NOTE

Isolation of IPN virus serotype VR-299 from turbot in Europe

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ABSTRACT: Serological and molecular properties of 2 birnaviruses isolated from Norwegian turbot \textit{Scophthalmus maximus} were examined. The viral RNA and polypeptides patterns, as well as neutralization test results, showed that these strains were closely related to infectious pancreatic necrosis virus, serotype VR-299. Infectivity trials indicated no pathogenicity for turbot. This is the first isolation in Europe of VR-299, the common North American virus strain of IPNV.

KEY WORDS: Birnavirus - IPNV - \textit{Scophthalmus maximus} - turbot

Serologically, IPNV is a heterogeneous aquatic birnavirus and all the isolates show some degree of cross-reactivity (Wolf 1988). Using polyclonal antibodies, 3 major serotypes (Sp, Ab and VR-299) plus 7 additional serotypes (He, Te, Ja, C1, C2, C3 and N1) have been established. These 10 serotypes constitute serogroup A; 4 other isolates from marine shellfish represent serogroup B (Hill & Way 1983, Christie et al. 1988).

All aquatic birnaviruses isolated from fish in the USA belong to serotype VR-299, while 4 serotypes (Ja, C1, C2 and C3) have been found in Canada. In Asia, isolates related to Sp, Ab or VR-299 serotypes have been reported. While the Ab serotype has been isolated from a variety of fish and shellfish species, most VR-299 serotype viruses from Asia have been limited to salmonids (Hedrick et al. 1985, Wattanavijan et al. 1988, Lipipun et al. 1989).

In Europe all the birnaviruses isolated from freshwater or marine fish belonged to serotypes Sp, Ab and He. The birnaviruses isolated in our laboratory since 1985 from trout, salmon and turbot with different European origins, or from our own production stock, belonged to serotypes Sp and Ab or were untypable strains (Ledo et al. 1990, 1992, Novoa et al. 1991, 1992). All these IPN viruses were isolated during routine virological surveys or from diseased fish collected for examination.

Until 1988, only IPNV serotypes Ab and Sp had been isolated from fish in Norwegian salmonid farms (Håstein & Krogsrud 1976, Krogsrud et al. 1989). A new IPNV serotype N1 has been reported from Norway (Christie & Havarstein 1989) which produced high losses in Atlantic salmon \textit{Salmo salar}, turbot \textit{Scophthalmus maximus} L. and halibut \textit{Hippoglossus hippocoglossus} L., and was recovered from scallops \textit{Pecten maximus} L. (Mortensen et al. 1990). In further epidemiological studies conducted in farms located along the Norwegian west coast, different IPN viruses were isolated and tested by ELISA using monoclonal antibodies. Most of them were also assigned to the new serotype N1 (Christie et al. 1990).

This report describes the first isolation of IPNV VR-299 from turbot in Europe. The identification and characterization of 2 birnaviruses (TB-311 and TB-160) indicated that they belonged to VR-299, the typical North American serotype of this virus.

Materials and methods. Virus isolation and physical-chemical characterization: During 1988 and 1989 imported turbot (adults and juveniles) were sent to our laboratories for virological analysis. Samples were processed for virus isolation following standard virological procedures (Amos 1985). The Chinook Salmon Embryo cell line (CHSE 214; Lannan et al. 1984) was used for all virus assays. The fish cells were cultivated at 15 °C in Eagle’s minimal essential medium (MEM) supplemented with 10% fetal bovine serum and containing 100 IU penicillin and 100 μg streptomycin ml\textsuperscript{-1}.

Viral RNA electrophoresis: The RNA segments of the 2 birnaviruses from turbot were extracted and analyzed in polyacrylamide gels. Concentrated virus (pelleted at 100000 \texttimes \textit{g} for 1 h) was treated with protei-
nase K (200 μg ml⁻¹) at 37 °C for 30 to 60 min. Samples were mixed with sample buffer, boiled for 5 min and analyzed in SDS-PAGE (Laemmli 1970) using 7.5 % polyacrylamide gels, that were run for 3 h at 100 V in a Mini Protean II Electrophoresis System (BIO-RAD, USA). The RNA bands were visualized by staining with silver nitrate. The reference strains of IPNV (Sp, Ab, VR-299 and N1) together with the strain TB-306, an atypical Ab related virus isolated from turbot (Novoa et al. 1992), were included in the same gel for comparison. The RNA segments of a characterized aquareovirus from turbot (TRV) (Dopazo 1991) were used as molecular weight markers.

**Analysis of virion polypeptides:** CHSE-214 cells were infected at a multiplicity of infection of 1 virus cell⁻¹ with the IPNV strains. After 1 h of adsorption, MEM with 2 % FBS was added and the uncovered plates were irradiated with UV light (44 ergs s⁻¹ mm⁻² at 25 cm) (Dobos & Rowe 1977). Five hours later, 10 μCi ml⁻¹ of [³⁵S]-methionine (sp. act. 1000 Ci mmol⁻¹) in methionine-free medium was added for 5 h. Then, the cells were washed 3 times with PBS and suspended in 200 μl of 0.1 N NaOH, 1 % SDS and 200 μl of sample buffer. The preparations were subjected to SDS-PAGE on a 12 % polyacrylamide gel for 5 h at 11 mA. The gel was treated twice with dimethylsulfoxide (DMSO) for 30 min, once with 20 % of 2,5-Diphenyloxazole (PPO) in DMSO for 3 h and then washed with water. The gel was then dried and autoradiography was done on X-ray film (Kodak, USA).

**Neutralization assays:** The possible relationship of the turbot isolates to 4 serotypes of IPNV (Sp, Ab, VR-299 and N1) was examined using specific rabbit antisera against the different serotypes. The neutralizing antibody titers were calculated by the method of Reed & Muench (1938) and expressed as the reciprocal of the highest dilution of antiserum protecting 50 % of the inoculated wells.

**Experimental infections:** Infectivity trials were conducted in order to determine the pathogenicity of the isolated strains (TB-311 and TB-160). For comparative purposes, the North American reference serotype VR-299 was included. Turbot (1 g), previously tested for virus, were injected intraperitoneally with 50 μl of several doses (8 × 10⁷, 8 × 10⁶, 8 × 10⁵ TCID₅₀ ml⁻¹) of each viral suspension; 8 fish were used per dose. The controls were inoculated with culture medium. Fish were maintained for 30 d at 15 °C in 25 l tanks using recirculated seawater with aeration. In all experiments the control and infected fish were treated the same. Dead and surviving fish were collected for virus assays.

**Results and discussion.** Two viruses (strains TB-311 and TB-160) were isolated from turbot and identified
as birnaviruses according to their physical and chemical characteristics: resistance to chloroform treatment, yellowish-green cytoplasm of infected cells stained with acridine orange which is indicative of a doubled-stranded virus, neutralization by antisera against IPN virus and a bisegmented RNA genome (Fig. 1). Whereas the TB-160 strain was obtained from apparently healthy fish of 3 to 4 g, TB-311 was found in haemorrhagic internal organs of brood stock, which suffered high mortalities.

The results of the neutralization assays are summarized in Table 1. High neutralization titers were obtained with VR-299 antiserum (14125 for TB-160 and 44668 for TB-311) and the reference strain VR-299. The 4 reference strains were neutralized more specifically by their respective antisera. The 4 selected reference serotypes of IPNV could be distinguished by the relative mobilities of the viral RNA as well as the viral polypeptides. The size of the RNA segments of the viral isolates from turbot were identical to those of VR-299 (1.9 and $2.1 \times 10^6$ daltons) (Fig. 1) and the polypeptides patterns of the turbot strains were also similar to those of VR-299 (Fig. 2).

No mortalities were recorded in turbot injected with
the 2 strains of birnavirus during 30 d (LD_{50} values higher than 8 \times 10^7 TCID_{50}), indicating that, under experimental conditions, they were not pathogenic for turbot. The North American reference strain VR-299 was also not virulent for turbot. Only in the case of the turbot strain TB-311 was virus recovered from survivors, where the titer of 10^3 TCID_{50} g^{-1} of tissue suggested that a carrier state had been established. Studies are being conducted to determine if other fish species are susceptible to these turbot isolates.

The isolation of 2 viruses from fish in Europe with the characteristics of IPNV serotype VR-299 alters the established distribution of the basic serotypes of aquatic birnavirus. Traditionally, the Sp and Ab serotypes were considered to belong to European waters whereas the VR-299 was considered a North American strain. However an IPNV virus from southern flounder Paralichthys lethostigma with the characteristics of the Ab serotype has been reported in the USA (McAllister et al. 1984), and viruses belonging to the serogroup II (usually found in marine shellfish) were isolated from salmonids and marine fish (Olesen et al. 1988, Ahne et al. 1989). To date, the birnaviruses isolated from fish and molluscs cultured in our area were predominantly Sp strains (Ledo et al. 1990, Dopazo 1991, Rivas et al. 1992). Moreover, birnaviruses recovered from imported stocks of turbot from European countries belonged to the Sp. Ab serotypes or were untypable strains (Ledo et al. 1990, Novoa et al. 1992, unpubl., data). Similar results were obtained when wild fish and sediments were monitored to detect reservoirs of IPN viruses (Rivas et al. 1992). Castric et al. (1987) associated the Ab serotype with mortalities of turbot in France. Therefore, the presence of the VR-299 serotype in turbot from farms surrounded by floating cages of salmon, in which this serotype was never detected, is striking. Although the origin of VR-299 in Europe is unknown, as is the case with the N1 strain (Mortensen et al. 1990), transmission from feral animals, i.e. molluscs, crustaceans, cannot be ruled out.

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