Glycoprotein E"nS of Pestiviruses Induces Apoptosis in Lymphocytes of Several Species

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The family of the Flaviviridae consists of three genera: flavivirus, pestivirus, and hepatitis C virus. Flaviviridae are small, enveloped viruses with diameters of 40 to 60 nm. The genome is a single positive-strand RNA molecule of 12.5 to 16.5 kb. The genomes of flaviviruses and hepatitis C virus encode two envelope-associated glycoproteins: E1 and E2 (9, 22). In addition to E1 and E2, pestiviruses contain an extra envelope glycoprotein, which has been designated E"m (3, 24). The viral function of glycoprotein E"m, which is secreted in considerable amounts into the extracellular environment (18), is not yet clear. Recently, E"m was identified as an RNase (10, 19). Classical functions of RNases are the processing of RNA in gene expression and the digestion of dietary RNA. RNases can also exert cytotoxic activities and can be involved in, e.g., antitumor and immunomodulatory processes (4, 16, 23). Since pestivirus infections induce leukopenia and immunosuppression (2, 5, 15, 24), pestivirus infections are characterized by leukopenia and immunosuppression. Our results suggest that E"m plays an important role in the pathogenesis of pestiviruses.

Abbreviations. BVDV, bovine viral diarrhea virus; cf, control fraction; CSFV, classical swine fever virus; EBIT, embryonic bovine tracheal; EMEM, Earle’s minimal essential medium; MAb, monoclonal antibody; ncpm, net counts per minute; PBS, phosphate-buffered saline; PBMC, peripheral blood mononuclear cells; Sk, swine kidney; ELISA, enzyme-linked immunosorbent assay; E"m production and purification. Purified E"m was prepared as described previously (10). Briefly, Spodoptera frugiperda cells were infected with recombinant baculovirus expressing glycoprotein E"m of strain C (1). Infected cells were incubated for 96 h and then lysed in buffer containing 30 mM Tris-HCl (pH 7.5)–10 mM MgCl2–1% Nonidet P-40. The lysed cultures were centrifuged to remove cell debris, and the supernatant was stored at -70°C.

Control fractions were prepared from cells infected with wild-type baculovirus (cf1) and from cells infected with recombinant baculovirus expressing glycoprotein E"m (cf2). The control lysates were chromatographed over column material similar to that used for the E"m lysates.

Hybridoma cells producing MAb C5 were kindly provided by G. Wensvoort, and ascites fluid containing MAb C5 was produced as described by Wensvoort et al. (26).

Isolation of lymphocytes and the lymphocyte proliferation assay. Blood from specific-pathogen-free animals housed in isolation units and from humans was used for isolation of lymphocytes. From each species, the same animal or person was bled once for each successive experiment. Blood (10 ml) was collected in heparin-coated tubes, diluted 1:1 in PBS, layered onto 5 ml of Ficoll-Paque (Pharmacia Biotech), and centrifuged for 20 min at 20°C. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min. Human and porcine blood, ovine blood, and bovine blood were centrifuged at 500 g for 10 min.

Each culture of the lymphocyte proliferation assays was set up in triplicate, and the assays were performed at least twice. To each well of round-bottom microtiter plates containing 25 μl of cell suspension (105 PBMC), 25 μl of a dilution of E"m, a similar dilution of cf1 or cf2, or cell culture medium was added. The cultures were incubated at 37°C in 5% CO2 humidified atmosphere. Then, 50 μl of concanavalin A (type IV-S; Sigma) in RPMI was added to a final concentration of 5 μg/ml per ml. The plates were incubated at 37°C for 5% CO2 humidified atmosphere and then pulsed with 0.4 μCi of [3H]thymidine in 10 μl of RPMI per well. After 16 additional h, the cells were harvested with an automatic 96-well harvester (Tomtec) onto glass fiber filters. Filters were dried for 1 h at 60°C, and radioactivity was measured in a liquid scintillation counter (1450 Microbeta Plus; Wallac). The results were expressed as mean ncpm of the different assays, where ncpm is counts per minute of the triplicate stimulated cultures minus the mean counts per minute of the nonstimulated cultures.

To confirm the specificity of the inhibition by E"m, a blocking assay with MAb C5, which neutralizes virus infectivity, was performed. E"m and MAb C5 were incubated for 1 h at 37°C in a humidified atmosphere before being used in the lymphocyte proliferation assay. An antihemovirus herpesvirus type 1 MAb, MAb 41, was used as a control.

Cell viability. Cell viability after the concanavalin A-induced lymphocyte proliferation was determined with nigrinosine staining. Aliquots of cells were mixed with an equal volume of 0.2% nigrosine in PBS. Tests were performed in triplicate, and the percentages of viable cells were determined microscopically.
Protein synthesis assaying. Inhibition of protein synthesis was used to assay the direct cytotoxic action of E\textsuperscript{\textsc{ens}} (12). PBMC were isolated as described above, washed twice, and resuspended in leucine-free RPMI, without fetal calf serum and antibiotic cocktail, at a concentration of 2 × 10\textsuperscript{6} cells per ml. To each well of a round-bottom microtiter plate containing 50 µl of cell suspension (10\textsuperscript{6} PBMC), 50 µl of dilutions of E\textsuperscript{\textsc{ens}}, similar dilutions of cf\textsubscript{2}, or leucine-free RPMI was added. The plates were incubated in a 5% CO\textsubscript{2} humidified atmosphere for 1 to 24 h, and then 10 µl of RPMI containing 0.4 µCi of [\textsuperscript{3}H]leucine was added. Incubation was continued for 1 h. The cells were harvested on DEAE filters with an automatic 96-well harvester (Tomtec). The filters were dried for 1 h at 60°C, and radioactivity was measured in a liquid scintillation counter (Wallac 1450; Microbeta plus). Tests were performed in duplicate. The inhibition of the protein synthesis of E\textsc{btr} and SK\textsc{6} (13) by E\textsuperscript{\textsc{ens}} was also studied. These adherent cell types were separately cultured in 165-cm\textsuperscript{2} cell culture flasks (Costar), EMEM, 10% normal calf serum, and 0.5% antibiotic cocktail. The semicontinuous E\textsc{btr} cells were originally isolated from bovine embryonic trachea in our laboratory and are used routinely. The normal calf serum was collected from specific-pathogen-free calves and was BVDV and BVDV antibody free. For the protein synthesis assay, the adherent cells were trypsinized, washed in cell culture medium, and resuspended and 5 × 10\textsuperscript{6} cells per well were dispersed in 96-well flat-bottom microtiter plates. After 24 h at 37°C, and with 5% CO\textsubscript{2} in regular cell culture medium, the attached cells were washed twice with leucine-free EMEM. To each well, 100 µl of 0.4 µM E\textsuperscript{\textsc{ens}}, a similar dilution of cf\textsubscript{2}, or leucine-free EMEM was added. The plates were incubated at 37°C in a 5% CO\textsubscript{2} humidified atmosphere. After a defined incubation time, 10 µl of PBS containing 0.4 µCi of [\textsuperscript{3}H]leucine was added and the incubation was continued for 2 h. Then, the cells were washed, trypsinized, and harvested on DEAE filters. The filters were dried for 1 h at 60°C and counted in a liquid scintillation counter (1450 Microbeta Plus; Wallac). The degree of protein synthesis after incubation of the cells with E\textsuperscript{\textsc{ens}} or the control fractions is expressed as a percentage of the protein synthesis of the cells in leucine-free cell culture medium.

Detection of apoptosis. Apoptosis of bovine lymphocytes was determined by the cell death detection ELISA (Boehringer Mannheim [catalog no. 1544 675]), which is based on the specific detection of mono- and oligonucleosomes in the cytoplasmic fraction of apoptotic cells. The apoptosis in 100 µl of cell suspension (10\textsuperscript{6} PBMC) was determined after 8 h of incubation with 0.4 µM E\textsuperscript{\textsc{ens}}, a similar dilution of cf\textsubscript{2}, or RPMI 1640. Similarly, apoptosis of E\textsc{btr} cells was measured, except that these cells were trypsinized, because they were attached to flat-bottom microtiter plates. The results are expressed as the enrichment factor, which represents the enrichment of mono- and oligonucleosomes in the apoptotic cells and is the quotient of the average absorbance of the test sample and the average absorbance of an untreated sample.

RESULTS

Immunosuppressive effect of E\textsuperscript{\textsc{ens}}. We determined the immunosuppressive effect of pestivirus envelope glycoprotein E\textsuperscript{\textsc{ens}} in vitro. Immunoaffinity-purified E\textsuperscript{\textsc{ens}} was found to have a strongly inhibitory effect on the concanavalin A-induced proliferation of bovine lymphocytes. Incubation of the lymphocytes with a concentration of 0.4 µM E\textsuperscript{\textsc{ens}} completely inhibited [\textsuperscript{3}H]thymidine incorporation, and even 0.01 µM E\textsuperscript{\textsc{ens}} resulted in a slight but significant inhibition.

The specificity of the immunosuppressive effect on lymphocyte proliferation was confirmed in two different ways. To exclude the possibility that contaminants in the purified E\textsuperscript{\textsc{ens}} fraction caused the inhibitory effect, cf\textsubscript{1} and cf\textsubscript{2} were tested in dilutions similar to that used for E\textsuperscript{\textsc{ens}} in the lymphocyte proliferation assays and did not inhibit lymphocyte proliferation (Fig. 1). Additionally, blocking assays were performed with MAb C5, which is neutralizing and is directed against E\textsuperscript{\textsc{ens}}. The immunosuppressive effect of E\textsuperscript{\textsc{ens}} could be blocked partially by MAb C5. Incubation of 0.2 µM E\textsuperscript{\textsc{ens}} with MAb C5 (0.4 mg/ml) before addition to the lymphocytes resulted in an increase in proliferation of from 1 to 35% of the maximum proliferation. Incubation of E\textsuperscript{\textsc{ens}} with antibody herpesvirus type 1 MAb 41 (0.5 mg/ml) did not result in an increase in lymphocyte proliferation. These results of the lymphocyte proliferation assays show that E\textsuperscript{\textsc{ens}} is immunosuppressive for bovine lymphocytes in vitro.

The immunosuppressive effect of E\textsuperscript{\textsc{ens}} was then studied in proliferation assays with human, porcine, and ovine lympho-
cytes. Proliferation of lymphocytes of these species was also completely inhibited by 0.4 μM E<sup>em</sup>. From these results, we conclude that E<sup>em</sup> is immunosuppressive for lymphocytes of species in which pestiviruses normally replicate as well as for lymphocytes of species in which pestiviruses normally do not replicate. 

**Cytotoxic action of E<sup>em</sup>.** The viability of bovine lymphocytes during the proliferation assay was studied by nigrosine staining of aliquots of cells. At the start of the assay, all wells contained more than 90% viable lymphocytes. After 4 h of incubation, the control wells contained 93% and the wells incubated with 0.2 and 0.4 μM E<sup>em</sup> contained 85% viable lymphocytes. After 3 days of incubation, the control wells still contained 93% and the wells incubated with 0.2 μM E<sup>em</sup> contained 71% viable lymphocytes. None of the lymphocytes in the wells incubated with 0.4 μM E<sup>em</sup> were viable at that time (Fig. 2). The nigrosine staining showed death of lymphocytes after incubation with E<sup>em</sup>, which suggested a cytotoxic effect of E<sup>em</sup>.

The kinetics of lymphocyte protein synthesis was used to quantify cytotoxic effect. The effect of E<sup>em</sup> on the protein synthesis of bovine lymphocytes was determined. A dose-dependent inhibition of protein synthesis that already became obvious after 4 h of incubation was observed. Incubation with 0.4 μM E<sup>em</sup> resulted after 8 and 24 h in decreases in protein synthesis of 41 and of 63%, respectively. Concentrations of 0.1 and 0.025 μM also resulted in a decrease in protein synthesis at all measured time points. A concentration of 0.006 μM E<sup>em</sup> significantly enhanced instead of inhibited protein synthesis at all time points (Fig. 3).

E<sup>em</sup>-induced inhibition of protein synthesis was not specific for bovine lymphocytes. Protein synthesis of human, porcine, and ovine lymphocytes was measured after 8 h of incubation with 0.4 μM E<sup>em</sup>, and these results were similar to the results with the bovine lymphocytes. After investigation of the species specificity, we also studied the cell type specificity of the cytotoxic effect of E<sup>em</sup>. We selected two epithelial cell lines in which pestiviruses normally replicate and measured protein synthesis after 8 h of incubation with 0.4 μM E<sup>em</sup>. In contrast to the results for the lymphocytes, we did not detect an obvious inhibition of protein synthesis in these cell types. In EBTr and SK6 cells, a decrease of 5% and an increase of 8%, respectively, were measured. Incubation with cf2 did not affect protein synthesis.

These results indicate that E<sup>em</sup> is cytotoxic for lymphocytes of different species and not for epithelial cells.

**Apoptosis.** Virus-induced apoptosis of infected cells has been described for several virus species, including pestiviruses (29). We showed that purified envelope glycoprotein E<sup>em</sup> induced apoptosis in bovine lymphocytes. Lymphocytes were incubated for 8 h with E<sup>em</sup> and then tested in the cell death detection ELISA. After incubation with E<sup>em</sup>, the amounts of mono- and oligonucleosomes in the lymphocytes, which indicate apoptosis of the cells, increased. The average enrichment factor after E<sup>em</sup> incubation compared to that for incubation in regular cell culture medium was 3.9 ± 1.2. The control fraction cf2 did not induce apoptosis, as shown by an enrichment factor of 1.0 ± 0.1. Although it has been described that the cytopathic effect of pestiviruses is mediated by apoptosis (29), we did not detect apoptosis of EBTr cells after E<sup>em</sup> incubation. The enrichment factor for mono- and oligonucleosomes in these cells was 1.1 ± 0.02.

**DISCUSSION**

In this study, we investigated the immunomodulatory properties of CSFV envelope glycoprotein E<sup>em</sup> in vitro. We used the E<sup>em</sup> produced in recombinant baculovirus-infected cells because it has the same characteristics as those of E<sup>em</sup> from CSFV-infected cells (10, 19). E<sup>em</sup> inhibited concanavalin A-induced proliferation of bovine lymphocytes very efficiently.
The reduction of the [3H]thymidine incorporation was dependent on the concentration of E\textsuperscript{\textit{rns}} in the lymphocyte cultures and could be partially blocked by a neutralizing MAb directed against E\textsuperscript{\textit{rns}}. Hulst (11) showed that E\textsuperscript{\textit{rns}} binds very strongly to the cell surfaces of bovine and porcine cells and that binding is probably irreversible. The binding of E\textsuperscript{\textit{rns}} to MAb C5 is probably less strong and probably reversible. The stronger avidity of binding of E\textsuperscript{\textit{rns}} for a cell surface receptor compared to that of MAb C5 might explain the only partial neutralization of the effect of E\textsuperscript{\textit{rns}}. Nigrosine staining of aliquots of stimulated lymphocytes, which indicates damage of the cell membrane, showed cell death after E\textsuperscript{\textit{rns}} incubation. The percentage of dead lymphocytes was dependent on the concentration of E\textsuperscript{\textit{rns}} and on the incubation time. To quantify the direct cytotoxic effect of E\textsuperscript{\textit{rns}} on the lymphocytes, we performed the protein synthesis assay. After 4 h of incubation, the inhibition of the protein synthesis was significant for all studied concentrations of E\textsuperscript{\textit{rns}}, except for the lowest concentration tested. This concentration had a stimulatory effect. After 4 h of incubation, the nigrosine staining did not yet show a significant increase in the percentage of dead cells. Thus, E\textsuperscript{\textit{rns}} induces a significant inhibition of the protein synthesis of lymphocytes before the cell membrane becomes permeable for nigrosine. Therefore, the cytotoxic effect of E\textsuperscript{\textit{rns}} is probably not caused by direct damage of the cell membrane. The hypothesis that E\textsuperscript{\textit{rns}} does not damage the cell membrane directly is further supported by the fact that a concentration of 0.006 \( \mu \)M E\textsuperscript{\textit{rns}} enhanced instead of inhibited protein synthesis.

Although CSFV infection is normally restricted to swine, no species specificity was detected for the cytotoxic effect of E\textsuperscript{\textit{rns}} against lymphocytes. The levels of inhibition of the lymphocyte proliferation and of protein synthesis were comparable for all tested species, including humans. It was rather unexpected that this viral protein had such a strong immunosuppressive effect on human lymphocytes because pestiviruses normally do not infect humans. It may be speculated that viruses that induce immunosuppression in humans carry a protein with a function similar to that of E\textsuperscript{\textit{rns}}.

Pestiviruses preferably replicate in lymphoid tissue but also replicate in epithelial cells. We used SK6 and EBT\textit{r} cells to investigate possible cytotoxic properties of E\textsuperscript{\textit{rns}} against epithelial cells. No cytotoxic action was detected against the epithelial cells. Hence, the cytotoxic action of CSFV E\textsuperscript{\textit{rns}} in vitro is directed mainly against lymphocytes and is not restricted to cells of the natural host. Pestivirus envelope glycoprotein E\textsuperscript{\textit{rns}} is defined as an RNase, and the catalytically important amino acid residues are highly conserved within the pestivirus genus (10, 19). The function of E\textsuperscript{\textit{rns}}, which is secreted from infected cells (18), in the viral life cycle is not yet clear. From other RNases, it is known that they can have different noncatalytic biological actions, in addition to ribonucleolytic activity. They have been shown to be involved in antitumor, neurotoxic, helminotoxic, and immunomodulatory processes (4, 14, 16, 23). A specific cytotoxic action of E\textsuperscript{\textit{rns}} against lymphocytes is consistent with the pathogenesis of pestivirus infections. Acute virulent and persistent CSFV infections are characterized by severe leukopenia and immunosuppression (21, 25). In other pestivirus infections, immunosuppression is also a common phenomenon (2, 3, 15).

Immune deficiency caused by apoptosis of lymphocytes has been described previously for a number of viruses such as human immunodeficiency virus (7), influenza viruses (8), and feline leukemia virus (17). Our results from the protein synthesis assay and nigrosine staining suggested that apoptosis in lymphocytes was induced by glycoprotein E\textsuperscript{\textit{rns}} of CSFV. Indeed, in the cell death detection ELISA, it was clearly shown...
that E\textsuperscript{ns} induced apoptosis in lymphocytes. To our knowledge, this is the first time that selective apoptosis of lymphocytes induced by a purified viral glycoprotein has been described.

In our experiments, E\textsuperscript{ns} induced apoptosis in lymphocytes and not in epithelial cells, but Hulst (11) showed that E\textsuperscript{ns} is involved in virus-cell interaction of several epithelial cells. It interacts with the cell surfaces of PK15, SK6, fetal bovine epithelium, SF21, and CL626 cells, probably by binding to a common cell surface receptor. Since E\textsuperscript{ns} only slightly inhibited protein synthesis of EBT\textsubscript{r} and even slightly enhanced protein synthesis of SK6 cells, we conclude that binding of E\textsuperscript{ns} to a cell surface does not necessarily give a cytotoxic effect or induce apoptosis. After binding of an RNase to a cell surface, several processes can occur. It can remain attached to the cell surface receptor without any effect on the cell. It can damage the cell membrane directly and cause cell necrosis (28). Activation of the cell surface receptor can possibly induce apoptosis (27).

After binding, the RNase could also be internalized by membrane penetration or endocytosis. Then, degradation of RNA membrane directly and cause cell necrosis (28). Activation of receptor without any effect on the cell. It can damage the cell surface does not necessarily give a cytotoxic effect or induce apoptosis. Hence, our results raise the following question. Does E\textsuperscript{ns} induce apoptosis by activation of a cell surface receptor or is it internalized after binding?

Further studies are needed to elucidate the mechanism by which E\textsuperscript{ns} acts on lymphocytes and to clarify the role of E\textsuperscript{ns} in infected animals. It may be hypothesized that E\textsuperscript{ns} is secreted in the blood of infected animals and consequently may affect lymphoid tissue throughout the body, as seen in acute infections with CSFV and in mucosal disease of cattle. Taken together, our results have shown that E\textsuperscript{ns} inhibits lymphocyte proliferation and protein synthesis due to selective induction of apoptosis in the lymphocytes of several species. This effect suggests an important role for E\textsuperscript{ns} in the leukopenia and immunosuppression caused by pestivirus infections and might have implications for therapeutic use of E\textsuperscript{ns} as an immuno-suppressive agent.

REFERENCES


