

antigen-dependent cells (ADC) is considered to be 620-2247. \* Corresponding author. Tel.: +34 (1) 620-2300; Fax: +34 (1) 1994). Thus, in mammals, *in vitro* proliferation of 0022-1759/97/\$17.00 Copyright © 1997 Elsevier Science B.V. All rights reserved.  
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doviruses (MacFarlan et al., 1984; Burkhardt et al., 1992). Some of those assays also included rabies 620-2247.

activities (Kontsekova et al., 1992; Kultubudin et have been used as an assay for mammalian T-cell 994). Some of those assays also included rabies 620-2247.

Antigen-induced immunoproliferation assays of 1992). Some of those assays also included rabies 620-2247.

leucocytes obtained from virally immunized or vac- 1992). Some of those assays also included rabies 620-2247.

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lymphocyte-like cell lines have yet been reported 1992). Some of those assays also included rabies 620-2247.

to date. To the best of our knowledge, no T- 1992). Some of those assays also included rabies 620-2247.

cell lines. Both of these have been obtained in 1992). Some of those assays also included rabies 620-2247.

T-lymphocyte-like cell lines have yet been reported 1992). Some of those assays also included rabies 620-2247.

from any fish. 1992). Some of those assays also included rabies 620-2247.

1990; Vallejo et al., 1991a,b) and B-cells (Miller et 1992). Some of those assays also included rabies 620-2247.

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Long-term cell lines of various leucocyte cell monocyte/macrophage series (Fasial and Ahme- clude, antigen-presenting cells of the cell lines which have been derived from fish in- Cell lines in such animals are poorly understood. of immunity in such animals are few examples of such cell lines in fish and, therefore, the functional aspects malians. However, there are few responses in mammals. Different aspects of the immune response in different types are valuable tools for the investigation of such long-term cell lines of the various leucocyte cell types are described in the present paper.

## 1. Introduction

**Keywords:** T-lymphocyte; Antigen-dependent cell proliferation; Viral hemorhagic septicemia virus; Glycoprotein G4; In vitro method; Trout

We describe a methodology to obtain the trout T-lymphocyte-like cell cultures showing *in vitro* antigen-dependent proliferation to research on fish vaccination and viral antigen-presentation mechanisms. That would facilitate further studies on the comparative and developmental immunology of lower vertebrates. It could also be applied to research on fish vaccination and viral antigen-presentation mechanisms.

For long-term culture, and to overcome the absence of inbred trout kidney and periodically used as viral antigen-presenting cells (called G4-pulsed Ad cells) to autologous ADC cultures. The methodology to obtain trout ADC lines constitutes a new tool that would facilitate further studies on the comparative and developmental immunology of lower vertebrates. It could also be applied to research on fish vaccination and viral antigen-presentation mechanisms.

Cells were obtained and maintained from each individual trout kidney and periodically used as viral antigen-presenting cells (called G4-pulsed Ad cells) to autologous ADC cultures. The methodology to obtain trout ADC lines constitutes a new tool that would facilitate further studies on the comparative and developmental immunology of lower vertebrates. It could also be applied to research on fish vaccination and viral antigen-presentation mechanisms.

Non-infected control trout kidney cells incubated in the presence of G4, confirming previously published observations, from which we used a yeast recombinant form of the glycoprotein G (G4) of VHSV. No similar ADC cultures could be obtained and maintained from each individual trout kidney and periodically used as viral antigen-presenting cells (called G4-pulsed Ad cells) to autologous ADC cultures. The methodology to obtain trout ADC lines constitutes a new tool that would facilitate further studies on the comparative and developmental immunology of lower vertebrates. It could also be applied to research on fish vaccination and viral antigen-presentation mechanisms.

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**Abstract**

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From the trout

An *in vitro* method to obtain T-lymphocyte-like cells



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an assay for T-cells. Anamnestic immune proliferative responses to rhabdoviruses in fish have been mostly studied in trout using whole virus (Chilmonczyk, 1978), isolated viral proteins (Estepa, 1992; Estepa and Coll, 1992a,b), recombinant viral protein fragments (Estepa et al., 1994b) and viral peptides (Lorenzo et al., 1995a,b). No leucocyte proliferation could be obtained from non-infected control trout kidney donors treated in parallel with all those VHSV antigens (Estepa et al., 1994b; Lorenzo et al., 1995a,b). Furthermore, trout individually restricted the selection of the rhabdoviral peptides stimulating the leucocyte anamnestic proliferation (Lorenzo et al., 1995b), suggesting the existence of peptide-presenting capacities and T-cell proliferation with similar characteristics to those of higher vertebrates (Burkhart et al., 1994; MacFarlan et al., 1984). However, to confirm this, ADC need to be isolated. To develop such ADC in trout, we have used the viral haemorrhagic septicaemia virus (VHSV), because of the important economic impact of these rhabdoviruses in the international salmoniculture industry. We have concentrated on the glycoprotein G of VHSV because trout neutralizing antibodies show exclusive specificity for this antigen (Bernard et al., 1983; Engelking and Leong, 1989; Estepa et al., 1994a,b; Gilmore et al., 1988; Lorenzen et al., 1990; Olesen et al., 1991), a recombinant form of G (G4) confers some protection against *in vivo* VHSV challenge (Estepa et al., 1994a,b), the G protein is expressed in the membrane of infected cells (Estepa et al., 1992) and the G protein and peptides derived from it are the strongest inducers of anamnestic lymphoproliferation (Estepa and Coll, 1992a,b; Estepa et al., 1994a,b; Lorenzo et al., 1995a,b).

Since neither inbred nor syngeneic trout populations are at present widely available (Ristow et al., 1995) and the histocompatibility groups of the trout are not yet well defined (Stet and Egberts, 1991), it was necessary to establish a primary individual trout haematopoietic cell culture from each donor trout in order to have enough supply of putative antigen-presenting cells during a year-long experimental period (Diago et al., 1993; Diago et al., 1991; Estepa et al., 1994a). The successful application of this technique has been the key to establishing the T-lymphocyte-like trout cell lines defined as ADC described in this report.

## 2. Materials and methods

### 2.1. Virus and recombinant G4 VHSV protein

The VHSV 07.71 isolated in France (LeBerre et al., 1977) from rainbow trout *Onchorynchus mykiss* (Walbaum) was grown in epithelial papilloma cyprine (EPC) cells and purified as described (Basurco et al., 1991). The recombinant fragment G4 (aa 9-443) devoid of the transmembrane region and part of the signal peptide, was cloned and expressed in the yeast *Saccharomyces cerevisiae* DC04 as previously described (Estepa et al., 1994b).

### 2.2. Production of trout survivors of VHSV infection

Outbred trout (0.5–2 g body weight) were first infected over 2 h at 12–14°C with  $10^6$  TCID<sub>50</sub>/ml of VHSV attenuated by 10 passages on EPC cells (Basurco and Coll, 1992). Under these conditions, trout survival was 10–30% ( $n = 4$ ,  $n$  = number of experiments). The trout surviving the infection were challenged 2 months later with  $10^6$  TCID<sub>50</sub>/ml of VHSV (2 h, 10–11°C). As in previous studies (De-Kinkelin, 1988), 5–24% ( $n = 4$ ) of the initial number of trout survived both infections and showed no signs of VHS. Survivor trout were used 4–6 months after the last VHSV challenge (Enzmann and Konrad, 1993; Lorenzo et al., 1995a,b).

### 2.3. Establishment of antigen-dependent cell (ADC) cultures

Leucocytes from individual trout kidneys were obtained as described (Estepa and Coll, 1992a,b), to avoid any possible mixed leucocyte reactions (Stet and Egberts, 1991). Trout kidneys were chosen as a source of leucocytes because cells responding to the VHSV antigens (Estepa et al., 1994b) and to the VHSV peptides (Lorenzo et al., 1995a,b) were known to be present in this organ. Furthermore, large numbers of trout leucocytes could not be obtained from any other organ in mature trout (Chilmonczyk, 1978). Briefly, the kidney was removed from trout previously cooled down to 4°C, killed by cutting off the head and blood allowed to drain for ~5 min. The kidney was cut into pieces, dissociated with a Pasteur pipette, decanted and washed in cell medium.



G4. After eliminating the excess of G4, the G4-pulsed Ad cells were treated with mitomycin to inhibit proliferation, and then frozen until use. The non-Ad cell population was distributed in 96-well plates containing 100  $\mu$ l/well of culture medium (25 000 cells/well) and then stimulated by the addition of 5000 autologous G4-pulsed Ad cells in 50  $\mu$ l of cell culture medium. After 5 days, colonies of dividing cells and scattered cells were observed throughout the wells. These proliferating cells were called antigen-dependent cells (ADC) and could be maintained for approximately 1 year by periodically adding G4-pulsed Ad cells. In the presence of a constant num-

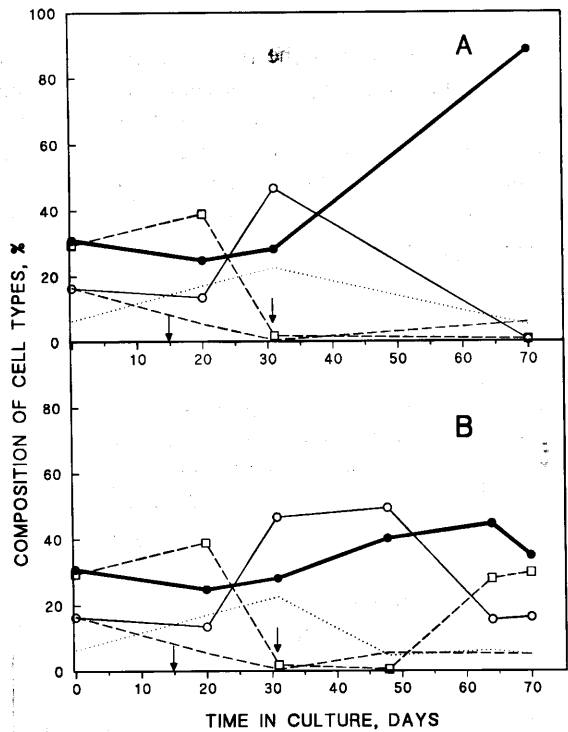


Fig. 2. Time course evolution of morphological cell types during the establishment of ADC cultures. The time course of the cellular types in the ADC cultures from T26 after the addition of its autologous G4-pulsed Ad cells was followed using cytocentrifuge preparations. After fixing and staining, the morphological cell types were classified using criteria defined previously (Estepa and Coll, 1992a,b): □—□, multinucleated; ○—○, eccentric nuclei; —, adherent cells; ····, large nucleated; ●—●, mature lymphocytes. About 200 cells were counted per slide and results were expressed as percentage of the total number of cells counted at each time point during culture. Time 0 and the vertical arrows indicate the time of addition of the G4-pulsed Ad cells. A and B show the time course evolution in two different experiments.

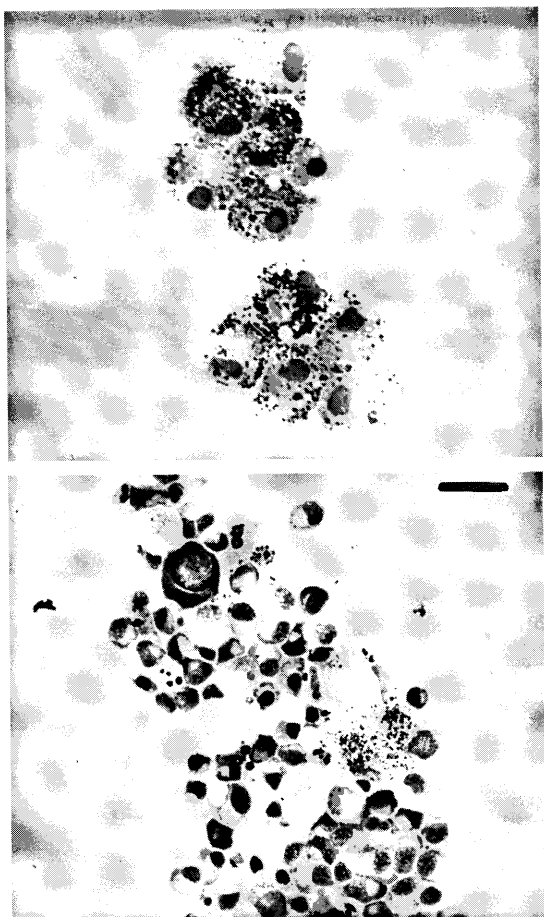


Fig. 3. Morphological appearance of G4-pulsed Ad cells (upper panel) and ADC (lower panel) in cytoslides. G4-pulsed Ad cells (25 000 cells/well) were added to resting ADC cultures (about 1 month after the last G4 stimulation). Cytocentrifuge preparations were obtained at 700 rpm (3 min), fixed for 10 min with 1% glutaraldehyde and stained with Toluidine blue in borate buffer. They were made permanent using Depex. Cytocentrifuge preparations of G4-pulsed Ad cells (upper panel) and proliferating ADC cultures, 15 days after the addition of G4-pulsed Ad cells (lower panel). Scale bar: ~10  $\mu$ m.

ber of G4-pulsed Ad cells, the ADC showed a cell-concentration dependence for proliferation that differed with the various trout used (Fig. 1). The minimum proliferation requirement for an initial ADC concentration has not yet permitted the cloning of these cells, despite numerous attempts.

After the first addition of G4 to whole kidney, many morphological cell types proliferated in the flasks (Fig. 2A,B). These cell types were of similar



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