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High-neurovirulence GDVII virus induces apoptosis in murine astrocytes through tumor necrosis factor (TNF)-receptor and TNF-related apoptosis-inducing ligand

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Abstract

We carried out a study to determine if the high-neurovirulence GDVII strain of Theiler’s murine encephalomyelitis virus (TMEV) and the demyelinating, low-neurovirulence BeAn strain induced apoptosis in murine astrocytes. Astrocytes, the major glial cell population of the central nervous system, were semipermissive for GDVII virus replication. Programmed cell death, demonstrated by apoptosis-specific caspase-3 protease activity, was maximal 8 h after GDVII infection at an m.o.i. of 1. Purified TMEV capsid proteins VP1, VP2, and VP3 did not induce apoptosis but antibodies to VP1 and VP2 inhibited it. Antibody inhibition of caspase-3 activity as well as flow cytometry experiments implicated TNF-related apoptosis-inducing ligand (TRAIL) and TNF-α-receptor (TNF-R) in apoptosis signaling. Conversely, TNF-α and the TRAIL-receptor were not upregulated. Furthermore, the number of functional TNF-α receptors, but not their affinity, was increased in apoptotic GDVII virus-infected astrocytes, as confirmed in binding experiments with 125I-labeled recombinant murine TNF-α.

In vivo studies showed that most of the cells loaded with the virus when injected in the brains of SJL mice were neurons but very few showed TUNEL costaining. Conversely, many of the apoptotic cells found were also positive for GFAP staining.

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Introduction

Programmed cell death or apoptosis is a highly conserved and controlled process that eliminates unwanted or damaged cells in multicellular organisms (Vaux and Korsmeyer, 1999). Induction of apoptosis in several cell types by viruses has been reported, including turkey spleen cells by avian adenovirus type II (Rautenschlein et al., 2000), mouse neuroblastoma cells by Langat flavivirus (Prikhod’ko et al., 2001), feline fibroblasts by feline immunodeficiency virus (Mizuno et al., 2001), HeLa cells by reovirus (Connolly et al., 2001), and Vero cells by avian coronavirus (Liu et al., 2001). The “altruistic suicide” of central nervous system (CNS) cells infected by viruses such as the alphaviruses, Semliki forest virus, and Sindbis virus, has also been demonstrated (Allsopp and Fazakerley, 2000).

Theiler’s murine encephalomyelitis virus (TMEV) is a picornavirus that persistently infects the murine CNS (Theiler, 1937). GDVII and BeAn viruses, representing the high- and low-neurovirulence groups, respectively, have been studied so far. Intracerebral inoculation of BeAn virus induces a chronic demyelinating disease in susceptible strains of mice that is reminiscent of human multiple sclerosis, whereas inoculation of GDVII virus causes an acute encephalitis with rapid demise (within 1 week) (Dal Canto and Lipton, 1976; Lehrich et al., 1976; Lipton, 1975).

BeAn virus induces apoptosis in cultured microglia but not in astrocytes (Zheng et al., 2001). Here we report that, consistent with the extensive cell death triggered within brain, GDVII virus is an inducer of apoptosis mainly in semipermissive astrocytes, although it also infects neurons upon intracerebral injection of mice. The apoptotic mecha-
nism involves tumor necrosis factor (TNF) receptors and the TNF-related apoptosis-inducing ligand (TRAIL), the same family of “cell suicide” inducers implicated in BeAn induction of apoptosis in other cellular systems (Jelachich et al., 1995, 1999; Jelachich and Lipton, 2001).

To demonstrate the pathological relevance of our in vitro results, we further established that intracerebral injection of GDVII virus induced apoptosis mainly in cerebral astrocytes around the injection site.

Results

Cytopathic effect and virus production in infected astrocyte cultures

As shown previously (Zheng et al., 2001), astrocyte cultures did not exhibit cytopathic effect (CPE) or loss of the normal polygonal flat morphology when infected with BeAn virus. Mock-infected cells maintained a flattened morphology with adherence to plastic. By contrast, GDVII virus infection induced CPE within 18–24 h in astrocyte monolayers. Although the percentage of infected cells is almost 100% in both primary and secondary cultures, as determined by the infectious center assay, the foci were more evident in secondary trypsinized cultures reaching 70–80% confluence than in primary, contact-inhibited cultures (not shown). Analysis of virus production by titration of infected astrocyte supernatants on BHK-21 cells demonstrated maximal titers of $5 \times 10^5$ PFU/ml in BeAn-infected astrocytes equivalent to the production of 0.2–1.2 PFU/cell. Titers two orders of magnitude higher ($10^7$), or 7–33 PFU/cell, were found in GDVII-infected cells (Fig. 1). Nonspecific binding of virus to the plastic of culture flasks without cells was not detected and the presence of residual virus remaining from the inoculum was ruled out (Fig. 1, circles).

Despite the low PFU output from BeAn-infected cells supernatants, our previous analysis by flow cytometry documented BeAn virus replication in the cytoplasm of astrocytes (Rubio and Martin-Clemente, 1999), and another recent study reported that BeAn virus is localized within the astrocytic cells, with little virus released into the supernatants (Zheng et al., 2001).

GDVII virus-induced apoptosis in astrocyte cultures

DNA laddering analysis has shown that BeAn virus does not induce apoptosis in astrocytes (Zheng et al., 2001). The ability of GDVII virus to induce apoptosis was assessed based on changes in caspase-3 activity, an enzyme with substrate specificity for the amino acid sequence Asp-Glu-Val1-Asp (Nicholson et al., 1995), since this enzyme is the main executioner caspase in a cascade of proteolytic cleavage events in dying cells. Astrocyte cultures infected at m.o.i.s of 1, 10, and 100 showed a significant increase in caspase-3 activity in comparison with mock-infected cultures ($P < 0.05$) (Fig. 2). This induction was specific since caspase-3 activity in cells infected at an m.o.i. of 10 and treated with 50 μM Z-VAD-FMK, the irreversible pan-caspase inhibitor, was not appreciably greater than that in mock-infected cultures. Caspase-3-specific activity was calculated based on a calibration curve using known amounts of chromophore p-nitroaniline (pNA), released from the substrate upon cleavage by the enzyme. Specific activity was calculated as picomoles of pNA liberated per hour per microgram of cell lysate extracts using the formula provided with the CaspACE kit. Mean specific activities were 96.5 ± 4.9 pmol/µg for GDVII-infected astrocytes and 24.3 ± 1.6

![Fig. 2. Caspase-3 activity in GDVII-infected astrocyte cultures.](image-url)

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pmol/µg for untreated cells. Maximal apoptosis was detected at an m.o.i. of 1. A well-defined positive control is provided by treating cultures with the strong apoptosis inducer Staurosporine (Fig. 2, +).

Analysis of the kinetics of caspase-3 induction in astrocyte cultures infected at an m.o.i. of 10 for periods ranging from 0 to 48 h revealed rapid induction of activity (4 h) that peaked after 8 h and decreased by 48 h (Fig. 3). We found around 30% of dead cells when the cultures were observed under phase-contrast microscopy at optimal conditions for apoptosis induction (i.e., 8 h at an m.o.i. of 1).

To exclude the possibility that cytotoxic factors in the BHK-21 cellular extract used as a source of virus induced the observed apoptosis, we tested various dilutions (from $10^{-2}$ to $10^{-5}$) of rabbit anti-TMEV antiserum, containing antibodies to the VP-1 and VP-2 capsid proteins (Clatch et al., 1987), for their ability to block apoptosis. Incubation with this antiserum completely abrogated the induction of caspase-3 activity (Fig. 4), indicating that apoptosis is dependent upon binding of the GDVII virus to receptors on astrocytes. This binding is presumably sterically inhibited upon binding of antibody to the virus. Preimmune rabbit serum incubated with virus at $10^{-2}$ dilution had no inhibitory effect on the apoptosis-inducing capacity of GDVII virus (Fig. 4).

**Purified virion proteins do not induce apoptosis**

Capsid proteins from CsCl-purified TMEV were isolated by reverse-phase high-pressure liquid chromatography (HPLC) on C-8 columns. VP1, VP2, and VP3 purified native protein were added to astrocyte monolayers in amounts corresponding to three-fold greater than the virus inoculum at an m.o.i. of 10, since the structural proteins are present in equimolar amounts in the TMEV capsid (Rozhon et al., 1985). None of the purified proteins induced significant caspase-3 activity (Fig. 5). The slight induction by VP2 might reflect some residual intact virus contamination in this particular preparation. Because the TMEV attachment site resides in a structure composed of both VP1 and VP3 proteins (Tyler, 1987), these two purified proteins were added together, as well as the three isolated proteins, without any appreciable induction of apoptosis (not shown). The lack of apoptosis inducing capability of the low-virulence BeAn strain in astrocytic cells is once more demonstrated (Fig. 5, BeAn).

**Upregulated TRAIL and TNF-R in infected astrocytes**

BeAn TMEV induces low-level secretion of TNF-α (Sierra and Rubio, 1993) and upregulates tumor necrosis factor-α-receptor (TNF-R) expression by 210% in astrocyte cultures (Aranguez et al., 1995). To determine whether GDVII virus-induced apoptosis involved signaling through TNF family members, the relative changes in expression of TNF-α and TRAIL and their respective receptors were examined in GDVII virus-infected cultures. Flow cytometry
revealed no increase in TNF-α and TRAIL-R in infected versus mock-infected cultures (not shown). Conversely, cytoplasmic TRAIL and surface TNF-R were both upregulated in GDVII-infected cultures (Fig. 6). Negative controls were provided by cultures mock-infected or infected with the nonneurovirulent strain BeAn (shaded profiles), or treated with control antibodies (thin lines). The increase in TRAIL and TNF-R was more evident in primary confluent astrocytes than in trypsinized, actively growing cultures (not shown).

TNF-α binding parameters in astrocyte cultures

Murine astrocytes exhibit a single binding site for TNF-α, and binding is specific and saturable since it was completely displaced by a 200-fold excess of unlabeled TNF (Aranguez et al., 1995). We compared the binding capacity of mock-infected astrocytes with GDVII-infected cultures (m.o.i. of 10) at 4 h when CPE was not detectable but caspase-3 was already induced (Fig. 3). As determined by Scatchard analysis, the $K_d$ of $^{125}$I-TNF binding in GDVII virus-infected astrocytes differed only slightly from that of mock-infected cells ($2.75 \times 10^{-10}$ M vs $3.0 \times 10^{-10}$ M). Nevertheless, 6385 receptors/cell were detected in infected cultures compared to 3258 receptor sites/cell in mock-infected cultures (Fig. 7). This 196% increase in the number of binding sites is essentially similar to a 208% increase previously reported for BeAn virus-infected astrocytes (Aranguez et al., 1995). The increase is consistent with the upregulated expression of TNF-R demonstrated by flow cytometry (Fig. 6).

Inhibition of apoptosis by antibodies

Analysis of caspase-3 induction in astrocyte cultures infected with GDVII virus at an m.o.i. of 10 in the presence of purified antibodies or antisera against TNF, TNF-R, TRAIL, and TRAIL-R revealed inhibition of enzyme activity only in the presence of anti-TNF-R and anti-TRAIL ligand (Fig. 8). However, the extent of inhibition never exceeded 40 to 60%, even at concentrations of antibody of 10 μg/ml, suggesting that both TRAIL and TNF-R provide different death-inducing signals. Normal rabbit serum or normal purified goat IgG (Fig. 8, −) does not induce any significant inhibition of caspase-3 activity. In one experiment, both antibodies were combined but no synergistic effect was found (not shown).

Identification of apoptotic astrocytes

Efficient TMEV growth has been detected previously in cultures of astrocytes (Rubio and Martin-Clemente, 1999; Zheng et al., 2001). Immunochemical staining of astrocytes in chamber slides for the astrocyte-specific GFAP marker...
revealed extensive cytoplasmic staining (Fig. 9A). TUNEL staining delineated relatively few nuclei (Fig. 9B), indicating that only some cells, that must be responsible of the caspase-3 activity, undergo apoptosis. Quantitative analysis of TUNEL-and GFAP-stained cells (Fig. 9C) using the analytical imaging station revealed a mean of 19% costained cells per field.

**Induction of apoptosis in the mouse brain**

To determine whether astrocytes as well as other brain cells undergo apoptosis after infection, the cortex, septum, nucleus accumbens, and anterior hypothalamus were examined 4–7 days after intracerebral inoculation of GDVII virus (2 × 10⁶ PFU). TMEV has been shown to infect astrocytes after intracerebral inoculation (Rubio and Martin-Clemente, 1999). Cytoplasmic GFAP (Fig. 10A) and nuclear TUNEL staining (Fig. 10B) on serial sections through the nucleus accumbens revealed colocalization of the two stains (Fig. 10C), indicating that GDVII virus induces apoptosis in astrocytes in vivo.

Consistent with previous results (Tsunoda et al., 1997), most of the infected cells in the brain were neurons based in morphological criteria as cells with big nuclei and round, nongranulated cell bodies with few processes. Both soma and neuropil were clearly labeled, as shown in the nucleus accumbens (Fig. 10D). This is not unexpected since previous studies have quantitated 15 × 10³ receptors per neuron compared with 2.5 × 10³ receptors per cell on astrocytes and oligodendrocytes (Rubio et al., 1990). Nevertheless, TUNEL staining was found in few infected neuronal nuclei, where the typical apoptotic chromatin condensation around the margin could be observed (Fig. 10E). Fig. 10F shows a stained nucleus, merged with the neuronal soma.

Control sections that omitted TdT for TUNEL, anti-TMEV, or anti-GFAP antibody were negative (not shown). Some remaining necrosis was found around the injection site in brains of control 6-week-old mice injected with 20 µl of DMEM, but no TUNEL staining was detected. No significant apoptotic events were detectable in healthy normal brains from age-matched SJL animals.

**Discussion**

Recent neurobiological evidence suggests a role for astrocytes, the major glial population of the CNS, beyond that of structural and trophic support of neurons (Nedergaard, 1994). Here we show that astrocytes undergo apoptosis when infected with the high-neurovirulence GDVII strain of TMEV as demonstrated by the induction of cysteine protease caspase-3 activity. The specific detection of the active form of caspase-3 is considered to accurately detect
apoptosis, free of interpretation bias and artifacts (Alnemri et al., 1996).

Our results are in accord with previous studies demonstrating that the GDVII virus is 50-fold more potent than the low-neurovirulence demyelinating BeAn strain in inducing apoptosis in BSC-1 African green monkey kidney cells (Jelachich and Lipton, 1996), and that BeAn induced apoptosis in microglia but not in mouse astrocytes, as measured by DNA laddering (Zheng et al., 2001). We also showed that, consistent with its neuropathogenic behavior in vivo, GDVII replicated and induced a cytopathic effect in astrocyte cultures.

Production of caspase-3 by astrocytes is a relatively early event, with maximal production at 8 h postinfection, slowly decreasing thereafter. The cell-permeable pan-caspase-specific inhibitor Z-VAD-FMK provided the control for the activity measured. The dose-dependent inhibition of caspase-3 by an anti-TMEV antiserum containing antibod-

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Fig. 9. Confocal images of astrocyte cultures infected for 24 h at an m.o.i. of 10 with GDVII. Staining for GFAP (A), TUNEL (B), or merged images (C) were shown. Scale bar, 30 μm.

Fig. 10. Sections of brain stained for GFAP (A), TUNEL (B and E), or replicating TMEV (D) at the level of the nucleus accumbens region 4 days after intracerebral inoculation of virus. In C and F, merged images of respective rows shows colocalization of staining in some cells. Scale bar in C and F, 30 μm.
ties to capsid proteins VP1 and VP2 demonstrated that the effect is not due to cytotoxic factors contained in the crude extract used as a source of virus.

Several viral proteins have been reported to induce or enhance apoptosis, including hepatitis C virus core protein (Zhu et al., 2001), hepatitis B HBx protein (Su et al., 2001), the 17-kDa protein from bursal disease virus (Yao and Vakharia, 2001), the adenovirus E4orf4 protein (Livne et al., 2001), the SV40 small t antigen (Gjoerup et al., 2001), and the matrix protein from vesicular stomatitis virus (Kopecky et al., 2001). Nevertheless, purified TMEV capsid proteins do not induce apoptosis in astrocytes consistent with the failure of TMEV VP2, unlike Coxackievirus VP2, to induce apoptosis in other cells (Henke et al., 2000). The two VP2 proteins differ in their high-variable regions located between amino acids 140 and 200 (Henke et al., 2001).

Apoptosis is usually triggered through pathways comprising “death” ligands and their cognate receptors (Pan et al., 1997; Pitti et al., 1996). Flow cytometry, used to detect altered expression of TNF-TRAIL family members in GD-VII-infected astrocytes, indicated a significant upregulation in TNF-R and TRAIL expression. A considerable background production of TRAIL in untreated astrocytes was also detected. Those results were more evident in primary than in secondary trypsinized cultures. Very low levels of TNF-α (up to 200 U/ml) were previously detected by ELISA in the supernatants of astrocytes infected with BeAn virus (Sierra and Rubio, 1993). TRAIL upregulation after infection with cytomegalovirus, reovirus, measles virus, and TMEV has been recently reported (Clarke et al., 2000; Jelachich and Lipton, 2001; Sedger et al., 1999, Vidalain et al., 2000).

Binding experiments with 125I-labeled recombinant murine TNF-α revealed an increase from 3258 specific and saturable TNF-R sites detected in uninfected astrocytes to 6385 binding sites per GDVII-infected cell, with the $K_d$ remaining almost unchanged. This 196% increase is similar to that previously reported to be induced by BeAn (Aranguez et al., 1995), despite the inability of BeAn to induce apoptosis. Thus, upregulation of TNF-R might not be the crucial factor for apoptosis signaling in this system.

Several studies have described inhibition of apoptosis by antibodies against TNF-α and TRAIL (Bermudez et al., 1999; Clarke et al., 2000; Kaplan et al., 2000; Kayagaki et al., 1999). In our hands, antibodies to TRAIL and TNF-R at concentrations from 10 to 0.1 μg/ml inhibited GDVII-induced caspase-3 activity by 60 and 50%, respectively (Fig. 8). However, TRAIL has been reported to be unable to bind TNF-R1 (Pitti et al., 1996), so that the possible interactions between the two signals in our system remain unclear. Astrocytes were the cellular source of signals since potential contamination by microglia, oligodendrocytes, neurons, or meningeal fibroblast was carefully ruled out in our cultures. Thus, astrocytes appear to generate signals for their own programmed death as reported for reovirus infection (Clarke et al., 2000).

Almost all astrocytes were infected with GDVII virus in culture and approximately 20% of such cells became apoptotic, as determined by TUNEL staining (Fig. 9). In the infected CNS, TUNEL staining of brain sections revealed apoptotic cells, many of which were also positive for GFAP staining (Fig. 10A–C). Thus, infection with GDVII virus caused programmed cell death in astrocytic cells also in the in vivo situation. By contrast, most of the cells loaded with GDVII virus were neurons, but very few showed TUNEL staining (Fig. 10D–F).

The inability of BeAn virus to induce apoptosis upon infection might be related to the persistence of the virus and its demyelinating effect on the CNS of susceptible mice. Despite the fact that we have studied the nonpersisting, nondemyelinating strain of TMEV, our results might bear on the pathogenesis of neurodegenerative disorders, especially since TRAIL has been reported to be upregulated in peripheral immune cells of patients with multiple sclerosis (Huang et al., 2000).

**Materials and methods**

**Mice**

SJL mice 4 to 6 weeks old were purchased from the Jackson Laboratory, Bar Harbor, MA, USA, and maintained on standard laboratory feed and water ad libitum in the Instituto Cajal animal care center.

**Viruses**

Baby hamster kidney cells (BHK-21) were grown at 37°C in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal calf serum (FCS) and penicillin-streptomycin (Gibco-BRL, Paisley, Scotland). For these studies, a strain of TMEV isolated in 1957 from a feral mouse in Belem, Brazil, called BeAn 8386, and the high-neurovirulence strain GDVII were used. After plaque purification and several passages in BHK-21 cells, cell cultures were infected for 48 h at 33°C, sonicated, and centrifuged to remove cell debris and viral particles were purified. The virus titer (10⁶ PFU/ml) was determined by a standard plaque assay with 1% Noble agar (Difco Laboratories, Detroit, MI) and staining with 0.2% crystal violet in 20% methanol.

**Astrocyte cultures**

Astrocyte cultures were prepared by mechanical dissociation of the cerebral cortex from newborn SJL mice (Rubio et al., 1990). The cortex was isolated under a dissecting microscope and cleaned of choroid plexus and meninges. Cell suspensions were filtered through 135-μm pore-size mesh into DMEM containing 10% FCS and penicillin–
streptomycin. After centrifugation, cells were filtered through a 40 μm nylon cell strainer (Falcon–Becton–Dickinson, Le Pont De Claix, France) and cultured in 25-cm² tissue culture flasks (Costar, Cambridge, MA) at 37°C. The medium was changed after 4 days in culture and subsequently two times a week for the entire culture period. Cultures were enriched in astrocytes by the removal of less adherent microglia and oligodendrocytes by shaking overnight at 37°C and 250 rpm in a table-top shaker (Thermo Forma, Marietta, OH). Cellular confluence was observed 10 days after plating, producing around 3 × 10⁶ cells per flask, showing a polygonal flat morphology. A mean of 98% astrocytes was confirmed by indirect immunofluorescence staining of methanol-fixed cultures using rabbit antialgial fibrillar acidic protein (GFAP) antiserum (Dakopatts, Glostrup, Denmark). The lack of noticeable mature oligodendrocytes and microglial/macrophage cells was determined using a guinea pig anti-myelin basic protein (MBP) antiserum prepared as described elsewhere (Rubio et al., 1990) and monoclonal anti-Mac-1 antibody (Serotec, Oxford, UK). Secondary fluorescein-labeled antibodies were purchased from Sigma Chemical Co. (St. Louis, MO).

**Viral proteins purification**

The purification of TMEV virion particles was performed by isopycnic centrifugation in CsCl gradients (Lipton and Friedman, 1980). Individual VP1, VP2, and VP3 virion proteins were purified by reverse-phase HPLC with and Aquapore RP-300 column (Browlee Laboratories, Santa Clara, CA) (Rubio and Martin-Clemente, 1999). Elution peaks were dialyzed, retaining all their immunological properties. Purity was checked by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) (Laemmli, 1970).

**Neutralization by antibodies**

Antiserum was produced in New Zealand White rabbits by subcutaneous and intramuscular injections at multiple sites. Four injections containing 300 μg of CsCl gradient-purified TMEV virus emulsified with complete Freund adjuvant (Difco) were given at 2-week intervals. The rabbits were bled 10 days after the last booster and antiserum was frozen at −20°C until used. The specificity of the antibody response was checked by Western immunoblotting using purified TMEV virion proteins separated by SDS–PAGE and showed that the antiserum contained antibodies against VP1 and VP2 capsid proteins. GDVII virus samples at different concentrations in culture medium were incubated with increasing dilution of the rabbit anti-TMEV antiserum for 30 min at 37°C. Thereafter, the above incubation mixtures were used for infection of astrocyte cultures and further determination of caspase-3 induction.

Affinity-isolated goat antibodies to mouse TNF-α and to soluble TNF-RI were purchased from Sigma. Rabbit polyclonal antiserum to TRAIL and rabbit-purified IgG to TRAIL-RI were acquired from Alexis Corp. (San Diego, CA). Different amounts of antibodies were added to the infected cultures and caspase-3 was determined 24 h later.

**Caspase-3 assay**

Caspase-3 (CPP32) activity was determined in astrocyte cell lysates using the CaspACE assay system kit from Promega (Madison, WI) according to the manufacturer’s protocol. Briefly, infected cell cultures were washed with ice-cold phosphate-buffered saline (PBS), lysed with cell lysis buffer, centrifuged at 15,000 g for 20 min after 2 cycles of freeze-thaw, and assessed for caspase-3 activity in 96-well polystyrene plates (Nunc-Immuno plates, Nunc, Roskilde, Denmark) based on absorbance at 405 nm. Apoptosis-positive controls were obtained by incubating cultures with 200 nM Staurosporine (Sigma) for 6 h.

**Flow cytometry**

Mock- and GDVII-infected cultures (m.o.i. of 10) were trypsinized, washed, and allowed to recover for 2 h at 37°C in complete medium. After washing cells with PBS containing 5% FCS and 0.2% sodium azide (Sigma), membranes were permeabilized with 0.3% Saponin (Sigma) for 5 min at room temperature. Cells were incubated with primary purified antibody or antiserum (diluted 1:200) for TNF-α, TNF-RI, TRAIL, and TRAIL-RI (listed under Neutralization by antibodies) for 30 min at 4°C. After two washes, the cells were incubated with the secondary FITC-conjugated goat anti-rabbit or donkey anti-goat antibodies (Cappel-Organon, Durham, NC) diluted 1:200. After two more washes, the cells were fixed in 1% paraformaldehyde and analyzed in a FACSCalibur (Becton–Dickinson, Palo Alto, CA). Data were evaluated using CELL Quest 3.1f, supplied with the instrument.

**TNF-α radioiodination**

Recombinant mouse TNF-α, free of protein stabilizers, was purchased from Innogenetics (Antwerp, Belgium) and labeled with ¹²⁵I (Amersham) using the chloramine T method (Greenwood et al., 1963). ¹²⁵I-rTNF-α and free iodine were separated on disposable PD-10 Sephadex G-25M columns (Pharmacia Biotech, Uppsala, Sweden). Specific activity was 203 Ci/mmol and the protein was 96% trichloroacetic acid (TCA) precipitable. The iodinated protein migrated as a single peak with an apparent molecular mass of 17 kDa in SDS–polyacrylamide slab gels.

**Binding assay**

Confluent astrocyte monolayers in multiwell (six-well) plates (Falcon–Becton–Dickinson) were used in the binding experiments. Cells were incubated with different amounts of ¹²⁵I-TNF-α with or without unlabeled TNF-α, as stated
in the text. The buffer used was PBS containing 0.1% bovine serum albumin (Gibco-BRL). After washing three times, cells were detached from the plastic surface with 2% SDS at 60°C and counted in a LKB-Wallac 1282 Compu-gamma counter (Sollentuna, Sweden).

Statistical analysis

The dissociation constant ($K_d$) and maximum binding ($B_{\text{max}}$) were calculated from the Scatchard plot of the binding assay data using the GraphPad Prism version 3.00 program. Data are given as mean values ($\pm$SD) of triplicate determinations from three independent experiments.

Intracerebral mice inoculations

Six-week-old SJL mice were anesthetized with Fluorthane and injections were made using a 25-μL Hamilton syringe at a site 1 mm right lateral and 2 mm rostral of the bregma. Twenty microliters of a suspension of GDVII virus (2 × 10^6 PFU) was infused at a rate of 1 μl every 5 s and the needle was maintained for an additional 5 s. Four to seven days after injections, brains were removed and samples were processed by immunochemistry. Acute encephalitis killed remaining, infected animals within a week after inoculation. Control injected mice received 20 μl DMEM.

Immunocytochemistry

Animals in all cases were perfused though the heart 4 days after intracerebral virus injection with 4% paraformaldehyde in PBS. After perfusion, brains were removed, immersed in the same fixative for 3 h, and left overnight in PBS. Vibratome sections of 30–40 μm were processed free-floating for immunohistochemistry. Sections or astrocyte monolayers in culture chambers (Lab-Tek Chamber slide, Nunc, Neperville, IL) were incubated with primary antibody (rabbit anti-TMEV or rabbit anti-GFAP, diluted 1:1000) followed by goat anti-rabbit Cy-3-conjugated antibody (rabbit anti-TMEV or rabbit anti-GFAP, diluted 1:100). After several rinses in PBS, sections were stained for TUNEL and examined in a Leica TCS NT confocal laser scanning microscope equipped with an argon/krypton-mixed gas laser with an excitation peak of 647 nm for Cy-3. Specificity was controlled by omission of the primary antibodies. Quantitative analysis was performed using an Analytical Imaging Station (Imaging Research Inc., Canada).

TUNEL assay

Apoptotic cells were detected using the TUNEL method. Samples were processed for TUNEL using a fluorescein in situ cell death detection kit following the manufacturer’s instructions (Boehringer Mannheim, Mannheim, Germany). Stained cells were visualized by fluorescence microscopy in the confocal laser microscope at an excitation peak of 488 nm. Method specificity was controlled by omission of terminal deoxynucleotidyl transferase (TdT) in the first step of the labeling.

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