Sequence analysis and expression of the M1 and M2 matrix protein genes of hirame rhabdovirus (HIRRV)*

T. Nishizawa1, G. Kurath2, J. R. Winton2,**

1Faculty of Applied Biological Science, Hiroshima University, Higashi-Hiroshima, 724 Japan
2Northwest Biological Science Center, 6505 NE 65th Street, Seattle, Washington 98115, USA

ABSTRACT: We have cloned and sequenced a 2318 nucleotide region of the genomic RNA of hirame rhabdovirus (HIRRV), an important viral pathogen of Japanese flounder Paralichthys olivaceus. This region comprises approximately two-thirds of the 3' end of the nucleocapsid protein (N) gene and the complete matrix protein (M1 and M2) genes with the associated intergenic regions. The partial N gene sequence was 812 nucleotides in length with an open reading frame (ORF) that encoded the carboxyl-terminal 250 amino acids of the N protein. The M1 and M2 genes were 771 and 700 nucleotides in length, respectively, with ORFs encoding proteins of 227 and 193 amino acids. The M1 gene sequence contained an additional small ORF that could encode a highly basic, arginine-rich protein of 25 amino acids. Comparisons of the N, M1, and M2 gene sequences of HIRRV with the corresponding sequences of the fish rhabdoviruses, infectious hematopoietic necrosis virus (IHNV) or viral hemorrhagic septicemia virus (VHSV) indicated that HIRRV was more closely related to IHNV than to VHSV, but was clearly distinct from either. The putative consensus gene termination sequence for IHNV and VHSV, AGAYAG(A), was present in the N-M1, M1-M2, and M2-G intergenic regions of HIRRV as were the putative transcription initiation sequences YGGCAC and AACA. An Escherichia coli expression system was used to produce recombinant proteins from the M1 and M2 genes of HIRRV. These were the same size as the authentic M1 and M2 proteins and reacted with anti-HIRRV rabbit serum in western blots. These reagents can be used for further study of the fish immune response and to test novel control methods.

KEY WORDS: Hirame · Rhabdovirus · Phosphoprotein · Matrix protein

INTRODUCTION

The initial isolation of hirame rhabdovirus (HIRRV) occurred in 1984 from moribund hirame (Japanese flounder) Paralichthys olivaceus and ayu Plecoglossus altivelis in Japan (Kimura et al. 1986). In addition, natural infections have been recorded among black seabream Milio macrocephalus, and mebaru Sebastes inermis (Sano & Fukuda 1987). The related fish rhabdoviruses, infectious hematopoietic necrosis virus (IHNV) and viral hemorrhagic septicemia virus (VHSV), are highly pathogenic for salmonids and certain species of marine fish. The 3 viruses were originally classified as 'other' members of the lyssavirus genus of the Rhabdoviridae (Wunner & Peters 1991), but are now regarded as unassigned vertebrate rhabdoviruses (Wunner et al. 1995). The HIRRV, IHNV and VHSV virions are composed of a negative-sense, single-stranded RNA genome and 5 structural proteins including a nucleocapsid protein (N), 2 matrix proteins (M1 and M2), a glycoprotein (G), and a polymerase (L). The M1 protein of the lyssaviruses is the equivalent of the polymerase-associated protein (P) of the vesiculoviruses (Wunner et al. 1995). The entire IHNV genome has been sequenced (Koener et al. 1987, Gilmore & Leong 1988, Morzunov et al. 1995, Q Inter-Research 1997 Resale of full article not permitted

*Sequence data in this article have been deposited with the DNA data bank of Japan (DDBJ) under Accession No. D45422
**Address for correspondence.
E-mail: jim_winton@nbs.gov

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Distinguishable from other fish rhabdoviruses by serum neutralization tests (Kimura et al. 1986), the structural proteins of HIRRV formed patterns following sodium dodecyl sulphate(SDS)-polyacrylamide gel electrophoresis or 2-dimensional gel electrophoresis that were similar, but not identical, to those of IHNV and VHSV (Nishizawa et al. 1991). The G, N, and M2 proteins of HIRRV were shown to share some antigenic determinants with IHNV and VHSV by western blot analysis (Nishizawa et al. 1991), and it has been suggested that HIRRV might be considered a serotype of IHNV (Winton 1992). Although both HIRRV and IHNV are widely distributed within Japan, HIRRV has not been found outside of Japan, and conversely, IHNV is believed to have been originally restricted to western North America.

We were interested in determining the relationships between HIRRV, IHNV, and VHSV at the molecular level; however, until recently (Nishizawa et al. 1995, Bjorklund et al. 1996, Kurath et al. 1997), there has been no sequence information available for the HIRRV genome. Here, we analyze the sequence data of the partial nucleocapsid protein gene and the 2 matrix protein genes of HIRRV, and compare them with the corresponding sequences available for IHNV and VHSV including the N-M1, M1-M2, and M2-G intergenic regions. In addition, we used an Escherichia coli expression system to produce recombinant M1 and M2 proteins in order to confirm the open reading frames (ORF) of the M1 and M2 genes, to better characterize the 2 proteins, and to create reagents for further studies on the function of the proteins and which can be used to test novel control strategies.

**MATERIALS AND METHODS**

**Virus growth and purification.** Hirame rhabdovirus (strain 8401-H) was grown in the epithelioma papulovirus cypri (EPC) cell line as described by Nishizawa et al. (1995). Virus was purified from cell culture fluid by centrifugation through a discontinuous gradient composed of 30, 35, and 20% (w/w) sucrose in STE (20 mM Tris, pH 7.4, 100 mM NaCl, 1 mM EDTA) buffer followed by centrifugation through a continuous gradient of 5 to 30% (w/w) sucrose in STE as described by Leong et al. (1981).

**cDNA cloning and sequence analyses.** The viral RNA was extracted from gradient-purified virions as described by Kurath & Leong (1985) and cDNA was synthesized, cloned, and sequenced as described by Nishizawa et al. (1995). Sequencing primers are listed in Table 1. Two overlapping cDNA clones, pH9 and pH21s1, were used that, together, represented a 2318 nucleotide region of the HIRRV genome containing the genes of interest. The DNA sequences were assembled and analyzed with the Mac DNASY Pro (Hitachi), MacVector (Eastman Kodak), and PCGENE (IntelliGenetics) software programs. Sequence data were taken from GenBank for the N, M1, and M2 genes of IHNV (strain K, X73872; strain WRAC, L40883), VHSV (strain FI13, X73873; strain 07-71, D00687, U02624 and U03502; and strain Makah, X59241, U02630 and U03503), and for the NS and M genes of rabies virus (strain CVS, X55727, D10499) and vesicular stomatitis virus (serotype New Jersey, K03387, M31884).

**Polymerase chain reaction (PCR) amplification and subcloning.** Two forward and 2 reverse primers (Table 1) were used for specific PCR amplification of the M1 and M2 genes of HIRRV using the plasmid DNA of clone pH21s1 as a template. The forward primers, FM1 and FM2, contained NdeI sites, and the reverse primers, RM1 and RM2, contained BamHI sites.

<table>
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<th>Primer</th>
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<td>Sense</td>
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<tr>
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<td>GCTGGAATTCCATCTCAAGGAACTTTCCTCTTGA</td>
<td>Reverse</td>
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Table 1. Primers used for sequencing and polymerase chain reaction (PCR) amplification of the M1 and M2 genes of hirame rhabdovirus
reverse primers, RM1 and RM2, contained EcoRI or HindIII sites at their 5’ termini, respectively. The PCR reactions were performed using Taq polymerase and 30 cycles of PCR, at 95°C for 1 min, 50°C for 1 min, and 72°C for 1 min. The PCR products were digested with Nde I and EcoRI or HindIII and ligated into the NdeI-EcoRI site (M1 construct) or into the Nde I-HindIII site (M2 construct) of the pET25b(+) expression vector (Novagen) which was used to transform *Escherichia coli* strain BL21(DH3) according to the instructions of the manufacturer.

**Expression of the target gene.** Clones of transformed *Escherichia coli* containing M1 or M2 gene expression plasmids were cultured at 37°C in LB medium (1% Bacto tryptone, 0.5% yeast extract, 1% NaCl, pH 7.5) containing 50 mg ml⁻¹ ampicillin. The cells were induced by resuspension in fresh LB medium containing 50 mg ml⁻¹ ampicillin and 1 mM isopropyl β-D-thiogalactopyranoside (IPTG). Following incubation at 37°C for 2.5 h, the cells were washed and resuspended in 50 mM Tris-HCl (pH 8.0) and 2 mM EDTA to produce the total induced fraction. After addition of 100 mg ml⁻¹ lysozyme and 0.1% Triton X-100, the cell suspension was incubated at 30°C for 15 min and then centrifuged at low speed (300 × g) for 15 min. The pellets and supernatants were used as insoluble induced and soluble induced fractions, respectively. These fractions were analyzed by 12% polyacrylamide gel electrophoresis (PAGE) under the reducing conditions of Laemml (1970). Western blots were performed as described by Towbin et al. (1979).

**RESULTS AND DISCUSSION**

Cloning of the genomic RNA of HIRRV strain 8401-H produced 2 cDNA clones that together represented a contiguous 2318 nucleotide region of the HIRRV genome. Sequencing both strands of the clones produced unambiguous data that showed the clones included a partial N and full-length M1 and M2 genes (Fig. 1). This sequence is available as GenBank accession number D45422. For the purposes of this paper, we refer to the HIRRV ‘genes’ as the entire sequence from the first nucleotide following the (A)₉ of the previous gene through the last A of the (A)₉ tract because the transcription start and termination signals of the fish rhabdoviruses remain experimentally unproven.

Nucleotides 1 to 812 included a partial ORF encoding 250 amino acids and a stop codon that represented the carboxyl-terminal portion of the N gene because of sequence similarities with the 3’ end (mRNA sense) of the N gene of IHNV (Morznov et al. 1995, Schutze et al. 1995) and VHSV (Bernard et al. 1990, 1992). Sequence identities between the equivalent regions of the partial ORFs of the N genes of HIRRV, IHNV, and VHSV are shown in Table 2. The nucleotide identities between HIRRV and the 2 strains of IHNV were higher than those between HIRRV and the 3 strains of VHSV that were analyzed, indicating that HIRRV was more closely related to IHNV than to VHSV. A CLUSTAL alignment of the deduced amino acid sequences of the partial N proteins of VHSV, IHNV and HIRRV showed several small (6 to 7 amino acid) blocks of highly conserved sequence in the central portion of the gene. The partial N gene of HIRRV showed approximately the same degree of sequence identity with the IHNV and VHSV strains as did the M1 and M2 genes; however, only a portion of the entire N gene was analyzed. The ORF of the HIRRV N gene was followed by a 62 nucleotide untranslated region, that included the putative transcription termination signal (discussed below).

The M1 gene of HIRRV occurred between nucleotides 813 and 1584 (Fig. 1). The longest ORF was 684 nucleotides in length (including the stop codon) and encoded a protein of 227 amino acids having a predicted molecular mass of 25779 Da (Nishizawa et al. 1995). The isoelectric point (pI) value of the deduced amino acid sequence was 8.68, which was more basic than that estimated by isoelectric focusing (pI 7.3 to 7.4, Nishizawa et al. 1991) probably due to differences in the degree of phosphorylation of the native form of the protein as produced by fish cells. Three potential serine/threonine phosphorylation sequences (SXXD/E and/or TXXD/E) occurred at amino acids 2, 122, and 186 of the deduced amino acid sequence. The M1 proteins of IHNV and VHSV are also basic and phosphorylated (MacAllister & Wagner 1975, Benmansour et al. 1994).

Sequence identities between the ORFs of the M1 genes of HIRRV, IHNV, and VHSV are shown in Table 2. As with the partial N gene ORF, the nucleotide identities between HIRRV and the 2 strains of IHNV were higher than those between HIRRV and the 3 strains of VHSV. Comparison of the deduced amino acid sequences of the M1 proteins of IHNV and HIRRV showed a highly variable region at the amino-terminus (amino acids 11 to 65) where only 13 of 55 (24%) of the amino acids were identical. This entire region was quite hydrophilic. An analogous region of the M1 protein of VHSV also showed significant variation among strains of the virus as noted by Benmansour et al. (1994). In the central portion of the M1 proteins of HIRRV and IHNV there was a highly conserved region (amino acids 108 to 147) where 39 of 40 (97.5%) amino acids were identical. The location of the conserved and variable blocks of the M1 protein of HIRRV appeared typical for rhabdoviruses.

Spiropoulou & Nichol (1993) described a second ORF within the P gene (equivalent to the M1 gene) of vesic-
Fig. 1. Nucleotide (nt) sequence of a portion of the N gene and the complete M1 and M2 genes of HIRRV. The major ORFs of the N gene (nt 1 to 750), the M1 gene (nt 850 to 1533), and the M2 gene (nt 1639 to 2220) and the corresponding deduced amino acid sequences are shown. The gene junction consensus sequences and the putative transcription start sites are underlined.

ular stomatitis virus (VSV), the prototype member of the vesiculovirus genus of the Rhabdoviridae. This second ORF encoded 2 nested forms of a short (55 and 65 amino acids), highly basic, arginine-rich, protein that was detected in the cytoplasm of infected cells using rabbit antibody raised against a peptide synthesized from the deduced sequence. In the M1 gene of HIRRV, a second, very short (25 amino acid) ORF coding for a highly basic (pI 11.91), arginine-rich (5 of 25 amino acids), protein occurred in an analogous position within the second reading frame. This ORF began at nucleotides 893 to 970, 21 nucleotides downstream from a putative transcription initiation sequence (AACA) internal to the main ORF. Other short ORFs of 10 to 20 amino acids were also present in the M1 gene. One of these, at nucleotides 1067 to 1117, was also in the second reading frame and occurred 14 nucleotides downstream from an AACA sequence. Similar short, highly basic, second ORFs of the M1 (P) genes of rabies virus and IHNV were noted by Spiropoulou & Nichol (1993) and by Morzunov et al. (1995); however, like the second ORF of HIRRV, their use remains unproven. It was interesting that the eukaryotic ribosome scanning model of Kozak (1986) predicts that the internal initiation of a second ORF is more likely when the primary start codon (AUG) is in a suboptimal context and the second AUG is in a more favorable context. This was true of the M1 gene of HIRRV where the primary ORF was only in fair context (G at position −3, T at +4) while the second and third ORFs were in a better context (A at position −3, C/T at +4). In contrast, a nearly optimal Kozak’s box occurs in the primary ORFs of the M1 and M2 genes of VHSV (Benmansour et al. 1994).

The M2 gene of HIRRV occurred between nucleotides 1585 and 2285 (Fig. 1). The longest ORF was 582 nucleotides in length (including the stop codon) and encoded a very basic protein of 193 amino acids having a predicted molecular mass of 21 615 Da and a pI of 9.7 (Nishizawa et al. 1995). The primary AUG was in a good context for translation (A at position −3). While 2 short ORFs of 23 and 25 amino acids were identified near the 3′ end of the M2 gene, these were in a poor Kozak’s context and there was no indication that they

Table 2. Pairwise comparisons of the open reading frame nucleotide sequences (upper right) and deduced amino acid sequences (lower left) of HIRRV, 2 strains of IHNV and 3 strains of VHSV. Percent identities are shown for the partial N (top), complete M1 (middle) and complete M2 (bottom) genes. Sequence data were taken from GenBank for the N, M1 and M2 genes of IHNV (strain K, X73872; strain WRAC, L40883) and VHSV (strain FI13, X73873; strain 07-71, D00687, U02624 and U03502; and strain Makah, X59241, U02630 and U03503).

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<th>Amino acid identity</th>
<th>HIRRV strain 8401H</th>
<th>IHNV strain K</th>
<th>IHNV strain WRAC</th>
<th>VHSV strain FI13</th>
<th>VHSV strain 07-71</th>
<th>VHSV strain Makah</th>
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<td>67.0</td>
<td>97.9</td>
<td>54.8</td>
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<tr>
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<td>74.2</td>
<td>55.8</td>
<td>55.5</td>
<td>56.6</td>
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<tr>
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<td>64.5</td>
<td>74.7</td>
<td>52.9</td>
<td>52.8</td>
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<td>74.7</td>
<td>52.9</td>
<td>52.8</td>
<td>54.9</td>
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<tr>
<td>VHSV strain Makah</td>
<td>59.7</td>
<td>64.5</td>
<td>74.7</td>
<td>52.9</td>
<td>52.8</td>
<td>54.9</td>
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of HIRRV was 65 nucleotides in length, considerably longer than the corresponding sequence of VHSV (35 nucleotides), but somewhat shorter than the untranslated region downstream of the M2 gene of IHNV (103 nucleotides).

The sequences AACABYMCT and AACAG have been described as consensus sequences for transcription initiation and/or capping signals of lyssavirus or vesiculovirus genes (Banerjee & Barik 1992, Bourhy et al. 1993). The sequence AACAB, which has been described as a potential transcription start signal for the VHSV N, M1, M2, and G genes (Bernard et al. 1990, 1992, Benmansour et al. 1994), was present in each of the 3 HIRRV N-M1, M1-M2, and M2-G intergenic regions (nucleotides 829, 832, 839, 1607, 2308 and 2311) downstream of the termination signals and before the primary initiation codon. It also occurred ahead of the potential second and third ORFs of the M1 gene (nucleotides 869 and 1049); however, the AACAB sequence is absent from IHNV gene junctions (Koener et al. 1987, Gilmore & Leong 1988, Morzunov et al. 1995, and it has not been experimentally proven that AACAB capping signals of lyssavirus or vesiculovirus genes (Banerjee & Barik 1992, Bourhy et al. 1993). The sequence AACAB, which has been described as a potential transcription start signal for the VHSV N, M1, M2, and G genes (Bernard et al. 1990, 1992, Benmansour et al. 1994), was present in each of the 3 HIRRV N-M1, M1-M2, and M2-G intergenic regions (nucleotides 829, 832, 839, 1607, 2308 and 2311) downstream of the termination signals and before the primary initiation codon. It also occurred ahead of the potential second and third ORFs of the M1 gene (nucleotides 869 and 1049); however, the AACAB sequence is absent from IHNV gene junctions (Koener et al. 1987, Gilmore & Leong 1988, Morzunov et al. 1995), and it has not been experimentally proven that the AACAB sequence functions as a transcription start signal in these rhabdoviruses of fish. In contrast, the conserved sequence, YGGCAG, was present immediately following the termination signals of nearly all of the known genes of HIRRV, IHNV, and VHSV. Only the comparable sequences in the M1–M2 junction of VHSV (strain 07-71) and the M2-G junction of VHSV
(strain Makah) are slightly altered to YGGCTC or YGGTAC, respectively (Benmansour et al. 1994). Thus, this sequence appears to be a more likely transcription initiation signal in the latter than in the former rhabdoviruses as noted by others (Morzunov et al. 1995, Bjorklund et al. 1996, Kurath et al. 1997), but this remains to be proven experimentally.

To confirm the identity of the putative ORFs and to learn more about the antigenic nature of the 2 matrix proteins, the M1 and M2 genes of HIRRV were subcloned into the pET25b(+) vector and expressed in *Escherichia coli* cells. The expression vectors encoding both the M1 and M2 protein genes were genetically stable in the host cells, and there was no significant difference in codon usage between the 2 genes. The results of SDS-PAGE analysis of the expression products from the M1 and M2 genes are shown in Fig. 2. While bacterially expressed proteins are not always identical to their native counterparts in size or in the degree of phosphorylation or glycosylation due to differences in protein processing and to the presence of amino acids encoded by sequences in the expression vector, in cells transformed with pET25b(+) encoding the M1 gene, an expressed protein with an apparently size identical to the native HIRRV M1 protein was detected. The expressed M1 protein of HIRRV was present in both the insoluble induced and soluble induced fractions. In contrast, no expression product of a size similar to the HIRRV M2 protein was detected in the total induced fraction of the cells transformed with pET25b(+) containing the M2 gene construct; however, a small amount of expressed M2 protein was detectable in the insoluble induced fraction. The identities of both expression products were confirmed by reaction with anti-HIRRV rabbit serum in western blot analyses (Fig. 2), verifying that the ORFs identified by sequence analysis encode the viral proteins in vivo.

The M1 and M2 expression products will be valuable reagents for further research on the function of these proteins in vivo and may provide reagents for experimental control methods. Fish surviving infection with IHNV have been reported to produce anti-M1 antibody in their sera (Mourich & Leong 1992, LaPatra et al. 1993, Ristow et al. 1993) and the M1 protein of rabies virus was shown to stimulate a strong cytolytic T-cell response in mice (Larson et al. 1992). Thus, the recombinant M1 protein of HIRRV might be an attractive candidate for further research as a potential vaccine or as a supplement to recombinant vaccines based upon the G protein.

It has been shown that the expressed M2 protein of VSV is capable of inhibiting transcription in vitro in the absence of any other viral component (Black & Lyles 1992, Black et al. 1993, 1994). If the M2 protein of HIRRV has a similar inhibitory function, this may in part explain why the expression level of the HIRRV M2 protein in *Escherichia coli* was quite low in comparison to levels obtained for the M1 protein. It would be interesting to test the recombinant M2 protein for ability to inhibit HIRRV transcription in vitro and in vivo as a potential control strategy.

The presence of an NV gene, conserved gene junction sequences, and similarities in the nucleotide sequences of the N, M1, M2, G and NV genes have led several investigators to conclude that HIRRV belongs to the same unique group of fish rhabdoviruses that includes IHNV and VHSV and that the 3 viruses comprise a novel genus within the *Rhabdoviridae* (Morzunov et al. 1995, Nishizawa et al. 1995, Bjorklund et al. 1996, Kurath et al. 1997). This relationship

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Fig. 3. Dendrogram produced by CLUSTAL analysis of the deduced amino acid sequences of the M1 genes of HIRRV (HRVM1); 2 strains of IHNV, from North America (IHNVKM1) and Europe (IHNVKM1); and 3 strains of VHSV, from Europe (0771M1, VHSFI13M1) and North America (MAKAHM1). The amino acid sequences of the NS protein (equivalent to phosphoprotein or M1 protein) of rabies virus strain CVS (RABCVSNS) and vesicular stomatitis virus serotype New Jersey (VSVNS) were used as representatives of the lyssavirus and vesiculovirus genera, respectively. Nearly identical dendrograms were generated by comparison of the partial N and the complete M2 amino acid sequences. Sequence data were taken from GenBank for the M1 genes of IHNV (strain K, X73872; strain WRAC, L40868) and VHSV (strain FI13, X73873; strain 07-71, U02624; and strain Makah, U02630) and for the NS genes of rabies virus strain CVS (X55727) and VSV serotype New Jersey (K03387).
was most easily seen in dendrograms generated by CLUSTAL analysis in which the N, M1 and M2 amino acid sequences of HIRRV were separately aligned with the homologous sequences of 2 strains of IHNV (European and North American), 3 strains of VHSV (2 European and 1 North American), rabies virus (strain CVS), and vesicular stomatitis virus (serotype New Jersey). The dendrograms showed that HIRRV was more closely related to IHNV than to VHSV and that these fish viruses are quite distinct from the prototype members of the lyssavirus and vesiculovirus genera (Fig. 3) in agreement with results of Morzunov et al. (1995) and Bjorklund et al. (1996). The nucleoprotein and both matrix proteins gave very similar dendrograms suggesting the evolution rate of the 3 genes is similar, although the M2 protein showed slightly less divergence (Table 2). These data also provide evidence that HIRRV is a distinct virus, rather than a strain of IHNV, because the amino acid sequence differences between the matrix proteins of HIRRV and their homologs in IHNV or VHSV were significantly greater than differences among even the most diverse strains of the North American and European isolates of VHSV (Basurco et al. 1995) or the North American and European isolates of IHNV (Nichol et al. 1995).

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