Acetylcholine Protection of Adult Pig Retinal Ganglion Cells from Glutamate-Induced Excitotoxicity

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PURPOSE. To determine which glutamate receptor (GluR) subtypes are responsible for glutamate-induced excitotoxicity in cultured adult pig retinal ganglion cells (RGCs) and to characterize the neuroprotective effect of acetylcholine (ACh) on pig RGCs.

METHODS. Adult pig RGCs were isolated from other retinal tissue by a modified panning technique using Thy 1.1 antibody. Isolated RGCs were cultured in control media and media containing glutamate, NMDA, or KA; glutamate and CNQX, MK-801, or AP-7; ACh, nicotine or muscarine; ACh and α-bungarotoxin (Bgt) or methyllycaconitine (MLA); and glutamate and choline or glutamate, choline, and MLA. To determine cell viability, cells were loaded with calcine and counted.

RESULTS. Ninety-eight percent of isolated cells were immunolabeled with Thy 1.1 antibody. Chronic exposure to 500 μM glutamate decreased the number of surviving large and small RGCs, compared to control conditions. This glutamate-induced excitotoxicity was mediated through both NMDA and non-NMDA GluRs. In neuroprotective studies, ACh, nicotine, and choline significantly reduced glutamate-induced excitotoxicity in adult pig RGCs through α-Bgt-sensitive nicotinic ACh receptors (nAChrRs).

DISCUSSION. This was the first report of a modified panning technique to isolate adult pig RGCs. Cell viability was relatively high using this method, and both large and small RGCs grew extensive neurites in culture. The finding that both NMDA and non-NMDA GluRs were involved in glutamate-induced excitotoxicity suggests that isolated pig RGCs provide a good model for glutamate receptor activity. ACh may be useful in protecting RGC from excitotoxic insults occurring in neurodegenerative diseases such as glaucoma. (Invest Ophthalmol Vis Sci. 2004;45:1531–1543) DOI:10.1167/iovs.03-0406

Excitotoxicity, neuronal cell death caused by excessive activity, is linked to various diseases of the central nervous system including the retina. In the retina, diseases associated with excitotoxicity include retinal ischemia, diabetic retinopathy, and glaucoma. Glaucoma is one of the leading causes of blindness in the world, affecting an estimated 66 million people. Although the fundamental cause of glaucoma is yet to be found, the primary risk factor associated with glaucoma is an increase in intraocular pressure. Several studies have identified an excess of the excitatory neurotransmitter, glutamate, in the vitreous humor. Previous studies have demonstrated that excess glutamate release in the eye leads to prolonged influx of nonspecific cations in retinal ganglion cells (RGCs) of the retina and triggers intracellular signaling cascades leading to apoptosis. RGCs are one of five main types of neurons in the vertebrate retina. The axons of the RGCs form the optic nerve, which acts to convey visual information to the brain. When RGCs die through excitotoxicity-induced apoptosis, loss of the visual field occurs.

Previous immunocytochemical, pharmacological, and in situ hybridization studies have demonstrated that RGCs contain both NMDA and non-NMDA ionotropic glutamate receptors (GluRs). In this study, we used cultured RGCs isolated from adult pig retina as a model for human disease, including glaucoma. Pig RGCs were used in this study, instead of the more common rodent model, because of the conflicting results reported using rodent preparations concerning the types of glutamate receptors involved in glutamate-induced excitotoxicity. For instance, in mixed rat retinal cultures, a number of studies have demonstrated that RGCs are susceptible to NMDA-induced cell death. However, other studies using the rat model have demonstrated that both NMDA and non-NMDA receptors are required for excitotoxicity; still other rat model studies have demonstrated that glutamate-induced excitotoxicity is due solely to non-NMDA receptors.

The purpose of this study was to characterize the specific GluR subtypes responsible for glutamate-induced excitotoxicity in adult pig cultured RGCs and to characterize a potential neuroprotective agent that promotes RGC survival. A growing body of evidence indicates that neuronal nicotinic acetylcholine receptors (nAChRs), in addition to mediating fast cholinergic transmission, may modulate other actions within the CNS. In particular, α7 nAChR has been linked to neuroprotection against glutamate-induced excitotoxicity in the brain. However, a potential neuroprotective role of acetylcholine (ACh) in the retina remains unexplored. In the retina, cholinergic neurons are a well-described population of amacrine cells known as starbursts, due to their unique morphology. This population of cholinergic neurons is equally divided in the retina, with one group in the inner nuclear layer of the retina and the other group lying in the ganglion cell layer. The cholinergic starburst amacrine cells receive strong excitatory input from bipolar cells and synapse onto RGCs.

In this study, we demonstrated that activation of α-bungarotoxin (Bgt)-sensitive nicotinic ACh receptors in adult pig RGCs was linked to neuroprotection against glutamate-induced excitotoxicity. The type of ACh receptors involved in this process was characterized.

METHODS

Dissociation and Panning Procedure

Pure RGCs were isolated from adult porcine eyes using a modified version of the two-step panning technique first described by Barres et al. Adult pig eyes were obtained immediately after slaughter from a local slaughterhouse and kept on ice for transport. Within 1 hour of
eye removal, retinas were removed from the eyes and subsequently dissociated and isolated using the modified panning technique. To separate retina from the eye chamber, the cornea and lens were surgically excised from each eye, and the underlying lens and vitreous humor were manually removed. The remaining eyecup containing retina was placed in modified CO2-independent medium (Gibco, Carlsbad, CA; cat # 18,045–088), maintained at 37°C, containing 4 mM glutamine, 10% fetal bovine serum (FBS), 5% antibiotic/antimycotic, and 4 mM HEPES. The retina was manually peeled out of the eyecup. Isolated retinas were subsequently chopped into smaller fragments and enzymatically treated with papain (27 μg/mg) for 20 minutes at 37°C. After 20 minutes, enzymatic treatment was inactivated by rinsing tissue in fresh 37°C CO2-independent medium containing 1 mg/mL DNase. Once enzymatically treated, retinal tissue was fully dissociated by gentle trituration using an unpolished Pasteur pipette. Dissociated cells were then incubated in a 150 × 15 mm Petri dish precoated with goat anti-rabbit IgG antibody (Jackson ImmunoResearch, West Grove, PA; cat #111–005-005; 0.5 mg in 10 mL of 20 mM Tris buffer) for 1 hour at 37°C to eliminate nonspecific binding. This was the first panning step. Goat anti-rabbit IgG antibody had been applied and evenly disubsed over the Petri dish the evening before retinal dissociation and kept at 4°C overnight. In the morning of the dissociation, IgG-treated Petri dishes were washed three times with PBS and washed once again with PBS containing 0.2% BSA for 20 minutes before removal and application of dissociated pig retina.

After 1 hour on the IgG plate, cells were transferred to another 150 mm × 15 mm Petri dish coated with mouse anti-rat Thy 1.1 antibody (BD Biosciences, San Diego, CA; cat # 55,4898; 12.5 μg in 10 mL PBS containing no magnesium chloride and no calcium chloride) bound to goat antigoat IgM (Jackson ImmunoResearch; cat #115–005-075; 0.36 mg in 10 mL of 20 mM Tris buffer). This was the second panning step. To prepare this second panning plate, an IgM 150 mm × 15 mm Petri plate was prepared 1 day before retinal dissociation using the same procedure described for the IgG plate. On the dissociation morning, unbound IgM was sucked off the Petri dish and Thy 1.1 was applied and maintained at 37°C before transferring cells from the IgG plate.

Cells remained on the IgM/Thy 1.1 plate for 1 hour at 37°C and the supernatant was subsequently discarded. RGCs bound to Thy 1.1 were released using 0.25% trypsin for 10 minutes at 37°C. After 10 minutes, trypsin activity was stopped with 1 mg/mL soybean trypsin inhibitor and cells were stained. At this point, the cell density of the dissociated RGCs was calculated with a hemocytometer, cells were diluted and plated at a density of 1 × 10^5 cells/mL into 24-welled culture dishes coated with poly-D-lysine/laminin and incubated at 37°C. The CO2-independent medium was modified to contain: 10% FBS, 5% antibiotic/antimycotic, 0.36 mg/mL goat antigoat IgM, 4 mM glutamine, 15 μg/mL growth factor (NGF), 500 μg/mL transferin, and 10 μg/mL insulin. Survival of cultured cells in this medium was significantly enhanced compared to cells cultured in Dulbecco's modified Eagle's medium (DMEM)-F12 medium in 5% CO2. In the modified CO2-independent medium, an average of 61% (SE ± 5) cells survived the dissociation process and remained healthy for a minimum of 3 days. This was compared to an average of only 42% (SE ± 3) cells that survived the dissociation process and remained healthy for a minimum of 3 days when cells were incubated in DMEM-F12 medium. Cell survival was determined by calcine fluorescence as described below.

After cells were allowed to settle for 2 hours, medium was replaced with fresh modified CO2-independent medium in each well. The first column of each 24-welled culture dish contained untreated control cells. The remaining five columns in each 24-well culture dish contained cells that were treated with appropriate agonists for antagonist screening. In these five columns contained their appropriate pharmacological agents for period of 1, 3, 5, or 7 days. After these time periods, cells were loaded with 2 μM of membrane-permeable calcine for 1 hour to label living viable cells.34 In some experiments, 10 μM calcine was used instead of 2 μM to label neurites as well as cell bodies. After the 1-hour incubation, living cells intensely fluoresce as a result of 495 nm excitation. Background fluorescence levels were inherently low with this assay technique because the calcine dye is virtually nonfluorescent before interacting with cells.

Microscopy was performed on a Nikon Diaphot epifluorescent research microscope illuminated by a 100-W mercury arc lamp with an excitation filter EX 510 to 590, dichroic mirror DM 580, and barrier filter BA590. Fluorescent cells were recorded by a Hamamatsu XC-77 CCD camera, captured using a Metamorph Imaging system (Universal Imaging, Downingtown, PA) and counted using Imagepro software (Media Cybernetics, Inc., Silver Spring, MD). The number of living RGCs in pharmacologically-treated wells was compared to the number of living cells counted in the wells containing untreated control RGCs to obtain a percent change from control. A minimum of five animals was used to generate all pharmacological results. From each animal, cells were plated onto a minimum of four individual wells for each agent tested. Statistical analysis was performed on data using an analysis of variance (ANOVA) followed by linear contrast. P < 0.05 was considered statistically significant. Statistical treatments were performed on data normalized to control values for each experimental series to minimize variation.

**Immunocytochemistry**

To ensure that all the panned cells were RGCs, some panned cells were processed immunocytochemically with an antibody specific for RGCs: Thy 1.1 (Dako, Carpenteria, CA; cat # MAB 107). Panned RGCs were plated onto coverslips and fixed in ice-cold 100% methanol for 10 minutes, washed three times with phosphate buffered saline (PBS), and blocked with 5% bovine serum albumin (BSA) in PBS. Primary mouse anti-rat Thy 1.1 antibody (1:100; BD Biosciences; CD90) was applied to fixed cells in 1% BSA in PBS overnight at 4°C. After 24 hours, coverslips were washed three times with 0.1% BSA in PBS and a secondary antibody linked to Alexa Fluor 488 (emission 494; Molecular Probes, Eugene, OR) was applied to cells for visualization at 1:100 in 1% BSA in PBS for 30 minutes at room temperature. After a secondary antibody treatment, coverslips were washed four times with 0.1% BSA in PBS. Coverslips were attached to slides using Slowfade antifade medium (Molecular Probes). Cells were photographed with a Hamamatsu XC-7CCD camera. To confirm that antibody staining was specific for the antigen, control slides were included in which the primary antibody was omitted.

**Pharmacological Treatment**

In some experiments, panned RGCs plated in columns 2 to 6 in 24-welled culture dishes were pharmacologically treated with glutamate, glutamate agonists, glutamate antagonists, ACh, ACh agonists, or ACh antagonists. In other experiments, a combination of these agents was applied. All agents were directly applied to individual culture wells to obtain a variety of agent concentrations. All agonists and antagonists were obtained from Sigma (St. Louis, MO): (agonists) glutamate, N-methyl-D-aspartate (NMDA), kainic acid (KA), ACh, choline, and nicotine; (antagonists) 6-cyano-7-nitroquinoline-2,3-dione (CNQX), 2-amino 7 phosphonoheptanoic acid (AP-7), 5 methyl-H,11-dihydro-5H-benzo[a,d]cyclohepten-5,10-imine hydrogen maleate (MK-801), α-hungarotoxin (α-Bgt), and methyllycaconitine, (MLA). Agonist and antagonist concentrations were determined from dose–response experiments. The minimal dosage that provided a maximum effect was used throughout the study.

**RESULTS**

**Immunoreactivity to Thy 1.1 Antibody by All Panned Retinal Cells**

The panning technique used in this study was specifically designed for isolation of adult pig RGCs. To verify that pig RGCs were isolated from all other retinal cell types, panned...
retinal cells were immunostained with primary antibody specific for retinal ganglion cells, Thy 1.1. In Figure 1, adult pig RGCs were isolated by the two-step panning technique described in the Methods section, plated, and immunostained with primary antibody Thy 1.1 (1:100) and secondarily labeled with Alexa Fluor 488. Figure 1A demonstrates a typical image obtained from a culture dish containing immunostained panned retinal cells using transmitted illumination. In Figure 1B, the same field of cells labeled with Thy 1.1 antibody are visualized using Alexa Fluor 488 fluorescence. As seen in this example, every cell in the field was immunostained with Thy 1.1 antibody. This was typical of results obtained from all panned retinal cells and supported the hypothesis that the panning technique used in this study exclusively isolated pig retinal ganglion cells. In 48 culture wells containing panned cells, an average of 98.2% (± 3) of cells stained with Thy 1.1 antibody. The percentage of cultured cells that stained with Thy 1.1 antibody did not change whether cells were analyzed for Thy 1.1 immunostaining 1, 3, or 5 days after culture. At no time was there any evidence of other retinal neurons or glial growth in culture wells containing panned cells. In 48 cultured wells containing immunostained pig RGCs, the ratio of large to small RGCs was 1:3. As evident from the Thy 1.1 immunostained RGCs, large and small RGCs were isolated and labeled. Large RGCs had displaced nuclei similar to large pig RGCs immunostained with alternate antibodies.37 Figure 1C illustrates a field of panned cells immunocytochemically processed when the primary antibody was omitted.

When cells were not panned, labeling with Thy 1.1 antibody (1:100) was restricted to a subset of cells. To generate Figure 1D, an entire retina was enzymatically treated, dissociated, plated, and cultured for 3 days without using the modified two-step panning technique to isolate retinal ganglion cells from other retinal cells. Under these conditions, a monolayer of cells covered each plated culture well within 3 days. After 3 days, cells were processed with Thy 1.1 antibody according to the immunocytochemical procedures outlined in the Methods section. Figure 1D illustrates the field of view under transmitted illumination. Figure 1E illustrates the same field of cells labeled with Thy 1.1 antibody and visualized using Alexa Fluor 488 fluorescence. Omission of the panning step resulted in a mixed culture where only a fraction of cells were immunostained with Thy 1.1 antibody. An overlay of Figure 1D and Figure 1E is illustrated in Figure 1F.

In Figure 2, RGCs were cultured for 3 days and loaded with 10 μM calcein instead of 2 μM calcein for 1 hour, which extensively labeled cell bodies as well as all processes. Large retinal ganglion cells exhibited large circular somata (30–45 μm in diameter) and typically had several fine smooth processes emerging from the cell body (arrows). The small RGCs had diameters <20 μm and were typically monopolar or bipolar with circular bodies (arrowheads), similar to results obtained from Luo et al.37

**Excitotoxic Effects of L-Glutamate and GluR Agonists on Adult Pig RGCs**

Chronically elevated glutamate in the vitreous humor of several species is associated with glutamate excitotoxicity and glaucoma.5–7 In Figure 3, the effect of chronically elevated glutamate on adult pig RGCs is illustrated. Figure 3A demonstrates the percentage of total panned pig RGCs that survive after...
exposure to a relatively high concentration of L-glutamate for 1, 3, 5 days, and 1 week. To obtain the data shown in Figure 3A, panned cells were exposed for various times to 500 μM L-glutamate. Each bar graph represents the mean percent of cells that labeled with calcein compared to control calcein-treated cells obtained from the same animal. Therefore, these graphs represent the mean percent of cells that survived experimental treatment. Data for all pharmacological studies were collected from 5 to 20 different animals.

After 1 day, there was a significant difference in survival between the groups of cells exposed to L-glutamate and control. As shown in Figure 3A, after only 1 day in culture, a mean of 85% of panned RGCs survived, compared to control conditions, and suggested that exposure of 500 μM glutamate for 1 day had an excitotoxic effect on 15% of panned RGCs. However, the largest difference between treated and untreated cells occurred by day 3 where a mean of 58% (±5) of cells survived in the presence of chronic 500 μM L-glutamate compared to untreated cells, suggesting that glutamate had an excitotoxic effect on a mean of 42% of panned RGCs. These results support the hypothesis that chronic exposure to high concentrations of glutamate has an excitotoxic effect on adult pig RGCs. Figures 3B and 3C demonstrate that there was a differential excitotoxic effect of chronic glutamate exposure on large and small RGCs. This was apparent by day 1 in culture, where there was a significant excitotoxic effect of L-glutamate on large RGCs compared to control conditions, but no significant difference observed in the percent of small RGCs compared to control conditions. However, after 3 days in chronic glutamate, there was a significant decrease in both large and small RGCs compared to control conditions, although the glutamate effect was greater on large RGCs compared to small RGCs (P < 0.05). The greater effect of glutamate on the larger RGCs compared to smaller RGCs remained through days 3, 5, and 7. Because glutamate exerted its maximal excitotoxic effect on large and small RGCs by the third day in culture, all cultured cells in subsequent experiments were exposed for 3 days.

In the described experiments, isolated RGCs were exposed to glutamate and agonists within a few hours of dissociation. Because the dissociation process can cause severe enzymatic and mechanical insults on the cells and can potentially affect expression of surface receptors involved in excitotoxicity or neuroprotection, experiments were repeated on RGCs that were cultured for longer periods of time in control medium before addition of any pharmacological agent. We found that the timing of agonist application to panned RGCs had no effect on cell survival. For instance, glutamate caused the same degree of excitotoxicity to panned RGCs whether it was applied to recently cultured cells (within 6 hours) or whether it was applied to cells allowed to settle for 24, 48, or 36 hours before glutamate application (data not shown). Due to this result, all pharmacological agents were applied to recently cultured cells for convenience.

The time-course of glutamate-induced excitotoxicity is demonstrated in Figure 4. As shown in this figure, a percentage of RGCs are destroyed during the dissociation and modified pan-
Glutamate has measurable excitotoxic effects on large RGCs at concentrations as low as 50 μM and had a maximal effect in the presence of 325 μM glutamate. The ED_{50} for the excitotoxic effect of glutamate on large RGCs was 325 μM. In Figure 5B, the dose-response curve for small RGCs was demonstrated. Glutamate had a threshold excitotoxic effect in the presence of 100 μM. Similar to the results obtained with large RGCs, a maximal excitotoxic plateau effect was measured on small RGCs in the presence of 500 μM glutamate. The ED_{50} value for small RGCs was 285 μM. In Figure 5C, the dose-response curve for total RGCs was demonstrated. A measurable threshold change in cell survival was measured in the presence of 50 μM, the maximal plateau effect was elicited in the presence of 500 μM glutamate, and the ED_{50} for the excitotoxic effect of glutamate on total pig RGCs was 302 μM. 500 μM glutamate was used for all subsequent experiments, as it represented the minimum effective dose that elicited maximal excitotoxic effects on both large and small RGCs.

RGCs contain both NMDA and non-NMDA type GluR subtypes. Since L-glutamate is the endogenous agonist for all retinal GluRs, experiments were performed to determine what type of GluR subtypes were responsible for the excitotoxic effect of glutamate on panned RGCs. Cells were cultured in the presence of the glutamate agonists, NMDA or KA (Fig. 6). A concentration range of KA or NMDA was applied for 3 days and results were compared between treated and untreated control cells. Ten μM KA had no significant effect on panned RGCs. However, concentrations as low as 50 μM KA significantly reduced the number of RGCs compared to controls and maximal excitotoxic effects occurred in the chronic presence of 100 μM KA, suggesting that KA receptors are at least partially responsible for glutamate excitotoxicity recorded from panned pig RGCs. Because maximal activation of KA receptors did not mimic the degree of excitotoxicity caused by 500 μM glutamate, it is possible that other GluR subtypes may be involved in glutamate excitotoxicity. Concentrations of NMDA as low as 100 μM significantly reduced the percent of panned RGCs by a mean of 17% compared to control conditions when cultured in media containing normal magnesium (Fig. 6A). However, unless cells are depolarized from their resting membrane potential, the voltage-dependent magnesium block characteristic of NMDA receptors would reduce the contribution of NMDA receptors to the excitotoxic effect of glutamate on pig RGCs. This was addressed by experiments where KA and NMDA were co-applied to panned pig RGCs (Fig. 6B). Under these conditions, KA can bind to KA/AMPA GluRs to open nonspecific cation channels and depolarize the RGCs. After depolarization and removal of the magnesium-induced voltage-dependent block of NMDA receptors, NMDA receptors should be activated. From the results summarized in Figure 6C, the combination of 100 μM KA and NMDA mimicked the total excitotoxic effect of 500 μM glutamate. Taken together, these results suggested that both KA and NMDA receptors are involved in the excitotoxic effect of glutamate observed on pig RGCs.

To confirm that both non-NMDA and NMDA GluR subtypes were involved in glutamate excitotoxicity in pig RGCs, experiments using specific antagonists were conducted. In Figure 7, various concentrations of the KA specific antagonist, CNQX,
were applied to panned adult pig RGCs before chronic exposure to 100 μM KA. Although 100 nM CNQX had no significant effect on KA-induced excitotoxicity, when cells were pre-treated in concentrations of CNQX equal or greater than a threshold dose of 10 μM, KA effects were eliminated (Fig. 7A). Figures 7B and 7C demonstrate that the competitive NMDA antagonist, AP-7, and the noncompetitive NMDA antagonist, MK-801, block the small but significant excitotoxic effect of NMDA on pig RGCs. Pretreatment of RGCs with 10 μM CNQX and 100 μM AP-7 or 100 nM MK-801 virtually eliminated the combined excitotoxic effect of KA and NMDA (Fig. 7D).

If glutamate-induced excitotoxicity is due to activation of a combination of NMDA and non-NMDA GluRs, it follows that a combination of the GluR antagonists specific for NMDA and non-NMDA GluRs should eliminate the effect of glutamate on panned RGCs. When cells were pretreated with CNQX, MK-801, or AP-7, glutamate-induced cell excitotoxicity was significantly reduced, but not blocked (Fig. 7E). However, when a combination of 10 μM CNQX and 100 nM MK-801 or 10 μM CNQX and 100 μM AP-7 were used, glutamate-induced excitotoxicity of panned RGC was completely eliminated. Taken together, these inhibition studies further support the hypothesis that glutamate-induced excitotoxicity of pig RGCs is mediated through a combination of NMDA and non-NMDA GluRs.

**Neuroprotective Effect of Acetylcholine on Panned RGCs**

Neuronal nAChRs play a role in neuroprotection of hippocampal and cortical neurons in the central nervous system. However, although nAChRs exist on vertebrate RGCs, it is not presently known if activation of these AChRs is linked to activation of a neuroprotection mechanism. In this study, the effect of ACh and nicotine on glutamate-induced excitotoxicity using panned adult pig RGCs were examined.

Figure 8 illustrates the summarized results obtained when panned pig RGCs were chronically exposed to 500 μM glutamate for 3 days in the presence of various concentrations of ACh (Fig. 8A) and the ACh agonists, nicotine (Fig. 8B) or muscarine (Fig. 8C). As evident from the data summarized in Figure 8A, when cells were pretreated for 2 hours in 5 μM ACh before addition of 500 μM glutamate, glutamate excitotoxicity in pig RGCs was completely eliminated. Similar results were obtained if cells were pretreated in 1 to 10 μM nicotine (Fig. 8B) and supported the hypothesis that ACh has a neuroprotective effect on glutamate-induced excitotoxicity in pig RGCs. However, when relatively high concentrations of ACh or nicotine (50 μM) were applied before glutamate, the significant neuroprotective effect was lost (right bars, Figs. 8A and 8B). This is likely due to nonspecific activity or toxic effects of these agents at high concentrations. When applied alone, 5 μM ACh or 1 μM nicotine had no significant effect on the survival of RGCs (data not shown).

Neuronal nAChRs are linked to the opening of nonspecific cation channels, whereas muscarinic ACh receptors are linked to activation of G-proteins and second messenger systems. To determine if muscarinic ACh receptors on pig RGCs are involved in neuroprotection of glutamate-induced excitotoxicity, panned RGCs were pretreated with various concentrations of muscarine for 2 hours before application of 500 μM glutamate.
Muscarine had no significant neuroprotective effect on panned RGCs, even at relatively high concentrations (Fig. 8C).

**Partial Mediation of ACh Neuroprotection through α7 Nicotinic ACh Receptors**

Eleven neuronal nAChRs can be pharmacologically grouped into α-Bgt-sensitive and α-Bgt-insensitive agents. Neuronal nAChRs of the α-Bgt-sensitive class contain α7 subunits and account for most of the α-Bgt binding in the vertebrate brain.40 Numerous investigators have examined the role of α7 nAChRs in neuroprotection against glutamate-induced excitotoxicity in several regions of the CNS,67,74,80 excluding the retina. In this study, experiments were performed on panned pig RGCs to determine whether the neuroprotective effect of ACh and nicotine is mediated through α7 α-Bgt-sensitive nAChRs. Panned RGCs were pretreated for 2 hours in various concentrations of α-Bgt before addition of 5 μM ACh and 4 hours before addition of 500 μM glutamate. Previous agents were not washed out as new additions were made, producing a final media cocktail containing all of the above-mentioned agents. Although 1 nM α-Bgt blocked the neuroprotective effect of ACh by 10%, 10 nM α-Bgt reduced its neuroprotective effect on panned RGCs by a mean of over 90% (Fig. 9A). When ACh was replaced with nicotine, α-Bgt had similar effects on blocking neuroprotection. As shown in Figure 9B, the neuroprotective effect of 1 μM nicotine was significantly reduced when panned cells were pretreated with 1 nM α-Bgt. However, when cells were incubated in 10 nM α-Bgt, the neuroprotective effect of nicotine on panned RGCs was blocked by a mean of 95% (Fig. 9B).

These initial pharmacological studies supported the hypothesis that α-Bgt-sensitive α7 nAChR subunits were responsible for the neuroprotective effect of ACh. However, much of the commercially available α-Bgt contains small amounts of contaminants that can affect other nAChR subunits. To address this issue, further pharmacological experiments were performed.

**FIGURE 7.** Glutamate antagonists prevent excitotoxicity. Bar graphs demonstrating the effects of: (A) CNQX on KA-induced excitotoxicity; (B) AP-7 on NMDA-induced excitotoxicity; (C) MK-801 on NMDA-induced excitotoxicity; (D) a combination of CNQX and MK-801 or AP-7 on the excitotoxic effect induced by a combination of KA and NMDA; (E) CNQX, MK-801, AP-7, a combination of CNQX and AP-7, or a combination of CNQX and MK-801 on chronic glutamate exposure. Each bar graph represents the mean ± SE (n = five experiments).
using the α7 antagonist, methyllycaconitine (MLA), and the α7 agonist, choline.

MLA is an α-Bgt-sensitive nAChR antagonist, highly selective for α7 nACh receptors when used at concentrations between 1 to 10 nM in most preparations, although it binds to non-α7 nAChRs at low concentrations in dopaminergic neurons of the substantia nigra and ventral tegmental area. At higher concentrations, MLA also affects other nAChR subunits that are likely to be expressed in pig retina. In inhibition studies, when panned RGCs were pretreated with various concentrations of MLA before ACh and L-glutamate, 10 nM MLA significantly reduced the neuroprotective effect of ACh and nicotine (Figs. 9C and 9D). However, inhibition of the neuroprotective effect was maximized when cells were pretreated with 100 nM MLA, which blocked the neuroprotective effect of ACh by a mean of 98% and that of nicotine by 100% (Figs. 9C and 9D).

The finding that a relatively high concentration of MLA was required to block ACh neuroprotection supported the hypothesis that multiple nAChR subunits are involved in the neuroprotective effect of ACh. This idea was further supported in experiments using choline (Fig. 10). After 3 days in culture, 75.8% (±3.2) of panned RGCs survived compared to 58% of RGCs that typically survive in the presence of 500 μM L-glutamate alone. In inhibition studies, pretreatment of cells in 10 nM MLA completely eliminated the neuroprotective effect of choline on glutamate-induced excitotoxicity in pig panned RGCs (Fig. 10). Taken together, these results strongly supported the hypothesis that ACh neuroprotection is partially mediated through α7 nAChRs, but additional nAChR subunits are also likely to be involved.

**DISCUSSION**

The present study is the first to isolate pure adult pig RGCs using a two-step panning technique with Thy 1.1 antibody. Ninety-eight percent of cultured RGCs were immunolabeled with Thy1.1 antibody, verifying the purity of the culture. This was in agreement with findings in other animal models and confirmed the specificity of the Thy 1.1 antibody for RGCs. Based on morphologic features, adult pig RGCs immunostained with the α7 staining antibody were identified.
with Thy 1.1 were subdivided into two groups: small and large cells, similar to those characterized by Luo and associates \(^3\) in a mixed retinal culture. Immunostaining with Thy 1.1 was localized on somas of both large and small pig RGCs with little staining on neurites. This predominance of staining on pig RGC bodies differs from the staining pattern previously described in intact retinas of chick, rodent, and cats,\(^3,5,51,52\) where Thy 1 primarily labeled the interplexiform layer and stained the ganglion cell layer to a lesser degree. The difference in Thy 1 staining patterns between these species could be due to a variety of factors. For instance, in the cat retina, Thy 1 was found in the inner nuclear layer as well as in the inner plexiform and ganglion cell layer and represents a retinal distribution pattern different from that seen in other species.\(^5,2\) Therefore, species difference could account for differences observed. In addition, pig RGCs were fixed in 100% methanol instead of the 4% paraformaldehyde used in all other studies of Thy 1 antibody. Different fixations have dramatic effects on immunostaining patterns in other systems\(^5,5\) and could explain the difference in staining pattern. Also, pig RGCs were incubated in primary antibody overnight compared to the 30-minute incubation time associated with other studies using antibodies against Thy 1, which could explain the different

**FIGURE 9.** The neuroprotective effect of ACh on glutamate-induced excitotoxicity is inhibited by \(\alpha\)-Bgt (A) and MLA (C). In (B) and (D), the effect of various concentrations of \(\alpha\)-Bgt and MLA on the neuroprotective effect of nicotine on glutamate-induced excitotoxicity is demonstrated. Each bar graph represents the mean ± SE (n = five experiments).

**FIGURE 10.** ACh neuroprotection is partially mediated through activation of \(\alpha7\) nAChR subunits. Bar graphs demonstrate the effect of various concentration of choline on glutamate-induced excitotoxicity and the effect of 10 nM MLA on choline’s neuroprotection. Each bar graph represents the mean ± SE (n = five to ten experiments).
immunostaining pattern present on pig RGC bodies compared to other species. Lastly, the difference in staining pattern may be due to the dissociation process. As seen in Figures 1 and 2, dissociated RGCs have fewer neurites than those identified in the intact retina. Fewer neurites and other changes that occur in the cells due to the dissociation process are likely to account for the labeling disparity.

The modified two-step panning technique allowed the determination of the concentration at which glutamate was toxic to adult pig RGC in vitro. A concentration of at least 500 μM glutamate induced maximal cell death of large and small RGCs after chronic exposure for 3 days. Large RGCs were affected first in a fashion similar to the pattern of loss seen in glaucoma.14 Significant loss of large RGCs to 500 μM glutamate occurred within 1 day, whereas significant loss of small RGCs to glutamate took 3 days. Also in agreement with in vivo studies,14,15,54 the loss of large RGCs was significantly larger than the loss of small RGCs at 1, 3, 5, or 7 days in culture. These results are in contrast to the study by Luo et al.,37 who examined the effects of glutamate on an unpanned mixed retinal culture. Under mixed culture conditions, there was a selective, dose-dependent loss of large RGCs due to excess glutamate exposure but small RGCs were resistant to glutamate. It is possible that the different culture environments used in the two studies explains this discrepancy. In the mixed retinal culture, the myriad of different cell types, including glia, may act to bind or transport glutamate away from the RGCs and effectively reduce the concentration of glutamate present in the culture. This suggestion is consistent with the finding that larger concentrations of glutamate were needed in the mixed retina culture system to induce large RGC death compared to the concentrations of glutamate used on panned RGCs.

The present study supported the hypothesis that both NMDA and non-NMDA receptors are involved in adult pig RGC excitotoxicity, based on the finding that a combination of KA and NMDA was required to mimic glutamate excitotoxicity. Neither KA nor NMDA alone was sufficient to account for the total excitotoxic effect of glutamate. KA had the largest effect on RGC survival compared to NMDA. However, because magnesium was present in the culture media, the result of NMDA alone on pig RGCs was diminished due to the voltage-dependent magnesium block characteristic of NMDA receptors.59 To better evaluate the contribution of the NMDA receptor on RGC excitotoxicity, KA and NMDA were co-applied. Under these conditions, KA would activate non-NMDA GluR channels, allowing permeation of nonspecific cations, depolarization of RGCs, and NMDA receptor voltage-dependent magnesium block and permeation of ions through NMDA receptor channels. Co-application of KA and NMDA accounted for 100% of the glutamate-induced excitotoxic effect. The combined excitotoxic effect of KA and NMDA was totally eliminated only when cells were pretreated with a combination of NMDA and non-NMDA specific antagonists.

In vivo, glutamate receptors on RGCs are largely found on the dendrites and not on the somata.36,57 Due to the dissociated in vitro system used in this study, it is likely that the expression patterns of glutamate and ACh receptors on panned RGCs differ considerably from that of RGCs in vivo. Therefore, agents in this study were applied to activate agonist-specific somatodendritic receptors as well as dendritic receptors. This raises a concern common to any in vitro system. If the dissociation process has changed the expression pattern of receptors and likely other cell properties, how can results from an in vitro system accurately represent what is happening in an in vivo system? From the results of this study, the data strongly support the hypothesis that both NMDA and non-NMDA receptors were involved in glutamate-induced excitotoxicity in pig RGCs. However, if the intact pig retina only contained non-NMDA receptors at dendritic synapses, it may be that only non-NMDA receptors played a role in excitotoxicity under physiological conditions. Experiments are currently underway to identify the receptor expression of pig RGC glutamate and ACh receptors in culture as well as in vivo.

Excitotoxicity

Both KA/AMPA and NMDA channels on vertebrate RGCs are permeable to calcium ions,58–60 which is a likely intracellular signal that acts to trigger downstream signaling pathways leading to apoptosis. Although the exact apoptosis signals in RGCs are unclear at present, a variety of such signaling pathways in other systems have been identified. These include, but are not limited to, signaling pathways linked to mitochondrial calcium overload with subsequent cytochrome C release,61 free radical production (nitric oxide, peroxide anion),62,63 p38 mitogen activated protein kinase (MAPK)-induced activation of MEF2 transcription factor,64,65 cytokine-induced activation of extra-cellular signal-regulated kinase (ERK),66 tyrosine phosphorylation of Janus kinase 2 and subsequent activation of AKT,67 and caspase activation with DNA fragmentation.68 Although it is not known which, if any, of these mechanisms are involved in RGC excitotoxic death, activation of both NMDA and non-NMDA receptors could create increases of intracellular calcium that are pathologic to the cells. Overall, large increases in intracellular calcium act as a second messenger to change the baseline activity of numerous enzymes that lead to apoptotic cascades and eventual cell death.

Neuroprotective Mechanisms

In previous studies, many laboratories have examined a variety of ways to prevent glutamate-induced excitotoxicity. Using pharmacological approaches, some studies have confirmed the importance of both NMDA and non-NMDA receptors in the mechanism underlying RGC death with the neuroprotective action of specific GluR antagonists.59,17,37,69 Other pharmacological agents that do not directly affect GluRs have neuroprotective effects, including adrenergic α2 agonists,4,70 beta 1 antagonists,71 and neurotrophic factors.72 In addition, other studies demonstrate that a low pH medium will protect retinal neurons from glutamate-induced delayed death through a proton modulation site on NMDA-operated channels73 and that NO-related species will react with critical cysteines on the NMDA receptor to prevent neuronal apoptosis.68,74 Nonpharmacological approaches to neuroprotection include preconditioning and activation of immunity. Preconditioning refers to strategies where subthreshold injuries induce an endogenous neuroprotective signal.69,75 Attempts are currently underway to isolate this endogenous signal. Other laboratories have demonstrated that certain types of immune activation positively correlate with retinal ganglion cell survival in retinal ganglion cell injury.67,77 It is hoped that this research may lead to a vaccine-like therapy for glaucoma.78

Although a variety of agents have a neuroprotective effect on RGCs in other systems, this study is the first to determine that relatively low concentrations of ACh and nicotine have a neuroprotective effect on panned adult pig RGCs against glutamate-induced excitotoxicity. The action of ACh and nicotine on RGCs was mediated exclusively through nAChRs, as muscarine failed to mediate any protection. The inefficiency of muscarinic receptors to mediate neuroprotection was also observed by Donnelly–Roberts et al.79 who found that the muscarinic receptor antagonist, atropine, did not block the neuroprotective effects of a cholinergic channel activator in rat primary cortical cell cultures.
Accumulating evidence from other studies suggest that the neuroprotective action of ACh in the brain is mediated through α-Bgt-sensitive α7 neuronal nAChRs, that transduce signals to phosphorylatedinositol 3-kinase to block β-amyloid-induced neurotoxicity. Activation of α7 nAChRs in the brain triggers JAK2 and activation of phosphorylatedinositol 3 (PI3)-kinase and Bcl-2, enzymes involved in cell survival which contribute to a neuroprotective effect. The neuroprotective role of α7 nAChRs in the retina was therefore explored.

Neuronal nAChRs represent a very heterogeneous family of ion channels. In the nervous system, nine different alpha subunits (α2–α10) and three different beta subunits (β2–β4) have been described. These subunits assemble in vivo to form various combinations, although α7 subunits form homopentamers in heterologous expression systems. In the brain, α7, α4, and β2 are the most abundant nAChRs, and there is growing evidence for α7 in the retina. Initial pharmacological studies supported the hypothesis that α-Bgt-sensitive α7 nAChR subunits were responsible for the neuroprotective effect of ACh; relatively low concentrations of α-Bgt inhibited the effect of ACh. However, because much of the commercially available α-Bgt contains small amounts of contaminants that can affect other nAChR subunits, further pharmacological experiments were performed using the α7 agonist, choline, and the α7 antagonist, MLA, to address this issue. Results demonstrated that although choline had a significant neuroprotective effect on RGCs, it only represented approximately one-half of the protective effect observed when cells were pretreated with 5 μM ACh. In inhibition studies, low concentrations of MLA that are specific for α7 nAChR subunits completely blocked the neuroprotective effect of choline. These results strongly suggested that α7 nAChR subunits were involved in ACh neuroprotection, but additional nAChR subunits were also involved. Further experiments are needed to identify and characterize the contribution of these other putative nAChR subunits.

How can activation of nAChR subunits lead to neuroprotection? Although the answer to this question is outside the scope of this article, it is likely that activation of nAChRs triggers a cascade of events ultimately inhibiting apoptotic pathways or enhancing cell survival pathways. The catalyst for triggering these cascades could be due to changes of membrane potential that occur when nAChR channels open, but more likely is due to ion permeation through activated nAChR channels. Nicotinic AChRs are nonspecific cation channels highly permeable to calcium ions. Calcium ions have been shown to affect a wide variety of intracellular second messenger cascades, including the MAP kinase pathways and PI3 kinase pathways, which are involved in ACh neuroprotection in other regions of the CNS. It is therefore likely that calcium permeation through calcium-permeable nAChRs triggers inhibition or activation of second messenger systems to inhibit apoptosis or enhance cell survival. In this scenario, high calcium influx through glutamate channels are excitotoxic, whereas calcium influx through nACh channels results in neuroprotection. How can calcium influx trigger two separate actions in the same cell? Perhaps the answer is due to cell compartmentalization that separates calcium-sensitive neuroprotective enzymes from calcium sensitive enzymes involved in excitotoxicity, or perhaps the amount of calcium influx through channels is the key. For instance, it is conceivable that a narrow range of calcium results in protection rather than excitotoxicity. Certainly, further experiments are needed to test these ideas.

In summary, this study was the first to demonstrate that the isolation of adult pig retinal ganglion cells using a modified two-step panning technique using Thy1.1 antibody. That chronic exposure of 500 μM L-glutamate induced excitotoxicity in cultured adult pig RGCs through a combination of both NMDA and non-NMDA GluRs was also determined. In addition, this study was first to demonstrate that an endogenous neurotransmitter found in the retina, ACh, has a neuroprotective effect against glutamate-induced neurotoxicity of RGCs and that ACh acts through α-Bgt-sensitive nAChRs, with involvement of α7 nAChRs. These findings suggest that ACh may be used in protecting RGC from excitotoxic and traumatic insults similar to those occurring in glaucoma. A better understanding of this neuroprotective pathway may lead to therapeutic intervention for glaucoma and a variety of other neuronal diseases in the retina and other regions of the CNS.

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