
inhibit *R. salmoninarum* after co-culture for as little as
1 d, which untreated macrophages cannot.

It appears that nitric oxide radical generation was
not involved in the retardation of *Renibacterium
salmoninarum* growth during the present study
because the inclusion of L-NMMA at concentrations
known to inhibit nitric oxide production in mammaian
and fish studies (Liew & Cox 1991, Wang et al. 1995)
had no impact. Rather, upregulation of the respiratory
burst appears to be responsible for *R. salmoninarum
'killing' by these MAF-treated cells. This study and the
results of Bandin et al. (1992) have shown that under
non-activated conditions the respiratory burst of trout
macrophages co-incubated with *R. salmoninarum* is
usually compromised during the course of a 6 d *in vitro*
infection, with maximal effects from Day 3 onwards.
However, in the present study, MAF treatment elev-
ated macrophage respiratory burst capacity during *in
vitro* culture with *R. salmoninarum*. That *R.
salmoninarum* was sensitive to *H₂O₂* exposure in a cell-free
system and that catalase abrogated growth inhibition
during the *in vitro* macrophage assay suggests that
MAF pretreatment permitted adequate *H₂O₂* genera-
tion by macrophages during the respiratory burst to
check *Renibacterium* growth, and that this effect was
likely to be a consequence of the bactericidal activity
of *H₂O₂* (Halliwell & Gutteridge 1989). A trend to
increased 'killing' was also seen in the cell-free system
when SOD was included, presumably due to the
increased *H₂O₂* levels produced following dismutation
of the generated superoxide anion. Although the
'killing' mechanism was *H₂O₂* dependent, *H₂O₂*
derivatives such as hypohalides, hydroxyl or singlet oxygen
species may also have been involved. The situation
described above parallels findings with some intracel-
lar pathogens of mammals such as *Brucella abortus*
(Baldwin et al. 1993) and *Cowdria ruminantium*
(Mahan et al. 1994).

An increased phagocytic uptake of *Renibacterium
salmoninarum* by activated macrophages may also
have contributed to this phenomenon since activated
tROUT macrophages typically have an increased phago-
cyotic uptake (Secombes & Fletcher 1992). This scena-
rio, however, assumes that killing of *R. salmoninarum*
can only occur intracellularly in fish macrophages.

DISCUSSION

The results presented indicate that rainbow trout
macrophages are capable of restricting *Renibacterium
salmoninarum* growth when activated by appropriate
cytokine stimulation *in vitro*, although whether the
observed growth inhibition is due to a bactericidal or
bacteriostatic mechanism is not clear and awaits fur-
ther study. Growth of *R. salmoninarum* was not inhib-
ited by untreated macrophages in the present study;
indeed, growth was better supported by intact
untreated macrophages compared with lysed macro-
phages, possibly as a result of microbicidal cytoplasmic
products released when macrophages were lysed, or it
could be that an intracellular microenvironment pro-
vided better conditions for growth. These findings con-
trast with the results of Bandin et al. (1992) who
showed that untreated trout macrophages could

![Graph](https://via.placeholder.com/150)

**Fig. 6.** Effect of catalase on *Renibacterium salmoninarum* growth inhibition when co-cultured for 7 d *in vitro* with un-
treated or MAF-treated macrophages, as measured by MTT
reduction. Data are expressed as a percentage of the optical
density values obtained with untreated macrophages in the
absence of catalase, and are means ±SE of triplicate fish

ntant. Introduction of catalase to untreated and MAF-
treated macrophages significantly enhanced bacterial
growth (*p* < 0.001) across all concentrations of bacteria
tested. In addition, the significant difference in *R.
salmoninarum* growth curves between control and
MAF-treated macrophages was no longer present
when catalase was introduced *in vitro*, suggesting that
the enhanced growth inhibition of *R. salmoninarum* in
response to MAF treatment is *H₂O₂* dependent.
fish bacterial pathogens such as *Aeromonas salmonicida* (Sharp & Secombes 1992), no data are currently available for *R. salmoninarum*. Nevertheless, large numbers of *R. salmoninarum* have been shown to be phagocytosed by normal rainbow trout macrophages following co-cultivation for 24 h (Bandin et al. 1995). As normal macrophages have low or no inhibiting effect on *R. salmoninarum* growth, phagocytosis of *R. salmoninarum* per se does not appear to relate to killing.

Previous studies using *Aeromonas salmonicida* have shown that immunisation of trout can prime for MAF release following a subsequent exposure to the homologous antigen (Marsden et al. 1994). Whilst it is not known whether MAF release can be primed with *Renibacterium salmoninarum*, it is clear from this study that, following activation, salmonid macrophages have the capability to suppress or eliminate *R. salmoninarum* infection. So are there any reasons why this may not occur *in vivo?* Temperature may be one important factor. Outbreaks of BKD in rainbow trout and Atlantic salmon follow a seasonal pattern in Scotland, occurring between February and September, rising to a peak in May (Bruno 1986a). Indeed, recovery from the disease during experimental infections has been shown to be temperature dependent (Bruno 1986b, 1988). In Scotland, freshwater temperatures do not reach 10°C until early/mid May, and rise to about 14°C in June/July. Since fish primary T cell responses, including production of MAF (Hardie et al. 1994a), are temperature sensitive (Bly & Clem 1992), it is likely that immediately prior to BKD outbreaks rainbow trout would have a restricted MAF-producing capacity on first encounter with *R. salmoninarum*. As temperatures rise during May/June, it may be possible to elevate MAF production, permitting macrophages to become activated and contribute to the clearance of bacteria.

As rainbow trout have the immunological armoury to combat *Renibacterium salmoninarum*, it should be possible to design vaccines which mobilise these defences in an appropriate manner. *In vivo*, the immune system can activate different mechanisms for attacking and eliminating pathogens. It is known in mammalian studies that helper T cells (TH) are required for both antibody and cell-mediated responses, and are composed of distinct subsets, distinguished by characteristic cytokine signatures (Del Prete & Romagnani 1994). Thus, TH1 cells secrete IFN-γ, IL2 and TNF-β, whereas TH2 cells secrete IL4, IL5, IL6 and IL10. Formation of the appropriate TH subset is fundamental for the resolution of disease states in mammals, and determines whether a cell-mediated or humoral (antibody) immune response predominates (Johnson & McMurray 1994, Surcel et al. 1994). Whilst virtually nothing is known about T cell subsets in fish, their immune system is remarkably similar to that seen in other vertebrates (Faisal & Hetrick 1992, Secombes 1994). Thus, by stimulating the equivalent of a TH1-like response in salmonids (i.e. cell-mediated immunity) resulting in MAF-stimulated macrophage inhibition of *R. salmoninarum*, it may be possible to induce protection. That antibody responses to various *R. salmoninarum* antigens are commonly not protective to the fish (Evelyn 1993) but indeed appear to promote the intracellular survival of the bacterium (Bandin et al. 1995), may indicate that TH2-like responses (i.e. humoral immunity) are being driven in many instances. Protection would not be elicited during a TH2-like response because suitable factors such as IFN-γ analogues (i.e. MAF) would not be available to activate macrophages. If TH subsets appropriate to *R. salmoninarum* were developed at permissive temperatures in salmonids, it would be expected that on restimulation by BKD infection, primed T cells would mount a MAF response at low environmental temperatures, with the potential to obviate spring outbreaks of BKD.

However, TH1 subset development is governed by the nature of primary antigen encounter as this determines subsequent antigen processing and presentation. Macrophages exposed to high doses of live or fixed *Mycobacterium bovis* are provoked into ‘exogenous’ pathways of antigen presentation which bias TH2 development and cell activation (Griffin & Buchan 1993). In contrast, recruitment of TH1 cells appears to be stimulated by endogenous processing of low doses of virulent or attenuated intracellular pathogens and IL12 signals from presenting macrophages (Bretscher 1992, Hsieh et al. 1993). These findings have prompted suggestions that cytokines, including IL2 and IFN-γ, could be used as immune-adjuvants or immunostimulants to propel TH1 formation where appropriate. Since many intracellular bacteria exhibit phagosome dependent expression of certain genes (Plum & Clark-Curtiss 1994) and intracellular growth is often a prerequisite to successful presentation and induction of T-cell mediated immunity (Berche et al. 1987), it is possible that key *Renibacterium salmoninarum* antigens may not have been expressed during previous attempts to produce vaccines employing whole broth culture preparations and extractions (Evelyn 1993). Thus, more sophisticated approaches to the isolation and presentation of immunologically pertinent antigens may permit immunoprophylaxis against this widespread and economically important fish disease.

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