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Characterization of Three Continuous Cell Lines from Marine Fish

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Abstract.—Three continuous cell lines were established: JSKG from gonads of Japanese striped knife jaw Oplegnathus fasciatus, KRE from embryos of a hybrid of kelp Epinephelus moara and red spotted grouper E. akaara, and PAS from the skin of greater amberjack (also called purplish amberjack) Seriola dumerili; these cell lines were passed 60, 89, 120 times, respectively. Although initially cultured in Leibovitz's L-15 medium, two of the cell lines, JSKG and PAS, exhibited optimal growth response in Eagle's minimum essential medium buffered with a combination of tris and sodium bicarbonate. These cell lines were initiated at a higher NaO concentration of 0.206 M but gradually adapted to the low NaCl concentration of 0.116 M after several subcultures. Optimum growth temperature was 25°C for JSKG and PAS cells, and 30°C for KRE cells. The modal chromosome number is 83 for the JSKG cell line, 92 for the KRE cell line, and 96 for the PAS cell line. Results for efficiency of plating indicate that all three cell lines are composed of transformed cells. Cell lines JSKG and PAS are susceptible to nine fish viruses, including channel catfish virus (CCV) and chum salmon virus (CSV). The KRE cell line is susceptible to CCV and fish rhabdoviruses of the vesiculovirus group. None of the cells showed cytopathic effect for Oncorhynchus masou virus (OMV) or Herpesvirus salmonis. Yields of infectious pancreatic necrosis virus (IPNV), infectious hematopoietic necrosis virus (IHNV), hirame rhabdovirus (HRV), and CSV were relatively low in these cell lines.

There is a growing interest in the aquaculture of marine fishes because of the high economic value of these fishes. Concomitantly, there is a high risk of viral diseases occurring in these fish species, especially in intensive marine aquaculture systems. Consequently, susceptible cell lines need to be established from marine fish for use in viral diagnostics. Since the establishment of the first fish cell line, most efforts have been concentrated on the development of cell lines from freshwater and coldwater sport and aquaculture species (Clem et al. 1961; Wolf and Quimby 1969; Wolf and Mann 1980). These cell lines are now being used in the isolation and study of economically important fish viruses.

This article describes the establishment of three cell lines derived from the gonad, embryo, or skin of three species of marine fish: Japanese striped knife jaw Oplegnathus fasciatus, a hybrid of kelp Epinephelus moara and red spotted grouper E. akaara, and greater amberjack Seriola dumerili.

Methods

Primary Cultures and Maintenance

Cell lines were initiated from the gonads of Japanese striped knife jaw (JSKG cell line), embryos of a hybrid of kelp and red spotted grouper (KRE cell line), and skin of greater amberjack (also called purplish amberjack; PAS cell line), according to the methods of Wolf and Quimby (1976). Gonad tissue was aseptically removed from the donor fish; embryo and skin tissues were dipped in a 50-mg/L solution of iodophor (popidone iodine, Meiji Seika) for 15 min. These tissues were finely minced to a size of about 1 mm³ and suspended in a phosphate-buffered saline (PBS) with NaCl to give a final concentration of 0.206 M, and 0.25% trypsin.
was added for digestion. The suspension was poured into a digestion flask and tissues were digested overnight at 5°C. Undigested tissues were separated by filtration through an 80-μm-mesh filter. Dispersed cell suspension was centrifuged at 5°C for 10 min at 200 \times gravity. The pellet was resuspended by gentle addition of several milliliters of the growth medium by pipette. Cell concentration was determined with a hemocytometer and adjusted to 3.5 \times 10^5 cells/mL with Leibovitz’s L-15 medium (GIBCO) supplemented with 20% fetal bovine serum (FBS, GIBCO). Cells were seeded into 25-cm² tissue culture flasks (Falcon) and maintained in Leibovitz’s L-15 medium with 20% FBS. Two antibiotics were added to the medium: 100 IU penicillin (Sigma) and 100 \μg streptomycin (Sigma) per milliliter. Sodium chloride in the medium was increased to obtain a concentration of 0.206 M, as recommended for establishing cell lines from marine fish (Clem et al. 1961). Primary monolayers were maintained in these media until the tenth passage, after which FBS in the medium was reduced to 10%. Likewise, after several passages, NaCl concentration was reduced to the normal concentration used in commercial media preparations, 0.116 M.

Detection of Cell Growth Response

Estimation of appropriate physiological and culture conditions for the three marine fish cell lines was based on cell growth responses. Responses of the cells were detected by the modified microplate staining technique. This technique was done by inoculating cells at a concentration of either 1 \times 10^5 or 2 \times 10^5 cells/mL in a 96-well microplate (Corning). After incubation, cells were fixed with 10% formalin for 30–60 min then washed thoroughly. Washed cells were stained with 0.1% crystal violet for 1–2 h, washed again, and air-dried (Nakajima et al. 1988). Absorbance of dried and stained microplates was read in a microplate spectrophotometer (Corona MTP-22) at 600 nm.

Analysis of variance was employed in statistical evaluation of data at significance levels of \( P \leq 0.01 \) and \( P \leq 0.05 \). Comparison of means was based on Duncan’s new multiple-range test (Wakimoto et al. 1984).

Growth Requirements

Culture media.—After several passes, JSKG, KRE, and PAS cell lines were tested for growth in three kinds of media: Eagle’s minimum essential medium with Earle’s salts (Eagle’s MEM, GIBCO), Leibovitz’s L-15 medium, and medium 199 (GIBCO). In Eagle’s MEM, three buffer systems were used: sodium bicarbonate alone (26 mM); sodium bicarbonate (8.9 mM) with tris (16 mM); and sodium bicarbonate (8.9 mM) with HEPES (14 mM). All media were supplemented with 10% FBS and antibiotics (100 IU penicillin and 100 \μg streptomycin per milliliter of the medium). After the initial seeding (1 \times 10^5 or 2 \times 10^5 cells/mL), all cell lines were incubated for 7 d at 20°C.

Temperature.—Optimum temperatures for growth of JSKG, KRE, and PAS cells were also determined. Cells were incubated at temperatures of 10, 15, 20, 25, 30, 35, and 40°C for 7 d. Medium for this experiment was chosen on the basis of results from the above experiment on suitability of culture media. Initial cell concentrations were the same as in the culture media test.

Sodium chloride.—Growth response of the fish cell lines to NaCl concentrations of 0.116, 0.171, 0.256, 0.341, and 0.512 M were likewise observed. Eagle’s MEM buffered with sodium bicarbonate and tris was adjusted to each of the above salt concentrations and used as growth medium for this test. Cells were incubated at 20°C for 7 d.

Chromosome Counts

Chromosome counts were determined for JSKG, KRE, and PAS cells at passage levels 57, 86, and 118, respectively. Semiconfluent and actively growing monolayer cultures of these cells, propagated for 24 h, were used. Chromosome counts were done according to the method of Earley (1975).

Plating Efficiency

Efficiencies of plating of JSKG, KRE, and PAS cell lines were determined at passage levels 57, 85, and 119, respectively. The ability of the cells to form colonies were determined after 15 d by the methods of Fryer et al. (1981). Epithelioma papulosum cyprini (EPC) cells, an established cell line from common carp Cyprinus carpio (Fijan et al. 1983), was used as the control.

Cytopathic Effects and Viral Replication

Cytopathic effects (CPE) in JSKG, KRE, and PAS cell lines were determined after infection with 11 fish viruses (Table 1). Stocks of the virus were prepared in chinook salmon embryo (CHSE-214) cells (Lannan et al. 1984), EPC cells, rainbow trout gonad (RTG-2) cells (Wolf and Quimby 1962), or
Table 1.—Fish viruses used for the production of cytopathic effect in JSKG, KRE, and PAS cell lines.

<table>
<thead>
<tr>
<th>Virus</th>
<th>Abbreviation (and virus type)</th>
<th>Source (or original reference)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Channel catfish virus</td>
<td>CCV</td>
<td>J. A. Plumb, Auburn University, Alabama</td>
</tr>
<tr>
<td>Chum salmon virus</td>
<td>CSV</td>
<td>Winton et al. (1981)</td>
</tr>
<tr>
<td>Eel virus of America</td>
<td>EVA</td>
<td>T. Sano, Tokyo University of Fisheries, Tokyo</td>
</tr>
<tr>
<td>Eel virus of Europe X</td>
<td>EVEX</td>
<td>T. Sano</td>
</tr>
<tr>
<td><em>Herpesvirus salmosis</em></td>
<td>H. salmosis</td>
<td>K. Wolf, National Fish Health Research Laboratory, Kearneysville, West Virginia</td>
</tr>
<tr>
<td>Hirame rhabdovirus</td>
<td>HRV</td>
<td>Kimura et al. (1986)</td>
</tr>
<tr>
<td>Infectious hematopoietic necrosis virus</td>
<td>IHNV NAV</td>
<td>J. R. Winton, National Fisheries Research Center, Seattle, Washington</td>
</tr>
<tr>
<td>Infectious pancreatic necrosis virus</td>
<td>IPNV VR 299</td>
<td>Yoshimizu et al. (1989)</td>
</tr>
<tr>
<td>Oncorhynchus masou virus</td>
<td>IPNV Ab</td>
<td>K. Wolf</td>
</tr>
<tr>
<td>Pike fry rhabdovirus</td>
<td>IPNV ChAb</td>
<td>K. Wolf</td>
</tr>
<tr>
<td>Spring viremia of carp virus</td>
<td>SVVCV</td>
<td>B. J. Hill, Fish Disease Laboratory, Weymouth, UK</td>
</tr>
</tbody>
</table>

* If the author of the original reference was also the source, the reference is noted.

channel catfish ovary (CCO) cells (Bowser and Plumb 1980a), and each virus was inoculated at a multiplicity of infection of 0.01 into two wells for each of the three marine fish cell lines in 24-well plates. Cell lines KRE, JSKG, and PAS were observed for CPE for at least 2 weeks. The CPE produced in these marine cell lines were compared with those produced in three freshwater fish cell lines: eel ovary (EO-2) cells (Chen and Kou 1981), fathead minnow (FHM) cells (Gravell and Malsberger 1965), and EPC cells.

Cell lines that developed CPE for IPNV VR299, IHNV ChAb, HRV, CSV, OMV, or H. salmosis were further tested for virus replication efficiency. Stocks of virus were prepared in either CHSE-214, EPC, or RTG-2 cell lines, and virus concentrations were determined by assay for 50% tissue culture infective dose (TCID50/mL; Reed and Muench 1938). Each virus was inoculated (multiplicity of infection = 0.01) into a monolayer of JSKG, KRE, and PAS in a 25-cm² tissue culture flask; each virus–cell line combination was prepared in duplicate. To allow absorption of the virus, flasks were incubated for 1 h at 15°C, except for those with CCV, which were incubated at 20°C. Thereafter, except for cell lines inoculated with H. salmosis and CCV, all samples were incubated at 15°C; the samples with H. salmosis were incubated at 10°C, and those with CCV were incubated at 20°C. After a determined number of days, 100 μL from each flask were pooled for each duplicate set of flasks, and samples were assayed for virus titers (TCID50/mL).

Results

Primary Cultures and Morphology

Primary monolayer cells were obtained from the gonad, embryo, or skin of the three species of marine fish (Table 2) and all are confirmed as epithelioid cells (Figure 1).

Growth Requirements

Culture media.—Growth responses of the three cell lines to three kinds of media and three buffer systems in Eagle’s MEM were evaluated. The JSKG and PAS cell lines showed significantly higher (P < 0.05) absorbance readings in Eagle’s MEM buffered with a combination of tris and sodium bicarbonate (Figure 2) than in other culture media. The KRE cell line showed a significantly higher (P < 0.05) absorbance reading in Leibovitz’s L-15 medium.

Temperature.—The KRE and PAS cell lines generally exhibited a wide range of temperature tolerance from 15 to 30°C; optimum growth was

Table 2.—Origin and morphology of the new cell lines from marine fish.

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Species</th>
<th>Tissue of origin</th>
<th>Morphology</th>
</tr>
</thead>
<tbody>
<tr>
<td>JSKG</td>
<td>Japanese striped knife jaw</td>
<td>Gonad</td>
<td>Epithelioid</td>
</tr>
<tr>
<td>KRE</td>
<td>Hybrid of kelp and red spotted grouper</td>
<td>Embryo</td>
<td>Epithelioid</td>
</tr>
<tr>
<td>PAS</td>
<td>Greater amberjack (=purplish amberjack)</td>
<td>Skin</td>
<td>Epithelioid</td>
</tr>
</tbody>
</table>
FIGURE 1.—Monolayers of three new cell lines from marine fish: (A) JSDK; (B) KRE; (C) PAS.
at 30°C for the KRE cell line and at 25°C for the PAS cell line. Neither of these cell lines could tolerate 35 or 40°C. Although the JSKG cell line grew optimally at almost the same temperature as the PAS cell line, JSKG cells could not tolerate 30°C whereas PAS cells could (Figure 3).

*Sodium chloride.*—The growth of JSKG and PAS cell lines was inversely proportional to increasing NaCl concentration in the medium (Figure 4). Growth was significantly higher ($P < 0.05$) at the lowest NaCl concentration (0.116 M), the usual concentration used in commercial media preparations. Highest growth response for KRE cells was attained by using an NaCl concentration of 0.171 M.

**Chromosome Counts**

All three cell lines were heteroploid and consisted mostly of 4n cells (Table 3). The diploid number of the Japanese striped knife jaw, the hybrid of kelp and red spotted grouper, and the greater amberjack is 48 (Ojima 1983). In 71 ran-
domly selected cells of the JSKG cell line, the counted chromosome number ranged from 72 to 97, and the modal number was 83; in the KRE cells (N = 63) the chromosome number ranged from 84 to 111, and the modal number was 92; in the PAS cells (N = 73) the chromosome number ranged from 60 to 108, and the modal number was 96 (Table 3; Figure 5). Chromosomes of these cells were mostly metacentric.

Plating Efficiency

The plating efficiencies of the three new cell lines were compared with the plating efficiency of the EPC cell line. Plating efficiencies of the tested cell lines were relatively high and were comparable to those of EPC cells (Table 4). These results suggest that JSKG, KRE, and PAS are composed of transformed cells.

Cytopathic Effect and Viral Replication

Development of CPE in JSKG, KRE, and PAS cell lines after inoculation with 11 fish viruses is presented in Table 5; for comparison, data of this type are also presented for three common freshwater fish cell lines. The JSKG and PAS cell lines supported replication of most of these viruses. None of these cell lines developed CPE after in-
occlusion with two herpesviruses, OMV and *H. salmons*. On the other hand, another herpesvirus, CCV, produced CPE in all three cell lines. The KRE cell line was susceptible only to fish rhabdoviruses belonging to the vesiculovirus group. Relatively low titers of IPNV VR299 and IHNV ChAb were observed in tested cell lines (Table 6; Figure 6). In the JSKG and PAS cell lines, CSV was initially detected 15 d after inoculation (Table 6) and CSV titers increased until day 23. The viral inoculum might have been too low to produce an immediately noticeable CPE.

**Discussion**

In this article we have described the establishment of three continuous cell lines from commercially important marine fishes. Since the 1980s, several cell lines from marine fish had been reported (Table 7). Although growth was initiated in Leibovitz's L-15 medium, growth responses of JSKG and PAS cell lines were higher in Eagle's MEM buffered with a combination of Tris and sodium bicarbonate. Nicholson et al. (1987) reported the use of Eagle's MEM in the establishment of cell lines from Japanese seaperch *Lateolabrax japonicus* and a grouper, *Epinephelus amblycephalus*.

Clem et al. (1961) revealed the beneficial effects of higher NaCl concentrations in initiating cell lines from marine fish species by the trypsinization method. Use of higher NaCl concentrations led to the successful propagation of the grunt fin (GF) cell line. For a cell line from yellowtail scad (also called omaka) *Caranx mate*, osmolarity was not a critical factor (Lee and Loh 1975) and the cells were successfully subcultured at normal NaCl.

**Table 3.—Modal chromosome numbers of the new cell lines from marine fish.**

<table>
<thead>
<tr>
<th>Cell line (pass number)</th>
<th>Species of origin (diploid number)*</th>
<th>Modal chromosome number</th>
</tr>
</thead>
<tbody>
<tr>
<td>JSKG (57)</td>
<td>Japanese striped knife jaw (2n = 48)</td>
<td>83</td>
</tr>
<tr>
<td>KRE (86)</td>
<td>Hybrid of kelp and red spotted grouper (2n = 48)</td>
<td>92</td>
</tr>
<tr>
<td>PAS (118)</td>
<td>Greater amberjack (=purplish amberjack) (2n = 48)</td>
<td>96</td>
</tr>
</tbody>
</table>

*Ojima (1983).*

**Table 4.—Plating efficiencies for the new marine fish cell lines and a freshwater fish cell line (EPC).**

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Cells seeded/25-cm² flask</th>
<th>Average number of cell colonies</th>
<th>Efficiency of plating (%)a</th>
</tr>
</thead>
<tbody>
<tr>
<td>EPC</td>
<td>10³</td>
<td>220</td>
<td>22.0</td>
</tr>
<tr>
<td></td>
<td>10²</td>
<td>23</td>
<td>23.0</td>
</tr>
<tr>
<td>JSKG</td>
<td>10³</td>
<td>110</td>
<td>11.0</td>
</tr>
<tr>
<td></td>
<td>10²</td>
<td>7</td>
<td>7.0</td>
</tr>
<tr>
<td>KRE</td>
<td>10³</td>
<td>298</td>
<td>29.8</td>
</tr>
<tr>
<td></td>
<td>10²</td>
<td>23</td>
<td>23.0</td>
</tr>
<tr>
<td>PAS</td>
<td>10³</td>
<td>176</td>
<td>17.6</td>
</tr>
<tr>
<td></td>
<td>10²</td>
<td>22</td>
<td>22.0</td>
</tr>
</tbody>
</table>

a (Number of cell colonies/cells seeded) × 100.
concentration of 0.137 M. Likewise, Nicholson et al. (1987) and Meguro et al. (1991) reported the establishment of marine fish cell lines without additional NaCl in the medium. In the establishment of the three cell lines in the present study, higher NaCl was used during primary propagation; however, increased levels of NaCl in the medium were not necessary in the successive transfers, and the cells adapted readily to the usual NaCl concentration used in commercial media preparations.

One of the advantages of cell lines that grow at a wide temperature range is their potential suitability for isolation of both warmwater and coldwater fish viruses (Nicholson et al. 1987). All three cell lines were able to grow at temperatures from 15°C up to 25 or 30°C and were also sensitive to viruses with optimum replication at 15°C. Likewise, the versatility of these new marine fish cell lines is evidenced by their ability to replicate CCV, which is a highly cell-specific virus, and a reovirus from chum salmon, CSV (Winton et al. 1981). CCV is known to replicate only in cell lines from ictalurid and related silurid species such as brown bullhead (BB), channel catfish ovary (CCO), and walking catfish kidney (KIK) cells, at high temperatures (i.e., 25–30°C; Bowser and Plumb 1980b; Wolf 1988). The JSKG and PAS cell lines produced initially low titers for CSV. These titers were not detectable until day 15 after inoculation, but they increased continuously until day 23 of incubation. Yoshimizu et al. (1988) reported replication of CSV in nonsalmonid freshwater fish cell lines. Studies conducted by Winton et al. (1981) showed replication of CSV at various degrees of CPE and moderate levels of virus titers in nonsalmonid lines. This article is probably the first report of CSV replication in marine fish cell lines. None of the three cell lines was able to replicate OMV and H. salmons. Both of these viruses are known to replicate only in salmonid cell lines (Yoshimizu et al. 1988).

Cell lines JSKG, KRE, and PAS were passed 60, 89, and 120 times, respectively, and chromosome counts revealed that these cell lines mostly consist of 4n cells. Moreover, plating efficiencies of the three cell lines that were established in this study suggest that all are composed of transformed
cells. Transformed cells are capable of growing from a dilute inoculum (Paul 1972), whereas non-transformed cells can form colonies with frequencies of only 0.1–2% after seeding at low densities (Fryer et al. 1981). Also it was reported that transformed mammalian cells are capable of producing clones of cells at high efficiency when seeded at very low densities in MEM-10 medium. Based on the above findings, the three marine fish cell lines that were established here can be considered as permanent lines. These cell lines were derived from marine species that are of economic importance, especially in Japan. Because these lines offer a wide range of susceptibility to most well-known virulent fish viruses, they can be useful tools in studying undetected viruses in marine species.

References

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(Plecoglossus altivelis). Diseases of Aquatic Organisms 1:207-217.
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