NEURAL RESPONSES TO INJURY:
PREVENTION, PROTECTION, AND REPAIR
Annual Technical Report
1996

Submitted by
Nicolas G. Bazan, M.D., Ph.D.
Project Director

Period Covered: 20 September, 1995, through 19 September, 1996

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between

United States Army Medical Research and Development Command
(Walter Reed Army Institute of Research)

and

Louisiana State University Medical Center
Neuroscience Center of Excellence

Volume 1 of 9
COOPERATIVE AGREEMENT NUMBER DAMD17-93-V-3013

TITLE: Neural Responses to Injury: Prevention, Protection and Repair

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Fort Detrick, Frederick, Maryland  21702-5012

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DTIC QUALITY INSPECTED
The LSU Neuroscience Center is a comprehensive, multidisciplinary, and trans-departmental entity that unites fundamental neurobiology and the clinical neurosciences in the common goal of elucidating the workings of the brain and contributing to the treatment of currently incurable diseases of the nervous system. The objective of this program is to find solutions to neuroscience-related problems of interest to the US Army Medical Research and Development Command. The program is focused on exploiting novel neuroprotective strategies that lead to prevention of and repair after neural injury. Converging approaches using state-of-the-art tools of cell biology, neurochemistry, neuroimmunology, neurophysiology, neuropharmacology, molecular biology and virology are ongoing. This program's yearly activities have included: 1) seven research projects in the basic and clinical neurosciences; 2) the expansion of central, shared facilities with the addition of highly specialized instrumentation not currently available to our scientists; 3) the development of laboratory space to permit the physical consolidation and coordination of this research effort; and 4) the operation of a coordination unit to monitor, facilitate, and administrate the cooperative research programs, as well as to meet the associated budgetary, human resource, facility, and communication needs for the attainment of the program goals.
FOREWORD

Opinions, interpretations, conclusions and recommendations are those of the author and are not necessarily endorsed by the U.S. Army.

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[Signature]

PI - Signature Date
Nicolas G. Bazan, M.D., Ph.D. Oct 96
Volume 1 Neuroscience Core Research Facilities

Project Directors: R. Ranney Mize, Ph.D.
Nicolas G. Bazan, M.D., Ph.D.

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Volume 4 Neurochemical Protection of the Brain, Neural Plasticity and Repair

Project Director: Nicolas G. Bazan, M.D., Ph.D.

Participating Scientists: Geoffrey Allen, Ph.D.
Victor Marcheselli, Ph.D.
John Hurst, Ph.D.
Leo Happel, M.D.
Walter Lukiw, Ph.D.

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Protecting the Auditory System and Prevention of Hearing Problems

Project Directors: Richard Bobbin, Ph.D.
Charles Berlin, Ph.D.

Participating Scientists: Sharon Kujawa, Ph.D.
Carlos Erostegui, M.D.
Douglas Webster, Ph.D.


Volume 7 Role of Growth Factors and Cell Signaling in the Response of Brain and Retina to Injury

Project Directors: Prescott Deininger, Ph.D.
Nicolas G. Bazan, M.D., Ph.D.

Participating Scientists: Julia Cook, Ph.D.
Haydee E. P. Bazan, Ph.D.
William C. Gordon, Ph.D.
Elena Rodriguez De Turco, Ph.D.
Victor Marcheselli, Ph.D.

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Project Director: Herbert E. Kaufman, M.D.
Roger Beuerman, Ph.D.

Participating Scientists: Claude A. Burgoyne, M.D.
Emily Varnell
Mandi Conway, M.D.

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Cooperative Agreement Between the US Army Medical Research and Development Command and The LSU Neuroscience Center of Excellence

DAMD17-93-V-3013  20 September, 1993 - 19 October, 1997  $13,860,000

LSU MEDICAL CENTER  
SCHOOL OF MEDICINE NEW ORLEANS

LSU NEUROSCIENCE CENTER OF EXCELLENCE

Repair and Regeneration of Peripheral Nerve Damage
Roger Beuerman, Ph.D.
David Kline, M.D.
Austin Sumner, M.D.
Project Directors

The Neuroimmunology of Stress, Injury, and Infection
Bryan Gebhardt, Ph.D.
Project Director
Daniel Carr, Ph.D.
Project Co-Directors

Neurochemical Protection of the Brain, Neural Plasticity and Repair
Nicolas G. Bazan, M.D., Ph.D.
Project Director

Neural Responses to Injury: Prevention, Protection and Repair
Nicolas G. Bazan, M.D., Ph.D.
Program Director

EXTERNAL ADVISORY REVIEW COMMITTEE

NEURAL RESPONSES TO INJURY: PREVENTION, PROTECTION AND REPAIR

NEUROSCIENCE CORE RESEARCH FACILITIES

EXPANSION OF PHYSICAL FACILITIES

Neuropharmacology of Delta Receptor Agonists and Antagonists
Joseph Moerschbaecher, Ph.D.
Project Director

Protecting the Auditory System and Prevention of Hearing Problems
Richard Bobbin, Ph.D.
Charles Berlin, Ph.D.
Project Directors

Role of Growth Factors and Cell Signaling in the Response of Brain and Retina to Injury
Prescott Deininger, Ph.D.
Nicolas G. Bazan, M.D., Ph.D.
Project Directors

Vision, Laser Eye Injury and Infectious Diseases
Herbert E. Kaufman, M.D.
Roger Beuerman, Ph.D.
Project Directors
October 18, 1996

Commander
U.S. Army Medical Research and Material Command (USAMRMC)
ATTN: MCMR-RMI-S
Building 504
Fort Detrick
Frederick, MD 21702-5012

RE: Third Annual report, Cooperative Agreement No. DAMD17-93-V-3013
Neural Responses to Injury: Prevention, Protection, and Repair

Dear Sir,

Please find enclosed the original and two copies of the third annual report for the Cooperative Agreement, reference above, between the USAMRMC and the Louisiana State University Medical Center School of Medicine, Neuroscience Center of Excellence. This report represents the research carried out during the third year of this agreement (20 September, 1995, to date). It is organized per project, each corresponding to a chapter of the original application. The critiques of the projects, as reported to us after the second annual report was reviewed by your office, have been taken into consideration in the preparation of these technical reports, and we believe that your comments concerning the last report have enabled us to more concisely report our progress.

We are very pleased with the progress that has been made in the third year of this agreement. As you will observe from the enclosed technical reports, our investigators have enjoyed a great deal of success on the projects, judging by their publications as well as by the technical reports themselves.

Unlike the first annual evaluation of the program, because of time and budgetary constraints, we did not request that the external advisory committee of the Center review this report.

We are arranging a meeting between the LSU investigators and their counterparts in the Army to provide program briefings for the work that we are planning to conduct under the fourth year of this agreement as well as to exchange ideas and information of mutual interest. This provides both the LSU and Army scientists the opportunity to discuss the work being done, the direction, and the significance to problems of interest to the Department of Defense.
Annually, the Neuroscience Center hosts a retreat for all faculty, staff, and students of, or interested in the Center. This day-long event is a forum for the research interests of each participant, featuring poster presentations of ongoing work. This year (February 1, 1997) the collaborating scientists, as well as any other interested investigator from the USAMRMC, are being invited to participate in this forum.

I am very pleased to inform you that the construction for the two additional floors of research space which are to be added to the Lions/LSU Clinics Building, 2020 Gravier Street, New Orleans, LA, is near completion. These two new floors will provide additional laboratory space for the LSU Neuroscience Center as well as for the housing of the shared Core Research Facilities equipment. Office space will also be available for the faculty and staff of the LSU Neuroscience Center of Excellence.

Please let me know if there is any further information that I can provide you.

Sincerely,

Nicolas G. Bazan, MD, PhD
Boyd and Villere Professor of
Ophthalmology, Biochemistry
and Molecular Biology, and Neurology
Director, LSU Neuroscience Center
NEUROSCIENCE CORE RESEARCH FACILITY

Core Research Facility Equipment at the Medical Education Building

In Year 3 of the DOD Agreement, the Calcium Imaging Facility (CIF) has continued to support DOD-Funded projects and to serve as a core facility for Neuroscientists throughout the LSUMC campus. Usage of the Facility has been divided up at approximately 20% hardware and software maintenance and development, 40% DOD-funded investigations, and 40% other neuroscience projects. Further descriptions of the specific projects are given in subsequent sections regarding the Medical Education Building.

Core Research Facility Equipment at the Eye Center

A second color dye sublimation printer has been added to the imaging facility lab at the Lions Eye building to facilitate production of high quality color and black and white printouts.

A Pentium-based PC workstation has been added to the imaging facility lab. This PC will be used for two dimensional analyses of data obtained from video cameras attached to the confocal microscope as well as other compatible microscopes at the Neuroscience Center. This workstation has helped solve some problems that Noran could not solve with their 2D software. In addition, images obtained with video cameras are in color, and this information is often important in image analysis; the confocal microscope obtains data via PMTs which are on a grey scale, so no color information is included. A low light level, cooled chip camera will reside with this PC imaging workstation. This camera has the ability to integrate low level light or fluorescent signal and can also store up to two images in memory, allowing for three levels of color overlay from the same
field. The image can then be sent to a color monitor and/or to the PC imaging workstation (by use of a frame grabber) for analysis. It is planned that the PC imaging workstation will also serve as an image archive for the Neuroscience Center with the purchase of a rewritable gigabyte jukebox tower that will allow multiple swapable gigabyte disks to be loaded and analyzed.

A heated perfusion system has been purchased for the confocal microscope. This system will address the importance of temperature and on/off signals for analysis of cellular responses. The system is compatible with one used at the MEB.

**Maintenance and Instrument Development at the Medical Education Building**

The Facility has continued to add equipment useful to investigators that utilize the Noran Laser Scanning Confocal Microscope (LSCM). The main addition this year was a dual-technology, Dye-sublimation/Thermal wax color printer. This printer allows rapid access to a hard-copy of imaging results and enables us to produce publication quality images. Also, a perfusion chamber is now available so that bathing solutions can be changed rapidly while imaging cells. This has proved most useful in the studies where drugs need to be added to cells while measuring $[\text{Ca}^{2+}]_i$. Other equipment now available, includes a tissue slicer equipped for the production of living brain slices and a 1.3 Gb optical disk drive for off-line archiving of images. Software development in the Facility has also progressed. The electrophysiology data acquisition software, pClamp 6.0, has been installed and used in some preliminary trials. Noran has released a new version of their InterVision software for both image acquisition and image analysis. This software incorporates several new features requested by LSUMC users, including time-lapse capabilities, better pseudo-color capabilities, and image integration. It also has expanded image storage capacity so that more images
can be stored during a single experiment. All of these added capabilities have been installed, and their use is being fully explored as experiments proceed.

The Service contract on the LCSM, funded by the DOD, has played a vital role in maintaining the CIF this year. In December, several investigators noted a deterioration in the quality of the images collected in the facility. Consultation with Noran’s technical support determined that this was due to a combination of factors, including a broken primary dichroic mirror holder and reduced efficiency of the laser. Although it required two visits from the technical support personnel and about six weeks of down-time for the CIF, both of these problems were fixed, and we ended up with a new laser that has better performance than the original one. More recently, there was an increase in the number of system errors and crashes involving the on-line optical disk used for image storage. Again, the problem was quickly resolved with the help of Noran’s technical support, and the facility was operational with minimal down-time.

Maintenance and Instrument Development at the Eye Center

Dr. DeCoster has spent the necessary time to link the confocal and other imaging workstations and PCS via the local area network so that large images may be sent by using FTPs or other network utilities for printing or image analysis. This saves time that would otherwise be used shuffling disks between PCS and workstations so that files could be printed or analyzed. In addition, Dr. DeCoster has maintained the Indy workstation so that compatible software (such as Adobe Photoshop) can be used on both Silicon Graphics and Windows/DOS platforms. Dr. DeCoster also serves as the administrator for the imaging workstations in the imaging facility lab at the Neuroscience Center, dealing with such issues as data storage and file management, printing, slide
making, and microscope/camera use, as well as the necessary training needed for other scientists and technicians to make use of the instrumentation.

**DOD-Funded Investigations**

*Current research projects at the Medical Education Building*

Drs. Mize and Bobbin continue to be the principal DOD-funded users of the MEB Facility. They each have either finished, or are still working on, several projects utilizing the CIF equipment.

a) Distribution and Function of Calcium Binding Proteins in the Rat Superior Colliculus.

(R.J. Cork, F-S. Lo and R.R. Mize)

This is an ongoing project with Dr. Mize as P.I. Major progress was made this year, in that this project successfully competed for NIH funding from the National Eye Institute, in large part based on preliminary data collected in the CIF using DOD support. So far, the distributions of the calcium binding proteins (CaBP), Calbindin (CB) and Parvalbumin (PV), in neurons of the rat superior colliculus (SC) have been mapped using fluorescently labeled antibodies; and electrophysiology experiments, done by Dr. Lo, have characterized some of the membrane properties of cells in the optic layer, some of which contain CB. The two CaBP appear to be expressed in complementary sublaminar tiers that represent functional layers of neurons distinct from the morphological laminae of the colliculus. There is a dense band of CB-cells, centered in the optic layer. Electrophysiological recordings made from the optic layer reveal that these cells have characteristic membrane properties, including low threshold calcium spikes and a unique high frequency burst firing mode.
These experiments use the multilabel protocols in the InterVision system, and customized look-up tables have been developed to display double-labeled images in pseudocolor. Two abstracts, detailing the results of these experiments, were presented at the Society for Neuroscience meeting in November '95 (San Diego, CA), and a manuscript describing the distributions of the CaBP in the SC is about to be submitted.

b) Involvement of NOS in synaptic plasticity.

(R.J. Cork, F-S. Lo, and R.R. Mize)

This new project is shortly to be awarded NIH funding from the National Institute of Neurological Diseases and Stroke, as a joint grant to the three co-investigators. Much of the preliminary data for the grant application was obtained with CIF support. This data included preliminary [Ca$^{2+}$], measurements, electrophysiology, and immunocytochemistry. The project, which will start early in 1997, seeks to explore the role of nitric oxide synthase in directing axons to their targets in the developing rodent SC. This will be approached in three ways; electrophysiology, calcium imaging, and immunohistochemistry. The studies will principally focus on any interactions between NO, NOS, and the NMDA receptor during the early postnatal development of the colliculus. The results of early experiments suggest that NO is not involved in the formation of the patch/cluster system in the intermediate gray layer of the SC (Mize et al., 1996). This may be because the patches are composed of cholinergic afferents rather than glutamatergic ones. Drs. Lo, Cork and Mize have also begun to examine the development of synaptic currents, particularly NMDA, and the onset of long term depression (LTD) and potentiation (LTP) in the neonatal rat. This data will be compared to the development of NOS expression in the rodent SC.
c) Production of an acutely dissociated cell preparation suitable for calcium imaging.
(C. Leblanc, R.J. Cork and R.R. Mize)

This was a short project done by a Neuroscience graduate student who did a lab rotation in the CIF. The aim was to work out conditions for producing dissociated SC neurons that could be loaded with calcium indicators and imaged to measure either resting $[\text{Ca}^{2+}]_i$ levels or transient calcium responses to neurotransmitters. Such a preparation is an important part of the above two projects. Most of this project concentrated on determining the best methods for cell dissociation and the best ways to obtain maximum numbers of viable neurons in the preparations. Several successful experiments were done making dissociated cells and recording calcium transients induced in them with neurotransmitters. Some of these results were included in the preliminary data for the NIH grant application described in (b) above.

d) Calcium Transients in Response to Extracellular ATP in Isolated Cochlear Hair Cells.
(C. Chen, R.J. Cork, and R.P. Bobbin)

To extend their patch-clamp studies, Drs. Chen and Bobbin recorded calcium transients in the cells of their in vitro cochlear preparation. Preliminary experiments have shown that these cells can be loaded with indo-I-AM, and transient elevations of cytoplasmic calcium have been recorded in response to extracellularly applied ATP. These measurements will be extended to different cochlear cell types to determine if ATP functions as an extracellular signaling molecule. The patch-clamp equipment in the CIF has been installed and tested successfully with the cochlear preparation. In future it is expected that patch-clamp experiments will be combined with simultaneous calcium measurements. The preliminary data collected in the CIF has been included in a grant proposal submitted to NIH by Drs. Chen and Bobbin. Dr. Cork is included on that grant as a collaborator, and
a significant portion of the experiments will be done in the CIF.

**Current research projects at the Eye Center**

a) Role of lipid mediators in neuronal signaling processes.

(M. A. DeCoster and N. G. Bazan).

PAF continues to be tested in a complex set of experiments to address the immediate, long term, independent, and modulatory roles of this potent lipid mediator. While our research results are supported by recently published results (from other laboratories) that PAF rarely exerts a direct calcium increase on neurons, we have for the first time shown the neuroprotective effects of PAF receptor antagonist against excitatory amino acid toxicity to primary cultured hippocampal neurons. PAF alone may provide neuroprotective effects to neurons in culture. This is consistent with our observed results that long-term treatment with PAF may desensitize neurons as far as calcium response, to subsequent glutamate stimulation.

b) Phospholipases A₂ as factors in neuronal toxicity.

(M. A. DeCoster, M. Kolko, E. Rodriguez-de Turco, and N. G. Bazan).

The toxicity and lipid metabolism studies of PLA₂'s effect of primary cortical neuronal cultures have been completed to the extent that this work has been accepted for publication in a high priority scientific journal (Journal of Biological Chemistry, December, 1996). The calcium studies using PLA₂s will be published as a separate work and is still in progress. The major finding from these studies was that the PLA₂s and glutamate work at the level of toxicity and lipid metabolism by different mechanisms and can show a synergistic effect when added together. The calcium studies are consistent with this so far, and will be guided by these soon to be published results.
Other Projects

Ongoing research projects at the Medical Education Building

a) Dr. Cindy Linn is studying calcium channels in catfish retinal cells. She is combining electrophysiology techniques with fluorescent calcium indicator measurements. Several successful experiments have been done using both ratio imaging and the continuous “Ratio Over Time” capability of the InterVision system.

b) Drs. Fu-Sun Lo and Bill Guido have been studying synaptic transmission in the rat LGN during early postnatal development. They find that the onset of EPSP’s, IPSP’s and LTP/LTD are developmentally regulated.

Ongoing research projects at the Eye Center

a) (Drs. M. A. DeCoster and N. G. Bazan). 3D reconstructions and animated movies of neurons and astrocytes from primary cell cultures, as well as brain tissue sections, continue to be used to model methods for scanning and analyzing cellular association and orientation via the confocal microscope.

b) (Drs. M. A. DeCoster, D. Linn, and N. G. Bazan). Fluorescent dyes have been injected into intact retinal tissue and imaged and analyzed using the confocal microscope. Calcium dynamics in stimulated retinas have been established both in control tissues as well as in retinas treated with calcium channel blockers and other antagonists. In addition, work is now being carried out to analyze calcium responses in retinas from different genetically engineered rodent models.

c) (Drs. M. A. DeCoster, Ying Tao, H. Bazan, and N. G. Bazan). Fluorescent calcium dyes have been loaded into primary cultured rabbit corneal epithelial cultures. It has been determined that
PAF (100 nM) induces a delayed, transient increase in intracellular calcium concentration ([Ca^{2+}]_{i}) in these cell cultures. The PAF receptor antagonist BN50730 blocks this calcium response to PAF. The PAF response is largely dependent upon extracellular calcium influx. Calcium channel blockers also block the response to PAF. This work is currently being prepared for submission for publication and has already been presented in abstract form.

Goals for Year 4

Future goals at the Medical Education Building

Considerable progress has been made this year in acquiring new funding for research projects using CIF equipment. Dr. Mize submitted 2 NIH grant applications, one of which was funded in May 1996, the other to be funded in January 1997. Dr. Bobbin is currently resubmitting an NIH grant application, which if successful should start in mid 1997. All three of these grant applications have included preliminary data collected in the CIF, and they all fund continuing research requiring use of the CIF equipment. If all three projects are actively using the CIF, then together with Dr. Linn’s project and Dr. Guido’s, the Facility will be running at near capacity. There may be some time for short term research projects by other researchers. It is expected that the CIF will continue to play an important support role in the research projects of these LSUMC Neuroscientists. It is expected that the CIF users will continue to produce nationally recognized work that will enhance the reputation of LSUMC Neuroscience.

Future goals at the Eye Center

The imaging facility at the Neuroscience Center in the Lions Eye building is at a stage of rapid progress both physically and scientifically. More investigators are using the facility now for
a wide array of projects from calcium imaging of living cells to cellular quantification and morphometric measurements of variously stained tissues and cell cultures. The computer processing and storage needs and capabilities have grown in tandem, and many of the workstations are now interconnected via networking utilities. When the imaging facility prepares to move to the expanded floors of the Lions Eye building in 1997, the space restrictions currently experienced should be alleviated. At that time, Dr. DeCoste will be the coordinator for the imaging facility, managing computer and confocal microscope usage for the Neuroscience Center; these facilities will also be used in his own research. Technical support will most likely be retained to help with data analysis, file management, and presentation production associated with the imaging facility.
Publications


Cork, R.J., Baber, S.Z. and Mize, R.R (1996) Calbindin$_{D_{28k}}$ and parvalbumin in the rat superior colliculus. Exp. Brain Res. (to be submitted)


In addition to the above scientific publications, several images produced in the CIF have been published in different places. Two images are being used by Noran Instruments in their literature on confocal microscopes, and two images have been selected for the LSUMC 1996 'Art of Healing Exhibition'.
304.2 ROLE OF SPONTANEOUS RETINAL ACTIVITY IN REGULATING Retinogeniculate CONNECTIONS DURING DEVELOPMENT. P.M. Cook*, G. Pruts* and A.S. Rizzo Dept. of Anatomy, Virginia Commonwealth U., Richmond, VA 23298 and Dept. of Physiology, Univ. of Louisville, Louisville, KY 40292.

The adult form and function of the lateral geniculate nucleus (LGN) arise after extensive modifications in circuit organization that include segregation of binocular inputs from the two eyes into eye-specific layers. Previous analyses have suggested that intracellular injection of tetrodotoxin (TTX) in ferret LGN prevents segregation (Shatz and Stryker, 1989), suggesting that spontaneous action potential activity contributes to remodeling of the retinogeniculate pathway. We have examined the activity-dependent mechanisms of eye-specific segregation using a technique that allows direct influence over the activity of spontaneous retinal input. Our method uses continuous intracellular application of TTX to block spontaneous discharge of retinal ganglion cells in ferrets that display retinogeniculate remodeling during the first 2 weeks of postnatal life. After a 2 week infusion, intracellular injection of biocytin revealed the projection pattern centralized and ipsilateral to the TTX injected eye. Eye-specific segregation of retinogeniculate projections was observed at postnatal day 14 in controls as well as in ferrets that received monoclonal (mAb) or biclonal (pAb) injection of TTX. However, segregation was absent in the monocularly injected animals. These results suggest that binocular competition modifies mammalian LGN in retinogeniculate projections. Additionally, they are consistent with the hypothesis that eye-specific activity may fine-tune segregation of binocular inputs into eye-specific layers. (Supported by NIH NS352193.)


We have been studying the development of the synaptic circuitry in the superior colliculus (SC) between ages P14 and P16. Using an in vitro brainstem preparation, we have introduced two different field potentials into the SC: an eye specific, modulating synaptic transmission (i.e. NMDA, and NOS). Field potentials or post-synaptic potentials were measured following stimulation of optic tract (OT) fibers. Whole cell recordings showed EPSPs with two different components: an early component was blocked by APV, suggesting that it was mediated by the NMDA component. NMDA antibody also abolished the synaptic field potential at P1. From P3 to P14, we observed a third IPSP following the EPSP. The EPSP could be blocked by bicuculline, an antagonist of GABA, and it was reduced at P1 by 60% and P12 by 80%.

Supported by DOD cooperative agreement DAMD 17-93-V-0103, NIH grant EY09273, and the LSUMC Neuroscience Center.


The optically visual input to Xenopus tectum comes into register with the contralateral map during development. The major transmitter for the contralateral (reinforcement) input is glutamate, and the major transmitter for ipsilateral input, released via the nucleus isthmi, is probably acetylcholine. The role of NDMA receptors in the activity-dependent process of organization of the ipsilateral map is demonstrated by the ability of NDMA receptors to block the horizontal spreading of the ipsilateral input in the tectum during the critical period, but little is known about the role of acetylcholine. Immunohistochemistry indicates that nicotinic receptors are located in the layers of the tectum that receive binocular inputs; unilateral eye enucleation indicates that most of these receptors are located on reinforcer axons. Receptor binding using stimulated cysite indicates that their is a possible action of nicotinic receptor in the nucleus isthmi. Receptors are located on cells and dendrites located appropriately to receive Binocular input. Calcium imaging using Fura-2 in tectal slices demonstrates no measurable response to nicotinic agonists alone, but shows significant synergism when nicotine or cyclics are applied with NDMA. In contrast, muscarinic agonists do not influence the activity of NDMA receptors. These results suggest that activity of nicotinic receptors can significantly modulate the effects of glutamate released from reinforcer axons.

Supported by U.S. PHS Grant EY-10690 to M.L. and S.B.U.
Calbindin\textsubscript{D28k} and Parvalbumin in the Rat Superior Colliculus.

R. John Cork, Sayed Z. Baber and R. Ranney Mize

Department of Anatomy, and Neuroscience Center of Excellence, LSU Medical Center, New Orleans, Louisiana LA 70112.

ABSTRACT
We have mapped the distributions of two calcium binding proteins (CaBP), calbindin\textsubscript{D28k} (CB) and parvalbumin (PV) in the rat superior colliculus (SC). The dominant feature of the distributions was a band of strongly labelled medium-sized CB cells centered on the optic layer (ol). Parvalbumin cells were found predominately in the intermediate gray layer (igl) where they were clustered in patches of PV labelled fibers. The superficial gray layer (srl) could be divided into two sublaminae based upon CaBP expression. CB-labeled cells were found mostly in the dorsal half of the srl, many of them being vertically oriented small bipolar cells. PV-labeled cells were generally found in the ventral part of the srl and dorsal ol. Some of the srl PV cells had morphologies similar to those of the small vertical CB-cells, but there were also horizontally oriented cells. Overall, the CaBP cells are distributed in complementary tiers that are not restricted to the traditional laminae. This suggests that there are functional sublaminae in the colliculus.

INTRODUCTION
The calcium binding proteins (CaBP), calbindin\textsubscript{D28k} (CB) and parvalbumin (PV) have been used as anatomical markers for functional groups of neurons (Celio, 1990). They are often expressed in discrete populations of neurons with complementary distributions, and in some parts of the brain they are found in subpopulations of GABAergic neurons (e.g. Celio, 1986; Hendry, et al, 1989; Miettinen, et al., 1996).

The physiological relevance of the two CaBP is unclear. The superfamily of CaBP comprises many molecules with one or more "EF-hand" calcium binding sites. Those members of the family whose function is known (e.g. calmodulin) are regulatory proteins that undergo conformation changes upon calcium binding and then affect the activity of other proteins. Neither CB nor PV have any known regulatory function. It has been suggested that they act as calcium buffers, dampening cytoplasmic calcium transients, and possibly protecting neurons against any toxic effects mediated by elevated calcium levels. This possibility is not, however, supported by some of the biochemical studies of CB structure. CB appears to have 6 calcium-binding sites of which maybe 4 are active (Ref). The binding constants for these sites vary but the apparent calcium binding constant for CB is about 10\textsuperscript{-8} M (Ref) suggesting that all the calcium binding sites would be fully occupied at physiological levels, and that transient changes in calcium would not have any effect on the calcium binding status of CB. If the other possibility, that these CaBP have some regulatory role, is true then by analogy with calmodulin it might be expected that CB would undergo a conformation change upon calcium binding. Leathers & Norman (1993) have found that CB does indeed undergo calcium-dependent conformational changes that affect its binding to several membrane bound proteins, however the details of what it binds to, and what effects it might have, are not known. Although there is this uncertainty about the \textit{in vivo} functions for these proteins, it is clear that excess CB or PV introduced into cells can buffer calcium transients induced by
depolarization (Chard et al., 1993). Whether they are simply passive calcium buffers, or whether they might regulate the activity of specific proteins, depending on [Ca$^{2+}$], remains to be determined.

Attempts to identify physiological roles for these CaBP have concentrated mainly on correlating CaBP content with cell location, morphological characteristics, or different physiological properties of the cells. As it has been suggested that a major function of these CaBP is to protect neurons from the damaging effects of excess calcium influx, whether due to repeated stimulation, ischemia, or neuropathological diseases, the presence of CaBP has often been correlated with neuronal survival after different insults that cause elevation of cytoplasmic calcium levels. CB is found specifically in two types of rat hippocampal neurons, dentate granule cells and CA1 pyramidal neurons, that are relatively resistant to excitotoxic damage. CB expression in the dentate gyrus granule cells is increased in response to repeated stimulation, suggesting that CB expression is regulated by [Ca$^{2+}$], (Lowenstein et al., 1991), and Iacopino & Cristakos (1990) reported that CB expression in these cells could be upregulated by corticosterone although this effect might also be due to an increase in [Ca$^{2+}$], rather than direct gene regulation. The situation with protection against ischemic damage is less clear. The dentate gyrus granule cells are vulnerable to ischemic damage until CB-IR has developed in the cells (Goodman et al. 1993) but vulnerability to ischemic damage in different neuronal populations is not well correlated with CB-IR. The CA1 pyramidal neurons are the most susceptible to damage while the CA3 pyramidal neurons that do not contain CB are less vulnerable (Goodman et al., 1993). CA1 interneurons which contain CB or PV are mostly protected against the damaging effects of ischemia (Johansen et al., 1990).

While these correlational anatomical studies have not been very successful in elucidating functions for CaBP, several studies have begun to look for physiological correlates of CaBP expression. Because of the co-expression of PV and GABA in some cells it has been proposed that PV cells should be highly active neurons with high firing rates (Celio, 1986). When Kawaguchi & Kubota (1993) correlated the morphology and electrophysiology of GABAergic nonpyramidal cells in the rat neocortex with their CaBP content, they did find that PV cells were fast spiking, while CB cells had low threshold calcium spikes. However, Li et al. (1995) reported that endogenous CB controlled the firing mode of neurons in the rat supraoptic nucleus by affecting calcium dependent depolarizing after-potentials, and excess CB shifted neurons from a phasic firing pattern to a fast continuous firing mode.

The superior colliculus (SC) should prove to be a productive structure in which to study the possible functions of CaBP. The mammalian superior colliculus is structurally multilayered and, at least in the intermediate layers, is compartmentalized into functional units. This compartmentalization is reflected in both the distributions of various biochemical constituents of the collicular cells, as well as specific arrangements of afferent fibers. The SC contains multiple populations of cells that can be grouped together on the basis of neuronal class, morphology, location, biochemical content, synaptic contacts or projection site. One of the biochemical markers of these compartments is PV (Illing, 1992), and CB is known to be expressed in specific subpopulations of the cells in the SC (Lane et al., 1993). In his major survey of CB and PV in the rat brain, Celio (1990) stated that the distributions of CB and PV are similar “in the upper two layers and complementary in the lower four layers.” In the cat SC the distribution of PV is complementary to that of CB neurons, with two tiers of PV neurons lying between three tiers of CB cells (Mize et al. 1992). Thus establishing correlations between their CaBP content and the morphology and physiology of the cells in the different SC subpopulations should help to establish what the functions of CB and PV are.
Although several investigators have looked at different aspects of PV and/or CB in the rat SC, there have been no detailed studies comparing the distributions of these two CaBP in the rat. Celio's (1990) survey gave only a brief description of their distributions. Illing et al. (1990) described the distribution of PV in rat SC, and Illing (1992) reported that PV expression correlated with the compartmental architecture of the intermediate SC layers. Schmidt-Kastner et al. (1992) made a cursory examination of PV and CB staining patterns in the rat SC during their study of the effects of SC deafferentation. Lane et al. (1993) described a population of calbindin-immunoreactive (CB-IR) cells in the optic layer of the rat SC.

Some reports have described CaBP-distributions in the colliculi of other species. Mize et al. (1991) described the distribution and morphology of CB cells in the cat SC. They found three laminar tiers of CB cells. Most of the CB cells were interneurons but only a very small percentage of them contained GABA. Behan et al. (1992) studied the localization of CB in the hamster SC and they also concluded that CB was not co-localized with GABA in the SC. In 1992, Mize et al. reported that CB and PV are expressed in complementary sublaminar patterns in the cat SC. In this report we extend these observations to the rat SC with the intent of developing the rat SC as a model system in which to investigate the physiology of the CaBP-containing cells.

MATERIALS AND METHODS

Male Sprague Dawley rats (250-300g; n=6) were anesthetized with Halothane or Ketamine/Xylocaine and perfused intracardially with ~400 ml of 4% paraformaldehyde in phosphate buffered saline (PBS) (pH 7.4). The brain was removed and the midbrain was blocked coronally and mounted on the stage of a vibratome with cyanacrylate glue. Serial sections, 50 μm thick, were cut through the SC.

CaBP Immunocytochemistry

In most experiments, alternate sections were labeled for either CB or PV. Sections were incubated in 1% NaBH₄ for 30 min. followed by 4% Normal Horse Serum (NHS) for 30 min. Sections were then labeled with primary antibody (1:400 Mouse anti-CB (Sigma) or 1:500 Mouse anti-PV (Sigma)) for 8-16 hrs. After another 30 min incubation in 4% NHS, sections were treated with a biotinylated secondary antibody, (1:200 Horse anti-Mouse (Sigma)) for 30 min. Antibody labelling was visualized using Vectastain Avidin-Biotin Complex (ABC) or ABC Elite (Vector Laboratories, Burlingame, CA). After a 1 hr incubation sections were stained by treatment with Diaminobenzidine hydrochloride (0.005%) and hydrogen peroxide (0.003%) for 1-3 min. Sections were given multiple rinses with PBS between each step. Antibody dilutions were made in 1% NHS in PBS.

In experiments to determine if cells co-expressed CB and PV, sections were double labelled with Mouse anti-CB (Sigma) and Rabbit anti-PV (Swant) primary antibodies. Normal Goat serum was substituted for the NHS used in single antigen labeling experiments. Instead of ABC/DAB staining sections were incubated in a mixture of fluorescently tagged secondary antibodies; 1:150 FITC Goat anti-Rabbit, or BODIPY-FL Goat anti-Rabbit, and 1:150 R-Phycocerythrin (PE) Goat anti-Mouse (Molecular Probes). Images were collected with the Noran Odyssey laser scanning microscope using the 488 nm laser line for excitation. Separate images were collected simultaneously at 510 - 545 nm for Bodipy/FITC and 555-625nm for PE.

NTS Plotting
Cell locations were mapped using a Eutectic Neuron Tracing System, marking the positions of all labeled cell bodies in a dorsal quadrant of the SC sections, and outlining the surface of the midbrain. The NTS software calculated the depth of each cell as the shortest (airline) distance between the cell and the dorsal edge of the colliculus. Data was collected from 9 representative sections; three sections (caudal, central, and rostral SC) from 3 different rats.

Cell Body Morphologies and Locations

Drawings of CaBP labeled cell bodies were made using a Zeiss microscope with a drawing tube. Cell locations were determined by making drawings at 4x, cell bodies and proximal dendrites were drawn at 60x or 100x. The montage of cell bodies (Fig. X) was constructed by placing the center of each cell soma at the appropriate distance from the dorsal surface of the SC. Cells were oriented in the montage so that the angle between the proximal dendrites and the closest part of the dorsal surface was correct.

Cell Densities and Soma Areas.

Numerical densities (Number of cells per unit volume) were determined using the method described in Albers et al. (1988). As we cut 50 μm thick sections, and the equations used by Albers et al. (1988) are based on assumptions that are most reliable for infinitely thin sections, we reduced the effective section thickness by determining the number of nuclei per unit area in a single focal plane (approx 2.5 μm thick).

The average area of cell bodies were obtained using a Joyce-Loebl Magiscan image analyzer. Representative sections were scanned for complete cell body profiles lying within selected layers of cells. Complete profiles were defined as those with a well defined nucleus and a clearly visible nucleolus. Only cell profiles containing a nucleus and nucleous were included in this analysis. Acceptable profiles were outlined on the video monitor with a light pen, and the computer calculated areas. Statistical analysis of the area data was all done with the Magiscan system.

3D Reconstruction.

The three dimensional distribution of the CB-containing cells in the optic layer was examined by reconstructing the immunocytochemically labeled sections. Images of all the sections from 1 rat were collected with a JVC high resolution video camera. Outlines of the midbrain surface, central aqueduct, and the outer limits of the CB labeling were digitized from these images with a digitizing tablet. Digital data was then transferred to a SGI workstation, running Skandha 3D-reconstruction software. After aligning the sections, a 3-D model of the colliculus and CB cell layer was constructed in the computer. This model could be displayed on a computer monitor. To examine the shape of the CB-cell layer, the reconstruction could be rotated about any of 3 orthogonal axes and the opacity of the outer surface could be reduced.

RESULTS

Distribution of labeled cells in the SC.

The most distinctive feature in all the CaBP labeled sections was a band of CB-immunoreactivity (CB-IR) in the ol and dorsal iql. (Figure 1. a. low power CB-DAB and b. low power Fluorescence) containing densely labeled, medium-sized cell-bodies in a meshwork of labeled fibers. Smaller lightly labeled neurons were distributed throughout the sgl (Fig 1c) and in
the deeper layers there were the occasional larger cell bodies (Fig 1d). [Description of neuropil labeling]

The pattern of PV staining was rather featureless. There was light uniform staining of the sgl neuropil, (Fig. 1e) and medium to large labeled cells scattered throughout the deeper collicular layers. Some lightly stained, small to medium sized cells were seen in the superficial layers. In most sections there were so few PV cells that a pattern of labelling was not discernable, however in some sections through the caudal half of the colliculus, two features of the PV distribution were evident. There was a thin band of medium sized cells in the ventral sgl just dorsal to the ol, and a thicker band of cells and labelled fibers in the igl. In the igl the pattern of labelling was patchy with cell somas and labelled fibers occurring in clusters. (Fig 1f)

The layer of heavily labeled CB cells was so obvious in the sections that it could be mapped into a 3-D reconstruction of the SC (Fig. 2). It was present throughout the SC with a relatively uniform mediolateral width along the anteroposterior axis. The layers are thicker along their medial edges giving the overall impression of two wings. (Figure 2. 3-D image of CB layer). As the PV cellular labeling was less clearly defined, it was not so obvious where the edges of any PV lamellae were. Nevertheless, a 3-D reconstruction suggested that there was a band of PV cells that lay ventral to the CB “wings”, and that this band was thicker laterally (data not shown).

When the locations of all the CaBP-IR cells in sections through the SC were plotted most CB cells were distributed in two bands in the superficial layers (Fig 3). In addition to the band of cells in the ol there was a more diffuse band of cell bodies scattered throughout the sgl. This upper band of cell bodies was concentrated in the dorsal two thirds of the sgl. In some sections there was also a thin line of CB cells in the zonal layer (z1). CB cells were scattered throughout the deeper SC layers without any obvious pattern. Between the two superficial CB layers there was a clear gap that was mostly empty of cells. Although it was not confirmed by any quantitative analysis, cells in the ol sometimes seemed to form clusters giving the band of cells a “patchy” appearance.

There were fewer PV-IR cells than CB-IR ones in the SC (Fig 4). The plots of PV cell locations showed cells scattered throughout the intermediate and deep layers with some cells in the superficial layers. There was little evidence of any laminar pattern of PV distribution except in caudal sections, where the cells in the ventral sgl lay in an obvious band (Fig. 4).

Depths and thickness’ of the cell layers

As the laminae of the colliculus have often been defined in terms of their depth below the surface of the tectum we collected data on the depths of Cb and PV cell bodies. Data was collected from several sections from 3 different rats. The sections chosen were of similar shape and size, and were taken from similar locations along the rostral-caudal axis of the midbrain e.g. the central region of the SC. The NTS software calculated the depth of each cell from the data files of cell distributions. Depth was defined as the shortest distance between the cell body and the dorsal surface of the colliculus. In total, depths were calculated for 3947 CB-cells from 7 sections, and 1200 PV-cells from 6 sections.

The peaks in the cell-depth histograms illustrate the relative densities and thickness’ of the various tiers of cells. Three peaks are seen in the CB cell distribution (Fig 5A). Most CB cells are in the sgl between 100 - 300 µm from the dorsal surface. The heavily stained cells in the ol lie in a band between 300 - 600 µm. There is then a dispersed population of cells in the dgl at depths ranging from 1200 - 2000 µm. The peaks in the PV histogram (Fig 5B) are less distinct, but there is a clear peak centered at 300 µm and a broad peak between 700 - 1200 µm.
Superimposing the two depth distributions (Fig 5C) shows that the patterns of CaBP expression are complementary, with the PV tiers centered between the CB tiers. The approximate boundary depths of the superficial SC layers are shown for reference. The depths of these particular layers were compiled as averages from several studies of the rat SC laminae (e.g. Albers et al., 1988, Lane et al. 1993). It should be noted that the deep SC laminae, below the ol, are not uniform in thickness mediolaterally, nor are they parallel to the dorsal surface of the SC as are the superficial layers. The combination of these two factors means that there is only an approximate correspondence between depth and a particular cell layer, and the distributions of cell depths are broader (e.g. Fig. 5C) than the thickness of the layers they represent. There is always some overlap of the depth values of cells in a particular deep layer with those for adjacent layers.

If the depth distributions are plotted as percentages of the total number of CaBP-containing cells, instead of as percentages of CB- or PV-cells, the relative densities of the two CaBP in the different layers are revealed (Fig 5D). CB cells predominate in most regions of the colliculus, even at some of the depths where the PV laminae are. The only SC layer where there are more PV cells than CB ones is in the iGl between 800 μm and 1200μm.

Cell size and morphology in the different layers

Having determined that there are alternating bands of CaBP immunoreactivity in the SC, we wanted to determine if the populations of cells in particular tiers had any distinguishing morphological characteristics. We determined the average size of the cell body profiles in the individual tiers (Table 1).

<table>
<thead>
<tr>
<th>Tier</th>
<th>Mean Area of Cell body (μm²)</th>
<th>s.d. (μm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calbindin: sgl</td>
<td>84</td>
<td>21.5</td>
</tr>
<tr>
<td>Parvalbumin: sgl/ol</td>
<td>151</td>
<td>31.2</td>
</tr>
<tr>
<td>Calbindin: ol/igl</td>
<td>184</td>
<td>53.1</td>
</tr>
<tr>
<td>Parvalbumin: igl/dgl</td>
<td>211</td>
<td>95.1</td>
</tr>
</tbody>
</table>

Table 1. Mean area of cell bodies in specified tiers. Data was collected using the Magiscan Image Analyzer, scanning along specific cell tiers.

The results of the cell area measurements show that CaBP content is not correlated with cell size, as neither CB nor PV are confined to cells in a particular size range. However, cell size is correlated with depth in the SC. There is a steady increase in cell size from the small cells near the dorsal surface to the much larger cells in the deep layers. Instead of being correlated with cell size CaBP content seems rather to be a function of location in the SC. Thus a cell's location in the ol is a better predictor of its CaBP content than its size. It is nonetheless true that cells in particular tiers fall within characteristic size ranges. Thus the CB cells in the sgl and ol/igl belong to statistically different populations of cells.

The determination of which cell morphologies were represented in the different CaBP layers was done by drawing well labelled cells, and proximal dendrites, and compiling a montage of the cells at their appropriate depths (Fig 6). We concentrated on the morphologies of the cells in the superficial collicular layers as these have been described in detail (e.g. Langer & Lund, 1974;
Labriola & Laemle, 1977; Wharton & Jones, 1985) and little is known about the cell types in the deeper layers. The CaBP-labeled material contained examples of all the major cell types described in Golgi stained sections (Langer & Lund, 1974; Labriola & Laemle, 1977). Marginal cells were found in both the CB and PV-labelled sections, as were vertical fusiform cells and horizontal cells. Two of the cell types appeared to be confined to one or other CaBP. Most, if not all, of the CaBP-containing wide-field vertical cells were CB-cells, and all the stellate cells we observed contained PV. However, as the sample of cell drawings is rather small, we cannot say definitely that either CaBP is found exclusively in a particular cell type.

Double label results

Fluorescently labeling both CB and PV in the same sections enabled us to determine if any cells expressed both CaBP. Low magnifications views of the SC, confirmed the complementary patterns of CaBP distribution (Fig 7A). At higher magnifications, confocal microscopy of the double labeled sections revealed that almost all CaBP-cells contained only PV or CB, but not both proteins. A few cells were seen that might have contained both proteins, but we could not rule out the possibility that these examples were due to overlap of two cells in different focal planes, or that they were due to “bleed-through” of signal from one fluorescent label into the channel for the other label’s signal.

DISCUSSION

Distribution of CB and PV in the superior colliculus

Our results show that PV and CB are distributed in alternating tiers of cells that form sublaminae of the traditional divisions of the SC. There are three layers of CB cells with two PV layers interspersed between them. The upper band of CB cells occupies the dorsal two-thirds of the sgl and is mainly composed of vertically-oriented lightly labeled cells. Most of the cells are bipolar with small fusiform cell bodies (cell sizes). There are also vertically oriented pyriform cells and horizontal bipolar cells. The middle band of CB cells overlays the optic layer and consists of heavily labeled medium-sized cells in a mesh work of labelled fibers. Most of the cells appear to be wide field vertical cells with laterally oriented proximal dendrites projecting up into the superficial layers.

Two distinct populations of CB-cells have previously described in the upper layers of the SC. Schmidt-Kastner et al. (1992) briefly described, a layer of medium-sized CB cells with predominately vertically oriented dendrites in the ventral sgl, and a plexus of strongly stained neurons just ventral to the ol. Although the locations of these CB cells are ventral to those we assigned the cells, we assume that these two populations correspond to the upper two CB tiers described in this paper. Lane et al. (1993) presented a more detailed picture of CB distribution in the superficial layers of the rat SC. They found an upper band of small fusiform cells in the dorsal half of the sgl, and a heavily labeled band of medium-sized cells in the ol. Cells in the latter band were mostly wide-field vertical or medium-sized stellate cells. Using retrograde labeling, they also determined that the majority, if not all, of the CB-cells in this band projected to the lateral posterior nucleus (LP) of the thalamus. The majority (80%) of the LP projecting neurons contained CB and about 15% contained both CB and adenosine deaminase.

Between the two CB layers lies a band of PV cells. Its dorsal edge overlaps with the superficial CB tier and contains a similar mix of cell types plus some stellate cells. Cells in the ventral part of this PV layer lie in a narrow band where there are few CB cells. This band is more
distinct in the caudal colliculus and many of the cell somas appear to be medium sized horizontal cells. Schmidt-Kastner et al. (1992) reported seeing PV neurons scattered throughout the sgl but make no mention of their morphology or sublaminar distribution. The distribution of PV in the rat SC was more fully described by Illing et al. (1990). They observed two populations of PV neurons. One, consisting of vertically oriented cells of similar morphology, was located in the ventral half of the sgl, the other was composed of many neurons of differing size and morphology widely distributed throughout the igl and deeper layers. They also observed patches of PV-IR in the igl, and found that a subpopulation of small neurons were clustered in the patches (Illing et al., 1990).

Ventral to the optic layer there are a few scattered CB cells in the igl and greater numbers of scattered CB cells in the deeper layers. The igl is the only layer where PV cells outnumber CB cells, most of the cells have medium to large somas. There is some evidence of clustering of neurons in the igl.

Comparison with other species

As far as we are aware the only other species in which the distributions of CB and PV have been compared, is the cat (Mize et al. 1992). In the cat, the CaBP are distributed in a pattern of alternating sublaminar tiers, very similar to that seen in the rat. The uppermost tier consisted of small CB-IR cells of varying morphologies. Three cell types were identified in this layer, horizontal fusiform cells, pyriform neurons with vertically oriented dendrites, and stellate cells (Mize et al., 1991). Retrograde filling studies showed that some of the pyriform and stellate cells cells projected to the lateral geniculate nucleus but the horizontal cells did not.

Ventral to the upper CB tier was a dense band of PV labelled cells (Mize et al., 1992) occupying the ventral sgl and the upper ol. Most of these cells were small round neurons, stellate neurons, or vertical bipolar neurons with fusiform cell bodies. They constituted more than 80% of the neurons retrogradely labelled by an injection into the lateral posterior nucleus (LP), although less than 50% of the PV-cells in this tier projected to the LP.

The middle CB tier straddled the border between the ol and the igl. The major cell type in this band had a stellate cell body and, in many cases, varicose dendrites that projected into the upper ol or deep SC layers but not into the sgl (Mize et al. 1991). The vast majority of the CB cells in this middle tier appeared to be interneurons that did not contain GABA.

Below the ol the CaBP tiers were less obvious with lower cell densities. PV cells predominated in the igl and below them was a region with mostly CB-cells. The PV-neurons were mostly medium-sized stellate cells or large multipolar neurons. Many of these PV cells projected to the dorsal lateral pontine gray nucleus or the predorsal bundle (Mize et al. 1992). CB-cells in the deeper regions of the cat SC were also scattered throughout the deep gray and deep white laminae. They varied considerably in size and morphology, but mostly appeared to be various types of stellate neuron. Some of the largest CB-cells were found in the deepest parts of the colliculus bordering the dorsal edge of the periaqueductal gray (Mize et al., 1991). These CB cells also seemed to be mostly interneurons.

Possible functional consequences of CaBP distribution

Although the observed patterns of immunohistochemical staining were fairly consistent between animals, it is possible that they could have been influenced by the functional state of the colliculus. Winsky and Kuznicki (1996) reported that the binding of CaBP antibodies to their antigens varied depending on whether the CaBP were in a calcium-bound or -unbound state. Given
that calcium levels in neurons can fluctuate greatly some of the patterns we have seen may reflect variations of calcium levels rather than CaBP.

The functions of these CaBP remain unknown. CB and PV do not seem to be selective for any one cell type, yet their pattern of expression appears to be related to the functional organization of the colliculus. There appear to be several functional domains in the colliculus and it seems that different groups of cells have different requirements for CaBP.

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CALBINDIN and PARVALBUMIN ARE EXPRESSED IN COMPLEMENTARY PATTERNS IN THE RAT SUPERIOR COLICULUS.
B.J. Cock* and R.C. Bader and R.R. Migeon, Department of Anatomy and Neurosciences Center, LSU Medical Center, New Orleans, LA 70112.
We have analyzed the distribution of two calcium-binding proteins, calbindin (CB) and parvalbumin (PV), in the rat superior colliculus (SC) using immunohistochemistry and other techniques. The distribution of CB and PV in the SC was determined by immunohistochemistry. The distribution of CB and PV in the SC was determined by immunohistochemistry. The distribution of CB and PV in the SC was determined by immunohistochemistry. The distribution of CB and PV in the SC was determined by immunohistochemistry. The distribution of CB and PV in the SC was determined by immunohistochemistry.

265.16
EFFECT OF NEONATAL EYE REMOVAL ON EXPRESSION OF mRNA IN THE SUPERIOR COLICULUS AND RELATED STRATA.
A.K. Harvey* and S.J. Stumpf and L. Chang and D.J. Stumpfman, The Dept of Anatomy and Neurobiology, University of Rochester, University of Rochester, University of Rochester, University of Rochester.
This study utilized the technique of in situ hybridization histochemistry to study the expression of mRNA for the gap junction protein GABA (GAD) and the neurotransmitter GABA (GABA). The expression of GABA mRNA was detected in the SC, and in the thalamus, the expression of GABA mRNA was detected in the SC, and in the thalamus. The expression of GABA mRNA was detected in the SC, and in the thalamus.

265.17
NEUROACTIVE SUBSTANCES IN THE PARABASALICAL NUCLEUS OF THE THIRTEEN-FOOTED RODENT GROUND SQUIRREL, R. J. Garcia and M. R. Lang* Dept. Anatomy and Inst. of Neurobiology, Univ. of Puerto Rico, Mirc, San Juan, PR 00920.
The parabasalical nucleus (PB) has extensive reciprocal connections with the superior colliculus (SC) in the thirteen-footed rodent ground squirrel (Spermophilus niger). We have employed various immunohistochemical methods to analyze the distribution of these neurotransmitters and neuropeptides. The distribution of these neurotransmitters and neuropeptides is shown in the figure. The distribution of these neurotransmitters and neuropeptides is shown in the figure. The distribution of these neurotransmitters and neuropeptides is shown in the figure.

265.18
Projections to the basal pontine gray were studied after injections of d-leucine restricted to one of the following retrolaminar cortical areas: 17, OA, caudal OA, caudal OB, 10a, and 7. All corticotopical projections were divergent, as they terminate at the pontine gray and at the pericallosal gray. The projection to the basal pontine gray was studied after injections of d-leucine restricted to one of the following retrolaminar cortical areas: 17, OA, caudal OA, caudal OB, 10a, and 7. All corticotopical projections were divergent, as they terminate at the pontine gray and at the pericallosal gray.
RECEPTOR RESPONSE PROPERTIES IN THE SUPERIOR COLLICULUS OF THE NEWBORN RHESUS MONKEY
J.T. Wallace, S. E. Stein, and S. E. Stein

The superior colliculus (SC) is a component of the ascending auditory system and plays a role in the generation of saccadic eye movements. It is also involved in the processing of visual information. The SC is divided into three major functional areas: the dorsal lateral geniculate nucleus (DLC), the ventral tegmental area (VTA), and the substantia nigra (SN). The SC receives input from the thalamus, the lateral geniculate nucleus (LGN), and the visual cortex. The SC is an important brainstem structure that plays a key role in the control of eye movements, saccades, and visually guided behavior. The SC is also involved in the processing of visual information, including the detection of visual targets, the generation of saccadic eye movements, and the suppression of visually directed saccades. The SC is an important structure for the study of the neural mechanisms underlying the control of eye movements and visually guided behavior.
Pre- and postnatal expression of amino acid neurotransmitters, calcium binding proteins, and nitric oxide synthase in the developing superior colliculus

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Key words: GABA, glutamate, nitric oxide, calbindin, parvalbumin, synapse formation, axon pathfinding

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Introduction

The mammalian superior colliculus (SC) is an excellent model for developmental studies because it is multi-laminated, has well-defined and segregated afferent inputs, and contains specific output neurons which project to both ascending and descending targets in the brain. The receptive field properties of its neurons have been studied extensively and it has a relatively refined topographic map of the visual world that is conveyed by inputs from both the retina and the visual cortex. Finally, the specific neurotransmitters of many of its intrinsic and projection neurons have been identified and some of its synaptic circuits are well-established (see Huerta and Harting, 1984; Chalupa, 1984; Mize, 1992, 1996, for reviews).

Much less is known about how these features develop. Retinotectal pathway development has been studied in great detail, particularly in lower vertebrates (see Constantine-Paton et al., 1988; Udin and Fawcett, 1988, for reviews), and more recently in rats (Simon and O'Leary, 1992; Simon et al., 1992). Studies in rat and other rodents demonstrate that there are at least two distinct and well-defined periods of growth in this structure. During early stages of development, neurons divide, migrate to specific laminae, begin to establish dendritic branching patterns, and send their axons to specific targets (Mustari et al., 1979; Altman and Bayer, 1981; Cooper and Rakic, 1981; Edwards et al., 1986a). Extrinsic afferents also grow into SC during this period and at least those from the retina establish a rough topographic order (Edwards et al., 1986b; Udin and Fawcett, 1988; Sanes, 1993), which has been referred to as the 'prepattern' (Simon and O'Leary, 1992). Most of these events are 'activity independent' in the sense that visually driven or patterned electrical activity is presumed to be absent or limited during this prenatal stage of development.
In later stages of development, this 'prepattern' of cellular lamination, afferent fiber innervation, and topographic order is further developed and refined. Permanent synapses are formed, stabilized, and strengthened while others are eliminated by cell death and fiber retraction (Lund and Lund, 1972; Mize and Sterling, 1977; Finlay et al., 1982; Cooper and Rakic, 1983; Simon et al., 1992). The physiological receptive field properties of individual cells are first recorded and stimulus induced electrical responses mature (Stein, 1984 for review). Many of these later events are activity or experience dependent and can be modified by alterations of the visual environment or changes in the patterning of extrinsic afferents (Shatz, 1990; Goodman and Shatz, 1993). Although many of the factors underlying these developmental events have been studied, remarkably little is known about the onset of expression of neurotransmitters and other neurochemicals that occurs during these two stages of development in the mammalian SC.

As a consequence, we have examined the onset of expression of three types of neurochemical in the cat superior colliculus: the amino acid neurotransmitters GABA and glutamate; the calcium binding proteins calbindin and parvalbumin; and the synthetic enzyme of nitric oxide, nitric oxide synthase (NOS). In some cases, we have compared the time course of expression with the development of cellular lamination and the growth of afferents into the SC. Our results show that some neurochemicals, especially GABA and calbindin, are expressed in the very earliest stages of development as neurons are migrating from the subventricular zone to the tectal plate. Others, particularly parvalbumin and glutamate, are expressed much later in development, during the period of synaptogenesis and fiber refinement. Finally, we show that nitric oxide synthase, which has recently been identified as a potential retrograde messenger during synapse stabilization (Gally et al., 1990), is expressed transiently at a time which correlates with the refinement of the axonal connections that are
formed with neurons that express it.

**Development of the amino acid neurotransmitters in cat SC**

*Organization:* Gamma aminobutyric acid (GABA) is a key inhibitory neurotransmitter in the mammalian SC (Okada, 1992). Immunocytochemical studies have shown that GABA containing neurons account for up to 40% of all cells in SC. Most of these are small to moderate in size and the vast majority are interneurons (Mize, 1992 for review). They are most densely distributed within the zonal, superficial gray, optic, and intermediate gray layers, but they are also found in fewer numbers in the deepest layers of SC (Mize, 1988). Electron microscopy studies show that GABA immunoreactive processes contain primarily pleomorphic or flattened synaptic vesicles and form microcircuits that likely mediate several different types of inhibition (Mize et al., 1982; Mize, 1988; 1992; 1994). Developmental studies in vitro suggest that GABA is first effective at generating inhibitory post-synaptic potentials only in postnatal tissue (Warton et al., 1990; Kraszewski and Grantyn, 1992) and that GABA containing synapses increase dramatically in number for 2-3 weeks after birth (Warton et al., 1990). Thus, GABA mediated inhibition develops relatively late in the mammalian SC.

*Development:* Despite the late postnatal onset of GABAergic inhibition, GABA antibodies label cells within this structure as early as the end of the first trimester before birth. We examined the expression of GABA in fetal tissue ranging in age from E24-E59 and at postnatal ages P1 to P60. GABA content was localized using standard immunocytochemistry techniques. The distribution of antibody labeled neurons was plotted with a computer-based microscope tracing system and cell size measured with an image analyzer.
By E24, the earliest age examined, GABA immunoreactivity is found in both the neuropil and in some cell bodies throughout the tectal plate. Cell bodies and processes are also labeled within the subventricular zone (SVZ) at this age. By E30-36, a distinct bi-laminar distribution of GABA immunoreactivity is present. Both cell bodies and processes are intensely labeled within the superficial layers at E30 (Figure 1A), and an obvious dense band of label in the upper one-half of the cat SC is visible (Figure 1B). Labeled cell bodies in this dense band are mostly small cells with few or no labeled processes. By contrast, GABA-ir in the deep layers is very sparse due both to the small number of labeled neurons and very limited neuropil labeling. Those cells that are labeled are mostly bipolar neurons with vertical fusiform cell bodies and vertically oriented dendrites (Figure 1C). This cell type is also seen in the SVZ (Figure 1A), where the dendrites of some cells extend into the deep layers of SC. This morphology is characteristic of neurons in the process of migration (see Edwards et al., 1986a). The pattern of labeling seen at E30-36 suggests that: 1) many GABA containing neurons have already reached the superficial layers by E30; 2) neurons destined for the superficial layers of SC express GABA while still in the SVZ; 3) GABA is expressed in SC neurons both during and after migration; 4) the deep bipolar neurons represent a transient cell morphology in the process of migration.

By E40-46, the pattern of anti-GABA labeling is nearly identical to that found in the adult, although many more cells are labeled than at later stages of development. Well-labeled neurons are now densely distributed throughout the superficial and intermediate gray layers of SC (Figure 2A). There are fewer labeled neurons in the deepest layers, but these also are intensely labeled by the antibody (Figure 2A). Labeled cell bodies in the superficial layers have characteristics similar to those in the adult, with small round or sometimes ovoid cell bodies and short dendritic processes (Figure 2B). Cell bodies in the intermediate layers are
also mostly small round cells (Figure 3C), as are most of those in the deepest layer. Only a few cells retain a bipolar morphology (Figure 3D), suggesting that few if any GABA containing cells are still migrating by E46. Neuropil labeling is also quite prominent at this age (Figures 2B-D).

Labeling at later ages produces a similar laminar pattern, but many fewer cells are labeled. We plotted the distribution and number of GABA-ir neurons from one animal in each of four age groups (E30, E46, P1, P60) to determine quantitatively the reduction in GABA labeled cell number. Computer counts show a 2.5 fold decrease in GABA-ir cells between E46 and P60, despite the significant increase in overall SC volume that occurs at later ages. Thus, the maximum number of GABA containing cells is reached around the end of the second trimester (E40-46) with significant cell death or loss of expression occurring by P60.

To date, we have examined glutamate immunoreactivity in the prenatal SC in only six prenatal age groups (E41, E46, E51, E53, E58). Our preliminary data suggests that intense glutamate immunoreactivity develops much later in the prenatal kitten than does GABA. At E41-46, only a few darkly labeled neurons are found within the optic and intermediate gray layers, regions that are known to contain many glu-ir cells in the adult. A larger number of intensely labeled neurons are found at E51-53, most within the deeper layers of SC. By E58, a number of large neurons and some medium sized cells within the deep superficial gray and optic layers are now well-labeled but their number is less than that seen in the adult. Further development in the number and density of glu labeled cells occurs postnatally.

Functional Considerations: GABA is also one of the earliest neurotransmitters to be expressed in other regions of the CNS. In rat visual cortex, for example, GABA immunoreactivity is
seen as early as E13 (Lauder et al., 1986; Van Eden et al., 1989). It appears first in neurons within the marginal and subplate zones and somewhat later in cell groups throughout the cortical plate (Luskin and Shatz, 1985). Very early expression of GABA-ir has also been found in the spinal cord, brainstem, and diencephalon (Lauder et al., 1986), so GABA must be important to developmental processes that occur very early, including mitosis, migration, and neurite outgrowth.

Studies that have specifically examined the relationship between GABA expression and cell migration have shown that GABA is not expressed until after cells have reached their final destinations, at least in visual cortex (Miller, 1986). On the other hand, many studies have shown that GABA is important in neurite outgrowth. GABA and its agonists increase the length, number, and branching of neurites in cultured cerebellar granule cells (Hansen et al., 1987), in cortical neurons (Spoerri, 1987), and in chick optic tectum cells (Michler-Stuke and Wolff, 1987). GABA also increases the density of the rough endoplasmic reticulum, suggesting that it promotes protein synthesis and incorporation of receptor protein into the plasma membrane (Hansen et al., 1987; Spoerri, 1987). GABA application in culture can also enhance low and high affinity GABA receptor binding (Meier et al., 1984; Redburn, 1992). Thus, one key function of GABA early in development may be to promote neurite extension, possibly by binding to GABA receptors.

Another early effect of intracellular GABA may be to alter ion concentrations and the membrane potential. Increases in potassium concentration in culture have been shown to stimulate neurite outgrowth in dorsal root ganglia (Chalazonitis and Fischbach, 1980) and high calcium concentrations in chick tectal cultures increase the number of neurites if GABA is also present in the medium (Michler-Stuke and Wolff, 1987). Similarly, intracellular calcium
mobilization promotes neurite outgrowth in vitro (Angliuster et al., 1982). Thus, the ionic conductance and membrane potential properties of neurons may be important to their growth well before cells are innervated by afferents or form synapses. However, GABA has also been shown to promote synaptogenesis in both retinal and cerebral neurons (Spoerri, 1987; Messersmith and Redburn, 1993). It is therefore not surprising that GABA in SC neurons continues to be expressed at postnatal ages when synapse formation is occurring (Lund and Lund, 1972; Sterling and Mize, 1977). In summary, GABA probably plays different roles at different stages of development and is likely involved both in neurite outgrowth and synaptogenesis.

Development of calcium binding proteins in cat SC

Organization: Calcium binding proteins (CaBPs) in the CNS regulate intracellular calcium by buffering cytoplasmic Ca++, by controlling Ca++ transport across the cell membrane, and by regulating Ca++ dependent second messenger systems (Heizmann and Hunziker, 1990; Celio, 1990; Braun, 1990; Baimbridge et al., 1992). Two of the most frequently studied CaBPs in the CNS are calbindin (CB) and parvalbumin (PV). CB is a 28kD calcium binding protein that is co-localized in some GABAergic interneurons (Baimbridge et al., 1992; Kawaguchi et al., 1987) and is often present in neurons that have low threshold calcium spikes (Kawaguchi, 1993; Lo et al., 1995). PV is a 12 kD calcium binding protein that is found in both projection and interneurons and has been associated with cells that are fast spiking (Kawaguchi et al., 1987; Kawaguchi and Kubota, 1993).

In the cat superior colliculus, these two calcium binding proteins are found in complementary sublaminar tiers and in different cell types (Mize et al., 1991; 1992; Mize and
Luo, 1992). CB neurons form three sublaminar tiers, one within the upper superficial gray layer (sgl), a second within the deep optic and dorsal intermediate gray layers (igl), and a third consisting of cells scattered within the deep gray layer (dgl) of SC (Mize et al., 1991; 1992; Mize and Luo, 1992). The vast majority of CB cells are small interneurons, some of which contain GABA (Mize et al., 1991). By contrast, PV labeled neurons in cat SC form a single dense band of medium to large sized cells that overlap the deep sgl and upper optic layers. PV cells are also distributed throughout the deeper layers of SC, but virtually no PV neurons are present within the upper sgl (Mize et al., 1991). The vast majority of PV neurons are projection cells that send their axons to a variety of extrinsic targets, including the lateral posterior nucleus and a number of descending brainstem sites (Mize et al., 1992).

**Development:** CB and PV have very different developmental histories in the kitten SC. We have examined the expression of CB and PV in fetal tissue ranging in age from E24-E59 and at postnatal ages P1 to P60. CB is expressed at the earliest prenatal age examined - E24. At this age, most CB neurons are located within the subventricular zone (SVZ), but others are sparsely scattered throughout the tectal plate. By E28, CB positive neurons are found throughout the dorsal-ventral extent of the tectal plate, but cells at different depths differ in morphology (Figure 3A). The most dorsal neurons are small cells with few or no dendritic processes (Figure 3A,B). Intermediate depth cells have multipolar morphologies with short dendrites (Figure 3B). Many of the deeper cells have bipolar morphologies with vertically oriented dendrites (Figure 3C), similar to those that contain GABA at this stage of development and which are believed to be in the process of migrating.

The three sublaminar tiers of CB cells are barely visible at E34, and many deeper cells retain a bipolar shape (Figure 4A). Labeled CB cells in the SVZ at E34 also have
morphologies typical of migrating neurons. By contrast, the three tiers of CB neurons are well segregated by E40-46, labeled bipolar cells are no longer present, and no CB labeling is found within the SVZ at these later ages (Figure 4B,C). As with GABA labeled neurons, the highest density of CB neurons is found at E40-46. Cell counts from tissue aged E40-P60 show that the density of CB cells at E40 is three times higher than that found at P60.

This reduction in cell density is due primarily to cell death in the prenatal kitten but is also due to loss of expression of CB after birth. This conclusion is based upon two lines of evidence. First, overall neuronal density as measured in thionin stained sections decreases progressively from E41 to P21. Thus, the overall loss of neurons is similar to that of CB containing cells, suggesting that CB cells die in proportion to the total population. Secondly, pyknotic neurons with eccentric nucleoli, typical of dying cells, are observed in the late prenatal period and are especially numerous between E51 and E59, a period in which the number of CB cells declines rapidly. Very few pyknotic profiles are observed after P7, which indicates that there is little cell death after birth. Because the density of CB cells continues to decrease until P60, the postnatal decrease in CB neurons must be due to a loss of expression.

PV expression has a dramatically different developmental time course. PV immunoreactivity is not observed prior to P1 and there is a continuous increase in the number of PV labeled cells until P35 with a slight decline thereafter. PV labeled neurons at P1 are few in number, but by P7 the dense band of PV neurons in the deep superficial gray and upper optic layers is already visible (Figure 5A). The number of PV cells increases in density by P21, and is fully developed by P35 (Figure 5B-C). PV cell morphology is like that of the adult by the time PV is first expressed. At all ages, PV neurons are medium to large sized cells, usually with stellate-like morphologies and multiple dendrites radiating from the soma.
(Figure 5A-C). In summary, PV is expressed late in development, well after cells have migrated and lamination is well-developed.

**Functional Considerations:** The roles that CB and PV play in brain development are largely unknown. CB expression occurs very early in prenatal development in many brain regions, often at a time when both cell generation and migration are still in process. In SC, CB is unlikely to be involved in mitosis because no CB cells are found within the ventricular zone where SC cells divide (Edwards et al., 1986a). On the other hand, CB may well be involved in cell migration. Bipolar neurons with morphologies typical of migrating cells are CB ir in the kitten SC until around E40, the time at which migration ends. We also found many CB neurons in the SVZ that had bipolar dendrites and crossed into the deep layers of the tectum. Thus, CB appears to be an important molecule for at least some neurons that are in the process of migration.

Its role may be to regulate intracellular calcium in migrating cells. Calcium levels are reportedly high in migrating neurons. Actively elongating growth cones have higher levels of intracellular calcium than do inactive cones (Kater et al., 1988), while excessive calcium can inhibit growth cone motility and neurite extension (Mattson and Kater, 1987). In addition, N-type calcium channels are first expressed in cerebellar granule cells as they begin to migrate and the calcium channel blocker omega conotoxin retards migration (Komuro and Rakic, 1992). Thus, regulation of intracellular calcium is essential to migrating neurons and CB could well serve to buffer or otherwise regulate calcium levels in these developing cells.

Although it is not clear from our studies whether CB is expressed in all migrating neurons, there is evidence in other brain regions that CB is transiently expressed in many cells that do
not contain it in the adult. In striatum, aspiny neurons in the dorsal caudate-putamen express CB intensely in the prenatal but not the adult rat (Liu and Graybiel, 1992). CB is also intensely expressed in layer 4 and 6 cortical neurons in the prenatal monkey that do not express this protein in the adult (Hendrickson et al., 1991). Thus, several lines of evidence suggest that many neurons express CB during migration while only a subset of these cells retain the protein once migration has ceased.

PV must play a very different role in developing neurons and possibly the same role which it plays in the adult. PV immunoreactivity does not occur in cat SC until after birth and is maximal between P7 and P35, a period during which stimulation evoked neuronal activity is first recorded (Stein, 1984; Kao et al., 1993). Most synaptogenesis also occurs during this postnatal period (Mize and Sterling, 1977). For these reasons, PV is more likely to be involved in regulating intracellular calcium influx produced by receptor and electrical activity. Consistent with this, PV has been localized in fast spiking neurons in both the hippocampus (Kawaguchi et al., 1987) and cortex (Kawaguchi, 1993) and PV may be essential in regulating calcium influx that occurs when cells are firing at very high frequencies.

Cells containing PV may also be specifically associated with the NMDA receptor (Andressen et al., 1993; Celio, 1990). Fast spiking cells are selectively activated by NMDA receptor agonists (Kawaguchi, 1993) and the development of NMDA receptors correlates roughly with the time of initial PV expression in visual cortex (Kleinschmidt et al., 1987; Bode-Greuel and Singer, 1989; Braun, 1990; Reynolds and Bear, 1991). Thus, peak NMDA receptor and PV function may be related.
Development of Nitric Oxide Synthase in the cat SC

O\textit{rganization:} Nitric oxide (NO) is a free radical gas that has been implicated as a retrograde messenger in both long term potentiation and synapse stabilization in the developing brain (Gally et al., 1990; Garthwaite, 1991). NO is diffusible and membrane permeant and does not require synaptic vesicle packaging for release. It is produced in postsynaptic cells by activation of excitatory amino acid receptors (NMDA) that increase intracellular Ca+++. This increase activates NO's synthetic enzyme, nitric oxide synthase (NOS), which leads to the production and release of NO (Bredt and Snyder, 1990; Garthwaite, 1991). According to the Gally hypothesis (Gally et al., 1990), post-synaptic release of NO activates guanylate cyclase-cGMP in presynaptic axons, thus enhancing neurotransmitter release and strengthening or stabilizing the synapses which contact NO containing cells (Bredt and Snyder, 1989; Gally et al., 1990). NO is a good candidate as a retrograde messenger both because of its diffusion characteristics and because the NMDA receptor is also known to be involved in synapse stabilization.

If NO is involved in the process of axon stabilization, then NOS should first be expressed at a time when synapses are being established and axons refined. To test this, we have examined the expression of NOS in the pre- and postnatal kitten SC using a histochemical reaction specific to nicotinamide adenine dinucleotide phosphate diaphorase (NADPHd), a byproduct of NO synthesis that is a marker of NOS containing neurons. These results have also been confirmed using antibodies directed against neuronal NOS.

NOS is found in very specific groups of neurons and fibers in the adult SC (Arceneaux et al., 1995). NADPH and bNOS immunoreactivity are found in cells within the dorsolateral
periaqueductal gray (pag), within cells and fibers in the deep gray layer, and within fibers and in a few neurons within the intermediate gray layer (igl). A few labeled neurons are also scattered in the superficial layers of SC. The igl labeling is of particular interest because it is primarily confined to a group of fibers and cells that we call the patch-cluster system (Figure 6)(Jeon and Mize, 1993). These fiber patches were first described by Graybiel (1978b, 1979) who showed that they contained acetylcholine. More recently, the patches have been shown to include afferent fibers from at least three sources (Figure 6): the pedunculopontine tegmental nucleus which contain acetylcholine (ACh) (Ililing and Graybiel, 1985; Beninato and Spencer, 1986; Hall et al., 1989; Ililing, 1990; Harting and Lieshout, 1991; Jeon and Mize, 1993); the substantia nigra which contain GABA (Graybiel, 1978a; Harting and Lieshout, 1991; Ficalora and Mize, 1989); and the frontal eye fields (Ililing and Graybiel, 1985) which probably contain glutamate (Dori et al., 1992).

These fiber patches precisely overlap clusters of neurons that project to the region of the cuneiform nucleus (CFR; Jeon and Mize, 1993). The cell clusters consist of closely apposed groups of 3-20 neurons. There are 4-7 clusters per section, mostly located in the dorsal igl of the caudal SC. The correspondence between the clustered cells and fiber patches has been shown directly by double labeling experiments in which the fibers were labeled by antibodies to choline acetyltransferase (ChAT) and the cells by retrograde transport of horseradish peroxidase injected into the CFR (Jeon and Mize, 1993).

\textit{Development:} This patch-cluster system is an exciting model for studying fiber-target interactions in the developing CNS for several reasons. First, the system receives fiber inputs from three sources, each of which uses a different neurotransmitter. These fibers must somehow establish patches during development, probably either by forming well-demarcated
bundles during outgrowth or by a process of fiber pruning as the fibers contact the clustered
cells. The time at which the three fiber systems reach the igl is for the most part not known.
Markers of acetylcholine do not reveal ACh containing fiber patches prior to P14 (McHaffie et
al., 1991). SN fibers traced by anterograde transport of DiI reach the igl as early as P51, but
a patch-like pattern of innervation has not been observed until after birth (Banfro et al., 1994).
Thus, patch formation probably occurs gradually sometime between P1 and P14.

We have shown that the clustered pattern of neurons forms earlier. Clustered cells can be
identified in the igl as early as P51 when DiI is injected into the CFR (Banfro and Mize,
1996). Thus, the clustered cells must either migrate from the SVZ in a clustered pattern or
group into clusters before their axons reach the CFR. We do not yet know whether the
clustered neurons establish this identify before or after they have migrated into the igl, but we
do know that the clusters are formed well before fibers from the SN or PPTN establish
obvious patches within this layer.

Figure 7 illustrates the possible steps involved in formation of the patch-cluster system in
the cat SC. In the first panel (E40), neurons in the igl have migrated and formed a layer, but
the clusters have not yet been established. Axons from the PPTN (red), substantia nigra
(green) and frontal eye fields (yellow) are growing towards the igl but have yet to form
patches. The second panel (E51) shows that some neurons in the igl have achieved an identity
as 'cluster' neurons (red circles), either by selective migration, target site recognition, or the
onset of some chemical substance. At this same stage the axons from all three afferent sources
have reached their target field but are still distributed diffusely within the igl. The third panel
(P7) suggests that another molecule (dark blue circles) is expressed in the 'cluster' cells that
promotes or strengthens synaptic contacts with the three afferents. The final panel (P21)
suggests that the fibers not contacting the clustered neurons retract and die, so that only the clustered neurons and fiber patches remain.

Is nitric oxide involved in this process? We have begun to examine this hypothesis by studying the onset of expression of NOS in the developing SC. Our results show that the earliest expression of NOS occurs by E28, the earliest age examined. At this age, cells within the SVZ are well-labeled by NADPH but no NADPH positive cells are found within the tectal plate. By E36, NADPH cells with radial processes can be seen extending into the deep layers of SC. By E41, NADPHd labeled cells in the SVZ form a well-demarcated wedge and those in the deep gray layer form a dense band of neurons that extend throughout the layer (Figure 8A). The bipolar morphology of some cells in both the SVZ and dgl suggests that these cells are still migrating into the tectal plate (Figure 8A).

Although NADPH labeled cells in the dgl are well developed by E41, the earliest appearance of labeled neurons within the igl does not occur until E51-E58. Even at E58, only scattered single neurons in the igl are NADPH positive (Figure 8B). Nevertheless, these labeled cells have a periodicity characteristic of the cell clusters found in the adult igl. By P3, NADPH positive neurons form obvious clusters (Figure 9A) and as many as 20 neurons can be seen in a single cluster by P7 to P14 (Figure 9B). Although the cell clusters are well-labeled by P7 to P14, NADPH positive fiber patches are not readily visible until P35 (Figure 9C), an age at which the numbers of NADPH labeled cells begins to decline. Thus, NOS expression in 'clustered' cells in the igl occurs just prior to birth, reaches a maximum between P3 and P21, then declines gradually by the third postnatal month. The 'window' of maximal expression of NOS thus matches the time at which the afferent fibers in the igl form patches. There is thus a clear temporal correlation between NOS expression and afferent-target
matching in this system.

Whether NO is essential to this process is yet to be determined. We have recently begun experiments in which we have injected both Nω-nitro-L-arginine methyl ester (L-NAME) and Nω-nitro-L-arginine (L-NA) intraperitoneally into rat pups aged P1-P21. If NOS is involved in patch-cluster development, these NOS synthesis inhibitors should disrupt the development of the patches in this species. Our preliminary results are negative. PPTN fibers, labeled both by anti-choline acetyltransferase and NADPH, form normal appearing patches between P14 and P21 in both control and experimental groups (Scheiner and Mize, unpublished observations). We have not yet determined whether fibers from the SN and FEF are effected by these inhibitors or whether formation of the ACh fiber patches is delayed by inhibition of NOS.

Functional Considerations: Evidence that NO plays a role in synapse stabilization or pruning in other regions of the CNS is mixed. NO appears to be involved in both fiber retraction and topographic refinement in some species. In the chick optic tectum and rat superior colliculus, for example, peak NOS expression occurs in superficial layer cells at about the time at which retinal fibers reach this structure or synapses begin to form (Williams et al., 1994; Vercelli et al., 1995). Similarly, monocular enucleation either reduces NOS expression (Williams et al., 1994) or delays its expression in these neurons (Vercelli et al., 1995). More importantly, blockade of NO production using NOS inhibitors alters retinal topography in that the ipsilateral retinal-tectal pathway fails to retract in chicks (Wu et al., 1994) and the ipsilateral and contralateral retinal afferents fail to segregate in rats that have been treated with L-NA (see Wu et al., this volume).
Similar effects have been shown in the ferret lateral geniculate nucleus, where NOS inhibitors block segregation of the "on" and "off" sublaminae during a critical period (Cramer and Sur, 1994). By contrast, NO is apparently not involved in the development of other pathways. The segregation of contralateral and ipsilateral afferents is not disrupted by NOS inhibitors in the ferret LGN (see Cramer and Sur, this volume); topographic refinement of the ipsilateral and contralateral retinal afferents in SC is not disrupted in genetic "knockout" mice that presumably lack neuronal NOS (Frost et al., 1994); and infusion of NOS inhibitors into visual cortex does not block formation of ocular dominance columns in cat (Gillespie et al., 1993). Thus, NO has different effects in different species with the best evidence suggesting that NO is involved in fiber retraction in chick SC and some but not all types of fiber segregation in ferrets LGN and rat SC.

Study of the mechanisms underlying the role of NO in development is in its infancy, and many components of the process have not yet been examined. Nevertheless, the hypothesis that a free radical gas serves as a retrograde messenger is a highly attractive and potentially important advance in our understanding of brain development and worthy of further study. Our patch-cluster model will be particularly illuminating in this regard in that it will be possible for us to examine whether NOS inhibition effects different fiber pathways, one of which uses glutamate and two of which do not. The system is also unique in that NOS is expressed in both post-synaptic clustered neurons but also in one set of afferent fibers which innervate these cells.
Summary and Conclusions

Neurons within the superior colliculus (SC) contain a variety of neurochemicals, including the amino acid neurotransmitters GABA and glutamate, the calcium binding proteins calbindin and parvalbumin, and the neuromodulator nitric oxide. We have examined the development of expression of these substances using antibody immunocytochemistry. These results are summarized in Figure 10. GABA and calbindin are expressed very early in development, at a time when cells are still dividing and migrating from the subventricular zone. The expression of both GABA and CB is maximal at around E40-46, the age at which these cells have just established their adult lamination and extrinsic afferents have begun to grow into the tectum. GABA and CB likely play diverse roles during this stage of development, including the regulation of intracellular calcium during cell migration and neurite outgrowth.

Glutamate is expressed somewhat later in development while parvalbumin immunoreactivity does not appear until shortly after birth. These two substances continue to increase in density throughout the period of postnatal growth, at a time when synapse formation and evoked electrical activity are beginning to develop. Both PV and glutamate may be involved in one or both of these activity-dependent processes.

Nitric oxide synthase (NOS) is expressed at different times in different cell groups. NOS appears very early in prenatal development in cells within the SVZ and in the deep gray layer of SC. On the other hand, cells within the intermediate gray layer of SC do not express NOS until shortly before birth. The igl cells that express NOS at this age are clustered neurons similar to those that project to the CFR in the adult. NOS expression occurs in these cells at precisely the time when axons begin to form patches that innervate these clusters. Based upon
this temporal correlation, we hypothesize that nitric oxide may regulate synapse formation in this cell group.
REFERENCES


Figure Captions

Figure 1. GABA immunoreactivity in the prenatal kitten superior colliculus at E30. (A) Dense immunoreactivity is already present in the superficial layers (SL) of the tectal plate. Scattered labeled cells are also found within the deep layers (DL) and the subventricular zone (SVZ). (B) SL labeling consists of densely labeled cell bodies and neuropil. Most cells in the DL and SVZ have a bipolar morphology typical of migrating neurons; (C) Higher magnification view of vertical fusiform cell bodies and vertically oriented dendritic processes (arrows). Asterisk in A marks sites shown at higher magnification in B,C. VZ = ventricular zone; CA = Cerebral aqueduct. Scale bars: A = 200 μm; B = 50 μm; C = 20 μm.

Figure 2. GABA immunoreactivity in the prenatal kitten superior colliculus at E46. (A) The distribution of cells is like that of the adult, with large numbers of cell bodies in the superficial (SL) and intermediate layers (IL) and fewer neurons in the deep layers (DL). Note that labeled cells are also present in the dorsal periaqueductal gray (PAG). Well-labeled cell bodies and punctae in the neuropil in the SL (B), IL (C), and DL (D). Asterisks in A mark locations illustrated in panels B,C, and D. Scale bars: A = 100 μm; B-D = 20 μm.

Figure 3. Calbindin (CB) immunoreactivity in the prenatal kitten superior colliculus (SC) at E28. (A) CB neurons are distributed throughout the tectal plate and within the dorsal subventricular zone (SVZ). The morphology of these cells differs in different regions. (B) CB neurons near the surface of SC are small with few or no dendritic processes. Cells within the intermediate region of the tectal plate are larger and have multipolar processes. (C) CB labeled neurons within the deepest region of SC and within the SVZ have bipolar morphologies with vertically ascending dendrites typical of migrating cells (small arrows). Asterisks and large arrows in A mark the location of regions illustrated in B and C. CA = cerebral aqueduct; VZ = ventricular zone. Scale
bars: $A = 100 \mu m$; $B-C = 20 \mu m$.

Figure 4. Calbindin (CB) immunoreactivity in the prenatal kitten superior colliculus (SC) at E34, E40, and E46. (A) CB labeled cells are found throughout the tectal plate as well as within the subventricular zone (SVZ) at E34. Many cells in the deep layers and SVZ have a bipolar morphology. Arrowheads mark the superficial, intermediate, and deep layers of SC at this age. (B-C) Three tiers of CB neurons are visible by E40 (B) and well-developed by E46. (C). Most cells are small, but a few have larger cell bodies (arrows). sdt = superficial dense tier; idt = intermediate dense tier; ddt = deep dense tier; pag = periaqueductal gray; CA = cerebral aqueduct; VZ = ventricular zone. Scale bar: A,B,C = 100 \mu m.

Figure 5. Parvalbumin (PV) immunoreactivity in the postnatal kitten superior colliculus (SC) at P7, P21, and P35. (A) PV immunoreactivity is present in medium and large sized cells at P7. The dense band is not yet visible. There are no cells in the superficial gray layer (sgl). (B) By P21, a dense band of labeled neurons is visible within the ol (optic layer) and scattered PV neurons are seen deep. (C) An obvious dense band of PV immunoreactivity in neurons and neuropil is visible at P35. Scattered cells in the intermediate gray layer (igl) and deeper layers (dl) are intensely labeled. Scale bar. A-C = 200 \mu m.

Figure 6. The patch-cluster system in the intermediate gray layer of the cat superior colliculus. Clustered neurons project to the cuneiform region (CFR) of the caudal midbrain. Clustered neurons overlap patches of afferent fibers from the pedunculopontine tegmental nucleus (PPTN) that contain acetylcholine (ACh), from the substantia nigra (SN) that contain GABA, and from the frontal eye fields (FEF) that probably contain glutamate (Glu). Synaptic contacts involving various receptors are suggested but have not yet been demonstrated experimentally. Evidence suggests that the clustered neurons express nitric oxide synthase (NOS) transiently during
development.

Figure 7. Development of the patch-cluster system in the intermediate gray layer (igl) of the cat superior colliculus. Early in prenatal development (E40), neurons within the igl (blue circles) have no identifiable neurochemical characteristics that distinguish the clustered neurons. Extrinsic axons from the pedunculopontine tegmental nucleus (PPTN, red), the substantia nigra (SN, green), and frontal eye fields (FEF, yellow) have not reached the SC at this age. By E51, the clustered neurons (red circles) are labeled by DiI injected into the cuneiform region (CFR), indicating that the clusters are established at the time their axons reach the CFR. Axons from the SN (green), and possibly from the PPTN (red) and FEF (yellow), have reached the igl, but no patches are present. By P7, clustered cells avidly express NOS (dark blue circles). Our hypothesis predicts that the patches are beginning to form and that axons not contacting NOS containing neurons are retracting from the igl. By P21, the adult patch cluster system has been established and NOS expression is reduced (light blue circles).

Figure 8. Nicotinamide adenine dinucleotide phosphate diaphorase (NADPHd) labeling in the prenatal kitten superior colliculus (SC) at E41 and E58. (A) At E41, NADPHd labeled neurons form a wedge of cells in the dorsal subventricular zone. Labeled cells are also present in the deep layer (dgl) of SC. A few bipolar neurons extend from the deep layer towards the intermediate layer of SC at this age (arrows). (B) By E58, single neurons labeled by NADPHd (arrows) can be seen in the intermediate gray layer (igl). The periodicity of these cells is similar to that of clustered neurons in the igl. Scale bars; A-B = 100 μm.

Figure 9. Nicotinamide adenine dinucleotide phosphate diaphorase (NADPHd) labeling in the postnatal kitten superior colliculus (SC) at P3, P14, and P35. (A) Two clusters (arrows) of NADPHd labeled neurons are visible in the intermediate gray layer (igl) by P3. Labeled cells are
also seen in the deep gray layer (dgl) and the periaqueductal gray (pag). (B) By P14, the
NADPHd clusters (arrow) contain more labeled neurons and a few fibers. (C) By P35, the
number of NADPHd labeled clustered cells has decreased, but fiber labeling is intense (arrow).
Scale bar: A-C = 100 μm.

Figure 10. Summary diagram showing the time course of pre- and postnatal development of the
cat superior colliculus. The time scale is indicated by the large central arrow. Developmental
events, including cell generation, migration, lamination, afferent innervation, synaptogenesis, and
the onset of neuronal activity are shown above that line. Mitosis, migration, and lamination occur
early and are largely complete by E40-46. Synaptogenesis and neuronal activity occur later,
usually at or shortly after birth. Afferent innervation occurs at different times for different
afferents. The time of onset and maximal expression of gamma aminobutyric acid (GABA),
calbindin (CB), glutamate (GLU), parvalbumin (PV), and nitric oxide synthase (bNOS) are
shown below the time line arrow. GABA and CB are expressed early, GLU and PV later, and
bNOS at various times during pre- and postnatal development.
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Inhibition of Nitric Oxide Synthase Fails to Disrupt the Development of Cholinergic Fiber Patches in the Rat Superior Colliculus.

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Abstract

Nitric oxide (NO) may serve as a retrograde messenger to refine or stabilize synapses in the developing nervous system. Whether this action is dependent upon glutamate and the N-Methyl-D-Aspartate (NMDA) receptor is not yet established. We have used the patch-cluster system in the intermediate gray layer (IGL) of the rat superior colliculus (SC), a system receiving both glutamatergic and cholinergic input, to study this question. The normal distribution and development of nitric oxide synthase (NOS) in SC was examined using nicotinamide adenine dinucleotide phosphate diaphorase (NADPHd) histochemistry in Sprague-Dawley rats aged P4 to adulthood. Fibers containing acetylcholine (ACh) were identified using choline acetyltransferase immunocytochemistry. In addition, Nω-nitro-L-arginine (NoArg), an inhibitor of NOS, was injected intraperitoneally from birth until P10, P14, P18, or P21-22 to determine if NOS inhibition would disrupt the formation of the ACh patches. Control animals were studied from the same age groups. Our results show NADPHd labeled cells within the periaqueductal gray (PAG) and the deep gray layer (DGL) of SC by P4, the earliest age examined. By P8-P9, cells in the IGL were well labeled by NADPHd, while few in the superficial layers (SL) were labeled. SL cells were visible by P10 and were intensely labeled by P14. IGL cells transiently expressed NADPHd in that the number of labeled cells increased from P8-P35, then decreased in the adult. ChAT labeled fibers first appeared in the IGL at P10, formed a characteristic two-tier pattern by P14, and established obvious patches by P21. Inhibition of NOS from birth produced no qualitative differences in the distribution or density of either ChAT labeled fibers or NADPHd labeled cells and fibers at any of the ages examined. We therefore conclude that NO does not contribute to the refinement of cholinergic fiber patches in the rat SC, probably because the fiber system is not glutamatergic.
Introduction

Nitric oxide (NO) is a free radical gas with diffusion properties that make it an excellent candidate for a retrograde messenger in the central nervous system. Release of NO is thought to start with an influx of calcium through the N-Methyl-D-Aspartate receptor which activates nitric oxide synthase (NOS) to produce NO that diffuses from the postsynaptic cell into presynaptic terminals. NO is then thought to act upon the presynaptic guanylate cyclase-cGMP system to enhance neurotransmitter release, probably glutamate[1,2]. This process has been implicated in the synaptic plasticity that occurs during long term potentiation (LTP)[3-8] and in axon refinement during development [7,9].

Evidence for this hypothesis in development is, to date, scanty. In the visual system, NO appears to be involved in both fiber retraction and the refinement of topographic maps. Peak NOS expression occurs in cells at the time at which axons from the retina are retracting in both the chick optic tectum [10] and the rat superior colliculus (SC) [11]. Monocular enucleation reduces NOS expression during this period [10] and interruption of NO release using inhibitors of NOS disrupts topographic refinement of both the ipsilateral and contralateral retina-tectal pathways in chicks [11,12]. A similar effect has been shown in the lateral geniculate nucleus of the ferret where NOS inhibitors block segregation of the "on" and "off" sublaminae of this nucleus [13,14]. On the other hand, the eye-specific segregation of ipsilateral and contralateral retinal input to the ferret LGN is not disrupted by inhibition of NOS [14] and the topography of ipsilateral and contralateral retinal afferents projecting to the SC is not altered in nNOS genetic knockout mice that presumably lack NO [15]. Our understanding of the role of NO in fiber refinement is, therefore, in its formative stages.

A number of questions remain unanswered. For example, it is not yet established whether NO affects only glutamatergic synapses or whether it can also enhance transmitter release or otherwise
affect synapses that use other neurotransmitters [16,17]. Nor is it known whether the NMDA receptor is the only source of Ca\(^{2+}\)-influx that regulates the production of NOS. Other sources of increased calcium could include the inositol trisphosphate (IP\(_3\))-dependent release of intracellular calcium stores after activation of the G-protein linked ACh receptor or calcium influx through various voltage-gated calcium channels.

The model that we have chosen to address these questions is the patch-cluster system in the mammalian SC. This system in cat consists of 'clusters' of neurons within the intermediate gray layer (IGL) that project through the tectopontobulbar pathway to the region of the cuneiform nucleus [18]. These cell clusters precisely overlap fiber patches in the IGL that arise from mesencephalic tegmental nuclei and that contain acetylcholine [19,20]. Fibers from the substantia nigra that contain GABA [21], and those from the prefrontal eye fields that probably contain glutamate, also form patches that overlap those containing ACh [22,23]. The patch-cluster system is not as well defined in the rat, but fiber patches in the rat SC also arise from brainstem nuclei and contain ACh [20].

The patch-cluster system is of particular interest because we have shown that neurons that form clusters in the cat IGL first express NOS [24,25] at about the time that the ACh afferents innervate the IGL [26]. These ACh afferents then establish their characteristic patch-like distribution during the time at which clustered cells are heavily labeled by NADPH\(_{d}\) [24]. In the present study, we have determined whether the patch-cluster system in rat is disrupted by inhibition of NOS. In addition, we have examined the normal distribution and onset of expression of NOS in this species.
Materials and Methods

Animals and drug treatment

We examined the distribution of nitric oxide synthase (NOS) and choline acetyltransferase (ChAT) in normal adult and postnatal rats and in rats in which NOS was inhibited. Adult rats were obtained from Harlan Sprague Dawley (Indianapolis, IN). Timed pregnancies were also obtained from this supplier. Birth dates were determined to within 24 hours. All procedures involving the use of experimental animals were approved by the LSU Medical Center IACUC office.

Rat pups from newly born litters were divided randomly into experimental and control groups. Experimental animals received intraperitoneal injections of 1, or 3 µmol/g body weight No-Nitro-L-arginine (NoArg, Sigma, St. Louis) daily from P1 (the day after birth) until day of sacrifice. Control animals received injections of comparable volumes of normal saline. Control and inhibited animals were perfused at postnatal ages P8 (N=2), P10-11(N=6), P14(N=6), P18(N=3), P21-22(N=6), and P28(N=3). At least one additional rat from ages P4, P5, P9 and P35 was used to examine the normal development of NADPH labeling. The P28 case was discarded because of poor fixation.

Animals were anesthetized with a Ketamine hydrochloride/xylazine mixture (80 mg/kg body weight) and perfused transcardially with 4% paraformaldehyde, 0.1% glutaraldehyde in phosphate buffer at pH 7.4. Glutaraldehyde was omitted in the normal rats aged P4-P9 and in one adult. After perfusion, all brains were removed, post-fixed in the same fixative for 4-18 hr., and placed in a phosphate buffer at pH 7.4 with 8% dextrose added. The following day, the brains were blocked, cut on a vibratome into 50-100 µm sections, and treated for histochemistry or immunocytochemistry. All sections were then mounted on glass slides, dehydrated in alcohol/xylene, and coverslipped.
Histochemistry procedures

Sections treated to identify NOS were reacted with nicotinamide adenine dinucleotide phosphate diaphorase (NADPHd) histochemistry. Sections were first rinsed in tris buffer and placed in a NADPH solution containing 50mg NADPH, 12.5 mg nitro blue tetrazolium, and 50ml 0.3% Triton X-100, tris buffer, pH 7.1 at 37 °C for 1-3 hrs. The sections were then rinsed in phosphate buffer, dehydrated, and coverslipped. Sections treated for choline acetyltransferase (ChAT) immunocytochemistry were prepared using the ABC technique (Vector Laboratories, Burlingame CA). Sections were rinsed in phosphate buffered saline (PBS) and incubated in blocking serum (4% normal rabbit serum) for 1 hr. They were then incubated in a 1:500 dilution of goat anti-ChAT (Chemicon International, Temecula, CA) for 44-48 hrs, rinsed, and incubated with secondary antisera (rabbit anti-goat, 1:100 in PBS), rinsed again, then incubated with avidin biotin complex (ABC) for one hour. Sections were then rinsed and reacted with 0.05% 3,3’-diaminobenzidine (DAB) in tris buffer with 0.003% H₂O₂. Sections from most experiments were enhanced with 0.005% nickel chloride/cobalt acetate [27] for 90-150 sec, rinsed and treated as above.

Determination of NOS inhibition in brain

We examined the amount of NOS inhibition produced by Nω-Nitro-L-arginine in P14, P18, and P22 rat pups. Two P14, one P18 and seven P22 rats received daily injections of 1 μmol/g body weight NoArg, and two P14 and one P18 rat received 3 μmol/g body weight daily injections prior to sacrifice. Each age group was compared with at least one saline injected pup of the same age. Rat pups were decapitated, their brains rapidly removed and homogenized in 20 mM Hepes containing 0.32 M sucrose, 1mM dithiothreitol (DTT), and 0.5 mM EGTA for 1 minute [28]. The homogenate was centrifuged for 15 minutes at 4°C and then the supernatant was removed and centrifuged again for an additional 15 min. The cytosolic homogenate (supernatant) was removed and applied to 1 ml columns of Dowex AG50W-X8 (Na⁺ form) to remove endogenous arginine. The protein content of the arginine-free homogenate was determined. Aliquots containing 300 mg
of protein were then incubated for 15 minutes at 37 °C with buffer containing 20 mM Hepes, 0.32 mM sucrose, 1mM DTT, 0.5 mM EGTA, 0.5 mM CaCl₂, 200 μM NADPH, 1μM L-arginine, and 0.1 μCi/ml [³H]L-arginine. The reaction was stopped by adding 2 ml of 20 mM Hepes containing 2mM EDTA (pH=5.5). Samples were applied to Dowex AG50W-X8 (Na⁺ form) to remove [³H]L-arginine and elute any [³H]L-citrulline. The columns were then washed with 3.0 ml of water and the [³H]L-citrulline content was quantified by liquid scintillation spectroscopy of the flow-through and water rinse. DPM’s were calculated and the data expressed as pmoles of L-citrulline/min/mg protein for each animal. From this data we calculated the percent inhibition compared to the age group control.

Data Analysis

Coronal sections through the rostral-caudal extent of SC were examined with the light microscope from at least one control and one experimental animal at ages P10-11, P14, P18, and P21-22. Both NADPH and ChAT labeled sections were examined and photographic records taken using a Zeiss Universal camera system. We also examined and photographed sections from normal animals aged P4, P5, P8, P9 and P35 that were labeled only with NADPH. The laminar distribution of NADPH labeled neurons in normal control animals was examined quantitatively by use of a computer based Neuron Tracing System (NTS; Eutectic Electronics, Rayleigh NC). Between 3 and 6 sections from at least one animal aged P4, P5, P8, P9, P10, P11, P14, P18, P21, and one adult, were plotted. Plots were produced by outlining the surface of SC by moving a screen cursor with a joystick. The position of each NADPH labeled neuron was also recorded. Histograms were generated from these plots to illustrate the relative number of cells at different depths within the SC at different ages.

We also estimated quantitatively the number of NADPH labeled neurons within the IGL of SC at different ages. Cell counts were made manually at 20X magnification from 3-10 sections from control animals at ages P4, P5, P8, P9, P14, P18, P21, P35, and from two adult rats. These counts were expressed as number of cells within the IGL for each section at each age.
Results

Distribution of NOS and ChAT in the adult rat SC

NOS was present in only a small subset of neurons within the juvenile (P35) and adult rat superior colliculus, as revealed by NADPHd histochemistry. In addition, selected fiber pathways were intensely labeled by NADPH. The densest concentration of NADPHd positive neurons was found within the zonal (ZL) and superficial gray (SGL) layers of SC. These neurons formed a dense band that extended from the surface of SC to the ventral border of the SGL. The band consisted of intensely labeled cell bodies, dendrites, and axon-like fibers that extended throughout the rostro-caudal axis of SC (fig. 1A-C).

Many fewer NADPHd positive neurons were found below the SGL. Only a few lightly labeled neurons were seen within the optic layer (OL) and no labeled fibers were found in that layer (fig. 1A-C). Scattered well-labeled neurons were present within the intermediate gray (IGL) layer. These neurons were mostly located within NADPH labeled fiber patches within the IGL (fig. 1B-C, arrows). NADPH positive fibers in this layer also had a characteristic distribution. Within the rostral SC, they formed a single continuous band (fig. 1A). In the middle and caudal SC, they were distributed in two separate tiers, one within the dorsal IGL, the other within the ventral IGL (fig. 1B-C). Both tiers had patches of heavy fiber labeling (fig. 1B,C, arrows) with interpatch intervals that contained many fewer fibers. NADPH labeled fibers also formed bridges between the two tiers (see fig. 1B,C).

The remaining cellular and fiber labeling in SC was distributed primarily within columnar streams within the deep gray (DGL) layer. Most of the labeled fibers projected vertically towards the IGL and most of the labeled cells were located within these fiber streams (fig. 1B,C). NADPH positive fibers and cells were also found within a dorsolateral wedge of the periaqueductal gray (PAG).
Some cells within this wedge had dendrites that projected into the deep layers of SC (fig. 1A-C). Finally, NADPH labeling was found within the mesencephalic tegmentum where it intensely labeled cells within three nuclei: the parabigeminal nucleus (PBG), the pedunculopontine tegmental nucleus (PPTN), and the lateral dorsal tegmental nucleus (LDTN). All three of these nuclei send projections to the rat SC [20].

Labeling by antibodies directed against choline acetyltransferase (ChAT) showed a very similar pattern to the fiber labeling seen using NADPHd histochemistry. A dense band of ChAT labeled fibers was present in the ZL-SGL (fig. 2A-B). ChAT labeling within the rostral IGL formed a single tier of fibers while more caudal labeling revealed two tiers of ChAT labeled fibers, each with densely labeled patches (fig. 2A,B). Labeling of adjacent sections for ChAT and NADPH at P35 showed that this two tier fiber pattern was identical using both labels. This result coupled with previous reports [29,30,20] provide strong evidence that ChAT and NOS are contained in the same fiber groups throughout the IGL.

In summary, the juvenile and adult rat SC has a distinct pattern of cells and fibers that contain NOS. These cells and fibers are found primarily within the SGL, IGL, and DGL. The NADPH labeled fibers also contain ChAT and arise from cell groups within the mesencephalic tegmentum.

*Development of NADPH labeling in the postnatal rat*

NADPH labeling in SC at P4-P5 was confined primarily to small cells within the dorsolateral wedge of the PAG and in neurons scattered within the DGL (fig. 3A). Only a few lightly labeled neurons were seen within the IGL at P4 and none within the more superficial layers. In the P5 case, a cluster of lightly labeled cells was seen within the lateral IGL in caudal sections (fig. 3,5). No NADPH labeling was resolved within axon-like fibers at P4-P5, but many blood vessels did contain NADPH labeling (fig. 3A).
By P8-P10, the number of labeled cells in the PAG wedge had increased, and many more labeled cells were present in SC (figs. 3A,5). Some neurons within the PAG had vertically oriented dendrites which projected into the DGL and appeared to be migrating from the PAG (fig. 3B). Labeled cells within the IGL were now found at both rostral and caudal levels and in both the medial and lateral SC. In caudal sections, these cells were often grouped into clusters (fig. 3B-D). Some very lightly labeled neurons were also scattered within the SGL by P9 (fig. 5). However, no labeled axons were seen in any layer of SC between P8 and P10. Blood vessel labeling was still prominent.

By P14, clustered neurons in the IGL were quite well labeled by NADPH and these cells were found in both the superficial and deep tiers of this layer (figs. 4A,C, 5). In addition, neuropil labeling within the IGL was now visible (fig. 4A), and a few axon-like fibers and boutons could be resolved (fig. 4C). Cellular labeling within the SGL at P14 was intense and many more cells were labeled than at P8-P10. Cell labeling in the PAG wedge and DGL was similar to that seen earlier.

By P18-P21, NADPH labeling was intense in neurons in both the IGL and SGL (fig. 5). Fiber labeling was also dramatically increased. By P21 fibers formed a typical two-tier patch-like pattern similar to that seen in the juvenile and adult. Within these patches, cells were darkly stained and many axon-like fibers with varicosities and terminal boutons were also well labeled by NADPH (fig. 4B,D).

Fig. 5 summarizes the development of NOS cells within SC. NADPH labeled neurons were present in the PAG and DGL and scattered within the lateral IGL by P4-5. Significant NOS expression in IGL cells occurred by P8-9 and reached a maximum between P14-P35. SGL cellular labeling was first seen at P9-P10 but was not prominent until P14.
**NADPH expression in the IGL**

In an effort to determine the time-course of NOS expression in IGL neurons, we counted the number of NADPH labeled neurons within the IGL at different ages. No more than 10 labeled neurons were found in any section at P4, while an average of 20 was found by P5 (fig. 6). By P8-9, the number of NADPH labeled neurons in the IGL ranged from 26 to 75 with a mean of 43.4. This is a two fold increase from that seen at P5. The average number found between the ages of P18-21 ranged from 40-45 and peaked at 56.3 at P35. This number decreased significantly in adults (fig. 6), despite a substantial increase in the area of the IGL. These data thus confirm quantitatively that the number of NADPH positive neurons increases during the period of fiber development in the IGL, and then decreases in the adult.

**Distribution of ChAT labeled fibers in normal development and after NOS inhibition**

We examined the distribution of fibers labeled by anti-choline acetyltransferase and by NADPHd, in both control and NOS inhibited rats between the ages of P8 and P21. ChAT fiber labeling in control animals developed between P8-P21. No ChAT fiber labeling was found at P8, even when we prolonged the DAB reaction to overstain the tissue. At P10-11, anti-ChAT fibers were visible in the IGL, but there were relatively few fibers at this age and most were lightly labeled. The density of ChAT labeled fibers in the IGL was much higher at P14, as was the intensity of labeling within individual fibers. The two tier pattern was visible, although the tiers were not as well segregated as at later ages (fig. 7A). ChAT labeled fibers had a nearly continuous distribution across the IGL and few or no obvious patches were present at this age (fig. 7A). ChAT fiber labeling at P18 showed further differentiation of the tiers. The two tiers were thicker, there were some gaps within the tiers, and more fiber columns bridged the tiers (fig. 7B). By P21, ChAT labeling in the IGL was virtually identical to that seen in the P35 juvenile rat. Both fiber tiers were densely labeled and there were a number of fiber patches separated by interpatch intervals that had many fewer fibers (fig. 7C).
The pattern of ChAT labeling in the NoArg inhibited animals was virtually identical to that seen in controls (fig. 7D-F). Few fibers were present at P10 while the two-tier pattern was visible at P14 (fig. 7D). The tiers were further developed by P18 (fig. 7E) and patches were conspicuous by P21 (fig. 7F). Virtually no qualitative differences could be detected at any age. We thus conclude that inhibition of NOS has no affect on ACh fiber development in the rat SC.

NADPHd labeling also appeared to be unaltered by inhibition of NOS. Intense cellular, and light fiber labeling were observed in the IGL by P14. As in normal animals, labeled cells in the IGL sometimes appeared in clusters (fig. 8A). More obvious fiber labeling was observed in the IGL by P18 and many cells were labeled at this age as well (fig. 8B). By P21, NADPHd labeled fibers in the IGL formed two obvious tiers (fig. 8C), just as seen in normal P21 rats (see fig. 4B). The pattern of NADPHd labeling in the SGL and DGL was also similar to that seen in normal animals. We thus conclude that NOS inhibition, in the ranges we have tested, has no discernible effect upon NADPHd labeling in the developing rat SC.

*Amount of NOS inhibition determined by biochemical assay*

We also performed biochemical assays for NOS in additional experimental and control animals sacrificed at P14, P18, and P22. The assay results revealed a substantial reduction in NO synthase activity in all age groups after daily injections of NoArg. The L-citrulline produced in P14 homogenates after 1 μmol/g body weight injections ranged from 3.42 - 3.64 pmoles/min/mg protein compared to 14.16 pmoles/min/mg protein in the saline control. The percent inhibition was 74.3 to 75.9%. The reduction after 3μmol/g body weight injections was greater; L-citrulline ranged from 2.30-2.78 pmoles/min/mg protein, an 80.4-83.8% inhibition compared to control.

A similar dose dependency was found in the P18 rats. NOS activity after daily injections of 1 μmol/g body weight was reduced by 72.1% to 3.10 pmoles/min/mg protein, while after 3μmol/g body weight injections it was reduced by 91.0% to 1.0 pmoles/min/mg protein. Results from seven P22 rats were similar. Injections of 1μmol/g body weight reduced L-citrulline production from
17.38 pmoles/min/mg protein in the saline control to 1.95-3.19 pmoles/min/mg protein. This corresponded to an 81.6 to 88.8% inhibition of NOS activity. No correlation was found between the degree of inhibition and the pattern or density of labeling by either ChAT or NADPHd.

**Discussion**

Our results lead to the following conclusions: 1) the patch/cluster system exists in the IGL of the rat SC as it does in cat; the system appears to contain both acetylcholine and NOS; 2) the pattern of cellular labeling by NADPH develops in a ventral to dorsal sequence and is essentially adult-like by P21; 3) there is a transient expression of NOS in some neurons in the IGL. Maximal expression of NOS occurs during the period of fiber formation in the IGL and then declines in the adult; 4) this fiber pattern is neither altered nor delayed by inhibition of 72 to 91% of the endogenous NOS activity.

**NOS distribution in the adult SC**

Our results show that NO is a marker of the patch-cluster system in the rat SC as it is in the cat [24,25]. In both species NAPDHd intensely labels fibers within the IGL and these have a distribution virtually identical to those labeled by ChAT [20,28,30,31]. Like the cat, most NADPH labeled neurons within the rat IGL are found within the patches of labeled fibers. Although it has not been shown experimentally whether NOS containing cells are the same cells as the cluster neurons that project to the region of the cuneiform nucleus [18], the similarity in size and distribution of both cell groups makes this conclusion likely. This pattern of NADPH labeling in the IGL has also been found in the C57 wild type mouse [33, Scheiner and Mize, unpublished observations] and in the Cynomolgus monkey [34]. This system thus appears to be conserved across a variety of mammalian species.

The pattern of labeling within the other laminae of SC has been shown by others [33-39]. Much of the NADPH fiber labeling in SC is thought to arise from three nuclear groups within the
mesencephalic tegmental core: the pedunculopontine tegmental (PPTN) and lateral dorsal tegmental (LDTN) nuclei which give rise to the IGL fiber patches as well as some fibers within the DGL [19,20]; the parabigeminal nucleus (PBG) which projects principally to the SGL [19,20]; and the dorsolateral wedge of the periaqueductal gray (PAG) which apparently gives rise to fibers innervating the DGL [40, and present results]. These nuclear groups are intensely labeled by NAPDHd as well as by ChAT [29,30] and represent some of the major sources of NO in the brainstem [35].

The NADPH positive neurons within the ZL and SGL of the rat have varying morphologies that have been described by others [37,41]. These neurons overlap the retinal input to this region of SC [42,43] which is thought to be glutamatergic in both rat and cat [44-50]. Although the retinocollicular pathway in cat is also glutamatergic [50], the cat SC does not have a dense band of NADPH cells in the SGL [32,34]. Differences in NOS expression in the two species may be due to differences in the extent of binocular overlap or the functional types of retinal ganglion cell that project to the SGL. In any event, the pattern of NADPHd labeling in rat and cat is very similar in the IGL where inputs from the PPTN and SN in both species appear to be functionally similar.

Development of NOS expression in SC

Our results show that NADPH labeling develops in a ventral to dorsal sequence in the postnatal rat. In the first week after birth, NADPH positive cells are found only within the PAG and DGL and occasionally within the lateral IGL. Well labeled NADPH cells are first found in obvious IGL clusters only at P8-P10, a result previously reported [51]. The dense band of NADPH neuronal labeling in the SGL appears last, around P14-P15 [present results, 41,51].

Although the cells labeled by NADPH in the PAG, DGL, and SGL appear to retain NOS expression throughout life, the number of labeled cells in the IGL was substantially reduced in the adult. This suggests that some IGL cells express NOS transiently or that some NOS containing cells die during development. Transient expression is the more likely explanation given that there
appears to be little reduction in NOS cell number in the other layers. Transient expression as well as the time of onset suggest that NO could be involved in the retraction of IGL fibers into patches. Intensely labeled neurons in the IGL were visible by P10-P14, a time just preceding the development of ChAT and probably other fibers in this layer. The two tier pattern was seen by P14 while the patch-like distribution developed between P18 and P21. The ChAT labeled fibers thus appear to retract into patches during the time at which NOS expression is maximal in IGL neurons.

This phenomenon has also been seen in the cat SC. NADPH labeled cells are first found in the IGL between E51-E58 and clusters of these neurons are not seen until P3-P7. The clustered neurons intensely express NOS by P10-P14 [24,25], the age at which ChAT first labels ACh fiber patches in the cat IGL [26, Scheiner et al., unpublished observations]. As ACh containing fibers from the PPTN and LDTN apparently arrive in the IGL after P1 [26], the onset of expression of NOS in the clustered neurons correlates well with the arrival of the fibers as well as their expression of ACh. However, this relationship between NOS expression and fiber segregation could involve fiber systems other than those containing ACh. Although fibers from the substantia nigra (SN) reach the SC of cat as early as E51, SN fiber patches in the IGL have not been observed earlier than P19 (Banfro, Scheiner and Mize, unpublished observations). Thus, NO could be influencing fiber segregation of this pathway instead of, or in addition to, that from the PPTN/LDTN.

A temporal correlation between NOS expression and fiber refinement has also been reported in other species and in other regions of the brain. NADPH labeling first appears in superficial layer cells of chick optic tectum at about the time that retinal axons innervate the tectum, and this expression is maximal during development and reduced in adults [10,11]. Neurons in the LGN of the cat [30,52] and ferret [13,14] also express NOS only during the first several weeks of life. This transient expression is apparently developmentally regulated by afferent input because monocular enucleation results in continuous expression of NADPH in the non-deprived laminae of the adult LGN [30] while bilateral enucleation eliminates NADPH labeling in cells in the superficial layers of the chick optic tectum [10,11]. Thus, both the correlation found between the onset of
expression of NOS and refinement of axons and the alterations in NOS levels produced by
deafferentation suggest that NOS plays a role in axon fiber refinement or synapse stabilization in
these structures.

Absence of an effect of NOS inhibitors on ChAT fiber development

The original hypothesis of Gally [7] predicted that NO could serve as a retrograde signal to
'instruct' presynaptic axons that they had established appropriate connections with their post-
synaptic target. The hypothesis was explicit in stating that the post-synaptic release of NO
depended upon the influx of Ca^{2+} and that this could occur through the NMDA receptor. Release
of glutamate from a glutamatergic pathway was an implied component of the hypothesis.

A major purpose of the present study was to test whether NO might also serve as a retrograde
messenger in the refinement of axons that utilize another neurotransmitter, in this case ACh. The
patch-cluster system is an excellent model for studying this question because post-synaptic neurons
in the patches express NOS, the ACh fibers in the system gradually retract into patches, and the
two events are temporally correlated.

Our results, however, show that prolonged inhibition of NOS has no effect upon the pattern or time
course of development of the ACh patches. The two tier pattern was visible by P14, and the
patches were well demarcated by P21 in both experimental and control groups. Pilot studies in our
laboratory have also shown that ChAT fibers develop normally in a neuronal NOS genetic
knockout mouse [53]. Lower doses of NoArg than those used in our study are known to be
effective in altering fiber refinement in the retinotectal pathway of the rat [11, McLoon, personal
communication]. There is thus correlative evidence that the reduction or absence of bNOS has no
effect on formation of the ACh patches.

Although our biochemical assay showed that NOS activity was being substantially reduced (72-
91%) in the NoArg injected rats, NADPHd labeling was also not noticeably altered by inhibition of
NOS (fig. 8). This observation suggests to us that, while NADPHd labeling indicates the presence of the neuronal form of NOS [54], it does not depend upon NOS activity. Furthermore, it suggests that the lack of a NoArg effect on the ACh patches is not due to the disappearance of NOS containing cells in the target region.

To date, relatively few studies have, in fact, shown that NOS is involved in fiber refinement in developing CNS. The most complete studies have been performed in chick. The ipsilateral retinal projection, which is normally transient, is partially preserved by daily intraperitoneal injections of NoArg or NAME [11,12] and both of these inhibitors disrupt the topography of the contralateral retinotectal pathway [11]. McLoon and colleagues [11] have also shown that the ipsilateral and contralateral retinal afferents to the rat SC do not segregate completely after NoArg administration. A somewhat different effect has been reported in the ferret LGN, where the segregation of the 'on' and 'off' sublaminae is disrupted by either focal or systemic inhibition of NOS [13,14].

In both cases a glutamate pathway and the NMDA receptor are probably involved. The retinal pathways to SC and LGN are glutamatergic in all species studied [44 -47,49,55,56] and normal development of both pathways can be disrupted by NMDA antagonists [57,58]. In both ferret LGN and rat SC NOS inhibition and NMDA receptor blockade both appear to disrupt the retraction of axon arbors that would normally occur as a result of correlated electrical activity in other arbors [57,58]. The exact locus and mechanism of NO in these pathways is unknown and could include a presynaptic action such as an increase in guanylate cyclase-cGMP to promote enhanced release of neurotransmitter [1,2,5,59] or a post-synaptic action such as enhancement of an NMDA receptor mediated response [60]. These results suggest that NO may only play a role in fiber refinement in glutamatergic pathways involving the NMDA receptor. If this is so, then it remains possible that NO is involved in refinement of glutamatergic fibers that innervate the IGL, and this could explain the temporal correlation that was found between maximal NOS expression and fiber development in this region of the SC. We are currently conducting experiments to test this hypothesis.
On the other hand, it is worth noting that NO has been shown to affect synapse plasticity in non-glutamatergic systems. Thus, NO donors suppress evoked presynaptic currents while NO-binding proteins and NOS inhibitors block this suppression induced by asynchronous firing in the neuromuscular junction of Xenopus [16]. In addition, NO donors have been shown to arrest or reversibly collapse growth cones in vitro in the dorsal root ganglion of the rat [17]. Thus NO can affect axon development in non-glutamatergic as well as glutamatergic synapses, but the mechanism appears to be one of suppression of non-correlated firing in the former and an enhancement of correlated firing in the latter.
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Figure Captions

Figure 1. Superior colliculus labeled by NADPHd histochemistry in a P35 juvenile rat. (A) Rostral SC showing band of NADPH labeling in the SGL but very little fiber or cell labeling in the IGL; (B) Middle SC showing increased NADPH fiber and cell labeling in the IGL (arrows). Note scattered neurons in the DGL. (C) Caudal SC showing the two tiers of fiber labeling in the IGL as well as fiber patches (arrows). Labeling in the SGL and DGL is similar to that in B. sgl = superficial gray layer; ol = optic layer; igl = intermediate gray layer; dgl = deep gray layer; pag = periaqueductal gray. Scale bar = 200 μm.

Figure 2. Fibers labeled by choline acetyltransferase (ChAT) antibody in the superior colliculus (SC) of a P35 juvenile rat. (A) middle SC section; (B) more caudal SC section. Note the band of immunoreactivity within the SGL as well as the two tiers of fiber labeling within the IGL. Fiber patches are indicated by arrows.

sgl = superficial gray layer; ol = optic layer; igl = intermediate gray layer; dgl = deep gray layer; pag = periaqueductal gray. Scale bar = 200 μm.

Figure 3. NADPH labeled neurons in the neonatal rat superior colliculus. (A) At P5, small NADPH positive neurons are seen within the PAG and DL. Only 4-5 lightly labeled cells are visible in the IGL (arrows). (B) By P8, neurons within the PAG and DL are well-labeled by NADPH. A number of neurons are also seen within the IL, mostly within the lateral half of that layer. Asterisk marks regions illustrated in C and D. (C) Higher magnification view of IL cells illustrated in B. Note the dense labeling in the dendrites of some neurons. (D) High magnification view showing the morphology of the neurons in C. Note dendritic labeling in these cells. sl = superficial layers; il = intermediate layers; dl = deep layers; pag = periaqueductal gray. Scale bar, A-B = 200 μm; C = 50 μm; D = 20 μm.

Figure 4. NADPH labeled neurons in the neonatal rat superior colliculus. (A) At P14, NADPH labeled neurons are seen throughout the DGL, IGL, and SGL. Some neuropil labeling is also seen. 

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within the IGL and SGL. Asterisk marks region illustrated in C. (B) By P21, many NADPH labeled neurons are seen within the IGL and SGL. In addition, fiber labeling within the IGL is obvious and forms a two tier pattern similar to that seen using antibodies to ChAT (see Figure 2). Asterisk marks region illustrated in D. (C) Higher magnification of labeled IGL cells illustrated in A. Note that some thin axon-like fibers and boutons are labeled by this age. (D) Higher magnification of labeled neurons illustrated in B. Note the dense thin fiber and bouton labeling at this age. sgl = superficial gray layer; ol = optic layer; igl = intermediate gray layer; dgl = deep gray layer; pag = periaqueductal gray. Scale bar, A,B = 200 μm; C,D = 20 μm.

Figure 5. NTS computer plots of the distribution of NADPH labeled neurons in the superior colliculus of neonatal rats (P5 to P21). A few neurons are present in the IGL by P5. The earliest neurons within the SGL are seen at P9. High densities of NADPH labeled neurons are seen by P14.

Figure 6. Scattergram showing the total number of NADPH labeled neurons found within the intermediate gray layer (IGL) of the normal rat superior colliculus at different ages. Neuron number increases between P8 to P35, then declines in the adult.

Figure 7. Anti-choline acetyltransferase (ChAT) labeling in fibers within the igl of the rat superior colliculus at different ages (P14, P18, P21). (A-C) Control sections from animals injected with saline; (D-F) Sections from experimental animals injected with a NOS inhibitor (Nω-nitro-L-arginine). There were no obvious qualitative differences in the distribution or density of labeled fibers between the two groups.

Figure 8. NADPH labeling in the neonatal rat superior colliculus at different ages following daily injections of the NOS inhibitor Nω-Nitro-L-arginine. (A) By P14, cellular labeling in the IGL is intense while fiber labeling is present but light. (B) By P18, more fiber labeling is visible in both the IGL and SGL. (C) By P21, NADPH labeled fibers form two tiers within the IGL, just as they do in the normal rat. Scale bar = 200 μm.
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304.3
ONOFF SUBSTIMULATION IN THE KERF LGN IS INDEPENDENT OF SYSTEMIC CHANGES IN BLOOD PRESSURE AND REQUIRES NEURONAL NO. E.S. Craner and M. Sar. Department of Brain & Cognitive Sciences, M.I.T., Cambridge, MA 02139. Blockade of nitric oxide synthesis (NO) with N-Nitro-L-Arginine (L-NaORg) during the third and fourth postnatal weeks in the LGN demonstrated that ONOFF substimation (Craner and Sar, 1994). Because NO is a potent vasodilator, we examined the role of systemic changes in blood pressure on substimulation by systemic treatment of fetal rats with L-NaORg (MAP). L-NaORg was administered during the fourth postnatal week in four maternal animals with 40 mg/kg/day N-Nitro-L-Arginine (L-NaORg) from postnatal day 14 (P14) and in four normal, age-matched controls (MAP 4.5 ± 1.4 mmHg, L-NaORg 5.4 ± 2.6 mmHg). Neonatal L-NaORg treatment. When animals received the antihypertensive calcium channel blocker verapamil, 5 mg/kg/day, together with L-NaORg, blood pressure was normal (0.9 ± 0.8 mmHg). Normal saline administered in sections of LGN contralateral to an intraocular injection of WGA-HRP; disruption of substimulation was similar to that seen in animals treated with L-NaORg alone, suggesting that LGN blockade-induced changes in retinogeniculate projections occur independently of changes in MAP. Substimulation in the LGN relies at least in part on the neuronal form of NO. Neuronal NO immunohistochemistry is similar to NOS and D-arginine phosphodiesterase in its transient expression during retinogeniculate innervation. In addition, ONOFF substimulation was blocked by blockade of the neuronal form of NO from P10 to P25 with 7-nitroindazole, which appears not to produce hyperthermia. While systemic changes in blood pressure are not involved in the segregation of substimulation, further experiments are necessary to examine the role of local changes in cerebral blood flow produced by NO release. Supported by EY10702.

304.4
ROLE OF SPONTANEOUS RETINAL ACTIVITY IN REORGANIZATION OF RETINOGENICULATE CONNECTIONS DURING DEVELOPMENT. P.M. Cook, G. Zufi, and A.S. Ramon, Dept. of Anatomy, University of Virginia, Richmond, VA 23298 and Dept. of Psychology, Univ. of Loughborigh, Loughborigh AB TK13M. The adult form and function of the lateral geniculate nucleus (LGN) arise after extensive modifications in circuitry that include segregation of axonal inputs from the two eyes into eye-specific layers. Previous studies have shown that transient inhibition of tetrodotoxin (TTX) in fetal cats prevents segregation (Shatz and Stryker, 1989), suggesting that spontaneous action potential activity contributes to remodeling of the retinogeniculate pathway. Now we have shown that transient inhibition of retinogeniculate projections during the first 2 week period of postnatal life. After a 2 week inhibition, intraocular injection of horseradish peroxidase (HRP) revealed the projection pattern contralateral and ipsilateral to the HRP injected eye. Eye-specific segregation of retinogeniculate projections was observed at postnatal day 14 in controls as well as in fetuses that received mannitol (3%) or bacitracin (1%) injection of TTX. However, segregation was aberrant in the noninjected eye. In a sample of 5 newborn cats, the noninjected eye occupied an approximately 50% larger volume of both LGNs than those from the TTX-treated eye. The total size of the LGN, however, remained unaltered and this suggests that the normal eye invaded normally normally occupied by the TTX treated eye. Moreover, a low density of axons from the contralateral eye were observed to terminate in layer 1 of the LGN, which normally receives only ipsilateral eye input. These results suggest that binocular competition modulates retinogeniculate projections. Additionally, they are consistent with the hypothesis that spontaneous retinal activity may fine-tune segregation of retinal afferents into eye-specific layers. (Supported by NSF BNS-9219193)

304.5
POSTNATAL DEVELOPMENT OF SYNAPTIC PLASTICITY IN THE KAT SUPERIOR COLLICULUS: LONG-TERM DEPRESSION IN THE SUPERFICIAL LAYERS. R.J. Cook, R.S. Le and R.R. Price. Anatomy Department and Neuroscience Center, Loyola University Medical Center, Maywood, IL 60153. We have been studying the development of the synaptic circuitry in the rat superior colliculus (SC) between ages P1 and P30. Using in vitro brainstem preparation, we have measured electrophysiological parameters of Iona and L-Type calcium channels and postsynaptic potentials, from the superficial layers of SC. We have also used immunocytochemistry and NADPH-diaphorase to map the development of various intrinsic and extrinsic properties involved in synaptic transmission (i.e. NMDA, and NO). Field potentials and post-synaptic potentials were measured following stimulation of optic tract (OT) fibers. Whole cell recordings showed EPSPs with two components, an early EPSP and a late EPSP. The early EPSP was blocked by APV, suggesting that it was mediated by the NMDA receptor. NMDA receptor-mediated LTP was also tested by stimulating the superficial layers at P1. From P3 some neurons also showed an IPSP following the EPSP. The IPSP could be blocked by bicuculline, an antagonist of GABA, receptor activation. LTP was also induced by P10, P20 (20 s, submaximal intensity) induced a long-term (29 min) depression (LTD) of the field potential and the IPSP. Neither APV (10 μM or 50 μM) nor bicuculline (10 μM) could prevent LTD induction in the OT. Interestingly, it is independent of NMDA or GABA receptor activation. Nitrendipine (5 μM), an L-Type Ca2+ channel blocker that blocks induction of LTD in the hippocampus, also failed to prevent LTD induction in the SC. The magnitude of the LTD decreased steadily from P1 to P8 and at P8/10 tonic OT stimulation induced a decrease in depression and long-term potentiation (LTP). Consistent with the idea that NOS promotes LTD, significant NOS expression was first observed in cells of the superficial layers at P5. Supported by DOD cooperative agreement DAMD 17-93-1-V-6131, NIH grant EY102973, and the LSUMC Neuroscience Center.

304.6
INHIBITION OF NITRIC OXIDE SYNTHASE FAILS TO DISRUPT THE DEVELOPMENT OF CHLORINICOLOR SIM PATECHES IN THE KAT SUPERIOR COLLICULUS. C.A. Schacter, R.J. Cook and R.R. Price. Dept. of Anatomy and the Neuroscience Center, Loyola University Medical Center, New Orleans, LA 70112. Nitric oxide (NO) has been proposed to be a retrograde messenger involved in synaptic refinement during development. The patch-cluster system in the intermediate gray layer (IGL) of the superior colliculus (SC) can be involved in the development of synapses (i.e. NMDAR1, and NOS). Field potentials and post-synaptic potentials were measured following stimulation of optic tract (OT) fibers. Whole cell recordings showed EPSPs with two components, an early EPSP and a late EPSP. The early EPSP was blocked by APV, suggesting that it was mediated by the NMDA receptor. NMDA receptor-mediated LTP was also induced by P10, P20 (20 s, submaximal intensity) induced a long-term (29 min) depression (LTD) of the field potential and the IPSP. Neither APV (10 μM or 50 μM) nor bicuculline (10 μM) could prevent LTD induction in the OT. Interestingly, it is independent of NMDA or GABA receptor activation. Nitrendipine (5 μM), an L-Type Ca2+ channel blocker that blocks induction of LTD in the hippocampus, also failed to prevent LTD induction in the SC. The magnitude of the LTD decreased steadily from P1 to P8 and at P8/10 tonic OT stimulation induced a decrease in depression and long-term potentiation (LTP). Consistent with the idea that NOS promotes LTD, significant NOS expression was first observed in cells of the superficial layers at P5. Supported by DOD cooperative agreement DAMD 17-93-V-6131, NIH grant EY102973, and the LSUMC Neuroscience Center.

304.7
ROLE OF RETINAL ACTIVITY IN MATURATION OF ELECTROPHYSIOLOGICAL MEMBRANE PROPERTIES AND SYNAPTIC RESPONSES OF FERRET LGN. A.S. Ramon and G. Prosky, Dept. of Anatomy, Virginia Commonwealth University, Richmond, VA 23298 and Dept. of Psychology, University of Loughborigh, AB TK13M. The mature form and function of the lateral geniculate nucleus (LGN) arise after extensive modifications in circuitry that, in the ferret, occur during the first postnatal month. We have shown that electrophysiological membrane properties and synaptic responses change markedly during this critical period of development (J. Neurosci. 14:2089, 1994; J. Neurosci. 14:2089, 1994; J. Neurosci. 14:2089, 1994; J. Neurosci. 14:2089, 1994). These changes, which appear to be coordinated to facilitate circuit remodeling, include potassium channels, calcium channels, and dendritic properties. The functional properties of amino acid receptors support this. Beta-2 adrenoceptors promote the development of the retinogeniculate pathway, which is dependent on the presence of retinal ganglion cells. Whole-cell recordings in the LGN slice preparation revealed that maturation of low-threshold calcium spikes and hyperpolarization-activated currents was outlasted by a 3 to 4 week-bivalent inhibition of tetrodotoxin. In contrast, synaptic properties were markedly affected. Transferred animals at P40 were found to display long duration NMDA-EPSCs that resembled those present in normal newborn animals; whereas the shorter-duration EPSCs seen in normal animals at similar age (9-30 days). In these animals, application of ifenprodil, which binds with higher affinity to heteromeric receptors containing the NR-2B subunit, blocked not only NMDA-EPSCs in newborn and TTX-treated animals at P40 in normal animals (9-30 days), but also exhibited a developmental switch in the subunit composition of the NMDA receptor is prevented by intravenous TTX. In conclusion, retinal activity may regulate development and change in synaptic properties of LGN neurons (NSF BNS-9421980).

304.8
CHLORINICOLOR PROCESSES IN XENOPUS TECTUM. M.I. Trima*, L. Pick, R. Lima & S.B. Ulin, Dept. of Physiology & Biophysics, SUNY, Buffalo, NY 14214. The ipsilateral visual input to the tectum comes into register with the contralateral map during development. The major transmitter for the contralateral (reinocuclal) input is glutamate, and the major transmitter for ipsilateral input, relayed via the nucleus isomusini, is probably acetylcholine. The role of NMDA receptors in the activity dependent process of organization of the ipsilateral map is demonstrated by the ability of NMDA receptor blockers to prevent the matching of the ipsilateral map to the contralateral map during the critical period, but little is known about the role of acetylcholine. Immunohistochemistry indicates that acetylcholine receptors are located in the layers of the tectum that receive binocular inputs. While eye enucleation indicates that most of those receptors are located on reinocuclal axons. Recent evidence suggests that activity is determined by the presence of nicotinic receptors in the tectum. Nicotinic receptors are located on cells and dendrites located appropriately to receive isomusini input. Calcium imaging using Fura-2 in tectal slices demonstrates no measurable response to nicotinic agonists alone, but shows significant synergism when nicotine or acetylcholine are applied with NMDA. In contrast, muscarinic agonists do not appear to interact with NMDA. These results suggest that activity of isomusini axons can significantly modulate the effects of glutamate released from reinocuclal inputs. Supported by USPHS Grant EY-10690 to M.I. Trima and S.B. Ulin.
MODULATION OF INTRACELLULAR CALCIUM DYNAMICS IN RAT HIPPOCAMPAL NEURONS BY PLATELET-ACTIVATING FACTOR. Mark A. DeCoster and Nicolas G. Bazan, Louisiana State University Medical Center, Neuroscience Center, New Orleans, LA 70112.

Platelet-activating factor (PAF) is a potent lipid mediator of several cellular responses. PAF is able to transiently increase intracellular calcium ([Ca^{2+}]) in several neuronal cell systems, mostly in cell lines and pituitary cells. PAF addition mobilizes [Ca^{2+}] in hippocampal neurons; however, this response was found in only 7.7% of the cells analyzed (Bito et al., Neuron, 9:285, 1992). In the present study confocal microscopy was used to monitor the possible modulatory effect of PAF on calcium dynamics in rat primary hippocampal neurons. In short-term experiments, [Ca^{2+}] was simultaneously monitored in multiple neurons; basal calcium oscillatory activity was observed in most cells. When 4 μM methylcarbamyl PAF (mcPAF) was added to these cultures, the average [Ca^{2+}] was increased slightly in cells and the range of calcium oscillations was increased approximately 3-fold. These changes were not observed when only bovine serum albumin (the vehicle for PAF) was added. However, in agreement with the results of Bito et al. (1992), we found that not all cells responded to PAF addition: in our studies, about 25% of the cells responded. Glutamate (80 μM) was added at the end of each scan as a positive control for neuronal viability. To investigate the possibility of longer term effects of PAF, we pretreated hippocampal cultures overnight with PAF, mcPAF, lysoPAF, or BSA and monitored [Ca^{2+}] changes induced by glutamate on the subsequent day. In general, we found that PAF (200-400 nM) and mcPAF (2-4 μM) pretreatment reduced the [Ca^{2+}] changes induced by low concentrations of glutamate (100-500 nM) when compared with cells pretreated with lysoPAF (2-4 μM) or the vehicle. In multiple experiments, when pretreatment with PAF was used, the [Ca^{2+}] changes caused by glutamate were reduced from sustained to transient increases. In two cases, the [Ca^{2+}] increases in response to 500 nM glutamate were completely reversed by PAF pretreatment. These inhibitory effects by PAF were overcome by higher glutamate (5 μM) concentrations. In conclusion, PAF induces calcium oscillations in hippocampal neurons. In addition, based upon the previous findings showing that PAF-induces glutamate release (Clark et al., Neuron 9:1211-6, 1992; Kato et al., Nature 267:175-9, 1994), our present results suggest that sustained PAF-induced glutamate release may produce glutamate receptor desensitization. (Supported by DAMD-17-93-V-3013 and NS 23002)

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NEUROTOXICITY AND MODULATION OF CALCIUM DYNAMICS IN RAT CORTICAL NEURONS BY PHOSPHOLIPASES. Miriam Kolko, Mark A. DeCoster and Nicolas G. Bazan. Neuroscience Center, LSU Medical Center, New Orleans, LA 70112-2234.

Phospholipases A2 (PLA2s) are important modulators of neuronal function. Ischemia and seizures promote an enhanced activation of PLA2 in brain that, in turn, leads to the accumulation of free arachidonic acid (Biochim. Biophys. Acta 218:1-10, 1970) and platelet-activating factor (Biochim. Biophys. Acta 963:375-383, 1988). Nonpancreatic secretory PLA2s (sPLA2s) participate in the inflammatory injury response. A receptor for sPLA2s has been cloned from muscle (Lambeau, et al., JBC. 269:1575, 1994). Two sPLA2s from snake venom (OS2 and OS3), bind tightly to this receptor. OS2 and sPLA2 from bee venom bind avidly to brain membranes, while OS3 does not. The present study evaluated the neurotoxic potential of these sPLA2s in rat cortical neurons in vitro using lactate dehydrogenase (LDH) release as the toxicity assay and the excitotoxic amino acid glutamate (80μM) as a control. Bee venom and OS2 (0.01-10 μg/mL) showed dose dependent toxicity while OS3 did not cause cell death. The toxicity experiments were combined with studies investigating the effect of the sPLA2s on intracellular free calcium concentration ([Ca2+]). The fluorescent calcium indicator fluo-3 was used with a confocal microscope to measure real-time calcium dynamics in these neurons. We observed basal oscillations in [Ca2+], in the imaged cultures with a 2.3 mM extracellular calcium concentration. Bee venom and OS2 dose-dependently (0.5-10 μg/mL) altered ([Ca2+]i) dynamics, while OS3 had no effect. Both bee venom and OS2 (0.5 - 10 μg/mL) obliterated calcium oscillations, and also decreased [Ca2+], to below baseline levels. Subsequent addition of glutamate to the cultures showed an immediate rise in the [Ca2+]. We did not see any calcium modulations in the cultures exposed to sPLA2 concentrations below 0.5 μg/mL, even though the sPLA2s were toxic to about 0.025 μg/mL. Using a cytofluorimetric assay we found that bee venom and OS3 destroy neuronal cell bodies and processes at concentrations as low as 0.025 μg/mL and are generally cytotoxic to neurons and glia at the high concentrations. In contrast, 80 μM glutamate damages only neurons. Since the neurotoxicity of bee venom and OS3 do not appear to correlate with calcium modulations at lower concentrations, a calcium-independent toxicity may be occurring at these low PLA2 concentrations. Furthermore, submaximally toxic concentrations of bee venom combined with 80 μM glutamate demonstrated higher toxicity levels than the two compounds added separately. Thus, LDH release from these studies resulted in more than additive activity, indicating that glutamate and bee venom are having a synergistic effect. Since OS3 is not affecting neuronal damage or the calcium dynamics in our experiments, these data support the observed binding properties of sPLA2s in muscle and brain tissue, and provide evidence for modulatory roles of PLA2 s in neuronal signal transduction. (Supported by DAMD-17-93-V-3013 and NS 23002).

NOTES:
248.3

EXPRESSION OF GLUTAMATE RECEPTORS OF RAT BERRERAL GRANULE CELLS DEPENDS UPON THE VOLUME OF CULTURE MEDIUM.

C. Calissano*, M. Puttfarcken, T. Cottrell, and P. Calissano. Dept. of Experimental Medicine, II University of Rome, and Institute of Neurobiology, CNR, Rome, Italy.

In a preliminary series of studies we have found that the response of cerebellar granule cells cultured in vitro for 8 days to a cytotoxic glutamate concentration of 100 μM was dependent upon the volume of medium in which neurons are grown. When cells are cultured in low volume (LV) (1.2x10^6 cells/1.2 ml/17 mm dishes) the glutamate EC50 for death of cultured cells is 10-20 μM. In contrast, when higher cultures are prepared in a high volume (HV) of 4.0 ml the cell death following glutamate treatment is reduced to 20-30%. Addition of a conditioned medium from LV cultures (containing 10% of HV cultures) markedly increases their response to the toxic glutamate treatment. In order to investigate this volume dependency and the action of CM on glutamate sensitivity, we have measured sodium currents, the currents evoked by AMPA/kainate and NMDA as well as the mRNA (1A, 1B, 2A, 2B, 2C) and the protein subunits forming the AMPA/kainate (Glur1, Glur2, Glur3) and NMDA (NRMDA1, NRMDA2B) receptors. We found that the glutamate resistant phenotype ensuing in HV culture conditions is accompanied by, and probably causally connected with, a lowered functional and physical expression of both well-functioning sodium channels as well as kainate and NMDA receptors.

We hypothesize that such volume dependency to the cytotoxic action of glutamate is connected with the extent of production and breakdown of a substance operationally defined as glutamate sensing activity or GSA.

248.4

ANTISENSE OXONUCLEOTIDES TO CALPAIN 1 mRNA PROTECT CULTURED HIPPOCAMPAL SLICES FROM NMDA-INDUCED PATHOPHYSIOLOGY.


We have shown that hippocampal slices exposed to NMDA (30 μM) develop a functional pathology characterized by a decrease in synaptic transmission, a loss of long term potentiation, and a decrease in the expression of the NMDA receptor subunits. Our hypothesis is that this pathology is mediated by an increase in calpain activity. To test this hypothesis, we developed an experimental model in which the activity of calpain is reduced by the treatment of hippocampal slices with an antisense oligonucleotide (5'-GAC AAC CCA TCG TCC GAG AGG A-3'). This oligonucleotide was designed to hybridize with the 3'-untranslated region of the calpain I mRNA, which is a major form of calpain I expressed in the hippocampus. The results of this study show that the treatment of hippocampal slices with the antisense oligonucleotide decreases the activity of calpain I, as measured by the cleavage of the substrate Suc-LLVY-AMC. This decrease in calpain activity results in an increase in the survival of hippocampal slices exposed to NMDA (30 μM) for 20 min. These results suggest that calpain activity is a major factor in the development of NMDA-induced pathology and that antisense techniques may be useful in the treatment of neurodegenerative diseases.

248.5

OVEREXPRESSION OF RECOMBINANT HUMAN CALPASIN D1, E, and F, AND ITS IMMUNOPROTECTIVE EFFECTS AGAINST EXCITOTOXICITY.

Oliver D. A. Evans, R. K. Gloor, and M. P. Goldman. Center for the Study of Nervous System Injury and Dept. of Neurology, Washington University School of Medicine, St. Louis, MO 63110.

Calpains are a family of cysteine proteases that play a critical role in cellular signaling and homeostasis. They are activated by intracellular calcium and are involved in a wide range of biological processes, including cell death and survival. In this study, we investigated the effects of overexpression of recombinant human calpains D1, E, and F on cellular survival and neurotransmission in cortical neurons. Our results suggest that overexpression of calpains D1, E, and F can protect neurons against excitotoxic injury induced by glutamate or kainate.

248.6

ROLE OF CALPAIN IN THE DEVELOPMENT AND RECOVERY OF EXCITATORY DENDRITIC INJURY IN VITRO.

B. D. Fedder, S. M. Goldman, and M. P. Goldman. Center for the Study of Nervous System Injury and Dept. of Neurology, Washington University School of Medicine, St. Louis, MO 63110.

In this study, we investigated the role of calpains in the development and recovery of excitotoxic dendritic injury in vitro. We found that overexpression of calpains D1 and E can protect neurons against excitotoxic injury induced by glutamate or kainate.

248.7

NMDA INDUCES CALPAIN-MEDIATED PROTEOLYSIS OF MICROTUBULE-ASSOCIATED PROTEIN 2 AND SYNAPTIC DYNAMICS IN CULTURED HIPPOCAMPAL NEURONS.

M. Mosqueda, B. T. Fadda, M. P. Goldman, and C. W. Xing. Center for the Study of Nervous System Injury and Dept. of Neurology, Washington University School of Medicine, St. Louis, MO 63110.

In this study, we investigated the role of calpains in the development and recovery of excitotoxic dendritic injury in vitro. We found that overexpression of calpains D1 and E can protect neurons against excitotoxic injury induced by glutamate or kainate.

248.8

EFFECT OF SECRETORY PHOSPHOLIPASES A1 AND GLUTAMATE ON VARIABILITY OF RAT CORTEX NEURONS AND CALCIUM DYNAMICS IN HIPPOCAMPAL NEURONS.


In this study, we investigated the role of calpains in the development and recovery of excitotoxic dendritic injury in vitro. We found that overexpression of calpains D1 and E can protect neurons against excitotoxic injury induced by glutamate or kainate.

248.9

SOCIETY FOR NEUROSCIENCE, VOLUME 21, 1995
444.1 INTERCELLULAR CALCICUM WAVES PROPAGATED VIA GAP JUNCTIONS IN NEURONS A.C. Charles and R.F. Tvedale. Dept Of Neurology, UCLA, Los Angeles, CA 90024 and Addiction Research Foundation and Dept Of Pharmacology, Univ Of Toronto, Canada M5S 2S1.

Spontaneous waves of increased intracellular calcium concentration were rapidly propagated over groups of primary mouse cortical neurons and immortalized hypothalamic (GT-1-1) neurons in culture. Ca²⁺ waves were propagated at a rate of 100-200 µm/sec over entire cultures of Ca²⁺-sensitive cells. Ca²⁺ waves were abolished by the removal of extracellular calcium and by TTX. Similar intercellular Ca²⁺ waves were induced by mechanical stimulation of a single cell. GT-1-1 neurons showed fluorescence recovery after photobleaching of a single cell, and intercellular Ca²⁺ waves were abolished by the gap junction blocker octanol. As a contrast, a different clone of the GT-1 neurons (GT-1-7) showed frequent spontaneous Ca²⁺ oscillations but no intercellular Ca²⁺ waves, no intercellular communication of the response to mechanical stimulation, and no fluorescence recovery after photobleaching of a single cell. These results show that neurons are capable of extensive Ca²⁺ signaling via gap junctions, and suggest that Ca²⁺ waves are the gap junction protein that enables intercellular Ca²⁺ signaling in GT-1 neurons. Intercellular Ca²⁺ waves in cultured neurons may represent a model for gap-junctional signaling between neurons in the developing nervous system, and between subsets of neurons in the adult brain.

444.3 ASSOCIATION OF TYPE I AND TYPE III INSULIN-1, 4,5 TRIOPHOSPHATE RECEPTORS.

P.C. Noda et al. Laboratoire de Neurobiologie, Johns Hopkins University, School of Medicine, 720 Rutland Ave., Ross 615, Baltimore, MD 21205. 1. Department of Physiology, University of North Carolina.

The Insulin 1,4,5-trisphosphate receptor (IP3R) is an intracellular calcium channel involved in coupling cell membrane receptors to calcium signal transduction pathways within the cell. The IP3R is believed to form a tetrameric structure to produce the calcium channel in endoplasmic reticulum membranes. Several isoforms (I, I1, I11, II, III) of IP3Rs have been identified which are coded by separate genes, and are expressed in many tissues with differing patterns of cellular expression. We have generated specific affinity purified polyclonal anti-peptide antibodies to each of the three isoforms. Western Blot analysis of INS-1E and A10 cells shows high levels of endogenously expressed type I and type III IP3R, but undetectable levels of type II IP3R. Immunoprecipitation experiments performed with cells from the three I type specific antibodies and Western Blotting with the type specific antibody. Both experiments yielded a band at 260 kDa, the appropriate size of both the type I and type III IP3R.

Immunocytochemistry performed on these cells with either antibody demonstrated similar ER staining patterns. The type III IP3R was absent from the secretory granules of AT20 cells. These data indicate that type I and type III IP3Rs can associate into a molecular complex.

444.5 CLONING AND SEQUENCING OF AN IP₃-RECEPTOR cDNA FROM LOBSTER OLFATORY ORGAN.

A.M. Maguire a, J.W. Farnes a, Brian A. Techie and B. M. Greenstein. Laboratory 1 and Dept. of Neuroscience and 2. Biology, Univ. of Florida, St. Augustine. FL. 32086.

Several lines of evidence suggest that Insulin 1,4,5-trisphosphite (IP₃)-receptors (IP₃R) occur in lobster olfactory neuronal and neuroendocrine cells, but it is not known about the structural similarities of these plasma membrane IP₃R to the better known intracellular IP₃Rs. IP₃, directly gates two types of ion channels in the plasma membrane of target epithelial receptor neurons (ORNs): Those changes functionally similar to IP₃R localized to endoplasmic reticulum (ER) and nuclear membranes in vertebrates (Fadool & Ache, Neurons, 9, 997; Han & Ache, PNAS, 91: 6204). We have characterized the molecular and functional properties of IP₃R from lobster olfactory organ neurons (ORNs) that are IP₃R transmembrane proteins, which may represent pre-synaptic membranes. Our findings suggest that IP₃Rs may play a role in modulating synaptic transmission and neuronal plasticity.

444.6 PLATELET-ACTIVATING FACTOR INDUCED INTRACELLULAR CALCIUM OSCILLATIONS IN RAT IPHOCAMPAL NEURONS. M.A. DiCicco, H.F. F. Bassani, a, and N. O. Bassi. LSU Medical Center, Neuroscience Center, New Orleans, LA 70112-0234.

As has been previously shown, we have found using confocal microscopy and fluorescent calcium indicators, that intracellular calcium concentration (Ca²⁺) oscillates spontaneously in rat hippocampal neurons in vivo. While addition of glutamate (GLU) to these hippocampal cultures causes distinct Ca²⁺ changes ranging from transient, single spikes (100-500 nM GLU) to sustained increases (10-20 µM GLU), glutamate is never observed to induce Ca²⁺ oscillations. We have investigated the ability of the potent lipid mediator platelet activating factor (PAF) to affect Ca²⁺ dynamics in hippocampal neurons. When 1 µM methacharylam PAF (mPAF) was added to the hippocampal neurons, the average Ca²⁺ increase was significantly in cells. Furthermore, the fluorescence values after mPAF additions was Bl8-fold higher than before additions, indicating an increase in oscillatory [Ca²⁺]. However, no oscillatory Ca²⁺ changes induced by mPAF were seen. Non-oscillatory Ca²⁺ changes were not seen in response to mPAF. In contrast, long-term effects of PAF treatment on hippocampal cultures appear to affect the majority of cells. Overnight treatment with PAF (200-400 µM) and mPAF (2-4 µM) reduced the neuronal Ca²⁺ changes induced by GLU the next day when compared with cells pretreated with hysPAF (2-4 µM) or the vehicle alone. In two cases, the Ca²⁺ increases in response to 500 nM GLU were completely inhibited by PAF pretreatment. Since PAF has been shown to enhance hippocampal excitatory synaptic transmission (Clark et al., Neuron, 9:1211, 1992), we postulate that induction of [Ca²⁺] oscillations by PAF may be an early signal of GLU release, resulting in GLU receptor desensitization. (Supported by SAR-16735-V-0131, 1992).
1476 MODULATION OF CALCIUM, NEUROTOXICITY AND ARACHIDONIC ACID RELEASE BY PHOSPHOLIPASE A2 TYPE II AND GLUTAMATE IN VITRO: M. Kolko, E.B. Rogers-deTurco, M.A. DeCoster, and N.G. Basacz, LSU Eye Center, 1553 T.C. Jeflin, New Orleans, LA.

Secretory phospholipase A2 Type II (sPLA2) may modulate neural function, under both physiological and pathological conditions. This 14 kDa enzyme, present in animals and in human platelets, is released upon depolarization or neurotransmitter stimulation. Moreover, ischemia induces sPLA2 gene expression in rat brain. We evaluated the effect of sPLA2 from bee venom (BV) alone and with glial cell line-derived neurotrophic factor (GDNF) on rat glial cells. BV (1000 ng/ml), alone, had no effect, nor did it affect basal oscillations in [Ca2+]i. However, BV dose dependently (0.025-10 μM) caused neurotoxicity, altered [Ca2+]i dynamics, and stimulated [NADPH]A release. GDNF (50 ng/ml), alone, did not induce neurotoxicity, selected L-mahacholinergic (L-mah) neurons, induced [Ca2+]i oscillations, but increased basal oscillations and significantly decreased BV-induced neurotoxicity. In contrast, the sustained [Ca2+]i response induced by BV, dose-dependently (0.25-5 μM), induced a transient increase in [Ca2+]i, followed by decreased basal oscillations and a significant decrease in [Ca2+]i. These results indicate that calcium-independent toxicity may occur at low sPLA2 concentrations and provide evidence for modulatory roles of sPLA2 in neural signal transduction. Supported by DAMO-17-19-5-V-2013.


Previously we reported that tumor-promoting phorbol esters stimulate phospholipase D (PLD) independently of protein kinase C activities in bovine lymph node lymphocytes. (Cao et al., Biochem. Biophys. Res. Commun. 171, 955-962, 1990; JIBID 217, 807-915, 1995). In the present study, we examined the effects of phorbol esters (PGE2), DP, D3, and H2 on PLD activity as measured by [1-14C]arachidonic acid incorporation into phosphatidylethanolamine (PE) in bovine lymphocytes. All PLD stimulations were measured in a dose-dependent manner in the concentration range between 0.1-10 μM. Prostaglandin E2 had the maximum stimulatory effect in the order of PGE2 > PGF2α > PGE2 > PGD2 > PC50. The PG-stimulated formation of PE was enhanced by staurosporine, a PKC inhibitor. When both were included, the effect of PGE2 and 12-O-tetradecanoylphorbol-13-acetate (TPA) on the PLD activation was additive. Furthermore, NaF, a G-protein activator, stimulated the PLD formation and this stimulation was enhanced by staurosporine. Interestingly, the stimulatory effects of NaF and GTPγS were not additive; however the formation of PE by NaF and TPA was additive. These results suggest that a similar signaling pathway of PLD activation. PLD activation and the stimulation by PG through PKC and PLA2 in lymphocytes may involve both G-protein-dependent and G-protein-independent signaling pathways. (Supported through NIH grant numbers HL83425 and AI-06547.)

Adequate vectors encoding either isoform of human glutamate decarboxylase (GAD67 or GAD65) under the control of the CMV promoter were constructed. Expression of GAD65 or GAD67 was verified by Western blot analysis of rat pituitary cells 48 hours after adenoviral-mediated (AdGAD) gene transfer. GAD65 expression was >90% of total GAD65, while GAD67 expression was 70% of total GAD67. Extracellular [GABA]a increased >25 fold (79 µM versus 3 µM) in neurons 9 days after AdGAD67 gene transfer. Intracellular [GABA]a also increased in these neurons (20 µM versus 0 µM) and remained as such from 3-9 days after vector addition. Intracellular glutamate concentration decreased almost 50% in neurons 3 days after AdGAD67 gene transfer, but returned to baseline (50-60 µM) by 9 days after vector addition. Similar [GABA]a and [glutamate] were observed in neurons following AdGAD65 gene transfer and in non-neuronal cell lines (GABA) and [glutamate] were determined by HPLC using a Fico-Tag column (Waters) for free amino acid analysis. These data suggest that glutamatergic neurons can produce and release GABA following AdGAD gene transfer in vitro. Conversion of glutamatergic neurons to those expressing both glutamate and GABA could yield therapies in a variety of neurodegenerative disorders.

INHIBITION OF MOSCARIC RECEPTOR BINDING BY PERNOSIDE, ORTHOVANADATE, METAVANDATE, AND AN ENDogenous INHIBITOR FROM ALZHEIMER'S BRAIN. H. Frey et al. have previously reported (Brain Res. 655, 153-160, 1994) that an endogenous inhibitor (<0.50% dilution) of 4,5GMP-antibody (4,5GMPA) is elevated in brain tissue from patients with Alzheimer's disease (AD). Further research with this inhibitor is in progress. Similar inhibition is irreversible and mediated by the thyl radical of glutathione which forms in the presence of the inhibitor. We now report the results of research on the effect of metavanadate (known to generate thyl radicals from glutathione), orthovanadate, and pervanadate on 4,5GMP-binding to the 4,5GMPAR. All three vanadium compounds inhibited binding in the presence of glutathione, with the order of decreasing potency and the concentration required for 50% inhibition (IC50) being: pervanadate (95 µM) > orthovanadate (250 µM) > metavanadate (452 µM). Omission of glutathione decreased the inhibition of the vanadium compounds from 2 to 6 fold. In contrast to our results with the endogenous AD inhibitor, preincubating the vanadium compounds with the 4,5GMPAR in the absence of glutathione at 37°C for 1 hour markedly decreased the IC50 values as follows:

pervanadate (13 µM) > orthovanadate (48 µM) > metavanadate (119 µM). Similarities of the vanadium compounds with the endogenous AD inhibitor were also noted, with the inhibition of each being blocked by Trolox, EDTA and Mn2+. Further studies are needed to identify the endogenous AD inhibitor. (Supported by Ramsey Foundation grant #N85S).


The service personnel during the Persian Gulf War were exposed to pesticides such as the insect repellent, DEET (N,N-diethyl-m-toluamide) and the herbicide, chlorpyrifos (0,0-diethyl O,3,5,6-tetrahydrophosphoronoxyphenoxyisopropyl). We propose to individual or component of benzene 5 days/week for 2 months to 5 mg PB Agarpl in water, ACH Agar plate, new, and 10 mg chlorpyrifos Agar plate in corn oil, etc. Binary treatments produced greater neurotoxicity than death induced by individual exposures. Neurotoxicity was further enhanced following exposure to these chemicals. PB decreased plasma benzo[a]pyrene (BaP) activity to 1% of control compared to 5% and 4% from chlorpyrifos and DEET. However, benzene exposure followed by chlorpyrifos treatment groups compared to individual dose groups. In contrast, only chlorpyrifos alone or in combination with other compounds produced a significant indication of brain acetylcholinesterase activity. It was significantly greater in ACH exposed given chlorpyrifos with PB or DEET or with both chemicals. We hypothesized that these components can cause synaptic neurotransmitter changes in the liver and brain that can contribute to the synergistic effects produced by PB. A study of the concentration of the lipoprotein (LDL)DEET and chlorpyrifos in circulation and their availability to the central nervous system. This increase in the effectiveness of these chemicals in the nervous system could then reach levels equivalent to lethal doses of individual components associated with neurologic deficits. (Funding provided by the Pernet Foundation.)

MODULATION OF CHRONIC NEUROTOXICITY BY COMBINED KAINATE AND GLUTAMATE TREATMENT OF HIPPOCAMPAL NEURONS. M. Clea, M.A. DeCoster, and N.G. Banaz. LSU Medical Center, Neuroscience Center, LA 70112-2234.

Acute treatment with excitatory amino acids, (EAA)s such as kainate (KA) and glutamate (Glu) cause neurotoxicity characterized by rapid second messengers such as intracellular calcium; however, cell death in this model is delayed until approximately 16-24 hours. While these mechanisms of EAA neurotoxicity are understood for acute treatment, less is known about the effects of chronic treatment with these compounds. We therefore incubated primary rat hippocampal neurons overnight with KA and G (30 µM to 3 mM). Neuronal injury was determined by lactic dehydrogenase (LDH) release. Dose-response curves showed different toxicity patterns for KA and G treatments. At lower concentrations KA was less toxic than Glu, but at concentrations higher than 300 µM, KA toxicity was 5 to 7 times of the control levels. KA attained a toxicity level at 2 to 3 times that of the baseline. Surprisingly, the toxicity of equimolar KA-G (300 µM 300 = 300 µM) treatment was similar in magnitude to that induced by glutamate alone. Thus, the expected additive effects of toxic equimolar KA-G treatment was not obtained. One possible explanation for these results is the reported neuroprotective effect of activation of the metabolic receptor by G; this action could mask the expected additive toxicity of KA-G treatment. A second possibility would involve the differential uptake of EAA s by astrocytes, which account for 10-15% of the cells in our culture. The role of both potential mechanisms will be investigated. (Supported by DAMD-17-93-V-0313.)


We demonstrated that in PC12 cells, the mechanism of stimulated acetylcholine release, inhibition by botulimum toxin A (BoNTa) was via interference of phospholipase A (PLA) stimulated arachidonic acid (AA) release from cell membranes as well as via a PLA inhibitor, a second specific PLA inhibitor. (8-hydroxy-2,4-dihydroxybenzyl 7,7-dimethyl-7,7-dihydroxybenzoic acid (E220) abolished [14C]AA and [14C]cholesterol release due to metals or Ca2+ but the Ca2+-independent PLA inhibitor, (2,4-dimethyl-2,5-dihydroxysteroid-3-1-naphthylamino)-2,4,6-pyren-3-one (HEL659) had no effect. The effects of Nias plus K+ were blocked by either E220 or the K+ type Ca2+ channel blocker; aconitoxin, but not the L-type Ca2+ channel blocker, nifedipine, indicating that these effects were dependent on Ca2+ influx via the neuronal type voltage-sensitive Ca2+ channels. BoNTA inhibition of [14C]AA and [14C]choline release was fully prevented by Nias plus 50-100 µM K+. These results indicate that Nias prevents the BoNTA effects via Ca2+ influx, PLA activation, and AA release; and provides additional support for our proposal of Nias in stimulated ACh release and its inhibition by BoNTA.


We have shown that suxamethonium (STP) induced lethality can be reversed by treatment with 4-aminopyridine (4-AP) if given at the time of respiratory arrest. The purpose of this study was to examine 4-AP's therapeutic window and to determine if its efficacy can be further enhanced when used at the time of respiratory arrest. The groups of unanesthetized guinea pigs were used in this study. These animals were chronically instrumented for the measurement of diaphragm EMG, esophageal EMG, Lead II ECG and B Cleo. All animals were given 100 µg STX and 5 µg Agarpl in water, 300 mg DEET Agarpl in water, etc. and 10 mg chlorpyrifos Agarpl in corn oil, etc. Binary treatments produced greater neurotoxicity than death induced by individual exposures. Neurotoxicity was further enhanced following exposure to these chemicals. PB decreased plasma benzo[a]pyrene (BaP) activity to 1% of control compared to 5% and 4% from chlorpyrifos and DEET. However, benzene exposure followed by chlorpyrifos treatment groups compared to individual dose groups. In contrast, only chlorpyrifos alone or in combination with other compounds produced a significant indication of brain acetylcholinesterase activity. It was significantly greater in ACH exposed given chlorpyrifos with PB or DEET or with both chemicals. We hypothesized that these chemicals can cause synaptic neurotransmitter changes in the liver and brain that can contribute to the synergistic effects produced by PB. A study of the concentration of the lipoprotein (LDL)DEET and chlorpyrifos in circulation and their availability to the central nervous system. This increase in the effectiveness of these chemicals in the nervous system could then reach levels equivalent to lethal doses of individual components associated with neurologic deficits. (Funding provided by the Pernet Foundation.)
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PAF INDUCES CYCLOOXYGENASE (COX-2) GENE EXPRESSION IN THE CORNEAL EPITHELIUM PARTIALLY BY A RECEPTOR-MEDIATED CALCIUM INFLUX. Y. Tao, H.E.P. Bazan, M.A. DeCoster, N.G. Bazan, L.S. Eye Center and Neuroscience Center, New Orleans, LA 70112

This study investigated the role and source of Ca²⁺ in the signaling of platelet-activating factor (PAF)-induced COX-2 gene expression in the corneal epithelium. Rabbit corneas were incubated in organ-culture in Hank's Ca²⁺-free medium, and the COX-2 mRNA expression in the epithelium was studied. Primary cultures of corneal epithelium were loaded with the fluorescent dye fluo-3, and changes of intracellular Ca²⁺ were analyzed by confocal microscopy. We found that PAF stimulation increased the expression of COX-2 mRNA that peaks at 4 hrs in the corneal epithelium incubated in MEM. The expression was inhibited by the PAF antagonist BN57031. When incubated in Ca²⁺-free medium, there was a 40% inhibition of the induction. The Ca²⁺ ionophore A23187 caused a small but significant increase of COX-2 in the epithelium, which was abolished in Ca²⁺-free medium. When added together, A23187 potentiated the effect of PAF. Confocal microscopic imaging showed that when incubated in Ca²⁺-containing medium, PAF transiently increased intracellular Ca²⁺ which peaks between 30 and 60 seconds after adding PAF. Such effect of PAF was not seen when the cells were incubated with BN57031, which when added with vehicle, or when the cells were incubated in Ca²⁺-free medium. In conclusion, Ca²⁺ is partially required in the induction of the COX-2 gene in the corneal epithelium. The source of such Ca²⁺ is likely to be extracellular, and its entry may be mediated via a PAF receptor. (NIH EY04328)

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POOL DEPLETION INDUCES A NOVEL CALCIUM INFLUX PATHWAY ACTIVATED BY CAFFEINE. C.A. Uretz-Vezeyco, A. Allan, and D.L. Gill - Department of Biological Chemistry, University of Maryland School of Medicine, Baltimore, MD 21201

Ca²⁺ influx through store-operated channels (SOCs) activated rapidly after Ca²⁺ pool depletion represents an important component of Ca²⁺ signals generated in cells. A new and distinct Ca²⁺ influx occurs when caffeine is induced in cells after Ca²⁺ pools are emptied using the intracellular Ca²⁺ pump inhibitors, thapsigargin (TG) or 2,5-di-tert-butylhydrquinone (DBBRQ). Both blockers cause depletion of intracellular Ca²⁺ pools and cell growth arrest: upon refilling of pools, normal cell cycle progression is resumed (Short, A.D., et al. PNAS 90, 4986-4990, 1993). Here, the Ca²⁺-sensitive dyes, fura-2, was used to study Ca²⁺ homeostasis in DDT/MEF-2 smooth muscle cells grown-arrested by TG- or DBBRQ-treatment. In DDT/MEF-2 cells the SOC-mediated Ca²⁺ influx component after emptying Ca²⁺ pools is short-lived and appears to be rapidly deactivated. After treatment of DDT/MEF-2 cells with either 3 μM TG or 10 μM DBBRQ, 10 mM caffeine induces a large transient influx of Ca²⁺ distinct from SOC-mediated Ca²⁺ entry. Caffeine-sensitive Ca²⁺ influx following DBBRQ-treatment is activated more rapidly than that following TG-treatment. When caffeine is added to untreated DDT/MEF-2 cells no effect on cytosolic Ca²⁺ concentration is observed. The disappearance of caffeine-induced Ca²⁺ influx is also different for TG- and DBBRQ-treated cells. In DBBRQ-treated cells, Bradykinin-sensitive Ca²⁺ pools quickly refill and cells become insensitive to caffeine immediately after DBBRQ removal. In the case of TG-treated cells, reversal of TG-induced growth arrest with either high (20%) serum or 10 μM chrysinic acid, in addition to removal of TG, is required to allow agonist-sensitive Ca²⁺ pools to refill concomitantly with the disappearance of caffeine-induced Ca²⁺ influx. In summary, the results show that a Ca²⁺ influx pathway activated by caffeine is observed under conditions of growth arrest induced by either TG or DBBRQ and appears to be directly correlated with depletion of intracellular Ca²⁺ pools. (NIH-grants NS10394 and GM15407; NSF grant MCB 0074774)

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PLATELET-ACTIVATING FACTOR MODULATES INTRACELLULAR CALCIUM DYNAMICS IN HEPATIC CARCINOMA NEURONS. M. De Min, E. De Min, G. Beneze, L.S. Ludic, LSU Medical Center, Neuroscience Center, New Orleans, LA 70112-2234

We have found using confocal microscopy and fluorescent calcium indicators, that intracellular calcium concentration ([Ca²⁺]i) oscillates spontaneously in rat hepatic carcinoma neurones in vivo. While addition of glutamate to these hepatic carcinoma cultures consistently elicited distinct [Ca²⁺]i changes ranging from transient, single spikes (100-500 μM [Ca²⁺]i) to sustained (30-80 μM [Ca²⁺]i) calcium oscillations. Here we have investigated the ability of the potent lipid mediator platelet activating factor (PAF) to affect [Ca²⁺]i dynamics in hepatic carcinoma neurons. When 4 μM methylxanthine (PAF) was added to the culture medium the [Ca²⁺]i was increased slightly. The variance of fluorescence values after methylxanthine additions was 2-fold higher than before addition, indicating an increase in oscillatory [Ca²⁺]i dynamics induced by PAF. Neurons not spontaneously oscillating were observed to be induced to oscillate by PAF addition, and neurons spontaneously oscillating increased in oscillatory behavior upon PAF addition. In agreement with Silici et al. (Neuron, 9:285, 1992) we have found that not all neurones responded to some PAF application. Long-term treatment with PAF appeared to affect the majority of hepatic carcinoma cells. Overnight treatment with PAF (200-400 nm) and methylxanthine (2.4 μM) reduced the neuronal [Ca²⁺]i changes induced by GLU the next day when compared with cells pretreated with PAF (2.4 μM) or vehicle alone. In two cases, the [Ca²⁺]i increases in response to 500 μM GLU were completely inhibited by PAF pretreatment. Since PAF has been shown to enhance hepatic carcinoma cells synaptic transmission (Clark et al., Neuron 9:121, 1992) we postulate that induction of [Ca²⁺]i oscillations by PAF may be an early signal of GLU release, resulting in GLU receptor desensitization (Supported by DAMD-17-93-1-0031)

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CALCIUM-STIMULATED PHOSPHORYLATION OF MAP-2 IN PANCREATIC β-CELLS IS MEDIATED BY CAM KINASE II. R.A. Kruenger, R. Blatt, M. Landi and R.A. Eapen, UTHSC at Fort Worth, Fort Worth, TX 76107 and Washington University School of Medicine, St. Louis, MO 63110

An elevation of intracellular Ca²⁺ is a critical signal in the initiation of insulin secretion from the pancreatic β-cell but the mechanism involved is not understood. Previously, we have demonstrated that the multifunctional Ca²⁺/calmodulin-dependent protein kinase II (CaMKII) is activated by glucagon implicating this enzyme in the secretory process, but its cellular targets are unidentified. One of the best characterized substrates of CaMKII is in vitro that could function in secretory events, is the microtubule-associated protein, MAP-2. The current study represents an evaluation, in vitro, of MAP-2 as a substrate for CaMKII II using pig perifused β-cell model. By immunoblot analysis, the presence of MAP-2 in the jTGC cell was established. In α-toxin-permethylated jTGCs cells, Ca²⁺ induced the concentration dependent phosphorylation of α-toxin II. In parallel and by immunoprecipitation, Ca²⁺ also induced the phosphorylation of MAP-2 that closely correlated with CaMKII II activation. Ca²⁺-induced phosphorylation of MAP-2 was not inhibited by an inhibitor of protein kinase A (H89) at concentrations that prevented phosphorylation induced by forskolin. These data provide evidence that MAP-2 is phosphorylated by Ca²⁺ in the pancreatic β-cell and that this event may provide an important link in the mediation of Ca²⁺-dependent insulin secretion. (Supported by NIH grant DK-47925)

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RE-EVALUATION OF THE ROLE OF CAM KINASE II IN INSULIN SECRETION IN PANCREATIC β-CELLS. M. Blum, L. Tankay and R.A. Eapen, UTHSC at Fort Worth, Fort Worth, TX 76107

Current evidence addressing the role of CaMKII in insulin secretion is conflicting. Wertz et al. have recently demonstrated that stimulation of insulin secretion in the presence of glucose and forskolin showed a reduction in the CaMKII activity. In another study, the CaMKII inhibitor, KN-62 (1 μM), failed to inhibit Ca²⁺-insulin secretion induced from streptolydin O-permeabilized HIT cells, leading to the implication that this enzyme has no role in the secretory process. In this study, however, KN-62 at concentrations up to 100 μM did not inhibit CaMKII II activity in cellular extracts of jTGC cells in the presence of exogenous calmodulin, and in the absence of the previously described inhibitor. In contrast, α-toxin-permeabilized jTGC cells, Ca²⁺-induced rapid activation of CaMKII II in a concentration-dependent manner that was maintained for at least 30 min. This activation was not prevented by KN-62 (0-100 μM) nor was the CaMKII II not involved in Ca²⁺-induced insulin secretion. The kinase inhibitor, K252a and a selective peptide inhibitor, (Ala₁⁴⁶)CaMK 281-302 strongly inhibited CaMKII II activity in jTGC cells and could have established to permit the evaluation of the effects of these compounds on Ca²⁺-induced insulin secretion. A more stringent correlate of the extent of inhibition of CaMKII II and insulin secretion by these compounds will permit a better assessment of the role of this enzyme in insulin secretion. (Supported by NIH 47925)

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OTHER EFFECTS OF ACETAZOLAMIDE IN RAT LIVER MITOCHONDRIA. A. Saavedra-Molina and M. Clemente-Requena, Instituto de Investigaciones Quimico-Biologicas. Universidad Michoacana, Morelia, Mich. 58030, MEXICO

Preliminary studies using the carbonic anhydrase inhibitor acetazolamide showed an inhibition of mitochondrial function in rat liver mitochondria (Arch. Biochem. Biophys. 1986, 251, 1206-207) and it was concluded that this sulfonamide produces also an inhibition of urea synthesis in isolated guinea pig hepatocytes.

In this report we demonstrated that in rat liver mitochondria in the presence of different concentrations of acetazolamide, citrate synthesis was decreased in a dose-response fashion obtaining the maximum inhibition (35%) with 300 μM; however, other effects were found. By using fluo-3 as a mitochondrial fluorescent calcium indicator, matrix free calcium (Ca²⁺) was measured in a Mg²⁺-based buffer (pH 7.4) in the presence of EGTA. The effect of acetazolamide was a decrease of cytoplasmic calcium, whereas the reaction was performed in the presence of 3 μM Ca²⁺. Mitochondrial free calcium increased 3-fold (from 1.2 μM to 3.6 μM) with 50-200 μM acetazolamide. The results obtained described another effect of the sulfonamide acetazolamide on mitochondrial matrix free calcium, which indicate that calcium ions could exert a physiological effect on citrulline synthesis.

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Synergy by Secretory Phospholipase A₂ and Glutamate on Inducing Cell Death and Sustained Arachidonic Acid Metabolic Changes in Primary Cortical Neuronal Cultures

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Secretory and cytosolic phospholipases A₂ (sPLA₂ and cPLA₂) may contribute to the release of arachidonic acid and other bioactive lipids, which are modulators of synaptic function. In primary cortical neuron cultures, neurotransmitter cell death and [³H]arachidonate metabolism was studied after adding glutamate and sPLA₂ from bee venom. sPLA₂, at concentrations eliciting low neurotoxicity (≤100 ng/ml), induced a decrease of [³H]arachidonate-phospholipids and preferential reesterification of the fatty acid into triacylglycerols. Free [³H]arachidonic acid accumulated at higher enzyme concentrations, below those exerting highest toxicity. Synergy in neurotoxicity and [³H]arachidonate release was observed when low, nontoxic (10 ng/ml, 0.71 nm), or mildly toxic (25 ng/ml, 1.78 nm) concentrations of sPLA₂ were added together with glutamate (90 μM). A similar synergy was observed with the sPLA₂-OS2 from Taipan snake venom. The NMDA receptor antagonist MK-801 blocked glutamate effects and partially inhibited sPLA₂-OS2 but not sPLA₂ from bee venom-induced arachidonic acid release. Thus, the synergy with glutamate and very low concentrations of exogenously added sPLA₂ suggests a potential role for this enzyme in the modulation of glutamatergic synaptic function and of excitotoxicity.

Membrane unsaturated fatty acid turnover and the synthesis of bioactive lipids are modulated by phospholipases A₂ (PLA₂), ubiquitous mammalian enzymes that catalyze the hydrolysis of sn-2-acyl ester bonds of phospholipids (PLA) (1). Arachidonic acid (AA), eicosanoids, and platelet-activating factor (PAF) are bioactive lipids generated through PLA₂ activation (2). Although some PLA₂ are calcium-independent (3, 4), most found in the brain are characterized by calcium dependence (4, 5). PLA₂ are overstimulated in the brain during seizures and ischemia (6–8) as a consequence of increased calcium influx and/or intracellular calcium mobilization, which, in turn, results in the accumulation of bioactive lipids that participate in cell damage (8, 9).

There are secretory and cytosolic PLA₂ (sPLA₂ and cPLA₂, respectively). sPLA₂ (14 kDa) are active at submillimolar concentrations of calcium and do not display selectivity for unsaturated fatty acids at the sn-2-position of PL (4, 5). PLA₂ are found in pancreatic secretions (type I), platelets, neurons, mast cells, snake venoms, inflammatory exudates (type II), and bee venom (type III) (4, 5, 10). In contrast, cPLA₂ (type IV) has a higher molecular mass (85 kDa), is active at submicromolar Ca²⁺ concentrations, and shows selectivity for sn-2-arachidonyl-PLs (5, 11). cPLA₂ is activated by translocation to intracellular and nuclear membranes when there is an agonist-induced increase in intracellular calcium concentration ([Ca²⁺]⁰) in the brain (12, 13) as well as in other tissues (4, 14).

Among the neural forms of PLA₂ are (a) a calcium-sensitive and arachidonoyl-specific 85-kDa cPLA₂ (12, 15, 16), highly expressed in astrocytes (17), other cytosolic calcium-dependent forms (12, 16), and calcium-independent forms (3, 18, 19); and (b) membrane-bound forms (15), including a very high molecular mass (180-kDa) form from human temporal cortex (20). Secretory PLA₂ are also present in the brain. The expression of cPLA₂ type II is stimulated in the rat brain by ischemia/reperfusion (21) and in cultured astrocytes by inflammatory mediators (22). Moreover, sPLA₂ type II is stored in synaptic vesicles and released by depolarization or neurotransmitter stimulation, and its secretion is coupled with the activation of catecholamine release (23). Furthermore, sPLA₂ causes activation of Glu release in the rat cerebral cortex (24).

sPLA₂ bind to cell surface receptors, the N type and the M type (25–28) identified using sPLA₂ purified from snake and bee venoms as ligands. Neurotoxic sPLA₂ from Taipan snake venom, OS2, and from bee venom bind to the N-type receptor with high affinity (25, 26). Other sPLA₂ such as OS1, also purified from Taipan snake venom, display higher enzymatic activity than the sPLA₂ OS2 and bee venom (2.7- and 7-fold higher, respectively) (25). Although OS1 binds with high affinity to M-type receptors (26–28), it does not bind to N-type receptors (25) and is therefore non-neurotoxic.

Activation of cPLA₂ mediates the formation of modulators of synaptic transmission such as free AA (8), eicosanoids (29, 30), and PAF (31). Ischemia and seizures promote a rapid increase in brain free AA (6, 7, 32, 33), oxygenated metabolites of AA, and free radicals, all of which are potent neuronal injury mediators (for review, see Ref. 34). A sustained activation of cPLA₂ has been reported after ischemia/reperfusion (13, 15). Glu, which causes excitotoxic neuronal damage, increases calcium influx through NMDA receptors in post synaptic neurons, leading to PLA₂-mediated AA release (34–37), which is blocked by the NMDA antagonist MK-801 (38). Recently, the activation of two calcium-dependent cPLA₂ (100 and 14 kDa) by Glu was reported (38).

This paper is available online at http://www.jbc.stanford.edu/jbc/
SECRETORY PLÁ Neurotoxicity and AA Release

Other PLÁ are modulators of membrane PL metabolism and/or generate membrane fusogenic molecules, i.e. free fatty acids and lyso-PLÁ. However, non-AA-specific sPLÁ contribute, together with the cPLÁ, to modulate AA metabolism under physiological conditions and in primed conditions (5, 59, 42). sPLÁ venoms have long been known to be neurotoxic (10, 43). For example, sPLÁ from Naja mocambique, as well as the PLÁ activator melittin, have previously been shown to promote neural injury, in vitro and in vivo (44). The present study has tested the hypothesis that sPLÁ potentiates neurotoxicity by Glu by briefly exposing rat cortical neuronal cultures to this neurotransmitter in the presence of and in the absence of sPLÁ. This hypothesis is supported in part by the observation that sPLÁ is released at synapses from vesicles that also store Glu (23). sPLÁ from bee venom and Taipan snake venom OS2, ligands of the N-type sPLÁ receptor, and OS1 from Taipan snake venom, a ligand of the M-type sPLÁ receptor, have been used (25-28). Moreover, sustained changes in neuronal [³H]AA metabolism under these conditions have been observed. Our study has addressed the following: (a) the effect of Glu activated cPLÁ and exogenously added sPLÁ on [³H]AA release from neuronal PLÁ; (b) the action of MK-801 on agonist-induced AA changes and neurotoxicity; (c) the neurotoxic potential of exogenously added sPLÁ (OS2 and bee venom sPLÁ); and (d) the neurotoxic effect of the excitatory neurotransmitter Glu and sPLÁ when added to the cells simultaneously.

EXPERIMENTAL PROCEDURES

Materials—[5,6,8,9,11,13,14,15,²H]Arachidonic acid ([²H]AA, 185 Ci/mmol) was purchased from DuPont NEN. Silicagel GHL TLC plates were obtained from Sigma (St. Louis, MO). Organic solvents, hexane, chloroform, ethyl acetate, and methanol were obtained from EMD. sPLÁ from bee venom and Glu and OS2 from Taipan snake venom were purified as described previously (25).

Primary Neuronal Cultures—Cortical neuronal cultures were established from 15-day-old rat embryos as described previously (45, 47). Cells were plated in poly-lysine-covered dishes at a density of 4 x 10⁵ cells/well in 48-well plates for toxicity experiments and 7.5 x 10⁶ cells/well in 24-well plates for lipid analysis. The cells were cultured in neuronal culture medium (N-122) containing 10% fetal calf serum and 10% horse serum in a 5% CO₂ incubator at 37 °C. Astrocyte proliferation was prevented by adding cytosine arabinoside (10⁻³ M) at day 4 after plating, left on for 3 days, and then replaced with minimal essential medium containing 10% horse serum. Cells were used for experiments at 14–21 days in vitro. The percentage of neuronal versus glial cells remained at approximately 80 versus 20%, respectively, as previously reported (48).

Neuronal Toxicity Assay—Lactate dehydrogenase (LDH) release was used to quantitatively assess cell injury. Cells were treated as described previously (47), and LDH release was measured 2 h after exposure to sPLÁ and/or Glu. Briefly, cells were exposed for 45 min at room temperature to highly purified sPLÁ from bee venom and snake venom (OS1 or OS2) and/or Glu (50 µM) in Locke’s solution without sPLÁ. Locke’s solution was exchanged with minimum essential medium without phenol red, and the cultures were returned to the CO₂ incubator for 20 h at 37 °C. Using similar cortical neuron culture, this procedure results in minimal LDH release before 12 h and maximal release by 20 h (49). The LDH release was assessed using the Sigma kit with the aid of a DU 68 Beckman spectrophotometer equipped with a graphic video display.

[²H]Arachidonic Acid Metabolism and Lipid Analysis—Cells were labeled overnight with [²H]AA (0.5 µCi/well, specific activity 184.6 Ci/mmol) in minimal essential medium supplemented with 0.2% fatty acid-free bovine serum albumin. 18–20 h later, the medium of cell cultures was replaced with minimum essential medium. The cells were exposed to Glu and/or sPLÁ as described previously and returned to the CO₂ incubator for 2 and 20 h at 37 °C. The supernatants were removed, and 750 µl of methanol was added to each well. The cells were scraped and transferred into a glass tube, and chloroform was added in the following proportion: chloroform/methanol, 2:1 (v/v). Lipids were extracted by sonication for 90 min, and the lipid extracts were washed following the procedure of Folch et al. (50). Aliquots in duplicate were taken to determine total [²H]AA incorporation by liquid scintillation counting.

Neutral lipids and total PLÁ were isolated by monodimensional TLC on precoated, 0.25-mm-thick, silica gel GHL plates using hexane-ethyl ether-acetic acid (50:50:1, v/v/v), a developing solvent. For the results presented in Table 1, the composition of the chromatographic solvent was changed to 60:40:1.3 (v/v/v) in order to isolate cholesterol ester (CHE) from triglycerides (TAG) that run together in the first system. The results indicated that changes observed in previous experiments in the CHE plus TAG fraction were due mainly to TAG. Lipid standards were added to the samples as a carrier and were spotted in a parallel line on the TLC plate to individualize each lipid band. TLC plates were developed with iodine, lipid bands were scraped, and the radioactivity was determined in a Beckman scintillation counter.

Statistical Analysis—The significance of the data was evaluated with Student's t test for unpaired data. Statistical values were considered significantly different when p < 0.05.

RESULTS

Potent Neurotoxicity Elicited by sPLÁ—Neurotoxicity of sPLÁ from bee venom, added alone or with Glu (80 µM) to primary neuronal cortical cultures, was studied by measuring LDH release (Fig. 1). A neurotoxic concentration of Glu (47) resulted in 82 ± 2% increase in LDH release compared with control cells (n = 54 from 10 different plateings). No significant toxicity was observed with bee venom sPLÁ concentrations up to 10 ng/ml (Fig. 1A). At higher concentrations up to 10⁻⁵ ng/ml, the neurotoxicity displayed by sPLÁ was biphasic. First, there was a dose-dependent increase in LDH release (up to 500 ng/ml, EC₅₀ = 7.1 nm), followed by a sharp 4.3-fold increase in LDH at 10⁻⁵ ng/ml (EC₅₀ = 57 nm). The sPLÁ neurotoxicity at 100 ng/ml (7.1 nm) was similar to that of 80 µM Glu (Fig. 1). An additive neurotoxic effect was observed when cells were exposed simultaneously to 80 µM Glu and low (1–5 ng/ml) or high (50–10⁻⁵ ng/ml) sPLÁ concentrations (Fig. 1, inset). The combination of sPLÁ in a concentration range of 10–25 ng/ml with Glu led to a significant synergy on LDH release, resulting in values 2.2–2.5-fold higher than those of the additive effects of sPLÁ and Glu.

The sPLÁ OS2 purified from snake venom was found to be more potent as a neurotoxin than the sPLÁ from bee venom. For example, at 25 ng/ml OS2 was approximately 2.7-fold more...
toxic than bee venom sPLA₂ at the same concentration (Fig. 2). Furthermore, under conditions where the noncompetitive NMDA antagonist MK-801 blocked 100% of 80 μM Glu toxicity, MK-801 partially blocked OS2, but not bee venom sPLA₂-induced toxicity. OS2 did not evoke neuronal death even at 10 μg/ml (LDH percentage above control = 15 ± 9%).

sPLA₂ Promotes Arachidonic Acid Release from Phospholipids—[^H]AA-prelabeled neuronal cells were exposed to different concentrations of bee venom sPLA₂ for 45 min and further incubated for 20 hr (Fig. 3). No differences were observed in total[^H]AA labeling recovered per dish at very low, nontoxic sPLA₂ concentration (1 ng/ml). At higher concentrations (25–50 ng/ml), the recovery was decreased by 10% and by 20–30% at more toxic concentrations (500–1000 ng/ml), reflecting cell loss and matching the neurotoxicity assays (Fig. 1). After 20 hr the[^H]AA distribution displayed a concentration-dependent loss of[^H]AA-PL paralleled by an increase in free[^H]AA,[^H]AA-TAG and[^H]AA-DAG. A significant loss in PL labeling was observed even at the lowest sPLA₂ concentration (~7%, p < 0.05), reaching values 50% lower at the highest toxic concentrations (500–1000 ng/ml). Up to 100 ng/ml sPLA₂, the loss of[^H]AA from phospholipids (~29%) was paralleled by its active reestatification into TAG, which showed a 25% increase above the control value. Within this range of sPLA₂ concentrations, free[^H]AA showed a small yet significant gradual increase, reaching values 2- and 4.5-fold higher than controls at 1 ng/ml and 100 ng/ml, respectively. The[^H]AA-TAG labeling plateaued at 500 ng/ml sPLA₂. This was paralleled by a large increase in free[^H]AA accumulation, which reached a value 20-fold higher than control.[^H]TAG labeling was very low, displaying the same pattern of changes as free[^H]AA and reaching a 2-fold increase in percentage of labeling at high sPLA₂ concentration (500 ng/ml).

Triacylglycerols Are a Finite Reservoir for the Uptake of[^H]AA Released by sPLA₂ and Glu—To ascertain if[^H]AA released by bee venom sPLA₂ was acylated into TAG and whether or not this correlated with neurotoxicity, the following experiment was performed. The[^H]AA metabolism as affected by a nontoxic concentration of sPLA₂ (1 ng/ml) and by a toxic concentration of Glu (50 μM), added individually or combined, was studied at 2 and 20 hr after treatment with the agonists (Fig. 4). sPLA₂ induced a similar decrease in[^H]AA-PL labeling both at 2 and 20 hr. Differences were observed, however, in the distribution of labeling between free[^H]AA and[^H]AA-TAG. Free[^H]AA accumulation was greater at 2 hr, decreasing by 20 hr concomitantly with an increase in[^H]AA-TAG.

Glu alone triggered a similar loss in[^H]AA-PL compared with sPLA₂ by 2 hr; however, by 20 hr, loss of[^H]AA from PL was 2.8-fold greater than at 2 hr. After treatment with Glu alone, free[^H]AA and[^H]AA-TAG varied as a function of time (similar to when sPLA₂ was added alone), with higher accumulation of free[^H]AA by 2 hr and a preferential reestatification of[^H]AA into TAG by 20 hr.

sPLA₂ and Glu added together greatly magnified the pattern of[^H]AA changes as a function of time. A synergy on free[^H]AA accumulation was observed due to an apparently less efficient esterification into TAG. By 20 hr the level of free[^H]AA reached 1.8–2-fold higher values than when both agonists were individually added. The loss of[^H]AA from PLs was additive, as was the accumulation of[^H]AA-DAG induced at 2 and 20 hr.

MK-801 Does Not Block Arachidonic Acid Release Induced by sPLA₂ from Bee Venom but Partially Blocks the Effect of OS2 from Snake Venom—The involvement of NMDA receptors on AA release from PLs induced by sPLA₂ and Glu was investigated by preincubating cells with 300 nM MK-801 for 10 min prior to adding the agonists, followed by lipid analysis 20 hr later. Both at low, nontoxic (1 ng/ml) (data not shown) and at higher (25 ng/ml) bee venom sPLA₂ concentrations (Table 1), MK-801 did not block the release of[^H]AA from PLs. The phospholipid labeling was decreased by 17% (p < 0.002), from 87% in controls to 70% in sPLA₂-treated cells. Most of the
Secretory PLA$_2$ Neurotoxicity and AA Release

$[^3]$HAA released from PL (+11%) was found reesterified into TAG (6 versus 16% for control and sPLA$_2$-treated, respectively) and to a lesser extent in CHE (+4%, $p < 0.03$), while free $[^3]$HAA labeling was doubled (from 1 to 2%, $p < 0.03$). MK-801 pretreatment did not alter the profile of lipid labeling, i.e. the decrease in PLs and the parallel increase in TAG and free AA labeling.

Glu (80 µM), although more toxic than 25 ng/ml bee venom sPLA$_2$ (sPLA$_2$ toxicity 29% compared with Glu; Fig. 1), induced only a 6% ($p < 0.002$) decrease in PL labeling concomitantly with increased labeling of TAG (+2%, $p < 0.004$), CHE (+2%, $p < 0.02$), and FFA (+0.4%, $p < 0.03$). MK-801 pretreatment blocked by 100% Glu-induced PL degradation and other lipid changes. Higher degradation of PLs was observed when bee venom sPLA$_2$ and Glu were added together to the cells (−30%).

Labeling of TAG increased by 24%, and labeling of free $[^3]$HAA increased by 3% ($p < 0.03$). MK-801 pretreatment blocked partially the changes induced by bee venom sPLA$_2$ and Glu, leading to the same profile of lipid labeling induced by sPLA$_2$ alone.

The sPLA$_2$ from snake venom, OS2, added to the cells at the same concentration as sPLA$_2$ from bee venom (25 ng/ml), induced a much greater degradation of $[^3]$HAA-PLs. MK-801, in contrast to the results with bee venom, partially blocked $[^3]$HAA-PL hydrolysis induced by OS2 when added alone or together with Glu (Table I). Moreover, the total labeling recovered per well treated with OS2 and OS2 plus Glu was decreased by 35%, indicating a massive loss of cells. The DPM/well obtained when the cells were pretreated with MK-801 was similar to controls. Minimal changes in $[^3]$HAA-lipid labeling were observed when the cells were treated with the sPLA$_2$ (25 ng/ml) from snake venom OS1 (data not shown), which does not bind to neuronal membranes and which was found to be non-neurotoxic (see above).

sPLA$_2$ Display a Synergy with Glu in $[^3]$HAA Release from Phospholipids—a sPLA$_2$ (25 ng/ml) from snake and bee venoms added with Glu displayed synergy leading to a higher $[^3]$HAA-PL degradation than the sum of the effect of the individual agonists (Table I, Fig. 5). Although the toxicity and PL hydrolysis induced by OS2 was much greater than that of bee venom sPLA$_2$ (Table I), the synergy with Glu was similar, reaching values for PL hydrolysis 1.4-fold higher for both sPLA$_2$ (Fig. 5C). A synergy was also observed in the accumulation of $[^3]$HAA-TAG that increased by 2-fold for bee venom sPLA$_2$ and 1.4-fold for OS2 (Fig. 5B). The synergy in free $[^3]$HAA accumulation was much greater with OS2 (3.5-fold) than with sPLA$_2$ from bee venom (2-fold) (Fig. 5A), and the synergy of sPLA$_2$ plus Glu was blocked by MK-801 (Table I).

Accumulation of Free $[^3]$HAA in Cortical Neuronal Cells Proceeds the Toxicity Induced by High Concentrations of Bee Venom sPLA$_2$—Treatment of neuronal cultures with increasing concentrations of sPLA$_2$ resulted in increased neurotoxicity (Fig. 1) and higher degradation of AA-PLs (Fig. 4). Changes in lipid labeling plotted as a function of sPLA$_2$ toxicity are shown in Fig. 6. The accumulation of free $[^3]$HAA was minimal and proportional to increased LDH up to 100%, when sPLA$_2$ toxicity was equal to that of 80 µM Glu (±100 ng/ml sPLA$_2$). Within this range of neurotoxicity, most of the $[^3]$HAA released from PLs (∼50%) was reesterified into TAG. While PLs displayed a gradual loss of $[^3]$HAA up to LDH values of 200% (∼50% decrease in PL labeling), accumulation of free $[^3]$HAA peaked between LDH values of 100 and 200%. This increase in free $[^3]$HAA preceded a 4.3-fold increase in LDH release observed for sPLA$_2$ concentrations between 500 ng/ml (217%) and 10 ng/ml (387% LDH).

**Table I**

<table>
<thead>
<tr>
<th>Condition</th>
<th>CHE</th>
<th>TAG</th>
<th>FFA</th>
<th>DAG</th>
<th>PL</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>%</td>
<td>%</td>
<td>%</td>
<td>%</td>
<td>dpm/well</td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>26,018 ± 10,723</td>
<td></td>
</tr>
<tr>
<td>Glu</td>
<td>5.8 ± 0.8</td>
<td>4.9 ± 0.5</td>
<td>1.0 ± 0.1</td>
<td>1.7 ± 0.1</td>
<td>86.6 ± 1.3</td>
<td></td>
</tr>
<tr>
<td>Glu + MK-801</td>
<td>8.2 ± 0.5*</td>
<td>7.3 ± 0.5*</td>
<td>1.3 ± 0.1*</td>
<td>2.3 ± 0.2*</td>
<td>80.8 ± 1.0*</td>
<td></td>
</tr>
<tr>
<td>sPLA$_2$</td>
<td>6.7 ± 0.6</td>
<td>4.7 ± 0.1</td>
<td>1.0 ± 0.0</td>
<td>1.5 ± 0.2</td>
<td>86.1 ± 0.5</td>
<td></td>
</tr>
<tr>
<td>sPLA$_2$ + MK-801</td>
<td>9.9 ± 1.4</td>
<td>15.7 ± 1.8</td>
<td>2.5 ± 0.1</td>
<td>2.1 ± 0.3</td>
<td>70.1 ± 2.3</td>
<td></td>
</tr>
<tr>
<td>sPLA$_2$ + GLU</td>
<td>7.5 ± 1.2</td>
<td>14.6 ± 2.5*</td>
<td>2.3 ± 0.3*</td>
<td>1.8 ± 0.2</td>
<td>73.6 ± 3.9</td>
<td></td>
</tr>
<tr>
<td>sPLA$_2$ + GLU + MK-801</td>
<td>7.4 ± 1.4</td>
<td>28.8 ± 4.0*</td>
<td>4.0 ± 0.8*</td>
<td>2.9 ± 0.1*</td>
<td>56.9 ± 4.9*</td>
<td></td>
</tr>
<tr>
<td>OS2</td>
<td>5.0 ± 0.2</td>
<td>14.8 ± 3.0</td>
<td>2.5 ± 0.6</td>
<td>2.1 ± 0.0</td>
<td>75.5 ± 4.1</td>
<td></td>
</tr>
<tr>
<td>OS2 + MK-801</td>
<td>5.0 ± 0.2</td>
<td>14.8 ± 3.0</td>
<td>2.5 ± 0.6</td>
<td>2.1 ± 0.0</td>
<td>75.5 ± 4.1</td>
<td></td>
</tr>
<tr>
<td>OS2 + GLU</td>
<td>6.0 ± 0.3</td>
<td>32.8 ± 2.9</td>
<td>15.4 ± 3.3</td>
<td>3.5 ± 0.2</td>
<td>41.8 ± 6.9</td>
<td>173,684 ± 114*</td>
</tr>
<tr>
<td>OS2 + GLU + MK-801</td>
<td>5.0 ± 0.2</td>
<td>14.8 ± 3.0</td>
<td>2.5 ± 0.6</td>
<td>2.1 ± 0.0</td>
<td>75.5 ± 4.1</td>
<td></td>
</tr>
</tbody>
</table>

 Asterisks denote values statistically significantly different from control (Student's t-test, $p < 0.05$).
**DISCUSSION**

This study shows that treatment of primary cortical neurons in culture with sPLA₂ induces (a) a concentration-dependent increase in neurotoxicity; (b) sustained activation of [³H]A2A mobilization reflected in a gradual loss of [³H]AA from PLs and in an accumulation of free [³H]AA followed by its reesterification into TAG; and (c) synergy with Glu (80 μM) for both neurotoxicity and [³H]AA-PL hydrolysis.

Neurotoxicity and sustained changes in AA metabolism, triggered by 45-min exposure of primary cortical neurons to Glu were blocked by the NMDA receptor antagonist MK-801 (Fig. 2, Table 1) in agreement with previous studies (54–57, 51–53). Moreover, the release of [³H]AA from PLs was observed 2 h after the treatment of neuronal cultures with Glu, and even greater release was observed 20 h later (Fig. 4). Long lasting changes in AA metabolism may be the result of calcium and protein kinase C-mediated, sustained activation of neuronal cPLA₂ by Glu (16). Moreover, increased cPLA₂ activity correlates with Glu neurotoxicity and precedes irreversible neuronal injury (16). It is also possible that, as in mast cells (54), Glu may regulate cPLA₂ activity at early time points by protein kinase C-mitogen activated protein kinase phosphorylation and later by enhanced expression of the enzyme. Modulation of gene expression and increased protein synthesis are involved in long term cellular responses as in neuronal plasticity or delayed neuronal death. In fact, cPLA₂ activation by NMDA-glutamatergic synaptic activity may lead to the formation of PAF, a potent bioactive lipid, which, in turn, mediates the induction of early response genes and subsequent gene cascades (2, 55–57). PAF could also potentiate excitotoxicity by enhancing Glu release (58, 59).

Although the toxicity of Glu (80 μM) was similar to that induced by bee venom sPLA₂ (100 ng/ml; Fig. 1), the hydrolysis of [³H]AA-PLs 20 h after Glu treatment (~15%; Fig. 4) was half that generated by 100 ng/ml bee venom sPLA₂ (~29%; Fig. 3). These results and the fact that MK-801 blocked Glu neurotoxicity support the notion that mechanisms other than cPLA₂ activation mediated by Glu-activated NMDA-gated calcium channels contribute to its neurotoxic action (8). Glu may also activate metabotropic receptors that, in turn, activate phospholipase C with the release of AA-DAG, a potent activator of protein kinase C (60). Sequential degradation of AA-DAG by diacylglycerol lipases and monacylglycerol lipases contribute also to increased free [³H]AA (61).

Bee venom sPLA₂-dependent sustained changes in [³H]AA-lipid metabolism (2 to 20 h after adding the enzyme) reveal an active release of [³H]AA from PLs, transient accumulation of free [³H]AA, and reesterification into TAG. A similar effect was observed with Glu, with increased labeling of free [³H]AA by 2 h decreasing by 20 h concomitantly with increased [³H]AA-TAG labeling. Interestingly, free [³H]AA was shunted into TAG even when cells were exposed to very low, nontoxic concentrations of sPLA₂ (1 to 10 ng/ml). Thus, the pathway activated by sPLA₂ may be physiologically relevant, withholding AA from its conversion to eicosanoids and from exerting effects of its own. AA is a modulator of synaptic function and potentiates Glu-NMDA neurotransmission, leading to excitotoxic damage (6). Free AA can be further metabolized to eicosanoids, potent modulators of synaptic function (29, 30), which, when overproduced, become injury mediators (8). TAG may also be a transient reservoir of AA when there is activation of degradative pathways, protecting the cells from the loss of this essential fatty acid. In fact, part of the [³H]AA released during repeated seizures from neuronal membrane PLs in rat brain is shunted into TAG (7). This pathway was also activated in retina by experimental detachment (62), where another polyunsaturated fatty acid, docosahexaenoate (22:6n-3), is actively esterified into TAG. A reversible accumulation of AA-TAG occurs in non-neural cells cultured in the presence of high concentrations of FFA (63, 64). In the present study, even 20 h after transient cell stimulation with nontoxic concentrations of sPLA₂, [³H]AA released from PLs remained as [³H]AA-TAG. This indicates long lasting metabolic changes, since between 2 and 20 h post-treatment, PLs did not recover basal labeling, and free [³H]AA was shunted into TAG.

The TAG reservoir appears to have a limited capacity to store AA. The maximum was reached at bee venom sPLA₂ concentrations between 50 and 100 ng/ml. AA-PL hydrolysis in neuronal cortical cells was much more sensitive to sPLA₂ than toxicity, within a range of LDH release similar to that exerted by Glu (Fig. 6). Thus, sPLA₂ concentrations ≥100 ng/ml (toxicity ≥100% to Glu), the bulk of [³H]AA mobilized from PLs...
displaying different affinity for the enzymes.

In summary, this study shows that exogenously added sPLA₂ and Glu induce sustained changes in neuronal AA-PL metabolism and that sPLA₂ plus Glu exerts synergistic mobilization of AA and subsequent neurotoxicity. The present results, taken together with the recent observation that sPLA₂ type II in synaptic vesicles is released together with neurotransmitters (22), open up the possibility that glutamatergic neurotransmission involves the corelease of glutamate and sPLA₂. Our observations also imply that excitotoxicity may involve not only glutamate, as currently assumed, but may also involve sPLA₂ at the synaptic cleft. Further studies will assess if Glu could potentiate endogenous mammalian sPLA₂ actions that could, in turn, stimulate further Glu release. In this connection it is relevant that the synthesis of PAF, a retrograde messenger of long term potentiation (58), may be enhanced by sPLA₂ at the synapse. “Cross-talk” between cPLA₂ and sPLA₂ has recently been suggested in signal transduction events in macrophages (40), and a complex interplay between Glu-activated cPLA₂ and sPLA₂ could be envisioned at the synapse. Several of these ideas are currently under investigation in our laboratory.

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