

Report of program activities.

THE THIRD ANNUAL REPORT

of

CLINICAL INVESTIGATIONS

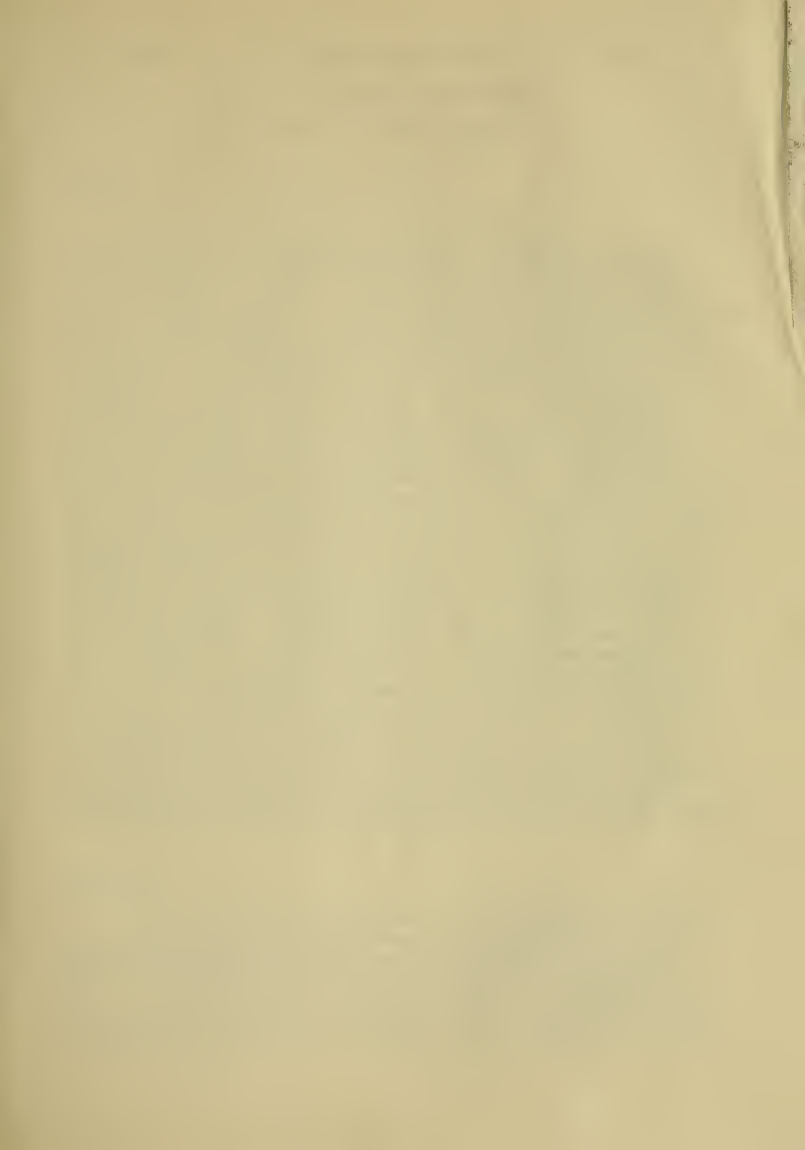
of the

United States, NATIONAL INSTITUTE OF NEUROLOGICAL DISEASES AND BLINDNESS

1955

National Institutes of Health, Public Health Service
U. S. Department of Health, Education, and Welfare

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National Institute of Neurological Diseases and Blindness

Clinical Investigations Annual Report

The Clinical Director's Report

General:

At the time of submission of the third annual report, the Clinical Investigation Unit of the National Institute of Neurological Diseases and Blindness has reached its total activation of three wards for a total of 78 beds. There were 357 inpatient admissions and 548 outpatient admissions. With the addition of Doctor Ludwig von Sallmann, the Branch of Ophthalmological Disorders rapidly expanded with the activation of six sections, recruitment of 13 investigators, and their supporting staff of technicians, etc. A corresponding increase in the number of research projects since 1954 shows a total of 98 projects for the Clinical Investigation Unit of this Institute. Many and varied neurological and ophthalmological diseases are under investigation with both clinical and applied basic research in such areas. Neurosurgery has been occupied with continuing studies of brain physiology and its relation to epilepsy, and, in particular, to seizures originating in the temporal lobe and the deep nuclei of the temporal lobe. In addition, studies in involuntary movement, pain, and the effects of surgical removals on the visual system have received intensive investigation. The Medical Neurology Branch is still continuing its studies on diseases of the lower motor neuron, demyelinating disorders, amyotrophic lateral sclerosis, and other diseases of the central and peripheral nervous system. The new Ophthalmological Branch has launched a broad program into the cause and basic mechanism underlying glaucoma, cataract, and inflammatory diseases of the orbit. The Epidemiology Branch has continued its field studies on amyotrophic lateral sclerosis and a pilot study of all neurological diseases in small closed medical communities. The Electroencephalography Branch has carried on studies of the correlation of metrazol-induced seizures, and the correlation of such with clinical and pathological material. This Branch is continuing its studies of experimental seizures utilizing injected penicillin and its studies on the thalamic and cortical connections.

Specific:

The Medical Branch, in its studies on neuromuscular disorders, has completed, with the aid of Doctor J. Godwin Greenfield, a basic Atlas of Muscle Pathology which has been accepted for publication by E. & S. Livingstone in Edinburgh, England. This pathology is based on approximately 5,000 slides from biopsies obtained from 127 patients with neuromuscular disorders. Specific pathological changes are described in afflicted muscle fibers and the nuclei. Positive

and negative clinical correlation have been presented in tabular form to demonstrate the basic pathology of neuropathies and myopathies. Regeneration of muscle has been demonstrated in patients with muscular dystrophy, and specific anatomical pathological changes are illustrated in dystrophia myotonica, but it has been found that it is impossible to differentiate muscular dystrophy from polymyositis on the basis of pathology alone.

Patients with muscular dystrophy have shown a decreased exchange of body potassium. Such an exchange, however, is a reflection of the decreased muscle mass of the body and not an inability of the muscle cell to use potassium. Dystrophic patients show an abnormal permeability to albumin tagged with Iodine 131. Such a permeability does not exist if the tag is on the red cell with Chromium 51. It is still not clear whether the protein is fractionated before or after leaving the blood stream. Studies in this regard are continuing. The finding of larger than normal amounts of oxyproteic acid in the urine of dystrophic patients by Doctor Curtis again points to some abnormality in protein metabolism.

The main emphasis of this Branch will now turn to the disease myasthenia gravis. It has been found that hexamethonium will relieve in part the symptoms of myasthenia gravis. Hexamethonium will also reverse the paralysis of the rat receiving either decamethonium or curare. The way in which this is done is not clear since hexamethonium presumably has no anticholinesterase activity. To further elucidate this, as well as the physiology of the altered end plate in myasthenia gravis, Doctors Shy and Li have been attempting to record from a single motor end plate while within a muscle fiber. They have been aided in this regard by Doctor Paul Fatt from Australia, and single end plate potentials have now been recorded for the first time in vivo from the mammalian muscle.

Doctor Tower and his group have demonstrated quantitatively an increase in aldol pentose excretion in patients with neuromuscular disease, but this also seems a reflection of the residual muscle mass and not of primary etiology in the disorder of muscular dystrophy. Doctor Horvath and Doctor Shy are continuing the studies of actomyosin threads and glycerinated fibers in muscle diseases. It would appear that normal human muscle is extremely sensitive to changes in ATP, potassium, and magnesium. Myotonic muscle, however, will contract over a broad spectrum of concentrations of these substances.

Doctor Tower and his group are continuing their studies of changes accompanying demyelinating disorders that confirm Kabot's findings that 65% of patients with multiple sclerosis demonstrate an increase in gamma globulin. Such an elevation, however, rarely precedes an onset of symptoms. Electrophoresis studies further reveal that an increase in gamma globulin occurs in many neurological disorders, with higher levels found in the cerebellar ataxias,

all of which discredits the specificity of gamma globulin in the diagnosis and prognosis of multiple sclerosis. A double-blind pilot study involving 81 patients with reference to the effectiveness of isoniazid was carried out concomitantly with the above studies. It was concluded that acute exacerbations were not prevented by isoniazid and that the therapeutic effect of isoniazid remained to be proved.

Doctor Tower's group is continuing its studies into the epileptic cortex and glutamic acid synthesis with the utilization of carbon 14 labeled asparagine. A new method for isolation of amino acids in the spinal fluid by changing them to their dinitro-phenyl derivatives and subsequent separation by column chromatography has been developed in Doctor Tower's and Doctor Curtis' laboratories. Similar studies with aspartic acid and gamma aminobutric acid are underway. In their studies as to the clinical effects of glutamine and asparagine on generalized seizures, Doctors Tower and Wells have surveyed 45 patients, of which 23 remained on medication. Three examples of toxicity manifested by nausea and vomiting were noted. In addition to decrease of seizures, an increase in alertness and well-being was noted in the 23 remaining patients. They hope to continue their studies utilizing also enterically-coated alpha-ketoglutarate. Quantitation of the amount of these substances in the cerebro-spinal fluid plasma is proceeding.

Doctors Wells and Tower are combining with Doctor Sokoloff in the studies of the effect of ventilation in "petit mal" seizures by use of Kety's radioactive krypton⁷⁹ method. Doctors Wells, Tower, and Haase are studying the concentration of B-12 in the spinal fluid. Doctor Tower is to be a co-author of a book of neurological chemistry to be published within the next year.

Doctor Chatfield is continuing his studies of temperature and its effect on the myoneural junction. He finds the lower temperatures at lower frequencies will initiate tetanus in hamster muscle. He finds but little relationship to the amplitude to muscle response and the tension developed. Doctor Chatfield has finished a book on clinical neurophysiology which will be published by Charles C. Thomas and Company.

Doctor Irwin and Mr. Wells have continued their studies despite the protracted illness of Doctor Irwin. They have carried on studies in the cross transfused head technique in relationship to respiratory and vasomotor response to central nervous system asphyxia. They find that the anterior spinal artery of the dog may carry enough blood flow to keep such centers functioning when all other blood supply is eliminated. They are continuing their studies in calcium metabolism in relation to neuromuscular blocking agents. They find calcium antagonistic to succinyl choline and that this antagonism is independent of adrenalin release and occurs in the presence of acidemia during hyperpnea.

Under the direction of Doctor Baldwin and his group, studies in the physiology of the cortex and its deep nuclei are continuing with utilization of patients with focal seizures, in particular, the temporal lobe. It would appear that patients with temporal lobe lesions have primary difficulty in awareness, attention, and perceptual analysis. Four patients yielded distinct memory patterns on stimulation of the temporal lobe. Studies are underway to isolate the vestibular isolation in the temporal lobe and initial studies seem to indicate that nystagmus can be elicited only in posterior temporal sections. Stimulation of the amygdaloid nucleus has given rise to motor and postural changes, in particular, in the face and upper extremity. The most consistent change in the amygdaloid stimulation is loss of consciousness. A variety of autonomic effects are being recorded both in man and higher primates in the amygdala and insula. Secondary sensory areas have been found to extend in the island of Reil and in the first and second temporal gyrus. Changes in blood pressure, pulse, temperature, and gastric mobility have been recorded in stimulations of the island of Reil. A new apparatus has been prepared to stimulate peripherally for recording evoked potentials on the cortex. Apparently difficulty in speech may arise from lesions in either temporal lobe, the dominant temporal lobe being more aphasic in character and the less dominant having difficulty in the handling of semantics. There seems to be a direct relationship between psychotic episodes and epileptiform activity in the temporal lobe in the majority of patients. Stimulation of the deep nuclei of the temporal lobe in higher primates shows first a widening of the palpebral fissure, a dilation of the pupil on the side of stimulation, the salivation, and, finally, the posturing characteristic of this area.

Doctor Frost and his section seems to feel that memory per se is not necessarily affected in temporal lobe lesions in both man and primates but that the main difficulty seems to lie in the attention and concentration, and, in particular, to perceptual relationships to other individuals. The Psychology Section is also initiating a long-term study in the affects of anti-epileptic agents on intelligence. A study of tissue removed at cortical surgery seems to reveal at least five types of lesions: 1) a pseudolaminar necrosis; 2) complete cortical necrosis; 3) a deposition of a peculiar type of lipid that stains with Sudan IV but also with thionine and is not dissolved in paraffin preparations. The other two changes found at pathology are neuronal damage with astrocytosis and meningo-cerebral cicatrix. Surgical studies for the relief of pain are continuing, utilizing the polygraphic analysis of patient response.

Much time has been spent in the Branch of Surgical Neurology in further elucidating neuro-ophthalmological pathways and their pathology. Attempts have been made to plot ganglionic changes in the retina from geniculate lesions with considerable success by

Doctor Van Buren. Studies are continuing into the primary pathology of papilledema in which characteristic swelling of the nerve endings have been seen on the retinal side of the lamina cribosa. Such swellings separate the lamina cribosa from Bruch's membrane and the fibers are caught in a sharp angulation around Bruch's membrane and it can be seen that this might well permanently damage the optic fibers. The characteristic visual field of a temporal lobe lesion is not homonymous and the effects are larger and more dense ipsilateral to the lesion. Lesions in the temporal lobe do not extend in their retrograde regeneration to all six layers of the lateral geniculate and it is suggested that perhaps macular fibers lie deeper in the geniculate. The first case of human moniliiasis in the eye is being reported by Doctor Van Buren.

The Section of Neuropathology has recently lost its chief and recruitment for a new head of this Section is in progress. Doctors Alvord and Brace have finished their studies of x-ray induced lesions of the central nervous system and have reported this in London, England, under the title "X-Ray Induced Pyknosis of Cerebellar Granule Cells in Guinea Pigs." They have shown considerable protection to radiation by pre-administered nembutal. Doctors Alvord and Goldstein of the Institute of Mental Health have continued their studies on artificial demyelination using some 450 animals. That portion of the brain homogenate that seems responsible has been isolated to a water soluble product. Doctor Alvord and Doctor Whitlock have finished their work on the startle response and, in particular, to the necessity of the sensory-motor area to startle response under light chloralose.

Doctor Li has extended over the past year his studies utilizing micro-electrodes of less than one-half micron in diameter. Such studies are being carried out in the human operating room as well as in the primate operating room. The functional properties of cortical neurons with particular reference to strychnine as well as the effect of recording responses of evoked neuronal discharges are under study. More recently Doctor Li, by this technique, has demonstrated that the nucleus ventralis lateralis has an inhibitory effect on the premotor cortex.

Doctor von Sallmann is continuing his studies that he started before joining the Institute of a systematic exploration of thalamic and hypothalamic nuclei and their effect on ocular pressure. He had previously reported an increase of intraocular pressure on stimulation of the ventral thalamus and dorsal hypothalamus. He now finds an increased ocular pressure on stimulation of the posterior hypothalamus and medial ventral thalamus. Accompanying this is a rise in the subcleral temperature. He has ruled out the cervical sympathetic and the external ocular muscles as a cause for the increased pressure. Occlusion of the internal carotid artery will

drop this pressure but it will rapidly return to its preocclusion height. It is abolished by the drugs Priscoline and Thorazine. Since general body functions such as blood pressure and pulse have effects on internal ocular pressure, the Branch of Ophthalmology has devised a multichannel transducer apparatus to try and exclude as many of these factors as possible. The external ocular muscles will be excluded by electromyographic techniques. Studies as to the peripheral receptors responsible for internal ocular pressure are being undertaken in reference to the innervation of the anterior chamber with particular reference to the trabecular network and its relation to Schlemm's canal.

Studies in the origin of cataracts by ionizing radiation, diet, and drugs constitute another large project in the Branch of Ophthalmology. Galactose cataracts are being studied in which particular emphasis has been placed on the histology of whole lens mounts. Apparently changes in the lens epithelium and the peripheral fibers precede the actual degeneration in the lens and both precede opacities. The feeding of myleran or the withholding of tryptophane from the animal will also precipitate such cataracts and these two methods will be used in future studies. In the studies of ionization changes in the lens, the mitotic index of the lens epithelium may be used as an index of radiosensitivity. Such an index is apparently the highest in the monkey. The ionization changes in the lens are remarkably similar to the degeneration with age and may be of value to the study of senile cataracts. Doctor O'Rourke and his group are studying the effects of steroids in uveitis. Horse serum may readily precipitate uveitis. The extent of the uveitis may be determined by four criteria: 1) keratic precipitates; 2) aqueous cellularity; 3) aqueous beam; 4) iris hyperemia. Using such criteria values, steroid drugs may be tested. It has been found that one-fifth a dose of prednisone is necessary to equal that of the other corticoids. However, ultimate clearing of this uveitis is not shortened by such a drug. Side effects, however, are less with such lower doses. Doctor O'Rourke is continuing his studies on ocular melanoma and is closely associated with the instrument development section of the Oak Ridge Laboratories. His primary interest is to differentiate ocular melanoma from other intraocular tumors. A collimated scintillation counter has been developed to aid in this study and a strain of mice are utilized in which an S-91 malignant melanoma has been induced. The amount of escape of P-32 in external ocular muscles, scleral tissues, and in the globe will be studied. Doctor Barnschein, Doctor Wanco, and Doctor Iser are continuing studies on the electroretinogram, and, in particular, the effects of photic stimulation in the rise rate to study accommodation in the optic nerve. They are also studying hypothermia on the electroretinogram and controlling their temperature evaluation from the cheek-pouch. Doctor Tanaka and Doctor O'Connor are continuing their studies of inflammatory diseases of the eye, in particular, trachoma and the E.K.C. virus. The finding that the A.P.C. virus may grow on con-

junctional epithelial cultures may do much to aid the study of trachoma. Labeled antibodies are being utilized to study the predilection to ocular toxoplasmosis. Doctor Gunkel is continuing his instrument development section and is further refining his magnetic self-recording screen and methods for better evaluation of flicker fusion fields. Doctor Resnik and his staff are now studying the proteins of the lens by fractionation through a base ion exchange resin and then further separation by ultracentrifuge and electrophoresis. Doctor Resnik has finished his study on alloxan and showed that it does not form a complex with metals as previously determined.

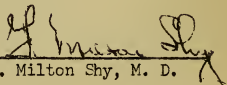
Doctor Kurland in the past year has continued his studies in amyotrophic lateral sclerosis. In an attempt to differentiate genetic from environmental factors, he carried on a study among the Chamorro's in California who migrated from Guam. It is Doctor Kurland's initial opinion that the incidence among the Chamorros will be as high as in their families back home. Doctor Kurland, in addition, went to Europe to the Archivo de Indias in Seville, Spain and to the records of the Jesuits in the Vatican Library. All the early references he could locate indicated that they were a healthy and long-lived group so his impression that amyotrophic lateral sclerosis was introduced by the Spaniards had been strengthened. This was further supported by the possibility that one of the Spanish islands off the mainland had an incidence much above normal. At the present Doctor Kurland is working in the Armed Forces Institute of Pathology where he is reviewing the cases of amyotrophic lateral sclerosis and related disorders to further understand such disease and in hopes of finding some lead to direct his search for the hypothetical underlying metabolic defect.

Doctor Ajmone Marsan, in addition to his heavy clinical responsibilities, has carried out, along with Doctor Rolston, a clinical electroencephalographic correlation of metrazol-induced seizure patterns. The MIS faithfully reproduced the patients' attack pattern in 85% of the cases and in 90% of the spontaneous attacks observed while off medication. Motor lateralizing signs are present in an unexpectedly high number of temporal cases. While visceral or non-visceral automatisms may be found in sites other than the temporal lobe, if the attack consists of automatisms alone the EEG is strikingly characterized by lack of changes or by unitemporal activation. If there is automatism plus aphasia, the EEG changes are usually bilateral temporal. Other correlations can be noted in project number 91. Doctor Ajmone Marsan is continuing his investigations in animals and the recruiting response in cortical and subcortical structures. He finds a progressive activation of two elements differing by the number of synapses, waning indicates a drop out of neurons of the shortest path. The recording of non-specific responses from a surface cortical electrode and from electrodes at different levels from within the cortex and underlying white matter and basal ganglia with a similar latency but with recruitment and waxing and waning seems to indicate that such

events must occur within the thalamus. Doctor Reiston and Doctor Ajmone Marsan are studying the effects of locally applied penicillin to the thalamic nuclei and the projection of the abnormal response. They also find that penicillin placed in the mesial thalamus and cingulate gyrus activates the spindle response.

With the calling together of scientists throughout the Country interested in cerebral palsy, Doctor Bailey has initiated a broad research program in this relatively little understood field. A preliminary protocol was formulated by this Committee which will guide the efforts of both intramural and extramural research teams. Most of the pathological material will ultimately pass through the National Institute of Neurological Diseases and Blindness where it will be reviewed by Doctor Anatole Dekaban, who has recently joined the Institute. Doctor Dekaban has, in addition, initiated a large pre- and post-natal study in cooperation with the Army and Navy Centers of Washington, with reference to various factors that may induce cerebral palsy. He is, in addition, working on the embryology of the mouse brain in anticipation of studying the effects of radiation at various stages of fetal growth and their correlation with lesions of the central nervous system. The Cerebral Palsy Program intramurally has been further strengthened by the addition of a pediatric neurologist, Doctor Gordon Millichap, who has been studying the effects of relaxant drugs on involuntary movements and rigidity. He finds initially that both may be increased with Reserpine or decreased with Flexin (2-amino-5-chloro-benzoxazole). In addition, he is studying the effects of carbonic anhydrase inhibitors on the cerebral seizures of childhood. He has initiated a study of the incidence of precocious puberty in cerebral palsied children.

These, then, are some of the specific studies that have been undertaken by the Clinical Investigation Unit of the National Institute of Neurological Diseases and Blindness. 39 manuscripts have been prepared or have been published during the past year, and 3 books are in the process of publication.


G. Milton Shy, M. D.

CLINICAL INVESTIGATION UNIT

PUBLICATIONS - 42

	<u>Manuscripts</u>	<u>Books</u>	<u>Total</u>
MEDICAL NEUROLOGY BRANCH:			17
Neurological Disorders	6	2*	
Neurochemistry	7	1	
Neurophysiology	1	1	
SURGICAL NEUROLOGY BRANCH:			11
Neurosurgical Disorders	5		
Neuropathology	1		
Experimental Neurosurgery	5		
CEREBRAL PALSY:	2		2
OPHTHALMOLOGY:	6		6
ELECTROENCEPHALOGRAPHY:	1	1	1
EPIDEMIOLOGY:	5		5
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	39	3	42

* Contributors from Surgical Neurology and Basic Research Unit.

TABLE I

Table of Organization of Clinical Investigations Personnel
(Personnel On Hand December, 1955)

Office of Clinical Director:

Clinical Director -- Dr. G. Milton Shy, M.D., M.S.,
M.R.C.P.
Administrative Assistant -- Mr. Elliot Brookman, B.A.
Secretary -- Mrs. Catherine Thomas
Secretary -- Mrs. Rae Myers

Branch of Medical Neurology: Dr. G. Milton Shy, Acting Chief
Neurological Disorders Service:

Clinical Associate -- Dr. Gunter Haase, M.D. Assoc. Prof. U. Oklahoma
Clinical Associate -- Dr. Kenneth Magee, M.D. Assoc. Prof. Michigan U.
Clinical Associate -- Dr. Glenn Drager, M.D. Assoc. Prof. Texas U.
Visiting Scientist -- Dr. J. G. Millichap, M.D. Mayo Clinic
Secretary -- Mrs. Aileen Poole
Secretary -- Miss Joyce Jenkins
Physical Science Aid -- Mr. William Matthews

Clinical Neurophysiology Section:

Neurophysiologist -- Dr. Paul Chatfield, M.D., Chief '55
Secretary -- Miss Dorothy Haase
Physical Science Aid -- Mr. Jeu S. Ming

Biophysical Applications Section:

Acting Chief -- Dr. G. Milton Shy, M.D., M.S., M.R.C.P.

Clinical Applied Pharmacology Section:

Pharmacologist -- Dr. Richard Irwin, Ph.D., Chief
Physiologist -- Mr. Jay B. Wells
Clerk-Typist -- Miss Alice Louise Cecil

Clinical Neurochemistry Section:

Neurochemist -- Dr. Donald Tower, M.D., Chief
Muscle Chemist -- Dr. Beni Horvath, M.D., Ph.D.
Chemist -- Dr. William Curtis, Ph.D.
Physical Science Aid -- Mr. John Phoenix
Physical Science Aid -- Mr. Edmund Peters
Clerk-Stenographer -- Mrs. Virginia Gray
Physical Science Aid -- Mr. Joseph Proctor
Research Associate -- Dr. Charles Wells, M.D.¹ 54

Surgical Neurology Branch: Dr. Maitland Baldwin, Chief

Neurosurgical Disorders Service:

Neurosurgeon -- Dr. Maitland Baldwin, M.D., M.S.,
F.R.C.S., Chief
Neurosurgeon -- Dr. John Van Buren, M.D., M.Sc. 54
Neurosurgeon -- Dr. Edward Laskowski, M.D.
Clinical Associate -- Dr. William Headley, M.D.?
Nurse -- Miss Karma Eastman
Secretary -- Miss Florence E. Beyer
Clerk-Stenographer -- Mrs. Maxine Reynolds
Clerk-Stenographer -- Mrs. Florence Heller
Supv. Histopath. Technician -- Mr. Fred Meiller

Experimental Neurosurgery Section:

Neurosurgeon -- Dr. Choh-luh Li, M.D., Chief
Med Bio Tech -- Mr. Alan Rowe
Physical Science Aid -- Mr. Phillip McGrath
Physical Science Aid -- Mr. Clarence Gilkes
Physical Science Aid -- Mr. Ulysses Robinson
Lab Animal Caretaker -- Mr. Otis Prince
Med Bio Tech -- Mr. Norman Mills
Animal Caretaker -- Mr. George Duvall

Experimental Neurosurgery Section (Cont.):

Med Bio Tech -- Mr. James Burch
Clerk-Stenographer -- Miss Vera Douglas
Nurse -- Miss Shirley Lewis

Clinical Psychology Section:

Psychologist -- Dr. Laurence Frost, Ph.D. St
Psychologist -- Miss Mildred Blevins

Clinical Neuropathology Section:

Acting Chief -- Dr. John Van Buren, M.D., M.Sc. St48
Histopathology Technician -- Mrs. Gladys Moore
Histopathology Technician -- Mr. Daniel Williams
Histopathology Technician -- Mr. Eustace Bourne
Clerk-Stenographer -- Mrs. Mary Di Pietro

Section on Embryological Neuropathology:

Embryologist -- Dr. Anatole Dekaban, M.D.
Histopathology Technician -- Miss Marie Kendall
Clerk-Stenographer -- Mrs. Grace Moore

Ophthalmology Branch:

Chief -- Dr. Ludwig von Sallmann, M.D.
Secretary -- Mrs. Frances Dearman
Clerk-Stenographer -- Mrs. Delma Snouffer
Clerk-Stenographer -- Mrs. Rose Daniele

Ophthalmological Disorders Service:

Clinical Associate -- Dr. James O'Rourke, M.D., Chief
Clinical Associate -- Dr. Gilbert Iser, M.D. ?
Clinical Associate -- Dr. George Goodman, M.D. ?
Clinical Associate -- Dr. Monte Holland, M.D. ?
Clinical Associate -- Dr. George O'Connor, M.D. ?

Ophthalmological Disorders Service (Cont.):

Clerk-Typist -- Mrs. Harriett Murphy

Ophthalmology Pharmacology Section:

Pharmacologist -- Dr. Frank Macri, Ph.D., Chief

Ophthalmology Physiology Section:

Visiting Scientist -- Dr. Hans Bornschein, M.D.

Visiting Scientist -- Dr. Theodo Wanko, M.D.

Physicist -- Dr. Ralph Gunkel, O.D.

Chemist -- Miss Patricia Grimes

Physical Science Aid -- Mr. Humphrey Patton

Ophthalmology Chemistry Section:

Chemist -- Dr. Robert Resnik, Ph.D., Chief

Chemist -- Mrs. Ann Wolff

Chemist -- Mr. Frank Suggs

Ophthalmology Bacteriology Section:

Visiting Scientist -- Dr. Chie Tanaka, M.D.

Bacteriologist -- Miss Mary Jane Cook

Physical Science Aid -- Mr. James Mitchell

Ophthalmology Histopathology Section:

Physiologist -- Mr. Leo Caravaggio

Histopathology Technician -- Mrs. Eleanor Collins

Branch of Electroencephalography:

Chief -- Dr. Cosimo Ajmone-Marsan, M.D., Ph.D.

Research Associate -- Dr. Bruce Ralston, M.D.

Technician -- Miss Maureen Benson

Technician -- Miss Carol Caswell

Technician -- Miss Helen Torrence

Secretary -- Mrs. George Ann Johnson

Physical Science Aid -- Mr. Samuel Cooper

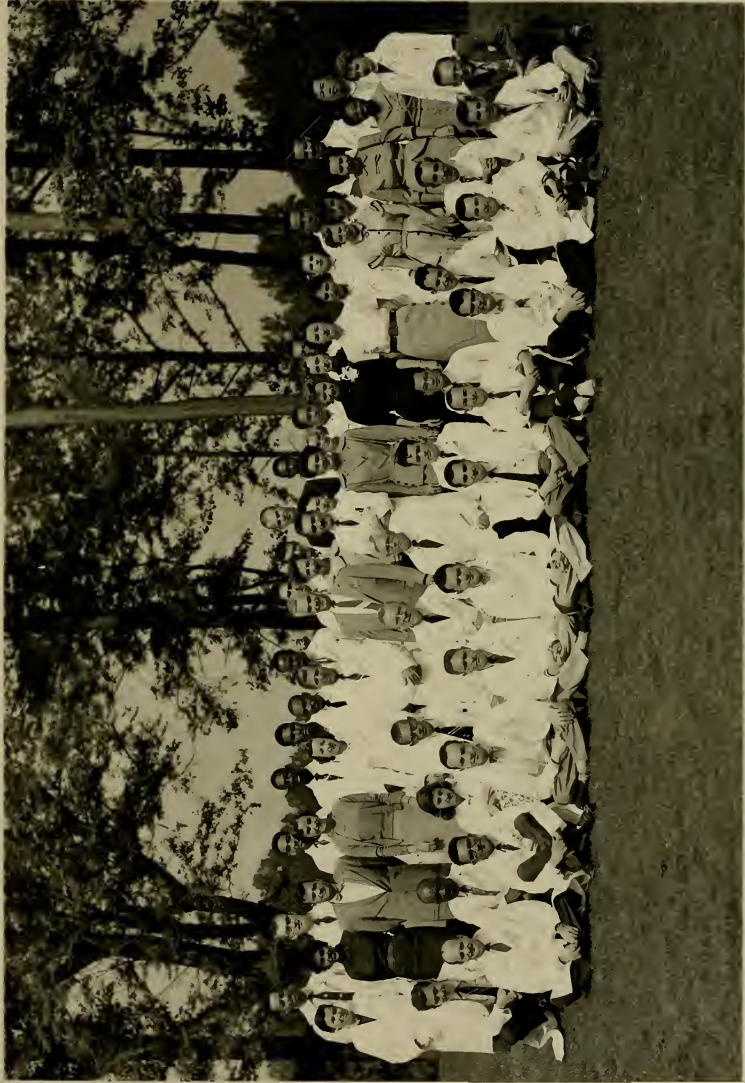
Assoc
Prof
U. Chi. Ill.

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Epidemiology Branch:

Epidemiologist -- Dr. Leonard Kurland, M.D., Chief

Clerk -- Miss Mary Kay



1955





Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Medical Neurology
BRANCH
3. Neurological Disorders Service
SECTION
4. - LOCATION
5. NINDB-1 (C)
SERIAL NO.
6. Studies in the Pharmacology of Myasthenia Gravis and Studies
PROJECT TITLE
in Prostigmine Tolerance
7. G. Milton Shy, M. D.
PRINCIPAL INVESTIGATOR
8. L. P. Rowland, M. D., Leonard Berg, M. D. and Richard L. Irwin, Ph.D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objective: In the treatment of myasthenia gravis, one item of essential information has long been lacking, and to a certain extent, a matter of controversy. It is known that in many myasthenics assistance cannot be afforded by large amounts of prostigmine, and, in others, symptoms may become so severe that the patient dies despite the administration of large amounts of this drug, which is presumably almost specific treatment for the disease. It has been the opinion of some investigators that this occurs merely because the patients are receiving an insufficient amount of prostigmine. Others have held the opinion that the patients are actually refractory to the drug; and some have feared to give large amounts of the drug because of possible adverse affects on muscle strength; i.e., the patient might become weaker rather than stronger. Because critical data on this problem are lacking, the following investigation is contemplated and underway.

Method employed: To myasthenic patients with varying degrees of severity of the disease, injections of prostigmine are given over a period of several hours. At first, only small amounts are used and this is gradually increased. This is continued until the patient achieves an optimal response in terms of increased strength. Concomitantly the likelihood for annoying side effects are observed. At this point, the injection is gradually increased to see how long the optimal state can be maintained and what happens beyond this point. Correlated with this study are measurements of serum cholinesterase and evaluation of the response of muscle to nerve stimulation.

Patient Material:

		<u>No.</u>	<u>Avg. Stay Days</u>
Admissions:	Adult males	2	23
	Adult females	4	135.8
	Children males	-	
	Children females	-	
Outpatients:	Number of patients	0	
	Number of visits	0	

Major findings: Recently this group of investigators have attempted to correlate all known information on myasthenia gravis into a plausible scheme and to attempt animal experimentation and subsequent human experimentation to demonstrate the plausibility of this scheme. If one searches for a depolarizing type of agent such as decamethonium in the myasthenic patient instead of nondepolarizing agent such as curare, then all research work in myasthenia gravis can be fitted into one scheme. Repetitive dosages of decamethonium will change the block from a depolarizing type of block to a competitive block. Some muscles are more sensitive to this than are others and these are primarily the muscles affected in myasthenia gravis. In muscles not so affected, however, an increased tolerance to decamethonium occurs as has been demonstrated by Churchill-Davidson in England, whereas those muscles affected to the point that a competitive block exists will become worse. It would be at this stage that myasthenia gravis would present. At this stage prostigmine will reverse the competitive block and we have done such in the laboratory. If one increases the competitive block to the point that only 5% of the twitch potential remains, then prostigmine or tensilon will not reverse this block. However, results in this laboratory demonstrate that hexamethonium will. At the point at which a depolarizing blockade becomes a competitive blockade, tetanus will no longer be held and will increase the blockade as shown in this laboratory. Curare itself acts as a competitive blockade. Further studies, then, must show if hexamethonium will act against curare as well as against decamethonium. If so, this will be the drug of choice in myasthenia gravis. This would also explain why the muscle of a myasthenic not affected by the disease is not resistant to decamethonium while those affected are not susceptible. It will also explain why some myasthenics react at some stages to prostigmine and not to others. Extracts of the thymus also give a blockade which resembles the depolarizing blockade and not a curarizing blockade.

Approximately 35 cats and 62 rats have been studied as to the affects of decamethonium blockade. From these studies it would appear that the decamethonium series rapidly changes from a depolarizing block to a competitive block which is indistinguishable from curare. At this point prostigmine will alleviate either a decamethonium block or a curare block as will hexamethonium.

It is therefore apparent that the only way to solve this problem is direct electrical recording off a single motor end plate and a single muscle fiber. This required complete retooling of the laboratory, which has now been accomplished, and with the aid of Doctor Paul Fatt from England we have been able to record motor end plate potential on the isolated sartorius muscle of the frog, from the muscles of the intact frog, and from the muscles of the intact rat. This has been the first time that motor end plate potentials have been recorded from the intact animal. The next stage will be recording from myasthenic muscle of the human patient.

Significance to neurological research: Primary studies on a few patients indicated that in some patients, at least, adverse symptoms will appear before myasthenia symptoms can be completely relieved, even when an attempt is made to control these side effects by the administration of specific antidotes. Among the adverse symptoms which have been noted is the decline in muscle strength, even greater than the weakness prior to the administration of prostigmine. If further study can corroborate these findings or further analyze them, that should have significant implications for the treatment of the disease and may cast some light on the nature of the disease.

Direct electrical recording from the motor end plate and its muscle fiber will be the only way that direct evidence may be found for a competitive or depolarizing substance in the myasthenic patient.

Proposed course of the project: This project will become the major research effort of the medical branch of this Institute in the coming year, and beds will be made available for myasthenia gravis patients if the initial studies are successful.

R.P.C. 2
December 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB-1 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH

13. NONE.

14. NONE.

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955:

Rowland, L.P., Korengold, M.C., Jaffe, I.A., Berg, L. and Shy, G. M. Prostigmine-Induced muscle weakness in myasthenia gravis patients. Neurology, Vol. 5, No. 2, February, 1955.

Rowland, Lewis P. Prostigmin-responsiveness and the diagnosis of myasthenia gravis. Neurology, Vol. 5, No. 9, September, 1955.

17. HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Medical Neurology
BRANCH
3. Neurological Disorders Service
SECTION
4. _____
LOCATION
5. NINDB-2 (C)
SERIAL NO.
6. Ionic Exchange in Neuromuscular Disease
PROJECT TITLE
7. G. Milton Shy, M. D.
PRINCIPAL INVESTIGATOR
8. Beni Horvath, M. D., Leonard Berg, M. D., and Donald Cummings
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objective: Proposed plan to determine the effect of potassium metabolism, in particular, intracellular concentrations of potassium in neuromuscular disorders, by the use of tracer elements and flame photometry and the utilization of specific pharmacological agents.

Method Employed: Clinical material will be subjects with muscular dystrophy, myasthenia gravis, myotonia, and myositis. Biopsies will be removed from such subjects under major nerve block. Our modification of the Boyle-Conway technique will be used for extracellular space. This procedure will continue to be controlled by conscientious objectors. Animal experimentation will be continued in rabbits, rats and guinea pigs, the experimentation on goats having now been terminated.

Patient Material:

	<u>No.</u>	<u>Avg. Stay Days</u>
Admissions: Adult males	11	31.3
Adult females	5	38
Children male	7	30.9
Children females	0	
Outpatient: Number of patients	0	
Number of visits	-	

Major Findings: Within the last year this unit has established a technique whereby the sodium and potassium of residual muscle and the amount of potassium exchanged in residual muscle can be determined in various diseases. In muscular dystrophy it has been demonstrated that potassium is low and sodium is high, and a regression curve may be drawn between the two. In spite of this, however, the potassium turnover of microcuries per milliequivalent

9. PROJECT DESCRIPTION (Continued)

is within the range of normal demonstrating this result to be the effect and not the cause of the disease. In myasthenia gravis this technique has been utilized to demonstrate that the defect in potassium lies in an inability to enter the cell or an excess leakage from the cell. The first has been related to the anticholinergic group of drugs and the second to the choline acetylase system. Initial results of this were reported at the Myasthenia Gravis Foundation Conference.

1) Residual muscle tissue of patients with muscular dystrophy contains an elevated sodium and reduced potassium concentration. Such concentrations may be correlated with the decrease of residual fiber water. There is a significant correlation between the lowering of the potassium concentration and the elevation of the sodium concentration.

2) The utilization of iodinated albumin demonstrates that the dystrophic patient either fractionates protein within his blood vessels or abnormally fractionates it from a vessel whose permeability is increased. Thus, the dystrophic patient may show concentrations up to 50% by using iodinated albumin, whereas such concentrations will be 8 to 10% when Chromium 51 tagged red cells are used. Since the same muscle that shows 50% vascularity with iodinated albumin shows but 10 neqs. of potassium per kg., this in reality then could not be explained on increasing vascularity since blood alone would furnish 25 neqs. of potassium. Further studies to find what happens to protein in muscular dystrophy are being carried out.

3) A method is described for the measurement of the amount of potassium exchanged in 22 hours in the whole human body. A procedure is presented to estimate the rate of potassium turnover in human muscle. By using the first of these methods muscular dystrophy patients appear to turn over less potassium in 22 hours than control subjects. The index of the potassium turnover of the muscle as measured by the second method is independent of the amount of potassium exchanged in 22 hours in the whole body. The index of potassium exchanged in dystrophic muscle is in the same range as normal muscle.

Significance to Neurological Research: It is anticipated that the role of potassium exchange is much more important to the general problem of myasthenia, myotonia and muscle wasting than the individual analysis of the potassium level of serum or urine at any one time. The relationship extends to the field of endocrine and metabolic disorders as well as specific neuromuscular disorders.

Proposed course of project: No more studies of ionic concentration or ionic exchange are contemplated. Further studies of the fate of injected protein in dystrophic patients will be carried out.

Analysis of NIH Program Activities

Project Description

- | | | |
|---|---|-------------------------------------|
| 1. <u>Neurology</u>
INSTITUTE | 2. <u>Medical Neurology</u>
LABORATORY OR BRANCH | |
| 3. <u>Neurological Disorders Service</u>
SECTION | 4. _____
LOCATION | 5. <u>NINDB-3 (C)</u>
SERIAL NO. |
6. An Investigation of the Relationship Between Blood and Spinal Fluid Glucose.

PROJECT TITLE

7. Leonard Berg, M. D.
PRINCIPAL INVESTIGATOR
8. G. M. Shy, M. D. and Donald Cummings
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Project: An investigation of the relationship between blood and spinal fluid glucose.

Objectives: 1. Elucidation of mechanisms of glucose transfer across blood-fluid barriers, of glucose utilization in brain and spinal fluid in normal and disease states.

2. Elucidation of details of cerebrospinal fluid function.

Methods Employed: The quantity of glucose and its degradation products in the spinal fluid will be followed during experimentally produced changes in the blood glucose. Experiments will be performed on suitable laboratory animals in their normal state as well as during various kinds of meningitis. A new experimental method, namely the use of radioactive glucose to follow glucose metabolism of spinal fluid, will be introduced after preliminary utilizing non-radioactive techniques.

Major Findings: Methods have been developed for the identification, elimination and isolation of glucose in plasma and spinal fluid. Isolation of the compound was necessary primary to the tracing of radioactive glucose because of the obvious necessity of separating the parent compound from radioactive degradation products for counting purposes. Thus far the work has been confined to in vitro experiments. The methods and yields appear to be satisfactory.

Significance to Neurology Research: It is expected that information will be gained relative to general problems of cerebrospinal fluid physiology. Advances in knowledge about mechanisms and treatment of meningitis and hydrocephalus are goals of this project.

Proposed Course of Project: Terminated because of investigator leaving Institute.

Analysis of NIH Program Activities

Budget Data

10. NINDE-3 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH.

13. NONE.

14. NONE.

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955:

None

17. HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

Analysis of NIH Program Activities

Project Description

1. Neurology
INSTITUTE
2. Medical Neurology
LABORATORY OR BRANCH
3. Neurological Disorders Service
SECTION
4. _____
LOCATION
5. NINDB-4 (C)
SERIAL NO.
6. Experimental Reduction of Reflex Activity in Special Disease States
PROJECT TITLE
7. M. G. Korengold, M. D.
PRINCIPAL INVESTIGATOR
8. _____
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Project: Experimental Reduction of Reflex Activity in Special Disease States.

Objectives: To determine response of reflex activity in specially prepared laboratory animals to various drugs in an attempt to achieve a reduction in this activity.

Methods Employed: With the use of suitable laboratory animals, experimental transection procedures are performed. Correlations are to be made between state of their reflex activity as influenced by various neuromuscular blocking agents. Of particular interest will be response to various curare and synthetic curare-like preparations, such as d-tubocurarine, repository tubocurarine, flakedil, succinyl choline, metubine, and syncurine. A study will be made of effects of such medication upon the reflex activity, as well as any associated side reactions involving other areas of the animal.

Major Findings: None.

Significance to Neurological Research: It would be of considerable value to learn more about the use of blocking agents in diseases in which spasticity is a major disability. This would be of value in elucidating further information about the nature of tone and factors that might influence it.

Proposed Course of Project: No change.

R.P.C. 2
December 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB-4 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH.

13. NONE.

14. NONE.

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955:

None

17. HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

Analysis of NIH Program Activities

Project Description

1. Neurology
INSTITUTE
2. Medical Neurology
LABORATORY OR BRANCH
3. Neurological Disorders Service
SECTION
4. _____
LOCATION
5. NINDB-5 (C)
SERIAL NO.
6. Protein Studies in the Hereditary Ataxias.
PROJECT TITLE
7. M. C. Korengold, M. D.
PRINCIPAL INVESTIGATOR
8. I. Jaffe, M. D. and C. Wells, M. D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Project: Protein studies in the hereditary ataxias.

Objectives: To determine the qualitative and quantitative protein components of the cerebral spinal fluid and their relationship to serum proteins and other examinations of protein normalcy.

Methods Employed: Patients are to be suitably screened in the Out-Patient Department for disease consistent with the diagnosis of Cerebellar Ataxia. These people will then be admitted as in-patients and blood studies consisting of total and fractional proteins, cephalin flocculation, and thymol turbidities will be run. Spinal fluid will be examined for its differential protein content by means of paper electrophoresis. Similar studies will be run on serum by means of paper electrophoresis, so that adequate comparisons can be made on the significance of protein abnormalities appearing in either area.

<u>Patient Material:</u>	<u>No.</u>	<u>Avg. Stay Days</u>
Admissions: Adult males	6	34.2
Adult females	2	25
Children male	1	6
Children female	-	
Outpatient: Number of patients	2	
Number of visits	2	

Major findings: Patients with hereditary cerebellar ataxia demonstrate the highest amount of gamma globulin in the spinal fluid that we have as yet determined. Since the presence of an increased gamma globulin in the spinal fluid has been stated to be specific for multiple sclerosis by some medical centers and since the average case of cerebellar ataxia is diagnosed by the uninitiated as multiple sclerosis, it would appear that these findings are of importance in two ways: 1) to show that an increase in gamma globulin is not specific to multiple sclerosis; and, 2) that this is the single neurological disease which gives the largest quantity of gamma globulin. The reason for the latter is not clear and a proposed course of this project will be toward elucidating protein metabolism in this disorder.

Significance to Neurological Research: To assist in learning new information about a system disease that may prove of value in correlating protein abnormalities in this condition with chemical abnormalities found in other demyelinating states. This may lead to previously undescribed abnormalities that may be of value in better understanding the nature of this disease and assist in providing a clue for possible future therapy.

Proposed Course of Project: A proposed course of this project will be toward elucidating protein metabolism in this disorder.

other diseases to be relatively easily recognized. In such cases, studied by many investigators over several decades, the areas of disagreement generally concern those of more precise etiology and pathogenesis rather than of observation and interpretation of clinical and microscopic abnormalities.

In many cases, however, there is a discrepancy between the clinical and histological observations. In certain of these, if the histological observations are clear-cut, and if the clinical observations are not unequivocal, one may well accept the histological diagnosis, recognizing that there are exceptions to any clinical rule and that the microscope might well reveal changes which otherwise would not be recognizable. Thus, distal muscular syndromes include not only the usual "neurogenic" muscular atrophies and dystrophia myotonica but also "inflammatory" and "degenerative" forms of "myogenic" disease. Furthermore, among the cases with proximal muscular weakness and wasting there are a few cases of "neurogenic" muscle atrophy, especially amyotrophic lateral sclerosis and "flabby infants", and of dystrophia myotonica.

In certain other cases, however, the discrepancy between the clinical and histological observations is less easily resolved. In our experience this is true of the proximal muscular syndromes in which an attempt is made histologically to distinguish between "inflammatory" and "degenerative" disorders: i.e., between various forms of "muscular dystrophy" and "myositis". We have not been able to reach any satisfactory solution to this difficult problem. At times one wonders whether the muscle biopsy is really helpful, but this pessimistic view cannot long be maintained: one cannot help but feel that further study of more cases will disclose the answers. In this situation we have found that the microscopist preferably should not know the clinical impressions: we have repeatedly seen how remarkably difficult it is to record unbiased observations if the clinical diagnosis is known. If accurate observations can be made, recorded, carefully analyzed and continuously reviewed, our present ignorance and confusion will be replaced by understanding as the result of continued investigations.

Significance to neurological research as can be seen from the results: Many controversial fields are present in the correlation of clinical disease and muscle pathology. This study has clarified the points listed above, but in addition, has raised additional controversies. An original classification of diseases of the lower motor neuron has been offered. An atlas with some seventy colored prints will be available in March to aid other investigators in their studies of these diseases.

R.P.C. 1 (b)
Dec. 1955

NINDB -6 (c)
SERIAL NO.

Proposed course of the project: The proposed course of this project will reside around the second objective: the elucidations of particular pathological changes associated with neuromuscular disease. In particular, this will reside in the fields of Polarizing Microscopy and Electron Microscopy. This is of particular interest in the proximal myopathies where structural detail may still be present with changes in the birefractility of the muscle fiber.

Analysis of NIH Program Activities

Budget Data

10. NINDB (C-2)
SERIAL NO.

11. Budget Data

12. BUDGET ACTIVITY: Research

13. NONE

14. NONE

Honors, Awards, and Publications

16. Publications

Atlas of Muscle Pathology. E. & S. Livingstone, Edinburgh,
England.

17. Honors and Awards

NONE

December 1955

Analysis of NIH Program Activities

Project Description

1. Neurology
INSTITUTE
2. Medical Neurology
LABORATORY OR BRANCH
3. Neurological Disorders Service
SECTION
4. _____
LOCATION
5. NINDB-7 (C)
SERIAL NO.
6. Study of the short-term effect of INH on the course of Multiple Sclerosis.
PROJECT TITLE
7. GeoR. Haase, M. D. and M. C. Korengold, M. D.
PRINCIPAL INVESTIGATORS
8. Kenneth R. Magee, M. D. and Leonard Berg, M. D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Project: Study of the short-term effect of INH on the course of Multiple Sclerosis, which were described as dramatic in a recent study. (The Effects of Isoniazid on Patients with Multiple Sclerosis, Kurtzke and Berlin, The Am. Rev. of Tuberculosis, Vol. 70, No. 4, October, 1954.)

Methods Employed: This was a clinical study. Patients who were clearly afflicted with multiple sclerosis were followed as in- and Outpatients. The total patient group was divided into two sub-groups, which were matched as nearly as possible. Employing the "double-blind method", one sub-group received INH while the control group received a placebo. Both groups were re-examined frequently to determine any alterations in the patient's status. The criteria for inclusion in the study included a clear-cut dissemination of lesions within the central nervous system and a dissemination in the onset of various unrelated symptoms in time. Such patients in whom the diagnosis was questionable were excluded from participation in the study.

<u>Patient Material:</u>	<u>No.</u>	<u>Average Stay</u>
Admissions: Adult males	7	90 days
Adult females	4	90 days
Children male	-	-
Children female	-	-
Outpatient: Number of patients	70	
Number of visits	200	

R.P.C. 1(a)
December 1955

NINDB-7 (C)
SERIAL NO.

Major Findings: Acute exacerbations were not prevented by the use of INH and beneficial effects of INH in multiple sclerosis were not proved.

Significance to Neurological Research: A major research project is at present underway in the Veterans' Administration based on the work of Kurtzke and Berlin as to the effects of INH in multiple sclerosis. This study was the first controlled double-blind study and does not support the findings of Kurtzke and Berlin.

Proposed course of Project: Terminated.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
2. Medical Neurology
BRANCH
3. Neurological Disorders Service
SECTION
4. -
LOCATION
5. NINDB-8 (C)
SERIAL NO.
6. Recording of Motor End Plate Potentials of Mammalian Muscles
PROJECT TITLE
Specifically in Normal and Myasthenic Man
7. G. Milton Shy, M.D., Choh-Luh Li, M.D., Paul Fatt, M.D.
PRINCIPAL INVESTIGATORS
8. Mr. Jay Wells
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objective: To determine if substances are present that compete with or block release of myoneuronal humoral substances in myasthenia gravis, or if depolarization exists.

Method employed: Micro-electrode recording from within the single muscle fiber by means of the technique described by Fatt and Katz will be utilized. The lumbricalis muscle will be the one recorded. The ulnar nerve will be stimulated in the forearm and the intramuscular potential recorded through a .5 micron diameter glass pipette led through a cathode follower. The motor end plate potential and the action potential will be recorded by an AC beam of the Dumont Scopè. The resting potential muscle will be recorded on a DC beam.

Patient Material: None to date.

Major Findings: Motor end plate potentials have been successfully recorded from curarized muscle of frog and rat to date, as well as the resting potential.

Significance to neurological research: If the substance causing myasthenia gravis is a curarizing material, then a given end plate potential would demonstrate no spectrum in it's amplitude. If the difficulty is in the release of acetylcholyne then a spectrum should be seen. The presence of a motor end plate potential without the simultaneous firing of the muscle is indicative that one of these mechanisms is in action.

Proposed course of project: A number of patients will be admitted with selective weakness to isolated muscles which will render recording from such muscles easy when nerves supply in such muscles are stimulated at some distance. If isolated motor end plate potentials are found, then various drugs will be given intra-arterially to try to change the electrical recording.

R.P.C. 2
Dec. 1955

Analysis of NIH Program Activities

Budget Data Sheet

10. NINDE-8 (C)
SERIAL NO.

11. BUDGET DATA

12. BUDGET ACTIVITY: Research

13. None

14. None

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955: None

17. HONORS AND AWARDS TO PERSONNEL DURING 1955: None

R.P.C. 1
Dec. 1955

Analysis of NIH Program Activities

Project Description Sheet

- | | | |
|---|---|-------------------------------------|
| 1. <u>Neurological Diseases and Blindness</u>
INSTITUTE | 2. <u>Medical Neurology</u>
LABORATORY OR BRANCH | |
| 3. <u>Neurological Disorders Service</u>
SECTION | 4. <u>-</u>
LOCATION | 5. <u>NINDS-9 (C)</u>
SERIAL NO. |
| 6. <u>Studies in Myoclonus.</u>
PROJECT TITLE | | |
| 7. <u>Charles E. Wells, M. D.</u>
PRINCIPAL INVESTIGATOR | | |
| 8. _____
OTHER INVESTIGATORS | | |
| 9. <u>PROJECT DESCRIPTION:</u> | | |

Objective: To further elucidate the nature of myoclonus, especially in regard to its clinical, electrical, and pathologic aspects; to assess the value of various drugs upon the myoclonus; and especially to establish this as a clear entity separate from other forms of abnormal movements.

Method employed: Patients will be studied by means of electromyography and electroencephalography, in addition to extensive clinical descriptions and evaluations. Movies will be made where possible to correlate clinical and pathologic findings.

Major Findings: Two patients with myoclonus have been studied within the past year by the methods described above correlating electroencephalographic and electromyographic changes with the clinical findings. In one of these patients, complete pathological correlation has also been possible.

Significance to Neurological Research: The subject of myoclonus is marked in the literature by a mass of unclear descriptions and indistinct terminology. It is hoped that a systematic study of the condition will lead to its clear definition as a clinical entity.

Proposed course of Project: It is proposed to study as many patients as possible with this condition during the next year, with the wish that perhaps five patients will be available for study.

R.P.C. 2
Dec. 1955

Analysis of NIH Program Activities

Budget Data Sheet

10. NINDB-9 (C)
SERIAL NO.

11. BUDGET DATA

	EST. EXPEND.	BUDGETED POSITIONS			MAN YEARS			PATIENT DAYS
		PROF.	OTHER	TOTAL	PROF.	OTHER	TOTAL	
FY 1956								
FY 1957								

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
REVIEW & APPROVAL TECHNICAL ASSISTANCE

-
13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO.

None

-
14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE IDENTIFY SUCH PROJECT.

None

-
16. PUBLICATIONS FROM THIS PROJECT DURING 1955:

None

-
17. HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

Analysis of NIH Program Activities

Project Description

1. Neurology
INSTITUTE
2. Medical Neurology
BRANCH
3. Neurological Disorders Service
SECTION
4. _____
LOCATION
5. NINDB-10 (C)
SERIAL NO.
6. Clinical Studies on Amyotrophic Lateral Sclerosis.
PROJECT TITLE
7. Leonard T. Kurland, M. D. and Kenneth R. Magee, M. D.
PRINCIPAL INVESTIGATOR
8. _____
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objective: A long term study to evaluate possible metabolic abnormalities in amyotrophic lateral sclerosis and to evaluate various agents as to their therapeutic efficacy in this disease.

Methods Employed: This will be a clinical study. A series of approximately 10 patients with amyotrophic lateral sclerosis will be admitted.

In addition to routine blood and cerebrospinal fluid studies, attempts will be made to determine the blood and cerebrospinal fluid amino acids to seek any abnormality. This will be done by the methods of paper chromatography.

Considering the possibility that amyotrophic lateral sclerosis may be a deficiency disease due to the failure of absorption of some vital food element from the stomach, one or more of the patients will be given repeated blood transfusions and parenteral amino acids and vitamins in an attempt to provide all necessary dietary elements without depending on gastric absorption.

Additional attempts at therapy will be made but their nature will depend on preliminary results of the investigation.

Major Findings: None.

Significance to Neurological Research: Amyotrophic lateral sclerosis is universally fatal. As very little is known concerning its etiology and no treatment exists, attempts to further our knowledge regarding this disease are of the utmost importancy and constitute one of the greatest needs of Neurology.

R.P.C. 2
December 1955

10. NINDB-10 (C)
SERIAL NO.

-
11. BUDGET DATA:

-
12. BUDGET ACTIVITY: RESEARCH.

13. NONE.

14. NONE.
-

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955:

None

17. HONORS AND AWARDS TO PERSONNEL DURING 1955:

None



Project Description Sheet

1. N.I.N.D.H. 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clinical Neurochemistry 4. _____ 5. NINDB-11 (C)
SECTION LOCATION SERIAL NO.
6. Glutamine and Glutamic Acid Metabolism in Normal and Epileptogenic Cerebral Cortex
PROJECT TITLE
7. Dr. Donald B. Tower
PRINCIPAL INVESTIGATOR
8. Dr. Maitland Baldwin, Dr. C. Ajmone-Marsan, Dr. Ellsworth Alvord, Dr. W. C. Curtis, Mr. John Phoenix, Mr. Edmund Peters
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION

- (a) Objectives of this project are to study the in vitro metabolism of glutamine and glutamic acid by slices of cerebral cortex from (1) human patients operated for focal epileptogenic cortical lesions and (2) experimental animals.
- (b) Methodology: Methods are essentially the same as those in the previous report on this project (1954-NINDB-15 (c)).
- (c) Patient material is obtained from NINDB patients admitted to the Surgical Neurology Branch and operated on for excision of focal epileptogenic cortical foci. Since some of these excisions require removal of overlying or adjacent uninvolved cortex, a small number of non-epileptogenic samples is being accumulated.
- (d) Major Findings: During calendar year 1955 studies on the defect in metabolism of glutamic acid by epileptogenic cerebral cortex have continued. One sample of human temporal cortex became available in which it was possible to follow glutamic acid metabolism in an epileptogenic focus and in an adjacent area of uninvolved cortex together with electrocorticographic and histological controls for each. Thus, in the same cortical area from a single patient the difference between epileptogenic and non-epileptogenic cortex could be clearly demonstrated:

Sample	Tissue Glutamic Acid		Tissue Glutamine		Tissue Ammonia	
	Level	Change	Level	Change	Level	Change
Non-Epileptogenic						
Initial	7.35		2.2		2.5	
60 min.	10.35	+3.0	3.75	+1.55	3.05	+0.55
Focal-Epileptogenic						
Initial	7.3		2.85		2.15	
60 min.	6.0	-1.3	4.2	+1.35	2.95	+0.8

These results are consistent with numerous other experiments on the respective samples obtained individually from different patients. No differences histologically could be demonstrated between the two samples despite the pronounced electrocorticographic and biochemical differences, confirming the concept of a biochemical lesion.

In 1954 it was shown that among others L-asparagine added to such tissue slices in vitro would correct this and other defects in epileptogenic cortex. These results suggested that L-asparagine could be metabolized by mammalian brain tissue. With the exception of asparaginases, known to be present in serum of herbivores (especially guinea pig) and in kidney and possibly liver of various mammalian species, and the recent finding by Meister that liver enzymes catalyzed transamination reactions with asparagine, no data is available to support the above contention.

¹⁴C-labelled asparagine has been obtained and is being utilized to study this possibility further. Considerable difficulty has been encountered in obtaining labelled asparagine. Through the cooperation of the Process Research Department, Merck and Co., a small amount of DL-asparagine-2, 3-¹⁴C was synthesized from ¹⁴C-maleic anhydride. When chromatographically pure material was incubated with slices of normal cat cortex under the usual experimental conditions, metabolism of asparagine could be clearly demonstrated. The results of this experiment suggest that D-asparagine cannot be utilized since about half the total radioactivity remained as asparagine by chromatography. The remainder was distributed, in order of descending activity, in aspartic acid, glutamic acid, glutamine, and possible traces in alanine and γ -aminobutyric acid, as shown by two-dimensional chromatography with radioautographs and actual counts of activity of each amino acid spot by a gas-flow Geiger counter. Since a 4-carbon compound was added and 5-carbon compounds (glutamic acid and glutamine) showed appreciable activity after incubation of slices, metabolism of the asparagine clearly occurred.

Also in 1955 a new convulsant agent, MEG, a glutamic acid derivative, was made available to us by the Medicinal Chemistry department of Merck and Co. This convulsant has been found by them to give an ideal assay method, using mice, for various anti-convulsants, including glutamine and asparagine. It has been found by us that this convulsant will produce severe, immediate seizures in cats. A study of the glutamic acid metabolism of cerebral cortex from such cats showed a most interesting result - a complete disappearance of glutamic acid but maintenance of glutamine during incubation. Thus, we now have two convulsant agents which affect glutamate metabolism in opposite ways: methionine sulfoximine,

which inhibits glutamine synthesis, and MEG, which inhibits glutamic acid maintenance.

- (e) Significance of this project: Since seizures are a major concern of NINDB as well as a major neurological problem to the population in general, knowledge of the mechanisms underlying them are essential for development of better and more rational therapy. More fundamentally, knowledge of seizure mechanisms will provide many of the factors which are concerned with neuronal function and activity in general and thus be basic to many problems of neurophysiology and neurological diseases. The defect in glutamic acid metabolism in human epileptogenic brain tissue is one of several components of a biochemical lesion of this tissue, which represents the first such lesion of a major neurological disease to be uncovered.
- (f) Proposed course of project: Continuation of present studies is underway. It is proposed to apply the studies with C¹⁴-labelled asparagine to human epileptogenic and non-epileptogenic material and to the two convulsant preparations cited above. Such studies should assist materially in elucidating the nature of the metabolic reactions taking place.

In addition it is contemplated that a more precise method for quantitative determination of amino acids in such tissue slices can be adopted as a result of methods developed in another project (NINDB-) from this section. By conversion of amino acids in tissue samples to the dinitrophenyl derivatives, separating individual amino acids by column chromatography, eluting the individual acids and reading each fraction colorimetrically, not only will more precise results be obtained but determinations can be extended to aspartic acid, γ -aminobutyric acid, alanine and other amino acids which cannot at present be quantitated. The above procedure will be tried on slices incubated in the standard fashion. If successful, confirmation of data by previous assay methods will be sought and extension of the studies to other amino acids applied.

Budget Data Sheet

10. NINDB-11 (C)
SERIAL NO.

11. BUDGET DATA:

FY 1956	ESTIMATED EXPENDITURES	BUDGETED POSITIONS			MAN YEARS			PATIENT DAYS
		PROF.	OTHER	TOTAL	PROF.	OTHER	TOTAL	
		2	2	4	0.3	0.2	0.5	*
FY 1957		2	2	4	0.3	0.2	0.5	*

* Patient material used is derived from patients admitted to other NINDB projects.

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
REVIEW & APPROVAL TECHNICAL ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 OR 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM 1)

Research and Development Divisions, Merck & Co., Inc.
Research Division, Wallace & Tiernan Co., Inc.

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO. (S) IF WITHIN NIH).

None.

Honors, Awards, and Publications Sheet

15. NINDB-11 (C)
SERIAL NO.

-
16. LIST PUBLICATIONS OTHER THAN ABSTRACTS FROM THIS PROJECT DURING
CALENDAR YEAR 1955:

Tower, D. B. "The Neurochemistry of Seizures" in Progress in Neurobiology: Neurochemistry V.1, Edited by S. R. Korey and J. I. Nurnberger; New York, Paul B. Hoeter, 1955. In Press.

Tower, D. B. "Neurochemical Aspects of Pyridoxine Metabolism and Function," Am. J. Clin. Nutrition. In Press.

-
17. LIST HONORS AND AWARDS TO PERSONNEL RELATING TO THIS PROJECT DURING
CALENDAR YEAR 1955:

None.

Received of the Treasurer of the
Board of Education

the sum of \$100.00
for the year ending
June 30, 1875

for the purpose of paying the
salary of the Superintendent
of Schools for the year ending
June 30, 1875

in full for the year ending
June 30, 1875

for the purpose of paying the
salary of the Superintendent
of Schools for the year ending
June 30, 1875

1875
No. 100

Project Description Sheet

1. N.I.N.D.B. 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clinical Neurochemistry 4. LOCATION (IF OTHER THAN 5. NINDB-12 (C)
SECTION BETHESDA SERIAL NO.
6. Clinical Evaluation of the Value of Glutamine or Asparagine for the clinical control of Seizures.
PROJECT TITLE
7. Dr. Donald B. Tower
PRINCIPAL INVESTIGATOR
8. Dr. Charles E. Wells, Dr. C. Aimone-Marsan
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

(a) Objectives of this project are to study the effectiveness of two naturally-occurring amino acid amides, L-asparagine and L-glutamine, in controlling seizures in epileptic patients and to observe their effects on epileptiform activity in the electroencephalogram of suitable patients. This project had its inception in the findings on human epileptogenic cortex in vitro that significant, chronic defects in metabolism of acetylcholine, glutamic acid and electrolytes occur which can be reversed by addition in vitro to such tissue of either glutamine or asparagine, neither of which appear to affect normal in vitro metabolism nor to exhibit any effects when administered in large quantities in vivo to normal subjects.

(b) Methodology and Patient Material: See 1954 project report (NINDB-).

(c) Major Findings: A total of 45 patients have been studied. Of these five represent preliminary patients, 17 have been discontinued for various reasons, and 23 remain active in the project.

All common types of seizures are represented in the above totals. All types appear to respond in similar fashion to amide therapy.

Results to date can be summarized as follows:

	<u>L-asparagine</u>	<u>L-glutamine</u>	<u>Both</u>	<u>Total*</u>
<u>Total on Therapy</u>	36	10	6	40
Male	24	9	6	27
Female	12	1	0	13
5-15 yrs.	17	6	3	20
16-40 yrs.	18	5	3	20
<u>Duration of Oral Therapy</u>				
Over 2 yrs.	2	0	0	2
1-2 yrs.	5	2	0	7
6-12 months	9	3	2	14
3-6 months	11	2	0	13
0-3 months	8	2	0	10
<u>Evaluation (Over 2 mcs.)</u>				
Essentially Seizure-free	4	0	0	4(10%)
Significant Seizure Reduction	17	4	1	22(56%)
No Change	7	5	1	13(34%)
Worse	0	0	0	0

EEG Improvement	10	4	0	14(35%)

Discontinuations have been made for the following reasons: No change after 3 or more months: 7; Side reactions: 2, Surgery: 1; Uncooperative parents or patient: 7 - total 17.

No toxicity has been encountered for periods exceeding two years. Repeated laboratory examinations have failed to reveal changes in hematological picture, hepatic or renal function or in routine blood chemistries, except for the expected elevation in B.U.N. and N.P.N. A number of cases with various common skin diseases and a few cases with previous history of anticonvulsant drug allergies have been given asparagine without incident.

Unpleasant side reactions have been encountered in only 3 patients (7.5%); consisting of persistent nausea and vomiting. Transient reactions of this nature have occurred in a number of other cases but have not persisted with continuing medication.

In the majority of patients placed on either amide, increased alertness and general well-being has been spontaneously reported regardless of effect on seizure-control. Among the school-age improved group, resumption of or improvement in schooling has been general and home and social situations have been much improved. The older age-group improved patients have been able to broaden their activities, to resume or do better at college, and to seek employment actively. One patient

has done such outstanding work in college (straight A average) since starting on amide therapy that he has been recommended for a Fulbright scholarship to continue his studies in languages.

In several instances unintentional omission of amide therapy has resulted in a prompt relapse to previous seizure frequencies with return to control after resumption of amide doses. In one case in particular this was particularly striking - the patient was hospitalized elsewhere for an unrelated condition; his dose of asparagine in chocolate milk was reduced to one-fifth through a misunderstanding and without the patient's knowledge; daily major and minor seizures ensued whereas before and after this episode on full dosage the patient has remained seizure-free.

3 patients have received DL-asparagine for periods up to several months. Results have been equivocal. 3 patients have received supplemental pyridoxine (Vitamin B₆) orally without effect on seizure frequency. Administration of α -ketoglutarate was tried in one patient but could not be tolerated because of taste and gastric irritation. One patient not otherwise included in this project received L-glutamine (parenteral preparation) intrathecally without untoward reaction.

In the course of studies on this group of patients several subsidiary observations have been possible. One patient has been studied because of high serum CO₂ content and found to be spontaneously hypoventilating. Diamox, a carbonic anhydrase inhibitor, had no effect on this patient's seizure-frequency despite an adequate biochemical response to the drug. In several cases where no change in seizure frequency was obtained with amide therapy and where previous anticonvulsant medication had been ineffective, relatively massive doses of Tridione proved very effective in eliminating seizures. In one case with a behavior disorder Reserpine is under trial in conjunction with asparagine. Good seizure control had been obtained with the latter and addition of Reserpine did not affect the control and appears to be controlling behavior.

- (d) Significance of this project would seem to be self-evident. It represents a new type of compound which has definite anticonvulsant properties, but which differs from all other anticonvulsants not only in its chemistry but in its lack of toxicity and lack of sedative or hypnotic action. It represents development of therapy based on a demonstrated biochemical lesion in human epileptogenic brain tissue. The number of problem cases referred to NINDB or contained in inquiries from neurologists and physicians in this country and abroad indicate that better, more effective therapy for seizures is in demand. With over a million and a half seizure patients in this country alone, it is a major health and economic concern. In addition this project is representative of the type of research for which the clinical investigation phases of NIH were created in that it has taken laboratory findings, applied them to patient material at the Clinical Center in a detailed manner seldom possible elsewhere, developed agents of considerable initial cost to the point where cost is being lowered to more reasonable ranges, and promoted trials elsewhere (as noted below).
- (e) Proposed Course of the project is to continue studies now underway. It is not planned at present to add further patients to the present group except as special occasions may arise. The possibility of obtaining an enterically-coated form of α -ketoglutarate is to be explored

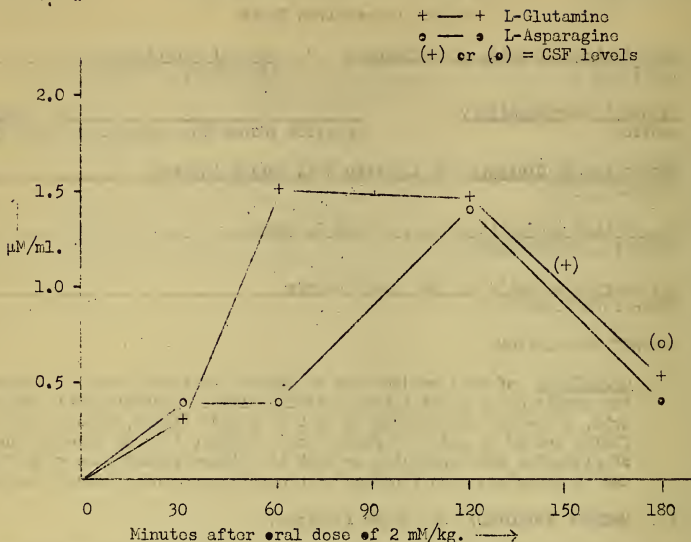
R.P.C. - 1e
December 1955

NINDB-12 (C)
SERIAL NO.

since it would be of interest to try this compound clinically because it is the immediate precursor of glutamic acid and might prove as effective as asparagine or glutamine. It is already produced synthetically and would ultimately be quite inexpensive if shown to be effective.

Much of the coming year will be devoted to studies of metabolism of asparagine and glutamine in these patients by methods developed in another project from this Section (NINDB-). These studies are designed to try to correlate dose with clinical response and to determine whether lack of response by some patients may be due to faulty absorption, too rapid elimination or the like.

AMIDE PLASMA "TOLERANCE" CURVES



Information obtained from determinations of the amide concentrations alone did not seem to offer as much value from a metabolic standpoint as determination of all possible compounds concerned. Therefore collection of patient data has been deferred pending modification of the method to permit simultaneous determination of glutamic acid, aspartic acid, glutamine and asparagine in the sample. This objective has been largely achieved. The method is as follows:

- (1) Sample: 1-3 ml. of blood plasma or spinal fluid are used. Blood is collected in heparinized tubes.
- (2) Deproteinization: The usual clinical methods of deproteinization were found to be unsatisfactory. Trichloroacetic acid could not be used because of its interference with subsequent steps in the procedure. Tungstate precipitation reduced recoveries to too great a degree. It was found possible to deproteinize the samples by passing the sample through a small column of Dowex-50 chromatographic cationic exchange resin. The proteins pass through while the amino acids are held on the column. The amino acids are then eluted with 0.8 N HCl in 55% ethanol and the eluant evaporated to dryness, using a Roto-Vac evaporator with mild vacuum and heating not over 55° C.

- (3) Conversion of Amino Acids to Dinitrophenyl Derivatives: The amino acid mixture is dissolved in a small volume of 2.5 N HCl and placed in an autotitrator unit in a water bath at 40°C. The solution is brought to pH 9.0 with alkali (usually NaOH). Excess fluorodinitrobenzene is added and the reaction continued for 80 minutes at pH 9.0 and 40°C with mechanical stirring. This procedure converts all amino acids and amines to their yellow-colored dinitrophenyl derivatives. Excess reagent is removed by extraction of the alkaline solution with ether.
- (4) Chromatographic Separation of Amino Acids: The aqueous solution is acidified with 1 ml. of 5N HCl and re-extracted with ether. The acid and neutral amine acids, including glutamic and aspartic acids, asparagine and glutamine, now go into the ether phase, leaving the basic amino acids in the aqueous phase. The ether phase is evaporated to dryness, using the Roto-Vac evaporator, and the residue taken up in 10 ml. of a 50-50 ether-chloroform mixture. This solution is then applied to a lightly packed Celite (diatomaceous silica) column, 40 cm. in length and 1 cm. in diameter buffered at pH 4.0 with phosphate buffer. 50-50 ether-chloroform is then run through the column. Six amino acids are held on the column. In their order from top to bottom they are: asparagine, glutamine, aspartic acid, serine, threonine and glutamic acid. All other amino acids in the acid fraction applied to the column are eluted in the first 10 ml. from the column.
- (5) Elution and Determination of Amino Acids: The six amino acids on the column are then eluted successively. Glutamic acid followed in order by threonine, serine and aspartic acid are eluted with 50-50 ether-chloroform and collected in individual fractions. Glutamine and asparagine are then successively eluted with 95-5 ether-chloroform. The six eluant fractions are evaporated to dryness and the residue taken up in 1% NaHCO₃ to a standard volume. The yellow-colored solutions are read at 360 m μ in a Beckman spectrophotometer against appropriate dinitrophenyl derivatives of each amino acid. The range of concentration which can be determined is 0.005-0.08 μ M (about 1-10 μ g.) in most cases.

The above procedure is to our knowledge the first which permits direct determination of glutamine, since all previous methods of this type involve hydrolysis of the glutamine to glutamic acid before determination. The above procedure is also to our knowledge the first to permit simultaneous determination of the four very closely related amino acid compounds, glutamic acid, glutamine, aspartic acid, and asparagine by a relatively simple, quantitative chemical method. Previous methods have generally involved the use of specific bacterial enzymes and manometric techniques, which are much more cumbersome and time consuming and less sensitive. The present method is also rather time consuming when all four compounds are determined, but can be speeded up considerably for determination of asparagine or glutamine alone.

The present method has a much wider application than that being used here. All amino acids in the original biological sample are

quantitatively converted to their dinitrophenyl derivatives. Three fractions are obtained in the procedure subsequently outlined: I. - basic amino acids, such as lysine, separated by the extraction with ether; II. - all acid and neutral amino acids except glutamic acid, aspartic acid, glutamine, asparagine, serine and threonine, separated by passage through the column without adsorption, and III. - the six amino acids held on the column. Fractions I and II could be separated by appropriate column and pH conditions into their individual components in a manner analogous to that used here for fraction III.

Many problems have been encountered in the development of this method which will not be detailed here. Because of the potential value of this method for many problems it was deemed essential to have every stage in the procedure thoroughly checked, reproducible and standardized for ready use by others. Therefore most of the time during calendar 1955 has been devoted to this phase of the project.

Preliminary runs by the above method of plasma samples from patients receiving oral asparagine therapy have been carried out. No significant changes in plasma glutamine or glutamic acid levels occur during the rise and fall of asparagine level (see graph above). Aspartic acid rises to a peak value about 25% of the asparagine peak value some 60 minutes after the asparagine peak, indicating some deamidation of asparagine during that period but not sufficient to account for the total asparagine determined.

- (e) SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: Significance of this project is in the interpretation of results being obtained in clinical therapy with glutamine and asparagine. With the use of any new compound for therapy it is important to be able to determine it in body fluids, to know the course of its metabolism and to be able to correlate dose, body level and clinical effect. With the quantities of amides being administered this becomes even more important.
- (f) PROPOSED COURSE OF THE PROJECT: The method developed is to be rechecked against the available microbiological methods for glutamic acid, glutamine and asparagine and then published.
- Patient samples (plasma and spinal fluid) will be run serially and in sufficient numbers to obtain the following data: (1) type or types of plasma and spinal fluid curves obtained from control patients given single oral doses of either glutamine or asparagine or both; (2) types of curves shown by seizure patients on daily doses of either or both compounds; (3) comparison of such data with clinical responses of patients; (4) determination of metabolic course of the compounds as reflected by plasma and spinal fluid changes.
- Urinary studies will be added to the above as soon as methods have been modified for this purpose. Semi-quantitation is already being obtained, using paper-chromatography of the dinitrophenyl derivatives of amino acids in the urine, but adaptation of the column method is being tried. Urine presents a difficult problem compared to other body fluids so that considerable work will be entailed in this phase.
- Finally adaptation of the method for tissue determination will shortly be instituted. Methods currently available and in use in this laboratory do not permit simultaneous quantitation of all amino acids present in the size of sample available, and for some of these amino acids no satisfactory quantitative methods are available. Such a method will be most useful in this project and in another project of this laboratory (NINDB-) concerned with brain amino acid metabolism.

December 1955

Analysis of NIH Program Activities

Project Description Sheet

1. N.I.N.D.B. 2. Medical Neurology
 INSTITUTE LABORATORY OR BRANCH
3. Clinical Neurochemistry 4. _____ 5. NINDB-14 (C)
 SECTION LOCATION (IF OTHER THAN SERIAL NO.
 BETHESDA)
6. Electrolyte Metabolism and Energy Metabolism in Normal and Epileptogenic Cerebral Cortex
 PROJECT TITLE
7. Dr. Donald B. Tower
 PRINCIPAL INVESTIGATOR
8. Dr. Maitland Baldwin, Dr. C. Aimone-Marsan, Dr. Ellsworth Alvord, Mr. Edmund Peters and Dr. Beni Horvath.
 OTHER INVESTIGATORS

9. PROJECT DESCRIPTION

- (a) Objectives of this project are to study the in vitro metabolism of electrolytes, sodium and potassium, and of energy-producing cycles and compounds by slices of cerebral cortex from (1) human patients operated on for focal epileptogenic cortical lesions and (2) experimental animals.
- (b) Methodology and Patient Material: See 1954 report (NINDB-18(C)).
- (c) Major Findings: Further studies in this project have been delayed pending development of suitable methods for obtaining data desired. Two problems are involved. The first concerns valid determination of tissue electrolyte concentration which, to be meaningful, must be expressed per unit of intracellular water. The latter requires measurement of the intracellular space for which no satisfactory method is currently available. We have learned that Engstrom in Sweden has developed but not yet reported an X-ray diffraction or absorption method which will accurately determine extra- and intracellular spaces in fresh tissue sections. We are therefore awaiting word on this possibility before resorting to the less reliable methods now available.

The second problem concerns the energy metabolism aspect of this project. The studies of Meister and co-workers have demonstrated that in purified enzyme systems, the synthesis of glutamine, which requires high-energy phosphate as ATP, is reversible (Ref. J. Biol. Chem. 209: 265, 1954). In this system, starting with glutamine, ATP is formed from ADP as glutamine is converted to glutamic acid. These experiments provide a more detailed concept of the situation which may exist in epileptogenic brain tissue. On incubation defects occur in metabolic systems which are energy-dependent. At the same

time glutamic acid is metabolized by the tissue, substantially reducing the level of this important intermediate. Correction of all defects occurs on addition of glutamine or asparagine. If Meister's results are applicable even to a limited degree, the addition of one or other amide could simultaneously replace metabolized glutamic (or aspartic) acid and provide extra ATP for cellular metabolism.

Two principal studies must be made in order to study the above possibilities. First the manner in which added glutamine or asparagine are metabolized by normal and by epileptogenic brain. Carbon-14 labelled amides as tracers for the addition can be expected to provide much of this information. Preliminary experiments with asparagine have already been carried out (see project NINDB-). However, in addition to amino acids, for which suitable chromatographic methods are available, the α -keto acid derivatives must also be examined since these provide the metabolic link between amino acids. Paper chromatography of the keto compounds has not been well worked out. During calendar 1955 much effort has gone into preparing authentic standard dinitrophenylhydrazones of all anticipated keto compounds and into determining suitable solvents for chromatography. When the studies are completed, it will be possible to trace metabolism, with the radioactive tracer, through the ketoacid compounds, as well as determining the amounts of such compounds by extraction of the chromatographically isolated spots.

The second study involves direct determination of ATP, ADP and Creatine phosphate levels in normal and epileptogenic brain tissue with or without added glutamine or asparagine. Specific muscle enzymes for each of these high-energy phosphate compounds have been developed in this laboratory for use in these studies.

- (d) Significance of this project is self-evident since it brings into these studies on epileptogenic cortex two factors, electrolytes and energy supplies, which are accepted as essential to neuronal activity and function, providing much evidence for the mechanism by which seizures can occur. If these findings can be clarified and extended, many of the missing links in the chain of circumstances underlining the seizure and concerned with neuronal activity in general can be set in place. This will represent a contribution not only to the problem of seizures but to neurophysiological mechanisms in general.
- (e) Proposed course: See under Major Findings.

Analysis of NIH Program Activities
Project Description Sheet

1. N.I.N.D.B. 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clinical Neurochemistry 4. _____ 5. NINDB-15(C)
SECTION LOCATION (IF OTHER THAN SERIAL NO.
BETHESDA)
6. Electrophoretic Studies of Human Cerebrospinal Fluid Proteins in
Demyelinating Diseases
PROJECT TITLE
7. Dr. William C. Curtis and Dr. Donald B. Tower
PRINCIPAL INVESTIGATORS
8. Dr. Charles E. Wells, Mr. John Phoenix and Mr. Edmund Peters
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

(a) Objectives of this project are to study the abnormalities in the protein fractions of cerebrospinal fluids from patients with multiple sclerosis in order to determine the significance of such abnormalities both diagnostically and prognostically in the clinical course of the disease.

(b) Methodology: Cerebrospinal fluid samples are obtained from NINDB ward or outpatients by lumbar puncture or during pneumoencephalography. Aliquots of these samples are examined for routine constituents.

Total protein content of the fluid samples is determined by refractometry or ultra-violet absorption or both. Each method is calibrated by Kjeldahl and gravimetric runs on spinal fluid samples.

Routine electrophoresis of all spinal fluid protein fractions is run by paper-strip electrophoresis or by micro-interferometric electrophoresis or both, as detailed previously (see 1954 NINDB-20 (C)). Where appropriate serum samples from the same patient are run simultaneously.

Fractionation of spinal fluid protein for individual study will be attempted in several ways: by electro-convection isolation, by counter-current distribution (if this can be adapted), and by column electrophoresis-chromatography. Separated fractions, especially gamma-globulins, will be studied by micro-interferometric electrophoresis for homogeneity and mobility. The separated fractions will also be studied by paper-strip electrophoresis with staining for associated lipid and carbohydrate by Sudan Black and Feulgen reagents respectively. Semi-quantitation of paper runs is carried out with a Spinco analytrol scanner.

- (c) Patient material: No patients have as yet been specifically admitted to this project. Patient material so far has been obtained from other NINDB project patients. As studies progress suitable patients with multiple sclerosis will be selected for serial study of their fluid patterns to permit correlation with the clinical state and progress of the disease.
- (d) Major findings: In calendar 1955 paper-strip studies begun in 1954 were continued. As a result of the study of some 60 patients, half of whom had multiple sclerosis, confirmation of the previous findings of Kabat et al (Ann. N.Y. Acad. Sci. 58: 613 (1954)) have been obtained. The percentage of fluids from cases of multiple sclerosis with elevated gamma globulins in both cases was about 65%. In both series early cases or cases with few and infrequent attacks the percentage of positive fluids was much lower. Serial sampling of given patients has tended to show persistence of elevation where such elevation was found initially. However, in both series elevated gamma globulin has been found in spinal fluids from cases with other neurological diseases, especially the hereditary cerebellar ataxias. Elevations are also known to occur in neurosyphilis and certain types of meningitis.

The results of this type of study clearly indicate that elevation of spinal fluid gamma globulin occurs in most patients with multiple sclerosis. However, this observation appears to be of little value diagnostically, since the elevation precedes clinical symptoms in too few cases to be of clinical use. It is also of little value prognostically since clear-cut variations in suitable selected patients have not been observed. And it is of little value etiologically since a number of apparently unrelated conditions show similar elevations and since the source of the gamma globulin is not indicated by such studies.

Thus, this approach appeared to offer little value in its continuation. After considerable effort it has still not been possible to demonstrate lipid or carbohydrate in association with spinal fluid protein fractions, although such are readily detected for serum by the same methods. It seemed, therefore, that a more basic approach to this problem is needed and the studies have been halted pending development of this phase (see below).

- (e) Significance of this project is self-evident, since it represents a detailed study into an established chemical abnormality in a disease, multiple sclerosis, which is a major problem not only from a public health standpoint but from a diagnostic, prognostic and therapeutic standpoint. Nothing is known about its course or the mechanisms underlying its cause. With this first clue of biochemical abnormality a step in the direction of covering these many problems can be made by such a detailed and long-term approach.

- (f) Proposed course: From serum and immunological studies it is now known that gamma-globulin is not a single entity but consists of a mixture of at least four to six types. The usual methods for demonstrating and isolating these types require too much protein to be applicable to spinal fluid studies. However, by newer physico-chemical techniques it seems possible that isolation of gamma-globulins from spinal fluid can be achieved in sufficient amounts to permit characterization by micro techniques.

Therefore, it is proposed that the following course of investigation be instituted during the coming year:

- (1) Perfection of methods for isolation of the gamma-globulin fraction from spinal fluid and serum, probably by electro-convection and/or column electrophoresis methods,
- (2) Comparison of (a) spinal fluid gamma-globulins with serum gamma-globulins from the same patient and (b) spinal fluid gamma-globulins from patients with and without elevations (ie, multiple sclerotics vs. non-multiple sclerotics), by various physico-chemical methods, primarily micro-interferometric electrophoresis, ultra-violet absorption, and if possible counter-current distribution. This phase is designed to determine whether serum and spinal fluid gamma-globulins are identical and whether the rise in gamma globulin in spinal fluids of multiple sclerotics is due to the same or different gamma globulin compared to that present in control patients.
- (3) With concentrated, isolated specimens of gamma-globulins paper-strip electrophoresis will be used to determine whether, with greater concentrations than those possible in usual spinal fluid runs, lipid and/or carbohydrate associated groups can be demonstrated.

It is possible that immunological methods may have to be resorted to to demonstrate differences between gamma-globulins if the above methods are not sensitive enough.

The above procedures will be applicable to other protein fractions as well. In the course of these studies isolation of al pha- and beta-globulins will also be undertaken for characterization by the same procedures.

Budget Data Sheet

10. NINDB-15 (C)
SERIAL NO.

11. BUDGET DATA

	ESTIMATED			BUDGETED POSITIONS			MAN YEARS			PATIENT
	EXPENDITURES	PROF	OTHER	TOTAL	PROF	OTHER	TOTAL	TOTAL	DAYS	
FY 1956		3	2	5	0.3	0.2	0.5		*	
FY 1957		3	2	5	0.3	0.2	0.5			

* Patient material used is derived from patients admitted to other NINDB projects.

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
REVIEW & APPROVAL TECHNICAL ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 OR 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM I)

National Institute of Arthritis & Metabolic Diseases-Dr. Saroff.

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS); IDENTIFY SUCH RESEARCH: (BY SERIAL NO(S) IF WITHIN NIH).

None

Project Description Sheet

1. N.I.N.D.B. 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clinical Neurochemistry 4. _____ 5. NINDB- 16 (C)
SECTION LOCATION SERIAL NO.
6. Pentosuria in Neuromuscular Disease
PROJECT TITLE
7. Dr. Donald B. Tower
PRINCIPAL INVESTIGATOR
8. Dr. G. Milton Shy, Mr. Edmund Peters, Mr. Milton Pogorelskin
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION

(a) Objectives of this project are to assess the importance, if any of pentosuria in patients with primary muscle disease and to identify the quantity and type of pentoses being excreted.

(b) Methodology: See previous report (1954-NINDB-19(C))

(c) Patient material is obtained from NINDB ward patients. Neuro-muscular disease patients are admitted to Medical Neurology for other studies and urine specimens provided for this project. Other patients with no muscle disease serve as controls and are derived from other NINDB ward patients with eye diseases, multiple sclerosis, seizures and the like.

(d) Major Findings: This project was completed during the first half of calendar 1955 and is to be published shortly. The results of the project can be summarized as follows:

(1) The quantitative excretion of free aldopentoses in 230 twenty-four hour urine specimens from 55 patients with and without muscle disease were studied by the para-bromaniline method. All patients were on fruit-free diets during study.

(2) The mean excretions of total free aldopentoses per 24 hours per kg. body weight were: (a) by patients with no muscle disease 3.6 μM for adults and 5.5 μM for children (under 15 years of age); and (b) by patients with myopathies 6.6 μM for adults and 9.6 μM for children. Differences between means of the two groups of adults or of children were statistically significant in both cases.

- (3) Identities of the pentoses excreted were established by 29 two-dimensional paper chromatograms of 24-hour urine specimens from 18 patients. Eight carbohydrate compounds were identified. The majority of specimens contained glucose, xylose, arabinose and ribose. A few exhibited spots identified as lactose, galactose, desoxyribose and a uronide (?glucuronic acid). No ketoses were detected in regular or concentrated specimens with the possible exception of fructose. No differences in the patterns of carbohydrates excreted were apparent between the group with and that without muscle disease.
- (4) Semi-quantitation of aldopentoses excreted was carried out by the spot area method on one-dimensional chromatograms of 14 24-hour urine specimens from 12 patients. Arabinose predominated in specimens from the group with no muscle disease, whereas ribose appeared to be significantly higher in specimens from the group with muscle diseases. Xylose appeared to be about the same in both groups.
- (5) The findings suggest that patients with muscular dystrophy, myotonia dystrophica, myasthenia gravis and muscular atrophy (?cause) excrete significantly higher amounts of free aldopentoses in 24 hours, largely accounted for as ribose, than patients with diseases such as multiple sclerosis, spinal cord disease, cerebral seizures and uveitis.
- (6) No correlation of these findings with clinical status of the muscle disease patients (based upon duration of disease, rate of progression, degree of muscle involvement and disability, histopathology, and electromyography) could be established.
- (7) Good correlation of μM pentoses excreted 24/hrs/kg. body weight with residual functional muscle mass (as measured by potassium exchanged per kg. body weight (Ke/Kg) obtained in a separate study (see 1954-NINDB-3(C)) was demonstrated for 22 adult patients for whom both sets of data were available. The coefficient of correlation was statistically highly significant. This finding suggests that, as residual functional muscle mass is reduced, the excretion of pentoses, mostly as ribose, increases in the urine.
- (8) Thus, the results obtained in this project relate pentosuria in muscle disease to the result of muscle wasting, probably related to the breakdown of nucleoproteins and nucleic acids and the inability of the muscle to salvage the ribose so liberated.
- (e) Significance of this project is primarily to establish definitively whether or not there is an abnormal degree or type of pentosuria in patients with primary muscle disease. Clues to the possible causes or mechanisms underlying muscular dystrophy in particular are virtually non-existent. It is important, therefore, to reach a conclusion about pentosuria, particularly since the literature is

now confused and incomplete on this point. If there is a significant abnormality here, it will be of great importance and warrant extension to studies of blood and muscle. On the other hand definite negative evidence will be of service in directing efforts elsewhere and clearing confusion in the literature. Since neuromuscular diseases are one of the principal programs of NINDB, this project is important as one phase of that study.

- (f) Proposed Course: This project is completed. No further studies are contemplated at this time.

Budget Data Sheet

10. NINDB-16 (C)
SERIAL NO.

11. BUDGET DATA

	ESTIMATED EXPENDITURES	BUDGETED POSITIONS			MAN YEARS			PATIENT DAYS
		PROF	OTHER	TOTAL	PROF	OTHER	TOTAL	
FY 1956	None	0	0	0	0	0	0	0
FY 1957	None	0	0	0	0	0	0	0

12. BUDGET ACTIVITY

RESEARCH ADMINISTRATION
REVIEW AND APPROVAL TECHNICAL ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 OR 1957; IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM 1)

None

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO(S) IF WITHIN NIH).

None

Project Description Sheet

1. Neurological Diseases and Blindness 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
Clinical
3. Neurochemistry 4. _____ 5. NINDB- 17 (c)
SECTION LOCATION (OTHER THAN BETHESDA) SERIAL NO.

6. Comparative Biochemistry of Smooth Muscle & Striated Muscle Actomyosin
PROJECT TITLE

7. Beni Horvath, M.D.
PRINCIPAL INVESTIGATOR

8. Mr. Joseph Proctor
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION:

- (a) Objectives: To characterize the uterine actomyosin in physico-chemical terms, preliminary to a study of actomyosin synthesis in the uterine muscle.
- (b) Methods Employed: Myosin and actin will be prepared from rabbit uterine muscle and striated muscle. Yield will be compared at different stages of development of the uterus. The myosin and actin so obtained will be characterized by viscosity, electrophoretic mobility, sedimentation constant etc. as well as by its reaction with ATP.
- (c) Major Findings: For the purpose of comparison between the two kinds of muscle an extraction procedure had to be established and standardized giving maximal yield for both type of muscles. The method of Dr. Albert Szent-Gyorgyi, originally devised for extraction of heart muscle, was found to give excellent yields of actomyosin from small samples of skeletal muscle also. This method has the added advantage over the method of Dr. Csapo employed earlier, that the extraction of the homogenized muscle takes only a few minutes rather than overnight. The new method uses Potassium Iodide instead of Potassium Chloride, in an alkaline solution.
- (d) Significance to the Program of the Institute: It is felt that studies of formation of myosin and actin may have a relationship to location of disease in muscular dystrophy. Quantitative extraction procedures are also essential for the study of proteins in Neuro-muscular Disease.

- (e) Proposed Course of Project: Adaptation of the above extraction procedure to the uterine muscles of rabbits, and isolation of actin and myosin from the actomyosin obtained by the method of A.G. Szent Gyorgyi. As the ovarian hormones estrogen and progesterone have a marked effect on the development of the uterus, the effect of these hormones will be studied on the time course of the protein synthesis in the uterine muscle, using the quantity of myosin and actin as an indicator.

Project Description Sheet

1. NINDB Clinical 2. Medical Neurology
3. Neurochemistry SECTION 4. LOCATION (IF OTHER THAN BETHESDA) 5. NINDB- 18 (C) SERIAL NO.
6. Alterations of Actomyosin Tensile Strength and Muscle Proteins in Neuromuscular Disease
PROJECT TITLE
7. Dr. Beni Horvath
PRINCIPAL INVESTIGATOR
8. Dr. G. Milton Shy and Dr. William C. Curtis
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION:

- (a) Objective: To determine alterations in tensile strength of Actomyosin threads from biopsy material of patients with neuromuscular disease. To estimate the quantity of Actomyosin in diseased muscles and to characterize in physicochemical terms the proteins of diseased muscles as compared to normals.
- (b) Methods: Muscle fibers obtained from patients afflicted with neuromuscular diseases were collected and preserved in glycerol at low temperature for future analysis. The tension developed by these fibers was measured in a specially built apparatus. The tension was studied as a function of the Potassium, Magnesium and ATP concentration of the medium. Samples of normal muscle were compared with samples of myotonic muscles and with samples of muscles suffering from muscular dystrophy and with muscle samples afflicted with dystrophia myotonica.
- Other samples also obtained on biopsy were collected and stored in the frozen state. These samples were extracted by the method of Dr. Szent Gyorgyi originally developed for the heart muscle employing an alkaline solution of Potassium Iodide. The actomyosin so prepared was studied in the form of threads. These threads contract in a suitable environment on the addition of ATP. Again the effect of the above variables was studied under identical conditions on the actomyosin prepared from muscles suffering from the above diseases and compared to normal actomyosin.
- (c) Patient Material: Muscle sample obtained from patients afflicted with neuromuscular diseases, under nerve block anesthesia.

- (d) Major Findings: On the glycerinated fibers it was observed that for normal muscles there is only a narrow range of ATP concentrations where the fibers develop maximal tension. On myotonic fibers and on fibers from muscles with myotonic dystrophy this range appears to be appreciably wider. Muscular Dystrophy fibers now being studied seem to resemble the myotonic ones more than the normals in the early stage of the disease. The effect of the other variables is less pronounced.

The same widening of the range of the optimal ATP concentration was also observed in the shortening experiments carried out on the threads prepared from actomyosin of the myotonic muscles.

- (e) Significance: No change from NINDB -22(C) 1954.
- (f) Proposed Course: Experiments along both lines will be extended to include other diseased states. Since it appears that myotonia may involve some alteration of the actomyosin molecule it seems to be desirable to characterize myosin and actin prepared from myotonic muscles by physico-chemical methods.

Project Description Sheet

1. N.I.N.D.B. 2 Medical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clinical Neurochemistry 4. _____ 5. NINDB-19(C)
SECTION LOCATION SERIAL NO.
6. The Nature and Characterization of Oxyproteic Acids, Pyrogens and
Similar Molecules
PROJECT TITLE
7. Dr. William C. Curtis
PRINCIPAL INVESTIGATOR
8. Dr. Donald E. Tower, Mr. John Phoenix, Dr. G. M. Shy & Dr. W. Windle
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION

- (a) Objectives of this project are to develop suitable physico-chemical methods for the isolation and characterization of oxyproteic acids, pyrogens and similar molecules. The chemical and physical natures of these compounds are poorly understood and their significance and mechanisms of action remain obscure.
- (b) Methodology: Isolation and purification of oxyproteic acids from urine and of the pyrogen, Piromen, will be carried out by ion exchange, column chromatography, foam fractionation and absorption techniques. Characterization studies will employ electrophoresis, diffusion, ultracentrifugation, sedimentation, velocity and equilibrium, selective adsorption and solubility studies to provide data on molecular size and structure.
- (c) Patient material will not be used for this project initially except for urine specimens from NINDB patients admitted to other projects.
- (d) Major Findings: In preliminary experiments, abnormally high concentrations of oxyproteic acids were isolated from the urines of patients with muscular dystrophy. Such patients also showed an abnormal loss or metabolism of injected human serum albumen tagged with radioactive I¹³¹ (see NINDB-). Since oxyproteic acids are thought to be intermediates in the breakdown of albumen, these two findings may be closely related and of significance to the disease.

More detailed studies have been delayed pending delivery of equipment on order for this project.

- (e) Significance of this project: This is a more basic type of study designed to shed some light on certain complex molecules. The oxyproteic acids occur normally in urine as products of in vivo degradation of proteins, especially albumen. No detailed study or characterization of these compounds has ever been reported. In neuromuscular diseases there is degradation of nerve and muscle tissue, apparently associated with increased excretion of oxyproteic acids. A comparison of excretion by normal persons and those with muscle disease in regard both to quantity and quality would be of considerable significance.

Similarly, pyrogens have been shown to be associated with inhibition of glial scarring in central nervous system regeneration. The physical and chemical properties of such compounds are poorly understood and their mechanisms of action remain obscure. Elucidation of some of these factors should prove of value not only in pyrogen therapy but also in the more fundamental problem of fever mechanisms.

- (f) Proposed Course: To initiate full scale studies outlined under methodology.

- (e) Significance of this project is most important and fundamental. No satisfactory chemical method for determination of microgram quantities of acetylcholine in biological specimens has ever been developed. Bioassay procedures are tedious, somewhat non-specific and often not sufficiently sensitive. In view of the importance of acetylcholine as a peripheral and also probably as a central transmitter of nerve impulses, there is a pressing need for better assay methods. This need is reinforced by the recent development by Udenfriend of a very satisfactory chemical method for Serotonin, a probable second transmitter in the central nervous system. Positive chemical identification of acetylcholine in nervous tissue has only been reported twice in 35 years. There is need for re-assessment of acetylcholine with a specific chemical method particularly as more and more possible transmitters are proposed.
- (f) Proposed Course of the project is to study the reaction in detail using a variety of flavonals (rutin, morin, quercetin, quercetrin, etc.) to determine its sensitivity, reproducibility and specificity. Depending upon the result of these studies adaptation of the method will then be attempted. The use of fluorimetric measurement of the final acetylcholine reaction product may require a red-sensitive fluorometer, which has been recently developed by Dr. Udenfriend (N.H.I.).

Project Description Sheet

1. N.I.M.H. and N.I.N.D.B. INSTITUTE 2. Lab. Research (NIMH) and Medical Neurology (NINDB) LABORATORY OR BRANCH
3. Section on Cerebral Metabolism (NIMH)
Section of Clinical Neurochemistry (NINDB)
SECTION
4. _____
LOCATION (OTHER THAN BETHESDA)
5. NINDB-21 (C)
SERIAL NO.
6. Cerebral Circulation in Minor Seizures.
PROJECT TITLE
7. Dr. Louis Sokoloff (NIMH) and Dr. Charles E. Wells (NINDB)
PRINCIPAL INVESTIGATORS
8. Dr. Donald B. Tower (NINDB) and Dr. C. Aimone-Marsan (NINDB)
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

(a) Objective: To measure cerebral blood flow in patients with minor cerebral seizures (petit mal type) during hyperventilation and during the occurrence of seizures.

(b) Methodology: Cerebral blood flow studies will be performed using the radioactive Krypton (Kr^{79}) method devised by Lewis, Sokoloff, Wentz, Wechsler, and Kety (Fed Proc. 14:92 (1955)) and used extensively by Dr. Sokoloff during 1954-1955 at the U.S. Naval Air Development Center, Johnsville, Pa.

Concentrations of radioactive Kr^{79} , not exceeding $100\mu\text{C./L.}$, will be breathed from a closed spirometer reservoir by the subject through a tightly fitting mask for a period not exceeding 20 minutes. With this method readings over the cerebral hemispheres with an extensively shielded scintillation counter will be possible about every 30 seconds. Simultaneously arterial and cerebral venous blood samples will be drawn for Kr^{79} content. From these data and the Fick principle cerebral blood flow can be calculated.

Cerebral blood flow will be determined by this technique in the resting state, during hyperventilation and during seizures induced by hyperventilation, photic stimulation or other suitable means. Simultaneous electroencephalograms will be recorded wherever possible.

Arterial and cerebral venous blood samples will be analyzed for $p\text{CO}_2$, CO_2 content and pH and for oxygen saturation during these studies.

- (a) Patient Material: Patients with suitable seizures will be admitted to NINDB wards for this study.
- (d) Major Findings: This project will begin in calendar 1956. The procedure has been studied and perfected over several years at the Johnsville Air Development Center under Dr. Sokoloff's direction and the radiation problems and hazards carefully worked out and approved at that laboratory. The proposal has been submitted to the Clinical Center Radiation Committee for approval.
- (e) Significance of this Project: The nitrous oxide method for measurement of cerebral blood flow requires a 10 minute observation, i.e. averaging of flow/min. over a 10 minute period. It is known from this method that hyperventilation in normal subjects and patients with generalized seizures (inter-ictal) results in decreased cerebral blood flow and that generalized seizures induced in monkeys result in marked increases in cerebral blood flow. No studies have been reported on effects of hyperventilation in patients with minor seizures (petit mal type) although it is this group which is most susceptible to the ictogenic effect of hyperventilation. In addition the relationship between cerebral blood flow and seizures has not been studied during actual seizures in man because of the time limitation imposed by the nitrous oxide method. With the radio krypton method accurate recording of such information would be of great fundamental significance.
- (f) Proposed Course: To implement the above project.

Project Description Sheet

1. Neurological Diseases and Blindness 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
Clinical
3. Neurochemistry 4. _____ 5. NINDB- 22 (C)
SECTION LOCATION (OTHER THAN BETHESDA) SERIAL NO.
6. Investigation of Vitamin B-12 Levels in Spinal Fluid
PROJECT TITLE
7. Dr. Charles E. Wells
PRINCIPAL INVESTIGATOR(S)
8. Dr. Guenther R. Haase and Dr. Donald B. Tower
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

- (a) Objective: To determine the normal spinal fluid levels of Vitamin B-12 in health and disease and to determine whether when given in large doses the vitamin is concentrated in the spinal fluid.
- (b) Methodology and Patient Material: Clinical material will be normal volunteer subjects and patients with a variety of neurological diseases (especially diabetes with neuropathy, pernicious anemia with combined system disease and the other peripheral neuropathies). Spinal fluid B-12 levels will be measured in these patients to determine the usual levels in normal patients and patients with neurological diseases. These subjects will then be given large doses of Vitamin B-12 parenterally for a period of one week (varying from 1,000 to 5,000 micrograms per day). At the end of this period a repeat lumbar puncture will be performed and the level of Vitamin B-12 will again be determined. It is hoped that in one or two normal subjects and in several patients it will be possible to do several lumbar punctures within the treatment period to determine a curve for the spinal fluid levels of the vitamin. Where practical vitamin levels in the blood will also be performed before and at the end of therapy. The Vitamin B-12 Assays will be carried out by the Lactobacillus microbiological method and will be performed initially at least by Merck and Co.
- (c) Major Findings: This project began in December 1955. Two patients have been run through the plan outlined above but reports of the results are not yet available.
- (d) Significance to Neurological Research: It is anticipated that spinal fluid levels of B-12 may rise considerably over a week's period with large doses of the vitamin, in contradistinction to the blood levels with such therapy. If such is the case, it would be of significance in determining the dosages to be employed in therapy with Vitamin B-12 and perhaps of more significance in demonstrating that the substance itself is concentrated within the spinal fluid.
- (e) Proposed Course: To carry out studies outlined above.

Budget Data Sheet

10. NINDB- 22 (C)
SERIAL NO.

11. BUDGET DATA:

FY	ESTIMATED	BUDGETED POSITIONS			MAN YEARS			PATIENT
	EXPENDITURES	PROF	OTHER	TOTAL	PROF	OTHER	TOTAL	DAYS
FY 1956		2	2	4	0.15	0.2	0.35	*
FY 1957		2	2	4	0.15	0.2	0.35	

*Patients drawn from other NINDB project.

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
REVIEW & APPROVAL TECHNICAL ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 OR 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM L)

Research Division, Merck and Co., Rahway, N.J., Dr. George Boxer

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO(S) IF WITHIN NIH).

None.

Project Description Sheet

1. N.I.N.D.B. 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
Clinical
3. Neurochemistry 4. _____ 5. NINDB- 23 (C)
SECTION LOCATION (OTHER THAN BETHESDA) SERIAL NO.
6. Textbook of Clinical Neurochemistry
PROJECT TITLE
7. Dr. Donald B. Tower
PRINCIPAL INVESTIGATOR
8. Dr. Maynard Cohen, (University of Minnesota)
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION:

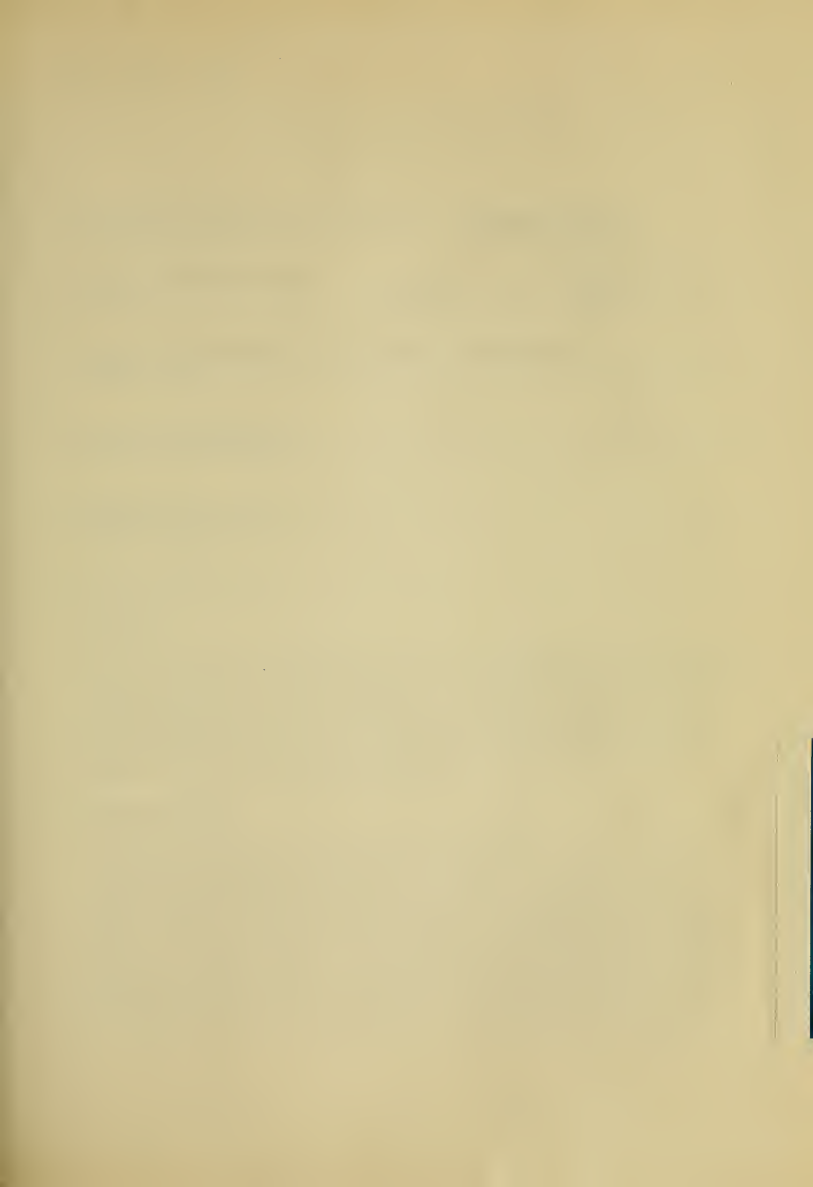
- (a) Objective: Writing of a book covering the aspects of neurochemistry of importance to the clinical neurologists and others in the field of neurology who may possess insufficient biochemical background yet wish to obtain a general grounding in this field.

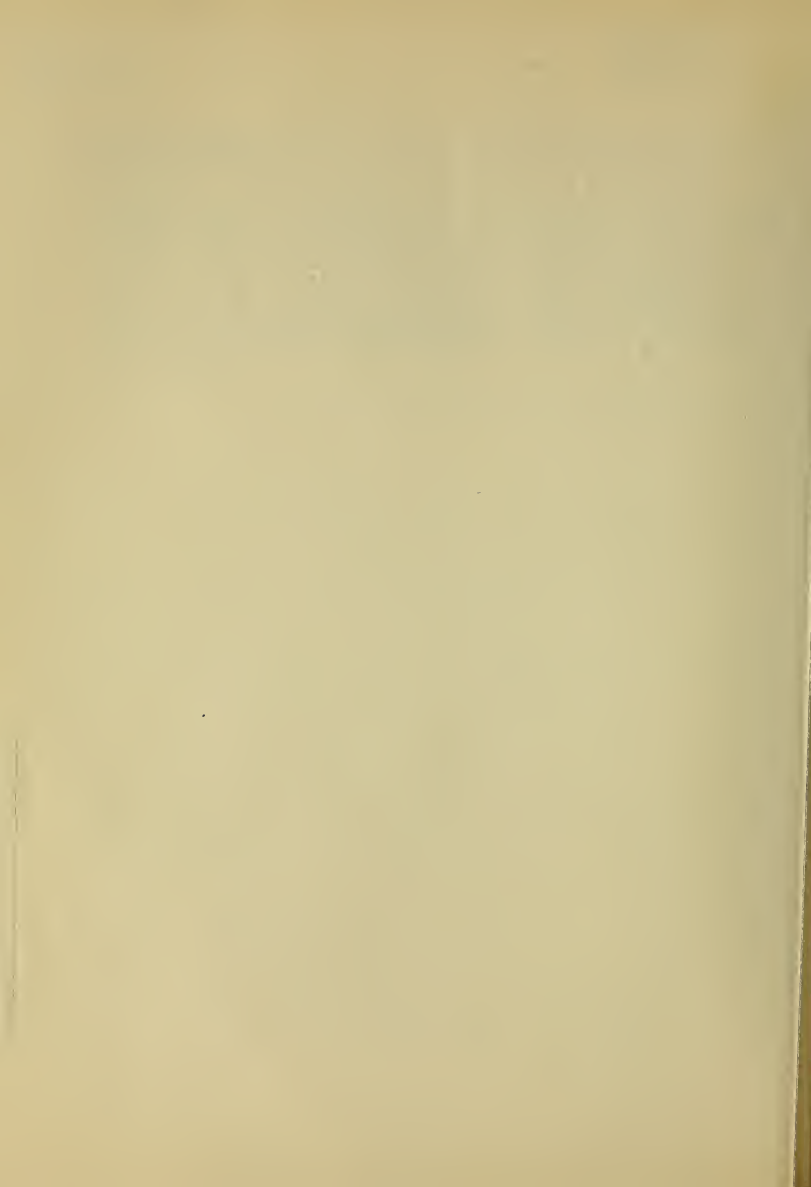
The book is intended to be 300-500 pages in length and to cover

- in as simple, diagrammatic terms as possible the major biochemical facts and views relating to the central and peripheral nervous systems and their correlation where known with clinical problems.

- (b) Significance: There have been three books on neurochemistry in the past 50 years. Thudichum's Chemical Constitution of the Brain first published 75 years ago appeared in revised form in German in 1901. This is a classic and contains much data still of value regarding the gross chemical make-up of gray and white cerebral tissue. In 1937 Page published a monograph on Chemistry of the Brain which brought Thudichum's data up to date and included what was then known of metabolism in brain tissue. This also is a classic but is much out of date, particularly regarding neuronal metabolism. In 1955 Elliott, Page and Quastel published Neurochemistry, a collection of 32 chapters by 32 authors covering most of the field. This book is actually valid up to 1951 only so that it is already somewhat behind current advances. In addition it is written primarily by and for investigators with biochemical background and not presented in any but the loosest organization. With a few exceptions there is little reference to clinical problems. The field of neurochemistry is experiencing growing interest from many areas of neurology, especially those in clinical neurology. A relatively short and moderately priced book directed to this latter group and organized in a logical basic to clinical progression will fill a much needed place.

- (c) Proposed Course: The collaboration of Dr. Maynard Cohen, in charge of Clinical Neurochemistry at the University of Minnesota, has been secured. The medical Publishing House of Paul B. Hoeber, Inc., New York has contracted to publish the book. As a result of preliminary review a short synopsis of some of the salient features of cerebral metabolism has been presented at a Symposium Interdisciplinary Research, University of Wisconsin and will be published in 1956. A one-day course on neurochemistry will be given by the investigators at the April 1956 meeting of the American Academy of Neurology. These two events will permit the authors to assess the approach most suitable for the intended "non-biochemical" audience. It is anticipated that the manuscript for the book can be completed by the end of calendar 1956.





METHODS EMPLOYED, Cont'd.

recording of muscular contraction. The potential changes from the strain gauge are led into a DC amplifier and then recorded photographically with a cathode ray oscilloscope. Two fine wire electrodes are also inserted into the muscle and after suitable amplification the gross muscle action potential is recorded on the second beam of the oscilloscope. These electrodes are also used to stimulate the muscle directly. The temperature of the muscle was originally measured with a thermocouple leading into a bridge circuit and recording galvanometer. A Brown recording potentiometer is now used for this purpose. In this way the mechanical and electrical events occurring in the muscle as the result of stimulation of the sciatic nerve can be studied as a function of the temperature of the muscle.

MAJOR FINDINGS

During the past year further pilot experiments were done in this project, using the rat as a test animal. Preliminary results have confirmed information already available, to wit that a lower frequency of stimulation is necessary to obtain a complete tetanus in a cool muscle than in one at normal body temperature. The relation of tension to temperature is a complex one and demands further analysis. There seems little relationship between the amplitude of the electrical response and the tension developed, and the necessity of recording the electrical events as a function of temperature has been questioned.

SIGNIFICANCE OF THE PROGRAM TO THE INSTITUTE

Since one of the major interests of this institute is the large and varied group of diseases of the neuromuscular system and since some of these disorders are allegedly affected by cold, it seems of interest to try to define more clearly the exact effects which temperature has on neuromuscular function.

PROPOSED COURSE OF PROJECT

It is planned to amplify the experiments which have already been performed on the rat and to continue these studies on hamsters arousing from hibernation, i.e. those automatically changing their body temperature from low to high values as well as in anesthetized non-hibernating hamsters whose body temperature can be altered over a wide range without the procedure being lethal to the animal. It is planned to study the effects of temperature on muscular contraction brought about both by stimulating the nerve to the muscle and the muscle directly. Since the method of recording muscular contraction in use has not proved entirely satisfactory, improved methods probably utilizing different apparatus will be sought for.

MAJOR FINDINGS

This project has just been started with the modification of an existing stereotaxic apparatus to fit the skull of the hamster. Further active research has had to be postponed for three months since Dr. Marshall is at present occupied with other research in collaboration with Dr. Aristides Leao, a Visiting Scientist.

SIGNIFICANCE OF THE PROGRAM TO THE INSTITUTE

It is felt that by studying the electrophysiology of the visual system, about which a great deal of information is already available, while varying the parameter of the body temperature of the animal, further information of interest may be obtained of interest both to neurophysiologists and ophthalmologists.

PROPOSED COURSE OF PROJECT

Outlined under "Methods".

Project Description

1. Neurological Diseases and Blindness 2. Medical Neurology
 INSTITUTE LABORATORY OR BRANCH
3. Clin. Neuropharmacology 4. _____ 5. NINDB-26(C)
 SECTION LOCATION (OTHER THAN LETHESDA) SERIAL NO.
6. Study of new chemical compounds with central nervous system effects.
 PROJECT TITLE
7. Richard L. Irwin, Ph.D.
 PRINCIPAL INVESTIGATOR
8. None
 OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: The study of pharmacologically active compounds that at first appear to be scientific curiosities is of importance for several reasons. Firstly some compounds of this type, when adequately characterized chemically and pharmacologically, find a useful place in therapeutics. Such substances have in the past also served as valuable research tools and have led to a greater understanding of physiological and biochemical mechanisms of disease processes. The objective of this project is to find compounds with central nervous system effects that have not been described and to investigate these compounds and others that have unusual or unexplained effects upon the central nervous system.

METHODS EMPLOYED: No method is known that will with certitude uncover pharmacologically active chemical substances. This project to date has been concerned with a search into folklore, anthropological literature and older scientific publications for reference to substances with pharmacologic activity that have not been previously or adequately investigated. Standard pharmacological procedures will be used when such substances are found and obtained.

MAJOR FINDINGS: To date reference to two substances have been found that are of interest. One of these compounds is reported to cause "madness" and is known as Felazlaz by some native peoples of the Sahara desert. Another reference to this compound indicated the name to be "Tebinna". Another group of compounds that have profound central nervous system effects that have received little attention is the Harmine alkaloids. These have been used as euphorics by Siberian peasants and have excitatory effects in animals that seem to be referable to the basal ganglia.

SIGNIFICANCE TO PROGRAM OF INSTITUTE: See objectives. This project is perhaps of more general interest than specifically related to an institute program.

PROPOSED COURSE OF PROJECT: To obtain, if possible, a supply of "Tebinna" (Felazlaz) and the harmine alkaloids for pharmacological testing and to continue the search for other active compounds.

Budget Data Sheet

10. NINDB - 26 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH

13. NONE

14. NONE

Honors, Awards, and Publications Sheet

15. SERIAL NO.

16. PUBLICATIONS OTHER THAN THIS PROJECT DURING 1955.

NONE

17. HONORS AND AWARDS TO PERSONNEL DURING 1955.

NONE

Project Description

1. Neurological Diseases and Elindness 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clin. Neuropharmacology 4. _____ 5. NINDB-27 (C)
SECTION LOCATION (OTHER THAN PETHESDA) SERIAL NO.
6. A study of respiratory and vasomotor responses to central nervous
system asphyxia.
PROJECT TITLE
7. Richard L. Irwin, Ph.D.
PRINCIPAL INVESTIGATOR
8. Jay F. Wells, M.S.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: To determine respiratory and vasomotor responses to asphyxia restricted to the central nervous system.

METHODS EMPLOYED: Responses were measured in anesthetized mammals using a vascularly isolated head in which neural connections remained intact.

MAJOR FINDINGS: The following observations were obtained from 22 dogs with vascularly isolated heads. The sudden occlusion of all arterial channels except the anterior spinal artery does not effect either vasomotor or respiratory function. When the anterior spinal artery is included in the occlusion an immediate vasopressor response ensues. This response is variable and is accompanied in some animals by cardiac arrhythmias. After an initial hyperpnea and polypnea, the occurrence of apnea, at approximately one minute after occlusion, was a constant finding in all dogs. When once established, apnea continued as long as the arterial channels remained occluded. When the arterial vessels were opened and blood flow resumed within ten minutes after the onset of apnea respiratory function returned to the pre-occlusion pattern.

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: Central nervous system function in relation to oxygen deprivation and carbon dioxide accumulation is inadequately understood. The number of neurological disorders which have parameters of function related to anoxia, asphyxia and hypercarbia attest to the importance of this type of study.

PROPOSED COURSE OF PROJECT: No immediate continuation of this project is contemplated. Experiments to separate the role played by anoxia from that of hypercarbia would be an obvious sequel to this project.

R.P.C. 2
December 1955 Analysis of NIH Program Activities
Budget Data Sheet

10. NIMDB - 27 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH

13. NONE

14. NONE

Honors, Awards, and Publications

15. SERIAL NO.

16. PUBLICATIONS OTHER THAN THIS PROJECT DURING 1955.

NONE

17. HONORS AND AWARDS TO PERSONNEL DURING 1955.

NONE

Project Description

1. Neurological Diseases and Blindness 2. Medical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clinical Neuropharmacology 4. _____ 5. NINDB-28 (C)
SECTION LOCATION (OTHER THAN ELTHESDA) SERIAL NO.
6. The antagonism of quarternary compounds to neuromuscular block.
PROJECT TITLE
7. Richard L. Irwin, Ph.D.
PRINCIPAL INVESTIGATOR
8. Jay E. Wells, M.S.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: This project is concerned with screening quarternary compounds for "de-curarizing" activity in anticipation of finding a compound that will be clinically useful in the treatment of myasthenia gravis.

METHODS EMPLOYED: Appropriate standard neuromuscular preparations.

MAJOR FINDINGS: NONE

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: The pharmacological treatment of myasthenia gravis during the past 20 years has centered upon compounds that have a pronounced anticholinesterase activity. Compounds that exhibit anticholinesterase effects only, such as physostigmine and the alkyl phosphonates, although somewhat efficacious in the treatment of myasthenia, have a low margin of safety or produce pronounced undesirable side effects at a clinical dose level. When a compound has anticholinesterase activity and also a direct stimulating effect upon the motor end plate, such as neostigmin or Mestinon (R), the clinical effectiveness is increased. This type of compound, with two distinct modes of action, and other similar ones, are invariably quarternary ions. It is also known that some quarternary compounds produce neuromuscular block. The pathological defect in myasthenia has often been proposed to be the circulation of a metabolite that reduces neuromuscular transmission by attachment to motor end plate receptors. It is unknown however whether the beneficial effect of neostigmin is dependent upon the displacement of a neuromuscular blocking metabolite from the motor end plate receptors or whether prostigmin acts by partially depolarizing the end plate membrane which would allow a smaller amount of acetylcholine to depolarize the end plate to a threshold level.

It has been shown in this laboratory, as well as in others, that certain quarternary compounds that have little anticholinesterase activity are somewhat antagonistic to a neuromuscular block. Hexamethonium, a bis-quarternary salt, is the best known example. This compound and other quarternary substance have found no clinical

application as antagonists to a neuromuscular block since they have pronounced ganglionic blocking action that precludes their use. In large doses they also exhibit a low grade neuromuscular blocking action. However, in homologous series of quaternary ions compounds usually exist which have little if any ganglionic or neuromuscular blocking activity. It is unknown whether these compounds have any antagonistic effect against a neuromuscular block. If a compound can be found that (1) has minimal ganglionic blocking action, (2) is capable of antagonism of a neuromuscular block and (3) has little affinity for acetylcholine receptors at the motor end plate then the possibility exists that such a compound would be beneficial clinically in the treatment of myasthenia. The feasibility of an active compound of this nature being of benefit is not precluded by the possibility that the pathological physiology of myasthenia gravis is not dependent on a circulating neuromuscular blocking metabolite since quaternary ions are also known to depolarize the motor end plate. If the myasthenic defect is, as has also been postulated, a deficiency in acetylcholine production, a quaternary compound that would partially depolarize the motor end plate would enable a smaller output of acetylcholine to effect adequate neuromuscular transmission.

This project, based on the above considerations, is part of the integrated muscle disorders program of this Institute.

PROPOSED COURSE OF PROJECT: To continue to test quaternary compounds for activity against a neuromuscular block.

Project Description

1. Neurological Diseases and Blindness 2. Medical Neurology
 INSTITUTE LABORATORY OR BRANCH
3. Clinical Neuropharmacology 4. _____ 5. NINDB-29 (C)
 SECTION LOCATION (OTHER THAN BETHESDA) SERIAL NO.
6. The development of a "myasthenic" animal for testing purposes.
 PROJECT TITLE
7. Richard L. Irwin, Ph.D.
 PRINCIPAL INVESTIGATOR
8. Jay B. Wells, M.S.
 OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: Myasthenia Gravis is a neurological disorder that somewhat resembles a state of chronic neuromuscular block. No adequate laboratory method has been described that will give information as to whether any drug will be effective in the treatment of myasthenia. The standard techniques to test for antagonism against a neuromuscular block are inadequate since the disease is not reproducible in a laboratory animal. The purpose of the present project is to develop a method for more accurate testing of drugs that may be of value in the treatment of myasthenia gravis.

METHODS EMPLOYED: The rabbit head drop method has been selected as a beginning point since this method is not complicated by either extraneous electrical stimuli or anesthesia. Rabbits have been chronically curarized in an attempt to reproduce somewhat the conditions found in the myasthenic patient and then tested for response to neuromuscular blocking compounds.

MAJOR FINDINGS: No suitable method has as yet been developed. The chronic curarization of rabbits by a depository preparation of d-tubocurarine did not appreciably alter the response of the animals to test doses of the same substance.

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: The development of a new method for screening compounds that would be pharmacologically active in the treatment of myasthenia would perhaps aid in finding substances of clinical value. If a method can be found to produce a "myasthenic like" state in a laboratory animal it is conceivable that new information may be obtained concerning the neuromuscular processes involved in myasthenia about which little is known at present.

PROPOSED COURSE OF PROJECT: The search for a new method to test substances that may be effective in the treatment of myasthenia gravis will be continued. Chronic "curarization" of test animals

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SERIAL NO.

by substances that have a different mode of action from d-tubocurarine will be investigated. The possibility that the inhibition of choline acetylase in vivo will produce a test animal that is somewhat similar to a myasthenic patient will be investigated. An attempt will be made to find compounds that are choline acetylase inhibitors.

Project Description

1. Neurological Diseases and Blindness 2. Medical Neurology
 INSTITUTE LABORATORY OR BRANCH
3. Clin. Neuropharmacology 4. _____ 5. NINDB-30 (C)
 SECTION LOCATION (OTHER THAN BETHESDA) SERIAL NO.
6. Neuromuscular blocking compounds: Peripheral and respiratory action related to calcium, potassium, magnesium, carbon dioxide and anesthetic gases.

 PROJECT TITLE
7. Richard L. Irwin, Ph.D.
 PRINCIPAL INVESTIGATOR
8. Jay B. Wells, M.S.
 OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: To determine how changes in the concentration of physiological ions and respiratory or anesthetic gases are related to the phenomenon of neuromuscular block.

METHODS EMPLOYED: During the course of this project a method has been devised to measure the respiratory effects of neuromuscular compounds under conditions that approximate those under which this type of compound is useful. The method is not dependent on extraneous nerve stimulation or artificial respiration. Peripheral actions are determined in standard nerve-muscle preparations in the rat, guinea pig, cat, rabbit and dog using alternating direct and indirect stimulation of the muscle.

MAJOR FINDINGS: The administration of calcium is markedly antagonistic to the peripheral as well as to the respiratory effects of the neuromuscular blocking compound succinylcholine. This has been found not to be due to a direct effect of calcium on muscle fibers or to the release of adrenalin by calcium. Indirect evidence suggests that cholinesterase activation by calcium is not involved. Calcium not only antagonizes a pre-existing neuromuscular block but also decreases the depth and duration of the block when calcium is administered prior to the blocking compound. Hypercarbia reduces both the duration and depth of the neuromuscular block produced by succinylcholine. The antagonism occurs only when acidemia has existed prior to the onset of the block and is thus the result of some physiological change that develops during hypercapnia. Anoxia, at least within the physiological limits of the cardiovascular system, has been found not to effect a neuromuscular block.

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: Myasthenia Gravis and familial periodic paralysis are neurological diseases in which both derangement of neuromuscular transmission and changes in ionic concentration in the blood may be involved. Until more knowledge on

the mechanisms of a neuromuscular block and its relation to ionic concentration is available the etiology and treatment of these diseases must necessarily remain empirical. It is therefore important to examine the phenomena of neuromuscular block in greater detail than is possible in the patients. The investigation of the neuromuscular block and the conditions which affect it is the problem with which this project is concerned.

PROPOSED COURSE OF PROJECT: The phase of the project concerned with succinylcholine, calcium and respiration is complete and is documented in manuscript form to be presented for publication. The next phase of this project will concern the effect of hypercapnia on the neuromuscular block produced by compounds which are not hydrolyzed by cholinesterases i.e. decamethonium and gallamine. The relationships between a neuromuscular block and ionic fluxes will be investigated.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness 2. Cerebral Palsy
INSTITUTE LABORATORY OR BRANCH
3. _____ 4. _____ 5. NINDB-31 (C)
SECTION LOCATION SERIAL NO.
6. The Evaluation of New Muscular Relaxant Drugs in the Treatment of Cerebral
PROJECT TITLE Palsy.
7. J. Gordon Millichap, M.D.
PRINCIPAL INVESTIGATOR
8. _____
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Project: The Evaluation of New Muscular Relaxant Drugs in the Treatment of Cerebral Palsy.

Objectives: (1) To establish reliable objective methods for the quantitative assessment of motor disturbance in children with cerebral palsy.

(2) To determine the value of new muscular relaxant drugs in the treatment of cerebral palsy.

Methods: Objective measurements are obtained of coordination, range, accuracy and rapidity of movement, muscle strength, maintenance of posture, and degree of involuntary movement. Test procedures include kymographic records of the effects of choreoathetoid movements on postural stability, dark ground photographic records of athetoid movements with timed exposure, Minnesota rate of manipulation and other manual dexterity tests, hand dynamometer, vital capacity, and maximum leg abduction measurements. Neurological, physical therapy and occupational therapy evaluations, speech recordings, and limited motion picture studies are obtained.

Measurements are repeated at three day intervals, and other evaluations are made during drug treatment in which controls are employed. Results are subjected to statistical analyses.

Analysis of NIH Program Activities

Patients under study:

Total: 10 children

In-patients: 4

Out-patients: 6

Significance to the Cerebral Palsy and Pediatric

Neurology Program: In patients with cerebral palsy, muscular relaxation and the reduction of involuntary movements are essential preliminaries to the development of voluntary muscle control and proper movement patterns. A muscular relaxant drug of proven value would be an important addition to the treatment of cerebral palsy, and an effective preventive of contractures and deformity.

Major Findings: Preliminary assessment of the results of screening studies in four patients suggests that the degree of motor disability is increased during treatment with Reserpine, and decreased by Flexin (2-amino-5-chloro-benzoxazole).

Proposed Course of Project: In view of the encouraging results with Flexin, the study will be extended to include a total of ten patients. They will be followed as out-patients.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness 2. Cerebral Palsy
INSTITUTE LABORATORY OR BRANCH
3. _____ 4. _____ 5. NINDB-32 (C)
SECTION LOCATION SERIAL NO.
6. The Anticonvulsant Action of Carbonic Anhydrase Inhibitors.
PROJECT TITLE
7. J. Gordon Millichap, M.D.
PRINCIPAL INVESTIGATOR
8. _____
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Project: The Anticonvulsant Action of Carbonic Anhydrase Inhibitors.

Objectives: (1) To determine the efficacy of acetazoleamide (Diamox) in the treatment of those seizures which are not markedly influenced by hyperventilation and acid-base changes. [It is generally postulated that Diamox influences seizure activity by means of the secondary metabolic acidosis consequent to the inhibition of renal carbonic anhydrase. However, in animal experiments, the drug has an anticonvulsant effect which is independent of its action on the kidney and which is correlated directly with the degree of inhibition of carbonic anhydrase in the brain (Millichap, J.G., Woodbury, D.M., and Goodman, L.S., 1955.) Control of seizures, other than those exacerbated by hyperventilation and alkalosis, should therefore be expected.]

(2) To confirm, by clinical experiment, the development of tolerance to Diamox which has been noted in animal investigations, and to determine the efficacy of ammonium chloride in the reduction of tolerance.

Methods: The results of Diamox therapy are correlated with the etiology, anatomical localization, and clinical and EEG patterns (including response to hyperventilation) of the seizures. Control medications and the double-blind-technique are employed, and the patients are reviewed at frequent intervals.

Patients Studied:

Total of 23 children.

In-patients: 2

Out-patient follow-up consultations: 120

Major Findings: (1) The effectiveness of Diamox has been established in the control of those major and minor seizures, not markedly influenced by acid-base changes.

(2) The development of tolerance, hitherto unreported, detracts from the practical value of Diamox as an anticonvulsant.

Significance to the Cerebral Palsy and Pediatric

Neurology Program: In children with cerebral palsy, the control of seizures is an essential preliminary to effective specific therapy, and there is urgent need for controlled evaluations of new and less toxic anticonvulsant agents. A study of the mechanism of the anticonvulsant action of Diamox should lead to an understanding of the physiological role of carbonic anhydrase in the nervous system and its relation to seizure activity.

Proposed Course of Project: The relatively low brain/plasma concentration ratio of Diamox limits its anticonvulsant activity (Millichap et al., 1955.) As a result of this finding, a more potent inhibitor has been produced and, if suitable patients are readily available, the compound will be evaluated clinically.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness 2. Cerebral Palsy
INSTITUTE LABORATORY OR BRANCH
3. _____ 4. _____ 5. NINDB-33 (C)
SECTION LOCATION SERIAL NO.
6. Studies on the Pathogenesis of the Kernicterus Syndrome
PROJECT TITLE
7. J. Gordon Millichap, M.D.
PRINCIPAL INVESTIGATOR
8. _____
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objectives: 1) To investigate the significance of bile staining of the brain in the production of the kernicterus syndrome. 2) To investigate the relation of liver insufficiency, hypoglycemia, hypothermia, and iron deficiency of prematurity to the development of kernicterus. 3) To determine possible methods of prevention.

Methods: 1) Respiration of rat brain slices, in the presence of bilirubin, is measured by use of the Warburg constant volume respirometer. The in vitro inhibitory effect of bilirubin is being studied in relation to a) the purity of the sample, b) temperature, c) pH and various constituents of the media, and d) the presence of iron and copper.

2) Rats with liver necrosis and hypoglycemia will be used in the investigation of the relation of liver insufficiency and hypoglycemia to the kernicterus syndrome. The production of kernicterus in these animals will be attempted by injection of bilirubin solutions, a procedure which has been found unsuccessful in normal rats. Respiration of brain slices from animals injected with bilirubin will be compared with that of animals not injected.

Preliminary Results: The inhibitory effect of bilirubin on brain homogenates in vitro (Day, R.L., 1955) has been demonstrated also on brain slices. To obtain this effect, however, it is necessary to use an unphysiological

media at a strongly alkaline pH. Under more physiological conditions, a significant degree of inhibition has not yet been obtained, despite staining of the tissue.

Significance to Cerebral Palsy Research Program: Though the incidence of the kernicterus syndrome has been reduced by exchange transfusion, a causal relation between bilirubin staining of the brain and the neurological sequelae of neonatal jaundice has not been established. The toxic effect of bilirubin, demonstrated in vitro, may be non-specific, and other factors associated with liver insufficiency, such as hypoglycemia and high serum ammonia levels, may be more significant. So that a better understanding of the nature of cerebral palsy may be obtained, and from the point of view of prophylaxis, it is important to elucidate the true causal factors in this condition.

Proposed Course of Project: No change.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness 2. Cerebral Palsy
INSTITUTE LABORATORY OR BRANCH
3. _____ 4. _____ 5. NINDB-34 (C)
SECTION LOCATION SERIAL NO.
6. Etiological and Clinico-pathological Correlations in Cerebral Palsy.
PROJECT TITLE
7. J. Gordon Millichap, M.D.
PRINCIPAL INVESTIGATOR
8. Neuropathologist to the Collaborative Cerebral Palsy Project.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Project: Etiological and Clinico-pathological Correlations in Cerebral Palsy.

Objectives: 1. To correlate prenatal, natal and post-natal factors generally considered of etiologic significance in cerebral palsy, with the clinical and pathological findings.

2. To determine the incidence of factors, other than neonatal jaundice, of possible significance in the etiology of the kernicterus syndrome.

Methods: (1) Records of the pregnancy, birth and preceding clinical history are obtained on cerebral palsy patients with a history of neonatal jaundice, and these will be compared with the records of all children referred for possible inclusion in other cerebral palsy research projects. Detailed examinations and investigations are performed, and suitable normal control patients will be studied similarly. So that the degree of reliability and completeness of the histories, consistent with a retrospective investigation, may be optimal, patients will be admitted to the study as soon after diagnosis as is possible.

Patients Studied:

Total: 24 children
Inpatients: 4
Out-patients: 20

Analysis of NIH Program Activities

Progress: Records have been obtained on 24 children with cerebral palsy, and assistance has been given in forming a cerebral palsy protocol which eventually will be employed in this study. The cooperation of local hospitals and health departments has been obtained for the referral of patients and records.

Significance to the Cerebral Palsy and Pediatric

Neurology Program: More precise etiological and clinicopathological correlations in cerebral palsy should lead to a better classification and understanding of this condition. The investigation of the kernicterus syndrome will help to elucidate the clinical significance of the results of laboratory research concerning bilirubin and brain respiration. (C.P. Project No. 802).

Proposed Course of Project: No change.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness 2. Cerebral Palsy
INSTITUTE LABORATORY OR BRANCH
3. _____ 4. _____ 5. NINDB-35 (C)
SECTION LOCATION SERIAL NO.
6. Precocious Puberty in Children with Cerebral Palsy.
PROJECT TITLE
7. J. Gordon Millichap, M.D.
PRINCIPAL INVESTIGATOR
8. _____
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Project: Precocious Puberty in Children with Cerebral Palsy.

Objectives: To determine the relation of skeletal and sexual precocity to organic brain damage in children with cerebral palsy.

Methods: (1) Children with cerebral palsy are investigated for signs of precocious puberty. A general physical examination and an x-ray study for bone-age are performed on all patients and, in those with positive findings, the urinary excretion of ketosteroids and corticoids is estimated. An endocrinology consultation is obtained to exclude causes other than neurogenic.

(2) The anatomical localization of the cerebral lesion in the patients with precocious puberty is compared with that in cerebral palsy patients who show no signs of precocity.

Patients Studied:

Out-patients: 12

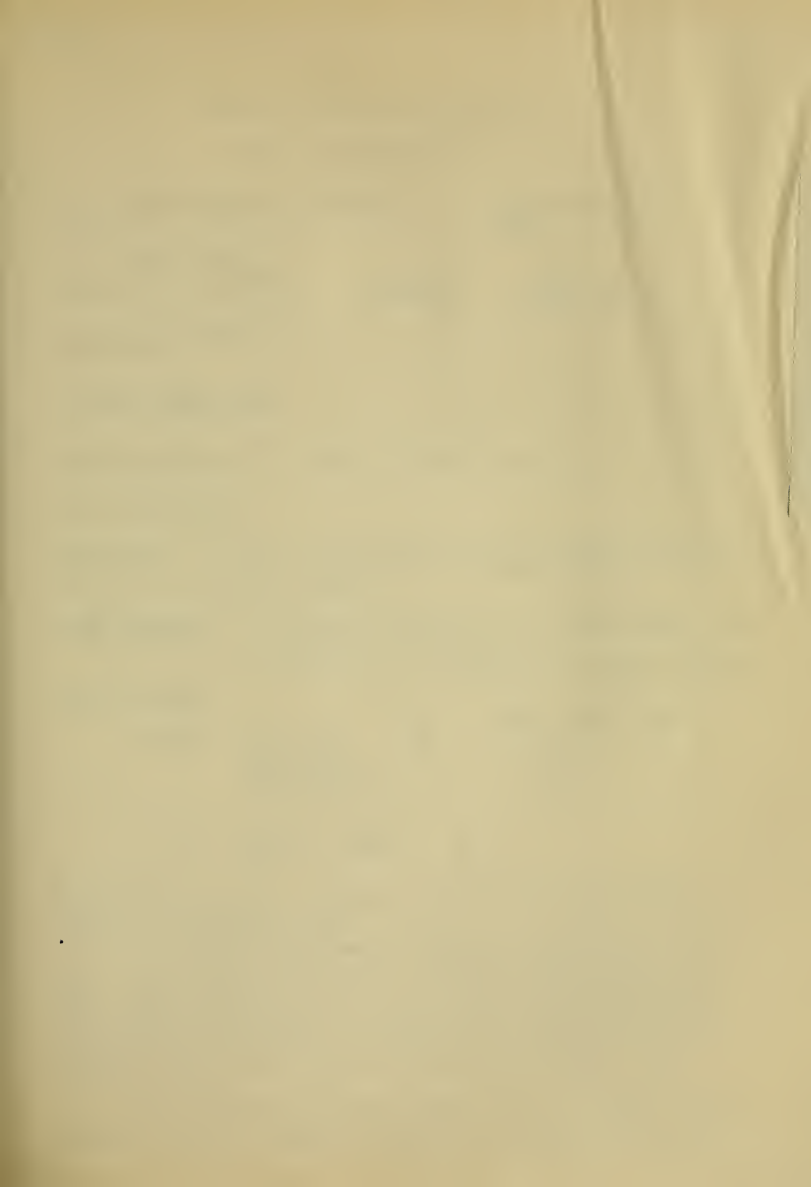
Analysis of NIH Program Activities

Significance to the Cerebral Palsy and Pediatric

Neurology Program: A knowledge of the relation of precocious puberty to the brain lesion in cerebral palsy would help to define cortical-hypothalamic connections, and the role of the cerebral cortex in endocrine function.

Major Findings: Of 12 cerebral palsy patients in this study, 5 showed definite evidence of precocity, and causes other than neurogenic were excluded.

Proposed Course of Study: Further patients will be studied as above and, in addition, patients with seizures and focal cortical lesions will be investigated.



Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Experimental Neurosurgery
SECTION
4. _____
LOCATION
5. NINDB-36 (C)
SERIAL NO.
6. Mechanisms of Memory
PROJECT TITLE
7. Maitland Baldwin, M. D.
PRINCIPAL INVESTIGATOR
8. Laurence L. Frost, Ph. D., and J. M. Van Buren, M. D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: To further our understanding of those functional representations within the temporal lobe of man and the higher primates which may subsERVE the mechanisms of memory.

Methods Employed: 1. Electrical stimulation of the human temporal lobe during operative exposure under local anesthesia.
2. Ablation of the temporal areas in higher primates.

Patient Material:

	No.	Average stay - days
Admissions: Adult Male	20	5.75
Adult Female	26	7.27
Children Male	3	17.33
Children Female	4	5.0
Outpatient: Number of Patients	21	
Number of Visits	28	

Major Findings: In four patients we have elicited responses which indicate that there may be neuronal patterns in the temporal lobe which subsERVE the memory mechanisms.

In six patients whose temporal lobes were ablated on one or the other side, we observed changes in the ability to record present experience. These changes were such that the patients could not describe or define immediate experience and in at least one of the cases included some disorientation in space. In these cases it is noteworthy that we ablated the most mesial temporal structures and that there was some evidence of damage of these structures on the side opposite to the surgical removal.

In all patients with chronic lesions in one or the other or both temporal lobes, there is some disturbance of the mechanisms which serve to provide awareness, attention, perception and analysis of the meaning

PROJECT DESCRIPTION (continued)

of present experience. These disturbances lead to a failure of recording of this present experience or they may also lead to an inaccuracy of recollection or an inaccurate recording of this experience.

We have ablated the temporal lobe on one or the other side in three chimpanzees. These animals are higher primates and observations based on their behavior are useful through analogy in the study of human behavior. In a previous report we described the first immediate results of these ablations which consisted of immature behavior. Despite this change in behavior the chimpanzees have relearned patterns essential to feeding, grooming and cage play, in one case at least. These further observations have been made over the period intervening between these reports.

Significance to Neurological Research: This project and those allied with it in the Section of Clinical Psychology, serve to provide some data on the physiological basis and perhaps on the structural basis for memory. A major effort is being directed towards the establishment of data which will provide an understanding of some of the structural bases for mental functions and therefore for mind.

Proposed Course of Project: We plan a continuation of our present study of those neuronal patterns in the temporal lobe which subserve memory. The combination of information derived from direct electrical stimulation of the temporal lobe in man with information derived from studies of humans and higher primates in which the temporal areas have been ablated should provide useful and perhaps significant information concerning the structural basis for mind.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness INSTITUTE
2. Surgical Neurology BRANCH
3. Neurosurgical Service SECTION
4. _____ LOCATION
5. NINDB-37 (C) SERIAL NO.
6. A Vestibular Representation in the Temporal Lobe PROJECT TITLE
7. J.M. Van Buren, M.D. PRINCIPAL INVESTIGATOR
8. Maitland Baldwin, M.D. and B. Ralston, M.D. OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: This study is undertaken to define the anterior boundary of the nystagmogenic area of the temporal lobe in man. It has been shown since the early portion of the 20th Century that lesions in the hemisphere of man produce accentuation of the quick phase of caloric nystagmus toward the side of the lesion. It has been subsequently demonstrated by Carmichael and Hallpike that this quick phase predominance toward the side of the lesion occurs whether cold water is irrigated in the contralateral or warm water is irrigated in the ipsilateral ear. The latter authors have further shown that the area responsible for this change lies in the posterior temporal region.

Because of the availability of patients in whom a measured quantity of temporal lobe had been removed for epilepsy, it was decided to follow out these studies to determine the anterior border of the portion of the cerebral cortex which affects the ocular response to caloric stimulation of the ears.

Method Employed: Prior to temporal lobectomy, a control examination was carried out and the same examination repeated following removal of measured portions of the temporal lobe. This examination consisted in positioning the patient with the neck flexed 30 degrees, and irrigating the external ear canals for 30 seconds with water 7 degrees above and 7 degrees below body temperature. Using a stop watch, the time of regular nystagmus was measured from the time the irrigation of the ear canal was commenced. In this way, a roughly quantitative measurement was obtained, which could be compared in the pre and post-operative periods.

PROJECT DESCRIPTION (Continued)

Patient Material:

	<u>No.</u>	<u>Average Stay - days</u>
Admissions: Adult Male	11	6.27
Adult Female	11	6.18
Children Male	3	17.0
Children Female	3	3.0
Outpatient: Number of patients:	10	
Number of visits:	10	

Major Findings: Although all of the material has not been fully correlated it appears that removal of the anterior 6 centimeters of the temporal lobe as measured along the sylvian fissure does not cause the appearance of nystagmus. Lesions, however, in the posterior temporal region, in the few instances where this has been invaded at operation, have produced the findings as mentioned above.

Significance to Neurological Research: The previous studies of this phenomenon have been by the use largely of cerebral tumors or, in a few cases, large destructive vascular lesions. The final localization of the phenomenon was arrived at by the method of super-imposition of the lesions upon a brain chart. The inherent weaknesses of this type of material and method of examination are apparent. It has therefore been felt of use to review this data using a more precise and quantitative cerebral lesion in man. In particular, we have been able to determine the anterior limits of this nystagmogenic area in man in the temporal lobe.

Proposed course of the Project: The majority of the patient work has now been finished. What remains is a correlation of the findings with a detailed review of the operative data.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Neurosurgical Service
SECTION
4. _____
LOCATION
5. NINDB - 38 (c)
SERIAL NO.
6. Visual Field Deficits Following Temporal Lobectomy
PROJECT TITLE
7. J.M. Van Buren, M.D.
PRINCIPAL INVESTIGATOR
8. Maitland Baldwin, M.D. and B. Ralston, M.D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: This study is directed toward the definition of the precise course of the visual fibers in the temporal lobe. The precise relationship between the fibers of the visual radiation passing anteriorly in the temporal lobe and the retinal areas which they supply has been previously worked out only in a gross manner by the use of patients with neoplasms of the temporal lobe.

The availability of patient material in which the temporal lobe has been removed for epilepsy offered a unique opportunity for the restudying of the problem in patients where the removal was made in a relatively quantitative fashion and in which the effects of inactivation by compression, as may be found about a cerebral tumor, could be eliminated.

Method Employed: Both prior and following operation the visual fields were examined both on the perimeter and on the tangent screen with a variety of test objects. Particular attention was paid to the density of the visual field defect. In each case this density was determined by the use of a test object as required up to 10 inches in diameter. The relationship of the quadratic field defect to macular vision was also carefully tested by the use of small white or small colored objects. The visual acuity in all cases was carefully tested. Every opportunity has been taken to reexamine patients at a considerable period following temporal lobectomy to rule out changes due to post-operative swelling which might appear in examinations made within the first month following operation.

PROJECT DESCRIPTION (Continued)

Patient Material:

	<u>No.</u>	<u>Average Stay - days</u>
Admissions:		
Adult Male	15	11.07
Adult Female	13	6.46
Children Male	3	17.0
Children Female	1	3
Outpatient:		
Number of patients:	10	
Number of visits:	10	

Major Findings: A complete evaluation of the findings so far has not been made. A preliminary impression is that there may be two types of visual field deficits. The most common is that of a non-homonymous upper quadrantanopia in which the largest and most dense defect appears in the eye ipsilateral in the temporal lobectomy. There are a few instances, but instances which have appeared with sufficient frequency to suggest that they are not merely chance happenings, in which the quadrantic field defect has proved to be homonymous. In no instance has the visual acuity been affected.

Significance to Neurological Research: The use of non-expanding destructive lesions of the temporal lobe in man has provided an unusual opportunity for the study of defects of the optic radiation in this area. The demonstration of two probable types of visual field defects in this area have been made. The significance of this difference at the moment is not apparent.

The non-homonymous nature of the field defects requires explanation in a differing spatial orientation of the optic radiation fibers from the opposite sides of the retina in the temporal portion of the optic radiation. Since in many cases the temporal lobectomy is accomplished by removal of gray and white substance from about the ventricle rather than by transection of the ventricle itself, it may be that these ipsilateral retinal fibers lie more laterally in the optic radiation and thus are more prone to damage than are the fibers from the contralateral retina which could lie closer to the ventricle and therefore be farther from the area of surgical damage. An alternative explanation would lie in the fact that instead of a lateral lamination there is an anteroposterior lamination in that the ipsilateral retinal fibers lie farther anteriorly in the temporal loop of the optic radiation than do the contralateral fibers. In this way removal of the temporal lobe from anterior to posterior would tend to strike first the ipsilateral fibers, then the contralateral, and thereby explain the larger defect in the ipsilateral eye.

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SERIAL NO.

PROJECT DESCRIPTION (Continued)

Proposed course of the Project: On the basis of the above unanswered questions, particular care is being taken in present temporal lobectomies to determine the degree to which the removal is tangential to the optic radiation and to what degree it is actually across the optic radiation, thereby entering the ventricle. It is hoped by further observation of patients with more detailed operative notes as to the nature of the lesion that the above incongruity may be elucidated.

In addition to the above studies, made purely on a clinical and operative basis, studies are being undertaken to reconstruct the lateral geniculate body in two patients who had died following temporal lobectomy. In this way, a correlation between the anatomical changes and those found at clinical examination of the patient may be achieved.

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Dec. 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB - 38 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH

13. None

14. None

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955:

None

17. HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

Analysis of NIH Program Activities

Project Description Sheet

- | | |
|--|---|
| 1. <u>Neurological Diseases and Blindness</u>
<u>INSTITUTE</u>
Neurosurgical Service | 2. <u>Surgical Neurology</u>
<u>BRANCH</u> |
| 3. <u>Experimental Neurosurgery</u>
<u>SECTION</u> | 4. _____
<u>LOCATION</u> |
| 5. <u>NINDB - 39 (C)</u>
<u>SERIAL NO.</u> | |
6. A Study of Functional Representation in the Amygdala of Man and the Higher Primates
PROJECT TITLE
7. Maitland Baldwin, M. D.
PRINCIPAL INVESTIGATOR
8. Laurence L. Frost, Ph.D., Charles D. Wood, M.S., and Shirley Lewis, R.N.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: To further understanding of the functions and functional connections of this important nucleus in man and the higher primates.

Methods Employed: 1. The electrical stimulation of the amygdaloid nucleus in patients under local anesthesia during operative exposure of the temporal regions.

2. The electrical stimulation of the amygdaloid nucleus in higher primates during operative exposure of the temporal regions and as chronic preparations after implantation of deep electrodes.

3. Electrographic recording from the temporal, parietal, frontal and deep temporal regions during these electrical stimulations.

4. As a result of the electrical stimulation of the amygdaloid nucleus in higher primates we have evoked afterdischarges from the nucleus to distant structures. We have recorded the spread of these after-discharges in some cases.

<u>Patient Material:</u>	<u>No.</u>	<u>Average Stay - days</u>
Admissions: Adult Males	18	5.61
Adult Females	23	7.09
Children Male	3	17.0
Children Female	4	5.0
Outpatient: Number of Patients	18	
Number of Visits	20	

Major Findings: The technique which we developed last year has been employed for electrical stimulation of the amygdaloid nucleus in patients during operation under local anesthetic. With this technique we have

Major Findings (continued):

evoked responses in thirty-six cases. These responses are motor, postural and autonomic. The motor responses are movements of face, jaws and upper extremities. The postural responses are posturing of the upper and lower extremities. The autonomic changes which occur are alteration of respiration and there is some change in sweating as well.

We have explored this nucleus with Horsley-Clarke and newer techniques in chimpanzees as well. In these animals we have implanted electrodes and developed chronic preparations after implantation. We have demonstrated that the anterior commissure provides a significant pathway of transmission between these nuclei in the monkey, and in both the monkey and the chimpanzee we have again elicited movements of face and jaw and autonomic changes as well as postural phenomena following these deep stimulations. There is a diffuse alteration in the cortical electrical activity which changes after the stimulation, and this is most marked in the parietal regions. During these stimulations we have observed the changes in behavior which occur. The most significant and frequent change is an alteration of consciousness which begins with the onset of stimulus and ends as it ends.

Significance to Neurological Research: The amygdaloid nucleus is an important and central mechanism within the temporal lobe. It has functions related to face and jaw movement, autonomic function, and behavior as well as the mechanisms of consciousness. We must understand it and its connections if we would understand temporal lobe epilepsy.

Proposed Course of Project: We are continuing the stimulation of this nucleus in man and in the primates. We will conduct a thorough study of its anatomical connections in the primates and we are studying the problems of behavior in man and the primates as well as the electrophysiological problems related to the stimulations in both man and the primates. Future studies of this kind should provide important information concerning the mechanisms of temporal lobe epilepsy and certain aspects of behavior and the mechanisms of consciousness.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
Neurosurgical Service
2. Surgical Neurology
BRANCH
3. Experimental Neurosurgery
SECTION
4. _____
LOCATION
5. NINDB-40 (C)
SERIAL NO.
6. Investigation of the Second Sensory Areas in Man
PROJECT TITLE
7. Maitland Baldwin, M. D.
PRINCIPAL INVESTIGATOR
8. J. M. Van Buren, M. D.
OTHER INVESTIGATORS
9. Project Description

Objectives: To outline the extent and nature of these sensory representations in man and if possible discover their functional significance.

Methods Employed:

1. Electrical stimulation of these areas under local anesthesia during operative exposure.
2. Recording of these results.
3. Electrographic recording during these stimulations.

Patient Material:

	No.	Average Stay - days
Admissions: Adult Male	14	6
Adult Female	17	5.82
Children Male	3	17
Children Female	7	9

Outpatient: Number of Patients: 17
Number of Visits: 19

Major Findings: We have evoked positive responses relative to second sensory areas in 53 patients. Some of these responses have been obtained from the upper bank and fissure of Sylvius, as well as the surface of the island of Reil. The latter category are infrequent, however. It is interesting that twelve of the cases showed sensory responses after stimulation of the lateral aspect of the first and second temporal convolution. These findings seem to indicate that there may be some sensory representation relative to this part of the temporal cortex. If this is true we have encountered yet another second sensory area. This area is as yet unreported and has not been described in the literature.

Significance to Neurological Research: This is a relatively unknown area of brain function. Knowledge of its function is important to our understanding of the sensory mechanisms of the brain and is essential to our knowledge of temporal lobe epilepsy since epileptogenic lesions often activate this area and their expression is not clearly understood.

Proposed Course of the Project: We must continue stimulations of these areas in man and reinforce the knowledge gained from these stimulations by evoked potential studies in the operating room. This work will be substantiated and reinforced by observations on the brains of higher primates as well.

With apparatus which we have developed in connection with another project, we will pursue these studies. This apparatus will permit us to evaluate the results of evoked potentials in these areas in both the human and chimpanzee. Such observations should provide more objective data and if this is forthcoming we may substantiate our observations concerning sensory representation in the temporal cortex.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
Neurosurgical Service
2. Surgical Neurology
BRANCH
3. Experimental Neurosurgery 4. _____
SECTION LOCATION
5. NINDB - 41 (C)
SERIAL NO.
6. An Investigation of the Autonomic Phenomenon which Follows Stimulation of the Temporal Areas in Man and the Higher Primates
PROJECT TITLE
7. Maitland Baldwin, M. D.
PRINCIPAL INVESTIGATOR
J. M. Van Buren, M. D., Laurence Frost, M. D., Charles Wood, M. S.,
8. Shirley Lewis, R. N.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: To further understanding of the central representations of autonomic function.

Methods Employed: 1. Electrical stimulation at operation.
2. Electrographic and electronic recording during this stimulation.

Patient Material:

	No.	Average Stay - days
Admissions: Adult Males	19	7.05
Adult Females	19	5.63
Children Male	3	17.0
Children Female	5	5.6

Major Findings: In patients at operation under local anesthesia we have observed changes in blood pressure, pulse, respiration and gastric motility following stimulation of the insula; following stimulation of the amygdaloid nucleus we have observed changes in respiration and in pulse rate.

In chimpanzees and in monkeys we have observed similar changes.

Significance to Neurological Research: Central representation of autonomic function is not clearly understood. These central representations which we have outlined are important in the understanding of the structural basis of emotions and in the understanding of the central control of the cardiovascular and respiratory systems.

PROJECT DESCRIPTION (continued)

Proposed Course of Project: We must outline the exact nature of these autonomic changes which follow central stimulation in man. We have entered mechanisms which are important in neurology as well as in the understanding of changes in the cardiovascular and respiratory systems.

We will continue electrical stimulation of these areas in man and the higher primates and attempt to make our recording of the results more exact by additional techniques, both in man and the primates.

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Dec. 1955

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Neurosurgical Service
Experimental Neurosurgery
SECTION
4. _____
LOCATION
5. NINDB-42 (C)
SERIAL NO.
6. A Study of the Evoked Potentials in the Somatosensory Cortex Following Stimulation of the Temporal Cortex (from peripheral stimuli and from stimuli applied to the surface of the temporal cortex).
PROJECT TITLE
7. Maitland Baldwin, M. D.
PRINCIPAL INVESTIGATOR
8. J. M. Van Buren, M. D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objectives: To outline the extent and nature of the somato-sensory representation in man and identify any functional relationship which may exist between it and the lateral temporal cortex.

Methods Employed:

1. Electrical stimulation of these areas under local anesthesia during operative exposure.
2. Recording of these results.
3. Electrographic recording during these stimulations.

Patient Material:

	No.	Average Stay - days
Admissions: Adult Male	1	4
Adult Female	1	6
Children Male	0	
Children Female	0	
Outpatient: Number of Patients:	0	
Number of Visits:	0	

Major Findings: During the period of this report we have developed an apparatus which will permit us to develop a peripheral stimulus and coordinate this development with cortical recording. It was necessary that this apparatus be developed as it is essential to the further pursuit of the objectives outlined for this particular project.

Significance to Neurological Research: The somato-sensory areas in man have not been outlined completely by evoked potential techniques. This is an essential step in the further understanding of this important area of functional representation within the cortex. There is no indication in present knowledge that the postcentral gyrus or somato-sensory

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Dec. 1955

NINDB-42 (C)
SERIAL NO.

Significance to Neurological Research (continued)

cortex is related to the lateral temporal cortex, nor is there any indication in present knowledge that the lateral temporal cortex subserves sensory function. Some of our findings obtained in another project suggest that such function is subserved here and this project may assist us in a substantiation of this evidence.

Proposed Course of the Project: We will apply the apparatus which we have developed to further studies of the human and higher primate cortices in order to outline the somato-sensory area and analyze any relationships between it and the lateral temporal cortex which can be observed through the use of these particular techniques.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
Neurosurgical Service
2. Surgical Neurology
BRANCH
3. Experimental Neurosurgery
SECTION
4. _____
LOCATION
5. NINDB - 43 (C)
SERIAL NO.
6. An Investigation of Speech Function Localized in the Temporal Areas of Man
PROJECT TITLE
7. Maitland Baldwin, M. D.
PRINCIPAL INVESTIGATOR
8. Laurence L. Frost, Ph. D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: To outline the extent of cortical speech representation in man and to understand the relative significance of the areas outlined in speech functions.

Methods Employed: 1. Electrical stimulation of the temporal and parietal cortex in patients during operations under local anesthesia.
2. Clinical examination of speech function before and after such operations.

<u>Patient Material:</u>	<u>No.</u>	<u>Average Stay - Days</u>
Admissions: Adult Male	24	9.67
Adult Female	23	6.35
Children Male	3	17.0
Children Female	6	5.5
Outpatient: Number of Patients	19	
Number of Visits	23	

Major Findings: During the period of this report we have continued our outline of the speech representation in human cortex. It is interesting that in some of these cases speech representation does not appear on the side opposite to the writing hand and in others it appears on the same side as the writing hand.

Individuals with temporal lobe lesions seem to have some difficulty in speech regardless of the side of the lesion. In the majority of cases, this difficulty is marked and obvious when the lesion is related to the side of dominance or major speech representation. However, in other cases, where the lesion is on the side opposite to that of major speech representation, there is also a difficulty in speech. In the first and larger group of cases the speech difficulty falls within the well-defined patterns of aphasia. In the second group it is more subtle

PROJECT DESCRIPTION (continued)

and is concerned with the ultimate semantics of speech. These patients have difficulty with the use of words so as to properly define a meaning. This difficulty does not appear where the word which supports or defines the meaning is used alone; as for example, if the patient is asked to name "pencil" after being shown this object, he does so with ease, but if he is asked to use the word "pencil" in a sentence so as to describe this object he has difficulty. These patients find it difficult to use words in adequate relationships but can use individual words adequately as unit symbols, but they cannot use groups of words as accurate symbols of thought or description.

Significance to Neurological Research: Speech is among the most complicated functions of the human brain. We know little of its intrinsic mechanisms. Some knowledge of these functions provides great assistance to the neurologist in diagnosis of brain disease and in the rehabilitation of those with brain lesions. This project relates to the one on vestibular and auditory function and also to the psychological study of speech. In combination these projects should provide information of value to the neurologist, psychologist and psychiatrist.

Proposed Course of Project: We will continue along present lines and add more precise methods of testing speech in the operating room as well as methods designed to outline the cortical representations of hearing and vestibular function.

We are beginning to observe the fragments of speech which occur during some cases of automatic behavior and we will continue observations of this distorted expression. It is interesting that individuals who owe their automatism to temporal lobe lesions may speak during their periods of automatic behavior if the essential temporal lesion is on the side of major speech representation.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Neurosurgical Service
Experimental Neurosurgery
SECTION
4. _____
LOCATION
5. NINDB - 44 (C)
SERIAL NO.
6. Studies of Involuntary Movements
PROJECT TITLE
7. Maitland Baldwin, M. D.
PRINCIPAL INVESTIGATOR
8. J. M. Van Buren, M. D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objective: There are certain diseases of the nervous system characterized by involuntary movements. At present, the precise cause of the majority of these debilitating conditions is unknown. Likewise, the precise action and sites of action of the various causal agents are not clear. However, the ultimate effects of many of these abnormal processes have been outlined. These effects have been analyzed through the various techniques of pathological anatomy.

If pathological anatomy can provide some definitive information, perhaps the correlation of these structural changes with clinical phenomena may provide further and equally useful information.

The most obvious clinical manifestation of these diseases is and has been the involuntary movements which seem to characterize their course.

Therefore, we have begun an analysis of these movements, with the hope that data thus acquired may be related to the definitive changes which have been reported as the ultimate effects of the disease processes.

In addition to our analysis of the involuntary movements, we hope to observe the changes in some of these movements which occur after certain accepted surgical treatments have been applied in selected cases.

Methods Employed:

1. Clinical Observation.
2. Photographic Techniques.
3. Analysis by illustration of movement phases.

Major Findings: The past year has been spent in the development of photographic and other means of illustrating the nature of involuntary movements in a small group of patients. This effort has provided us with a useful technique. In the use of this technique we find that the movements studied can be analyzed and this analysis indicates the essential characteristics of the movement pattern.

Major Findings (continued): In three cases we observed involuntary movements. It was necessary to remove the temporal lobe in each of these cases. This removal was followed by changes in the involuntary movements. Indeed, in two cases the movements ceased. In the other case, there was some change in the movements as well as in the rigidity.

Present knowledge does not serve to explain these findings. It may be that the amygdaloid nucleus has some relationship in the abnormal functions which underly some of these movement patterns.

Significance to Neurological Research: Diseases which cause involuntary movements are not understood and cannot be treated effectively. These studies will constitute one small step towards an increase in understanding.

Proposed Course of Project: The techniques of observation will be applied to selected patients and some of these patients will be treated by interruption of pathways which subserve motion. Wherever possible we will make anatomical analyses of the available material and correlate these findings with the clinical observations. Likewise we will begin correlation of this data with observations made in the laboratory on experimental animals.

Patient Material:

		No.	<u>Average Stay - days</u>
Admissions:	Adult Males	0	
	Adult Females	2	150
	Children Male	1	18
	Children Female	0	
Outpatients:	Number of Patients	1	
	Number of visits	1	

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
Neurosurgical Service
2. Surgical Neurology
BRANCH
3. Experimental Neurosurgery
SECTION
4. _____
LOCATION
5. NINDB-45 (C)
SERIAL NO.
6. Human Cortical Physiology
PROJECT TITLE
7. Maitland Baldwin, M. D.
PRINCIPAL INVESTIGATOR
8. J. M. Van Buren, M. D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: To outline the extent and nature of functional representations in the cortex of man.

Methods Employed: 1. Electrical stimulation of the human cortex under local anesthesia during operative exposures.
2. Observation of the apparent effect of selected cortical ablations in man and higher primates.
3. Electrographic recording during stimulation of the exposed human cortex and peripheral recording designed so as to analyze autonomic and motor responses which may follow this stimulation.
4. Electrical stimulation of the cerebral cortex in higher primates under analagous conditions with cortical and peripheral recording.

<u>Patient Material:</u>	<u>No.</u>	<u>Average Stay - Days</u>
Admissions: Adult Male	22	9.86
Adult Female	24	7.79
Children Male	6	24.67
Children Female	8	9.88
Outpatient: Number of Patients:	20	
Number of Visits:	24	

Major Findings: We have stimulated the human cortex in the frontal, temporal, parietal, and occipital regions. The majority of these stimulations have been undertaken under local anesthesia so that the subjective as well as objective responses might be observed and recorded. The majority of these stimulations have been undertaken over the wide area of the temporal cortex, since the majority of our cases suffer from temporal lobe lesions and the area of the lesion indicates the area of exploration and stimulation. We have elicited motor and sensory responses from the temporal cortex during the past year. These responses

PROJECT DESCRIPTION (continued)

which were evoked in man have also been tested in the laboratory and we have elicited motor responses from the temporal cortices of both monkeys and chimpanzees.

We have observed a change in cortical activity which seems to follow stimulation of the amygdala in the human and also in the chimpanzee. In the human, electrical stimulation of the amygdaloid nucleus is followed by a relative suppression of activity over the temporal and parietal cortices.

Significance to Neurological Research: Accurate observations of human cortical physiology have a wide clinical and scientific application. The analysis of these observations in terms of "brain maps" has served to integrate the various functions in terms of the entire cortex. Thus, through compilation of brain maps which summarize large numbers of responses, one may obtain ideas as to concentration of function within certain cortical areas.

Proposed Course of Project: We will continue the gentle electrical stimulation of the human cortex. During the coming year these stimulations will provide more data because we have now obtained new apparatus for the simultaneous recording of multiple peripheral responses and also we are now in a position to substantiate some of the information gained by direct cortical stimulation through the use of the evoked potential technique.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
Neurosurgical Service
2. Surgical Neurology
BRANCH
3. Experimental Neurosurgery
SECTION
4. _____
LOCATION
5. NINDB-46 (C)
SERIAL NO.
6. Epileptogenic Mechanisms in the Brain of Man
PROJECT TITLE
7. Matland Baldwin, M.D.
PRINCIPAL INVESTIGATOR
8. C. Ajmone-Marsan, M.D., J.M. Van Buren, M.D., Bruce Ralston, M.D., /
OTHER INVESTIGATORS
D. B. Tower, M. D., and E. C. Alvord, M.D.
9. PROJECT DESCRIPTION

Objectives: To study causal mechanisms of epileptic seizures in man:

- a. To study the characteristics of epileptogenic areas in the brain of man.
- b. To study the characteristics of the lesions which predispose to epileptogenic changes in the brain of man.
- c. To study the clinical expression of these epileptogenic areas in man.
- d. To study the electrographic characteristics of epileptogenic activity in the brain of man.
- e. To study the approved methods of surgical therapy for these lesions and develop new therapeutic methods.
- f. To study brain function as it is exposed in the extravagant experiments devised by these lesions.

Methods Employed:

- a. Clinical neurological examination.
- b. Special radiographic examination.
- c. Electrographic examination.
- d. Electrocortigraphic examination.
- e. Electrical stimulation of the lesion exposed at operation.
- f. Selective isolation of the lesion at operation.
- g. Cinematographic and sound recording.
- h. Histological and chemical examination.

Patient Material:

	<u>No.</u>	<u>Average stay - days</u>
Admissions:		
Adult Male	55	18.93
Adult Female	48	20.17
Children Male	23	42.65
Children Female	19	21.26
Outpatient:		
Number of patients:	76	
Number of visits:	124	

PROJECT DESCRIPTION (Continued)

Major Findings: During the past year we have studied the motor and autonomic phenomena of temporal lobe seizures in the human and in the chimpanzee. From these studies we now conclude that an epileptogenic lesion in one or the other temporal lobe sooner or later expresses its presence through movements of face and jaws, and these movements also begin on the side of the lesion. Thus observation of the initial movements in a temporal lobe seizure may serve to lateralize the essential epileptogenic lesion. It seems clear that these movements become bilateral in the majority of cases but in the cases we have observed in the chimpanzees which we have used as experimental preparations, these movements began on the same side as the essential lesion.

In almost all patients and in the majority of experimental animals, a temporal lobe seizure actually begins with a change in the width of the palpebral fissures. This widening is bilateral and is soon followed by a change in pupillary size. Thereafter the movements of face and jaw begin. It may be significant that the face usually changes in color as the pupils dilate and before salivation occurs. Salivation actually occurs in most cases of temporal lobe seizures. The first salivation begins at the same time as the movements of the face. At this time the secretion is slight and the mixture is mucoid. As the attack continues the quality of the secretion changes and it becomes watery. These autonomic changes may be accompanied by another, which is an alteration in respiration. This usually occurs at the same time as the movements of face and jaw begin and it is an inspiratory arrest. All of these autonomic phenomena are significant, and we have reproduced them experimentally in the chimpanzee and observed them frequently in the sequence described as they occurred in our patient material.

Posturing of the upper extremities is characteristic of the expression of epileptic activity in one or the other temporal lobe. This usually begins on the side opposite to the lesion and it occurs after the movements of face and jaws and coincident with the second type of salivation. We use the term "posturing" because the involved extremity becomes hypertonic before any movement occurs, and subsequently movements either of hand or arm are superimposed on a posture of either flexion or extension.

Many temporal lobe seizures are characterized by automatic behavior. This behavior is either ictal or postictal, and indeed it is difficult to place it within the sequence of the attack. However, in our cases it has usually occurred at the end of the attack or is a continuation of an attack characterized by the phenomena which we have described above. In a very few of the cases the entire attack consists of this bizarre behavior. In some cases speech is an outstanding feature during this alteration of consciousness. These are cases in which the essential lesion lies in the temporal lobe

PROJECT DESCRIPTION (Continued)

on the side of major speech representation. This speech is expressed either as jargon or as sentences inappropriate to the immediate environmental circumstances. In one case the patient replied to a direct question. When he was asked what he was doing, he said he was looking for something. However, his search was fruitless and he did not speak again.

We have reproduced fragments of this bizarre pattern in chimpanzees and recorded it in color movies. This reproduction occurred after the establishment of an afterdischarge in one amygdaloid nucleus and subsequent to the production of movements of face and jaws, autonomic changes, and posturing phenomena which we have described above. The animal plucked at himself, looked about the room, and seemed to search for something.

In the surgical treatment of temporal lobe seizures, we have achieved a greater understanding of the significance of the deeper structures in the production of these seizures. We have observed cases at operation in which the abnormal discharges were recorded infrequently from the temporal surface and with increasing frequency as we approached the deeper structures in the surgical dissection. In these cases we have also had an opportunity to record directly from amygdale, hippocampus and hippocampal gyrus and these recordings have provided examples of the epileptogenic activity at its apparent source.

In the course of such dissections, we have been able to modify the original Penfield technique so as to facilitate this operative procedure.

We have tested the effects of hypothermia on epileptic discharge in the amygdaloid system in chimpanzees, and we have also tested the effect of hypothermia on epileptic discharge which originated in the supplementary motor area in one human case. In the experimental cases, hypothermia seemed to dampen this discharge and reduce its spread. In the human case, the hypothermia did not materially alter the activity of the epileptogenic lesion. In the experimental animals we were dealing with the amygdaloid or limbic system, and in the human, with a more or less discrete cortical lesion lying on the lateral aspect of the convexity. It may be that this variation in effect gives some indication of variation in characteristics of these separate and distinct neuronal areas.

In seizure cases characterized by a spike-and-wave discharge, we have tested the effects of some of the drugs commonly used in the reduction of symptoms due to basal ganglia disease. These drugs seemed to alter the extent and range of the motor phenomena which characterized the expression of the spike-and-wave discharge. They did not alter the frequency of the attacks and in no way changed the fundamental alteration of consciousness which characterized each of

PROJECT DESCRIPTION (Continued)

these seizures.

We studied the effects of thorazine and serpasil in cases of temporal lobe and centrencephalic seizures. These drugs do not affect the frequency of attacks but they do tranquilize the patient.

We have observed the psychotic phenomena which sometimes accompany or occur as a result of epileptic discharge in one or the other temporal lobe. In three cases of temporal lobe seizures, we withheld anticonvulsant medication for a brief period. During this period each of these cases became confused, irrational, and incoherent. One case became aggressive and another patient showed unusual alteration in mood. In each patient there was evidence of psychosis and this was confirmed by competent psychiatric opinion. These psychological aberrations disappeared after resumption of anticonvulsant medication. They were coincident with epileptic discharges referable to one or the other temporal region.

In one case the frequency of the attacks was directly proportional to the degree of amnesia. This was a case of temporal lobe seizures in which the attack consisted entirely of a feeling of "having been there before". The deja vu phenomena was followed by a brief period in which consciousness was altered, the face changed color, the pupils dilated and the palpebral fissures widened. The entire duration of the attack never exceeded one minute. This patient's ability to record immediate and present experience, and at least her ability to recollect experiences of the immediate past, was radically altered by the frequency of these attacks. The number of attacks never exceeded five per day and they were usually widely spaced throughout a twenty-four hour period. As they decreased in frequency her recollective facilities improved and she was able to retain information derived from direct observation and audition. She could not retain this information adequately during the periods of frequent seizures.

In cooperation with the Department of Social Work we have begun and are continuing a study of the social factors which influence the family, community or institution which contain individuals who are epileptic.

In cooperation with the section of Clinical Psychology we are studying and have studied personality characteristics in these cases.

Significance to Neurological Research: The clinical observations which we have described above may be significant in the localization and lateralization of temporal lobe lesions which are epileptogenic. They also serve to further our understanding of functional representation within the human and higher primate temporal lobe. The effects of hypothermia on the epileptogenic lesion in the limbic system have been not noted previously. The fact that hypothermia seems to suppress epileptogenic discharge in this system and leaves it unaltered or

PROJECT DESCRIPTION (Continued)

actually accelerates it when it originates in the cortex, may be important in the further understanding of the special characteristics of the limbic system as compared to the overlying cortical areas.

Our observations concerning the use of artane and similar substances in centrencephalic epilepsy, bear further elaboration and may lead us to a better understanding of the motor phenomena which characterize these seizures.

The psychotic episodes which we have observed as an outward expression of epileptic activity in the temporal lobe are significant because their occurrence demonstrates that epileptic discharge in this region may cause psychosis. Such a psychosis presents a social, psychiatric, and legal problem and its origin must be clearly understood. It can be reversed and will disappear under effective treatment. When it occurs it resembles some form of adult schizophrenia and may be mistaken for an acute schizophrenic turmoil.

Memory difficulties usually follow temporal lobe seizures which are characterized by *deja vu* phenomena or by other psychical illusions. This is demonstrated in one case which we outlined above.

Proposed course of the Project: We will continue the clinical studies. With new methods of electrographic recording and stimulation, we must expand our area of observation in the human operating room and on the ward. In the primate laboratory we can now create chronic epileptogenic preparations in the chimpanzee through the use of implantation electrodes which can be controlled (remotely) by radio impulses. In this way we hope to study the behavior of these animals, before, during and after the incidence of these discharges in their temporal lobes, and also analyze the seizure patterns which occur as the result of these discharges. These applied-scientific techniques will be reinforced by observations of a more basic nature which we can accrue from the projects on second sensory areas, somato-sensory localization, and amygdaloid function. All of these projects can be integrated so as to give us a greater knowledge of the phenomena which characterize epileptogenic discharge in the temporal regions. In therapy, we have improved our surgical techniques and we have tested old drugs in new fashions. We hope to devise new surgical techniques and improvise newer pharmacological treatments.

Analysis of NIH Program Activities

Budget Data

10. NINDB-46 (C)
SERIAL NO.

11. BUDGET DATA

12. BUDGET ACTIVITY: RESEARCH

13. NONE.

14. NONE.

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955:

17. HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

R.P.C. 1
Dec. 1955

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Neurosurgical Service
SECTION
4. _____
LOCATION
5. NINDB-47 (C)
SERIAL NO.
6. Pain Mechanisms
PROJECT TITLE
7. J.M. Van Buren, M.D.
PRINCIPAL INVESTIGATOR
8. None
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: A study of patients suffering from painful lesions has been undertaken to provide material for the following studies which will be further elaborated as future evidence indicates: (1) Evaluation of the standard surgical procedures for the relief of pain. (2) Evaluation of standard medical procedures for the relief of pain including some of the newer medications such as stildamidine for the relief of trigeminal neuralgia. (3) Evaluation of objectively recorded autonomic responses by the use of a polygraph in patients with varying degrees of pain. In this study the anticipatory autonomic response to a known and forewarned painful stimulus will be evaluated as well as an attempt made to obtain some quantitative measurement of pain threshold in patients both free of and suffering from pain. The conjunction of these studies with the above mentioned clinical studies may give more objective evidence to the evaluation of these treatment programs. (4) The section of sensory pathways (pain pathways) affords the opportunity for study of patients with these surgical lesions. The advantage of study in a surgical rather than a neoplastic or diffuse vascular lesion is obvious, in that the lesion is more precise in type and is usually of known dimensions.

Methods Employed: The methods to be used in treatment of pain problems are to include the standard surgical procedures for the relief of pain, either in the peripheral nerve, dorsal root, or central pain pathways. The medical treatment of pain will likewise follow well approved and standard patterns.

PROJECT DESCRIPTION (Continued)

The new feature which will be added to the study of pain is by the use of a polygraph in which blood pressure by an auscultatory method (no arterial puncture) electrocardiogram, respirations, skin resistance, the pressure variations from a plethysmograph and pressure variations in a hollow viscus such as bladder and/or stomach may be recorded. By the simultaneous recording of these data it is hoped that further insight may be gained into the autonomic responses to the painful stimulus in man. At the present time, considerable difficulty in instrumentation is being encountered. Apparently an instrument of this type for simultaneous recording of many modalities has not previously been constructed so that the investigator is undertaking the fabrication and collection of the proper recording instruments. This has delayed progress on the project considerably.

Additional studies of the patient following a central nervous system lesion of known dimension will be undertaken on a clinical level. Problems of interest concern the central pathways for sweating in man which as far as the author is aware have not been elucidated above the upper cervical cord. Another problem of basic nature in which the lesions could be used, is for the study of the pathways of conduction of the muscle spindle afferent impulses.

Patient Material:

		No.	<u>Average Stay - days</u>
Admissions:	Adult Male	4	12.25
	Adult Female	9	36.22
	Children Male	0	
	Children Female	0	
Outpatient:	Number of patients:	15	
	Number of visits:	33	

Major Findings: Work so far has been directed toward instrumentation.

Significance to Neurological Research: There is preliminary evidence that the autonomic responses to pain and other stimuli in patients suffering pain is enhanced. This will be investigated as possible basis for objective assessment of the individual's response to pain.

Proposed course of the Project: The major hurdle at the present time consists in the proper preparation of the polygraph for autonomic recordings. For this reason the major patient load for this project has been held up.

PROJECT DESCRIPTION (Continued)

types of lesions: (1) Pseudolaminar necrosis. This lesion is loss of cerebral substance including neurons and glia, limited largely to the central layers of the cerebral cortex. In this area the cerebral substance appears spongy, and is frequently completely destroyed. Lesser changes consist in a loss of neurons in this area. This change is apparently secondary to interference with the fine penetrating blood vessels from the surface of the cerebral cortex. Collateral Golgi stains of cerebral cortex to show the endothelium of the capillaries demonstrates that the maximum density of the capillary network lies in these central layers of the cortex, probably in layer III to upper V. From this, the inference perhaps follows that this area of the cerebral cortex requires the greatest blood supply. It has also been shown by other work that in the lower portion of the cortex a junction is made between the blood supply from the surface and from the white substance. Hence, it is apparent why this area of cerebral cortex is the most vulnerable to mild anoxia.

(2) Complete cortical necrosis. This lesion is apparently a continuation of the above-mentioned process in which none of the cerebral cortex remains.

(3) Fats and lipid deposition. In many epileptogenic lesions in which there is lack of severe pathological change phagocytes filled with lipid which will stain with sudan IV as well as scharlach have been found in the perivascular spaces and as well at times free in the cerebral substance apparently at a distance from vessels. The origin of this lipid is obscure as in some cases there is no obvious breakdown of myelin in tissue.

(4) Neuron damage. Damage to the neurons has appeared in many of the sections. The most common change is that of the dense pyknosis of the cell, shrinkage of the cell in decreasing its lateral dimensions and tortuosity of the apical dendrite. It is considered that in many instances this is a traumatic artifact since there frequently is no surrounding increase in the pericellular satellites. A similar change has been shown in a recent paper by Cecile Vogt, published in the Annals of the First International Congress of Neuropathology.

(5) Fibrous astrocytosis of the cortex. In this change, best demonstrated on the metallic impregnations, fibrous astrocytes appear in layers of the cortex in which they are normally absent. They are normally present, of course, in the molecular layer, and occasionally in the VIth layer of the cortex.

(6) The meningo-cerebral cicatrix. This is the classical brain scar in which there is a union of the fibrous tissue elements of the dura with the fibrous framework of the cerebral cortex itself, namely the tissue about the blood vessels. There is general hypertrophy of these structures, a fibrillary astrocytosis and loss of fine capillary supply to the affected cerebral cortex. Or, there may be a considerable quantity of debris arising from myelin destruction, and old blood pigment which is deposited in the phagocytes.

PROJECT DESCRIPTION (Continued)

Significance to Neurological Research: Although the basis of epilepsy as yet remains obscure, basic studies on the structural changes of epileptic lesions are necessary for an illucidation of its mechanism.

Proposed course of the Project: It is hoped by further study in this direction, particularly with cytochemical methods, that further knowledge of basic importance may be uncovered in this direction.

R.P.C. 2
Dec. 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB-48 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH

13. None

14. None

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955:

None

17. HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Neurosurgical Service
SECTION
4. _____
LOCATION
5. NINDB-49 (C)
SERIAL NO.
6. Effect of Tumors upon Central Nervous System Function and Structure.
PROJECT TITLE
7. J.M. Van Buren, M.D.
PRINCIPAL INVESTIGATOR
8. M. Baldwin, M.D. and Philip Rubin, M.D.
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION

Objectives: This study has a dual aim: 1) to carry out physiological-anatomical correlations in man, 2) to evaluate the effects of newer methods of treatment for tumors of the central nervous system. In the present program attempt will be made to utilize the intrusion of disease upon the central nervous system of man as an "experimental" lesion.

Although much can be accomplished through animal experimentation in determining the function of structures within the nervous system it is apparent that the need for confirmation of these findings in man is ever present. In addition, studies of function which cannot be properly evaluated in animals (e.g., sweating) must be carried out in man. Although the fruits of a study of this type are of widespread nature, particular attention has been placed upon three major features: (1) To note the effects of neoplasm upon the function of the optic pathways of man and particularly upon the correlation of the visual fields and the anatomical changes in retina, geniculate body and calcarine cortex. (2) To study, particularly in cases of slowly expanding lesions of the upper brain stem, the effects of these lesions upon personality, consciousness and the electroencephalogram. Since the majority of these lesions are of terminal nature, opportunity therefore will arise for clinical-anatomical correlation.

PROJECT DESCRIPTION (Continued)

Methods Employed: The evaluation on a clinical basis of the position and nature of the tumor will be determined as can best be done by history, neurological examination, contrast examinations, and surgery.. The symptoms under consideration will be evaluated using largely clinical methods, although it is hoped that the polygraph needed for autonomic evaluation will be of aid in this direction. In the post-mortem examination, careful serial examination of the involved structures will be undertaken both grossly and microscopically and charts prepared from this material. By the use of these charts and the method of overlapping areas it is hoped that features of localizing anatomical significance may be discovered.

Patient Material:

		<u>No.</u>	<u>Average Stay - days</u>
Admissions:	Adult Male	17	24.88
	Adult Female	15	19.07
	Children Male	1	19.0
	Children Female	4	37.25
Outpatient:	Number of patients:	12	
	Number of visits:	33	

Major Findings: With regard to the visual field pathways in man, it has been found that a lesion affecting the pre-geniculate pathways will cause degeneration in the retina supplied by these fibers which is obvious on microscopic examination. In cases in which there has been partial loss of ganglion cells within the retina it is possible to correlate visual acuity and visual field with two-dimensional reconstructions of the retina. At the present time data regarding the effects of brain tumors upon the level of consciousness, personality and EEG are too few for evaluation. It is too early to evaluate the results of therapy.

Significance to Neurological Research: Neurophysiological data in animals is extensive but research of this type if it is to be helpful in the evaluation or treatment of human disease must be confirmed in man. Since critical evaluation of vision, personality and consciousness cannot be made in animals and the human EEG is species specific in type, re-evaluation of these phenomena in man is necessary. The new surgical-radiotherapeutic techniques have not as yet been carefully investigated in any considerable series of tumors of the central nervous system.

Proposed course of the Project: The accumulation of a large number of cases for a study of this type in which overlapping areas of involvement must be charted is imperative before results of any reliability can be achieved. Thus our present problem is one of collection of data.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness INSTITUTE
2. Surgical Neurology BRANCH
3. Neurosurgical Service SECTION
4. _____ LOCATION
5. NINDB-50 (C) SERIAL NO.
6. The Pneumographic Effects of Non-Expanding Epileptogenic Lesions PROJECT TITLE : of the Temporal Lobe.
7. J.M. Van Buren, M.D. PRINCIPAL INVESTIGATOR
8. Maitland Baldwin, M.D. and E. Alvord, M.D. OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: This study has been undertaken to elucidate the effects of atrophic lesions upon the outlines of the temporal horn. There have been no previous studies of the lateralizing and localizing value of minor changes and the configuration of the temporal horn in epilepsy.

Methods Employed: The anatomy of structures outlining the temporal horn were reviewed in the embryo and in normal adult human brains. The variations of normal were reviewed in this connection. In 41 cases of proven epilepsy in the temporal lobe the pneumographic evidence was reviewed and the temporal horns measured in a systematic manner.

Patient Material:

		No.	<u>Average Stay - days</u>
Admissions:	Adult Male	18	-
	Adult Female	23	-
	Children Male	-	-
	Children Female	-	-
Outpatient:	Number of patients:	0	
	Number of visits:	0	

PROJECT DESCRIPTION (Continued)

Major Findings: It became apparent that the features outlining the temporal horn consisted of the amygdala, the hippocampus, the roof of the temporal horn (formed largely by the tapetum), and the collateral eminence. Depending on the position of the collateral eminence, four types of normal temporal horns can be distinguished on the sagittal views of the temporal horn. In addition, instances were found in which adhesions had occurred between the lateral border of the hippocampus and the lateral wall of the temporal horn. This might be a cause for lack of filling of the lateral cleft. This defect was not recognized, however, in the clinical material.

After carefully considering the problem, it appears that we were in no position to determine the limits of normality in any of the measurements of the temporal horn. Consequently, the evidence as obtained from history, physical examination, electroencephalography and the findings at craniotomy, including electrocorticography were taken as the absolute criteria for lateralization of the epileptic discharge in the temporal lobe. Arbitrary criteria were then applied to all of the measurements of the temporal horn. These arbitrary criteria were varied to obtain the maximum degree of agreement between the findings of what was considered abnormality on this basis and the actual lateralization of discharge within the temporal lobe. The criteria finally arrived at as indicating enlargement and therefore significant atrophy within the temporal lobe were (1) over 3 mm. greater height of the temporal horn on the affected side, or (2) over 4 mm. in width of the lateral cleft on the affected side. In cases in which the lateral cleft on both sides was greater than 4 mm., the larger one was selected as showing abnormality. Using these criteria alone, of the 28 cases considered to show sufficient filling of the temporal horns for evaluation, 12 showed no asymmetry, 13 showed evidence of dilatation of the temporal horn on the side from whence the epilepsy arose, while 3, on the basis of this criteria, were lateralized to the wrong side.

Significance to Neurological Research: It had been felt that of the criteria used for lateralization and localization of epileptic discharge in seizure discharge from the temporal region, that the pneumoencephalogram usually provided the least information. On the basis of this present study, and better criteria for evaluation of abnormality in the temporal horn, it has been found that in cases with adequate filling nearly 50% show some degree of enlargement of the temporal horn of sufficient degree to suggest lateralization from the pneumographic evidence alone.

Proposed course of the Project: This project has been finished and the report is in press with Acta Radiologica.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness INSTITUTE
2. Surgical Neurology BRANCH
3. Neurosurgical Service SECTION
4. _____ LOCATION
5. NINDB - 51 (C) SERIAL NO.
6. A Histological Study of Papilledema using the Golgi Method in Man. PROJECT TITLE
7. J.M. Van Buren, M.D. PRINCIPAL INVESTIGATOR
8. None OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objective: It is the object of this work to further advance knowledge as to the precise changes occurring in the eye in papilledema. Although sporadic papers regarding the cytopathology of papilledema have appeared in the past 50 years, in only one instances (Cone and MacMillan) have attempts been made by the use of specific metallic methods to stain completely the structures undergoing change of papilledema. As far as is known to the author no studies with the Golgi method have been made previously in adult human autopsy material.

Method Employed: It is of primary importance to fix the retina immediately following death. In order to accomplish this, concentrated formalin is injected within the globe at the bedside. By the time the post-mortem examination is undertaken, the retina has been completely fixed and the posterior part of the eye can be removed without damaging the retina. Fixation is then continued in 10% formalin until it is convenient to undertake the Golgi staining. The staining method is a combination of potassium bichromate and potassium acetate mixture in which the eye is incubated at 37 degrees for 10 days to 2 weeks. It is then transferred to a warm 1% silver nitrate solution for 48 hours. The eyes are then rapidly dehydrated in absolute alcohol and transferred immediately to thin celloidin in which they are kept for 48 hours. Following this they are transferred to thick celloidin for a few minutes, then transferred to a mounting block, thick celloidin poured over the specimen, and the whole specimen and the mounting block immediately immersed in chloroform for hardening. Thus the eyes are not infiltrated but rather included in the celloidin. This inclusion is sufficient to permit sectioning at 100 micron intervals which is needed for this method.

PROJECT DESCRIPTION (Continued)

Patient Material:

	No.	Average Stay - days
Admissions: Adult Male	2	-
Adult Female	1	-
Children Male	0	
Children Female	0	
Outpatient: Number of patients:	0	
Number of visits:	0	

Major Findings: To date, the following has been apparent: There are definite, well defined grape-like swellings upon the optic nerve fibers in the optic papilla. These in all instances have appeared on the retinal side of the lamina cribrosa. A careful study of the fiber structure of the lamina cribrosa itself has been undertaken and indications are that in addition to bending away from the cavity of the eye with papilledema that there is also a separation of lamina cribrosa from the margin of Bruch's membrane. The nerve fibers have also been stained at the margin of Bruch's membrane, and their confinement by the sharp edge of this membrane is apparent. They make a very sharp detour around this edge, and it is not difficult to see how they might be injured by the increased pressure on this sharp margin.

Significance to Neurological Research: The sharp angulation of the peripheral retinal fibers over the margin of Bruch's membrane may possibly find its clinical counterpart in the constriction of the peripheral portions of the visual field. Further study must be undertaken on this subject. The precise cause and structural concomitants of papilledema have not been elucidated to date in the literature.

Proposed course of the Project: At the present time the eyes from only three patients have proved suitable for study. More material is needed before detailed results can be advanced.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness INSTITUTE
2. Surgical Neurology BRANCH
3. Neurosurgical Service SECTION
4. _____ LOCATION
5. NINDB-52 (C) SERIAL NO.
6. The Histological Investigation of a Case of Retinal Moniliasis. PROJECT TITLE
7. J. M. Van Buren, M.D. PRINCIPAL INVESTIGATOR
8. None OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objective: The chance finding of a case of retinal moniliasis and the availability of the retina for detailed examination has prompted a thorough study of the histological features. As far as is known, no previous case of monilia infection of the retina has been recorded.

Method Employed: The eyes were obtained at post-mortem some 5 hours following death. They are fixed in formalin, imbedded in paraffin, and then serially sectioned by the investigator so that the small monilia lesion would not be overlooked. Routine histological stains were carried out including the PAS which demonstrated the monilia hyphae within the granulation tissue.

Patient Material:

	<u>No.</u>	<u>Average Stay - days</u>
Admissions: Adult Male	-	-
Adult Female	1	-
Children Male	-	-
Children Female	-	-
Outpatient: Number of patients:	-	-
Number of visits:	-	-

Major Findings: The lesion in the retina was thought to have had its origin just below the internal limiting membrane, presumably being an embolic infection which had lodged in a small vessel. It had depressed the subjacent retina, considerably thinning out the ganglionic cell layer and the internal molecular layer. Exudate had broken through the internal limiting membrane and had spread in the subhyaloid space. Indeed, leucocytes were present over the papilla some distance from the initial infection.

R.P.C. 1 (a)
Dec. 1955

NINDB-52 (C)
SERIAL NO.

PROJECT DESCRIPTION (Continued)

Significance to Neurological Research: Although the case is an isolated one, the fact that it is apparently new to the ophthalmological literature suggested that it was worthy of review and report.

Proposed course of the Project: Additional histological work remains to be done, literature review must be continued and the report written in final form.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness INSTITUTE
2. Surgical Neurology BRANCH
3. Neurosurgical Service SECTION
4. _____ LOCATION
5. NINDB-53 (C) SERIAL NO.
6. Study of the Adult Human Retinal Cytology using Golgi and Chromatic Methods. PROJECT TITLE
7. J.M. Van Buren, M.D. PRINCIPAL INVESTIGATOR
8. None OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objective: This study of normal retinal cytology is undertaken as a basis for the evaluation of degenerative lesions in the retina. As far as can be determined, no previous work has been carried out using the Golgi method in the adult human retina. A considerable portion of the available reports of retinal cytology are confused by post-mortem artifact. By the use of early retinal fixation these artifacts may be eliminated.

Method Employed: The study will be carried with human retinas fixed within half an hour of death by intra-ocular injection of formalin. Some will be stained by the Golgi and others by chromatic methods.

The Golgi method will be used primarily for study of the conformation of individual retinal elements, while the chromatic methods will be used primarily for the study of retinal cyto-architecture. In the latter connection, serial sections will be made of the eyes, and from these, 2 dimensional reconstructions of the retina and its various cell fields can be made.

Patient Material:

	No.	<u>Average Stay - days</u>
Admissions: Adult Male	2	-
Adult Female	2	-
Children Male	-	-
Children Female	-	-
Outpatient: Number of patients:	-	-
Number of visits:	-	-

PROJECT DESCRIPTION (Continued)

Major Findings: To date, perhaps the most important finding has been that human post-mortem material has proved suitable for Golgi staining. As far as is known, a Golgi method suitable for routine post-mortem work has not been previously employed. At the present time, the nature of various components of the retina is being investigated by this means.

Using the chromatic method, variations in the types of ganglion cells have been found from the central fovea to the periphery, and the varying thickness of the ganglion cell layer as described by other investigators has been confirmed. At the present time, only four retinas serially sectioned have been obtained in which visual fields were done prior to death. The correlation between this retinal degeneration and the visual field is of primary importance. It is planned to convert retinal distances as measured off in the various cell fields into visual angle. From this, a simultaneous charting of the retinal cell fields and the visual fields as determined clinically can be carried out. It is hoped that this anatomical-clinical comparison may prove fruitful.

One interesting variant has been found on the chromatic stains which consists in the not infrequent displacement of ganglion cells and other retinal elements outside of their commonly accepted zones.

Significance to Neurological Research: Although Golgi studies of retinæ in low animals has been carried out, it seems desirable to confirm the cytological findings in man. Such an opportunity now presents itself.

A further study in the normal variants in the human retina is necessary to determine the limits of normal.

The correlation between the anatomical configuration of the retina and the visual field is obviously of basic importance to the proper evaluation of visual field studies.

Proposed course of the Project: More human autopsy material is desirable. The methods as devised above have proved satisfactory and must be applied to more cases. Further reconstructions must be made in cases of serially sectioned retinæ presently available.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Neurosurgical Service
SECTION
4. _____
LOCATION
5. NINDB-54 (C)
SERIAL NO.
6. The Effect of Lesions of the Optic Pathways upon the Retina.
PROJECT TITLE
7. J.M. Van Buren, M.D.
PRINCIPAL INVESTIGATOR
8. None
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objective: Study will be made of the effects of degeneration upon the areal pattern of ganglion cells in the retina by the use of reconstruction technique. Attention will also be paid to other retinal layers to determine the degree of transsynaptic and transneuronal degeneration. As far as the author is aware, only one case of retinal degeneration following a lesion of the pregeniculate pathway has been reported in man. In no instances have retinal changes from postgeniculate changes been recorded in man, although Polyak has recorded one instance to the author's knowledge in a monkey. Human material is presently available for evaluation, and lesions have been placed in macaques some 12 months ago, which are presently ready for study.

Method Employed: The eyes were fixed by intra-ocular injection of formalin, marked for orientation on the slide, imbedded in paraffin and serially sectioned. Degenerated areas were marked out with ink on the slides, the slides projected at 35x magnification and traced upon paper. The degenerated areas on the tracing were measured, then from these measurements a two-dimensional reconstruction made. Conversion of retinal distances to degrees of visual angle permitted the superimposition of the areas of degeneration upon the clinically recorded visual field.

Patient Material:

	No.	<u>Average Stay - days</u>
Admissions:		
Adult Male	1	10
Adult Female	-	
Children Male	-	
Children Female	-	
Outpatient:		
Number of patients:	-	
Number of visits:	-	

PROJECT DESCRIPTION (Continued)

Major Findings: In both monkeys and a human case it has been found that lesions of the pre-geniculate optic path are followed by a definite degeneration in the retina. If, indeed, complete section of an optic tract is carried out, this degeneration will encroach upon one-half of the fovea and extend above and below this foveal point in a vertical straight line. By the use of reconstruction studies in the human case, the conversion of retinal distances into visual angles, and the simultaneous charting of degenerative areas in the visual field, it has been possible to compare the anatomical with the clinically determined findings. In like fashion, it has been possible to compare the visual acuity with the degree of retinal degeneration.

Significance to Neurological Research: These studies of purely destructive regions of the geniculate optic pathway are intended to elucidate the pathological changes in the retina following such a lesion. A clinical anatomical correlation is believed of basic importance for the understanding of the significance of visual field studies.

Proposed course of the Project: Although there is no lack of animal material, every possible advantage will be made of additional human material as it presents itself. It is felt that the major emphasis must be upon human material as this is an unexplored field.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Neurosurgical Service
SECTION
4. _____
LOCATION
5. NINDB-55 (C)
SERIAL NO.
6. The Effect of Lesions of the Optic Pathways upon the Geniculate Body.
PROJECT TITLE
7. J.M. Van Buren, M.D.
PRINCIPAL INVESTIGATOR
8. None
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objective: Since the visual field changes following temporal lobectomy have been carefully observed in another study of this series, it has been thought of interest to study the geniculate representation of this visual field change. As far as the author is aware, no previous studies of the geniculate body following temporal lobectomy in man have been carried out.

Method Employed: The brains from two patients in which the temporal lobes were removed were fixed in 10% formalin, the slices containing the geniculate bodies separated, imbedded in celloidin and serially sectioned, keeping sections at 500 micron intervals. The sections were stained for Nissl substance and myelin. The sections were then photographed at 50 x magnification, and the entire geniculate body printed at this size. The degenerative lesion, therefore, could be made out on the photographic prints and its precise dimensions and details made out by simultaneous comparison under microscope. By this means, the layering and the lesion could be outlined with pencil, subsequently drawn over in india ink, and then rephotographed to diminish the size to a degree that would permit publication. If it is desired later for demonstration purposes, 3 dimensional model can be made.

Patient Material:

	No.	Average Stay - days
Admissions:		
Adult Male	2	-
Adult Female	-	
Children Male	-	
Children Female	-	
Outpatient:		
Number of patients:	-	
Number of visits:	-	

PROJECT DESCRIPTION (Continued)

Major Findings: It has become apparent that following temporal lobe lesions, degeneration does not extend to all six layers of the geniculate body, as has been postulated with lesions in the occipital lobe. Since the visual field changes do not affect the macular portion of the visual field, it gives some suggestion that the macular representation may lie somewhat deeper within the geniculate body than the portions subserving upper quadrantic gaze which are destroyed in temporal lobectomy.

Significance to Neurological Research: This study is intended as a further approach to clinical-anatomical correlation of visual function in man.

Proposed course of the Project: The above conclusions were reached on a study of one case only. The other case remains to be charted and reconstructed. It is hoped that it will be possible to obtain a third case for confirmation. Later on, when some chimpanzees with temporal ablations are autopsied, further confirmation of the degeneration in the anthropoid ape will be obtained.

Analysis of NIH Program Activities

Project Description Sheet

- | | | |
|--|--|--------------------------------------|
| 1. <u>Neurological Diseases and Blindness</u>
INSTITUTE | 2. <u>Surgical Neurology</u>
BRANCH | |
| 3. <u>Experimental Neurosurgery</u>
SECTION | 4. _____
LOCATION | 5. <u>NINDB-56 (C)</u>
SERIAL NO. |
6. Microelectrode Studies of the Cerebral Cortex in Experimental Animals and Man.
7. Choh-luh Li, M. D.
8. Maitland Baldwin, M. D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Project: Development of methods for recording electrode activity of single cells from the human and animal cortex.

Objectives: To develop methods which can be adapted in the human operating room for the recording of electrical activity of single cortical cells, particularly in epileptic patients, with the purpose of studying the mechanism of epilepsy.

To apply these methods to the human cortex and to the cortex of experimental animals. After this application to test the effects of stimulation of the deep-lying nuclei on single unit activity in the cortex to which these methods were applied.

Methods Employed: Microelectrodes made of glass capillary pipettes with an inside diameter of less than half a micron are used for recording electrical activity from single cells.

This microelectrode technique was applied with a number of variations. The effect of strychninization was studied in experimental animals and the effect of recruiting responses was also observed. Likewise, this technique was so adapted that it might be used to study the action of resting potentials of cortical neurones.

Major Findings: The major findings of this project during the past year are:

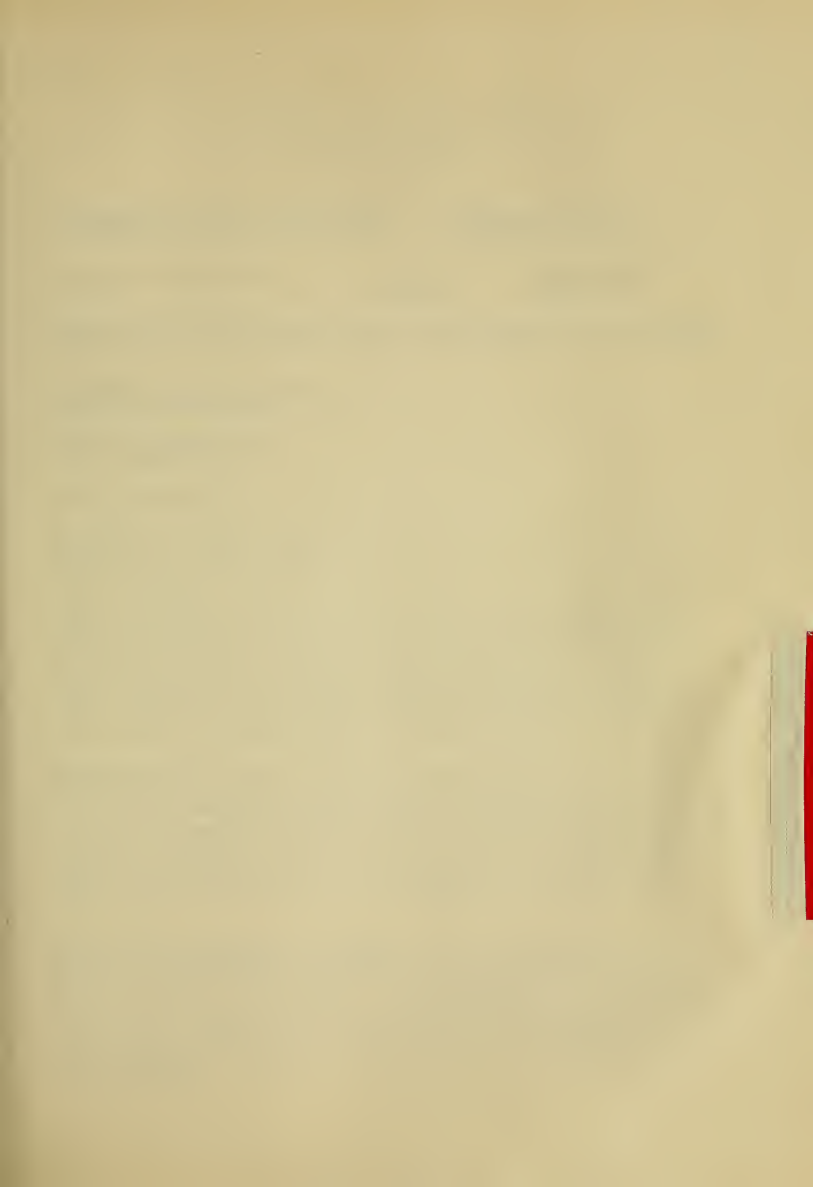
- (a) Recording of single unit potential activity from the human cortex in two cases and from the chimpanzee cortex in two of these experimental animals. Furthermore, we have studied the functional properties of cortical neurones with particular reference to strychninization, as well as the effect of recruiting responses of evoked neuronal discharge in experimental animals.
- (b) More recently, we have studied the action and resting potentials of these single cortical units and observed the facilitatory effect of stimulation of unspecific thalamic nuclei on cortical sensory neuronal responses, and finally, the inhibitory effect on cortical moto-

Major Findings (continued)

neuronal activity of stimulation of thalamic nuclei on single motor units in the cortex of experimental animals. We have also observed that single units in the motor cortex are inhibited after deep stimulation of certain thalamic nuclei.

Significance to Research: The study of single unit potentials and the characteristics of single neuronal units in the higher primate nervous system has great significance to the better understanding of function. The single cortical unit is basic in the relationships of the various cortical areas and its individual characteristics have not been studied. Furthermore, these individual characteristics as observed in the normal human and experimental animals may lead us to further conclusions concerning the activity of such neurones under the abnormal conditions of the epileptogenic cortical lesion.

Proposed Course: During the next year we will study the effects produced in single cortical units following deep stimulation of thalamic and other nuclei. We will also study the human epileptogenic cortex with the microelectrode techniques and we will study normal and epileptic preparations in man and the higher primates.





Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Clinical Neuropathology
SECTION
4. _____
LOCATION
5. NINDB-57 (C)
SERIAL NO.
6. Relation of Supraspinal Structures to an Experimental Startle Reaction
PROJECT TITLE
7. Ellsworth C. Alvord, Jr., M.D.
PRINCIPAL INVESTIGATOR
8. David C. Whitlock, M.D.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: To determine the neuroanatomical and physiological substrata of the startle reaction.

Methods Employed: Acute and chronic lesions of the central nervous system of cats have been made in various locations to determine the effects of such lesions upon the generalized muscular contractions which can be reflexly evoked by various suddenly applied stimuli in cats under chloralose anesthesia. Other experiments have been performed utilizing a stereotaxically oriented electrode to explore the C.N.S., both by recording evoked electrical potential from and by applying single electrical stimuli to localized regions of the C.N.S.

Major Findings: In brief, it has been found that the sensory motor area of the cerebral cortex influences the sensitivity of the animals to the startle reflex while under light chloralose anesthesia. This was tested by removal of the sensory motor area. Then while under chloralose anesthesia, the testing of the response to tactile stimuli were carried out a week later. It was found that the threshold was raised on the portion of the body corresponding to the area of de-cortication.

Significance to Neurological Research: Study of the startle reaction provides basic information concerning the organization of the nervous system, particularly with respect to its more diffusely organized functions, and is of importance in the understanding of reflex activity in general and perhaps of reflexly induced epileptic seizures in experimental animals and patients. Since chloralose is frequently used in neurophysiological experiments, further knowledge of its actions is also important.

R.P.C. 1 (e)
Dec. 1955

NINDB-57 (C)
SERIAL NO.

PROJECT DESCRIPTION (Continued)

Proposed course of the Project: At the present time, the paper is being prepared by the investigators for publication. Dr. Alvord has transferred his work to Baylor University.

R.P.C. 2
Dec. 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB-57 (C)
SERIAL NO.

12. _____
BUDGET ACTIVITY: RESEARCH

Honors, Awards and Publications

16. _____
PUBLICATIONS OTHER THAN ABSTRACTS FROM THIS PROJECT DURING 1955:

None

17. _____
HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Clinical Neuropathology
SECTION
4. _____
LOCATION
5. MINDB- 58 (C)
SERIAL NO.
6. Pathogenesis of "Allergic Encephalomyelitis"
PROJECT TITLE
7. Ellsworth C. Alvord, Jr., M.D.
PRINCIPAL INVESTIGATOR
8. Norman Goldstein
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: To determine the cause and the mechanism underlying the development of disseminated encephalomyelitis.

Methods Employed: Brain and various extracts of brain are prepared in a water and oil emulsion with killed acid-fast bacilli. Following subcutaneous inoculation into guinea pigs, the animals are examined for clinical evidence of weakness, incoordination, or other neurological disorder. The spinal cord and brain are examined histologically for presence of characteristic perivascular inflammation sometimes with demyelination.

Major Findings: A total of 450 animals have been studied. 50 animals were killed two days following the injection of the brain and brain extract material in order to study the early lesions. The remaining 400 animals served as control and were injected with different brain fractions and sacrificed at the end of the 30 day period. The earliest lesions which were seen on the second day consisted in perivascular inflammatory response usually in the lower brain stem and spinal cord. The incidence of inflammatory lesions rose considerably after this time as was expected. At the present time the experimental work has been completed.

Significance to Neurological Research: Study of the etiology and pathogenesis of "allergic" encephalomyelitis in experimental animals will yield information of importance to patients treated with vaccines containing brain material, as in anti-rabies inoculations, and may also yield pertinent information on diseases which appear histologically quite similar and may be related, as in other post-vaccinal and post-infectious encephalomyelitis. Its relation to multiple sclerosis and other demyelinating diseases remains to be determined.

R.P.C. 1 (a)
Dec. 1955

NINDB- 58 (C)
SERIAL NO.

PROJECT DESCRIPTION (Continued)

Proposed course of the Project: The findings in the cases of early allergic encephalomyelitic lesions are being correlated by Dr. Kenneth McGee for publication. The results in the other 400 animals used to evaluate the effectiveness of various chemical fractions of the brain in producing allergic encephalomyelitis is being reviewed by Dr. Kies at NIMH and Dr. Alvord at Baylor University.

Analysis of NIH Program Activities

Budget Data

10. NINDB - 58 (C)
SERIAL NO.

12. _____
BUDGET ACTIVITY: RESEARCH

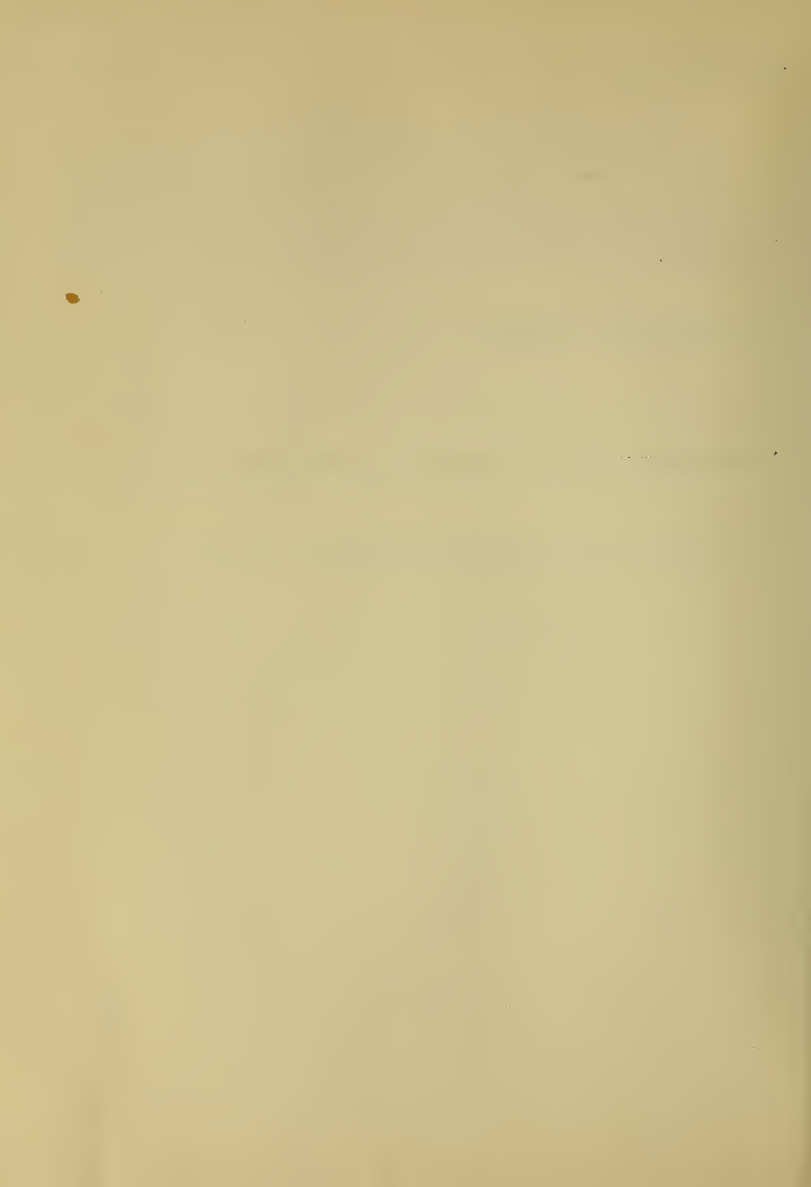
Honors, Awards, and Publications

16. _____
PUBLICATIONS OTHER THAN ABSTRACTS FROM THIS PROJECT DURING 1955:

None

17. _____
HONORS AND AWARDS TO PERSONNEL DURING 1955:

None



Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Clinical Neuropathology
SECTION
4. _____
LOCATION
5. NINDB-59 (C)
SERIAL NO.
6. Neuropathologic Investigation of Focal Epilepsy and other Neurological Diseases
PROJECT TITLE
7. Ellsworth C. Alvord, Jr., M.D.
PRINCIPAL INVESTIGATOR
8. Dr. Brace
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives: Experiments have been performed to determine the effects of high doses of X-irradiation on the nervous system of guinea pigs.

Major Findings: 50 animals have been received for histopathologic study with particular emphasis on changes in the cerebellar granule cells. Interest centers especially about the time course of these changes, their variation with different doses of X-ray delivered to different parts of the head and body, and their modification by drugs such as nembutal.

Significance to Neurological Research: The experiments in animals provide an opportunity to analyze the means by which the brain reacts to various injurious agents such as X-radiation, which are being used in increasing dosage for therapy in human neoplastic disease.

Proposed course of the Project: This paper has been finished and was presented at the Second International Neuropathological Congress in London under the title of "X-Ray Induced Pyknosis of Cerebellar Granule Cells in Guinea Pigs." It was also submitted to the Journal of Neuropathology and Experimental Neurology.

Analysis of NIH Program Activities

Budget Data

10. NINDB - 59 (C)
SERIAL NO.

12. _____
BUDGET ACTIVITY: RESEARCH

Honors, Awards, and Publications

16. _____
PUBLICATIONS FROM THIS PROJECT DURING 1955:

1) X-Ray Induced Pyknosis of Cerebellar Granule Cells in Guinea Pigs.
Submitted to the Journal of Neuropathology and Experimental Neurology.

17. _____
HONORS AND AWARDS TO PERSONNEL DURING 1955:

None

R.P.C. 1
Dec. 1955

* Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Clinical Neuropathology
SECTION
4. _____
5. NINDB-60 (C)
SERIAL NO.
6. Neuropathologic Investigation of Focal Epilepsy and Neurological Diseases
PROJECT TITLE
7. Ellsworth C. Alvord, Jr., M.D.
PRINCIPAL INVESTIGATOR
8. John Lilly
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objective: To determine the relative degree of damage to the cerebral cortex by repetitive electrical stimulation with different wave forms of electrical current.

Major Findings: One monkey after 7 weeks of daily prolonged stimulation by electrical currents of special wave characteristics showed moderate gliosis of the cortex with good preservation of the neurons. Other animals are now being prepared in attempt to control the effects of operative trauma and of stimulation by different electrical wave forms to test the hypothesis that repetitive electrical stimulation need not necessarily destroy the neurons.

Course of the Project: Studies are being presently continued by Dr. Lilly.

R.P.C. 2
Dec. 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB - 60 (C)
SERIAL NO.

11. BUDGET DATA

12. BUDGET ACTIVITY: RESEARCH

13. NONE

14. NONE

HONORS, AWARDS AND PUBLICATIONS

16. PUBLICATIONS OTHER THAN ABSTRACTS FROM THIS PROJECT DURING 1955:

17. HONORS AND AWARDS TO PERSONNEL RELATING TO THIS PROJECT DURING
CALENDAR YEAR 1955:





Project Description

1. Neurological Diseases and Blindness 2. Surgical Neurology
 INSTITUTE LABORATORY OR BRANCH
3. Clinical Psychology 4. _____ 5. NINDS-61 (C)
 SECTION LOCATION (OTHER THAN PETHESDA) SERIAL NO.
6. Functioning of the Temporal Lobe in the Recording of Experiences.
 PROJECT TITLE
7. Laurence L. Frost
 PRINCIPAL INVESTIGATOR
8. Maitland Baldwin, Mildred Blevins
 OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: To determine the role of the temporal lobe with its structures and central connections, in learning, recollection, recall, and reproduction.

METHODS EMPLOYED:

- a. Psychological tests and specific measures of learning administered to patients with temporal lobe seizures prior to and following removal of one or the other temporal lobe. These same tests administered to patients with bilateral temporal lobe damage and to patients with damage in cerebral areas other than the temporal lobe.
- b. Observation of and tests on monkeys and chimpanzees subjected to bilateral temporal lobe removal.
- c. Observation of imitative learning and general retentiveness in infant monkeys who have had both temporal lobes removed.

PATIENT MATERIAL:

		No.	Average Stay - days
Inpatient:	Adult Male	14	47
	Adult Female	10	46
	Child Male	2	49
	Child Female	3	45
Outpatient:	Number of patients:	10	
	Number of visits:	15	

MAJOR FINDINGS:

- a. Psychological tests indicate that memory, so commonly defined, is not altered or rendered non-functional by the presence or removal of a damaged temporal lobe. What seems more likely is that functions of attention and concentration are changed in such a manner that the patients attention fluctuates rapidly and that it is difficult for him to maintain a state of concentration. A test devised to measure the recording and comparing of experiences indicates that many patients with cerebral pathology, cortical or sub-cortical, have difficulty in so doing when compared with a "normal" group.

- b. Observation of monkeys and chimpanzees following unilateral and bilateral temporal removals reveals nothing in the way of decreased ability to record prior experiences. Rare "oral" tendencies as reported by Klüver and Bucy have been transient. Some alteration in the recognition of "textures" may be present.
- c. Infant monkeys have had their temporal lobes removed bilaterally. When reestablished in the general monkey colony, no differences were noted between them and their fellows. They learned and imitated readily. There was no outstanding "tameness."

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: Epileptic patients in general frequently complain of a memory loss, and temporal lobe patients in particular occasionally have a "memory" aura; as well, "memory" responses are occasionally obtained during stimulation at operation. This project hopes to be able to define memory and to learn the part played in the recording of experiences by the temporal lobe; as such it is related to the project: Mechanism of Memory, Branch of Surgical Neurology.

PROPOSED COURSE OF PROJECT: Continue the present line of research with greater emphasis placed on bitemporal lesions, and lesions of other than temporal areas.

Project Description

1. Neurological Diseases and Blindness 2. Surgical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clinical Psychology 4. LOCATION (OTHER THAN PETHESDA) 5. NINDB-62 (G)
SECTION SERIAL NO.
6. The Role of the Temporal Lobe in Perception.
PROJECT TITLE
7. Laurence L. Frost
PRINCIPAL INVESTIGATOR
8. Mildred Ebevins, Maitland Baldwin
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: An attempt to determine the role of the temporal lobe in the perception of visual, auditory, and tactual stimuli.

METHODS EMPLOYED:

- a. Standard psychological and neurological tests of perception administered to patients with temporal lobe electrographic foci. The same tests administered to patients with electrographic foci elsewhere in the cerebral hemispheres, or diffuse electrographic disturbances, and to patients and controls who have no abnormal cerebral findings.
- b. Two tests devised, one primarily for perception, one primarily for learning.
- c. Conditional discriminations with primates and electrical stimulation of the temporal lobe and its hypothecated connections.

PATIENT MATERIAL:

	No.	Average Stay - days	
In patient:	Adult Male	20	46
	Adult Female	22	45
	Child Male	8	48
	Child Female	9	45
Outpatient:	Number of patients	15	
	Number of visits	22	

MAJOR FINDINGS:

- a. Perception as measured by standard psychological and neurological tests indicates that cerebral injury, focal or diffuse, precludes accurate perception. The more gross the injury the more unclear the perception. It is becoming apparent that temporal lobe foci tend to alter in a subtle manner the perception of relationships to other individuals. This is different in crucial ways from perception and relationship problems faced by others with cerebral and sub-cortical injury.
- b. The special test devised for perception indicates that some

patients with focal cortical injury may show an initial facility for more fluid perception which becomes increasingly rigid in the patient is found to make more broad interpretations of his perceptions. This data does not seem true of those with diffuse cerebral damage or those with apparently normal cerebral function.

The test devised primarily for learning indicates that auditory perception of those with cerebral damage may be altered, particularly in the recognition of auditory stimuli.

c. Stimulation of an hypothecated connection between the primate temporal lobes (the anterior commissure) apparently grossly altered the monkey's perception of his environment for the duration of the stimulation. This same result has been seen on occasion following stimulation of the depths of one temporal lobe.

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: Perception and its interpretation plays a major role in personality. Since perception is seemingly altered in a subtle manner in patients with temporal lobe damage. The definition of this alteration is basic to the understanding of the temporal lobe. As well this project is allied to projects titled: Mechanism of Memory and Visual Field, Defects following Temporal Lobectomy. These latter two projects are directly from the Branch of Surgical Neurology.

PROPOSED COURSE OF PROJECT: Continuation of the present lines of inquiry, modified as further investigation dictates. It will be necessary to gather information on perception from a greater number of patients with focal electrographic changes elsewhere in the cerebral cortex and sub-cortical structures.

Project Description

1. Neurological Diseases and Blindness 2. Surgical Neurology
 INSTITUTE LABORATORY OR BRANCH
3. Clinical Psychology 4. _____ 5. NINDB-63 (C)
 SECTION LOCATION (OTHER THAN BETHESDA) SERIAL NO.
6. Speech Areas in Man
 PROJECT TITLE
7. Laurence L. Frost
 PRINCIPAL INVESTIGATOR
8. Maitland Baldwin
 OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: Localization of brain areas involved in the production and interpretation of speech. Determination of types of speech errors associated with dominant hemisphere lesions.

METHODS EMPLOYED: Speech testing of conscious patients while the exposed brain is being electrically stimulated. Speech testing of patients with dominant hemisphere lesions. Speech testing of patients during auras and in the post-ictal period.

PATIENT MATERIAL:

		No.	Average Stay - days
Inpatient:	Adult Male	1	48
	Adult Female	3	47
	Child Male	0	
	Child Female	1	48
Outpatient:	Number of patients	0	
	Number of visits	0	

MAJOR FINDINGS: There has been no thorough evaluation of data. Such evaluation as there has been indicates that of the expression speech difficulties, nominative speech is the most severely affected by discharging lesions of the dominant hemisphere. It is probable that receptive difficulties are present but the data are too few for adequate evaluation.

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: One of the programs of the Branch of Surgical Neurology is functional localization in the brain, to which this project specifically applies. This project is allied with one entitled: An Investigation of Speech Function Localized in the Temporal Areas of man of the Branch of Surgical Neurology.

PROPOSED COURSE OF PROJECT: Continuation of collection of data until sufficient is gathered to warrant conclusions.

Budget Data Sheet

10. NINDB-63 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH

13. NONE

14. NONE

Honors, Awards, and Publications

15. _____
SERIAL NO.

16. PUBLICATIONS OTHER THAN THIS PROJECT DURING 1955.

NONE

17. HONORS AND AWARDS TO PERSONNEL DURING 1955.

NONE

R.P.C. 1 (a)
December 1955

NINDB-64 (C)
SERIAL NO.

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: Personality defects are commonly seen, in those with abnormal temporal lobes (whether seizures result or not). This project is related to one entitled: Mechanism of Memory, Branch of Surgical Neurology.

Project Description

1. Neurological Diseases and Blindness 2. Surgical Neurology
 INSTITUTE LABORATORY OR BRANCH
3. Clinical Psychology 4. _____ 5. NINDB-65(C)
 SECTION LOCATION (OTHER THAN BETHESDA) SERIAL NO.
6. Intellectual Status with Drug Change in the Cerebral Palsied Child.
 PROJECT TITLE
7. Laurence L. Frost
 PRINCIPAL INVESTIGATOR
8. Mildred Elevins, J. Gordon Millichap
 OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: To determine changes in measured intelligence as various anti-convulsant drugs are used on children with cerebral palsy.

METHODS EMPLOYED: Standardized intelligence tests of the appropriate chronological level are administered to children with cerebral palsy following changing and reestablishing them on various anti-convulsant medications, principally DIAMOX.

PATIENT MATERIAL:

	No.	Average Stay - days
Inpatient:	Adult Male	0
	Adult Female	0
	Child Male	3 27
	Child Female	4 25
Outpatient:	Number of patients	14
	Number of visits	17

MAJOR FINDINGS: Project recently started and data too few for analysis.

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: To aid in evaluation of anti-convulsant medication particularly of cerebral palsied children to better aid in their schooling.

Budget Data Sheet

10. NINDB-65 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY: RESEARCH

13. NONE

14. NONE

Honors, Awards, and Publications

15. _____
SERIAL NO.

16. PUBLICATIONS OTHER THAN THIS PROJECT DURING 1955
NONE

17. HONORS AND AWARDS TO PERSONNEL DURING 1955
NONE

Project Description

1. Neurological Diseases and Blindness 2. Surgical Neurology
INSTITUTE LABORATORY OR BRANCH
3. Clinical Psychology 4. _____ 5. NINDB-66 (C)
SECTION LOCATION (OTHER THAN BETHSUDA) SERIAL NO.
6. Intellectual and Social Evaluation of Severely Mentally Retarded
Cerebral Palsied Individuals. _____
PROJECT TITLE
7. Laurence L. Frost
PRINCIPAL INVESTIGATOR
8. Anatole Dekaban
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

OBJECTIVES: To determine more accurate methods of evaluating the social adjustment and intellectual capacity of severely mentally retarded cerebral palsied individuals.

METHODS EMPLOYED: Standardized tests of intelligence and social maturity are administered. The tests analyzed to determine which parts of the tests provide the most valid and reliable estimate of intellectual and social functioning. Using these data further test items are elaborated in order to refine existing tests.

PATIENT MATERIAL:

		No.	Average Stay - days
Inpatient:	Adult Male	2	14
	Adult Female	0	
	Child Male	4	14
	Child Female	0	
Outpatient:	Number of patients	0	
	Number of visits	0	

MAJOR FINDINGS: The project has just been instituted and no findings are available.

SIGNIFICANCE TO THE PROGRAM OF THE INSTITUTE: To provide a more adequate measure of the intellectual and social abilities of severely retarded brain damaged individuals.

PROPOSED COURSE OF PROJECT: Continue the present line of research.

Budget Data Sheet

10. NINDE- 66 (C)
SERIAL NO.

11. _____
BUDGET DATA

12. _____
BUDGET ACTIVITY: RESEARCH

13. _____
NONE

14. _____
NONE

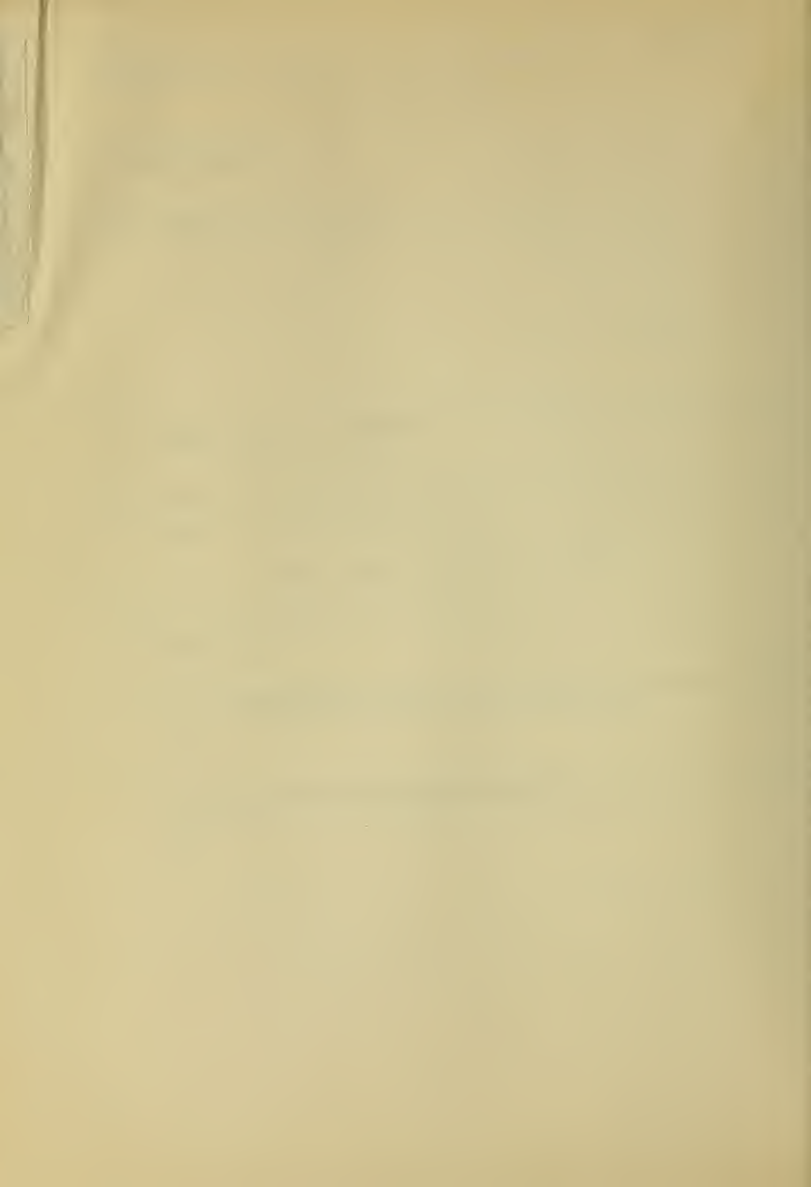
Honors, Awards, and Publications Sheet

15. _____
SERIAL NO.

16. _____
PUBLICATIONS OTHER THAN THIS PROJECT DURING 1955
NONE

17. _____
HONORS AND AWARDS TO PERSONNEL DURING 1955.
NONE





National Institute of Neurological
Diseases and Blindness
Surgical Neurology Branch
Section on Developmental Neurology

The Section on Developmental Neurology has been recently formed. The first three months were mainly utilized for the organization of clinical investigation relating to the congenital and early acquired disorders of the CNS and also for the organization of the research neuropathological laboratory. Our staff is not yet complete, and certain items of laboratory equipment must yet be delivered.

Two research projects have just begun; three others are in the preparatory stage as outlined below. Two of the latter projects are rather extensive, and because of the material necessary they call for the participation of other city hospitals. This is imposing special problems and unavoidable delays.

The outline of the five projects is enclosed.

Anatole Dekaban, M. D.

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R.P.C. - 1 (a)
Dec. 1955

NINDB-67 (C)
SERIAL NO.

PROJECT DESCRIPTION (Continued)

Proposed Course of the Project: This work has just begun. The accumulation of data of such material is necessary and must continue before analysis of findings can be attempted.

ANALYSIS OF NIH PROGRAM ACTIVITIES

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Developmental Neurology
SECTION
4. _____
LOCATION
5. NIMDB-68 (C)
SERIAL NO.
6. Preparation of the Horizons of the Normal Development of the Major Structures of the CNS in Mice.
7. Anatole Dekaban, M. D.
PRINCIPAL INVESTIGATOR
8. Marie Kendall, B. A.
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objectives: The main purpose of this work is to supply norms of the gross and microscopic structure of the CNS of mice during progressive stages of prenatal and postnatal development.

Methods Employed:

1. The CNS of mice at 16 progressive developmental stages will be dissected.
2. Representative sections will be prepared and stained.
3. Main structures will be identified and marked on low power microphotographs.

Mice Material:

Stock colony of white albino mice.

No. of Stages in Processing

4

Major Findings: This project has just begun. Brains of eight mice representing four horizons are dissected and are being further processed.

Significance of Neurological Research: An experimental approach to congenital malformations of the CNS is necessary to help understand certain obscure malformations occurring in humans. The provision of norms of the central nervous system has to precede the experimental production of congenital malformations, as there does not exist any proper guide in the form of an atlas or of a satisfactory reference during consecutive stages of the development.

R.P.C. - 1 (a)
Dec. 1955

NINDB-68 (C)
SERIAL NO.

PROJECT DESCRIPTION (Continued)

Proposed Course of the Project: This work has just begun. The processing of sections from the CNS of mice and the preparation of horizons must continue until completion of all stages.

ANALYSIS OF NIH PROGRAM ACTIVITIES

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Developmental Neurology
SECTION
4. _____
LOCATION
5. NINDB-69 (C)
SERIAL NO.
6. Experimental Production of Congenital Malformations of the CNS.
7. Anatole Dekaban, M. D.
PRINCIPAL INVESTIGATOR
8. Possibly
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objectives: A project is in preparation which relates to the production of congenital abnormalities of the CNS in mice using various noxious agents.

Methods Employed:

1. X-ray radiation of mice during different stages of pregnancy.
2. Utilization of certain enzyme inhibitors during different stages of pregnancy.
3. Gross, skeletal, and microscopic examination of the obtained specimen.

Material: White albino mice.

Findings: This project is in preparation.

Significance to Neurological Research: Morphogenetic and pathological analysis of congenital abnormalities of the CNS at different stages of development will be utilized as an adjunct in explaining similar abnormalities occurring in humans.

Proposed Course of the Project: Collection of material must precede analysis of findings.

ANALYSIS OF NIH PROGRAM ACTIVITIES

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Surgical Neurology
BRANCH
3. Developmental Neurology
SECTION
4. _____
LOCATION
5. NINDB-70 (C)
SERIAL NO.

Principal Investigator from NINDB - Anatole Dekaban, M. D.

PROJECT DESCRIPTION:

In a process of preparation is a joint project with participation of Walter Reed Army Hospital, the Naval Medical Center, and possibly Georgetown University Hospital. The purpose of this investigation is a progressive clinical study of infants who were exposed to undue stress during intrauterine life or during the process of birth. This project cannot be outlined in detail until formally approved by the heads of the respective institutions and until subsequent discussions with the chiefs of the departments concerned will be held.

ANALYSIS OF NIH PROGRAM ACTIVITIES

Project Description

- | | | |
|---|---|---|
| 1. <u>Neurological Diseases and Blindness</u>
<u>INSTITUTE</u> | 2. <u>Surgical Neurology</u>
<u>BRANCH</u> | |
| 3. <u>Developmental Neurology</u>
<u>SECTION</u> | 4. _____
<u>LOCATION</u> | 5. <u>NINDB-71 (C)</u>
<u>SERIAL NO.</u> |

Principal Investigator: Anatole Dekaban, M. D.

PROJECT DESCRIPTION:

In a process of preparation is a project relating to gross and microscopic examination of the CNS of human embryos, fetuses and stillbirths as well as infants dying early in life. The material will be obtained from the Walter Reed Army Hospital, the Naval Medical Center, and possibly Georgetown University Hospital. The detailed outline of this project has also to await its formal approval by the heads of these Institutions.

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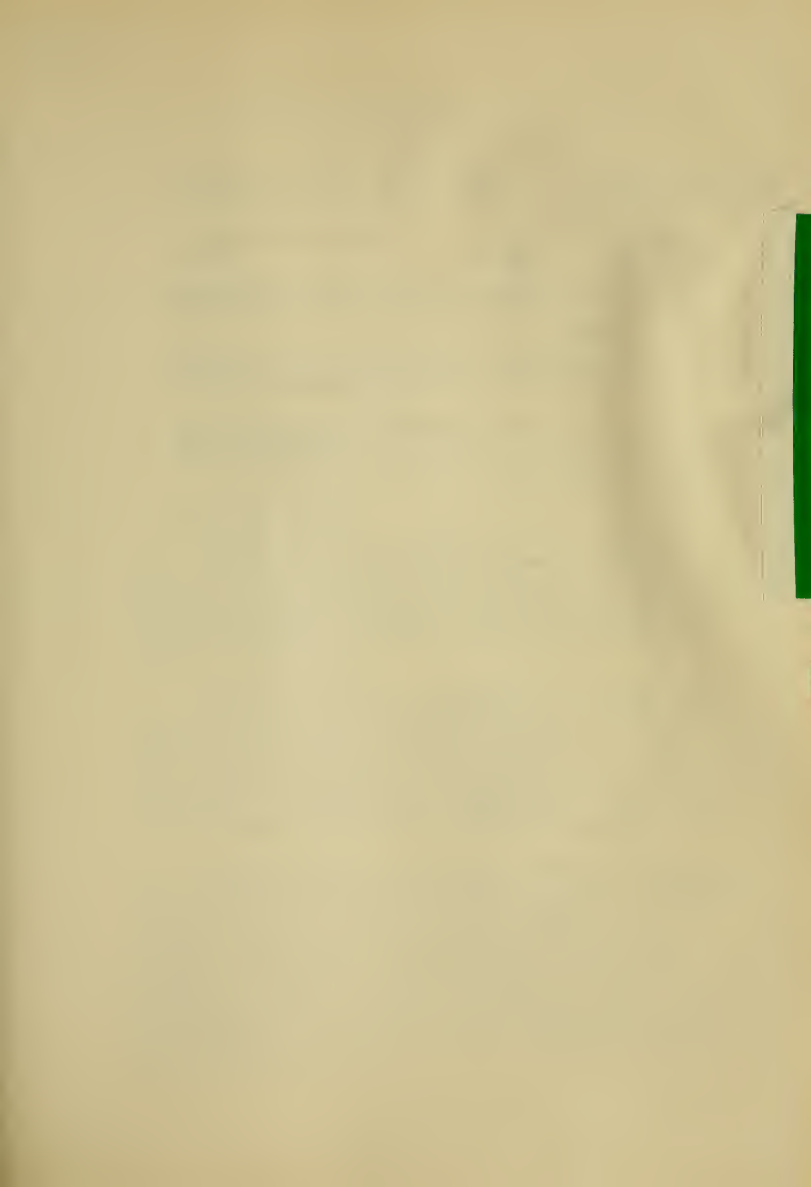
PHILOSOPHY DEPARTMENT

PHILOSOPHY 101

LECTURE NOTES

BY [Name]

These notes are based on the lectures given by [Name] in the Philosophy 101 course during the semester of [Year]. The notes are intended to provide a summary of the main points of the lectures and to serve as a study guide for students. The notes are not intended to be a substitute for the lectures themselves, and students are encouraged to attend the lectures and to read the assigned texts.



Analysis of NIH Program Activities

Project Description

- | | | |
|--|---|--|
| 1. <u>Neurological Diseases and Blindness</u>
INSTITUTE | 2. <u>Ophthalmology</u>
LABORATORY OR BRANCH | |
| 3. <u>Ophthalmology Physiology</u>
SECTION | 4. _____
LOCATION | 5. <u>NINDB - 72 (C)</u>
SERIAL NO. |
| 6. <u>Studies on Central Nervous System Control of Intraocular</u>
PROJECT TITLE | | |
| _____ Pressure | | |
| 7. <u>Ludwig von Sallmann, M.D. and J. Frank Macri, Ph.D.</u>
PRINCIPAL INVESTIGATOR(S) | | |
| 8. <u>Theodor Wanko, M.D., Hans Bornschein, M.D.</u>
OTHER INVESTIGATORS | | |
| _____ & Miss Patricia Grimes | | |
| 9. <u>PROJECT DESCRIPTION:</u> | | |

Objectives: Central nervous influence on intraocular pressure has been the subject of discussion for many years. On the basis of ill-defined clinical observations, it was postulated that the hypothalamus might be involved in disturbances of intraocular pressure regulation. Though this postulate was advanced only minor attempts have been made to establish the validity of this concept.

The present endeavour is to examine this question in a systematic manner under controlled conditions. Of the various sites in the brain which are known to influence autonomic regulation and possibly the intraocular pressure, the hypothalamus and adjacent structures were first examined for two reasons: (1) considerable information is available on this area on other functions, i.e., blood pressure, pupillary size, etc; (2) this area is presumed to be a center of autonomic integration.

In previous communications on this subject it was reported that stimulation of the ventral thalamus and dorsal hypothalamus resulted in isolated rises and falls of intraocular pressure, whereas stimulation of the ventral hypothalamus induced a sympathetic mass discharge with concomitant intraocular pressure changes. In this work no attempt was made to analyze the mechanism of intraocular pressure changes or to identify the receptors, effectors or pathways responsible for these responses. The object of the present study is a critical analysis of the above in addition to mapping of positive stimulation points in other parts of the brain.

Methods Employed: A modified Horsley-Clarke stereotaxic instrument, multichannel Sanborn recording units, a Leeds and Northrup sensitive temperature recorder, a dual beam oscilloscope for electromyography and scaler with a rate meter for recording blood volume changes in the anterior segment of the eye after injection of P-32 labeled red cells; have been assembled for the above mentioned experiment. The experimental animal will be the cat.

Major Findings: Results obtained to date by stimulation of various areas of the posterior hypothalamus and the medial ventral thalamus on intraocular pressure indicate the following: the rise of intraocular pressure is accompanied by a rise of subscleral temperature; the rise is not mediated through the cervical sympathetic nerve; it is not due to mechanical pressure exerted by the striated extraocular muscles; occlusion of the common carotid artery during continuous stimulation causes first a transient fall of intraocular pressure in the homolateral eye, followed by a sustained rise to the pre-occlusion level. Subscleral temperature changes in these cases follow a pattern similar to the intraocular pressure responses; the effect on the intraocular pressure could be abolished by Priscoline and Thorazine.

Significance: Studies of central nervous control of the intraocular pressure present a new phase of glaucoma research. It was brought forward in this year's Josiah Macy Foundation Conference on Glaucoma that experimental evidence for such regulation may have direct clinical implications.

<u>Patient Material:</u>	<u>Number</u>	<u>Average Stay</u>
		<u>Days</u>
<u>In-Patients:</u> Adult Males	1	9
Adult Females	1	57
Children: Males	0	
Children Females	0	
<u>Out-Patients:</u>	Number - 10	
	Number of visits - 31	

Significance: Eye pressure responses to autonomic drugs can be elicited by effects on inflow of the aqueous by contraction of the extraocular muscles and particularly by volume changes in the intraocular vasculature. In the present study the influence by neurovascular responses is emphasized. Results of this work should have implication on the problem of eye pressure pathology (glaucoma).

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Ophthalmology
LABORATORY OR BRANCH
3. Ophthalmology Pharmacology
SECTION
4. _____
LOCATION
5. NINDB-74 (C)
SERIAL NO.
6. Studies on Diet- and Drug-Induced Experimental Cataract
PROJECT TITLE
7. Ludwig von Sallmann, M.D.
PRINCIPAL INVESTIGATOR
8. J. Frank Macri, Ph.D. and Mr. Leo Caravaggio
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objectives: Information obtained by the cytological examination of whole mounts of the lens epithelium after exposure of eyes to ionizing radiation suggested similar studies on diet- and drug-induced lens damage. To find the role played by the lens epithelium in the development of galactose, myleran and tryptophane deficiency cataracts is the immediate objective of this study. Cytological examination of the lens epithelium in combination with histological studies on paraffin sections of such lenses will allow the investigators to determine the sequence of pathological events which ultimately lead to gross opacification of the lens. In particular it will be shown whether injury to the germinative zones of the lens epithelium or primary damage to the lens fibers causes the development of lens opacification.

Methods Employed: The technique of cytological and histological examinations of the animal lenses is the same as described in Project NINDB-_____ in the experiments on ionizing radiations. Experiments on rats and rabbits have been carried out utilizing this procedure.

Major Findings: In galactose lens injury a steep increase of cell division in the lens epithelium precedes the swelling and degenerative changes of superficial lens fibers near the lens equator. Both changes occur before lens opacities can be seen with the biomicroscope. The results of feeding experiments with myleran are incomplete. Tryptophane deficient rats will be available for histological examination in the near future.

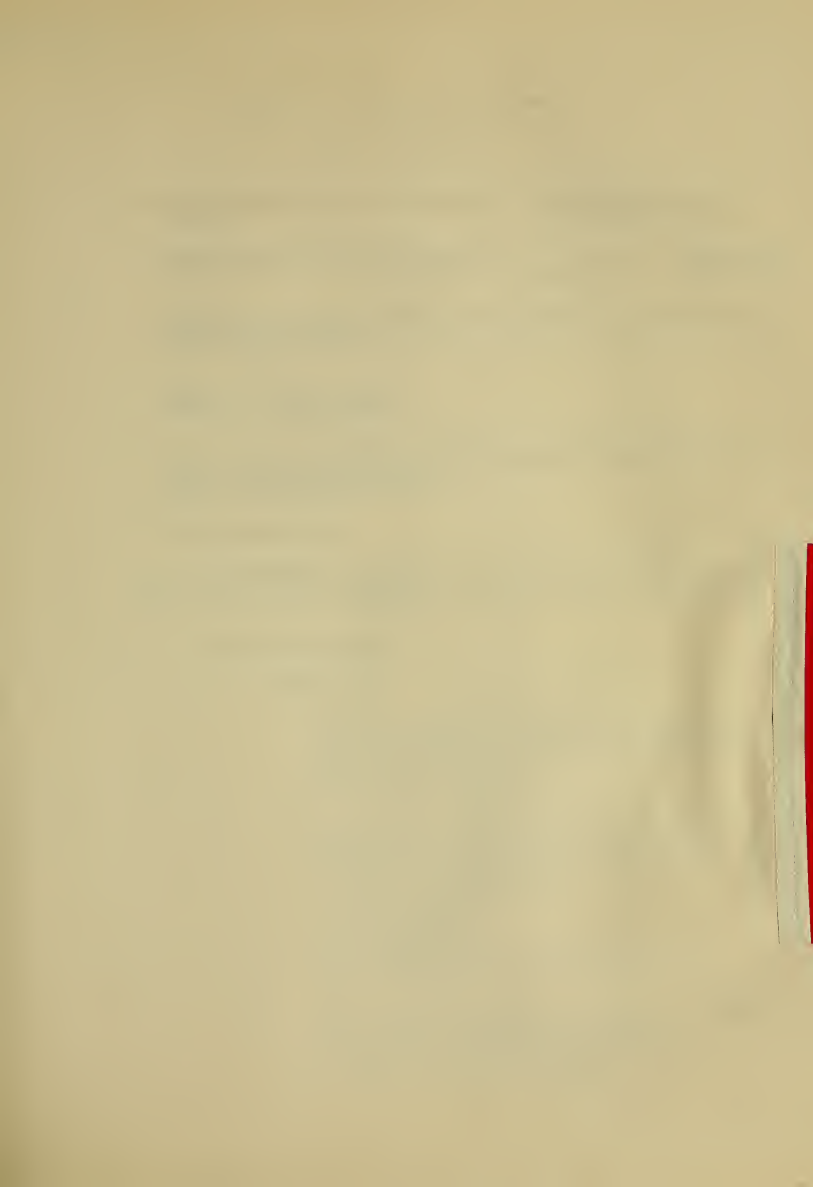
Significance: By systematic histological examination both in vivo, by biomicroscopy and particularly by cytological and histological examinations at various time intervals after introduction of the cataractogenic agents or deficiencies, it can be expected that similarities and differences in the pathogenesis of various types of experimental cataracts will be established which in the future might be correlated with chemical studies. Knowledge of the pathogenesis of various types of cataract might permit a rationale for therapeutic intervention.

The experiments with P-32 are conducted on litter mates (rabbits) with various time intervals allowed to elapse before sacrificing the animals.

Major Findings: The mitotic rate in the lens epithelium of the mouse per 100,000 cells is the highest (101), that of the rhesus monkey the lowest (2.9), among the species under study. The radio-sensitivity of these lenses appear to depend directly upon this function. Duration of post irradiation depression of mitosis does not seem to reflect the degree of radiation sensitivity; nor does the number of degenerative cells express necessarily the degree of radio vulnerability of the lens. In order to clarify the situation, it is necessary to study both epithelium and lens fibers in prolonged experiments. This part of the work is well under way but not completed.

The mitotic index of the lens epithelium varies greatly with age and parallels in limits the radio-sensitivity of various age groups. The degenerative changes of the cells in old age (rats) are strikingly similar to those induced by ionizing radiation. At the present time human lenses removed for cataract are being studied to compare the cytology of senile cataract with that of radiation cataract.

Significance: The described studies might lead to a better understanding not only of radiation induced lens damage but also to the pathogenesis of senile cataract.



Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness INSTITUTE
2. Ophthalmology Branch LABORATORY OR BRANCH
3. Ophthalmological Disorders Service SECTION
4. _____ LOCATION
5. NINDB-76 (C) SERIAL NO.
6. Evaluation of Newer Steroids in the Treatment of Experimental and Clinical Uveitis. PROJECT TITLE
7. James F. O'Rourke, M.D. PRINCIPAL INVESTIGATOR(S)
8. Gilbert Iser, M.D.; Monte G. Holland, M.D.; George Goodman, M.D. and G. Richard O'Connor, M.D. OTHER INVESTIGATOR(S)
9. PROJECT DESCRIPTION:

Objectives: A comparison of prednisone anti-inflammatory effects with those of cortisone and hydrocortisone in experimental and clinical uveitis therapy.

Methods Employed:

Basic Study.

1. Induction of horse serum uveitis in previously sensitized animals followed by the intramuscular injection of either prednisone, cortisone, hydrocortisone or ACTH.
2. An estimation of the uveitis response to each separate steroid treatment regimen based on grading of lesions according to the severity of:
 - (a) keratitic precipitates
 - (b) aqueous cellularity
 - (c) aqueous beam
 - (d) iris hyperemia.Control animals treated with intramuscular saline injection.
3. Series of lesions treated to demonstrate steroid effects both in blocking the onset of subsequently induced uveitis and in the treatment of established lesions.

Clinical Studies. Prednisone is employed in the therapy of anterior and posterior uveitis, plus optic and retrobulbar neuritis. Observations of the speed and adequacy of response following local and systemic therapy will be compared to the results of previous steroid therapy, in each individual case. The basic uveitis studies for etiologic diagnosis include x-ray, skin testing with indicated antigens and specific blood studies for hematologic and blood chemistry alterations.

<u>Patient Material:</u>	<u>Number</u>	<u>Average Stay</u> <u>Days</u>
In-Patients: Adult Males	6	56
Adult Females	4	27
Children Males	0	0
Children Females	0	0
Out-Patients:		
Number	18	
Number of Visits	246	

Major Findings:

1. Basic studies of animal uveitis response to prednisone therapy indicate a potency approximately five times that of other steroids. Ultimate clearing time of inflammation, however, is not greatly shortened, nor is the onset of anti-inflammatory effect noticeably hastened.

2. On clinical trial prednisone has not to date exceeded the anti-inflammatory effects of cortisone or hydrocortisone given in therapeutic doses. The dose required per patient, however, is much lower and the incidence of serious side effects was negligible.

3. The specificity of response noted in several patients to only certain steroids raises the possibility of a tissue selectivity.

4. Prednisone has been found effective when combined with specific anti-toxoplasmic chemotherapy in the treatment of granulomatous uveitis in several instances. Thus has an overlapping of projects between the toxoplasmosis study and the steroid evaluation project been effected.

Significance to Institute Program: Two factors of general clinical interests have emerged: (1) the inability of prednisone to exceed the response which follows other steroid therapy, and (2) the selectivity of response to various steroids among the patients studied. These factors may well apply to other areas of the Institute in which such treatment is employed.

Encouraging results noted on the combination of steroid therapy with specific chemotherapy has special reference to uveitis studies in Ophthalmology. As such they will motivate further comparative evaluation studies with currently accepted regimens of uveitis treatment.

Proposed Course of Project: Continued clinical evaluation of all steroids both alone and in combination with specific chemotherapy is contemplated in the treatment of granulomatous uveitis.

Analysis of NIH Program Activities

BUDGET DATA SHEET

10. NINDB-76 (C)
SERIAL NO.

11. BUDGET DATA

	ESTIMATED EXPENDITURES	BUDGETED POSITIONS			MAN YEARS			PATIENT DAYS
		PROF	OTHER	TOTAL	PROF	OTHER	TOTAL	
FY 1956								
FY 1957								

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
REVIEW & APPROVAL TECHNICIAN ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM 1)

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO.(S) IF WITHIN NIH)



2. Basic Studies. Tissue analyses of S-91 malignant melanomas of mice are carried out on acid digests of tumor and normal tissues, at various time intervals following the intravenous injection of radioactive isotopes. Calculations of the relative uptake and retention of the isotope by tumor tissue are based on the counts per minute obtained from suitably prepared dry samples of the tissue digests.

Simultaneous activation analysis studies of animal tumor tissue by neutron bombardment are underway at the Oak Ridge National Laboratory, Division of Analytical Chemistry, in an effort to determine which inorganic ions are present and operative in melanoma metabolism.

<u>Patient Material:</u>	<u>Number</u>	<u>Average Stay</u> <u>Days</u>
In-Patients: Adult Males	1	22
Adult Females	3	20
Children Males	1	1
Children Females	0	0
Out-Patients:		
Number -	3	
Number of Visits -	8	

Major Findings: The progress of this project thus far is to be reported in three categories: (1) equipment, (2) clinical studies, and (3) basic studies.

1. Equipment.

Geiger-Mueller Counters - Acquisition of two special probes designed for ocular counting makes possible the application of both external and surgical (retrobulbar) techniques.

Scintillation Counter - Parts for special collimated scintillation counter and gamma spectrometer system have been assembled with significant economy by collaboration of efforts with the Oak Ridge National Laboratory, Physics Division. The principles of focusing collimation and pulse height analysis operative in the system permit greater resolution and exclusion of scattered radiation in gamma ray counting than was previously possible.

Gamma source uptake and localization studies will be carried out on both animal and human subjects in an effort to evaluate the performance of this equipment for tumor localization.

2. Clinical Studies. Both external and retrobulbar ocular P-32 counting services are now available for suitable patients. To date many anatomical and physical limitations of beta source counting have become apparent: both primary and metastatic ocular malignancies have escaped detection on the basis of P-32 uptake in our studies. This has served to motivate a continued search for suitable gamma emitting sources. Two papers are in preparation dealing with such limitations of P-32 counting.

3. Basic Studies. Radioactive phosphorus uptake by normal animal eyes and by S-91 malignant melanoma include the following categories:

A. Normal eyes.

- (1) The percentage of administered P-32 present in the animal globe at one- and 24-hour intervals following intravenous injection (completed).
- (2) The role of extraocular muscle and fascial tissues on in vivo P-32 counting (completed).
- (3) Scleral absorption ratio of intraocular P-32 emission (completed).
- (4) The effect of intraocular and extra-ocular inflammation on P-32 uptake (in progress).
- (5) P-32 uptake of component ocular tissues (pending).

B. Animal S-91 Melanoma Studies.

- (1) The Differential Absorption Ratio of P-32 in malignant melanoma of mice (completed).
- (2) Activation analysis of S-91 melanoma, plus gamma and beta spectrometric studies for inorganic ion makeup of melanoma (in progress at Oak Ridge).
- (3) Uptake of gamma emitting isotopes by S-91 mouse melanoma (pending).

Significance to Institute Program: The construction and evaluation of newer tumor localizing equipment will yield results applicable to other NINDB studies dealing with malignant disease.

Clinical studies of non-malignant ocular disease by the use of isotopically tagged antibodies will in all probability overlap tumor localization procedures with regard to equipment required for counting of high resolution.

Proposed Course of Project: 1. Continuation of P-32 counting techniques applied to referred patients, pending the completion of basic studies on gamma emitting sources. 2. The evaluation of gamma uptake by animal tumors. 3. The improvement of gamma counting resolution.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
2. Ophthalmology
LABORATORY OR BRANCH
- Ophthalmology-
3. Physiology SECTION
4. _____ LOCATION
5. NINDB-78 (C)
- The Influence of Hypothermia on the Electrical Responses of the
6. Visual System.
PROJECT TITLE
7. Hans Bornschein, M.D.
PRINCIPAL INVESTIGATOR(S)
8. Gilbert Iser, M.D., George Goodman, M.D. and Ralph Gunkel, O.D.
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION:

Objectives: To determine the influences of hypothermia on the electroretinogram using a hibernating animal (hamster).

Further steps will be the comparison of the effects of hypothermia and anoxia on the electroretinogram and the electrical activities of the retinal ganglion cells and optic nerve.

Methods Employed: The electroretinogram in the anesthetized hamster will be picked up with a special contact lens electrode during stimulation with single light flashes and/or flicker light of various intensities, and in various states of adaptation. Measurements will be made with a twin beam oscilloscope and appropriate preamplifiers. The body temperature will be varied by circulating water through copper coils between which the animals are placed. The temperature of the head will be measured in the cheek-pouch. The effect of acute anoxia (ligation of aorta) will be studied at various temperature levels.

Significance to Institute Program: In the last few years the human electroretinogram has gained clinical importance, although the diagnostic value is still a matter of discussion. The electroretinogram is the result of various potentials of different signs, the origin and significance of which are controversial. In order to understand the clinical electroretinogram it is necessary to obtain more basic data along the lines of analysis of the components of the electroretinogram.

Proposed Course of Project: At first it is proposed to study only the effect of hypothermia on the electroretinogram (using the hamster), with variation of the stimulus conditions (intensity etc.) Further studies will include the recording of retinal ganglion cell activity and/or optic nerve impulses. This latter work will probably have to be done in larger animals such as the cat, with controlled circulation.

Analysis of NIH Program Activities

BUDGET DATA SHEET

10. NINDB-79 (C)
SERIAL NO.

11. BUDGET DATA

	ESTIMATED EXPENDITURES	BUDGETED POSITIONS			MAN YEARS			PATIENT DAYS
		PROF	OTHER	TOTAL	PROF	OTHER	TOTAL	
FY 1956								
FY 1957								

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
REVIEW & APPROVAL TECHNICIAN ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM 1)

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO.(S) IF WITHIN NIH)

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Ophthalmology
BRANCH
3. Physiology SECTION
4. _____
LOCATION
5. NINDB- 80 (C)
SERIAL NO.
6. Evaluation of Electroretinography as a Useful Clinical Tool.
PROJECT TITLE
7. Gilbert Iser, M.D., and George Goodman, M.D.
Principal Investigators
8. _____
Other Investigators

9. PROJECT DESCRIPTION:

Objectives: Electroretinography is the study of the electrical potentials elicited from the retina by a light stimulus. It affords the ophthalmologist an objective index of retinal function. Karpe (1945) suggested the use of a single flash light stimulus after several minutes of dark adaptation as the standard procedure in clinical studies, and this is the method which has been used by clinical investigators. However, there are many limitations to the use of the Karpe method. Large areas of central and peripheral retina may be damaged and yet the electroretinogram remains normal. In addition, as it is usually recorded in the dark-adapted eye, the electroretinogram is dominated by rod potentials and poorly reflects cone function.

The success obtained with subjective flicker fusion frequency tests in revealing minimal retinal lesions suggests the study of the objective flicker fusion frequency by electroretinographic means. Furthermore, recent experiments with the use of a high intensity flickering stimulus have shown that the resulting photopic flicker electroretinogram is primarily a response of the retinal cones. This indicates that the high intensity flicker electroretinogram may be affected abnormally by macular disease and diseases of the cone system where the single flash electroretinogram is not.

The present study was undertaken to determine whether the flicker electroretinogram would provide more diagnostic information than the standard single flash electroretinogram in ocular disease and other choreoretinal affections.

Methods Employed: Patients are tested while resting in bed. The Grass Photoc Stimulator is used as a flashing light source, and the patients are tested with stimuli of varying intensity, frequency, and wavelength. The potentials are picked up simultaneously from both eyes with contact lens electrodes, and the binocular recordings are made on a Grass eight-channel direct writing electroretinographic machine.

<u>Patient Material:</u>	<u>Number</u>	<u>Average Stay Days</u>
In-Patients: Adult Males	0	0
Adult Females	1	39
Children Males	0	0
Children Females	1	38
Out-Patients: Number	12	
Number of Visits	43	

Patients suitable for study are drawn from those referred to the Ophthalmology Branch for investigation. Patients with normal eye findings are used as controls and those with retinal detachment, hereditary, inflammatory and vascular retinopathies are studied for diagnostic changes. Approximately 75 patients were studied electroretinographically during the past year.

Major Findings: During the past nine months the equipment was assembled and tested. The electroretinographic tracings on normal controls are comparable to those obtained by other investigators.

Patients with retinal disease were tested with scotopic and photopic single flash techniques. A depressed electroretinogram was found in cases of severe myopia and diabetic retinopathy, extensive posterior uveitis, retinal detachment, and marked color deficiency. The degree of electroretinographic abnormality appears to be dependent upon the area of retina involved and the type and degree of structural or functional damage which has occurred.

Preliminary investigations with flicker electroretinography were carried out using high intensity white light stimuli. The amplitude of the flicker waves and their fusion frequency are decreased in cases with diffuse retinal disease. Further studies with the flicker electroretinogram and the use of different stimulating intensities and wavelengths are under way.

Significance to Institute Program: Ophthalmologists have long sought more accurate methods of determining retinal function, especially macular integrity, in the presence of opacities of the ocular media. The answer to this search may lie in the development of new techniques of electroretinography.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness 2. Ophthalmology Branch
INSTITUTE LABORATORY OR BRANCH
3. Ophthalmology Physiology 4. _____ 5. NINDB-84 (C)
SECTION LOCATION SERIAL NO.
6. Improvement in the Design and Construction of Ophthalmic
Instruments.
PROJECT TITLE
7. Ralph D. Gunkel, B.S., O.D.
PRINCIPAL INVESTIGATOR(S)
8. _____
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

Objectives: To make improvements in the instruments used in ophthalmic examination or treatment.

Specifically: To make final improvements in the magnetically controlled self-recording tangent screen device mentioned in the last report.

To adapt a non-varying, non-flickering light source to the Bausch and Lomb retinal camera for greater uniformity in pictures than is now possible with the carbon arc light.

To develop a clinically useful flicker fusion field testing device and technique.

To develop other devices, instruments, or techniques as needs and situations arise.

Major Findings: The tangent screen device has probably been improved to its ultimate state of refinement and could be produced commercially in its present form. The marking device formerly causing some difficulty has been simplified to a pneumatic plunger and bulb which does not fail to function. Since a uniform illumination of seven foot-candles is desired for standardizing central visual fields, it was considered necessary to design a suitable light source. This was done, using a 16-inch circular fluorescent tube with a variable shutter arrangement. This also provides for more sensitive tests by reducing the illumination to less than seven foot-candles if desired.

A 100 watt zirconium arc lamp has been adapted and installed in place of the carbon arc lamp on the Bausch and Lomb retinal camera. It has been found to give usable Kodachrome pictures with one-half and one-fifth second exposures. It is currently being tested with Anscochrome film, which it is believed will give better pictures with shorter exposure time.

Instruments have been assembled and some work has been done toward developing a new flicker-field technique. A Sylvania glow modulator tube with its square wave generator, lens, and filter system appear to elicit more definite responses from the subject than were formerly possible.

A transparent plastic electrode holder has been constructed for electroretinography and found to be satisfactory. It provides positive contact, visible electrodes, and eliminates the need for using tape as was formerly done.

Numerous requests have been received and met for the construction of plastic shields, pads, and lens holders individually designed for special cancer patients.

Significance to Institute Program: It is believed that the tangent screen device developed here provides the most convenient, rapid and reliable means yet available for the measurement of central visual fields. Its use is now accepted as standard procedure in our Eye Clinic.

While it is not possible to strictly evaluate devices or ideas contributed to electroretinography and other projects, it is believed that they are considered definitely helpful.

Proposed Course of Project: A project of this type must have sufficient flexibility to permit collaboration with new workers and in new projects. Present plans call for continuation of this project along lines similar to those of the past year, with modifications to meet needs and situations as they arise.

(a) The agar-diffusion method for the detection of anti-Toxoplasma precipitins in the eye: In addition to the presence of neutralizing antibodies (Dye Test Antibodies) and complement-fixing antibodies to Toxoplasma, it is thought that there are also precipitins for Toxoplasma in immune sera. The purpose of the agar-diffusion method is to detect precipitin reactions under conditions where small volumes of antibody-containing fluids, such as aqueous humor, are used. It is thought that in infectious diseases of the uveal tract, antibody formation takes place in the eye in excess of that which reaches the interior of the eye through permeation of the blood-aqueous barrier. Precipitins present in the aqueous humor should be detectable when samples of aqueous are allowed to diffuse through a semi-solid clear agar gel toward an opposing body of antigen molecules. Opaque lines of precipitate should be seen at the interface between the diffusing antigen and antibody whereas little as 10 micrograms per ml. antibody nitrogen is present.

(b) The detection of labelled Toxoplasma antibody in the chorioretinal lesions of living eyes: Anti-Toxoplasma rabbit serum with a dye test titer of 1:250,000 has been prepared. This method involves the labelling of such antibody with fluorescein, an azo-dye, or radioactive iodine. Antibody so labelled and injected parenterally can be expected to localize in the eye lesion at the site of antigen liberation. Here it should be detected by the use of ultraviolet light, direct ophthalmoscopy, or a Geiger counter respectively.

<u>Patient Material:</u>	<u>Number</u>	<u>Average Stay</u> <u>Days</u>
In-Patients: Adult Males	5	17
Adult Females	11	27
Children Males	1	31
Children Females	4	24
Out-Patients: Number	42	
Number of Visits	292	

Granulomatous uveitis cases are obtained by physician referral. Screening for the presence of high Toxoplasma dye test titers or toxoplasmin skin reactions can be done before the selection of the patient. Those patients with evidence of anterior involvement of the eye are of particular interest for aqueous study.

Major Findings: The presence of precipitins in Toxoplasma antisera of known high dye test titer was determined using the standard tube dilution method. The reaction seems to be pH sensitive, occurring most readily at pH 6.5. It does not occur in the presence of active complement. No discrete precipitin

lines have been observed in agar as yet, but the buffer system and electrolyte content may have to be altered to obtain optimal reactions.

Rabbit gamma globulin preparations of high purity have been made through the use of an electro-convection apparatus. There is evidence that antibody activity is preserved through this particular method of preparation.

Supra-choroidal injections in the rabbit's eye have been made using the technique of Vogel. It is planned to establish posterior pole infections in the rabbit's eye preparatory to the use of labelled antibody.

Significance to Institute Program: There is good evidence that at least in the earlier or acute phases of ocular toxoplasmosis, treatment with Pyrimethamine, Sulfa drugs, and the anti-inflammatory agents such as ACTH or Cortisone has resulted in the halting of a progressive uveitis and the restoration of useful vision. If a more specific diagnosis could be made at the onset, much valuable time might be saved for the patient if he is treated with anti-Toxoplasma drugs, as opposed for example, to six months of anti-tuberculous therapy or one year of tuberculin desensitization. Many cases of ocular toxoplasmosis have undoubtedly gone undiagnosed because of the lack of even the more rudimentary serological aids to diagnosis.

The agar-diffusion technique also offers a good possibility for the diagnosis of other types of uveitis using a series of many different antigens. In this regard it might be a truly fundamental gain in the otherwise highly dubious current etiologic diagnoses of uveitis.

Proposed Course of Project: To proceed with the labelling of the purified rabbit gamma globulin, first with fluorescein and then with the other afore-mentioned agents. To test the predilection of labelled antibody for the artificially induced Toxoplasma lesion in the rabbit's iris. To test its predilection for posterior pole lesions in the rabbit and then in man.

We will continue to observe and treat patients with presumed toxoplasmic uveitis, with special emphasis on the information to be gained from paracentesis and examination of the aqueous humor, response to drug therapy, and eventually for detection of toxoplasmic lesions in situ.

Present Status: Active cases are now available for investigation. Purified rabbit anti-Toxoplasma globulin has been prepared. Precipitins to Toxoplasma have been demonstrated in serum by the serial tube dilution method, but not as yet by the agar-diffusion method.

Analysis of NIH Program Activities

BUDGET DATA SHEET

10. NINDB- 81 (C)
SERIAL NO.

11. BUDGET DATA

ESTIMATED EXPENDITURES	BUDGETED POSITIONS		MAN YEARS			PATIENT DAYS
	PROF	OTHER	PROF	OTHER	TOTAL	
FY 1956						
FY 1957						

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
 REVIEW AND APPROVAL TECHNICIAN ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S). (ITEM 1)

Doctor Leon Jacobs of the National Microbiological Institute, Laboratory of Tropical Diseases, is carrying on animal, tissue culture, and epidemiological studies on Toxoplasma gondii in cooperation with our branch. He furnishes facilities and supervision for the toxoplasma dye test, complement fixation test, and for the production of human skin-test antigen from the peritoneal exudate of infected mice. He supplies us with high titer rabbit anti-serum. He performs the standard dye test on gamma globulin fractions which are to be tested for antibody activity. He handles the inoculation of animals with suspected infectious samples of aqueous humor and other tissues. Miss M. K. Cook, who is in the employ of the Ophthalmology Branch of NINDB, is directly associated with Doctor Jacobs in his laboratory. In addition to assisting with the dye test and routine animal work, she is pursuing experiments on the tissue culture of Toxoplasma in the rabbit retina.

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES, OR FUNDS) IDENTIFY SUCH RESEARCH: (BY SERIAL NO(S) IF WITHIN NIH.)

This complements animal experimentation carried on by Doctor Don Eyles at the Memphis Public Health Service facility.

3. Reproduction of experimental trachoma in monkeys or apes.

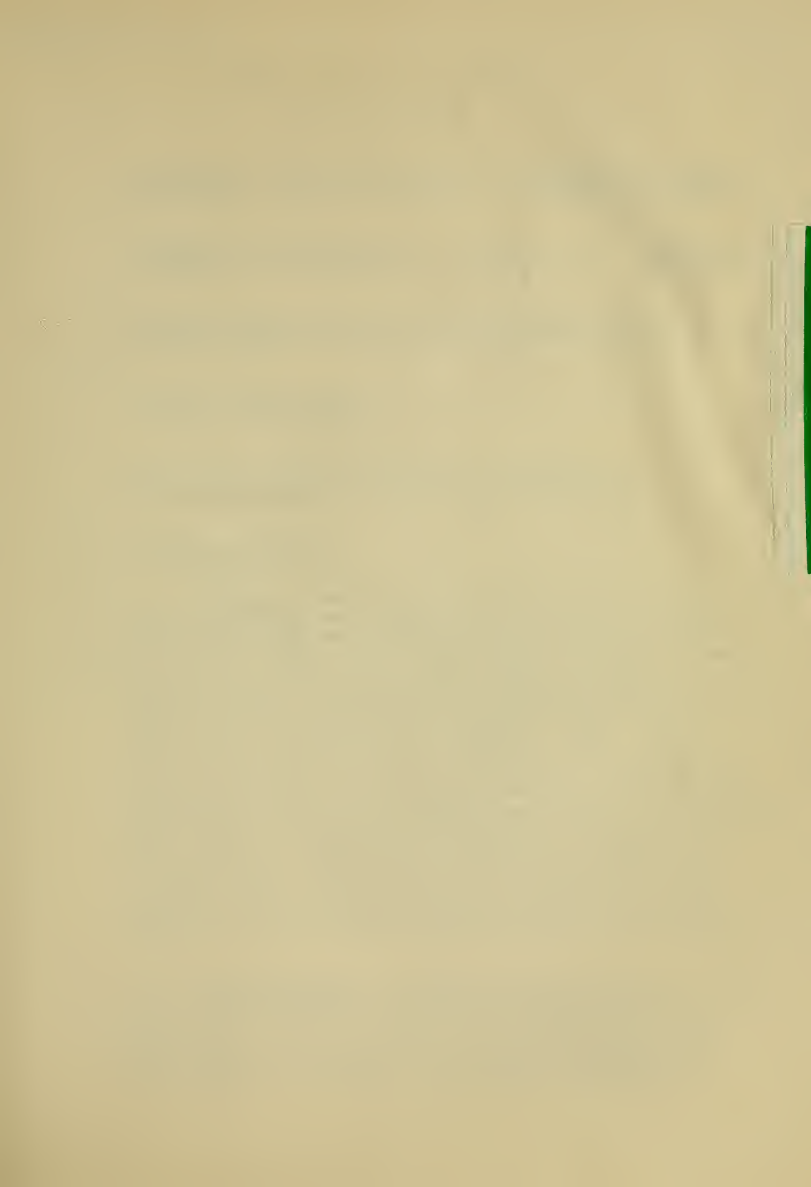
In attempting to culture viruses from cases of uveitis, aqueous humor samples will be inoculated into tissue cultures of epithelial cells, and cytopathic changes will be observed.

Patient Material: Material from active trachoma with numerous inclusions and active cases of epidemic keratoconjunctivitis are preserved by dry freezing and transported by air from Doctor Yukihiko Mitsui, Kumamoto, Japan.

Major Findings: The cytopathogenic effects of A.P.C. virus in the conjunctival epithelial cells resemble those described for He La cells. Maintenance medium for the cultivation of the A.P.C. virus in the conjunctival epithelial cell culture was found to be a mixture of 10% horse serum and 90% Eagles He La growth media.

Significance to Institute Program: Trachoma is a universal problem. Temporary control can be obtained with antibiotics and improvements in sanitary conditions. Cultivation of the trachoma virus becomes an urgent demand and will aid in ultimate control of the disease. The trachoma committee of W.H.O. is interested in promoting studies in this area.

The significance of virus culture studies on uveitis might be very great since a definite etiological diagnosis can rarely be made in present day studies of uveitis. If virus cultures were positive, much of the guess-work in current diagnosis might be eliminated by a laboratory test.



methylene blue staining will selectively demonstrate the nerves and permit a close study of the anatomy and the morphology of the nerve endings in the trabeculum and outlet channels.

The studies are to be carried out on human eyes when they become available. It is contemplated that monkeys, cats, rabbits, dogs and birds will be the experimental animals.

Major Findings: As the study is at present in a preliminary stage, it is not possible to state any conclusive findings. However, eyes of several cats, rabbits and monkeys have been studied, using the methods outlined and it has been possible to trace nerves running into the trabeculum, which apparently terminate there. To fully delineate the morphology of the delicate nerve endings will not be possible until the technique of staining and impregnation has been perfected.

Significance to Institute Program: If the innervation of the components of the trabeculum can be established on morphological grounds it will offer a new piece of evidence indicating how nervous impulses may be mediated in regulating the intraocular pressure. It would be of fundamental importance in particular with reference to the neurovascular mechanisms involved in regulating the intraocular pressure and also have significance in reference to the glaucoma problem. Evidence has steadily accumulated indicating that the trabeculum, Schlemm's canal and collector channels are extremely important in the regulation of the normal intraocular pressure and also the probable site of resistance to aqueous outflow in glaucoma. Through the systematic study of the innervation of these structures, it is hoped that a morphological basis will be found to complement the understanding of the basic physiology of the outflow mechanism.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Ophthalmology
LABORATORY OR BRANCH
3. Ophth. Chemistry 4. _____ 5. NINDB-85 (C)
SECTION LOCATION SERIAL NO.
6. A Study of the Proteins of the Lens
PROJECT TITLE
7. Robert A. Resnik
PRINCIPAL INVESTIGATOR
8. Frank G. Suggs and Ann Wolff
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION:

Many investigators have studied the proteins of the lens. These studies have not resulted in a clearly understood representation of what these proteins are chemically, and in fact different preparations are reported by different workers. Some reports have gone so far as to suggest that there are more than the four lens proteins which have been characterized by Kormer in his classical studies. No detailed studies of the chemistry of these proteins have been published and only recently has there been an attempt at determining the molecular weight of one of these proteins.

Objective: The fractionation of the lens proteins into homogeneous components is the ultimate goal. Any one fraction that is obtained will be used for the determination of molecular weight and in dye binding studies. These data will be used to characterize the normal proteins.

Methods employed: Fractionation will be attempted by the use of a cellulose base ion exchange resin. The ultracentrifuge and electrophoresis will be used to determine homogeneity. Characterization of these proteins by comparison with other proteins will be carried out by the technic of equilibrium dialysis. These methods will be supplemented by any that will be particularly useful as the work progresses.

Major Findings: Preliminary attempts at fractionation with ammonium sulfate did not provide any homogeneous fractions as evidenced by a number of peaks in the sedimentation pattern. The ion exchange studies have not been started.

Significance to Institute Program: While much data are available on the size, shape, and electrical and chemical properties of other proteins, the data for lens proteins are rather skimpy and incomplete. Through studies, somewhat similar to those planned here, a better understanding of the roles of proteins has resulted in many instances. It seems desirable to carry out studies of this fundamental nature on lens proteins for this reason.

Proposed Course of Project: The ion exchange resin to be used and the necessary equipment is now available. The initial efforts will be aimed at separating, on a micro scale, proteins from rat lenses. The empirical procedure that is to be developed will be used on a larger scale for obtaining significant (or macro) quantities of one or more proteins. The comparison of normal and pathological proteins in addition to studying species differences will be included as a part of this project.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Ophthalmology
LABORATORY OR BRANCH
3. Opth.Chemistry
SECTION
4. _____
LOCATION
5. NINDB-86 (C)
SERIAL NO.
6. The Chemistry of Alloxan
PROJECT TITLE
7. Robert A. Resnik
PRINCIPAL INVESTIGATOR
8. Ann Wolff
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION:

The lens of the eye contains proteins. It is reasonable to assume that clouding of this structure is a result of changes in these proteins. It is known that the optical properties of proteins may be altered in the process of denaturation. This often manifests itself by a change in the availability of free sulfhydryl groups of the protein. In instances of cataract formation the free sulfhydryl groups, present in protein and glutathione, are diminished in number. Such changes occur in experimental cataracts induced by alloxan. Although it is true in all probability that the ocular changes that result in diabetes mellitus and in alloxan diabetes are due to the diabetes itself, since the alloxan is known to oxidize sulfhydryl compounds, it is possible that, as a result, it could indirectly deplete the lens of this substance. Two suggestions have been proposed in an attempt to explain the toxic manifestations of alloxan: 1. Alloxan is toxic because it complexes with essential metals. 2. Alloxan is toxic because it reacts preferentially with the sulfhydryl groups of glutathione and protein.

Objectives: To study the reaction of alloxan with metals and sulfhydryl groups to determine whether these reactions might possibly be interpreted as having any relation to the toxicity of alloxan.

Methods Employed: One means of identifying substances is by comparison of their absorption spectra. The shape of the curve may be used as a qualitative means of identification, while the intensity of the spectrum may be used as a quantitative determination of the substance. In addition, any shift in the wavelength of the absorption maximum is indicative of a reaction taking place. In the case of reactions involving metals, complex formation is said to take place if a ligand (the complexing agent) reacts with a metal. These changes are also detectable by changes in the pH titration curve of the ligand when titrated in the presence and absence of metal.

Major Findings: Part 1. It has been proven that alloxan does not complex with metals as evidenced by experiments with copper. Alloxanic acid can and does complex with metals. This ligand is formed from alloxan by a base catalyzed rearrangement. Titrimetric and spectrophotometric evidence for this has been obtained.

Part 2. The reaction product between alloxan and glutathione has not been identified, but evidence has been obtained to obviate the possibility that the toxicity of alloxan can be attributed to the formation of a product with an absorption maximum at 305 mu. Substances with identical absorption spectra (or with a maximum at 305 mu) as the alloxan-glutathione adduct and which are not themselves known to be toxic have been studied. Thus violuric acid and the reaction product between dimethyl alloxan and glutathione absorb at 305 mu, but neither violuric acid nor dimethylalloxan are toxic.

The 305 chromophore has been proposed to contain the grouping, $-N=C-C=O$. A structure which is theoretically possible by analogy with the reaction of alloxan with free α -diamines has been proposed.

Significance to Institute Program: In the past, speculation without evidence has been made as to what the mechanism of the reaction of alloxan with physiological systems might be. This has been used by various groups in an attempt to explain the toxicity of alloxan. It has been established in this work that the toxicity of alloxan cannot be so explained. That is, it does not complex with metals and compound 305 may be obtained with the non-toxic, dimethylalloxan, and glutathione.

Proposed Course: Part 1. This is essentially complete. There are some aspects that are yet to be worked on but these involve some basic studies which will be deferred for the present. The nature of these future studies will be an analysis of the titration curves and an investigation of the effect of copper on the base catalyzed rearrangement of alloxan to alloxanic acid.

Part 2. This has not been completed. Studies involving reaction of N-substituted peptides with alloxan are contemplated.

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Ophthalmology
LABORATORY OR BRANCH
3. Ophth.Chemistry 4. _____
SECTION LOCATION
5. NINDB-87 (C)
SERIAL NO.
6. The Effect of Sex Hormones on the Formation of Cataracts Induced by Galactose.
PROJECT TITLE
7. Robert A. Resnik
PRINCIPAL INVESTIGATOR
8. _____
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION:

It has been well established that diets rich in galactose when fed to experimental animals result in the formation of cataracts. In addition, cataract formation occurs in infants and young children with idiopathic galactosemia. This condition, i.e. the formation of cataract, is peculiar to galactose of the hexoses. In the laboratory the susceptibility of rats to the effects of galactose seems to be confined to immature animals. Older animals appear to be refractory to the effects of galactose. In view of these observations it would appear desirable to determine the effect of castration on the metabolism of galactose.

Objectives: To determine whether the development of galactose induced cataracts is related to the sexual maturation of rats.

Methods Employed: Immature rats of both sexes are allowed to develop to maturity after castration is performed in the 4th to 6th week of age. The animals are then fed a 25% galactose diet and observed for the formation of cataracts. A group of controls of both sexes will be run in the same experiment.

Major Findings: This project was activated recently. No results are available as yet.

Significance to Institute Program: While it is known that diets high in galactose are able to produce cataracts in rats the reason for this effect is not known. In addition, why it is confined to the immature animals is not clearly understood. Information that might indicate some relationship between sexual maturation and the effect of galactose on the lens would be of interest both from the academic and practical points of view.

Proposed Course of Project: Until the results of the initial experiments are known the experiment as outlined in the section on methods will be used as such. It is contemplated to use very old animals in this experiment if possible.

9. PROJECT DESCRIPTION (Continued)

Method employed: An unusual opportunity to investigate the problem has been afforded by the extremely high prevalence of ALS among the Chamorros (natives) of Guam and the other Mariana Islands in the Western Pacific. Unusual or suspicious findings would be investigated more thoroughly at the Clinical Center.

a. Clinical description of patients on Guam and at the Mayo Clinic, with an analysis of the principal clinical features, is still under way. An investigation was carried on at the Mayo Clinic to determine if any variation in gastric cytology by chymotrypsin lavage could be detected.

b. Clinical Pathology: The clinical pathology program described last year has been temporarily discontinued because of lack of personnel available for this study on Guam. Through this program various blood and spinal fluid biochemical studies and genetic typing were to be conducted.

c. Pathological Anatomy: Review of series of ALS cases at the Armed Forces Institute of Pathology under the supervision of Dr. Webb Haymaker.

Material collected in Guam and from Mayo Clinic patients is also being examined by Dr. Sayre. Particular interest in the United States group is centered on those patients with a positive family history of motor neurone disease.

d. Epidemiology and Genetics: In California, a comparison of the incidence of ALS of Chamorros now living in that state and their families still in Guam is being carried out.

Comparisons of familial incidence among patients and a random sample "control" group on Guam are still being determined. In these latter groups a comparison of possible environmental etiologic factors is being investigated.

Studies of nutritional status are being conducted by Mrs. Whiting among Chamorros here in this country as well as on Guam and other Mariana Islands.

A search was made of records in the Archivo de Indias in Seville and the Government Archives in Madrid (Spain) and in the Vatican Library in Rome (Italy) to determine if ALS was present in the Mariana Islands prior to the arrival of the Spanish or was introduced after the Spanish arrived in the islands. This information would help determine how important it was to search for a possible exogenous factor.

9. PROJECT DESCRIPTION (Continued)

e. Therapeutic Studies: A number of possible agents are being tested among the patients on Guam and have been tested on a series of patients observed at the Mayo Clinic.

Major findings: Amyotrophic lateral sclerosis appears to have become highly prevalent in the Mariana Islands after the arrival of the Spanish explorers about 1670. Amyotrophic lateral sclerosis, on Guam, which is clinically and pathologically identical with the disease as we know it elsewhere, does not appear to be affected by pregnancy and presumably does not affect the course of pregnancy. No unusual blood groupings, biochemical observations, or therapeutic results can be reported for the population of Guam. The gastric cytology study at the Mayo Clinic was unsuccessful and will be repeated. Electromyographic results in the Guam area are similar to those observed among ALS patients elsewhere.

Although the study is incomplete, there are findings which suggest that the prevalence of ALS is high among Chamorros residing in California.

A review of the reported findings of Ask-Upmark of Uppsala, Sweden, in "Amyotrophic Lateral Sclerosis Observed in Five Persons After Gastric Resection" was made and it is my impression that Ask-Upmark's description of a correlation of ALS and gastrectomy was not quite so. After careful examination of the case records in his laboratory, there was some question regarding the diagnosis in several patients. Since gastrectomy had been performed on over 1,000 patients in the series he studied, the few cases in which ALS developed could be a chance happening. This would tend to de-emphasize the importance of gastrectomy or gastric malformation in the etiology of ALS.

Significance to neurological research: In addition to providing clinical data for an important disease of the nervous system, the study illustrates the value of combined field epidemiologic and clinical approach.

Clarification of the etiology of the disorder would be expected to improve the opportunity for therapeutic success.

Proposed course of the project: No change. Several of the above projects are still under way. The study in California will be continued in the summer of 1956.

R.P.C. 2
Dec. 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB- 88 (C)
SERIAL NO.

11. BUDGET DATA

12. BUDGET ACTIVITY: Research

13. U. S. Navy, Bureau of Medicine and Surgery, U. S. Naval Hospital
on Guam

Government of Guam: Department of Medical Services

Mayo Clinic

14. None

R.P.C. 2
Dec. 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB - 89 (C)
SERIAL NO.

11.

12. BUDGET ACTIVITY: Research

13. Mayo Clinic

National Office of Vital Statistics, U. S. P. H. S.

Office of Vital Statistics, Canada

14. None

Honors, Awards, and Publications

16. PUBLICATIONS FROM THIS PROJECT DURING 1955: None

17. HONORS AND AWARDS TO PERSONNEL DURING 1955: None



BRANCH OF ELECTROENCEPHALOGRAPHY

The activity of the Branch has substantially increased during 1955.

EEG examinations for routine diagnosis and for research show a 60% increase over those performed in the previous year.

A total of 1276 examinations were carried out on patients referred to the Branch from our Institute and from other Institutes as follows:

N.I.N.D.B.	1065
N. C. I.	123
N. H. I.	30
N.I.A.M.D.	29
N. M. I.	27
N.I.M.H.	<u>2</u>
Total	1276

Of the above total, 232 examinations were performed on out patients.

Electrocorticographic studies during surgical exposure of the cerebral cortex were performed on 38 epileptic patients.

The major research project in the clinical field remains the study of the physiopathology of the epilepsies with particular emphasis on the group of focal lesions within the temporal lobe. This project was described in detail in the last yearly report. It was stated that this is a long range project and the necessity for collecting a large number of patients was emphasized, together with the need for a careful and long follow-up of operated subjects before drawing any definite conclusion.

The second project is a corollary of the preceding one. It deals with the diagnosis and classification of the various types of epileptics by means of activation procedures. It has been found that the slow injection of Metrazol, carried up to the point of eliciting a clinical seizure, yields a wealth of electrographic and clinical information which, when properly interpreted, proves to be extremely useful in clarifying the diagnosis. Interesting observations on the electrographic correlates of numerous clinical signs could also be made during this study.

A third project is still under way also in the field of the epilepsies (anti-convulsant therapy). It concerns the EEG evaluation of possible effects of new types of medicaments, and it is carried out in collaboration with the Section of Neurochemistry.

The question of seizures in cases of cerebral palsy and the possibility of their surgical treatment is the object of another project. The outline of the electrographic criteria for the classification of these rather complex clinical cases is the chief aim of our study.

A new project has been recently started in collaboration with the N.I.M.H. and the Institute of Living (Hartford, Conn.). It concerns the study of old age changes in the brain and in particular, the relationship between the cerebral blood flow and electroencephalographic changes.

The experimental research in animals deals with the thalamo-cortical relationship and with the problem of spread of epileptic discharges. The subject of one project is the study of the "recruiting response" in cortical and subcortical structures, while another project deals with the mechanism of "spindles" which characterize the electrical activity of the brain in certain experimental conditions.

In the field of experimentally induced epileptogenic lesions two projects are under way. The purpose of these studies is to gain further information on the different electrographic behavior in cases where the lesion is produced at a subcortical level on one hand, and in those in which the lesion is located in various cortical regions, on the other.

Preliminary results of three of the above-described projects were presented in the course of this year at different Scientific meetings.

During the past year the staff of the Branch of Electroencephalography has been increased and includes at the present time three EEG technicians and one technician in experimental neurology. In addition, a research associate has joined our Branch during the last six months, collaborating very actively on several of the projects.

Methods employed and patient material: This main project is, therefore, subdivided into the following sub-sections:

- A) EEG diagnosis of temporal lobe epilepsy.
- B) Electrographic study of temporal lobe discharges.
- C) Electrographic-clinical correlations of epileptic seizures.
- D) Sleep and temporal lobe epilepsy.
- E) Electrographic manifestations of subcortical epileptogenic lesion.
- F) Clinical-pathological correlations in cases with parasagittal and temporal EEG abnormalities.
- G) Cortical physiology in man (electrocorticographic study of spontaneous, evoked, and abnormal activity with particular emphasis on the temporal region).

Routine EEG examinations are carried out on all epileptic patients admitted by the Neurosurgical Branch. Numerous tracings are obtained in various conditions (after withdrawal of medication, during induced sleep, during spontaneous or induced seizures) using a minimum of 21 electrodes and various methods of localization. If the patient is eventually selected for surgical treatment, the electrical activity will be recorded from its exposed cortex both at rest and following electrical stimulation of various cortical and subcortical structures, before and after surgical ablation of the abnormal tissue. EEGs are then taken at regular intervals in the post-operative period.

Major findings: This is a long range project in which positive and useful results can only be obtained with a large number of cases and with pathological and clinical correlations. The clinical evaluation of surgical treatment of epilepsy requires a minimum of one to three years follow-up and most of the conclusions in regard to electroencephalographic criteria for physiopathology and diagnosis will obviously depend upon the surgical results. This limitation applies particularly to the sub-sections A, D, E, F, and partly to sub-sections B and C. Interesting observations were however made (sub-sections B and G) on the electrical activity of the exposed insula where epileptiform discharges can often be recorded. The behavior of paroxysmal abnormalities following cortical undercutting, and their relation with electrically induced afterdischarge was also investigated. The effect of arousal, acoustic stimulation and speech upon the spontaneous activity of various regions of the temporal cortex was studied.

Significance to the program of the Institute: The main program of the Neurosurgical Branch of the NINDB concerns the temporal lobe epilepsy. This particular research project of our Branch is a primary adjunct to the research of the Neurosurgical Branch.

Proposed course of project: Fundamentally unchanged.

R.P.C. - 2
December 1955

Analysis of NIH Program Activities

Budget Data

10. NINDB-90 (C)
SERIAL NO.

12. BUDGET ACTIVITY: Research

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM I)

Branch of Neurological Surgery - NINDB

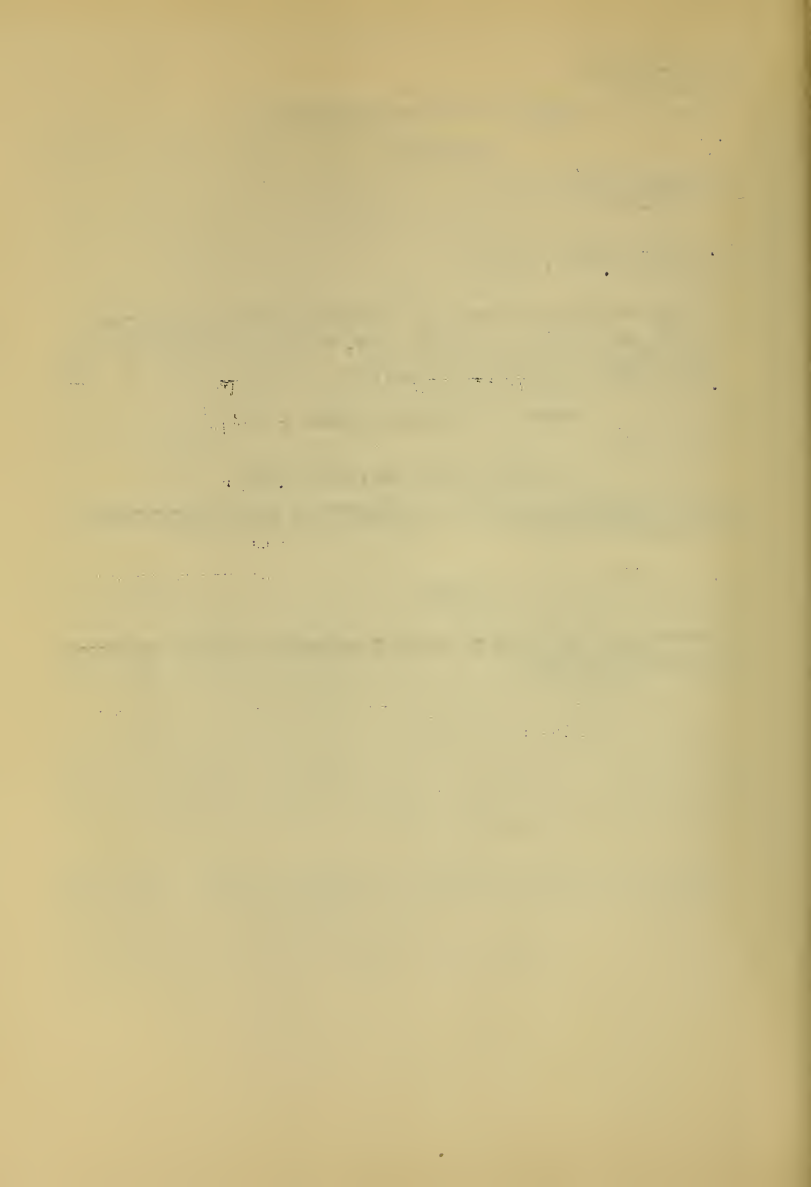
Honors, Awards, and Publications

16. LIST PUBLICATIONS OTHER THAN ABSTRACTS FROM THIS PROJECT DURING CALENDAR YEAR 1955:

None

17. LIST HONORS AND AWARDS TO PERSONNEL RELATING TO THIS PROJECT DURING CALENDAR YEAR 1955:

None



Major findings: The following information was obtained by the study of Metrazol-induced seizures (MIS).

- a) The MIS faithfully reproduces the patient's attack pattern (while on medication) in 85% of the cases and in over 90% the spontaneous attacks observed while off medication and at the time of the Metrazol activation. In only 4% was the MIS wholly unlike the habitual patient's attack.
- b) The average amount of Metrazol for a MIS is 6.55 mgs/Kg \pm 3.26. There is no correlation between threshold value and age, sex, or frequency of attacks. There is a slight correlation between threshold and duration of seizures in years and between threshold and type of epilepsy. The chances for the MIS to be a major generalized convulsion and/or an aspecific attack are not necessarily increased by increasing the dose of Metrazol.
- c) The value and reliability of various clinical lateralizing signs occurring in a MIS could be established. Aphasia and arm and leg movements are the most while eyes turning the less reliable signs.
- d) Motor lateralizing signs are present in an unexpectedly high number of temporal cases while visceral or non-visceral automatisms can be observed occasionally in cases with a parasagittal lesion. Aphasia predominates in bitemporal lesions. Consciousness may be completely unimpaired in a high percentage of seizures originating from the parasagittal region. The incidence of major generalized attacks markedly increases from unitemporal, to bitemporal to parasagittal to centrencephalic epilepsy.
- e) Cases of temporal lobe epilepsy.
 - i) If the MIS consists of automatism alone, the EEG is strikingly characterized by lack of changes or by unitemporal activation.
 - ii) If there is automatism plus aphasia the EEG changes are usually bilateral temporal.
 - iii) If there is automatism plus local motor phenomena, the EEG changes are focal or bilateral temporal (if the same clinical pattern takes place with no EEG changes, the lesion could be in the parasagittal area).
 - iv) If there is automatism plus local motor sign and a major seizure the EEG pattern may vary but there is a preponderance of bitemporal changes.
 - v) Automatism plus major seizures always are accompanied by EEG changes which can be focal temporal or diffuse.

- f) Other similar correlations were made for MIS in non-temporal epileptogenic lesions.
- g) The various data obtained during MIS in well-established clinical cases could be satisfactorily applied to a number of cases in which the clinical-EEG diagnosis and classification was uncertain or not established, and valuable information was obtained in a high percentage of cases.
- h) Comparison of the results obtained by the study of a MIS and by the study of only the early paroxysmal changes elicitable with Metrazol indicates the superiority of the former which yields a substantially higher number of useful results.

Significance to the program of the Institute: See project 1.

Proposed course of project: Collect a larger number of activations within the next year. Compare the effect of various activations in the same patient and study the modifications of the seizure pattern and of the EEG before and after surgical treatment.

A more accurate study of the electrical activity of various cortical regions will eventually be achieved by using a 16 channel electroencephalograph.

Analysis of NIH Program Activities

Budget Data Sheet

10. NINDB-91 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
REVIEW & APPROVAL TECHNICAL ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM I)

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO(S) IF WITHIN NIH)

Analysis of NIH Program Activities

Budget Data Sheet

10. NINDB-92 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY:

RESEARCH ADMINISTRATION
REVIEW & APPROVAL TECHNICAL ASSISTANCE

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM I)

NIMH and Institute of Living, Hartford, Conn.

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO(S) IF WITHIN NIH)

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness
INSTITUTE
2. Electroencephalography
LABORATORY OR BRANCH
3. _____
4. LOCATION (IF OTHER THAN BETHESDA)
5. NINDB-93 (C)
SERIAL NO.
6. Electrographic study in cases of cerebral palsy with seizures
PROJECT TITLE
7. C. Ajmone Marsan
PRINCIPAL INVESTIGATOR
8. M. Baldwin, A. Dekaban, G. Millichap
OTHER INVESTIGATORS

9. PROJECT DESCRIPTION

Objectives: To establish diagnostic and prognostic criteria in regard to the outcome of seizures in case of massive cerebral lesions when medical treatment has failed and the patient is a possible candidate for surgery.

Methods employed and patient material: Serial EEG examinations are carried out on patients admitted by the branch of Neurological surgery. The electrographic study is performed in different conditions and with various activating procedures. Particular emphasis is placed on localizing devices and in differentiating cortical from sub-cortical lesions. In cases of surgical operation, the exposure of an entire hemisphere provides unparalleled opportunities for electrical exploration of various cortical areas and of a number of subcortical structures.

Major findings: A limited number of patients have been studied in this relatively recent project. A long clinical and electrographic follow-up are needed in order to assess the validity of any offered interpretation.

Significance to the program of the Institute: This project is an important complement to similar researches for the coming year in the section of Neuropathology and in the Branch of Neurological Surgery of the NINDB.

Proposed course of the project: Accumulate wider information through a larger number of patients with a longer post-operative follow-up. At least 2-3 years project period.

Analysis of NIH Program Activities

Budget Data

10. NINDE-93(C)
SERIAL NO.

12. BUDGET ACTIVITY: Research

Honors, Awards, and Publications

16. LIST PUBLICATIONS OTHER THAN ABSTRACTS FROM THIS PROJECT DURING
CALENDAR YEAR 1955:

Problems in the EEG Analysis of Epileptic Activity in Patients
with Massive Cerebral Lesions. - C. Ajmone Marsan and M. Baldwin.
EPILEPSIA - in press

17. LIST HONORS AND AWARDS TO PERSONNEL RELATING TO THIS PROJECT DURING
CALENDAR YEAR 1955:

None

R.P.C. - 1
December 1955

Analysis of NIH Program Activities

Project Description

1. Neurological Diseases and Blindness
INSTITUTE
2. Electroencephalography
LABORATORY OR BRANCH
3. SECTION
4. LOCATION (IF OTHER THAN BETHESDA)
5. NINDB-94(C)
SERIAL NO.
6. Electroencephalographic control of anticonvulsant therapy.
PROJECT TITLE
7. C. Ajmone Marsan and D. Tower
PRINCIPAL INVESTIGATORS
8. C. E. Wells
OTHER INVESTIGATORS
9. PROJECT DESCRIPTION

Objectives and patient material: These are extensively described in project NINDB-12(C). The EEG picture is emphasized in this part of our project. Each patient receives a series of EEG examinations before the beginning of the treatment in order to evaluate the type and amount of paroxysmal abnormalities. Following the beginning of medication, the patient is re-examined approximately once a week for various months.

Major findings: The results of this study are still to be evaluated. The preliminary impression is that the clinical improvement is not paralleled by a similar electrographic improvement, and on the whole, most of the patients who seem to do well clinically still show an abnormal tracing where the paroxysmal discharges are still present with characters similar to those observed in the pre-medication records. It must be noted, however, that - at least electrographically - the selection of patients for this study was made on the basis of very abnormal records.

Significance to the program of the Institute: The possibility of determining with objective criteria the effect of an anticonvulsant treatment presents obvious advantages. The persistence of an abnormal tracing in cases of definite clinical improvement makes the prognosis more guarded than in cases of full normalization of the EEG accompanying the clinical improvement. Possibly only certain

R.P.C. - 1 (a)
December 1955

NINDB-94(C).
SERIAL NO.

types of EEG abnormalities are susceptible to improvement and consequently only certain types of cerebral seizures can definitely benefit from the treatment under study, even though some transitory diminution in the number of seizures may be observed in a relatively high percentage of cases.

Proposed course of the project: To elaborate the various EEG findings, from a quantitative and, possibly, qualitative, standpoint and to correlate them with the clinical ones of project NINDB 12(C).

R.P.C. - 2
December 1955

Analysis of NIH Program Activities

Budget Data

12. BUDGET ACTIVITY: Research

Honors, Awards and Publications

16. LIST PUBLICATIONS OTHER THAN ABSTRACTS FROM THIS PROJECT DURING
CALENDAR YEAR 1955:

None

17. LIST HONORS AND AWARDS TO PERSONNEL RELATING TO THIS PROJECT DURING
CALENDAR YEAR 1955:

None

(those with a larger number of synapses) can be supposed to have a more restricted field of distribution but to be very effective within this limited field. Shorter paths may excite a larger number of cells but in order to do this they require considerable temporal summation. The "recruiting" would, therefore, depend upon a progressive activation of two main groups of elements differing from each other by the number of synapses involved and the "waning" phase could be due to dropping out of the neurons excited by the shortest paths.

When recording the non-specific response from a surface-cortical electrode and from a needle electrode placed at different levels within the cortex itself, the underlying white matter, radiations, various basal ganglia and eventually the thalamus, not only the latency tends to remain the same, but recruiting and waxing and waning do occur in structures where obviously no cortical neurons are involved, and also after these have been depressed (cortical cocainization) or eliminated (cortical ablation). It logically follows that all of these events must originate subcortically and probably within the thalamus itself.

The transformation of polarity undergone by the response in the depth, its relatively unchanged amplitude and some of the mentioned observations can be explained by postulating the presence of two interrelated but separated electrical fields, one originating from fibers (positive deflection) the second from cell activity, a situation somewhat similar to that visualized by Grey Walter (1953). Thus, the transition from a high amplitude negative, to a lower amplitude negative-positive, to a high amplitude positive deflection throughout the cortex and underlying white matter can be interpreted as the result of the algebraic sum of two waves of opposite polarity with predominance of either one above or below a certain level; this level would not signify the passage from "sink" to "source" but the borderline between the two fields.

Significance to the program of the Institute: The system under investigation seems to be involved in certain types of epileptic disorders. A deeper knowledge of its physiology could probably throw some light on the complex mechanisms underlying psychoparetic manifestations, alterations of awareness and attention commonly found in epileptics.

Proposed course of project: Continue the investigation of this system with particular reference to a) inter-relationship with specific system, b) effect of CNS stimulants.

Analysis of NIH Program Activities

Project Description Sheet

1. Neurological Diseases and Blindness 2. Electroencephalograph
INSTITUTE LABORATORY OR BRANCH

3. SECTION 4. LOCATION (IF OTHER THAN BETHESDA) 5. NINDB-96 (C)
SERIAL NO.

6. Spindle mechanism and experimental epileptic lesions.
PROJECT TITLE

7. B. Ralston
PRINCIPAL INVESTIGATOR

8. OTHER INVESTIGATORS

9. PROJECT DESCRIPTION

Objectives: To study the possible role of the spindling mechanism in the transmission of focal epileptic discharges.

Methods employed: Creation of acute epileptic foci by the use of Penicillin in various cortical and subcortical regions and the changing of the transmission from these foci to other areas by the effect on the spindle formation. Experimental animals - cat.

Major findings: Foci produced in certain areas such as the mesial thalamus and the cingulate gyrus have an activating effect on the spindle formation and also synchronize these discharges over wide cortical areas. There is also an alteration in the frequency of the spindling to a pathological degree.

Significance of the Program to the Institute: It is possible that the discharges of so-called centrencephalic and subcortical epilepsy are the result of focal lesions "broadcasting" via this mechanism.

Proposed course of project: To continue and extend these observations on the cat and to make additional ones on the monkey, and to study the effects of various anesthetic and other pharmacological agents on the spindle formations.

Analysis of NIH Program Activities

Budget Data Sheet

10. NINDB-96 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY:

RESEARCH	<input checked="" type="checkbox"/>	ADMINISTRATION	<input type="checkbox"/>
REVIEW & APPROVAL	<input type="checkbox"/>	TECHNICAL ASSISTANCE	<input type="checkbox"/>

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM I)

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO(S) IF WITHIN NIH)

R.P.C. - 1(a)
December 1955

NINDB-97 (C)
SERIAL NO.

Significance to the program of the Institute: One of the main clinical projects is the EEG study of focal (temporal in particular) epilepsy. The possibility of establishing criteria for selecting cases with cortical from those with subcortical lesions would be of the greatest significance and practical use.

Proposed course of project: Continue the study and extend it to monkey with particular emphasis upon the cingulate, supplementary motor and temporal areas.

Analysis of NIH Program Activities

Budget Data Sheet

10. NINDB-98 (C)
SERIAL NO.

11. BUDGET DATA:

12. BUDGET ACTIVITY:

RESEARCH /x/ ADMINISTRATION / /
REVIEW & APPROVAL / / TECHNICAL ASSISTANCE / /

13. IDENTIFY ANY COOPERATING UNITS OF THE PUBLIC HEALTH SERVICE, OR OTHER ORGANIZATIONS, PROVIDING FUNDS, FACILITIES, OR PERSONNEL FOR THIS PROJECT IN EITHER 1956 or 1957: IF COOPERATING UNIT IS WITHIN NIH INDICATE SERIAL NO(S) (ITEM I)

14. IF THIS PROJECT RESEMBLES, COMPLEMENTS, OR PARALLELS RESEARCH DONE ELSEWHERE IN THE PUBLIC HEALTH SERVICE (WITHOUT INTERCHANGE OF PERSONNEL, FACILITIES OR FUNDS), IDENTIFY SUCH RESEARCH: (BY SERIAL NO(S) IF WITHIN NIH)



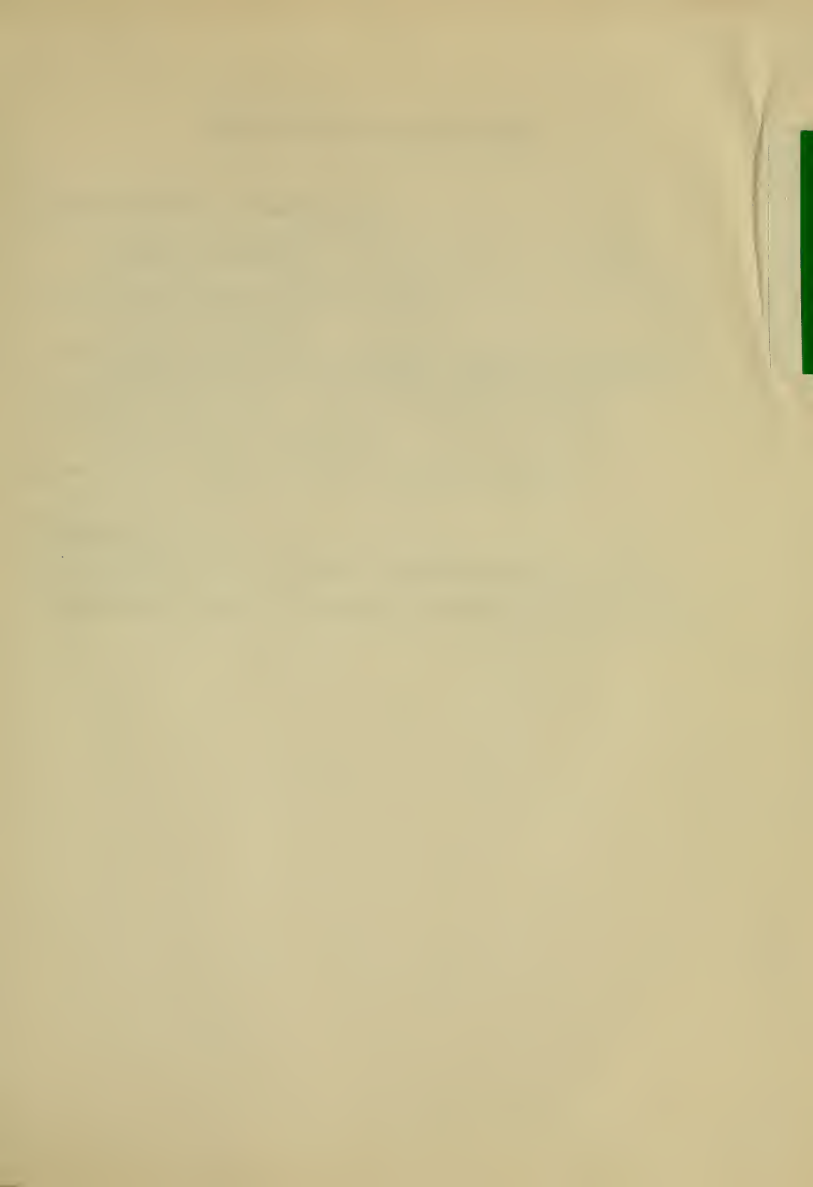
$$\begin{array}{r} 467 \\ 266 \\ \hline 201 \end{array}$$

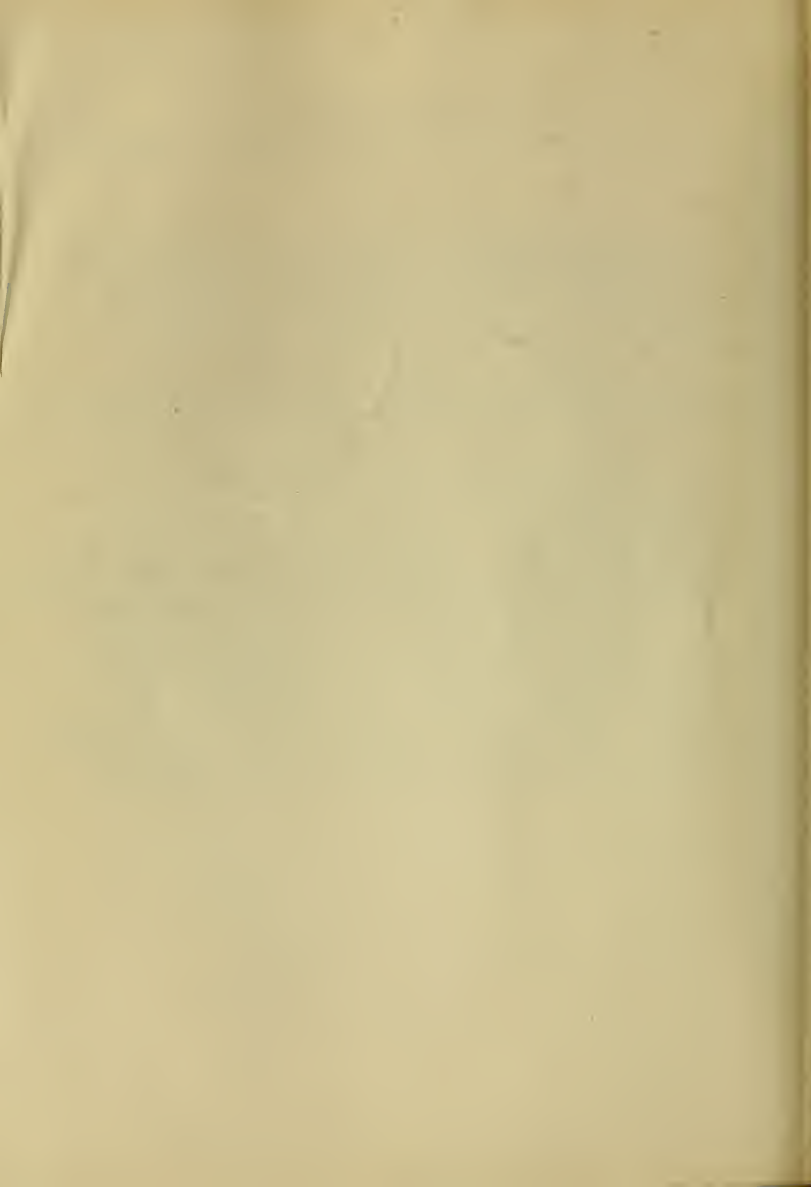
Services Given By The
Clinical Investigative Unit of the
National Institute of Neurological Diseases and Blindness

Although the primary function of each Institute of the National Institutes of Health is that of the research projects contained in this report, nevertheless, each Institute must give of its skills to aid other Institutes and the Clinical center to understand further the mechanism of diseases of the patients under their care.

Approximately 386 beds are activated in the Clinical Center at the present time, and this Institute has rendered 247 neurological consultations, 59 neurosurgical consultations, and 1,103 ophthalmological consultations.

The Electroencephalographic Laboratory of this Institute has continued to service all Institutes of the National Institutes of Health and ran 1,276 electroencephalographic records and 38 electrocorticographic records during the past year. Fifteen major surgical operations have been performed for Institutes other than NINDB. Personnel in our various laboratories, such as Neuropathology, work in close cooperation with the general pathological laboratories of the Clinical Center and during the year 1955, 617 cases were received in the Laboratory of Clinical Neuropathology. The Neurochemistry Laboratory of this Institute has also continued to do the spinal fluid proteins for the Institute. Such services tend to keep the clinical investigative unit more broadly orientated than is possible within the narrow confines of the research program. The unit, therefore, is pleased to be able to carry out these functions in addition to its research activities.





AWARDS, RECOGNITIONS, AND ACTIVITIES

Staff appointments to Universities:

- Doctor G. Milton Shy, Associate Clinical Professor of Neurology,
Georgetown University
- Doctor Maitland Baldwin, Associate Clinical Professor of Neurosurgery,
Georgetown University
- Doctor Donald B. Tower, Associate Clinical Professor of Neurology and
Associate Professor of Biochemistry, Georgetown University
- Doctor Cosimo Ajmone-Marsan, Instructor in Clinical Neurology,
George Washington University
- Doctor Kenneth R. Magee, Instructor, Georgetown University

Hospital Consultants:

- Doctor G. Milton Shy, U. S. Medical Naval Center, Bethesda, Maryland
- Doctor Maitland Baldwin, U. S. Medical Naval Center, Bethesda, Maryland

MEMORANDUM FOR THE RECORD

1. Subject: [Illegible]

2. Reference: [Illegible]

3. Summary: [Illegible]

4. Recommendation: [Illegible]

5. Remarks: [Illegible]

6. Disposition: [Illegible]

Approved: [Illegible]

7. Date: [Illegible]

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