A gene from *Renibacterium salmoninarum* encoding a product which shows homology to bacterial zinc-metalloproteases

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A genomic library constructed from *Renibacterium salmoninarum* isolate MT444 DNA in the plasmid vector pBR328 was screened using *Escherichia coli* host strain DH1 for the expression of genes encoding putative virulence factors. A single haemolytic clone was isolated at 22 °C and found to contain a 3.1 kb HindIII fragment of inserted DNA. This fragment was present in seven isolates of *R. salmoninarum* which were examined. Western blots of extracts from clones exhibiting haemolytic activity were performed with antisera raised against either cellular or extracellular components of *R. salmoninarum* and failed to identify any additional proteins compared to control *E. coli* containing pBR328. However, minicell analysis revealed that a polypeptide with an apparent molecular mass of 65 kDa was associated with a haemolytic activity distinct from that previously described for *R. salmoninarum*. The nucleotide sequence of the gene encoding this product was determined and the amino acid sequence deduced. The product was 548 amino acids with a predicted molecular mass of 66,757 Da and a pI of 5.57. The deduced amino acid sequence of the gene possessed strong similarities to those of a range of secreted bacterial zinc-metalloproteases and was tentatively designated hly. Neither protease nor lecithinase activities were detectable in *E. coli* recombinants expressing gene hly. Haemolytic activity was observed from 6 °C to 37 °C for erythrocytes from a number of mammalian species and also from fish. Gene hly was expressed in *E. coli* as a fusion protein consisting of maltose-binding protein at the N-terminus linked to all but the first 24 amino acids, largely constituting the putative signal peptide, of the N-terminus of Hly. The soluble fusion protein was produced and purified by affinity chromatography. Antiserum raised against the purified fusion protein was used to probe Western blots of cell lysates and extracellular products from seven isolates of *R. salmoninarum* cultured under conditions of iron-sufficiency or iron-restriction. The results indicate that the availability of iron modulates the expression of the hly gene.

Keywords: *Renibacterium salmoninarum*, bacterial kidney disease, BKD, metalloprotease, gene expression

**INTRODUCTION**

*Renibacterium salmoninarum* is the causative agent of a chronic, granulomatous infection known as bacterial kidney disease (BKD) which affects farmed and wild salmonid fish worldwide (Evelyn, 1993). BKD is a particularly difficult disease to treat; there is as yet no effective vaccine, antibiotics are often ineffective and the bacterium is transmitted vertically within the salmonid egg. *R. salmoninarum* is a Gram-positive, slow-growing, fastidious, intracellular pathogen which is difficult to culture *in vitro* and this may be the major reason why little is known about the virulence or pathogenicity of the bacterium.
organism (Evenden et al., 1993). There are as yet no defined or minimal culture media available, and cultures of the bacterium may take at least 6–8 weeks to grow and are prone to contamination (Evelyn, 1993).

Many studies of *R. salmoninarum* have concentrated on the major secretory antigen, also known as antigen F (Getchell et al., 1985), haemagglutinin (Daly & Stevenson, 1990) and p57 (Wiens & Kaattari, 1991). This protein is the predominant cell surface and extracellular protein (ECP) of the bacterium and has been associated with hydrophobicity and virulence (Bruno, 1988, 1990). However, the characterization of other components which may be of importance to the progression of infection is required in order to provide an improved understanding of BKD. In particular, molecules which may be associated with the invasion of and survival within host cells, or which modulate the host immune response, such as haemolysins or toxins, have been identified as being of considerable importance to the progress of infections caused by other intracellular pathogens (Finlay & Falkow, 1989). As part of a comprehensive molecular study of *R. salmoninarum*, previous work in our laboratory has involved the cloning and expression of a haemolytic protein from the pathogen (Evenden et al., 1990).

In this study, we present the entire nucleotide sequence of a gene encoding a second haemolytic protein from *R. salmoninarum* isolate MT444 which is distinct from that previously described. As a first step in the process of understanding the role of this molecule, a fusion protein of the enzyme has been produced to provide a reliable source for further studies which will not depend upon *in vitro* cultures of the organism. This report describes the construction of the gene fusion and the production of antisera to the purified fusion protein.

**METHODS**

**Bacterial strains and plasmids.** *R. salmoninarum* isolates MT414, MT417, MT420, MT425, MT444, MT452, MT450, MT451, MT452,910019 and the type strain ATCC 33209 (Table 1) have been previously described (Grayson, 1993). *Yersinia ruckeri* strain 12/6 and *Aeromonas salmonicida* strain CM30 were obtained from Dr C. J. Rodgers, Fish Disease Laboratory, Ministry of Agriculture, Food and Fisheries, Weymouth, UK. *Escherichia coli* strains DH1 [recA1 endA1 gyrA96 thi-1 hsdR17 (rK-m-K sii-M-m') rpsL20 (ompW15 Tn10)] and XL1-Blue [recA1 endA1 gyrA96 thi-1 hsdR17 supE44 relA1 lac (F' proAB lacIqZAM15 TnlO] were used as hosts for plasmid vectors. Minicell analysis of plasmid-encoded polypeptides was performed with strain DS410 (lac rpsL minA minB). In some cases, strains Y1089 [Δ(lac)U169 proA Δ(lac) araD139 strA hflA150::Tn10(ter)] (pMC9 amp' ter') and Y1090 [Δ(lac)U169 proA Δ(lac) araD139 strA supF mcrA trpC22::Tn10(ter) (pMC9 amp' ter')] were used. The plasmid vectors pBR328, pUC18, pMAL-p and pMAL-c have been described previously (Guan et al., 1987; Maina et al., 1988). Restriction analyses and plasmid constructions were performed according to standard techniques (Maniatis et al., 1982).

**Media and reagents.** *R. salmoninarum* was cultured in Mueller–Hinton medium supplemented with 0·1% L-cysteine hydrochloride (MHCb) at 15°C and the availability of iron was restricted in cultures of *R. salmoninarum* by the addition of 200 μM ethylenediamine di-(o-hydroxyphenylacetic acid) (EDDHA) previously shown to be sub-inhibitory (Grayson, 1993). *E. coli* strains were grown in Luria–Bertani (LB) medium at either 22°C or 37°C unless otherwise stated, and 0·2% glucose was added when required. For screening clones, colonies on LB agar (1·5%, w/v) plates were overlaid with 0·75% agarose containing either 1% (w/v) casein, 0·1% (w/v) gelatin or 20% (v/v) fresh, washed erythrocytes of rabbit, sheep, horse, rat, mouse and rainbow trout, and compared with *E. coli* DH1 containing pBR328. Duplicate plates were incubated at 6°C, 15°C, 20°C, 25°C, 30°C and 37°C for up to 4 days and compared at regular intervals with control plates. Brain heart infusion broth (Difco) supplemented with thiamin (0·5 μg ml–1) was used for minicell analysis. Antibiotics were used at the following final concentrations: 100 μg ampicillin ml–1; 25 μg chloramphenicol ml–1; 15 μg tetracycline hydrochloride ml–1. Restriction endonucleases and ligase were purchased from Northumbria Biologicals and were used according to the manufacturer’s instructions. All other chemicals and reagents were purchased from either Sigma or BDH.

**SDS-PAGE and Western blotting.** SDS-PAGE was performed as described by Laemmli (1970). Electrophoretic transfer of proteins to nitrocellulose was performed as described by Towbin et al. (1979). Wistar rats were immunized subcutaneously with 100 μg maltose-binding protein (MBP)–Hly fusion protein emulsified in Freund’s complete adjuvant, and then boosted 3 weeks later with 100 μg protein emulsified in Freund’s incomplete adjuvant. The rats were bled 3 weeks after the last injection. Control sera were gathered prior to immunization. Rabbit antisera to MBP was obtained from New England Biolabs. Western blots were probed with primary antibodies and specific binding was detected by immunoperoxidase staining of horseradish-peroxidase-labelled swine anti-rabbit immunoglobulins or rabbit anti-rat IgG. Preparation of *R. salmoninarum* cell lysates and extracellular products for use in immunoblot analyses has been described previously (Grayson, 1993).

**Minicell analysis.** Minicell analysis was performed as described by Dougan & Kehoe (1984). Plasmid-encoded polypeptides were specifically radiolabelled by adding 20 μCi (740 kBq) Tran-35S-label methionine and cysteine (ICN Flow) to a suspension of purified minicells and incubating for 30 min. The labelled minicells were harvested, separated by SDS-PAGE, and their proteins electrophoretically transferred to nitrocellulose prior to staining the blots for total protein using colloidal gold (Aurodye; Janssen Biotech) and subjecting them to autoradiography using Kodak X-OMAT film.

**Southern blot hybridization.** Bacterial genomic DNA was isolated according to standard methods (Maniatis et al., 1982). Total genomic DNA was cleaved with HindIII endonuclease and the fragments produced were separated by electrophoresis at 50 V on a 0·75% agarose gel. The DNA was transferred to nitrocellulose filters (Southern, 1975) and probed for a specific complementary sequence using a DNA probe which had been labelled using a non-radioactive DNA labelling and detection kit according to the manufacturer’s instructions (Boehringer Mannheim). Filters were prehybridized at 68°C for 2 h, hybridized with 0·5 μg digoxigenin-labelled probe at 68°C for 24 h, washed and dried. The presence of the probe was detected with anti-digoxigenin phosphatase conjugated antisera and visualized by enzyme-substrate colour reaction.
Table 1. Source and origin of R. salmoninarum isolates used in this study

All isolates were provided by Dr D. W. Bruno, SOAFD Marine Laboratory, Aberdeen, Scotland, UK, except isolate 910019, which was supplied by Dr C. J. Rodgers, MAFF Fish Disease Laboratory, Weymouth, England, UK.

<table>
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<th>Origin</th>
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<td>MT 241</td>
<td>-</td>
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Fig. 1. Restriction map of pTHG1 DNA deduced from single enzyme digests. The 3.1 kb insert of R. salmoninarum DNA is represented by the heavy line. Also indicated are the Sall deletion and the PvuII deletion which were used in the minicell analysis of pTHG1 encoded proteins. ori, origin of replication; cam', gene encoding chloramphenicol resistance; amp', gene encoding ampicillin resistance; tef', gene encoding tetracycline resistance.

Nucleotide sequencing. From the restriction map of pTHG1 (Fig. 1) it was decided to subclone the following fragments for sequencing: (i) HindIII–PvuII ~ 623 bp, (ii) PvuII–PstI ~ 727 bp and (iii) PstI–PstI ~ 520 bp. The DNA sequence of the cloned R. salmoninarum DNA was determined from double-stranded plasmid templates by dideoxy-chain termination using the Sequenase Version 2.0 kit (United States Biochemical). Double-stranded templates were denatured and the sequencing reactions were carried out according to the manufacturer's instructions. Using pUC18 primers, sequencing reactions were prepared from subclones consisting of vector pUC18 containing fragments of R. salmoninarum DNA. Oligonucleotide forward and reverse primers were purchased from Promega; [35S]dATP [≥ 1000 Ci mmol−1 (37000 GBq mmol−1)] was purchased from Amersham. The labelled reaction mixtures were separated under denaturing conditions by electrophoresis on either 6% (w/v) or 4% (w/v) polyacrylamide gels of 55 cm in length (Macrophor 2010, LKB), the gels dried and subjected to autoradiography. Both DNA strands of each fragment were completely sequenced, with a minimum of four forward and four reverse reactions (using both dGTP and dITP) for any given region.

Nucleotide and amino acid sequence analysis. Analyses of the nucleotide and deduced amino acid sequence were performed on DNASTar software using the PROSCAN and ALIGN programs. The best 200 scores of identity were obtained and the top 30 of these, as well as a variety of other selections, were further compared by pairwise and multiple amino acid sequence alignments with the translated R. salmoninarum protein sequence. Multiple sequence alignments and comparisons were carried out with the Clustal algorithm within the MEGALIGN program of DNASTar. The percentage identity of matches was scored using a gap penalty of 3, deletion penalty of 6 and a standard probability of acceptable mutation. The prediction of α-helix, β-sheet, turn and coil regions (Chou–Fasman, Garnier–Robson algorithms), hydrophilicity (Hopp–Woods algorithm), Eisenberg correlation of amphipathic structure with secondary structure, Karplus–Schulz chain flexibility, surface probability profile (Emini algorithm), and Jameson–Wolf antigenic index were also calculated.

Identification and purification of the fusion protein. A 1.7 kbp Ebol fragment was excised from the bly gene and ligated in-frame to the 3' end site of plasmids pMAL-c and pMAL-p.
Fusion proteins were purified according to the instructions provided by New England Biolabs. Briefly, cultures of *E. coli* XL1-Blue containing the fusion plasmid were grown at either 22 °C or 37 °C to an OD$_{550}$ of 0.5. A sample of cells was removed; to the remaining culture 2 mM IPTG and samples was added to induce expression and samples removed after 1, 2, 3 and 4 h. All of the following steps were carried out either on ice or at 4 °C to minimize protein degradation. For affinity purification, cultures were induced for 2 h prior to harvesting the cells. These were resuspended in 20 mM Tris/HCl, 200 mM NaCl, 1 mM EDTA, 1 mM NaN$_3$, 1 mM PMSF, pH 7.4. The sample was frozen overnight at −20 °C, thawed in cold water, sonicated for 3 min (Ultrasonic Processor; Heat Systems Ultrasonics), centrifuged and the supernatant (crude extract) removed and retained. The pellets were resuspended in the above buffer and resuspended in an amylose resin column. The diluted extract containing the fusion protein, stored on ice, was added to the column and then the column washed free of all protein other than the specifically bound MBP fusion protein. The fusion protein was eluted with SDS-PAGE buffer and electro-phoresed by SDS-PAGE. Gels were stained for protein or transferred to nitrocellulose. Compared to minicells containing pBR328 alone, minicells containing pTHG1 possessed only the 65 kDa protein indicating that the 46 kDa molecule was transcribed from the SalI end of the 3.1 kb HindIII DNA insert in pTHG1. Minicells containing pTHG201 produced a 49 kDa protein encoded by pTHG201 probably represented a serendipitous in-frame fusion of the remainder of the gene encoding the 65 kDa protein with the phenotype proved to be too unstable for any further meaningful manipulation. The haemolytic activity of clone H3 was observed from 6 °C to 37 °C for erythrocytes from all species except rabbit, although only weak haemolysis was recorded against sheep erythrocytes. It was also observed that clones grown at 37 °C were unstable and rapidly lost their haemolytic phenotype and consequently clones were routinely cultured at 22 °C. Clone H3 displayed no evidence of degradation of either casein, gelatin or L-α-phosphatidylcholine when compared with identically treated controls. *E. coli* strains Y1089 and Y1090, both of which possess mutations preventing the production of the Lon protease, were transformed with either pTHG1 or pBR328; stable clones were isolated at 22 °C and assayed for haemolytic activity as described above. The absence of the Lon protease did not affect the haemolytic phenotype encoded by pTHG1.

**RESULTS**

### Cloning of a haemolytic determinant from *R. salmoninarum*

Screening of the pBR328 HindIII gene bank in *E. coli* DH1 resulted in the identification of a single clone, designated H3, which possessed a haemolytic phenotype. Clone H3 was found to contain plasmid DNA with a single 3.1 kb HindIII insert. The recombinant plasmid was designated pTHG1. Fig. 1 shows a restriction map of this plasmid. In an attempt to increase the yield of recombinant protein the 3.1 kb HindIII fragment was also subcloned in the plasmid vector pUC18 but the phenotype proved to be too unstable for any further meaningful manipulation. The haemolytic activity of clone H3 was observed from 6 °C to 37 °C for erythrocytes from all species except rabbit, although only weak haemolysis was recorded against sheep erythrocytes. It was also observed that clones grown at 37 °C were unstable and rapidly lost their haemolytic phenotype and consequently clones were routinely cultured at 22 °C. Clone H3 displayed no evidence of degradation of either casein, gelatin or L-α-phosphatidylcholine when compared with identically treated controls. *E. coli* strains Y1089 and Y1090, both of which possess mutations preventing the production of the Lon protease, were transformed with either pTHG1 or pBR328; stable clones were isolated at 22 °C and assayed for haemolytic activity as described above. The absence of the Lon protease did not affect the haemolytic phenotype encoded by pTHG1.

### Detection and identification of the haemolytic product encoded by pTHG1

Cells and extracellular products harvested from clone H3 were resuspended in SDS-PAGE buffer and electrophoresed by SDS-PAGE. Gels were stained for protein or transferred to nitrocellulose. Compared to control *E. coli* DH1 containing pBR328 alone, no additional proteins could be detected on either SDS-PAGE gels or Western blots probed with rabbit antisera raised against *R. salmoninarum* ATCC 33209 whole cells, ECPs or cell walls. In addition, no haemolytic activity was associated with cell extracts or ECPs of clone H3. In order to simplify the identification of the haemolytic protein encoded by pTHG1, the plasmid was cleaved with either SalI alone or SalI and PvuII and then religated. Clones containing a SalI deletion of pTHG1 (designated pTHG101) were then confirmed for sensitivity to tetracycline and found to be haemolytic when compared with controls *E. coli* DH1 containing pBR328. In contrast, clones containing both the SalI and PvuII deletions of pTHG1 (designated pTHG201) which were confirmed as sensitive to chloramphenicol were non-haemolytic using the same assay. Minicell analysis of the products encoded by each of the plasmid derivatives was performed in order to identify the haemolytic component. The autoradiograph of minicells which had been separated by SDS-PAGE and transferred onto nitrocellulose showed only plasmid-encoded proteins including those conferring antibiotic resistance (Fig. 2). Compared to minicells containing pBR328 alone, minicells containing pTHG1 possessed at least four additional proteins of molecular masses 65, 46, 43 and 36 kDa. Minicells containing pTHG201 possessed only the 65 kDa protein indicating that the 46 kDa molecule was transcribed from the SalI end of the 3.1 kb HindIII DNA insert in pTHG1. Minicells containing pTHG201 produced a 49 kDa protein. On the basis of these results it was decided that haemolytic activity was conferred by the 65 kDa protein and that the gene encoding this protein was located at the PvuII end of the 3.1 kb HindIII DNA insert. pTHG201 contained at most 1.3 kbp of insert DNA which would be available for transcription, which would be insufficient for the translation of a 49 kDa truncated protein. The 49 kDa protein encoded by pTHG201 probably represented a serendipitous in-frame fusion of the remainder of the gene encoding the 65 kDa protein with the remainder of the cam gene and under the control of the cam promoter. The PvuII site of the cam gene is located 114 bp downstream of the initiation codon. This would be sufficient to code for 38 amino acids of about 4 kDa molecular weight with the additional 45 kDa being encoded by insert DNA. Sequencing of the *R. salmoninarum* DNA substantiated these initial findings.

The cloned 3.1 kb HindIII fragment is present in different isolates of *R. salmoninarum*

Southern blot hybridization was carried out by digesting with HindIII, chromosomal DNA from a number of isolates of *R. salmoninarum* including the type strain (ATCC 33209) as well as strains of *A. salmonicida*.
R. salmoninarum haemolysin

5.0 -

kDa

3.0 -

kDa

2.0 -

kDa

1.0 -

kDa

0.0 -

kDa

Fig. 2. Analysis of minicells containing either pBR328, pTHG1, pTHG101 or pTHG201. Approximately 50 µg protein from each sample was separated by SDS-PAGE using a 10% polyacrylamide gel, transferred onto nitrocellulose and autoradiographed to reveal plasmid-encoded polypeptides labelled with [35S]cysteine and [35S]methionine. Tracks: 1, pTHG201; 2, pTHG101; 3, pTHG1; 4, pBR328. The positions of molecular mass standards are indicated in kDa. The arrow indicates the position of the 65 kDa protein associated with haemolytic activity. Antibiotic resistance proteins are present at approximately 26 kDa (chloramphenicol acetyltransferase), 29 and 32 kDa (β-lactamase) and 36 kDa (tetracycline resistance).

Fig. 3. Southern blot of R. salmoninarum, A. salmonicida and Y. ruckeri DNA (5 µg) digested with HindIII. The DNA was probed with a digoxigenin-labelled 3.1 kb HindIII fragment of R. salmoninarum DNA. Lanes: 1, 0.05 µg probe; 2, R. salmoninarum strain ATCC 33209; 3, R. salmoninarum isolate MT414; 4, R. salmoninarum isolate MT417; 5, R. salmoninarum isolate MT420; 6, R. salmoninarum isolate MT444; 7, R. salmoninarum isolate MT452; 8, A. salmonicida strain CM30; 9, Y. ruckeri strain 12/6; 10, R. salmoninarum isolate MT425. The positions of DNA molecular size markers are indicated in kb.

Sequence of the gene encoding the haemolytic determinant

Plasmid pTHG1 includes a 3.1 kb R. salmoninarum MT444 chromosomal DNA insert which contains the structural gene that encodes a haemolytic product. This 3.1 kb fragment was characterized by restriction enzyme analysis (Fig. 1) and subcloned into the plasmid vector pUC18 as a series of three contiguous fragments. The sequencing templates, which were designated pURS1, pURS2 and pURS3, constituted 1.87 kb of pTHG1 insert DNA. Haemolytic activity was not detected in E. coli XL1-Blue transformants containing either pURS1, pURS2 or pURS3. A complete restriction map of the DNA sequence showed that the predicted restriction sites matched the observed endonuclease cleavage pattern. The correct orientation of the contiguous DNA fragments which formed the sequencing templates was confirmed by digesting pTHG1 with HincII to release a 980 bp DNA fragment. The nucleotide sequence was found to contain a possible open reading frame (ORF) of 1647 bp starting at nucleotide 132 and ending at nucleotide 1778 with a G+C ratio of 55.5%. Following translation, this putative ORF would code for a polypeptide of 548 amino acid residues with a calculated molecular mass of 66757 Da and a pI of 5.57 (Fig. 4). The ATG triplet encountered at base pair 132 is preceded by a putative ribosome-binding site (AGGAA) located seven nucleotides upstream from this site. Possible -10 and -35 promoter regions are indicated (Fig. 4). Confirmation of the computer predictions of promoter region, transcriptional start site and initiation codon awaits corroborating transcriptional data. There were four sequences of imperfect dyad symmetry which were found to extend from nucleotides 10 to 80, 47 to 155, 189 to 298 and 1793 to 1830. The last sequence, which is underlined in Fig. 4, resembles a rho-independent termination sequence being rich in C and G and also T bases at one end (Platt, 1986).
Fig. 4. For legend see opposite page.
Polypeptide sequence identity search of the PIR database using the proscan option of the DNAStar program revealed that the deduced protein sequence possessed strong similarities to a range of secreted bacterial zinc-metalloproteases, which are sometimes known as neutral proteases. All of these enzymes are members of a diverse family of zinc metalloendopeptidases which are recognized by a unique signature, His-Glu-X-X-His, and are sometimes known as neutral proteases. The similarities were substantial and highly conserved with respect to the residues forming the catalytic site of the proteases. *Identical residues; - conservative amino acid substitutions. Rsal, R. salmoninarum; Bst, B. stearothermophilus; Bcal, B. caldolyticus; Bmeg, B. megaterium; Bthe, B. thermoproteolyticus; Bcer, B. cereus; Bpol, B. polymyxa; Bs, B. subtilis; Bamy, B. amyloliquefaciens; Lmo, L. monocytogenes; Paer, P. aeruginosa; Vchol, V. cholerae; Vprot, V. proteolyticus; Lpneu, L. pneumophila; Ec, E. carotovora.

A polypeptide sequence alignment of metalloproteases of Bacillus, Erwinia, Legionella, Listeria, Pseudomonas and Vibrio from the PIR database and Renibacterium metallohaemolysin and showing the highly conserved residues which form the catalytic centre of the enzymes. *Identical residues; - conservative amino acid substitutions. Rsal, R. salmoninarum; Bst, B. stearothermophilus; Bcal, B. caldolyticus; Bmeg, B. megaterium; Bthe, B. thermoproteolyticus; Bcer, B. cereus; Bpol, B. polymyxa; Bs, B. subtilis; Bamy, B. amyloliquefaciens; Lmo, L. monocytogenes; Paer, P. aeruginosa; Vchol, V. cholerae; Vprot, V. proteolyticus; Lpneu, L. pneumophila; Ec, E. carotovora.

Fig. 5. Amino acid sequence alignment of metalloproteases of Bacillus, Erwinia, Legionella, Listeria, Pseudomonas and Vibrio from the PIR database and Renibacterium metallohaemolysin and showing the highly conserved residues which form the catalytic centre of the enzymes. *Identical residues; - conservative amino acid substitutions. Rsal, R. salmoninarum; Bst, B. stearothermophilus; Bcal, B. caldolyticus; Bmeg, B. megaterium; Bthe, B. thermoproteolyticus; Bcer, B. cereus; Bpol, B. polymyxa; Bs, B. subtilis; Bamy, B. amyloliquefaciens; Lmo, L. monocytogenes; Paer, P. aeruginosa; Vchol, V. cholerae; Vprot, V. proteolyticus; Lpneu, L. pneumophila; Ec, E. carotovora.
with the \textit{malE} gene encoding MBP. A blunt-ended \textit{EheI} site located at nucleotide 203 (Fig. 6) provided an in-frame fusion when ligated to the \textit{Stul} site of the pMAL poly linker. This manipulation was expected to yield 524 amino acids of the \textit{R. salmoninarum} protein with a molecular mass of 63842 Da fused to MBP. A recombinant plasmid, designated pTHG203 (Fig. 6), containing an in-frame fusion between \textit{hly} and \textit{malE} was constructed and expressed in \textit{E. coli} XL1-Blue. \textit{E. coli} clones producing the fusion protein (MBP–Hly) were identified by Western blotting using rabbit anti-MBP antiserum. MBP–Hly was produced cytoplasmically in \textit{E. coli} using pMAL-c only, because of the difficulty in isolating clones expressing the fusion in the pMAL-p vector. Hly was fused to MBP starting from Ala-25, which is four amino acids upstream of the predicted signal peptide cleavage point at Ala-28. Construction of recombinant plasmid pTHG203 allowed the production of a fusion protein composed of the entire MBP molecule (molecular mass 42.7 kDa) fused at the C-terminus, with a short factor Xa cleavage site, to 524 amino acids (molecular mass 63842 Da) derived from \textit{R. salmoninarum} haemolysin (Hly). The small-scale pilot experiment showed after 2–4 h induction that MBP–Hly was produced at a high level and mainly in a soluble form (Fig. 7). Because considerable breakdown and instability was observed at 37 °C, IPTG induction was carried out at 22 °C. Inducing at 22 °C in conjunction with a second sonication of the insoluble cell material greatly improved the stability and yield of MBP–Hly. The MBP–Hly protein was purified from the cytoplasm of \textit{E. coli} containing recombinant plasmid pTHG203 by using a one-step affinity chromatography procedure as described in Methods. Purified MBP–Hly migrated as a single band.
on SDS-PAGE with an apparent molecular mass of 105 kDa, which was similar to the predicted molecular mass of 106542 Da. Protein concentrations were estimated by the method of Bradford (1976) and yields of 50 μg (ml culture)^{-1} were obtained.

**Expression of hly is modulated by the availability of iron**

Western blots of MBP-Hly which were probed with antisera from rats immunized with purified MBP-Hly showed good recognition of the fusion protein (Fig. 7). Cell extracts or ECPs were obtained from seven isolates of *R. salmoninarum* which had been cultured under either iron-restricted or iron-sufficient conditions, separated by SDS-PAGE and blotted onto nitrocellulose. Western blots of sonicated *R. salmoninarum* cell extracts derived from iron-sufficient cultures which were probed with rat anti-MBP-Hly revealed the presence of strongly immuno-reactive bands of approximately 70 and 25 kDa with a weakly stained 48 kDa band in all isolates (Fig. 8a). However, these bands were barely discernible on Western blots of cell extracts from iron-restricted cultures (Fig. 8b). No bands were detected by the MBP-Hly antiserum on Western blots of ECPs from either iron-restricted or iron-sufficient cultures.

**DISCUSSION**

This study has described a novel haemolysin from *R. salmoninarum*. Gene *hly* has been found to be present in each of the seven isolates of *R. salmoninarum* so far examined and the nucleotide content was 55.5% G+C, consistent with the overall 55.5% G+C of the *R. salmoninarum* chromosome (Banner et al., 1991). Gene *hly* encodes a product which confers a haemolytic phenotype on *E. coli*. The active component, as identified by minicell analysis, possessed an apparent molecular mass of 65 kDa and was active against erythrocytes from a number of animal species, including rainbow trout but not rabbit, from 6 °C to 37 °C. These features are reminiscent of those of a membrane-active toxin. The results of other researchers investigating the immunological and enzymic properties of ECPs or whole cells from *in vitro* cultures of *R. salmoninarum* have failed to identify any haemolytic or cytotoxic components although proteolytic activities have been demonstrated (Bandín et al., 1991; Griffiths & Lynch, 1991; Rockey et al., 1991). The possible contribution of an unidentified toxic component to the pathology of the disease has been suggested (Bruno & Munro, 1986; Turaga et al., 1987). It is notable that Bruno & Munro (1986) provide the only report of β-haemolysis by *R. salmoninarum* isolates cultured *in vitro* and this activity was recorded against rabbit erythrocytes. In addition, Bruno & Munro (1986) found that β-haemolysis declined following routine subculture of *R. salmoninarum* with an accompanying decline in the virulence of the organism. Evenden et al. (1990) reported the cloning of a gene from *R. salmoninarum* encoding a product which was capable of lysing rabbit erythrocytes. The product of gene *hly*, on the other hand, was not found to be active against

**Fig. 8.** The detection of epitopes of MBP-Hly in cell extracts of seven *R. salmoninarum* isolates which had been cultured in either (a) MHC or (b) MHC supplemented with 200 μM EDDHA. Preparations were separated by SDS-PAGE using a 10–15% gradient gel, transferred onto nitrocellulose and then probed with a rat antiserum raised against MBP-Hly. All tracks contained an equivalent amount of protein (2 μg). Tracks: 1, isolate 910019; 2, isolate MT452; 3, isolate MT425; 4, isolate MT420; 5, isolate MT417; 6, isolate MT414; 7, strain ATCC 33209. The positions of protein molecular mass standards are marked (kDa).

rabbit erythrocytes, and furthermore did not degrade either casein or gelatin. The instability of clones in which the gene is expressed at 37 °C or when inserted into the pUC18 vector, and the difficulty in maintaining cultures of *E. coli* which contained pTHG1, either stored at -20 °C or at 4 °C, provide an indication of the toxicity of the protein to *E. coli* cells. Because of the difficulties experienced with high level expression of the protein and lack of a means of detecting the protein in cell extracts or ECPs of either *E. coli* clones or *R. salmoninarum* cultures, further biochemical characterization was considered to be impossible at this stage.

Zinc-dependent endopeptidases form a heterogeneous family, with widely differing specificities and sensitivity to
inhibitors (Jongeneel et al., 1989). It has been suggested that variation in the spacer regions between the catalytic zinc ligands and the residues forming the substrate-binding site could reflect differences in substrate specificity and the mechanism of catalysis (Vallee & Auld, 1989). A better understanding of the structural requirements for enzyme activity has been constrained by the limited availability of protein sequence and tertiary structure information (Jongeneel et al., 1989). In the absence of such information it is impossible to predict what degree of specificity the haemolysin of R. salmoninarum may exhibit. However, many of the proteases with which it shares considerable similarity have been well characterized and found to possess broad substrate specificity. The elastase of Pseudomonas aeruginosa, for example, has been shown to be active against elastin and collagen (Heck et al., 1986), human IgA and IgG (Döring et al., 1981) and several complement components (Hong & Ghebrehiwet, 1992).

The failure to detect proteolytic activity against casein and gelatin substrates suggests that the R. salmoninarum haemolysin either requires some other factor for activity or differs from many other zinc-metalloproteases and is only active against specific substrates. Mengaud et al. (1991) and Domann et al. (1991) reported similar difficulties in detecting any strong proteolytic activity in either E. coli clones expressing the Listeria monocytogenes metalloprotease or in L. monocytogenes culture medium and suggested that this may have been due to a lack of processing in vitro.

This study has demonstrated the immunological detection of epitopes of MBP–Hly in extracts derived from R. salmoninarum cultured in vitro and has therefore confirmed the validity of this approach to the study of a particularly difficult pathogen. It should be noted that R. salmoninarum does not utilize starch or maltose (Goodfellow et al., 1985) and Western blots of R. salmoninarum cultures which were probed with rabbit anti-MBP antisera showed no evidence for the recognition of cross-reactive epitopes. The availability of rat anti-MBP–Hly antisera will therefore prove useful in future research. The immunological detection of epitopes of MBP–Hly revealed a number of bands in the cell extracts of all seven isolates of R. salmoninarum which were examined. No components were detected in the ECPs. It seems likely that the various bands represent breakdown products of the 67 kDa Hly protein. However, the possibility exists that they represent cross-reacting epitopes on other proteins. Many zinc-metalloproteases are secreted as preproenzymes and the mature protease is produced following cleavage of the propeptide (Simonen & Palva, 1993). In addition, these bands were almost undetectable under iron-restricted conditions suggesting that the production of Hly by R. salmoninarum is greatly reduced when iron is no longer freely available. Whether the regulation of the production of Hly occurs at the transcriptional level or as a consequence of post-transcriptional factors such as the induction or suppression of other genes with a regulatory function is unknown. The precise contribution of metal ions such as Fe\(^{3+}\), Fe\(^{2+}\), Zn\(^{2+}\) and Ca\(^{2+}\) to the regulation of R. salmoninarum growth and the production of specific components, such as Hly, awaits the development of a defined growth medium and better elucidation of the metal ion requirements of R. salmoninarum. The lack of immunologically detectable Hly in the ECPs of R. salmoninarum cultures suggests that Hly was not secreted under the culture conditions employed in this study. However, the presence of a signal peptide at the N-terminus of Hly which bears a strong resemblance to other prokaryotic signal sequences suggests that this protease would be expected to be exported across the R. salmoninarum cell wall. It may be that Hly remains attached to or associated with the cell wall and thus does not form a readily detectable part of the ECPs.

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**REFERENCES**


