Characteristics of a new reovirus isolated from epizootic ulcerative syndrome infected snakehead fish

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ABSTRACT: Epizootic ulcerative syndrome (EUS) has been infecting a wide range of fishes in the South and Southeast Asia for the last 2 decades. One reovirus-like agent (snakehead reovirus, SKRV), isolated from an EUS-infected snakehead fish and investigated in the present study, is the only reovirus so far isolated from an EUS-infected fish. SKRV was characterised by the presence of a double-stranded RNA genome with icosahedral symmetry and double capsid. The virus had an average size of 71 nm, a buoyant density of 1.36 g ml⁻¹ in CsCl and lacked a lipid-containing envelope. Apart from the above, the presence of a segmented genome and structural proteins falling into 3 specific size classes confirmed that the virus belongs to the family Reoviridae. SKRV differed from aquareoviruses by the lack of a cytopathic effect (CPE) with syncitium formation and in the segmentation pattern of RNA genome. The resistance to pH (3.0 to 9.0) and heat treatment and inability to multiply in mammalian cell lines and haemagglutinate human ‘O’ red blood cells (RBCs) differentiated SKRV from the rest of the similar genera in the family Reoviridae. Serological comparison indicated the antigenic distinctness of the isolate from selected American and European aquareoviruses. SKRV grew well in SSN-1 and SSN-3 cells at 25 to 30°C but not in the most common Aquareovirus susceptible coldwater fish cell line—CHSE-214.

KEY WORDS: EUS · Reovirus · Snakehead fish · Characterization

INTRODUCTION

Epizootic ulcerative syndrome (EUS) has been recognised as the single most potentially damaging disease of fishes in South and Southeast Asia for the last 2 decades. The disease infects a wide variety of fishes, both freshwater and brackishwater species, and snakeheads have by far been the most severely affected fish. A number of infectious agents has been isolated from EUS-infected fish (Roberts et al. 1994). During an investigation of EUS in Thailand, a reovirus-like agent was isolated from an EUS-infected fish following a co-habitation experiment (Roberts et al. 1994). The family Reoviridae includes non-enveloped, double-stranded RNA viruses, icosahedral in structure, that may appear spherical with a size of 60 to 80 nm in diameter and consist of several protein layers surrounding the inner core (Murphy et al. 1995). The precise morphology of the virions varies depending on the genus. Further characteristics consist of 10,11 or 12 segments of linear double-stranded genome depending on the genus with a total $M_r$ (relative molecular ratio) of 12 to $20 \times 10^6$. The $M_r$ of these segments range from 0.2 to $3.0 \times 10^6$. Fish viruses belonging to the family Reoviridae are of comparatively recent origin compared to many other groups of fish viruses. The family Reoviridae presently has 9 genera, one of which is the new genus Aquareovirus that includes the reoviruses isolated from fish (Murphy et al. 1995). Since the first isolation of a reovirus from golden shiners Notemigonus crysoleucas in 1977 in the USA, more than 30 aquareoviruses have so far been isolated from fish and shellfish worldwide (Lupiani et al. 1996). The present study was aimed at characterisation of a reovirus-like agent recovered from the above-described characteristically EUS-infected snakehead fish.

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MATERIALS AND METHODS

Viruses and cells. The snakehead reovirus (SKRV or T9231) was isolated from a characteristically EUS-infected striped snakehead fish *Channa striata* in Thailand during an epizootic in 1992 (Roberts et al. 1994) following a co-habitation trial. EUS was induced in 6 healthy striped fish by co-habitation with 6 characteristically EUS-infected striped fish. All 6 of the EUS-infected fish died within 5 d of co-habitation trial. Tail ulcerations were developed in the healthy striped fish 12 d post co-habitation. The ulcerated tissues from the integument and muscles of the tail lesions were sampled for virological examination by inoculating the tissue homogenate onto striped snakehead cells (SSN-1, Frerichs et al. 1991) and bluegill *Lepomis macrochirus* fry cells (BF-2). The cytopathic effect (CPE) was developed in SSN-1 cells incubated at 25°C after 8 d. Virus was further passaged onto fresh SSN-1 cells and CPE observed after 2 d. Stock virus was prepared by growing SKRV in SSN-1 cells and, once CPE was complete, tissue culture fluid was harvested, clarified by centrifugation at 1000 × g for 15 min and stored at –70°C in aliquots.

Other viruses used in the study for characterization of SKRV were blotched snakehead virus (BSNV, John & Richards 1999), golden shiner virus (GSV, Plumb et al. 1979) and catfish reovirus (CRV, Amend et al. 1984) from the USA (provided by Dr R. P. Hedrick, School of Veterinary Medicine, University of California, USA) and tench reovirus (TNRV) and chub reovirus (CHRV) (Ahne & Kolbl 1987) from Europe (provided by Dr W. Ahne, Institute of Zoology and Hydrobiology, University of Munich, Germany).

SSN-1 cells were used for propagation of SKRV. BSNV were inoculated and propagated in SSN-1 or BF-2 cells. GSV was grown on *Epithelioma papillosum cyprini* (EPC), fathead minnow *Pimephales promelas* cells (FHM) or chinook salmon *Oncorhynchus tshawytscha* embryo cells (CHSE-214). Brown bullhead *Ictalurus nebulosus* cells (BB) were used for propagating CRV. TNRV and CHRV were grown and titrated using EPC or FHM cells. Cells were maintained in Leibovitz-15 (L-15) or Eagle’s minimum essential medium (EMEM) (Gibco BRL, Paisley, UK), supplemented with 10% foetal bovine serum (Gibco BRL). All the cells were maintained at 25°C with 5% FBS after inoculation with T9231, BSNV, GSV, CRV and at 20°C after TNRV and CHRV inoculation.

Purification of the viruses. SKRV was concentrated by ultracentrifugation following propagation of the virus in SSN-1 cells grown in 500 cm² triple layer flasks (Nunc, Roskilde, Denmark). When CPE was extensive, culture fluid was harvested and clarified by centrifugation at 2000 × g for 30 min to remove the cellular debris. Supernatant and pelleted cells were then processed separately. Approximately 600 ml of collected supernatant was high speed clarified at 12,000 × g for 10 min in an SW-28 rotor in a Beckman L80 ultracentrifuge (Beckman, Fullerton, CA, USA) before pelleting the virus at 100,000 × g for 90 min in an SW-41 Ti rotor (Beckman). The virus pellets were pooled and resuspended in 2 ml TNE buffer (0.01M-Tris HCl, 0.1M-NaCl, 0.001M-EDTA, pH 7.5). Pelleted cells were resuspended in 4 ml TNE buffer and subjected 3 times to freeze/thawing in liquid nitrogen. Cell associated virus was extracted by treatment with trichlorotrifluoroethane (Freon, Sigma, Poole, UK). An equal volume of freon was added to the cell suspension and mixed well using a vortex mixer at full speed for 3 min. After centrifugation at 2000 × g for 10 min, the aqueous phase was removed and freon phase was re-extracted using an equal volume of TNE buffer as above. Aqueous phase extracts were pooled and virus pelleted by ultracentrifugation at 100,000 × g for 90 min. Pelleted virus was resuspended in 1 ml TNE buffer. Combined virus preparations from culture supernatant and cell pellet were pooled and purified by isopycnic banding in CsCl gradients.

Polyethylene glycol (PEG, MW 8000, Sigma) was used to concentrate CRV, GSV, TNRV, CHRV and BSNV. CRV was propagated in BB cells by simultaneous inoculation of virus and cells in 500 cm² triple layer flasks. Tench and chub reoviruses were propagated in the same way in EPC/FHM cells. GSV was grown in preformed, confluent CHSE-214 cells in 175 cm² flasks after infecting at a multiplicity of infection (MOI) of 0.01 to 0.1. The virus was allowed to adsorb for 1 h at room temperature before adding maintenance medium containing antibiotics (50 IU ml⁻¹ penicillin, 50 µg ml⁻¹ streptomycin and 50 µg ml⁻¹ kanamycin). BSNV was propagated in preformed monolayers of BF-2 cells as above in 175 cm² flasks in L-15 medium supplemented with antibiotics.

When the CPE became extensive, remaining cells were scraped into the culture medium and the culture fluid clarified at 2000 × g for 30 min. Cell pellets were freon extracted as described above except that the virus recovered after freon extraction was added to the clarified supernatant without ultracentrifugation or directly overlaid on to the top of a CsCl gradient. Approximately 800 ml of infected tissue culture fluid was processed for virus concentration using PEG as described earlier (John & Richards 1999).

Concentrated virus resuspended in TNE was layered on top of a preformed 20 to 40% discontinuous CsCl gradient and isopycnically banded by centrifuging at 130,000 × g for 17 h in an SW-41 Ti rotor. The virus bands recovered by side puncturing the centrifuge tubes were pooled and resuspended in TNE buffer and
repelleted at 100,000 × g for 90 min. The purified virus pellet was dissolved in 200 µl TNE buffer and stored at −70°C until use.

**Electron microscopy.** SKRV was grown on SSN-1 cells (24 to 48 h), fixed in Karnovsky’s fixative (Karnovsky 1965) and the cell pellets were held overnight in 0.1 M cacodylate buffer at 4°C. Following post-fixing in 0.5% osmium tetroxide for 1 h and dehydration by acetone, the pellets were impregnated at 60°C for 48 h with epoxy resin (Araldite CY212, Agar Scientific Ltd, Stansted, UK). Ultra thin sections (80 nm) of cell pellets were cut using a Reichert Ultracut E microtome (Reichert, UK) (Hayat 1989). The sections after staining with uranyl acetate (Watson 1958) and lead citrate (Reynolds 1963) were examined and photographed using the transmission electron microscope at 80 kV. For negative staining, 5 µl of purified virus suspensions was stained by methylamine tungstate. Grids were examined and photographed using the transmission electron microscope at 80 kV.

**Growth in mammalian cell lines.** The ability of SKRV to multiply in mammalian cells was determined by using BHK-21 (baby hamster kidney) and Vero (green monkey kidney) cell lines. BHK-21 cells were grown in EMEM and incubated at 25 and 37°C. Vero cells were maintained in DMEM (Dulbecco’s modified Eagle's medium, Sigma) with 10% FBS and antibiotics (50 µg ml−1 penicillin, 50 µg ml−1 streptomycin, 50 µg ml−1 kanamycin and amphotericin 2.5 µg ml−1) at 37°C. Flasks were observed and examined using photographs for the development of CPE.

**Biophysical and biochemical characteristics.** The type and nature of nucleic acid of SKRV were analysed using a DNA inhibitor, 5-ido-2 deoxyuridine (IUDR) and acridine orange staining (Rovozzo & Burke 1973). Resistance of SKRV to chloroform treatment was tested according to the method of Feldman & Wang (1961). SKRV was tested for the ability to withstand heat treatment at 56°C for 2 h and pH treatment at pH 3 and 9 for 30 min. Density of the virions were found out following isopycnic centrifugation of semipurified virus in cesium chloride gradient as before. At the end of the run, fractions were collected using a Beckman fraction recovery system (Beckman). The refractive index of each fraction was determined using an ABBE refractometer (Bellingham & Stanley Ltd, Kent, UK) after titrating the fractions in SSN-1 cells. The density and titre of each fraction were plotted against the respective number of fractions and the density of virions was extrapolated from the graph against fractions that gave the highest titre.

**Analysis of structural proteins.** Structural proteins of SKRV were analysed by SDS-polyacrylamide gel electrophoresis (PAGE) using a BRL Mini-V 80 system (BRL Life Technologies UK Ltd, UK) along with BSNV, CRV, GSV, TNRV, CHRV and broad range molecular weight standards (BioRad, Richmond, CA, USA). Structural proteins of the viruses were resolved by discontinuous polyacrylamide-SDS slab gels (Laemmli 1970) of 10% acrylamide prepared according to the general procedures of Hames & Rickwood (1990). Electrophoresis was conducted at 200 V and 200 mA for 45 to 55 min and the gels were stained in 0.1% Coomassie brilliant blue. Molecular weights of the virion proteins were determined according to the method of Hames & Rickwood (1990).

**Analysis of nucleic acids.** Nucleic acid from purified virions was extracted using Proteinase K-SDS digestion and phenol-chloroform extraction according to Burleson et al. (1992). RNA present in the solution was quantified by measuring the optical density in an Ultrospec UV spectrophotometer (Pharmacia Biotech, Sweden) at 260 nm. Viral RNA was analysed by agarose gel electrophoresis according to Sambrook et al. (1989) with minor modifications. Agarose gels of Seakem agarose (1.5%) (Flowgen, Leicestershire, UK) and Metaphor agarose (3%) (Flowgen) were prepared in MOPS buffer (0.2 M 3-N-morpholino-2-hydroxypropane sulfonic free acid, 0.05 M sodium acetate, 0.005 M EDTA, pH 7.0) with formaldehyde. Viral RNA samples and molecular mass markers (RNA molecular mass markers II and III, Boehringer Mannheim, Germany) were electrophoresed on a horizontal gel electrophoresis apparatus (Pharmacia Biotech) at 1 or 1.5 V cm−1 (14 to 20 h). The gels were stained in 1 µg ml−1 ethidium bromide for 30 min and destained in DEPC (diethylpyrocarbonate, BDH, Poole, UK) treated deionised water before photographing using Polaroid films or scanning using Gel Works advanced ID software on a Gel-doc system (UVP, UK). Molecular weights of the viral RNA bands were calculated from a semi-logarithmic standard curve of molecular weights plotted against migration distance of the markers.

**Serological studies.** The ability of SKRV haemagglutinate human ‘O’ type erythrocytes (Sigma) was determined by the method of Kuchler (1977). Polyclonal rabbit antisera against the viruses was prepared in New Zealand white rabbits as described by Lilley & Frerichs (1994). Cross-neutralising activity of the antisera of SKRV, CRV, GSV and CHRV was tested by the α procedure against SKRV, CRV, GSV, TNRV and CHRV (Rovozzo & Burke 1973). Briefly, all viruses were 10-fold serially diluted and mixed with equal volumes of the antisera at appropriate dilutions. Mixtures were incubated in 96-well plates at room temperature for 60 min and 100 µl of appropriate single cell suspension in maintenance medium was added. The plates were
then incubated at appropriate temperatures and monitored for CPE over 10 d and neutralisation indices were calculated as the ratio of the virus titre without antiserum to the virus titre in the presence of antiserum.

RESULTS

Cytopathology

The CPE of SKRV in SSN-1 were compared with the CPE of aquareoviruses. SKRV did not produce typical syncitia with multiple nuclei in SSN-1 cells similar to those found in other aquareoviruses in the respective cell lines (Fig. 1). SKRV infection in SSN-1 cells progressed with the formation of focal areas of cell destruction in the monolayer due to aggregation and rounding up of granular refractile cells forming cell clumps. CPE in the form of holes in the cell sheet usually progressed for about 6 to 10 d to involve the entire monolayer, at which time the cells in clumps became detached from the substrate, leaving a few still attached to the culture surface. SKRV was unable to grow and cause any CPE in the most common aquareovirus susceptible cold-water fish cell line, CHSE-214, and in both BHK-21 and Vero cells during the period of observation.

Electron microscopy

Like other aquareoviruses, SKRV particles had icosahedral symmetry with double capsid structure. The virions in cell culture had an overall diameter of 68 ± 1.8 nm with a size range of 62 to 77 nm (n = 19). Characteristic cytoplasmic inclusion bodies of variable size and shapes (some reaching a length of 3.6 µm) with relatively electron-dense areas of granular matrix were found scattered in the cytoplasm without any special membranous envelope at different stages of virus formation. Some of the viroplasms were found to be fully formed with a paracrystalline array of virus particles filling the entire area of the inclusion body (Fig. 2).

Unenveloped spherical particles having a mean diameter of 71 ± 1.2 nm (range 64 to 77 nm, n = 18) and double capsid structure were observed in the negatively stained preparations of SKRV (Fig. 3). The virions had an inner diameter of 43 to 48 nm and capsid thickness of 12 to 15 nm. Morphologically, SKRV was similar to the other aquareoviruses CRV, GSV, TNRV and CHRV.

Biophysical and biochemical characteristics

Lack of inhibition of SKRV in SSN-1 cells growing in 50 µg ml⁻¹ IUDR-incorporated culture medium and the presence of yellowish green cytoplasmic inclusions following acridine orange staining of virus-infected cell cultures indicated the presence of a double-stranded DNA genome.

Fig. 1. (a) Normal SSN-1 monolayer. (b) SKRV growing in SSN-1 cells showing formation of focal areas of cell destruction in the monolayer due to aggregation and rounding up of granular refractile cells forming cell clumps.
RNA genome (Rovozzo & Burke 1973). Infectivity resistance to chloroform treatment confirmed the absence of a lipid-containing envelope (Feldman & Wang 1961). SKRV was resistant to heat treatment at 56°C for 2 h and was also stable at acid (3.0) and alkaline (9.0) pH for 30 min (Rovozzo & Burke 1973). Isopycnic centrifugation of pelleted virus at 130,000 \( \times \) \( g \) for 17 h in 20 to 40% discontinuous CsCl gradient and assay of recovered fractions in SSN-1 cells established a buoyant density of 1.36 g ml\(^{-1}\) for infective particles.

**Analysis of structural proteins**

Polypeptide composition of SKRV was similar to other American and European aquareoviruses CRV, GSV, TNRV and CHRV (Fig. 4, Table 1). Structural proteins of SKRV were resolved into 5 components falling into 3 size classes (large, medium and small) of polypeptides characteristic of the family Reoviridae. Estimated molecular masses of the proteins of SKRV were as follows: VP1 = 154, VP2 = 140, VP3 = 65, VP4 = 47 and VP5 = 38 kDa.

**Analysis of nucleic acids**

The electropherotype of genomic RNAs of all the viruses analysed fell into the general nucleic acid organisation pattern of the family Reoviridae, having 3 size classes of genomic segments (Fig. 5a). The RNA of SKRV analysed for 20 h in 3% Metaphor agarose gel further resolved the small genomic segments (Fig. 5b). Estimated molecular masses of the SKRV genome segments are shown in Table 2. Unlike the 3 aquareoviruses CRV, GSV and TNRV, smaller segments of SKRV RNA did not resolve into 5 components characteristic of the genus Aquareovirus. The smallest RNA segment of SKRV had an estimated molecular mass of \( 0.67 \times 10^6 \) whereas the smallest segments of CRV and GSV were 0.54 and 0.52 \( \times 10^6 \) respectively (as obtained in the present study). Comparative analysis of the electropherotype of SKRV with that of CRV, GSV and TNRV indicated that the SKRV genome probably contains only 10 RNA segments that ranged in molecular mass from 2.93 \( \times 10^6 \) (4.39 kbp) to 0.67 \( \times 10^6 \) (1.01 kbp) Da. The total molecular mass of the SKRV genome was approximately 16.38 \( \times 10^6 \) (24.56 kbp).

**Serological characteristics**

SKRV, GSV and CRV were tested for their capacity to haemagglutinate human ‘O’ type red blood cells (RBC). At a final dilution of 1% RBC after addition of virus, none of the tested virus isolates showed agglutination of human ‘O’ erythrocytes at room temperature, 25°C.

Neutralisation indices (NI) of the test involving antiserum against SKRV, GSV, CRV and CHRV and respective viruses and TNRV showed specificity of SKRV antisera evident from the high homologous NI and little cross reaction with any of the other viruses tested (Table 3). SKRV was also not neutralised by any of the other antisera tested. Results from the cross neutralisation studies indicated that SKRV is antigenically unrelated to the American and European aquareovirus isolates GSV, CRV, TNRV and CHRV.
Table 1. Estimates of molecular mass of viral structural proteins of SKRV, reovirus type 3 and selected aquareoviruses. SKRV was analysed by SDS-PAGE in 10% acrylamide concentration.

<table>
<thead>
<tr>
<th>Virus</th>
<th>Virus polypeptides (molecular mass in kDa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>VP1</td>
</tr>
<tr>
<td>Reo 3α</td>
<td>155-λ1</td>
</tr>
<tr>
<td>Reo 3β</td>
<td>137/143-λ1</td>
</tr>
<tr>
<td>SKRV</td>
<td>154</td>
</tr>
<tr>
<td>CRV</td>
<td>132</td>
</tr>
<tr>
<td>GSV</td>
<td>137</td>
</tr>
<tr>
<td>13P2</td>
<td>135</td>
</tr>
<tr>
<td>CSV</td>
<td>137</td>
</tr>
<tr>
<td>AFRV</td>
<td>137</td>
</tr>
<tr>
<td>SRV</td>
<td>145</td>
</tr>
<tr>
<td>LSV</td>
<td>139</td>
</tr>
<tr>
<td>SBR</td>
<td>130</td>
</tr>
</tbody>
</table>

*aFrom Joklik (1983a); *b*virion proteins ≤ 2% (λ3-135, μ1-80, μ2-70 and σ1-42 kDa) not included; *c*from Nibert et al. (1996); *d*from the present study; *e*from Winton et al. (1987); *f*from Varner & Lewis (1991); *g*protein not included; *h*Marshall et al. (1990); *i*from Hsu et al. (1989); *j*Samal et al. (1991b); *k*Subramanian et al. (1994) (2 additional minor proteins [VP3-126 kDa, VP4-73 kDa] also reported)
Table 2. Estimated molecular mass of RNA segments of SKRV analysed in 1.5% agarose gels. Molecular masses of segments were calculated from semi-logarithmic plot of molecular weight versus migration distance of the markers

<table>
<thead>
<tr>
<th>Segment</th>
<th>SKRV</th>
<th>CRV</th>
<th>GSV</th>
<th>Reovirus type 3</th>
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<tr>
<td>L1</td>
<td>2.93</td>
<td>2.40</td>
<td>2.50</td>
<td>2.50&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>L2</td>
<td>2.83</td>
<td>2.40</td>
<td>2.50</td>
<td>2.70&lt;sup&gt;d&lt;/sup&gt;</td>
</tr>
<tr>
<td>L3</td>
<td>2.78</td>
<td>2.30</td>
<td>2.20</td>
<td>2.50</td>
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<tr>
<td>M1</td>
<td>1.69</td>
<td>1.60</td>
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<td>1.60</td>
</tr>
<tr>
<td>M2</td>
<td>1.65</td>
<td>1.50</td>
<td>1.50</td>
<td>1.70</td>
</tr>
<tr>
<td>M3</td>
<td>1.51</td>
<td>1.30</td>
<td>1.40</td>
<td>1.60</td>
</tr>
<tr>
<td>S1</td>
<td>0.88</td>
<td>0.90</td>
<td>0.90</td>
<td>0.92</td>
</tr>
<tr>
<td>S2</td>
<td>0.74</td>
<td>0.88</td>
<td>0.80</td>
<td>0.76</td>
</tr>
<tr>
<td>S3</td>
<td>0.71</td>
<td>0.61</td>
<td>0.66</td>
<td>0.64</td>
</tr>
<tr>
<td>S4</td>
<td>0.67</td>
<td>0.41</td>
<td>0.40</td>
<td>0.61</td>
</tr>
<tr>
<td>L1&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.40</td>
<td>0.35</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Total</td>
<td>16.38</td>
<td>14.70</td>
<td>14.71</td>
<td>14.73</td>
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</tbody>
</table>

<sup>a</sup>From Winton et al. (1987); <sup>b</sup>calculated at mol. wt of 6.67 x 10^5/kbp; <sup>c</sup>from Shatkin et al. (1968); <sup>d</sup>from Ginsberg (1988)

DISCUSSION

SKRV used in the present study was a single reovirus-like agent isolated from a characteristically EUS-infected striped snakehead fish. The nature of cytopathology induced by SKRV in homologous SSN-1 cells was different from the typical syncitia formation caused by aquareoviruses in respective cell lines (Winton 1989, Hetrick et al. 1992). The CPE of SKRV in SSN-1 cells, which was characterised by focal areas of cell destruction in the monolayer due to aggregation and rounding up of granular refractile cells forming cell clumps and lack of a plaque-like syncitium formed out of cell fusion, was similar to the CPE induced by mammalian reoviruses in cell cultures (Gomatos et al. 1981, Chen & Jiang 1984, Lupiani et al. 1989, Samal et al. 1990, Varner & Lewis 1991). Multiplication of cells escaping lytic infection resulting in regeneration of the monolayer after SKRV infection was observed occasionally in SSN-1 cells. A similar observation was reported in the case of GSV in FHM cells (Plumb et al. 1979) and turbort reovirus in EPC cells (Lupiani et al. 1989). The refractory nature of Vero and BHK-21 cell lines to SKRV infection indicated the host specificity of the virus, which helps to differentiate it from reoviruses of mammalian origin (Rosen et al. 1979, Joklik 1983c) and orbiviruses (Gorman et al. 1983). Aquareoviruses were also, in general, found unable to grow in cells of mammalian or avian origin (Meyers 1979, Hsu et al. 1989, Samal et al. 1990, Dopazo et al. 1991).


Resistance to heat treatment at 56°C for up to 2 h has shown the similarity of SKRV to orthoreoviruses (Drayna & Fields 1982). Unlike orthoreoviruses, heating to 60°C rapidly inactivates the orbiviruses (Murphy et al. 1995). Aquareoviruses exhibit variations in the sensitivity to heat treatment (Plumb et al. 1979, Winton et al. 1981, Chen & Jiang 1984, Lupiani et al. 1989, Varner & Lewis 1991). Reoviruses are also characterised by their resistance to pH, which varies among different genera (Murphy et al. 1995). Of the 5 different genera of reoviruses infecting vertebrates (Orthoreovirus, Orbivirus, Rotavirus, Aquareovirus and Coltivirus), orbiviruses and coltiviruses are characterised by a loss of infectivity at low pH in contrast to the stability of reoviruses at low pH (Gorman et al. 1983). Avian reovirus strains are also resistant to pH 3.0 treatment (Kawamura et al. 1965). Aquareoviruses are generally stable over a wide pH range (Meyers 1979, Plumb et al. 1979, Winton et al. 1981, Chen & Jiang 1984, Ahne & Kolbl 1989, Lupiani et al. 1989, Varner & Lewis 1991). Rotaviruses vary in their stability to pH treatment (Holmes 1983). Retention of infectivity by SKRV after heat and pH treatment shows that it is characteristically similar to the vertebrate members of the family Reoviridae (except orbiviruses and coltiviruses). As is typical of the members of the family Reoviridae which have vertebrate hosts, complete virions of SKRV had a buoyant density of 1.36 g ml⁻¹ in CsCl (Smith et al. 1969, Holmes 1983). Buoyant density of complete virions of aquareoviruses was also reported to range from 1.34 to 1.36 g ml⁻¹ (Winton et al. 1987, Hsu et al. 1989, Varner & Lewis 1991).

Table 3. Cross-neutralisation indices among SKRV, GSV, CRV, CHRV and TNRV. Neutralisation indices (NI) expressed as logarithmic difference in virus titres in the presence and absence of antiserum. NI values equal to or greater than 1.7 indicate serological similarity. Dilution of antiserum used in the test given in parentheses.

<table>
<thead>
<tr>
<th>Virus strain</th>
<th>Antiserum</th>
</tr>
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<tbody>
<tr>
<td>SKRV&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.5</td>
</tr>
<tr>
<td>CRV&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.44</td>
</tr>
<tr>
<td>CHRV&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.79</td>
</tr>
</tbody>
</table>

<sup>a</sup>From Winton et al. (1987); <sup>b</sup>calculated at mol. wt of 6.67 x 10^5/kbp; <sup>c</sup>from Shatkin et al. (1968); <sup>d</sup>from Ginsberg (1988)
Structural proteins of SKRV were resolved into 3 size classes characteristic of the family Reoviridae (Joklik 1983c). Although the electropherotype of SKRV in SDS-PAGE was similar to other aquareoviruses, individual molecular masses of the λ class polypeptides were slightly different. SKRV had 5 structural polypeptides of molecular mass ranging from 154 to 38 kDa with 2 large (λ1 and λ2), 1 medium (μ1) and 2 small (σ1 and σ2) virus structural proteins. Comparable electrophoretic profiles with the structural proteins falling into 3 size classes (λ, μ and σ) with similar molecular mass ranging from 145 to 32 kDa were present in aquareoviruses (Hedrick et al. 1984, Winton et al. 1987, Hsu et al. 1989, Marshall et al. 1990, Samal et al. 1990, 1991a).

Although a detailed study of the gene coding assignments and radiolabelling of viral proteins would be required to completely characterise the structural and non-structural proteins of SKRV, with the available information on viral polypeptides of orthoreoviruses with a similar genomic segmentation pattern, SKRV structural proteins were found similar to the existing classification pattern of the proteins of mammalian reoviruses. Virions of reovirus type 3 Dearing strain consists of 9 polypeptide species distributed in 3 size classes having a molecular mass ranging from 155 to 34 kDa (Joklik 1983c). Later studies have found that the molecular mass of the reovirus structural proteins ranged from 145 to 41 kDa (Nibert et al. 1996). Of the 9 structural proteins, 4 polypeptides were present in the virions in less than 2% each. Investigations of striped bass aquareovirus polypeptides and gene coding assignments have identified the presence of 7 structural polypeptides and 5 non-structural polypeptides associated with the virus (Subramanian et al. 1994).

Among the 5 structural proteins of SKRV, μ1 (65 kDa) and σ2 (38 kDa) are present in the highest concentrations. Proteins of the same size groups make up more than 60% of the mass of the reovirus type 3 Dearing strain and form the major structural proteins of the Orthoreovirus capsid (Smith et al. 1969). Similar size proteins (VP3 and VP5) also form the major structural proteins present in larger quantities in many aquareoviruses (Winton et al. 1987, Samal et al. 1991a). The nature of polypeptide composition of SKRV is different from orbiviruses that have 4 major and 3 minor polypeptides (Gorman et al. 1983, Joklik 1983c, Joklik 1983c, Hedrick et al. 1984, Winton et al. 1987, Samal et al. 1991a, Nibert et al. 1996), it could be inferred that the SKRV electropherotype has close similarity with that of the reovirus type 3 Dearing strain. The strength of the bands was also taken into consideration in drawing this conclusion, as the purified virions are reported to contain equimolar quantities of these 10 species of double-stranded RNA segments (Shatkin et al. 1968). The total size and M∞ of the SKRV genome (16.38 × 10⁶, 24.56 kbp) are similar to that of the reovirus prototype strain T3D (type 3 Dearing) which has a total size of 23.549 bp (Wiener & Joklik 1989). Genome size of SKRV was, however, different from the orbiviruses, which has a total genome length of only 19.2 kbp and molecular mass of 13 × 10⁶ (Roy 1996).

Similar to avian reoviruses, SKRV also differs from mammalian reoviruses in their inability to haemagglutinate human ‘O’ RBC erythrocytes (Kawamura et al. 1965, Joklik 1983c). One of the evident characters that differentiated SKRV from avian reoviruses bearing similar characteristics was the absence of syncitium formation in the cell lines infected with SKRV. A complete lack of antigenic relatedness between SKRV and the aquareovirus isolates in the α-cross neutralisation study indicates that SKRV shares no common neutralising epitopes with aquareoviruses and shows the generic difference between them (Ginsberg 1988). Since we do not have a polyacrylamide gel of the viral RNA to unambiguously prove the presence of a 10-segmented genome for the SKRV, this isolate could be considered as a new strain of Aquareovirus at present. However, if this isolate does contain only10 RNA segments, SKRV would be the first report of the isolation of a reovirus from a poikilothermic host that belongs to the genus Orthoreovirus.
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