Comparative evaluation of bacterial culture and two ELISA techniques for the detection of *Renibacterium salmoninarum* antigens in salmonid kidney tissues

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**ABSTRACT** Soluble antigens from *Renibacterium salmoninarum* (Rs) incorporated into immunostimulating complexes (iscoms) induced polyclonal antibodies directed to the heat-stable, 57 kDa antigen complex of Rs. The antibodies were used in an enzyme-linked immunosorbent assay (ELISA) for the detection of Rs antigens in salmonid kidney samples. The negative-positive threshold value in the ELISA was determined to be 0.10 based on the analyses of 408 samples from rainbow trout *Oncorhynchus mykiss*, Arctic char *Salvelinus alpinus*, Baltic salmon *Salmo salar*, and brown trout *Salmo trutta*, all negative for Rs by bacterial culture. The Rs organism was isolated on selective kidney disease medium (SKDM) after up to 12 wk of incubation from 45% of the samples collected from fish farms having enzootic bacterial kidney disease (BKD), and from fish in populations with clinical signs of BKD (n = 260). Soluble antigens of Rs were detected with the polyclonal ELISA in 50% of these samples. In a comparative evaluation of 3 diagnostic techniques (bacterial culture, the polyclonal ELISA described here, and a commercially available ELISA based on monoclonal antibodies), 167 kidney samples were individually analyzed. The results of the diagnostic tests (positive or negative) were identical for 95% of the examined samples. A higher sensitivity of the polyclonal ELISA when compared with the monoclonal method was indicated in this study.

**KEY WORDS**: Bacterial kidney disease · Diagnostics · ELISA · *Renibacterium salmoninarum* · Iscom

**INTRODUCTION**

*Renibacterium salmoninarum* (Rs) is the causative agent of bacterial kidney disease (BKD), a chronic systemic infection of salmonid fishes (Fryer & Sanders 1981) which can be transmitted vertically (Evelyn et al. 1984a, b) as well as horizontally (Bell et al. 1984). As chemotherapy and vaccination have only limited effects, the control of BKD is dependent on sanitary methods and a continuous health monitoring of farmed and feral fish populations.

In Sweden, the first case of BKD was diagnosed in 1983. Since then, the causative agent has been detected, with different degrees of disease manifestation, from about 50 fish farms, mainly on the east coast. Rainbow trout *Oncorhynchus mykiss*, the species commonly reared in Swedish fish farming, seems to be more resistant to BKD than Arctic char *Salvelinus alpinus* and Baltic salmon *Salmo salar*.

The Rs agent is a slow-growing, Gram-positive short rod that survives and multiplies intracellularly in the host (Young & Chapman 1978). Confirmatory diagnostic methods used in the control program for clinical BKD in Sweden since 1985 are Gram staining, immunofluorescent antibody staining of smears and the peroxidase-antiperoxidase immunohistochemical technique (Jansson et al. 1991). Fish populations have routinely been tested by bacterial culture to detect car-
MATERIALS AND METHODS

**Bacteria.** The bacterial strains included in this study are listed in Table 1. Strains of Rs isolated from fish with clinical signs of BKD were grown on charcoal agar. KDM-C agar (Daly & Stevenson 1985) or on selective kidney disease medium, SKDM agar (Austin et al. 1983), supplemented with 10% heat-inactivated sheep serum instead of calf serum, at 15°C. All other bacteria were grown on horse blood (5%) agar at 20°C or 37°C.

**Polycylonal antibody preparations.** Antigens for iscom preparation were prepared from Rs strain 4/86 (Table 1), cultivated on KDM-C agar for 15 d as follows. Bacteria were suspended in peptone-saline (0.1% peptone, Difco laboratories, Detroit, MI, USA) and centrifuged at 2500 g for 20 min. The pellet was resuspended in distilled water containing 2% of the detergent MEGA 10 (Hi!dreth 1952, Hanatani et al. 1984) and incubated at 37°C with mild agitation for 1 h. The suspension was sonicated and centrifuged (3000 x g) for 10 min. The protein content of the supernatant was quantified by the method of Bradford (1976), and iscoms were prepared according to the principles of Morein et al. (1984). Briefly, the supernatant was mixed with 1-3-phosphatidyl choline 1,2-dipalmitoyl, cholesterol (1% of each) and the glycocide Quil A (0.2%). The mixture was dialysed against phosphate-buffered saline, pH 7.2 (PBS) for 5 h at room temperature and then at 4°C. Antigens were also extracted from the bacterium by washing in distilled water (pH 5.5) according to the hemagglutinin water-extraction method described by Daly (1989).

Rabbits were injected intramuscularly with 300 µl of the iscom preparation (21 µg protein) or with 50 µg of the extracted hemagglutinin in Freund’s complete adjuvant and booster-injected 3 and 5 wk later, in the same manner except that Freund’s incomplete adjuvant was used for the booster injections with the hemagglutinin. Blood was collected after an additional 2 wk and 1 and 2 wk thereafter. Repeated booster injections were administered every 10th week. Blood samples collected before the first immunization were used as a control. Rabbit IgG was purified from serum on a Protein G column (Pierce, Rockford, IL, USA), dialysed against PBS and distilled water and then freeze-dried. The effectiveness of the resulting rabbit IgGs as capture antibodies in ELISA was compared by the Mann-Whitney Wilcoxon test (Altman 1991).

**SDS-PAGE and Western-immunoblotting.** The Rs strain 4/86, was inoculated into a semidefined medium (Embley et al. 1982) and incubated with agitation at 17°C for 4 wk. Bacterial growth was harvested, diluted, and treated as described for the preparation of samples for the polyclonal ELISA. The supernatant was boiled
Table 1. Bacterial strains isolated from fish and other sources that were tested by the polyclonal ELISA (rabbit anti-Rs iscoms as capture antibodies). CCUG: Culture collection of the University of Göteborg, Sweden; NVI-N: National Veterinary Institute, Oslo, Norway; NVI-S: National Veterinary Institute, Uppsala, Sweden.

<table>
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<th>Source</th>
<th>Species</th>
<th>ELISA result</th>
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for 5 min in a Tris-Cl buffer, pH 6.8, containing 10% (w/v) sodium dodecyl sulphate (SDS) and 50 mM dithiothreitol. The proteins were analysed by polyacrylamide-gel electrophoresis in the presence of SDS in 12% polyacrylamide [Mini-Protean II System, Bio Rad]. Polypeptide bands were visualized with 0.1% Coomassie blue R-250 solution. Western-immunoblotting was performed to detect antibodies to the different proteins in the supernatant by first separating 25 μg of sample and 5 μg of BioRad standard in a 4% SDS stacking gel and in a 12% resolving gel at 200 V for 45 min. Proteins were transferred to a nitrocellulose (NC) filter for 1.3 h at 70 V. The filter was blocked with 5% milk powder (Semper, Stockholm, Sweden) in Tris-Cl buffer, pH 7.5 (TBS) for 1 h and then washed with TBS containing 0.5% Tween-20 (TTBS). This was followed by a 2 h incubation with the prepared rabbit IgG, diluted 1:50 in TTBS containing 1% milk (AB-buffer). The incubated NC filters were washed twice with TTBS and then incubated again for 2 h in peroxidase-conjugated swine anti-rabbit IgG (Dako A/S) diluted 1:500 in AB-buffer. They were then washed in 2 changes of TTBS and 1 change of TBS. Finally, the reaction was developed with 2,3-diamino benzidine-tetrahydrochloride (Sigma, St. Louis, MO, USA) in 50 mM Tris-HCl, pH 7.6 with 0.5% H2O2.

Experimental fish. Baltic salmon (2-yr-old) at an average weight of 32 g were successively acclimatized to a temperature of 9 ± 1°C in 250 l tanks supplied with continuously flowing well water and were fed commercial trout dry food (Dansk Orredfoder A/S, ECOlife 17, Brande, Denmark). Fish originated from a local fish farm, regularly health monitored as part of the Swedish Fish Health Control Programme, and had no history of BKD. Fish were challenged by immersion in oxygenated water containing Rs bacteria (strain 15/67/92; 1 × 10⁸ colony forming units ml⁻¹). The fish were observed daily and dead fish collected. Surviving fish were killed by an overdose of MS 222 (Sandoz, Ltd, Basle, Switzerland; 0.75 ml l⁻¹) and sampled 4 and 8 wk post-infection.

Sampling procedures. A total of 733 samples came from farmed and feral fish (mainly Baltic salmon Salmo salar) and included both 1- and 2-yr-olds as well as broodstock: 304 samples originated from rainbow trout (64 of these were experimentally challenged as described above), 229 samples from Arctic char, 58 from Baltic salmon (including 45 experimentally challenged as above), and 42 from brown trout. All fish were examined for gross clinical signs of BKD, and kidney samples were taken aseptically into sterile bags for bacterial cultivation and for ELISA, mainly at the fish farms. For the comparison between cultivation, polyclonal ELISA, and the monoclonal ELISA kit, kidney samples were divided into 6 pieces and 2 of
these were randomly placed in each of 3 bags, in order to compensate for a possible nonrandom distribution of bacteria or bacterial antigens in sampled kidney tissue.

**Bacterial culture.** The kidney samples were mechanically homogenized (Stomacher Lab Blender 80, Seward Laboratory, London, UK) in peptone-saline, 1 part kidney, 3 parts PBS, and 25 μl HemoDe per ml homogenate in a stomacher. The samples were heated 1 part kidney, 3 parts PBS, and 25 μl HemoDe per ml homogenate in a stomacher, and treated as described for the bacterial samples. Homogenates were prepared with a 1:1 ratio (w/v). Ten μl of the homogenate was spread on SKDM agar, horse blood (5%) agar, and KDM-C agar. SKDM and KDM-C agar plates were incubated in plastic bags at 15°C and examined weekly. SKDM plates were incubated for up to 12 wk. The long incubation time for SKDM was necessary since visible colonies of Rs appeared between the 6th and 11th week in several low-grade infected samples. Benediktsdóttir et al. (1991) found the highest prevalence of positive samples during the 6th and 9th week of incubation. In order to minimize problems with samples contaminated with bacteria or fungi and thus optimize the cultivation method, each kidney sample was spread onto the whole surface of an agar plate. Blood agar plates were incubated at 20°C and examined after 4 d. Pure culture of isolated bacterial fish pathogens were identified by conventional methods including biochemical and serological tests. Growth of Rs was confirmed by Gram staining and immunofluorescence (monoclonal antibody against Rs, Micrologix International Limited, Sidney, BC, Canada) on typical colonies from SKDM agar. Strains of Rs were further verified by catalase and oxidase reactions and finally with API-zym (bioMérieux, Marcy-l’Étoile, France; Austin & Austin 1987). Samples were processed immediately at the laboratory, with the exception of 52 samples (from brown trout, rainbow trout, and Baltic salmon) which were kept frozen at −20°C for up to 1 mo prior to cultivation.

**Sample preparation for polyclonal ELISA.** Samples for the polyclonal ELISA were prepared according to the method of Gudmundsdóttir et al. (1993).

**Bacterial samples (Table 1):** The bacteria were diluted to an optical density (OD) of 1.0 at 660 nm in PBS, and 25 μl of the solvent HemoDe (terpene and butylated hydroxy anisole, Fisher Scientific, Pittsburgh, PA, USA) was added for each mL of suspension. The samples were carefully mixed and heated to 98 ± 2°C in an autoclave for 15 min. After centrifugation at 2500 g for 20 min at 4°C, the aqueous phase was collected and used in the ELISA.

**Kidney samples:** Homogenates were prepared with 1 part kidney, 3 parts PBS, and 25 μl HemoDe per ml homogenate in a stomacher. The samples were heated and treated as described for the bacterial samples. Kidney samples were analysed directly by the ELISA or were frozen at −20°C for later assay.

**Polyclonal ELISA.**

**Coating and blocking:** Microtiter plates (Nunc 4-42404, Roskilde, Denmark) were coated overnight at 4°C with rabbit IgG (10 μg ml⁻¹) prepared against Rs iscoms diluted in 0.05 M carbonate buffer, pH 9.6 (200 μl well⁻¹). Initially, plates were also coated with rabbit IgG prepared against Rs hemagglutinin (HA) and with normal rabbit Ig at the same concentration and in the same manner as described above. Afterwards, 4 washes were carried out with saline (0.9% NaCl; 0.001 M PBS) containing 0.5% Tween 20. All wells were blocked with 1% bovine serum albumin (Sigma a-4503, St. Louis, MO, USA; 200 μl well⁻¹) in distilled water for at least 30 min at room temperature (RT) before the samples were added.

**Samples:** A volume of 50 μl PBS containing 0.5% Tween 20 (PBS-Tween) was added to all wells, followed by addition of 50 μl of supernatant from each prepared sample to each of 2 wells. A negative control prepared from a pool of kidney samples from 20 Rs-negative rainbow trout (checked by bacterial cultivation) was tested in 4 wells on every microtiter plate. Dilutions of a HemoDe-treated Rs suspension were tested as positive controls (2 antigen concentrations, PC 1 and PC 2). PBS-Tween was added to the first row of every plate and the resulting background OD value was subtracted from all sample values. The microtiter plates were incubated at RT for 2 h.

**Conjugate:** Affinity-purified, peroxidase-labelled goat anti-Renibacterium salmoninarum (Kirkegaard & Perry Laboratories Inc., Gaithersburg, MD, USA) diluted in PBS-Tween to a concentration of 0.2 μg ml⁻¹ was added to all wells (100 μl well⁻¹) and incubated at RT for 1 h.

**Substrate:** The enzyme reaction was visualized using tetramethyld benzidine (Kebo, Stockholm, Sweden) at a concentration of 0.1 mg ml⁻¹ in 0.1 M acetate buffer, pH 6.0, containing H₂O₂ (0.006%). The reaction was stopped after 10 min with 1 M H₂SO₄ and the OD was measured by spectrophotometer at 450 nm.

**Negative-positive threshold:** The negative-positive threshold of the ELISA was defined as the mean OD values plus 3 standard deviations of samples from clinically healthy fish, tested by bacterial culture to be negative.

**Control of false positive reactions:** All samples exceeding the negative-positive threshold value in the ELISA were checked for false positive reactions by repeating the ELISA. Microtiter plates were coated with the same rabbit IgG prepared against Rs iscoms but also with normal rabbit IgG (10 μg ml⁻¹, Dako A/S, Glostrup, Denmark), 1 well of each for every sample. The ELISA was subsequently performed as previously described.
**Interassay correction:** Interassay variations in the ELISA were corrected by a method modified from Pascho et al. (1987): All values were multiplied by a correction factor, 1 - A, for each microtiter plate tested. A is defined according to the following formula:

\[ A = \frac{1}{2} \left[ \frac{PC_{1}}{PC_{1} + PC_{2}} \right] \]

where PC1 is the absorbance values of positive control 1 (the mean value of duplicates) at each plate; PC2 is the absorbance values of positive control 2 (the mean value of duplicates) at each plate; PC1v is the mean absorbance values of positive control 1 in all plates tested; PC2v is the mean absorbance values of positive control 2 in all plates tested.

**K-Dtect ELISA.** A commercially available monoclonal ELISA kit marketed as K-Dtect (DiaXotics, Wilton, CT, USA) was used in accordance with the manufacturer’s instructions. Briefly, kidneys were diluted in sample dilution buffer (SDB; 1 g kidney: 1 ml SDB), homogenized, and centrifuged. The supernatants were added to duplicate wells of a microtiter plate that had been coated with monoclonal antibodies against a soluble 57 kDa antigen of Rs (p57). The plate with samples, positive controls with varying concentrations of p57 antigen, and negative controls, was incubated for 1 h at RT before the plate was washed to remove unbound material. A biotinylated monoclonal antibody directed to a different epitope of p57 antigen was then added to the wells for a 1 h incubation at RT. After washing, streptavidin linked to horseradish peroxidase was added for incubation for 30 min. Finally, the plate was washed, the chromogen ABTS and H2O2 were added, and the OD was measured with a spectrophotometer. The washing solution and SDB were diluted prior to use. All other working solutions of reagents were included in the test kit.

**RESULTS**

**Specificity of rabbit IgG**

The iscom-induced rabbit anti-\textit{Renibacterium salmoninarum} IgG was superior as capture antibodies in the ELISA when compared with the HA-induced antibodies (Table 2). Differences in OD values, using the 2 antibody preparations for capture in samples collected from experimentally challenged rainbow trout and salmon, were found to be significant by the Mann-Whitney Wilcoxon test (p = 0.01).

No cross reactivities of the iscom-induced rabbit IgG with other fish-pathogenic bacterial strains were found using the polyclonal ELISA. All strains of Rs (13 different isolates) gave strong extinction values. Among the non-fish pathogens tested, a positive OD value was obtained with a bacterial preparation of \textit{Arthrobacter globiformis} (Table 1). In Western blot, major immunoreactive bands of the Rs antigen preparation were detected with the rabbit anti-\textit{Renibacterium salmoninarum} iscom IgG near 57 kDa, as well as 2 immunoreactive bands of lower molecular mass. An identical profile was obtained with the polyclonal antibodies produced after immunization with the water-extracted HA of the bacterium (Fig. 1).

**Polyclonal ELISA standardization**

Based on the OD values of kidney samples (n = 408) from rainbow trout, Arctic char, Baltic salmon, and brown trout collected from fish negative for Rs by bacterial culture, the negative-positive threshold value in the polyclonal ELISA was determined to be 0.10. All
kidney samples originated from fish farms or fish populations considered to be free from Rs infection. Initially, 3 dilutions (i.e. undiluted homogenate, 1:2, and 1:4) of kidney samples were tested, both with the rabbit anti-Renibacterium salmoninarum IgG and with normal rabbit IgG, to determine optimum background OD levels. Acceptably low background OD values were observed at the 1:2 dilution of the prepared sample. The use of 4 washings between every incubation step in the ELISA instead of 3 washings, as was used initially, further lowered the background OD value. In these samples, when only 3 washings were used, the negative-positive threshold value was set at 0.14 (74 samples collected from farms with clinical BKD and 77 samples from populations considered to be free of Rs). A Tween-20 content of 0.5% was used throughout in this study. Although the ELISA procedures were standardized, day-to-day variations occurred, especially in samples showing high OD values. To compensate for this variation, all values (except values from the experimentally challenged fish) were corrected by the modified Pascho et al. (1987) statistic. The mean of the negative control samples was reduced from 0.019 to 0.015 by this correction, but the status of positive or negative values remained unchanged. Observed distributions of OD values from samples classified as negative by bacterial culture are summarized in

![Graph showing distribution of polyclonal ELISA optical density (OD) values from samples classified as negative by bacterial culture](image)

Fig. 2. Observed distribution of polyclonal ELISA optical density (OD) values from rainbow trout Oncorhynchus mykiss, Arctic char Salvelinus alpinus, Baltic salmon Salmo salar, and brown trout Salmo trutta, which were diagnosed as negative by cultivation (no live Rs were detected in the samples; n = 408)
of the results in Table 3a, the specificity of the polyclonal ELISA method was set at 99.7%. According to Table 3b, the sensitivity was calculated to be 95.8% when compared with cultivation. However, several samples only showed positive by ELISA. ELISA values of samples with an overgrowth of contaminating microbes on SKDM were excluded in the calculations.

**Comparison of SKDM, polyclonal ELISA, and K-Dtect ELISA**

Cultivation of Rs on SKDM was compared with both the ELISA method described here and a commercially available ELISA method for samples from 175 fishes (Table 4). It was not possible to interpret 8 samples, with identical classification by the 2 ELISA methods in cultivation because of contamination of the SKDM agar. An agreement between all 3 methods occurred in 159 of the remaining samples (41 positive and 118 negative). Four kidneys, negative in K-Dtect, were positive with both cultivation and the polyclonal ELISA, and an additional 2 kidneys were positive only with the polyclonal ELISA. Rs was not cultivable from 1 sample showing positive according to both the ELISA methods whereas one sample was positive only by cultivation.

**DISCUSSION**

Previous studies of immunostimulating complexes (iscoms), especially those containing antigens from viruses, have demonstrated the advantage of this antigen preparation over more traditional preparations (Morein et al. 1984, Lövgren et al. 1987). The ability of the iscoms to stimulate immunogenicity depends

<table>
<thead>
<tr>
<th>Cultivation (SKDM)</th>
<th>Polyclonal ELISA</th>
<th>K-Dtect</th>
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<tr>
<td></td>
<td>Pos</td>
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<tr>
<td>129 Neg</td>
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<td>8 Contaminated*</td>
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* Cultivation results not readable due to overgrowth by bacteria or fungi

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**Table 4:** Detection of *Renibacterium salmoninarum* in salmonid kidney samples (n = 175) by cultivation on selective agar (SKDM) and by 2 ELISA methods (polyclonal ELISA and K-Dtect)
mainly on a favorable presentation of the antigens, but is also dependent on the iscoms' function as an adjuvant (Dalsgaard 1978). Iscoms consist of Quil A, a glycoside which forms micelles which build complexes with cholesterol, into which antigens are incorporated by hydrophobic interactions. Several authors have described the hydrophobic nature of the cell surface of Rs (Daly & Stevenson 1987, Bruno 1988), and a capsule structure was recently demonstrated (Dubreuil et al. 1990) which may contain valuable specific diagnostic antigen(s).

In the present study, we used solubilized antigens extracted by a mild detergent for incorporation into iscom structures visible in electron micrographs (data not shown). Rabbits that were immunized with this immunogen produced antibodies with a similar profile in Western blot to those produced towards water-extracted HA antigens. The iscom-generated antibodies reacted primarily with the antigen F (Getchell et al. 1985), which in earlier studies has been described as a heat-stable antigen of 57 kDa, but also as having 2 lower molecular mass antigens that may be degradation products of antigen F (Griffiths & Lynch 1991). Antibodies produced with the iscom immunogen reacted with 13 geographically diverse isolates of Rs and were chosen for the ELISA because they were found in initial tests to be more effective as capture antibodies than the HA-induced antibodies. No cross reactivity against other fish-pathogenic bacteria has been detected. Cross reaction was detected against Arthrobacter globiformis, a bacterium earlier described as related to Rs (Stackebrandt et al. 1986). This bacterium is easily cultivated on blood agar, but has not been reported to have been isolated from fish tissues.

On the basis of analyses of 408 samples from rainbow trout, Arctic char, Baltic salmon, and brown trout classified as negative for BKD, the negative-positive threshold value in the present ELISA was determined to be 0.10. In accordance with earlier studies, no species-specific differences in OD values were observed among the negative samples (Gudmundsdottir et al. 1993, Meyers et al. 1993). One sample did exceed the threshold value in this group, but no growth of bacteria was detected when the kidney sample was cultivated on blood agar or KDM-C. Therefore, it seems unlikely that other bacteria were cross-reacting. One single fish sampled in the same population on the same occasion showed clinical indications of bacterial infection (ascitic fluid in the peritoneum, petechial hemorrhages of the heart, and pale liver). However, it was not possible to verify BKD by the immunofluorescent antibody technique or cultivation. The negative result of cultivation from the ELISA-positive fish might be explained by a higher sensitivity of the ELISA method, by an unequal distribution of bacteria, or the presence of bacterial antigens alone in the sampled kidney tissue.

The sensitivity of the present ELISA method was compared with that for cultivation on SKDM of kidney samples collected from fish farms classified as Rs-infected or with fish showing clinical indications of BKD as well as from fish experimentally challenged with Rs by immersion. The Rs organism was isolated on SKDM from 68 out of 69 fish with macroscopic signs of BKD (swollen, greyish kidney). All of these samples were also classified as positive by the ELISA. Among samples collected from fish with weak or no clinical indications of BKD, 44 samples were shown positive by both methods. A discrepancy between the 2 methods was seen in 22 samples (Table 3b), which might again be caused by an unequal distribution of low numbers of bacteria or the presence of bacterial antigens alone in the kidneys. Twelve more cases were positive according to the ELISA as compared with the SKDM performed under optimal conditions (contaminated agar plates excluded). Therefore it can be concluded that the present ELISA method is comparable with or has even greater sensitivity than cultivation, which is in accordance with earlier studies (Pascho et al. 1987, Griffiths et al. 1991, Gudmundsdottir et al. 1993). From a practical point of view, the ELISA method also appears more reliable than culture as the latter is susceptible to contamination problems when sampling is carried out in the field.

Results produced by all 3 methods were identical in 95% for 167 kidney samples tested by cultivation on SKDM, polyclonal ELISA, and K-Dtect. Six samples from a fish farm having enzootic BKD were negative by the K-Dtect but were positive by polyclonal ELISA. Four of these samples were also positive on SKDM. These data indicated a higher sensitivity of the polyclonal ELISA method when compared with the K-Dtect. This is not surprising, because several different epitopes for capture are possible using polyclonal antibodies that are not recognized by monoclonals. The existence of antigenic heterogeneity of Rs surface proteins has previously been indicated by immunogenic analysis (Bandin et al. 1992). The p57 antigen is released into fish tissues during infection (Túraga et al. 1987, Rockey et al. 1991), which may increase the accessibility of antigens for the capture antibodies. This may explain the higher sensitivity reported with the use of ELISA than with the use of cultivation on agar medium. The detection limit of polyclonal ELISA determined from serial dilutions of a washed broth culture of Rs added to homogenized kidney material, was $9 \times 10^5$ bacteria g$^{-1}$ of kidney, in accordance with the results of Bandin et al. (1996). In naturally infected fish, however, kidneys estimated to contain $10^3$ Rs g$^{-1}$ of kidney were found positive by polyclonal ELISA (data
not shown). Testing of more Rs-negative samples will probably lower our threshold level because higher background OD values obtained in the initial part of this study were included in the negative-positive threshold value of 0.10. A consequence of a reduction of this value would, of course, be a lower detection limit.

In conclusion, we found polyclonal ELISA, using the iscom-induced rabbit anti-Renibacterium salmoninarum IgG for capture, to be useful for routine screening for Rs in fish populations in the Swedish Fish Health Control Programme. The sensitivity of the method was comparable with and possibly greater than that of cultivation under optimal conditions. Further, ELISA was more convenient for the screening of wild brood fish.

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