Extracellular Synthesis of cADP-Ribose from Nicotinamide-Adenine Dinucleotide by Rat Cortical Astrocytes in Culture

Ludmila Pawlikowska, Susan E. Cottrell, Matthew B. Harms, Ya Li, and Paul A. Rosenberg

Department of Neurology and Program in Neuroscience, Children's Hospital and Harvard Medical School, Boston, Massachusetts 02115

cADPR is an endogenous calcium-mobilizing agent that in vertebrates is synthesized from nicotinamide-adenine dinucleotide (NAD) by bifunctional enzymes with ADP-ribosyl cyclase and cADPR hydrolase activity. ADP-ribosyl cyclase and cADPR hydrolase activity have been reported in the brain, but the cellular localization of these activities has not been determined previously. In the present study, selective culturing techniques were employed to localize ADP-ribosyl cyclase activity and cADPR hydrolase activity to astrocytes or neurons in cultures derived from rat embryonic cerebral cortex. ADP-ribosyl cyclase activity was determined by incubating cultures with 1 mM NAD in the extracellular medium for 60 min at 37°C and measuring formation of cADPR by bioassay and by HPLC. Astrocyte cultures and mixed cultures of astrocytes and neurons had mean specific activities of 0.84 ± 0.06 and 0.9 ± 0.18 nmol cADPR produced/mg protein/hr, respectively. No detectable ADP-ribosyl cyclase activity was found in neuron-enriched/astrocyte-poor cultures. cADPR hydrolase activity was detectable by incubating cultures with 300 μM cADPR for 60 min at 37°C and assaying loss of cADPR or accumulation of ADPR. The demonstration of extracellular ADP-ribosyl cyclase and cADPR hydrolase activities associated with astrocytes may have important implications for the role of extracellular cADPR in signal transduction and in intercellular communication in the nervous system.

Key words: cADP-ribose; NAD; ADP-ribosyl cyclase; cADPR hydrolase; astrocytes; extracellular enzymes; signal transduction; calcium

Received April 12, 1996; revised May 31, 1996; accepted June 13, 1996.

This work was supported by National Institutes of Health Grants NS 26830 and NS 31353, and a Mental Retardation Core Grant to Children's Hospital.

Correspondence should be addressed to Dr. Paul Rosenberg, Enders Research Building, Department of Neurology, Children's Hospital, 300 Longwood Avenue, Boston, MA 02115.

Copyright © 1996 Society for Neuroscience 0270-6474/96/165372-10$05.00/0

The Journal of Neuroscience, September 1, 1996, 16(17):5372–5381
MATERIALS AND METHODS

Tissue culture. Three kinds of embryonic rat cortical cell cultures were used for this study. Astrocyte-rich (AR) cultures contain 90% astrocytes and 10% neurons. Neuron-enriched/astrocyte-poor (AP) cultures contain 20% astrocytes and 80% neurons. Astrocyte cultures do not contain any neurons, but may include small numbers of other cell types such as oligodendrocytes. AR and AP cultures derived from embryonic Sprague Dawley rats (Charles River Laboratories, Wilmington, MA) were pre-
pared as described previously (Rosenberg, 1991). Tissue was harvested from 16-d-old embryos, dissociated with 0.027% trypsin and plated on poly-L-lysine- (AP) or collagen and poly-L-lysine- (AR) coated glass coverslips in 35 mm dishes (5 coverslips/dish) or in coated wells in 24-well plates (Costar, Cambridge, MA). The growth medium was 8:1:1 DMEM/Earle's balanced salt solution (EBSS) at 22°C. Initial experiments were performed with ammonium formate (0.1 g/ml) pH adjusted to 4.0 with formic acid (0.5 ml/min). Detection was by inhibiting after 4 d for 48 hr and replacing the medium on day 7 with AP medium (Rosenberg, 1991). This medium was not subsequently changed in AP cultures to maximize survival, and the cultures were placed on water-saturated filter paper pads to prevent water loss. Astrocyte cultures were made as described previously (Levison and McCarthy, 1991). In brief, cerebral cortex tissue was harvested from neonatal rat pups and mixed glial cultures prepared. Tissue was dissociated with 2.5% trypsin and triturated in Basal Medium Eagle's supplemented with 10% bovine serum albumin (BSA), 1-251 radioiodinated with 50-100 x g, resuspended in BME-C, plated in tissue culture flasks, and grown for 12 d at 37°C with media changed after 3 d and every 2 d thereafter. Astrocyte cultures were then prepared from the mixed cultures. Other cell types were removed by shaking at 260 rpm for 1.5 hr at 37°C, removing the medium, and rinsing twice with BME and once with a divalent cation-free saline, pH 7.5, composed of (in mM): glucose 11, HEPES 20.2, trisodium citrate 10.2, KCl 4, NaCl 110, plus 2 mg/l phenol red (GHCKS). The adherent astrocyte cells were then dissociated with 0.25% trypsin in GHCKS. BME-C was added to inactivate the trypsin, and the suspension was centrifuged at 100 x g for 10 min. Supernatant was discarded and cells were resuspended in BME-C and plated on coverslips in dishes or in wells as for AR cultures. The astrocyte cultures were then maintained with three medium changes per week.

The age of the cultures used in experiments ranged from 21 to 35 d for AR and AP cultures, and 12 (after reaching confluence) to 70 d for astrocytes.

Media. Complete BHSS was from Sigma (St. Louis, MO). Calcium-free BHSS was prepared by supplementing calcium-, magnesium-, and bicarbonate-free BHSS with anhydrous magnesium sulfate (0.098 g/ml) and sodium bicarbonate (0.35 g/ml).

NAD incubation experiments. Cultures were incubated with wells (0.25 or 0.5 ml) or dishes (1.3 ml or 1.5 ml) of BHSS at room temperature or with agitation at 37°C for up to 120 min. Initially, in some experiments, calcium-free BHSS was used. Collected media were replaced with BHSS [for monitoring the effect of the incubation conditions on cells by microscopy and lactate dehydrogenase (LDH) assays] or with 2% SDS for (protein assays).

When samples from experiments done in complete BHSS were assayed for calcium-releasing activity in the sea urchin egg homogenate bioassay, medium calcium was removed after incubation by shaking sample with Chelex 100 resin (50 mg/ml) (Bio-Rad, Hercules, CA) for 1 hr, centrifuging 1 min at 12,000 rpm, and transferring the supernatant to a fresh tube.

HPLC assays. A 3.9 x 150 mm-Bondapak C18 reverse-phase column (Waters, Milford, MA) was connected behind a 5 x 50-mm PL SAX-100 strong anion exchange column (Polymer Laboratories) with an appropriate guard cartridge protecting each column (Kim et al., 1993; Lee et al., 1993).

Initial experiments were performed with ammonium formate (0.1 M), pH adjusted to 4.0 with formic acid (0.5 ml/min). Detection was by absorbance at 260 nm. The following retention times were observed: cADPR, 6.2 min; AMP, 9.4 min; ADPR, 10.6 min; NAD, 15.2 min; and adenosine, 20.4 min.

Subsequently, for the studies of cADPR hydrolytic activity, we used an ammonium formate gradient (67.6-250 mM), with a flow rate of 0.5 ml/min. Initial conditions were 67.6 mM ammonium formate, in which cADPR was maintained, then ammonium formate was increased to 250 mM over the next 12 min. The column was taken from 100% 250 mM to 100% 67.6 mM ammonium formate over the next minute and then equilibrated from 19 to 25 min in 67.6 mM ammonium formate before the next sample was applied. Sample volume was 50 µl. Guanosine (100 µM) was added to each sample as an internal standard, and for each substance measured, external standards were incorporated into each run. Quantitation was achieved using Beckman System Gold software by comparing the peak height of each substance being measured with the peak height of the internal standard in each sample, and comparing the values obtained with standard curves constructed of ratios of external standards with the internal standard. The following retention times were observed in this system: cADPR, 4.4 min; guanosine, 8.4 min; and ADPR, 12 min.

Preparation of sea urchin egg homogenates. The protocol of Dargie et al. (1993) was followed. Lytechinus pictus sea urchins (Marinus, Long Beach, CA) were injected intracoelomically with KCl (0.5 M) to induce spawning, and eggs were collected into artificial seawater (ASW) composed of (in mM): NaCl 460, MgCl 2, MgSO 4, CaCl 2, 10, KCl 10, NaHCO 3, 2.5, pH adjusted to 8.0 with NaOH. Approximately 50 ml of eggs was collected from 25 females, and a sample was tested to confirm fertilizability by sperm. Eggs were dejellied by exposure to ASW, pH 5, and washed once in ASW, pH 8.0, twice in calcium-free, EGTA-supplemented ASW [as above except EGTA (1 mM)], twice in calcium-free ASW, and once in intracellular medium (ICM) composed of (in mM): potassium gluconate 250, N-methylglucamine 250, HEPES 20, MgCl 2, 1, pH 7.2). Eggs were spun down 1 min at 1000 rpm and supernatant discarded after each washing. The final volume after compaction during washing was 20 ml. The eggs were resuspended in ICM (150 ml) freshly supplemented with leupeptin (25 µg/ml), aprotinin (20 µg/ml), benzamidine (10 mM), and phospho-
creatine (8 mM), soybean trypsin inhibitor (100 g/ml), and ATP (1 mM). The egg suspension was homogenized in a Darton homogenizer with type A Teflon pestle (Thomas cat # 3431D76) and placed on ice.

Calcium-release bioassay. A modified version of the protocol of Galione et al. (1993) was used. For each day's assays, a fresh tube of 10% homogenate was thawed at 20°C, diluted 50% with supplemented ICM containing phosphocreatine (10 mM), creatine phosphokinase (10 U/ml), ATP (1.0 mM), omoglycerin (1 µg/ml), antimycin (1 µg/ml), and NaN 3 (1 mM), and incubated 1 hr at 20°C. The dilution and incubation were repeated two more times, and the fluorescent calcium indicator Fluo 3 was added to 3 µM after the final dilution. The final concentration of homogenate was 1.25%.

Fluorimetry was done in a Hitachi F-2000 fluorescence spectrophotometer set for 490 nm excitation and 535 nm emission wavelengths with both slits at 5 nm. Preincubated homogenate (0.25 ml) was used for each assay. The homogenate was placed in a 1.0 x 0.5-mm glass cuvette, stirred continuously with a magnetic stir bar, and maintained at 20°C with a circluion water bath. Dishes (10 x 8 cm) were filled with a Hamilton glass syringe after baseline fluorescence was established, usually at 30 or 50 sec. For measurement of cADPR in experimental samples, standard curves of response to cADPR standards in incubation media based on initial rate of fluorescence increase were constructed.

Fluorometric assay for ADP-ribosyl cyclase. A sensitive assay for ADP-ribosyl cyclase was implemented based on the work of Graef et al. (1994, 1996) showing that nicotinamide guanine dinucleotide (NADG) is a substrate for the enzyme, and that the product cyclic guanosine diphospho-
"
over a period of 4 min using a kinetic plate reader (Molecular Devices, Menlo Park, CA).

Protein assay. A bicinchoninic acid reagent protein assay (Pierce Chemical Company, Rockford, IL) kit using a bovine serum albumin (BSA) standard was used to assay the amount of protein present in wells and dishes used for experiments. Postexperiments cultures solubilized with 2% SDS and stored at 4°C were used. Data obtained were used to calculate ADP-ribosyl cyclase and cADPR hydrolase activity per milligram of protein. For 35 mm dishes, protein content was 0.496 ± 0.08 mg for AP cultures (n = 6 and 2 culture dates), 0.96 ± 0.17 mg for astrocyte cultures (n = 8 and 3 culture dates), and 1.19 ± 0.15 mg for AR cultures (n = 12 and 5 culture dates).

For experiments measuring hydrolase activity, 12 mm glass coverslip cultures in wells were used to decrease the volume of medium required for the experiment. In these cultures, the protein was determined using the Bradford dye-binding protein assay (1976) with a BSA standard (Bio-Rad). In these experiments, the cell layer was solubilized in NaOH (0.1 M). Protein content was 54 ± 7 and 99 ± 19 μg for astrocyte and AR cultures, respectively.

Data analysis. Statistical variation was determined by calculating SD in multiple data points from a single experiment or SEM where data from multiple experiments were pooled. Statistical comparisons were performed using Student’s t test or ANOVA followed by the Tukey–Kramer test for multiple comparisons, using the Instat2 program from Graphpad Software.

Materials. All reagents and drugs were obtained from Sigma except IP3 and heparin (Calbiochem, La Jolla, CA), HEPES, and ATP (Boehringer Mannheim, Indianapolis, IN), Chelex 100 (Bio-Rad), Fluo 3 (Molecular Probes, Eugene, OR), and methanol (Fisher, Orangeburg, NY). cADPR was purchased both from Amersham (Arlington Heights, IL) and Sigma.

EXPERIMENTAL RESULTS

Three types of cultures differing in cell composition were used: neuron-enriched/AP (20% astrocytes, 80% neurons), AR (90% astrocytes, 10% neurons), and astrocyte (100% astrocytes). AR cultures contain ~30 times the number of astrocytes as AP cultures (Rosenberg, 1991). Both AP and AR cultures are rich in synapses, which are exposed to the extracellular medium in AP cultures but sequestered by a near-confluent layer of astrocytes in AR cultures (Harris and Rosenberg, 1993).

Implementation of the egg homogenate calcium-release bioassay

We implemented the sea urchin egg homogenate bioassay developed by Lee and colleagues (Clapper et al., 1987) and modified by Galiane et al. (Galiane et al., 1993) to look for extracellular ADP-ribosyl cyclase activity associated with neurons and astrocytes in culture. Representative traces and a typical standard curve of Fluo 3 fluorescence increase caused by calcium release in response to injected authentic cADPR are shown in Figure 1, A and B. The homogenate also released calcium in response to IP3 (1 μM) (Fig. 1C). Calcium release in response to NAD (10 μM) was low and delayed (Fig. 1D) as reported previously (Clapper et al., 1987). This difference in the kinetic characteristics of homogenate response to cADPR and to NAD made it possible to distinguish between them at NAD concentrations up to 1 mM in the medium exposed to cells (10 μM final concentration in homogenate). We also confirmed previous observations that IP3 (1 μM) desensitized the homogenate to subsequent injections of IP3, but not to cADPR (100 nM), and that, conversely, cADPR desensitized the homogenate to cADPR but not to IP3 (Clapper et al., 1987; Dargie et al., 1990).

Demonstration of cADPR formation in cultures using bioassay

AR well cultures were incubated 30 min in calcium-free HBSS containing NAD (100 μM) (calcium-free medium was used initially, because medium samples were ultimately to be tested for calcium-releasing activity). A distinct calcium-releasing effect was caused by the 100 μM NAD incubation medium both with 15 min (Fig. 2, trace 2) and 30 min (Fig. 2, trace 3) exposure to cultures that was different from the response observed with the control solution (Fig. 2, trace 1).

We found that media samples retained some calcium-releasing activity in homogenate desensitized to both cADPR and IP3, indicating the presence of a calcium-releasing agent other than these two compounds. The source of this residual calcium-releasing activity in the incubation samples was then investigated and was found to be nicotinamide, which previously has been shown to have calcium-releasing activity in the sea urchin egg homogenate bioassay (Clapper et al., 1987). We found that nicotinamide (100 μM) produced an increase in fluorescence of 8 ± 1.2 (mean ± SEM) fluorescence units equivalent to the increase produced by 2.7 ± 0.3 nM cADPR (24 determinations in 3 separate experiments). Taking this information into account, the experimental protocol was modified. The incubations were carried out in 35 mm dishes instead of 24-well plates to increase the tissue to volume ratio (wells contain ~0.3 μg protein/μl, based on minimal medium volume at 300 μl; dishes contain ~0.6 μg protein/μl based on minimal medium volume of 1.5 ml), the NAD concentration was increased to 1 mM, and incubation time was increased to 60 min. These changes substantially increased the amount of calcium-releasing activity produced by the cultures. Two injections of nicotinamide (100 μM) were used to eliminate the response to nicotinamide present in the media samples (which could at most be 1 mM in the media, diluted to 10 μM in the homogenate). In fact, nicotinamide (10 μM) produced no significant effect on the fluorescence of the homogenate after previous treatment with nicotinamide (100 μM). Nicotinamide was therefore routinely injected before assay of media samples.

Astrocyte and AR media samples elicited calcium release in homogenate pretreated with heparin (to block IP3 responses) and to nicotinamide (Figs. 3A, 4A), indicating the presence of a calcium-mobilizing agent other than IP3 and nicotinamide. Response to astrocyte and AR samples was almost completely abolished in homogenates desensitized to nicotinamide and to cADPR (Figs. 3B, 4B), indicating that the sample activity was attributable to nicotinamide and to cADPR only. The astrocyte and AR samples also desensitized homogenate to further calcium release by cADPR but not to calcium release by IP3 (Figs. 3C, 4C). In summary, once the effect of nicotinamide was suppressed, culture media samples were shown to desensitize homogenate to cADPR and, conversely, cADPR was shown to desensitize homogenate to the media samples. These results strongly support the hypothesis that cADPR was present in the media of cultures incubated with NAD. A time course of the synthesis of cADPR from NAD in astrocyte cultures in medium with and without calcium is shown in Figure 5. Maximal extracellular concentrations of cADPR were attained by ~60 min, and no evidence of clearance of cADPR was observed.

Determination of the specific activity of cADPR synthesis in cortical cultures

To investigate whether cADPR synthesis activity was localized to neurons or astrocytes, we determined the ADP-ribosyl cyclase activity in AP, AR, and astrocyte cultures. Cultures were incubated with NAD (1 mM) in HBSS (with calcium), and samples of the media were assayed for calcium-releasing activity at 60 min, after calcium had been removed from the medium by the use of Chelex ion-exchange resin. AR and astrocyte media samples appeared to have
comparable activity: 0.84 ± 0.06 nmol cADPR/mg protein/hr for astrocyte cultures (n = 3; mean ± SEM) and 0.96 ± 0.18 nmol cADPR/mg protein/hr (n = 5; mean ± SEM) for AR cultures (Table 3). No calcium-releasing activity distinguishable from that caused by the NAD present was observed in the AP cultures (n = 2).

Because AP cultures contain less protein than astrocyte cultures or AR cultures, we were concerned that the production of cADPR by these cultures might be significant but below the level of detection of the bioassay. To pursue this, we used a sensitive fluorometric assay based on the work of Graeff et al. (1994, 1996) that showed that NGD is a substrate for ADP-ribosyl cyclase, yielding a fluorescent product, cGDPR. Figure 6 shows data from one representative experiment of three that were performed. When astrocyte cultures and AR cultures were incubated with NGD (100 μM), which is not fluorescent, they caused a robust time-dependent increase in fluorescence in the extracellular medium, which is attributable to the accumulation of cGDPR (Graeff et al., 1994). In contrast, AP cultures caused a very small increase in the fluorescence of the medium that was significantly different (p < 0.05) from the control values [NGD (100 μM) in EBSS that was not exposed to cultures] in two of three experiments. In three experiments, the mean increase in fluorescence intensity in the medium exposed to AP cultures at 60 min was 4.4 ± 0.2% (mean ± SEM) of the fluorescence intensity appearing in the medium exposed to AR cultures and 4.8 ± 0.4% of the fluorescence intensity appearing in the medium exposed to astrocyte cultures. In these three experiments, the protein content was 42 ± 8, 177 ± 12, and 154 ± 6 μg in the AP, AR, and astrocyte cultures, respectively.

**Evidence for an extracellular localization of ADP-ribosyl cyclase activity**

The fact that formation of cADPR could be demonstrated by adding NAD to the extracellular medium suggested that ADP-ribosyl cyclase activity associated with cortical cultures was itself an extracellular enzyme activity. To pursue this further, we ex-
posed cultures to NAD (1 mm) in the extracellular medium with and without 0.25% Triton X-100, and compared ADP-ribosyl cyclase activity and LDH in the extracellular medium (Table 1). The presence of Triton X-100 in the medium increased the LDH activity 31.3 ± 13 fold (n = 3) without significant effect on the ADP-ribosyl cyclase activity.

Demonstration of cADPR production in astrocyte cultures by HPLC

We sought to confirm the production of cADPR in astrocyte cultures directly by HPLC. Astrocyte dishes and wells were incubated at 37°C with complete HBSS containing NAD (1 mm) and media samples were assayed using a two-column HPLC system (Kim et al., 1993) (see Materials and Methods). A peak, which was absent in media controls and in media of cultures incubated with HBSS only, appeared at the retention time for cADPR of 6.2 min (Fig. 7A, upper trace, downward arrow).

To confirm the identity of this peak as cADPR, samples were placed in boiling water for 30 min, which converts cADPR quantitatively to ADPR but does not affect ADPR (Kim et al., 1993). This treatment degraded the 6.2 min peak in the medium sample (compare upper trace with lower trace in Fig. 7A at downward arrow). Furthermore, we confirmed that authentic ADPR was unaffected by boiling (Fig. 7B, upper chromatogram), whereas authentic cADPR was quantitatively converted to ADPR by boiling (Fig. 7B, middle chromatogram). The elution of authentic cADPR before boiling is shown in Figure 7B, lower chromatogram. In addition, we verified that authentic cADPR coeluted with the putative cADPR (data not shown).

cADPR hydrolysis in cortical cell cultures

ADP-ribosyl cyclase activity has been reported to copurify with cADPR hydrolyase activity in several different cell types, and particularly in mammalian cells with extracellular activity (Franco et al., 1993; Lee et al., 1993; Takasawa et al., 1993b), leading to the proposal that NADase is a bifunctional enzyme (Kim et al., 1993). We therefore investigated the presence of cADPR hydrolyase activity in the cortical cultures. Cultures were incubated for 60 min at 37°C in HBSS containing cADPR (300 μM), and media samples were assayed by HPLC. We found a small but significant decrease in the concentration of cADPR in the medium exposed
to astrocyte and mixed cultures compared with medium that had been incubated under the same conditions but without cells (Table 2). These changes in ADPR concentration correspond to a rate of hydrolysis of cADPR of 167 and 111 nmol/mg protein/hr, respectively (Table 3). There was an expected concomitant although smaller increase in ADPR concentration that also was significant (Table 2). These changes in ADPR concentration correspond to a rate of hydrolysis of cADPR in the astrocyte and mixed cultures of 24 and 18 nmol/mg protein/hr, respectively, considerably less than that calculated on the basis of changes in cADPR concentration (Table 3). In media exposed to AP cultures, there was neither a significant decrease in cADPR nor an increase in ADPR, as expected given the absence of ADP-ribosyl cyclase activity in these cultures.

**DISCUSSION**

The objective of this study was to demonstrate, quantify, and localize ADP-ribosyl cyclase and cADPR hydrolase activity in rat cortical cultures. The evidence for ADP-ribosyl cyclase activity in cortical cell cultures presented in this study rests on the selective desensitization properties of the sea urchin egg homogenate bioassay and a demonstration of cADPR in media samples by HPLC. When sea urchin egg microsomes are exposed to either cADPR (100 nm) or IP$_3$ (1 μM), they become desensitized and no longer release calcium in response to subsequent injections of the same agent, but retain a response to the other agent. Injection with heparin (250 μg/ml) also results in desensitization to IP$_3$. The desensitization criterion led us to rediscover the calcium-releasing activity of nicotinamide (Clapper et al., 1987), and made it possible to account for and almost completely eliminate residual non-cADPR and non-IP$_3$ calcium-releasing activity in experimental samples. Homogenate desensitized to both nicotinamide and IP$_3$ released calcium in response to media samples from AR and astrocyte cultures incubated with NAD (Figs. 3A, 4A), whereas homogenate desensitized to nicotinamide and cADPR no longer did (Figs. 3B, 4B). Furthermore, multiple injections of media samples desensitized the homogenate to further calcium release by authentic cADPR, but not by IP$_3$ (Figs. 3C, 4C). These results provide strong evidence of the presence of cADPR in the extracellular medium.

We observed a large increase in fluorescence after heparin...
injection (Figs. 3A, 4A). Heparin is known to activate ryanodine receptors in a calcium-dependent manner, and the fact that we saw a fluorescence increase with heparin suggests that the calcium concentration in the homogenate was high enough (>20 nm) to have permitted heparin-activated release of calcium mediated by ryanodine receptors (Ehrlich et al., 1994). The responses to the culture media were smaller after heparin injection (compare responses in Figs. 3A, 4A with responses in Figs. 3C, 4C). It seems most likely that this difference is attributable either to partial desensitization of the ryanodine receptor by heparin, to partial depletion of the cADPR-sensitive calcium stores, or to nonlinearity of the Fluo 3 fluorescence response.

Additional confirmation of the production of cADPR by cortical cultures was sought by HPLC. No cADPR was detected in NAD standards or in the media of cultures incubated with HBSS alone. A peak with the retention time characteristic of cADPR did appear in a time-dependent manner in media samples from cultures incubated with NAD, strongly suggesting the presence of cADPR (Fig. 7A). The specific susceptibility of the 6.2 min peak to boiling (Kim et al., 1993), analogous to the behavior of authentic cADPR but not ADPR (Fig. 7A, B), as well as coelution with authentic cADPR, offered additional proof of its identity as cADPR.

In summary, three different pieces of evidence support the identification of cADPR in the medium of cortical cultures incubated with NAD: (1) the medium caused release of calcium in sea urchin egg homogenates desensitized to nicotinamide and IP₃, but not in homogenates desensitized to nicotinamide and cADPR; (2) the medium desensitized homogenate to further release of calcium by cADPR but not by IP₃; (3) an HPLC peak is present in the medium that has the same retention time as cADPR. This peak is not present in medium of cultures incubated without NAD, nor is it present in NAD; this peak is selectively degraded by boiling, which also degrades authentic cADPR but does not affect ADPR, and it coelutes with authentic cADPR.

The ADP-ribosyl cyclase activity appears to localize to astrocytes. No calcium release and therefore no detectable cADPR formation was observed in the medium of AP cultures incubated with NAD (1 mM). This result suggests that embryonic neurons in culture do not have detectable ADP-ribosyl cyclase activity. Previously, it has been shown that embryonic neurons in culture do not possess detectable β-adrenergic receptors (Trimmer et al., 1984), whereas adult cortical and hippocampal neurons clearly do (Madison and Nicoll, 1982, 1986a,b). Therefore, the absence of ADP-ribosyl cyclase activity from embryonic neurons in culture may be attributable to an effect of growing the cells in tissue culture or to a developmental phenomenon. In fact, CD38 has been demonstrated immunocytochemically in the cell bodies and dendrites of neurons in the adult human cerebellum and cerebral cortex (Mizuguchi et al., 1995).

The ADP-ribosyl cyclase activity in astrocytes of 0.94 ± 0.06 nmol cADPR/mg protein/hr determined in this study is lower than previously reported values of 4.5 ± 0.1 nmol/mg/hr for salamander brain, 20.6 ± 2.8 nmol/mg/hr for chick embryonic brain, 62.7 ± 6.2 nmol/mg/hr for dog brain (Lee and Aarhus, 1993), and 300 nmol/mg/hr reported for pig brain (Gu and Sih, 1994). There are several possible explanations for this discrepancy. One is that the cited studies were performed using adult animals, and therefore a developmental difference might account for the difference in activity as well as for the apparent lack of activity associated with neurons, cited above. By this reasoning, the increase in activity seen in whole adult brain might be attributable to a developmental increase in activity associated with astrocytes or to the emergence of activity associated with neurons. Almost all of the activity seen in culture is extracellular, and thus it also is possible that with development, a significant increase in intracellular activity occurs. Furthermore, there may be significant regional differences in the expression of ADP-ribosyl cyclase activity in neurons and astrocytes in the brain, and cells other than neurons and astrocytes (e.g., oligodendrocytes, endothelial cells, microglia) might exhibit high activity. Finally, there may be significant species differences.

We directly demonstrated cADPR hydrolyase activity in the cultures by incubating them with cADPR (300 μM). Hydrolyase activity has been found associated with cyclase activity in all other tissues except Aplysia, leading to the suggestion that mammalian NADase is a bifunctional enzyme (Kim et al., 1993). In previous studies, the cADPR hydrolyase activity of the NADase CD38 was

---

**Table 1. Effect of permeabilization of cultures on extracellular ADP-ribosyl cyclase and lactate dehydrogenase activities**

<table>
<thead>
<tr>
<th></th>
<th>ADP-ribosyl cyclase activity (nmol/mg protein/hr)</th>
<th>Lactate dehydrogenase activity (U/ml medium)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBSS</td>
<td>1.25 ± 0.06</td>
<td>1.1 ± 0.2</td>
</tr>
<tr>
<td>HBSS + 0.25% Triton X-100</td>
<td>1.19 ± 0.04</td>
<td>36.4 ± 14.7</td>
</tr>
</tbody>
</table>

AR cultures were incubated with 1 mM NAD in HBSS with or without 0.25% Triton X-100 for 1 hr at 37°C in a shaking water bath, after which medium samples were bioassayed for cADPR activity. Data represent mean ± SEM from three separate experiments.
determined to be ∼10 times the ADP-ribosyl cyclase activity (Franco et al., 1993; Inageda et al., 1995). Similar results were obtained with a purified NADase from splenic lymphocytes (Kim et al., 1993). Therefore, the results obtained in this study are qualitatively consistent with what has been seen before. However, cADPR hydrolase activity in the astrocyte and AR cultures appears to be ∼100-200 times the ADP-ribosyl cyclase activity, based on loss of cADPR from the extracellular medium (Table 3). This could be attributable to the involvement of an NADase other than CD38, to modulation of CD38 resulting in different relative synthetic and hydrolytic rates, or to the hydrolytic action of an enzyme other than cADPR hydrolase on cADPR. For example, we have acquired evidence for the existence of a novel extracellular cyclic nucleotide phosphodiesterase in cortical cultures (Rosenberg and Dichter, 1989; Rosenberg and Li, 1994), and it is possible that this as-yet uncharacterized enzyme might possess some activity against cADPR. Perhaps a more reliable estimate of cADPR hydrolytic activity may be obtained by basing it on the accumulation of ADPR (Table 3). In this case, cADPR hydrolase activity would be 29 and 19 times the synthetic activity, values quantitatively similar to those found previously for a splenic NADase (Kim et al., 1993).

The results presented here suggest that most of the ADP-ribosyl cyclase activity in cells derived from embryonic rat cerebral

Figure 7. Confirmation by HPLC of cADPR production by astrocyte cultures. Astrocyte dish cultures were incubated with complete HBSS with NAD (1 mM) at 37°C for 120 min. Media samples (50 μl) were run on a two-column HPLC system (PL-SAX column followed by a C18 reverse-phase column) (see Materials and Methods) and assayed by absorbance detection. The selective degradation of cADPR by boiling for 30 min was used as an additional test for cADPR. The solid upward arrows indicate the time of injection of sample. A, Degradation by boiling of 6.2 min peak in astrocyte incubation medium sample by boiling. Upper trace, Astrocyte medium sample (120 min incubation) before boiling. A peak is present at 6.2 min, indicated by downward open arrow, which is the retention time of cADPR. Lower trace, Astrocyte medium sample (120 min incubation) after boiling for 30 min. Boiling has removed the peak at 6.2 min, indicated by downward open arrow. B, Boiling for 30 min degrades authentic cADPR to ADPR but does not affect ADPR. Upper trace, ADPR (5 μM) standard after 30 min boiling. The retention time is 10.8 min, the value previously determined for unboiled ADPR. Middle trace, cADPR (2 μM) standard after boiling for 30 min. The cADPR peak at 6.2 min is no longer visible; instead, a peak has appeared at 10.8 min, which is the ADPR retention time. Bottom trace, cADPR (2 μM) standard before boiling for 30 min. cADPR elutes as a sharp peak at 6.2 min.
Table 2. Evidence for cADPR hydrolase activity in cultures derived from rat cerebral cortex

<table>
<thead>
<tr>
<th></th>
<th>cADPR</th>
<th>ADPR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>234 ± 14</td>
<td>9.9 ± 1.9</td>
</tr>
<tr>
<td>n = 12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Astrocytes</td>
<td>216 ± 22</td>
<td>12.6 ± 2</td>
</tr>
<tr>
<td>n = 8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>p &lt; 0.05</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AR cultures</td>
<td>212 ± 10</td>
<td>13.5 ± 1.1</td>
</tr>
<tr>
<td>n = 8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>p &lt; 0.05</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Cultures were incubated with 300 μM cADPR for 60 min at 37°C, after which samples of the medium were taken for direct assay of cADPR and ADPR by HPLC. For comparison, to control the possibility of nonenzymatic hydrolysis, medium containing 300 μM cADPR was incubated under similar conditions without cells. At the end of the incubation period, medium was collected and assayed by HPLC. Analysis of significance was by Student’s two-tailed unpaired t test. The mean protein of astrocyte cultures in these experiments was 54 ± 7 μg, and for mixed cultures it was 99 ± 19 μg per 12 mm coverslip culture. Concentrations are given in micromolars. Data are pooled from four separate experiments, and number of determinations (n) are given that were performed.

Table 3. ADP-ribosyl cyclase and cADPR hydrolase activity in cultures derived from rat cerebral cortex

<table>
<thead>
<tr>
<th></th>
<th>Astrocytes</th>
<th>AR cultures</th>
<th>AP cultures</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADP-ribosyl cyclase</td>
<td>0.84 ± 0.06</td>
<td>0.96 ± 0.18</td>
<td>ND</td>
</tr>
<tr>
<td>cADPR synthesized/mg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>protein/hr (mean ± SEM)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>cADPR hydrolyse (1)</td>
<td>167</td>
<td>111</td>
<td></td>
</tr>
<tr>
<td>(nmol cADPR degraded/mg protein/hr)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>cADPR hydrolyse (2)</td>
<td>24</td>
<td>18</td>
<td>ND</td>
</tr>
<tr>
<td>(nmol ADP accumulation/mg protein/hr)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For determination of ADP-ribosyl cyclase activity, cultures were incubated with 1 mM NAD in HBSS for 1 hr at 37°C in a shaking water bath, after which samples were bioassayed for cADPR activity (see text). cADPR hydrolyase rates are given based both on cADPR loss and on ADPR gain in extracellular medium, based on data shown in Table 2. ND, Not detectable.

cortex is extracellular and that this extracellular activity is associated with astrocytes but not neurons. However, all that is known about the biological role of cADPR as a mobilizer of calcium release presumes an intracellular site of action. The cADPR produced by extracellular cyclase could therefore have a novel biological function, could somehow be transported inside the cell to exert its known effect there, or could be involved in activation of NADases, which themselves might play a key role in signal transduction (Kim et al., 1994).

REFERENCES


