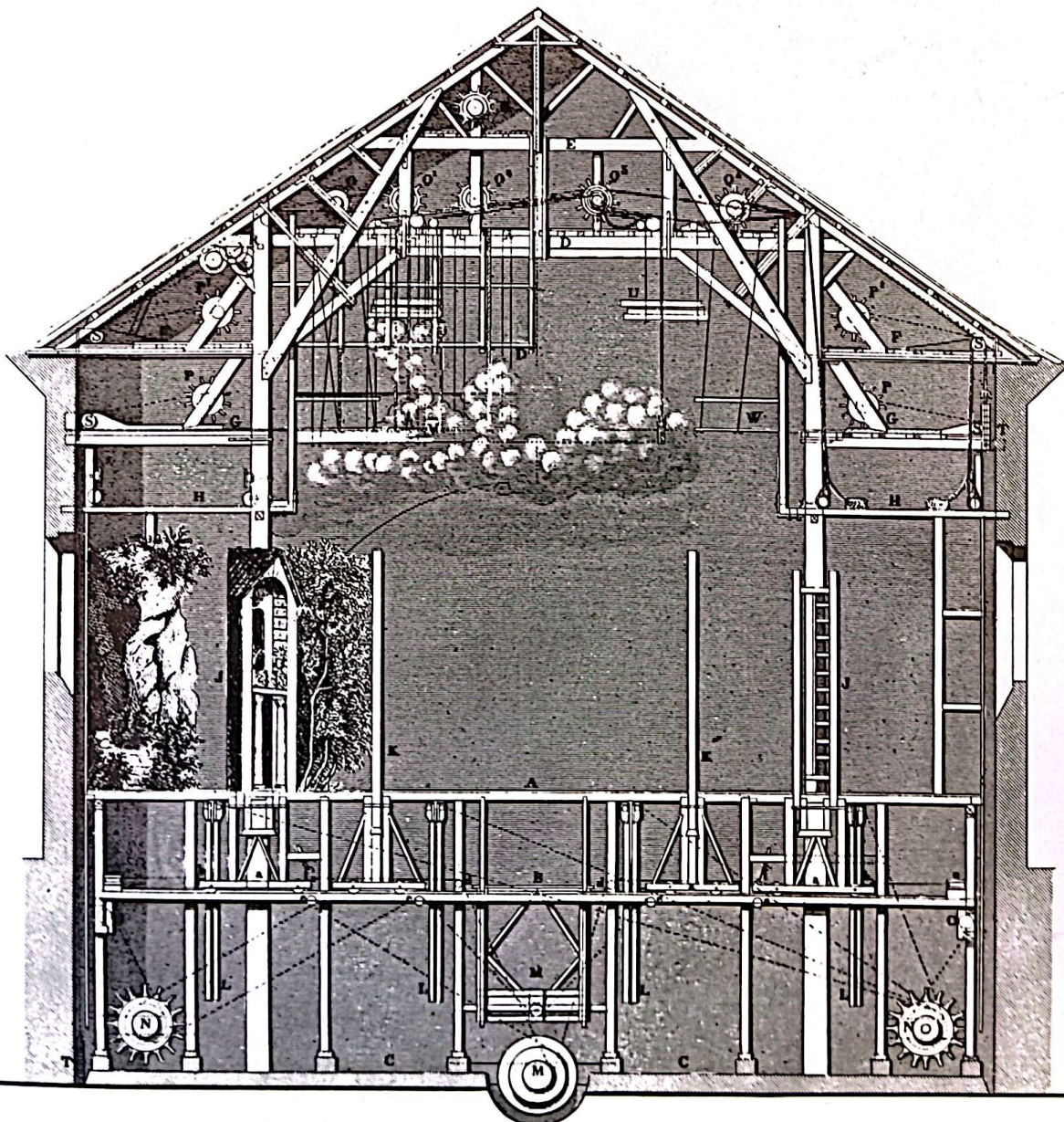


Marian Cleeves Diamond

ENRICHING HEREDITY

The Impact
of the Environment
on the Anatomy of the
Brain



Are our mental capacities determined at birth by our genes or can they develop in response to our environment? In the classic debate over whether anatomy is in fact destiny, any evidence of a physiological basis was regarded as evidence against environmental influences. Now, Dr. Marian Diamond, one of the foremost researchers of the anatomy of the brain, forges an important bridge between the two sides of the age-old nature/nurture controversy by demonstrating that *anatomy itself* can be changed by the environment.

Bringing together nearly three decades of extensive research, including previously unpublished data, she provides the evidence that, at any stage of life — from prenatal to old age — an enriched environment can improve brain size. Based on her pathbreaking studies with rats — whose brains resemble our own in many aspects of form and functioning — this breakthrough book shows how access to stimulating objects produces a thicker cortex and larger neurons, which, as other researchers have demonstrated, facilitate faster, more accurate problem solving — a key component of intelligence.

Enriching Heredity documents the additional benefits of an enriched environment in overcoming the negative effects of stress and of protein deficiencies caused by dietary imbalances. Perhaps most provocative are Diamond's findings that the offspring of mammals placed in enriched environments during pregnancy have larger brains than their mothers.

Diamond emphasizes that just as a stimulating environment can effect brain growth, the reverse also holds true: her data indicate that impoverished surroundings not only inhibit brain growth, but can actually decrease the size and number of its cells. Noting as well how male and female brains compare and contrast, she illuminates differences in their physical structure and development, and documents the effects of sex hormones in shaping brain anatomy.

(Continued on back flap)



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Marian Cleeves Diamond



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Dedicated with Love to
my mother, Rose, and my father, Montague,
and to Dick and Arne
for reasons they would understand



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PREFACE

The neural basis of behavior is a subject that has fascinated me for over four decades. As a young girl, I used to look at people and wonder what was actually going on behind their eyes. How was it possible for people to think, and what was responsible for the process? A considerable number of us who study neurobiology took our first steps in the field of psychology, where we hoped to answer questions dealing with the fundamental processes responsible for mammalian behavior. Though many present-day investigators continue to concentrate their efforts on behavioral processes, others study the structural and chemical components of the brain in an attempt to understand its functions. We are now aware that ions and molecules form brain cells and the connections among them, which in turn produce the mind and behavior. Utilizing this basic knowledge, thousands of scientists today are working at many diverse levels—behavioral, structural, chemical, molecular-biological—to unravel the mysteries of human behavior.

This book draws on 27 years of research from my laboratory dealing with environmental influences primarily on the anatomy of the mammalian forebrain, primarily that of the rat. One major aim of the book is to bring together at one time this body of information gathered in a semistepwise fashion over the years. It can now serve as a foundation on which to build a global picture, on the one hand, or a more refined one, on the other. These anatomical findings can also provide other investigators with information about the plasticity of brain regions at different ages and can be used as guidelines for more sensitive types of measures by chemists or molecular biologists. Some of the material has not been published previously; we are introducing it now to strengthen the larger story. As the data were gathered into a single body, patterns of a developmental sequence became evident that were not clear in separate, scattered publications. Over the years, results were coming in at

a rapid rate, and now I have had the opportunity to synthesize and correlate their meaning with greater insight. Facts that were discovered 20 years ago have now been integrated with our more recent data.

This book is intended for anyone who wishes to learn about the effects of different types of environmental influences on mammalian forebrain structures: both interested lay people and neuroscientists in all categories, from neuroanatomists, neurochemists, and neurologists to psychiatrists and social and behavioral scientists. Gynecologists and obstetricians, in particular, may gain valued insight into the effects of sex steroid hormones on cortical structures. Finally, this book can supplement a course dealing with neuroanatomy and behavioral biology or serve as a reference source.

The majority of the anatomical studies mentioned here have been completed in my laboratory with all of us working together: technicians, undergraduate and graduate students, postdoctoral fellows, and colleagues. These people have always been included as authors on the publications, and their names are found in connection with their research studies in the list of references for this book. As a rule, they are also mentioned within the text in conjunction with their work. The behavioral studies related to the early anatomical work were conducted by Professors David Krech and Mark Rosenzweig and Dr. Edward Bennett in the Department of Psychology at Berkeley. However, since the late 1960s, both the behavioral and anatomical studies have been confined to our Department of Physiology-Anatomy. I have not included all of the reported investigations dealing with environment and the brain. Emphasis here is on brain anatomy and the environment, and the results are primarily those from my laboratory, because these data have been collected to tell a particular story. However, when appropriate, some methods and results of experiments from other laboratories dealing with forebrain anatomy, chemistry, and behavior are woven into the text.

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Five minutes after Laura Wolff, senior editor of The Free Press in New York, walked into my office in Berkeley, I knew I wanted to work with her to produce this book. Not only is she clear thinking and wise but she is compassionate as well. Thank you, Laura, for a priceless collaboration and friendship. My copy editor, Richard Mickey, provided careful editing beyond compare. I am certain that part of his brain concerned with editing is enlarged into the upper one percent category of intelligent human beings. To Eileen DeWald, managing editor of The Free Press, I also am deeply indebted. What a team of excellence! Thank you one and all.



1

CAN WE CHANGE OUR BRAINS?

The nature-nurture controversy in regard to human behavior is an ageless one, extending from the times of Plato and Aristotle. The controversy has continued over the centuries because of a lack of relevant data to support the belief that the environment interacts with inherited characteristics on the one hand, or that heredity is all-powerful to the exclusion of the environment, on the other. The differences which exist among individuals as well as those between groups have been cited by proponents of both views. Until recently it was primarily sociologists, psychologists, and educators who emphasized the importance of the environment—specifically, the social and political inequities affecting the aged, women, the poor, and members of minority groups, to name but a few—in influencing behavior. Others believed that human behavior was the result of the divine will or biological predestination rather than environmental factors. Now, however, with the impressive advances in the science of genetics, many biologists are justifiably inclined to stress the importance of heredity. But with the studies presented in this book, we have evidence to support the view that the environment plays a role in shaping brain structure. These are the first controlled laboratory studies to demonstrate that various types of experiential environmental conditions can alter the anatomy of the outer layers of the mammalian brain, which in turn affects the learning ability of the organism.

This does not mean, though, that others had not previously predicted that such changes could occur. As early as 1815 Spurzheim asked whether tissues could increase with exercise, because blood is carried in greater abundance to parts that are excited, and nutrition is performed by the blood. He thought that this principle might certainly apply to the brain

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as well as to the muscles. Charles Darwin in 1874 noted that the brains of domestic rabbits were considerably smaller than those of the wild rabbit. He wrote that the differences might be attributed to their having been closely confined during many generations so that their intellect, instincts, sense, and voluntary movements were not exerted. In other words, the domestic animals did not have to face the natural threats of life in the wild. These are but a few early examples predicting that the structure of brains can change depending upon the input from the environment.

We now have evidence to illustrate the details of the anatomical changes that do occur with modifications in the environment. This evidence addresses many of the questions that concerned the early sociologists and educators, including the effects of the environment on the young as well as the elderly, sex differences, and the effects of nutritional deprivation, isolation, or crowding. It is now clear that the brain is far from immutable.

Documented speculations about brain changes in response to factors in the environment are not new. For example, in 1819, it was reported that an Italian anatomist, Malacarne, postulated that experience could alter brain structure (2). A century later, in 1911, a Nobel Prize-winning anatomist, Ramon y Cajal, suggested that "cerebral exercise" could establish new and more numerous connections between nerve cells in the brain (3). He realized, with the knowledge available at the time, that no new nerve cells would be formed after birth in the cerebral cortex, but he proposed that cells become bigger from use. Just over 50 years later, our neuroanatomy group at Berkeley measured a greater cortical thickness and larger nerve cell dimensions in the brains of rats that had spent some time in stimulating, interesting environments than in the brain of their brothers living in impoverished conditions. These were the initial experiments that set the stage for the production of this book.

The effort providing the background for our environmental studies extended over several decades. In the 1920s, in the psychology department at Berkeley, Professor Robert Tryon noted that some rats ran mazes better than others (4, 5). He wondered whether he could establish a maze-bright strain by breeding the animals that ran the mazes well—and similarly, whether he could breed a maze-dull strain. After several generations of inbreeding, he did create his two desired strains, one maze-bright and one maze-dull. Over the following years, the two strains were maintained separately from each other in the department colony.

In the 1950s two Berkeley scientists, a psychologist, David Krech, and a Nobel-laureate chemist, Melvin Calvin, asked themselves whether the brain chemistry of these two strains of rats might differ. Before they could try to answer their question, they had to decide which chemical would be most significant to measure. At the time, few neurotransmitters in the brain were known; one was the chemical, acetylcholine. The two scientists—Krech and Calvin—reasoned that if the animals were “bright,” they should have more of this chemical to facilitate transmission of an impulse across the synapse. Acetylcholine is very unstable and difficult to quantify, but its hydrolyzing enzyme, acetylcholinesterase, is stable at room temperature (at least up to 6 hours) and, thus, more easily measured. The scientists hypothesized that if there were more acetylcholine in the maze-bright animal, then there should also be more acetylcholinesterase to break down the acetylcholine after it had served its function. With this idea to work from, it was now possible to begin a team effort to study brain chemistry and behavior. Calvin suggested that Bennett, a neurochemist from his laboratory, collaborate as the chemist; and Rosenzweig, a psychologist, joined Krech to complement the psychology team. With time, the hypothesis of Krech and Calvin proved to be correct; there was more acetylcholinesterase in the brains of the maze-bright animals than in those of the maze-dull. This was the first evidence of a specific positive correlation between brain chemistry and learning ability (6).

In 1949, prior to these studies at Berkeley, Hebb, at McGill University in Montreal, had hypothesized that animals living in enriched environments early in life develop permanent brain changes that enhance problem-solving capabilities (7). Hebb had based his hypothesis on the fact that rats that had been used as house pets, and thus had experienced enriched living conditions, were better at running mazes than were confined, caged rats. In the early 1950s, several psychologists, stimulated by Hebb’s hypothesis, began to ask how much enrichment in an adult animal was necessary to produce maze-learning abilities superior to those of a nonenriched animal. They learned that rats that had been exposed to enriched, rather than impoverished, environments early in life were better maze learners upon reaching young adulthood.

A logical next question was whether animals living in stimulating environments differed from animals living in isolated conditions not only in behavior but also in brain chemistry. The Berkeley team of scientists drew on the reasoning and techniques they had used to study the maze-bright and maze-dull strains and applied them to the brains of animals

that had experienced enriched or impoverished conditions. They found that the brain acetylcholinesterase concentration was greater in the environmentally enriched rats than in the impoverished ones (8).

While working at Cornell University in upstate New York in the 1950s, I read about these studies that correlated behavior and brain chemistry and wondered if there might be measurable structural differences in the brains of rats with dissimilar behavior and chemistry. For example, did the maze-bright and maze-dull animals differ in morphology of the cerebral cortex? Were nerve cell processes restructured into new patterns in these strains of animals? Could we somehow demonstrate a connection between brain anatomy and learning? My resulting excitement about various possibilities was intense. This initial curiosity then led me on continuously for the next several decades, being rekindled with each set of new data. As with most research, each answer led to a dozen more questions; to stop at any one point was tremendously dissatisfying.

Several preliminary studies were undertaken before we embarked on the long series of investigations dealing with the anatomy of the developing and aging brain and the environmental influences which changed the basic structural patterns. The first dealt with the localization of acetylcholinesterase in various cellular layers of the cortex in maze-bright and maze-dull animals. After identifying enzyme differences between the two groups, we counted cerebral-cortical nerve and glial cells in several maze-bright and maze-dull strains of rats to learn how the number of cells correlated with the enzyme distribution. We found no significant correlations. Cells were counted in acetylcholinesterase-high and -low strains of rats as well. Again no significant findings were obtained. Undoubtedly, differences were present at the level of the connections between cells, but from our results, not in cell number. But these initial experiments helped organize our thoughts for the major thrust of our anatomical experimental work over the years ahead. We had a better idea of what kind of morphological measurements could help us answer the unending questions about the brain and the environment.

This book will present the knowledge gained from such experiments in two main sections: (1) normal forebrain development and aging, and (2) modifications of forebrain development and aging due to environmental factors.

For many years after we began to study the effects of the environment on the cerebral cortex, it was never clear how the data fit into the lifetime continuum of brain development and aging. No baseline for the dimensions of the rat cortex, for example, was available for the young, adult, and old-aged animal. Roger Sperry, the Nobel laureate from California

Institute of Technology, once said, "Marian, all you are doing with your enriched environments is stimulating the maturation of the cortex." We did not know whether he was right or not. Were the stimulating environmental conditions increasing a growing, maturing cortex, or a cortex which had reached a plateau, or a decreasing, shrinking cortex? When does the cortex stop growing, and how does it age under "normal" laboratory conditions? In order to answer these questions, we accumulated, over a 7-year span, information on the patterns of development and aging in the male and female cortex and other forebrain structures. We wished to learn about the cell populations in the cortex over a lifetime. Not only was it important to examine the cortex as a whole, but we wondered whether the right and left cortices followed similar patterns during development and aging because new information was accumulating about functional differences in the two hemispheres of human beings. Would structural differences help us to understand the basis of the functional aspects?

In recent years a great deal of evidence has been offered indicating that the separate hemispheres of the brains of human beings serve different functions. It has been said that the right hemisphere is concerned especially with spatial mapping, processing information holistically and simultaneously, and with artistic and musical functions, whereas the left is involved in more analytical, sequential processing and in the production and understanding of language. Admittedly, such precise functions cannot be attributed to locations in the rat cortex, but with the knowledge that differences between the hemispheres exist, we thought it imperative to look at the asymmetrical structure of the two hemispheres and to learn how the separate patterns are maintained during the lifetime of the animal. By examining asymmetrical patterns in the rat, we hoped to gain an understanding of factors responsible for creating asymmetry.

Since sex differences in thickness of the outer layers of the brain were becoming more obvious with each type of measurement, the role of the sex steroid hormones in creating these differences demanded attention. Markedly different patterns of asymmetry in thickness were noted between the male and female rats. Aging had a specific effect on asymmetry. In order to learn what structures were responsible for these thickness differences, neuron and glial cell counts were made on samples of the right and left cortices in males and females. Were the numbers of neurons and glial cells partially responsible for thickness differences?

Investigators have reported that both male and female rat cortices contain estrogen receptors. We were interested to learn whether these receptors were equally concentrated in the hemispheres of the male and

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female rats. If not, was it possible that they played a role in establishing brain asymmetry?

It was becoming clear that cortical asymmetry was related to sex hormones. But in 1980 some French scientists found that asymmetry was also related to other factors. These investigators had discovered that lesions in the left cerebral cortex affected the immune system differently from lesions in the right cortex. However, they did not confine their lesions to specific regions in each hemisphere of the cortex, but performed extensive cortical removal. They did not make refined lesions because they were using the cortex as a control and had not expected to alter the immune system with decortication. We then attempted to benefit from their findings and tried to localize more specifically which cortical areas were playing a role in this reported cortical immune response. One important question arose from their results: If we could alter the cortex with stimulating environments, could we then someday carry out experiments which would strengthen the immune system with "enriched" conditions?

The results of our experiments dealing with cerebral development and aging indicated that mammalian cortical nerve and glial cells are subject to structural changes due to age, sex hormones, and immune response. But what about other forebrain structures? We learned that areas such as the hippocampus, entorhinal cortex, amygdala, and corpus striatum differed considerably from the cerebral cortex in their developmental and aging patterns. Each will be discussed briefly to indicate the importance of referring to a specific forebrain region rather than to the whole brain when dealing with growth and aging studies.

Once we had accumulated fundamental information, or baselines, on the "normal" laboratory rat forebrain, we were prepared to investigate how alterations in the environment could induce modifications in the structure of rat brains. We were interested to learn how the thickness of the young cortex was affected by enriched and impoverished living conditions. Once we had established that, we made more refined anatomical measurements, such as neuron number and size, glial cell number, and blood vessel size. We examined the effects of enrichment during the cortex's natural growing process, and then throughout the long period of natural decline in cortical thickness.

During the first postnatal month—the growing period of the cortex—the pups had to live with their mothers in either enriched or impoverished conditions. The experimental manipulations we made and the measurements we took during this time indicated whether these conditions were affecting maturation or not.

The most common experimental design used for decades in our laboratory was that provided for the already-weaned rats, those that no longer lived with their mothers. For the majority of these experiments, three basic environmental conditions were used: enriched (12 rats per large cage, 70 × 70 × 46 cm, plus toys); standard colony (3 rats per small cage, 20 × 20 × 32 cm, with no toys); and impoverished (1 rat per small cage and no toys). The cerebral-cortical thickness was measured on rats that had lived in these conditions for varying periods of time ranging from 1 day to 160 days. We were interested to learn whether the cortex could change both for animals as young as 14 days of age and for animals as old as 904 days of age, the latter roughly equivalent in the lifespan of rats to 90 years for a person. Not only were cortical thickness modifications measured in the brains from animals living in various environmental conditions, but nerve and glial cell counts were taken as well. The dimensions of the nerve cells were measured, including the size of the nerve cell body and its nucleus, the number and length of dendrites, the number and distribution of spines, and the length of the postsynaptic density. We also examined the internal and external skull dimensions to determine whether the anatomical and neurological changes within the brain affected the skull.

Since there has been much interest in training one side of the brain separate from the other, we wished to determine whether our multisensory environmental conditions altered the cortical thickness of one hemisphere differently from the other.

Both males and females were exposed to their enriched or impoverished conditions for similar periods of time. In addition, the brains of males living only with males during the enrichment period were compared with those of males living with females. Cortical thickness measurements were made on these groups of rats.

Nutritional effects on brain development and aging are of ongoing concern as we attempt to learn about conditions that best facilitate brain function. Our laboratory has collaborated with one investigator who carried out a set of nutritional experiments dealing with protein-deficient mothers and their nutritionally and environmentally rehabilitated offspring. Environmental enrichment proved to be an important factor even with the nutritionally deficient animals.

While the above nutritional experiments were in progress, we became interested in the relation between glucose metabolism and environmentally enriched conditions. In collaboration with Dr. Carolyn Smith at the National Institute of Mental Health in Bethesda, Maryland, we carried out a study to learn whether the metabolic rate of the brains in

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rats in enriched environments was greater than in the standard colony animals. The results from this experiment were quite the opposite from our predictions and reminded us of the importance of doing experimental work to substantiate hypotheses. One cannot always foresee the many variables involved.

In the fall of 1985, we were invited to China to lecture, and I wondered what experiment with our rats might be of interest to our hosts. With their problem of overpopulation, perhaps the effect of crowding during brain development would be of interest. We conducted an experiment for which we used 36 animals in the enriched cage instead of the usual 12 and later measured the cortical thickness changes in them compared with the standard animals.

Is it possible that factors such as air ions could affect the outer layers of the brain? We were most fortunate to have one of the world's experts on air ions, Albert Krueger, working down the corridor at Berkeley (one of the precious advantages of being at this remarkable university), and we collaborated with him and his group on experiments dealing with the effects of air ions on cerebral-cortical structure and chemistry. Once we learned that ions did alter the cortex, we became interested in their effect on the immune system, specifically in the way they alter white blood cell counts. The complexity of the many interactive factors in the body and the environment was becoming more evident and intriguing.

Most of our efforts have focused on understanding the changes produced by environmental enrichment and impoverishment in the brains of animals exposed to those conditions. However, we have also been intrigued by the question of the possible transmission of the changes from these animals to their offspring. In other words, was it possible to detect morphological changes in the brains of future generations? Admittedly, this question suggested a possible Lamarckian view, but several factors could be involved which needed to be explored.

All of our brain measurements—whether of cortical thickness, dendritic branching, dendritic spines, or synaptic length—have little meaning without an index of behavioral changes. Our laboratory has not maze-tested the enriched or impoverished rats. But psychologists from many other laboratories have studied such rats in several different kinds of mazes and have found evidence of behavioral changes that correlate well with our anatomical brain results. (The data will be discussed in Chapter 9.)

Our ultimate goal in studying the brains of rats is to gain a better understanding of the human brain. Obviously, for ethical and practical reasons, we cannot completely control the environmental input into hu-

man brains, nor are we able at present to make precise anatomical measurements of identical areas of the brains of human subjects over time. Even the scanning techniques of computerized tomography (CT) and positron emission tomography (PET) do not give sufficient resolution, and besides, these methods are not safe to use for repeated measurement of living, healthy human brains. In the future, the noninvasive technique of magnetic resonance imaging (MRI) may offer more discriminating measures for detailed anatomical study of the healthy human brain. In the meantime, we use animal models; we can control the environment with animal subjects, and we can very precisely measure cortical dimensions on preserved animal brain tissues.

One question which is invariably raised about the study of rat brains is, How are the results applicable to humans? Needless to say, one cannot make direct extrapolations from the many variables involved, but several of the basic principles which are established from rat work can be applied to humans. For example, whether we are dealing with rats, cats, dogs, monkeys, or human beings, the brain consists of nerve cells and glial cells. Investigators have shown that the number of neurons in a single column of cortical nerve cells is the same in rats, cats, dogs, monkeys, and man (1). Furthermore, in all these species most nerve cells have branches that we call *axons* and *dendrites*. It is the pattern and the quantity of these branches, i.e., the complexity of the circuitry, that account for some of the differences among species. The cell bodies and branches of the neurons communicate with each other through functional junctions called *synapses*. As far as we know at present, the numerous chemical neurotransmitters located in the billions of synapses in the brain are of a similar nature in rats and man.

Nerve cells are designed to receive stimuli, store information, and transmit impulses, and this is true in both rats and man. Furthermore, much of what we know about the human reproductive system and its hormonal relationships has been established from studies with rats. In essence, most of what we have learned about the normal functioning of the human nervous system and its associations with the reproductive hormones has been learned from animals.

The kinds of brain measures used on our rats and the accompanying results are basic to the understanding of neural structure and function whether we study rats or man. Our findings may have wide enough application to encompass mammalian nerve cells in general. Both the human brain and the rat brain are very immature at birth, especially in the cerebral cortex, the structure which has received most of our attention. As will be described in more detail in Chapter 2, the cortex grows

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at an explosive rate immediately after birth, but the patterns of development are different for males and females—a finding which cannot be easily measured in humans with present technology.

There are many reasons to believe that data collected from rat brains can be useful in establishing guidelines for studying events occurring in the human brain. Accepting the vast complexity of the human brain, I only hope that this collection of information from controlled rat experiments and from a few studies on human brains will eventually serve in directing others toward a better understanding of the potential of the human brain . . . and thus, to greater efforts to improve upon the human condition.

2

NORMAL FOREBRAIN DEVELOPMENT AND AGING

The massive right and left cerebral hemispheres account for about 85% of the total human brain and about 45% of the rat brain. The outer few millimeters of nerve cells on these hemispheres constitute the cortex, commonly referred to as the cerebral cortex ("cortex" means "bark"). In actuality, there are three divisions of the cortex: the three-layered archicortex; the five-layered paleocortex; and the six-layered neocortex. The neocortex was of greatest interest to me not only because it is the seat of higher cognitive functions but because it is one of the last structures to develop embryologically and is one of the most recent phylogenetically. The neocortex of the human brain is a structure whose abilities are unique in the animal kingdom. (The term cortex or cerebral cortex will be used to refer to neocortex unless otherwise specified.)

Before we could make sense of our measurements of the cortices of enriched and impoverished rats, we first had to map the normal developing and aging pattern of the cerebral cortex for use as a standard for comparison. To obtain this standard, or baseline, we measured the thickness of the cortex on both female and male Long-Evans rats that had lived 3 per cage for varying periods of time to attain different ages. In our early work (1) we had dealt mainly with rats between 25 and 105 days of age and did not know whether the cortex was normally increasing or decreasing during this period. We realized what an important consideration we had overlooked and proceeded to gather data from the developing and aging cortex.

First, let us look at the cortex through data from both hemispheres combined to obtain the overall cortical developmental events. Figure 1 illustrates the location of brain samples removed for microscopic examination of the frontal, somatosensory, and occipital cortices, from anterior to posterior. The numerical designations according to Krieg (2) are placed on the left side of the hemisphere: Very brief functional descriptions are offered: area 10 represents the frontal cortex, an area in the human brain used for sequential planning; area 4 represents the motor cortex; areas 3 and 2 designate general sensory areas; areas 18 and 18a are visual association areas; and 17 is the primary visual cortex. Area 39 represents a multisensory integrative area.

Figure 2 shows a transverse section of the rat brain illustrating divisions of the occipital cortex as an example and how the thickness was measured on lines extending from the surface to the underlying white matter. The sections were cut at 20 micra (a micron or μm is one one-thousandth of a millimeter). We have learned that several cellular features

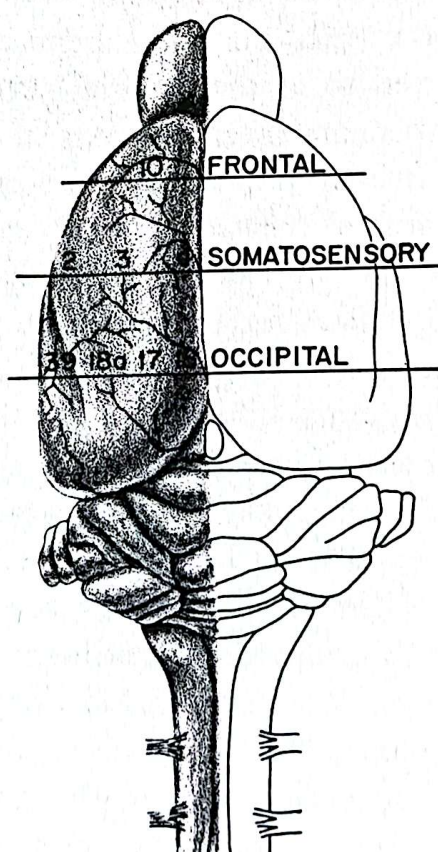


FIGURE 1. Dorsal view of rat brain indicating regions sampled for study. (Right) Frontal, somatosensory, and occipital sections are illustrated. (Left) Numerical designations of Krieg (2).

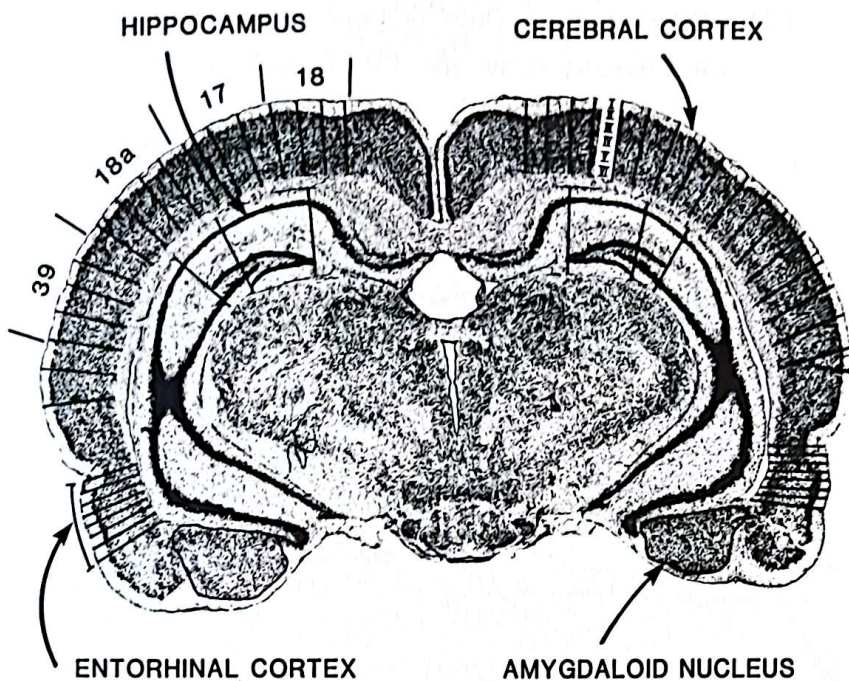


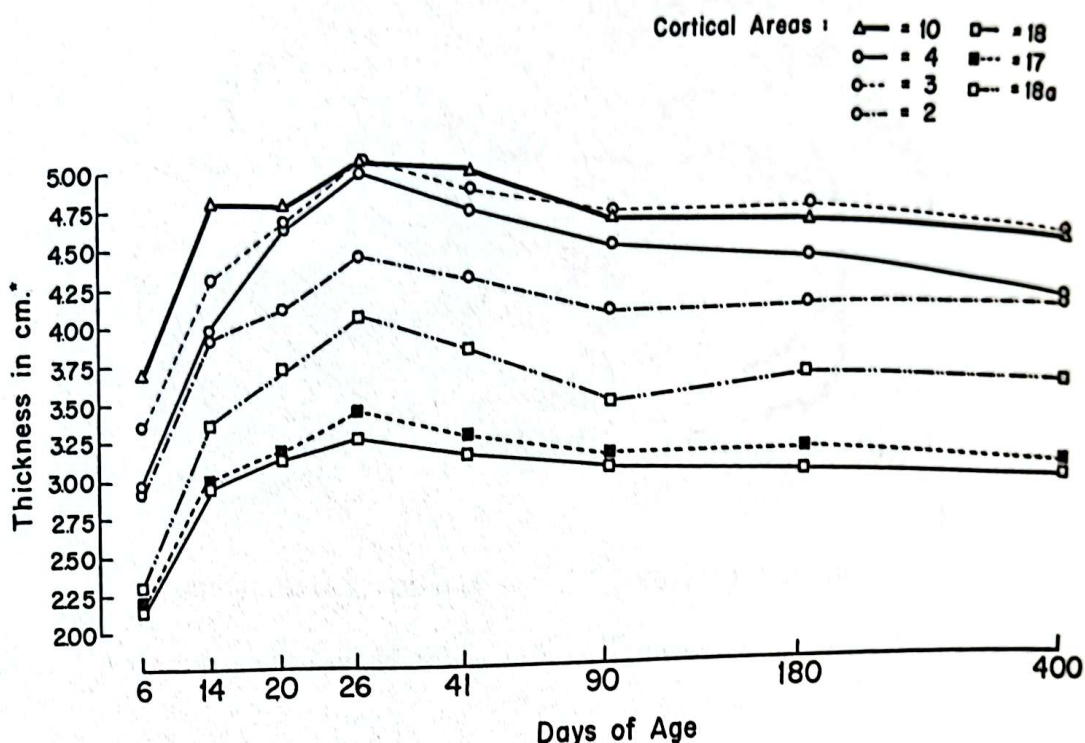
FIGURE 2. *Transverse or coronal section of rat brain indicating areas studied.*

account for cortical thickness, such as nerve cell density and size and glial cell number. The cortical cell layers I through VI are also shown.

Differences between Male and Female Cerebral Cortices

Figure 3 represents a growth curve of the frontal, somatosensory, and occipital cortex from 6 days to 400 days of age based on an average of 15 to 17 male rats per age group (3). This figure shows that the most rapid growth of the male rat cortex occurs between 6 and 10 days after birth. The cortex continues to grow until it reaches a peak, and a general decrease in thickness begins, sometime between the ages of 26 and 41 days. The early postnatal cortical increase in the male rat amounts to almost 45% before the peak is reached. Though not shown on this graph, a 9% decrease occurs between the ages of 41 and 650 days. After day 650 all cortical areas continue to decrease, but the occipital cortex decreases more steeply than the other regions until the last measurement at 904 days. These developing and aging cerebral cortical data fall into two basic slopes on the graph: a positive one before 41 days of age and a negative one throughout life after 41 days. Our challenge is to see whether environmental input can alter these slopes.

Cortical Thickness (cm) — Development and Adulthood in Male,
Long-Evans Rats ($N = 15-17/\text{age group}$)

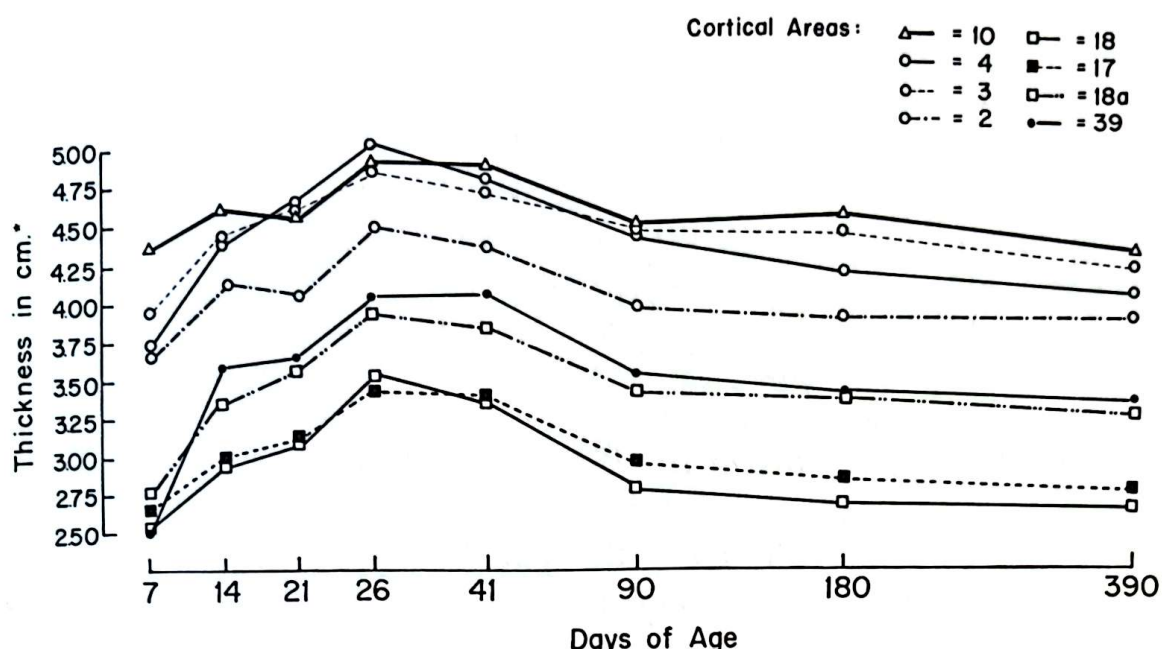


*enlarged microscopic units

FIGURE 3. Cortical thickness (cm): development and adulthood in male Long-Evans rats ($N = 15-17/\text{age group}$).

In turning to the female cerebral cortex (Figure 4), we find a different developmental pattern from that of the male. The female frontal lobe is already fairly well developed at birth and grows only 15% in the first few weeks until it reaches a peak and then begins to decrease. The female somatosensory cortex is also more further developed at birth than is that of the male. In the female a major sensory integrative area, area 39, grows by 45% during the first week alone and does not reach its peak until close to the age of 45 days (see Figure 4). We do not have this precise measurement for the male. The two areas—frontal (10) and somatosensory, show clearly that the female cerebral cortex has a developmental pattern quite different from the male. In general, the female's cortical thickness is greater at birth than the male's, but by 3 weeks the male's cortex is thicker than the female's and the difference in thickness holds as the animals age. The occipital cortex in the male becomes markedly thicker than that of the female. Yanaid (4) found that the rate of neonatal proliferation of cortical cells in the male is slower than in the female. He suggests that such a phenomenon may represent "delayed maturation" of the male brain as compared with the female. It may be

Normal Forebrain Development and Aging 15
 Cortical Thickness (cm) — Development and Adulthood in Female,
 Long-Evans Rats (N = 10-20/age group)



*enlarged microscopic units

FIGURE 4. Cortical thickness (cm): development and adulthood in female Long-Evans rats (N = 10-20/age group).

assumed that females have a higher rate of proliferation of brain cells at a prenatal stage.

If natural selection operates in such a way as to ensure the reproduction of the species, it must promote the survival of young members of the species. It is possible that the female cortex is more highly developed at birth to ensure a better start. The human female, for example, ovulates only about 420 times during her life, whereas the human male discharges over 2 hundred million sperm with each ejaculation. The fact that she has fewer opportunities to reproduce suggests the importance of the early well-being of the female as she adapts to her environment.

Just as different regions of the female cortex develop at different rates, they also age differently, at least up until the age of 390 days—the age of the oldest group we have yet measured. In females between the ages of 26 and 390 days, cortical areas 4 and 18 decrease in thickness by as much as 22 to 31%. At the same time areas 10, 3, and 2 age only by half as much, by 11 to 15%. During the second week, we have seen that the frontal lobe increases by 6%, at the same time that area 39 increases by 45%. In the male all areas of the cortex increase by 28 to 46% during this same period of time. It appears that, on the whole, the separate

regions of the female cortex develop and age in a less uniform pattern than those of the male.

As stated earlier, most of our work has dealt with the rat brain, a fact which evokes the usual cautions about extrapolating from rats to man. And indeed there are differences on many levels. In the cerebral cortex of the rat, neuronal cell division is complete shortly after birth; it is followed immediately by profuse dendritic branching. Our results have shown that this rapid cortical development in the rat occurs during the first 40 to 45 days postnatally, with slightly different patterns in the male and the female. In contrast, cell division in human brains continues for about 1 year postnatally (5). But in human brains, also, rapid dendritic growth takes place during the first 3 to 4 postnatal years; as Dobbing and Sands point out (6), at least five-sixths of human brain growth by dendritic branching is postnatal and "in this respect humans resemble rats more closely than formerly thought."

Some very recent results on the developing human cortex support the notion that human brain growth follows the general developmental pattern seen in the rat cortex. From over 100 children with various neurological disorders, investigators selected 29 children (age 5 days to 15 years) who had suffered transient neurological events that did not measurably affect their normal neurodevelopment (7). The brains of these children were studied with 2-deoxy-2-[¹⁸F]fluoro-D-glucose and positron emission tomography (PET scan) to plot the functional development of the brain. By examining the resulting human growth model, we learned that the absolute values for cortical glucose uptake were low at birth and rapidly increased up to the age of 3 to 4 years. These high rates of glucose uptake were maintained until about the age of 10 years, when they began to decline (8).

However, the authors of this study grouped their data on 3- to 8-year-olds into a single sample. When we examine separately the actual data points on their graphs, it appears that the cortex is beginning to decrease after the age of 3 to 4 years, and so including the 3- to 8-year-olds into one sample masks the initial stages of the decrease. Looking at the measurements for each year, we see a pattern similar to that found in the rat. The basic cortical developmental pattern shows a rapid increase after birth followed by a decline one month later in the rat or about 10 years later in the human brain. These new findings with the human cortex provide additional support for using rat data as a guideline to mammalian cortical morphology during developing and aging. We can therefore obtain knowledge from animal models about the human cortex.

Asymmetry and Sex-Hormonal Influences on the Cerebral Cortex

Even though scientists were aware from about the middle of the nineteenth century that the two halves of the human brain were linked to some specific behaviors such as speech and awareness of body image, most rat brain scientists did not consider brain asymmetry in their chemical or anatomical studies until recently. It was not until 1975, when we were studying the normally developing and aging Long-Evans rat cortex, that we decided to make detailed comparisons between the hemispheres (9). We learned that in the male rat the right cerebral cortex was thicker than the left.

The male's right cortex was also thicker than the left in the S₁ strain (maze-bright) rats from the Berkeley psychology colony. [By using two strains, we established that the cortical asymmetrical pattern in the male rat brain was not peculiar to only one strain of animals.] To determine the cerebral-cortical thickness, we measured three representative transverse forebrain samples; in accordance with our usual procedures, we obtained our representative sections by utilizing subcortical landmarks for maintaining constancy. For example, the frontal cortical sample was taken immediately anterior to the genu of the corpus callosum; the somatosensory sample, at the crossing of the anterior commissure; and the occipital sample, at the level of the posterior commissure. These landmarks were chosen to allow us to compare regional degrees of asymmetry. (The regions included 10, 4, 3, 2, 18, 17, 18a and 39, according to Krieg's designations; see Figure 1). In order to understand whether asymmetry is present in the young and continues throughout the lifetime of the animals, we measured sections taken from male rats from birth to 904 days of age (the age of the oldest rats we have had).

Figure 5 presents the percentage differences between the right and left cerebral-cortical thicknesses in preweaned (6 to 20 days), adult to middle aged (90 to 400 days), and old aged (900 days) male Long-Evans rats, using 7 to 15 rats per age group and a total of 93 animals. The newborn male showed a similar right-greater-than-left pattern, but the data are not presented here. It is clearly evident that at every age and in every region except one, the right cortex is thicker than the left, with the differences ranging from 1% to 8%. Out of the 49 areas measured in all age groups the differences were statistically significant in 31 of them, or 63%. Some of the statistically nonsignificant differences were in areas which did show asymmetry in some animals. But the failure of

**Percent Difference Between Right and Left
Cerebral Cortical Thickness in Young Adult
and Old Aged Male Long-Evans Rats**

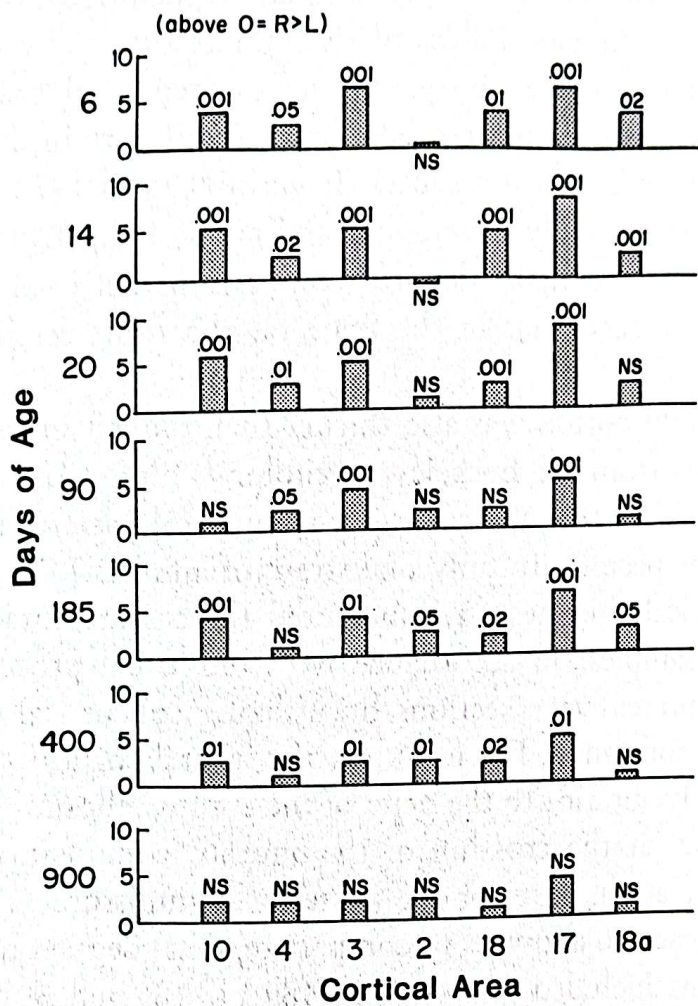


FIGURE 5. Percent differences between left and right cerebral-cortical thickness in young, adult, and older-aged male Long-Evans rats.

some, especially the very old, animals to exhibit any significant asymmetry partially explains the relatively low overall percentage difference in asymmetry.

For example, area 3, a general sensory area, and area 17, the primary visual cortex, showed the most marked right-greater-than-left differences consistently at every age. On the other hand, there was an apparent lack of asymmetry in area 2—another general sensory area, which in the rat receives input from the whiskers and from other sensory receptors—until the animal reaches 185 days of age, at which time it shows a statistically significant difference. Such a finding is intriguing in light of the marked asymmetry in the area adjacent to area 2, namely, area 3. Both regions are reportedly somatosensory areas. These data indicate that the sensory input from the whiskers, or the way that input is processed, differs from

other sensory input in that it requires similar handling by both the right and left hemispheres.

In the male Long-Evans rat, the clearly defined asymmetry or laterality in some regions appears to be governed by testosterone. When the cortex was measured at 90 days of age on rats from which the testes had been removed at birth, the sections through the frontal and parietal cortex showed that the left cortex had become thicker than the right, a pattern similar to that of the intact female. However, in the occipital cortex, the right side retained its original dominant size. From these results, it becomes evident that some regions of the male cortex are governed by testicular hormones, though in others, perhaps, the thickness is determined more by genetic factors. Evidently, cortical asymmetry, as noted in the young and adult male rats, is essential for specific behavioral patterns during most of the animal's life. However, in the very old male rat (900 days), the right-dominant pattern is no longer statistically significant. Even the occipital cortex, which retains its laterality when the testes have been removed at birth, shows no significant right dominance in the very old animal.

After studying the asymmetry in the Long-Evans rats, we utilized data from male rats of the S₁ strain which were obtained from some of our other experiments. Again the right hemisphere was on the average thicker than the left in most areas. Areas 18 and 2 were the exceptions; there the two hemispheres were equal. In areas medial 10, 4, 3, 17, 18a, and 39, the hemispheric differences were statistically significant. By presenting these more detailed data from two strains it is possible to determine which areas demonstrated consistent asymmetrical patterns. Data from the two strains were similar for all areas except area 18, a visual association area; the results for area 18 were right greater than left in the Long-Evans strain and right equal to left in the S₁ strain.

In comparing the growth rates in the right and left hemispheres from 6 days to 90 days in the Long-Evans strain, we found the patterns to be very similar. They essentially paralleled each other. In comparing the left and right cortices in the S₁ strain between different age groups ranging from 55 days to 105 days, we observed a less consistent asymmetry pattern. For example, between 55 and 64 days of age the right hemisphere increased; it then decreased until 105 days of age. The left hemisphere did not follow a parallel developmental course in areas medial 10, 4, 17, 18a, and 39. Since the S₁ strain was bred specifically for maze-bright experience, it is possible that the right-dominant visual cortex was especially susceptible to stimulation at a time near sexual maturity.

A separate but similar study was made of the right and left cortical

thickness patterns in the cerebral cortex of the Long-Evans female rat. We found quite different asymmetrical patterns from those seen in the Long-Evans male (10). (See Figure 6; also see note 49 for statistical tests used on our data.) The female did not show the strong asymmetries. (In sampling the female brains, we did not use as great an age span as with the males, only 7 to 390 days. The 900-day-old animals in the male study were a recent addition; we are at present attempting to raise the females to 900 days.)

In our earlier studies with 54 measurements of right-left differences in the female cortex, few statistically significant differences were found, but the left cortex was thicker than the right in 35 of the measurements, a 65% difference. We have since studied additional age groups (11). Using female brains from 18 to 41 days and as a result including 9 new measurements, making a total of 63 measurements, we found the right

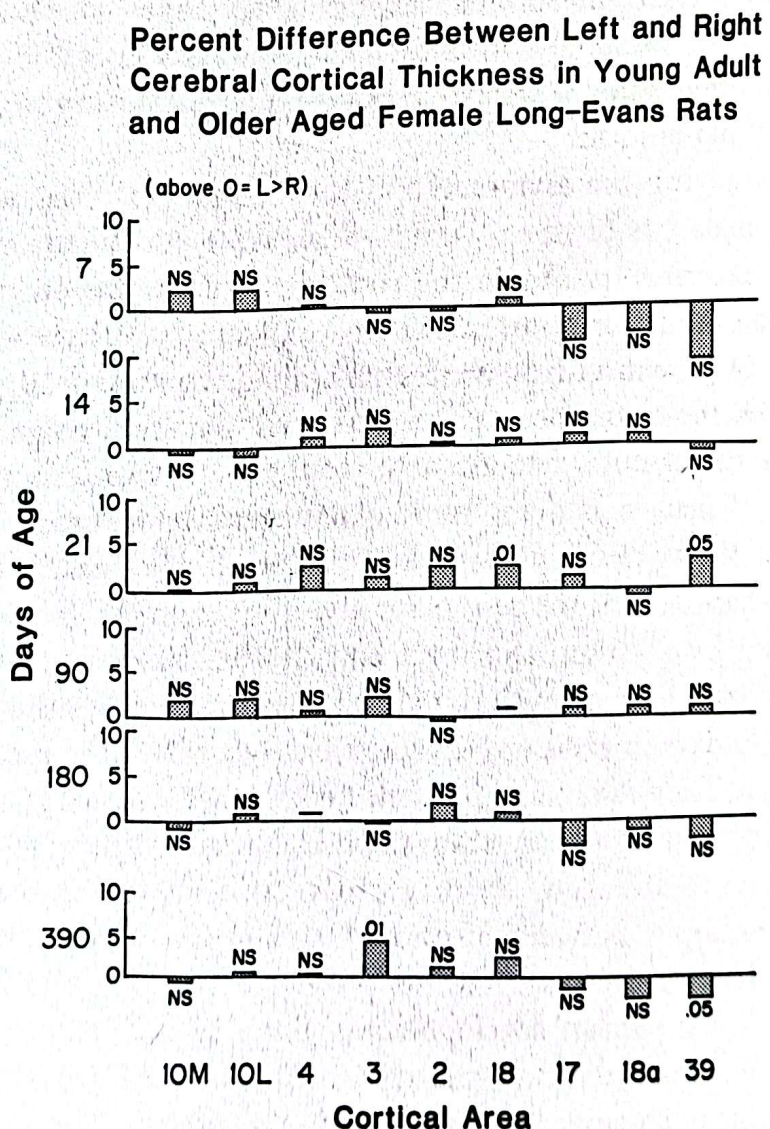


FIGURE 6. Percent difference between left and right cerebral-cortical thickness in young, adult, and older-aged female Long-Evans rats.

to be nonsignificantly greater than the left in 65% of the cases. From examining all these female data, it appears that the female cortex is not significantly asymmetrical. Yet, evidence to be discussed later shows that, for example, the estrogen receptor concentration and neuron and glial cell counts do follow the left-dominance pattern in the female, at least for certain age groups.

In comparing the male and female asymmetrical cortical patterns, sex differences are evident. The male shows quite significant right dominance in many areas, while the female shows no significant dominance. Our data suggest that the male loses significant cerebral dominance in old age, which may indicate that his brain becomes more like that of the female. Could these changes in cerebral dominance patterns be partially responsible for the decline in aggression seen in the older male? It has been reported that the older human male has more "domestic qualities" and prefers to stay closer to the "hearth" than the younger male (12). Once again the rat data offer directions for future studies of male human brains.

Most male rats are superior to most females in visual-spatial ability, which may be related to the greater importance of spatial perception for territoriality in the male (13, 14). Right structural dominance in the visual-spatial region of the cortex provides a basis for a key to understanding male behavior. In fact, Greer Murphy (15), working in my laboratory and utilizing the asymmetrical guidelines offered by the male rat, examined 31 human visual cortices and found the same trend as seen in the rat. He measured the volume of area 17 and learned that the right cortex was about 5% ($p < 0.05$) greater than the left.

The male's greater asymmetry in other cortical regions as well as the visual cortex may give him an ability to focus his attention more intensely, and this ability may be related to testosterone levels. As shown previously, a severe reduction in testosterone due to removal of the testes alters the male cortical dominance pattern in certain regions.

Our data showing that female rats lack significant asymmetry in the cerebral cortex are paralleled by recent findings regarding the human female brain. Here two fiber bundles connecting the two halves of the forebrain, the large corpus callosum (16, 17, 18) and the smaller anterior commissure (19), are measurably larger in the female than in the male. The corpus callosum, consisting of about 300 million fibers, connects the right and left cerebral cortices. Male and female differences have been demonstrated on preserved material from both fetal and postnatal human brains. Other investigators, utilizing magnetic imaging on live human brains, do not agree with these findings (20). But I have examined carefully de Lacoste's study on the fixed corpus callosum and have observed

the sex differences on the enlarged photographs of the anterior commissure, which connects not only right and left cortices but the olfactory bulbs as well. Both these examinations lead me to agree that these structures are larger in the human female than in the male.

Speculations with regard to the meaning of these female brain data are enticing. The female rat results revealing no significant cortical asymmetry and the human female reports showing a thicker connecting link between the hemispheres indicate that it is important for a more balanced distribution of information to pass between the female's two cortices. What might be an advantage for a lack of prominent asymmetry in the female? Possibly marked asymmetry might make the brain less flexible in responding to new situations. Asymmetry might be more of a hindrance than a benefit to the female, confining her range of behavior so that protection of the immature young would be too limited as they explore their new surroundings. The structure of the female cortex suggests that it allows for a "free flow" of information to occur between the two sides of the brain, providing a more diffuse response to the input from the environment.

This lack of a pronounced cerebral asymmetry in the female's cortex may be the "natural" state, but we have shown that internal environmental influences, such as levels of sex hormones, can alter her basic patterns. For example, in the brains of female rats that had been ovariectomized at birth and were later studied at 90 days of age, we found that the right cortical mantle was thicker than the left in 7 out of 9 regional comparisons, with significant differences in areas 17, 18a, and 39. Specifically, the right hemisphere was greater in area 17 by 3%, in area 18a by 5%, and in area 39 by 5%. In other words, without her ovaries for 90 days, her cortical thickness pattern became very similar to that of the male at 90 days of age with his testes intact. These data support our hypothesis that the ovarian hormones play a role in establishing and maintaining cerebral dominance.

By once again examining the data in Figure 6, we see a right-dominant pattern developing in the occipital cortex of the 180-390 day old female. These data suggest that by obtaining further old-age data, we may encounter an even stronger right-dominant pattern in this region of the brain. Does the female brain become more like the male with aging? It has already been shown that the 904-day-old male loses his significant cortical dominance with aging, thus becoming more like the female. If all goes well, we will have new data on the cortex of the very old female in the not too distant future.

With the knowledge that sex steroid hormones can alter cortical

structure and that the hemispheric asymmetry patterns are different in male and female rats, we continue to pursue an understanding of asymmetry in the cortex. It has been demonstrated that estrogen receptors are present in the cerebral cortex of both sexes for the first 3 weeks of postnatal life (21, 22, 23). But no one has reported whether the concentration of these receptors differs between the right hemisphere and the left. If the sex hormones do play a role in establishing hemispheric differences, knowledge of estrogen receptor concentration is important.

We found, in newborn rats that estrogen receptors are more highly concentrated in the male left cortex than in the right. Exactly the opposite pattern characterizes the female cortex. These data were discovered by using radioactive estradiol to determine the concentration of the receptors in the cortex during the first month of postnatal life (24). The results presented in Figures 7 and 8 clearly indicate that the concentration of these receptors is highest immediately after birth, rapidly decreases during the first month, and then essentially disappears.

We postulate that the presence of estrogen receptors during the development of the cerebral cortex may play a role in establishing the hemispheric asymmetrical patterns, for the following reason. In previous ex-

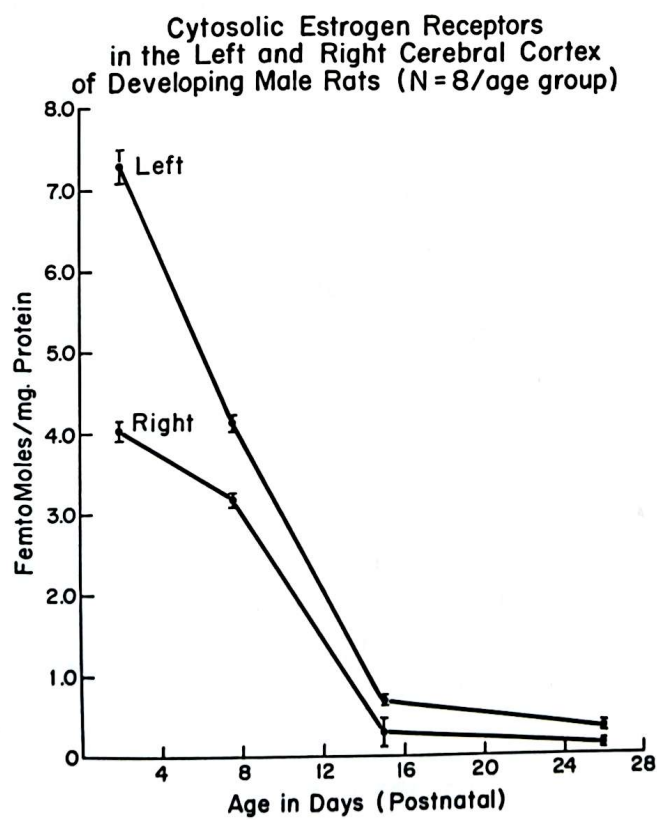


FIGURE 7. Cytosolic estrogen receptors in left and right cerebral cortex of developing male rats ($N = 8/\text{age group}$).

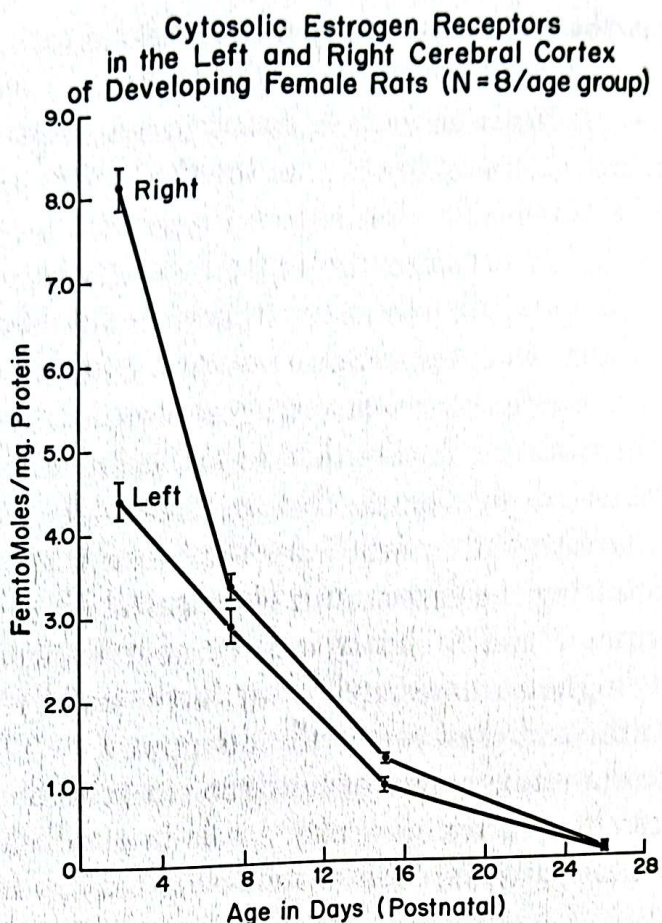


FIGURE 8. Cytosolic estrogen receptors in left and right cerebral cortex of developing female rats ($N = 8/\text{age group}$).

periments, we demonstrated that exogenous estrogen administered to adult female rats decreased cortical thickness (25). If estrogen acts on early postnatal cortical cells in much the way it does on adult cortical cells, then one might hypothesize that the hemisphere with the greater number of estrogen receptors will have the thinner cortex. This is precisely what we have found. Estrogen receptors are found in different concentrations in the right and left cortices in male and female rats.

In order to study the estrogen receptors in this first study, total samples of dorsal cortex were excised from each hemisphere. The kind of large tissue sampling used did not indicate a precise cortical location of the receptors. More recent preliminary results (26) have shown that the female right somatosensory cortex has a higher concentration of estrogen receptors immediately after birth than the right frontal or occipital regions.

Having found these sexual dimorphic patterns of asymmetry in the cerebral cortex, we then wished to learn whether similar dimorphism existed in several subcortical areas which have rich fiber connections with

the cortex. If some structures do not possess asymmetry, it might offer a clue to the reasons for asymmetry in the areas where it does exist.

On the same tissues we had used for collecting the cortical-thickness development and aging data reported in 1975 and 1985, we turned to several subcortical regions and measured the thickness of the hippocampus, a region dealing with recent-memory processing, sexual behavior, and spatial mapping; the area of the amygdaloid nucleus, a region concerned with emotional behavior and reward systems; and the area of the corpus striatum, a structure which modifies movement as well as pain thresholds (see Figure 2. The corpus striatum is not shown in this posterior section. It is found more anteriorly beneath a frontal cortical section. Fig. 1)

On drawings from projected microslide images, we measured the hippocampal thickness (see Figure 2) on a total of about 100 male rats ranging from 6 to 900 days of age. Figure 9 illustrates the percentage of right-left differences in hippocampal thickness in rats up to 400 days of age (27). It is evident from this figure that in the male rat, the right hippocampus is significantly thicker than the left for the first 21 days of life. During the period of early sexual maturity, the significant differences between the right and left hippocampi disappear, only to appear again at the age of 185 days. Measurements on the 904-day-old male rat hippocampus indicate that the significant asymmetrical pattern is lost at this time. That is, the right hippocampus is still thicker than the left, but the

The Percent Difference Between Right and Left Dorsal Hippocampal Thickness in Long-Evans Rats

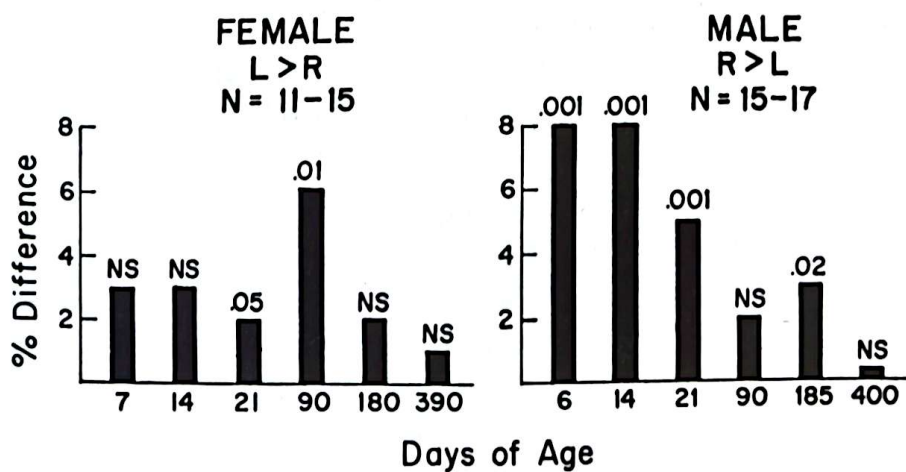


FIGURE 9. Percent difference between right and left dorsal hippocampal thickness in Long-Evans male and female rats.

differences are not statistically significant. The Long-Evans rat becomes sexually mature before 60 days of age, and it is about this time that the asymmetrical pattern in the male hippocampus first becomes reduced.

The reason why the right hippocampus is larger than the left, especially at the beginning of the animal's life, is not known. But we were curious to learn whether testosterone played a role. For this experiment, the testes were removed at birth, and the brains were studied 90 days later. The results showed that the left hippocampus was significantly larger than the right, but only by 2% ($p < 0.05$) in the 18 male brains measured. At 90 days of age in the intact animals, the right hippocampus was thicker than the left by 2%, but the difference was not significant. Perhaps 90 days is too long to wait to take the measurements. In the future, it will be necessary to remove the testes at birth and then examine the right-left differences in the hippocampus during the early period of the animal's life.

Because the right hippocampus and the right cerebral cortex of the intact male are each thicker than the left, the visual-spatial roles of the two areas may be closely related. O'Keefe and Nadel (28) have proposed an internal cognitive map theory correlated with hippocampal function. They hypothesized that the cells in the hippocampus encode the animal's specific position in its environment. We have already addressed the role of the right posterior cortex in visual-spatial acuity. The evidence points in the direction of a possible close functional relationship between these two forebrain structures during the early stages of the animal's life.

In the female, the right-left differences in hippocampal thickness were the opposite of those found in the male. The left hippocampus was found to be thicker than the right in female animals from 7 to 390 days of age with 11 to 15 animals per age group (Figure 9), but only at 21 and 90 days of age were the differences statistically significant. Thus, marked asymmetrical differences became more obvious at the time of female sexual maturity, again suggesting the possibility of a relationship between sex hormones and asymmetry.

In order to test that possibility, hippocampal thickness was measured in female rats that were ovariectomized at day 1 and autopsied at 90 days of age and was compared with the hippocampal thickness of 90-day female controls. In both groups the left hippocampus was thicker than the right. These data showed no evidence of a gonadal hormonal influence on the female asymmetry. Thus, removal of the ovaries at birth appears to have less of an effect on the hippocampus than on the cerebral cortex. It may be that the hippocampus, a phylogenetically older structure, does not alter its structure as easily as does the more recent cerebral cortex.

The marked asymmetry seen in the young animals declines with age in both the male hippocampus and the male cortex. The female also experiences a reduction in asymmetry in the hippocampus as she ages: from 90 days to 390 days her hippocampus showed a decreasing left-dominant pattern.

The next structure which was examined for right-left differences was the amygdaloid (almond-shaped) nucleus (Figure 2). This nucleus consists of numerous individual nuclei and is associated with many different functions, including expression of rage, aggression, sexuality, and reward mechanisms.

The amygdaloid nucleus is unique so far in our measurements in that it shows both strain differences and right-left pattern differences. In the Long-Evans male rats, the amygdaloid nucleus was symmetrical (29). The left-versus-right percent differences were nonsignificant in every case from the age of 6 days to 400 days. This was the first forebrain structure in the male Long-Evans rat where we did not find asymmetry. However, the male amygdaloid nucleus in the S₁ strain of rats showed significant asymmetry, with the right greater than the left (8 to 9%, $p < 0.01$). Replication measurements verified this strain difference. None of the scientists with whom we have discussed these findings have yet suggested an interpretation.

In summary, our experiments revealed asymmetry patterns in the male and female forebrain structures measured, with but one exception: the amygdaloid nucleus. In general the male showed marked right dominance early in life and the female displayed either no significant dominance or a left-dominant pattern.

Forebrain asymmetry in certain brain regions is not unique to the rat. There are isolated reports on brain asymmetry as one moves up the phylogenetic scale. A group of nerve cells called the habenula (Latin, "rein," the strap of a bridle) in the posterior thalamus of amphibians and fish has been noted to be asymmetrical (30). In the cat, fissural pattern differences exist in the hemispheres (31). In the great apes, the end of the Sylvian fissure on the surface of the hemispheres is longer on the left side than on the right (32). Such asymmetry is also evident in the human brain. The occipital lobe protrudes farther posteriorly on the left than on the right, and the posterior horn of the lateral ventricle is longer on the left side than on the right.

Though there is evidence of asymmetry throughout the animal kingdom, we do not fully understand why the brain developed this way. We can shed some light on the subject by examining language areas in the human brain. Motor speech activities are primarily governed by the left

hemisphere in most right handed people. Hence, the left hemisphere is referred to as the dominant hemisphere for speech. The area in the right hemisphere comparable to the one on the left for motor speech, processes the emotional component of speech. Thus, this example suggests that lateralization allows for specialization of function. Both hemispheres deal with speech but each contributes a separate component to the process. Undoubtedly, as we learn more about each hemisphere individually, we will understand more about how integration of their outputs occur.

Our work has shown that the sex hormones can alter the asymmetry patterns in male and female rat brains. We may better understand the significance of brain asymmetry once we have had a chance to study behavioral deficits after the asymmetry patterns have been reversed for a considerable period of time. We will also need to study the effects of asymmetry on other body systems. We used to believe that there were paired structures in the brain to serve as a safety factor in case of the destruction of one part, but now it appears that the paired brain structures combine two different functions that are eventually integrated for more elaborate processing. Possible asymmetry exists to allow one side to function more efficiently independent of the other prior to integration. It is to the advantage of the animal—whether rat or human—for one side to develop rapidly, namely, the side governing the function most important for the animal's survival. For example, we could suppose that the visual-spatial role of the right cortex is essential for maximum survival of the male; then we could explain why nerve cells are present in the right cortex of the male than in the left. Similarly, since left-hemisphere dominance shows up early in the female, we could hypothesize that the early development of language skills contributes to her survival.

Cortical Structure and Altered Sex Hormone Levels

In light of what we knew of the impact of sex hormones on forebrain asymmetry, we were encouraged to consider their significance in the development of the cortex as a whole, including its cells and synapses. Knowledge of the relationship between gonadal hormones and the cerebral cortex is not as extensive as knowledge of the relationship between these hormones and other regions of the brain. The hypothalamus and preoptic area of the brain are important in mediating the effects of gonadal hormones on their target organs. Gonadal steroids influence sexual differentiation of the hypothalamus at a critical period in the development

of the animal. In rats, the first 5 days after birth represent the critical time in the development of the hypothalamus. During this period, the pattern of the male hypothalamus is established by the action of testosterone that has been converted to estrogen. If the male is castrated at birth, the hypothalamus retains the female pattern. In adult female rats, the hypothalamus regulates the secretion of gonadotropic hormones in a cyclic fashion, resulting in the rhythm of ovulation and formation of corpora lutea in the ovary. Knowledge about the function of the hypothalamus in males and females has accumulated steadily over the years and is fairly well accepted. However, few investigators have examined the effects of gonadal steroids on the physiology and anatomy of the cerebral hemispheres.

We first became interested in the interaction of the ovarian hormones with the cerebral cortex when we were tabulating the results from experiments dealing with pregnant rats in the late 1960s and early 1970s. We had not studied the brains of female rats prior to this time, because we wished to avoid the additional variables that might be introduced by the estrus cycle. Actually, we learned about the influence of the sex hormones on the female cortex quite by accident. We were interested in the pups from parents living in different types of environments. In order to carry out the study, it was essential to mate males and females that had lived in either enriched or impoverished conditions. A puzzle emerged from the experiments involving the female rats living in enriched or impoverished conditions during pregnancy: we learned that the usual cortical thickness differences between the enriched and impoverished animals were not present. Upon close observation of the data, it was apparent that during pregnancy the brain of the impoverished female had increased to dimensions equal to those of the enriched animal. With this finding, we then began to study more closely the relationship between the ovarian hormones and the development of the cerebral cortex.

Information that is slowly accumulating shows that estrogen plays a role in shaping the structure of the cerebral cortex. Estrogen has been reported to alter the incorporation of amino acids into the neurons of female animals. Litteria and Thorner report that chronic injections of estradiol in adult ovariectomized rats depressed incorporation of tritiated amino acid into proteins in the cerebral cortex (33), as did injections given at 24 hours and 72 hours after birth (34). However, others find the opposite effect: single injections of estrogen into ovariectomized animals increased incorporation of tritiated amino acids into proteins and increased the protein content of the cerebral cortex (35, 36). It is possible that the dosage and time factor could be responsible for these opposing

results. Nevertheless, estrogen clearly plays a role in cortical metabolism, though it may not be as pronounced as its role in other areas.

In the same year that we published our paper on the effect of pregnancy on the rat cerebral cortex (37), Presl et al. (38) found estrogen receptors in the cerebral cortex of young rats from 5 to 50 days of age. A perinuclear localization of the receptors was identified. In the same year, Kato et al. (39) found that at birth the concentration of estrogen receptors in the anterior pituitary was about 3 times as great as in the cerebral cortex; at day 45 the estradiol concentration was lower in the cerebral cortex than in any other part of the brain. These results indicate that estrogen receptors do exist in the cortex, but there are fewer of them there than in other regions.

Although estrogen receptors disappear from the cortex after the first postnatal month, we found that the brains of 116-day-old pregnant rats had thicker cortices than those of nonpregnant animals. Was it possible that estrogen could influence cortical structure in older animals even in the absence of receptors? A series of experiments were planned to learn whether the removal of the ovaries at various times during the life of the female altered cortical structure. The times chosen were (1) at birth; (2) during the early sexual reproduction period, at 90 days of age; and (3) later in life, at 300 days of age.

For the study of the impact of ovariectomy at birth (40), two females from each litter were ovariectomized and two served as sham controls, their ovaries having been exposed surgically but not removed to expose both sets of rats to similar stress. For each of the experiments, the females lived in standard colony cages for a 90-day period following the surgery—the first 21 days with their mothers, and then three ovariectomized females (or three controls) per cage.

The ovariectomized rats were found to have a thicker cortex in certain areas than the control, intact rats. The area which showed the most consistent increase was area 4 (see Figure 1). In the first experiment, this area was thicker in the ovariectomized rats than in the controls by 3% ($p < 0.05$), and in the replication experiment by 6% ($p < 0.05$) (40). That area 4 should show such responses to ovarian hormonal deprivation was unpredicted, because this area is reportedly associated with motor function. Admittedly, it is known that on the average females do not possess as great a muscular strength as males. These results suggest that a dampening of the motor cortex in response to ovarian hormones may diminish the output from the cortex to the nerves innervating the skeletal muscles, thus reducing the strength of contraction.

In order to determine whether changes had occurred in the neuronal

dimensions of area 4 as well as in its cortical thickness as a result of the reduction in ovarian hormones, we measured the area of the neuronal perikarya and nuclei. For these measurements, the tissues were photographed and the cells were magnified ($1500\times$) so the outlines could be accurately traced with a planimeter. We learned that the perikarya and nuclei were larger in the ovariectomized rat than in the control by 12% ($p < 0.01$) and 6% ($p < 0.01$), respectively. In addition, the control rats had a 17% greater cell density ($p < 0.001$), signifying fewer dendrites, than the ovariectomized rats. Here was the first evidence that removal of the ovaries at birth changed not only the thickness of the cortex in area 4 three months later, but the dimensions of the nerve cells as well.

In an attempt to determine more specifically whether these changes were occurring in the upper or lower cortical layers, the cortex was divided into equal halves between layers II and VI. We learned that the lower half changed dimensions significantly more than the upper half. Sheridan (41) later demonstrated that estrogen receptors were predominantly located in layers V and VI in the 2-day-old rat. Despite differences in methodologies and in the ages of the rats used in the two experiments, both our findings and Sheridan's indicated that estrogen was affecting the neurons in the lower layers of the cerebral cortex.

The fact that ovariectomy at birth changed the structure of the cortex attests to the significance of the ovarian hormones. But perhaps their significance is not lasting; perhaps they play their part in cortical formation during the animal's early growth. To consider this possibility we had to study rats whose reproductive organs were sexually mature (as they usually are by 60 days of age) before we removed the ovaries. For our experiment, female Long-Evans rats were ovariectomized at 90 days of age and lived without their relatively high concentrations of ovarian hormones for 90 days, or 3 months, before cortical thickness measurements were made. In these rats, which were 180 days old when we made the measurements, half of the measures showed the ovariectomized rats' cortices to be thicker than the controls'. But there were no significant differences.

In the third experiment of this type, ovaries were removed from rats at 300 days of age in an attempt to learn the effect of diminished ovarian hormones on the cortex of female animals one-third of the way through their normal lifespan. There were no significant differences between these older ovariectomized rats and their controls, but in 13.5 (1 was equal) out of 18 areas measured in the right and left cortices, the ovariectomized animals had thicker cortices than did their controls.

The above results indicate that the cerebral cortex is most responsive to ovarian hormonal deprivation in the early stages of postnatal life, but that some influence is evident in the older animals, where slight increases in cortical dimensions are noted after ovarioectomy. If the older animals had lived for longer periods of time without their ovarian hormones, one wonders whether the differences would have become more marked.

Since ovarian hormones are used as contraceptives, we have good reason for wanting to understand the effects of these hormones on cerebral-cortical structures and functions. The rat results provide evidence that ovarian hormones can alter the outer layers of the brain. More recent experiments indicate that ovarioectomy influences cortical cells at the synapse, the specialized apparatus for transmitting an impulse from one nerve cell (the presynaptic nerve cell) to another (the postsynaptic cell) across an intercellular space called the synaptic cleft. A synapse is composed of synaptic vesicles; presynaptic electron-dense projections; the presynaptic membrane; the synaptic cleft; and the postsynaptic membrane and its accompanying electron-dense material, which makes up the "postsynaptic density" (see Figure 10). When an organism receives input to one of its senses, the information is recorded as a nerve impulse, which is conducted to the synapse. Since the synapse does not involve a physical contact between neurons, a chemical carrier—a neurotransmitter—is generally required to bridge the synaptic cleft. Synaptic vesicles in the presynaptic terminal store quantities of the transmitter. Energy for the release of the transmitter is generated in the mitochondria of the terminal. Binding of the neurotransmitter to receptors of the postsynap-

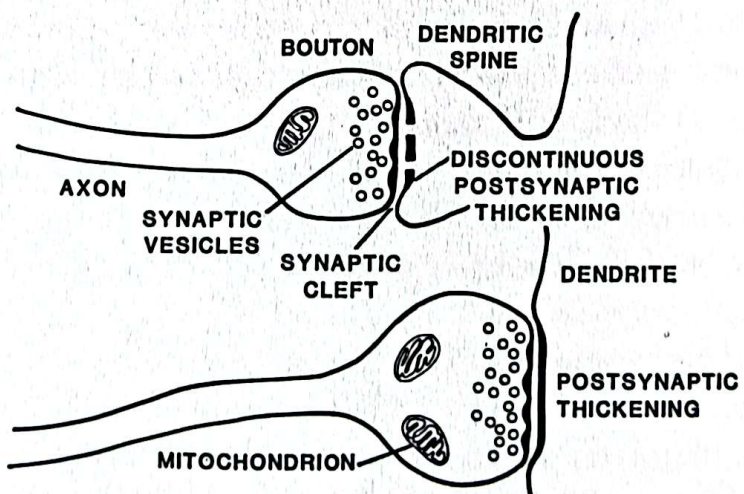


FIGURE 10. Synapse, (below) on a dendrite shaft and (above) on a spine.
Axons are drawn to show the synaptic vesicles and both the presynaptic and the postsynaptic thickenings. Upper synapse has discontinuous postsynaptic thickening.

tic membrane produces changes in this structure that enable the impulse to continue into the adjacent neuron.

Pfenninger and Rees (42) demonstrated that during the formation of the synapse, the postsynaptic density is the first of all the synaptic membrane specializations to develop. With this finding, it was reasonable to consider the presence of an altered postsynaptic density to be a qualitative indicator of a possibly altered synapse and a criterion for quantification of its dimensions. Many investigators have measured the length of the postsynaptic density as representative of the length of the synapse with the assumption that it affects the overall efficacy of the synapse. More recent evidence has supported the hypothesis that discontinuities in the postsynaptic density may characterize more mature synapses.

Peters and Kaiserman-Abramof (43) took serial cross sections of synapses and observed cases in which a single synapse exhibited both a postsynaptic density with gaps and one with no gaps. One might consider that split synapses were simply immature, incompletely formed synapses. However, since the gaps have been observed exclusively in the larger, and therefore probably more mature, synapses, this possibility seemed unlikely to those of us studying postsynaptic densities. Furthermore, evidence by Dyson and Jones showed that the number of synapses with discontinuities in the postsynaptic density increased with maturation (44). Greenough et al. investigated the frequency of split synapses in the cortices of rats reared in enriched and nonenriched environments (45). These authors termed the gaps "subsynaptic plate perforations" and found that the relative frequency of the perforations increased dramatically in rats between 10 and 60 days of age. They also observed that rats reared in more complex environments had postsynaptic thickenings with a significantly greater number of perforations than rats raised in isolated conditions. (The synapses in the occipital cortex were measured to obtain this finding.)

That the discontinuous synapses increased with maturation and with stimulating environments suggested that this type of synapse did represent a more highly developed or more mature structure. Also, as previously mentioned, ovariectomy increased cortical thickness as well as the dimensions of the neuronal soma and nucleus. Therefore, in our experiments dealing with the effects of ovariectomy on cortical structural changes, it appeared most reasonable to choose discontinuous synapses for our next measurement.

In our attempt to learn whether the increased cortical dimensions after ovariectomy represented a more mature cortex, we not only quantified the discontinuous synapses, but we also analyzed the curvature of

the synapses as a function of maturation. Dyson and Jones distinguished among positively curved, negatively curved, and flat junctions between pre- and postsynaptic bindings (44). Positively curved synapses were defined as those in which the postsynaptic density curved into the presynaptic terminal; negatively curved synapses were those in which the postsynaptic density curved toward the postsynaptic ending. These researchers found an increase in the frequency of flat junctions with age. They also discovered that a larger proportion of the synaptic curvatures were negative in young animals. With these facts, along with evidence that anesthesia produced a preponderance of negatively curved synapses, Jones and Dyson proposed that negatively curved synapses indicated non-function, and positively curved synapses, function. They also suggested that the flattening that occurred in more mature synapses represented an adaptation through which a "finer control of the neural network system" was accomplished.

In order to obtain synapses for our study, 7 littermate pairs of 1-day-old rats were separated. Under cryogenic (cold) anesthesia, one was ovariectomized and the other served as a sham-operated control. At 90 days of age, the animals were coded so the investigator was unaware of the treatment each rat had received. On photoelectron-micrographs, synapses were analyzed in layer II of the right medial occipital cortex, or area 18 (46). The total magnification was 34,000 \times . Only axospinous synapses were analyzed. Both continuous postsynaptic thickenings and discontinuous synapses on dendritic spines were quantified. For the frequency of occurrence of the split synapses, chi-square tests were used; these revealed that the ovariectomized animals had significantly more split synapses (20%, $p < 0.05$) than the sham-operated rats. The discontinuous postsynaptic densities occurred in both groups, but more frequently in the ovariectomized one.

A significant difference was observed in the frequency of occurrence of positively and negatively curved synapses between the sham-operated and ovariectomized animals. The ovariectomized group had significantly fewer negatively curved synapses (63%, $p < 0.05$) and more positively curved synapses (19%, $p < 0.05$) than the sham-operated control. Accepting Dyson and Jones's hypothesis concerning the meaning of synaptic curvature, we can conclude that the ovariectomized animals had more functioning synapses (positively curved) and fewer nonfunctioning synapses (negatively curved) than their sham-operated littermates. Though no one has established any definitive functional implications of the difference between discontinuous, positively-curved synapses and negatively-curved synapses, these results on the effects of ovariectomy on cerebral

cortical structure all support the available information indicating that a cortex from an ovariectomized rat is in a more mature state of development than one from an intact female.

Hormone Replacement

Ovarian hormone replacement seemed a logical next step to us in continuing our search for the relationship between cortical development and ovarian hormones. First, female rats that had been ovariectomized at birth were given estrogen in the form of ethinylestradiol, starting at day 40 and continuing until day 90 (dosage: $1\mu\text{g}/\text{kg}$) (47). The results obtained from cortical thickness measurements indicated that the ethinylestradiol-treated rats had a significantly thinner cortex than the oil-injected sham ovariectomized controls. The measured cortical areas with significant differences, ranging from 4 to 6%, were areas 4, 3, 18, and 17. Again, area 4 showed a response to ovarian hormones, as it had with ovariectomy, but, as might be predicted, in the opposite direction. In a preliminary study, the same dose of ethinylestradiol from day 69 to day 90 had produced no significant differences in cortical thickness. Evidently either 21 days is not a sufficiently long period for cortical change to occur, or the age at which hormone replacement begins plays a crucial role, or possibly both age and length of exposure are important.

Progesterone is known to have an antiandrogenicity effect, so one might hypothesize that progesterone might increase cortical dimensions. To study the effect of progesterone on the cortex, rats ovariectomized on day 1 were treated with progesterone from day 40 to day 90 (dosage: $2\text{ mg}/\text{kg}$) (47). In comparing the cortical thickness of the ovariectomized rats receiving progesterone with the sham-ovariectomized rats, we found that the cortex was thicker as a consequence of progesterone treatment. In general, progesterone had an effect opposite to that of ethinylestradiol. The cortical areas which showed significant increases (ranging from 3 to 5%) were areas 10M, 10L, 4, 3, 2, and 18. In the previous studies with ovariectomized rats without hormonal supplements area 4 showed a consistent response to altered ovarian hormonal levels. With progesterone, this area increased by 3% ($p < 0.05$) compared with the intact controls. According to a one-way analysis of variance test, area 4 demonstrated a significant level of difference from the control in each of the experiments with changed concentrations of ovarian hormones, whether ovariectomy alone or ovariectomy with ethinylestradiol or progesterone.

On the tissues from the ovariectomized rats that we had used for cortical thickness studies, we also measured both the hippocampus and the pyriform cortex (see Figure 2). There was no significant difference

in hippocampal thickness between ovariectomized females *not* given replacement hormones and the control. However, the hippocampus was significantly thinner, by 3% ($p < 0.01$), in the ovariectomized rats receiving ethinylestradiol than in the controls. As occurred in the cortex, the estrogen compound reduced the thickness of the hippocampus; but progesterone had no significant measured effect on this part of the forebrain. The ovariectomized females and the ethinylestradiol-treated females showed no significant differences from the controls in the pyriform cortex. The pyriform cortex was not measured in the progesterone-treated rats.

In summary, these studies clearly indicate that the structure of the rat cerebral cortex and hippocampus can be significantly altered by various levels of female sex steroid hormones under certain conditions. Therefore, we have reason to believe that the normal course of cerebral development, and some of the differences between the male and female brains, are shaped by the sex hormones.

Growth and Aging Patterns in Subcortical Structures

Not all forebrain structures develop and age at the same rate. This section will deal with regions of the forebrain that differ from each other in development and aging patterns—including the hippocampus, the entorhinal cortex, the amygdala, and the corpus striatum (see Figure 2); corpus striatum, however, is not shown, as mentioned previously in Chapter 2 (48). Attempts to understand these normal distinctive aging styles in the rat forebrain allow us insights into regions that may be involved in abnormal aging. In addition, some regions of the forebrain are more susceptible to environmental influences than others, as will be seen in Chapter 4. By examining areas with different aging patterns and different responses to the environment, we may find some clues to the interaction of aging and environmental stimulation and, in turn, shed light on possible neural patterns of disease processes.

The occipital cortex, located in the back or posterior region of the cerebral cortex (Figure 1), processes visual information; damage to this area results in partial or complete blindness. This region is offered as an example of the phylogenetically more recent six-layered neocerebral cortex.

The entorhinal cortex (see Figure 2) is also known as the *transitional cortex*, because this five-layered paleocortex arises in the phylogenetic scheme between the archicortex and the neocortex. The entorhinal cor-

tex constitutes an important site of convergence for information from many sensory systems and for pathways concerned with memory processing.

The hippocampus, named for its physical resemblance to a sea horse, is found deep below the cerebral cortex and represents the three-layered archicortex (see Figure 2). A single, precise role of the hippocampus has not been pinpointed, but present information suggests that it is involved in such functions as learning and memory processing, emotional reactions, sexual behavior, and spatial mapping.

Both the hippocampus and the entorhinal cortex are of particular interest at present in our society, now that more people are living to old age. These are sites of abnormal formations, such as neurofibrillary tangles and plaques, which are characteristic (and indeed are the only certain diagnostic features) of the dreaded Alzheimer's disease. There is apparently no adequate animal model for this disease. But by observing the developing and aging patterns of these regions in an animal (because we cannot make such precise measurements in human brains), one can gain an insight into possible areas of specific variations.

The amygdaloid nucleus (see Figure 2), consisting of a conglomerate of nuclei, is so named because it appears almond-shaped in the human brain. This nucleus is located within the temporal lobe and is of interest because it plays a part in regulating such functions as aggression, sensory integration, responsivity to reward, and sexuality.

The corpus striatum is commonly called the basal ganglia, because it consists of masses of nerve cell bodies found in the base of the cerebral hemispheres. Its growth and development deserve attention because of its role in skeletal muscle control (it programs the inception and termination of motor activity) and because of its modification of pain. In addition, it has high quantities of various neurotransmitters compared with other areas of the forebrain, such as the cerebral cortex, suggesting a continuously active functional role.

In order to determine the growth and aging curves of these forebrain structures, we measured the thickness of the occipital and entorhinal cortices, as well as the hippocampus, on transverse histological sections of brains from male Long-Evans rats. The rats lived as littermates with their mothers until 22 days of age in small colony cages ($34 \times 20 \times 20$ cm) and then were housed in groups of three or four in similar colony cages. Seven to 16 animals were studied in each of the following age groups: 26, 41, 108, 650, and 904 days of age. The measurements of the area of the amygdaloid nucleus were made on 9 to 15 rats per age group at 6, 26, 55, 90, 185, and 400 days of age. For the measurements of the area

of the corpus striatum, 7 to 15 animals per age group were used at 6, 10, 14, 20, 26, 41, 55, 77, 90, and 108 days of age.

The results from the occipital and entorhinal cortices and from the hippocampus are presented in Figure 11. Here it is clearly demonstrated that the hippocampus continues to grow slowly from 26 to 904 days of age. The increase is gradual throughout this time, providing an overall increase of 10%. Both the entorhinal and occipital cortices decrease with aging, however, but their patterns of aging differ. Sometime between 26 and 41 days, the occipital cortex begins to decrease steadily until 650 days of age, by approximately 11%. Between 650 and 904 days of age, there is a steep decline of another 11%, giving a total decrease between 26 and 904 days of about 21%.

Though the entorhinal cortex also decreases over time, its most dramatic decrease, by 8%, occurs between 26 and 41 days of age. From 41 to 904 days there is only another 5% reduction in thickness. While the occipital cortex is changing so dramatically between 650 and 904 days of age, the entorhinal cortex is not showing any significant change. Thus, it is clear from Figure 11 that these three regions of cortex follow quite different patterns during aging (when the animals live three to a cage in a controlled laboratory environment).

The growth and aging pattern of the amygdaloid nucleus offers yet

Thickness of entorhinal cortex, occipital cortex and hippocampus with aging

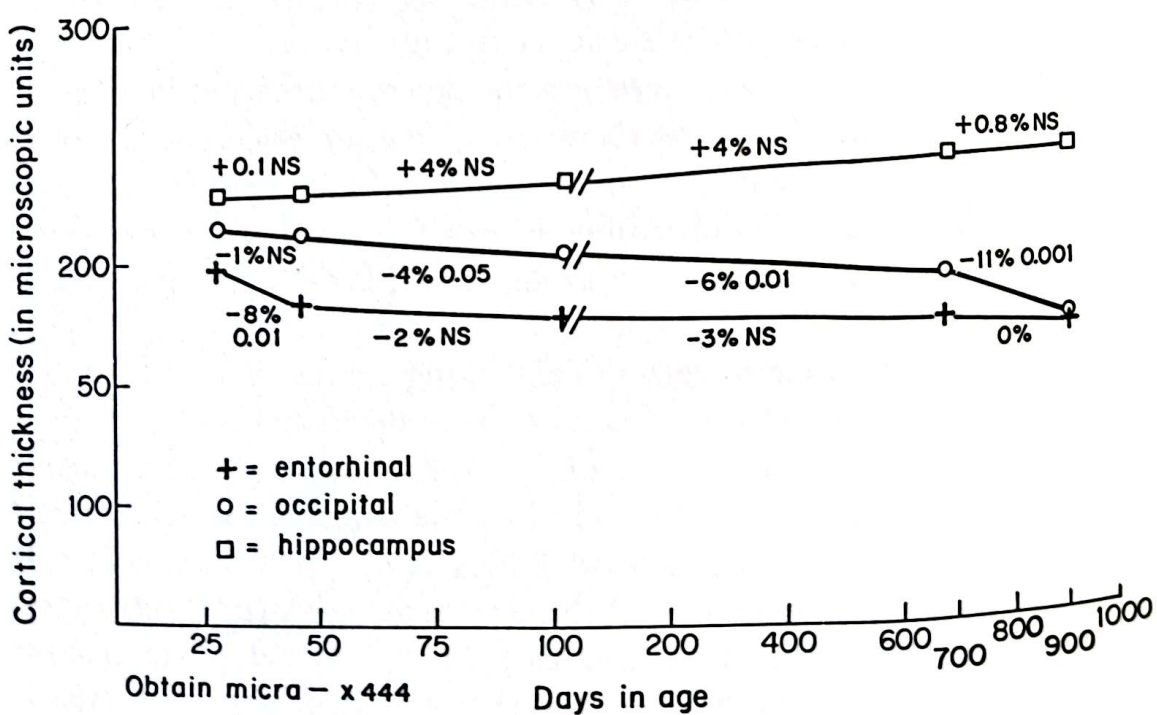


FIGURE 11. Thickness of entorhinal cortex, occipital cortex, and hippocampus with aging.

a different picture. After reaching a growth peak between 26 and 55 days of age, the amygdaloid nucleus decreases until sometime between 90 and 185 days of age, when it once again continues to increase. After 185 days it begins to diminish again until 400 days, the oldest age for which we measured this structure. Its aging pattern appears independent of those of the other regions measured.

The corpus striatum was not measured for as extensive a period as were the other forebrain regions. Yet, for the short period for which its dimensions were recorded, it also follows an aging pattern different from the surrounding regions. The corpus striatum increases greatly from 6 to 20 days of age, and from then on the increase is more gradual until 77 days of age, when it appears to peak. Though more points at later dates are needed to confirm this apparent peak, the development and aging pattern of this region is different from that of the others.

Because the various regions of the forebrain age at different rates, it is not precise enough to speak about an aging brain without defining the region involved. Whereas the hippocampus is steadily increasing in dimensions throughout a lifetime, the entorhinal cortex and occipital cortex display decreasing trends. These latter, more recently evolved cortices of five and six layers, are possibly more subject to the consequences of aging as they are to the environmental conditions to be mentioned in Chapter 4, than is the older three-layered cortex. The amygdaloid nucleus offers yet another aging pattern quite different from the three cortical regions. We have yet to identify the factors that influence these patterns.

3

DEVELOPMENT AND AGING OF CORTICAL NERVE CELLS AND GLIAL CELLS

It is well known among life scientists that not only are brain nerve cells formed at rapid rates during embryogenesis, but large numbers die during this time as well. Some investigators have stated that as many as 50 to 65% of the newly formed nerve cells are lost before birth. Most lay people believe that they are losing nerve cells in the brain as they age. Postnatal brain cell loss is a real concern. In fact, a very common question asked by a lay audience is, How many cells do I kill everytime I take my martini? Not only can we not answer this question with accuracy, but until recently we could not answer questions about brain cell loss with "normal" aging. Only by examining the brains of animals in controlled environments could we obtain a reasonable answer to this last query. The usual factors, such as diet, genetic background, air quality, and the general environment, all have to be taken into consideration. In order to understand how the environment alters brain cells, it is first essential to obtain measurements of cell populations in standard laboratory conditions.

We learned that nerve cell counts in rats are fairly unstable during the first 108 days of postnatal life. It may have been that neuronal death was continuing after birth for a short period or that the cells were moving to different locations. But after 108 days and until 904 days of age, the nerve cell number per unit area was remarkably stable. In other words, the aging rats were not losing a significant number of neurons from the cerebral cortex. Our results indicate that if an animal is living

in a healthy environment throughout its lifetime, the number of nondividing cells exhibits unusual consistency after early adulthood. The counts of the glial cells—the structural and metabolic support cells of the nerve cells—followed patterns very similar to those of the neurons.

In order to create a field large enough for the cells to be identified and counted by several investigators, we took photomicrographs of 1 mm^2 areas of the medial occipital cortex (area 18; see Figure 1), a visual association area (1). Each area was enlarged to create a field of about 1 m^2 (a single field is reproduced in Figure 12). Area 18 was chosen for our detailed cellular analysis because it appeared to be the most responsive to our varied external environmental conditions. It was the first area to demonstrate a structural change when the rats had been enriched for only 4 days.

By photographing four adjacent, overlapping columns of the cortex from the surface down to the underlying white matter (at a magnification of $640\times$), we gave ourselves an adequate sample of tissue for cell counts. A montage was created from the individual pictures, and a clear sheet of Plexiglas was placed over it (Figure 12). In this manner, the cell types could be identified and marked with a colored wax pencil directly on the

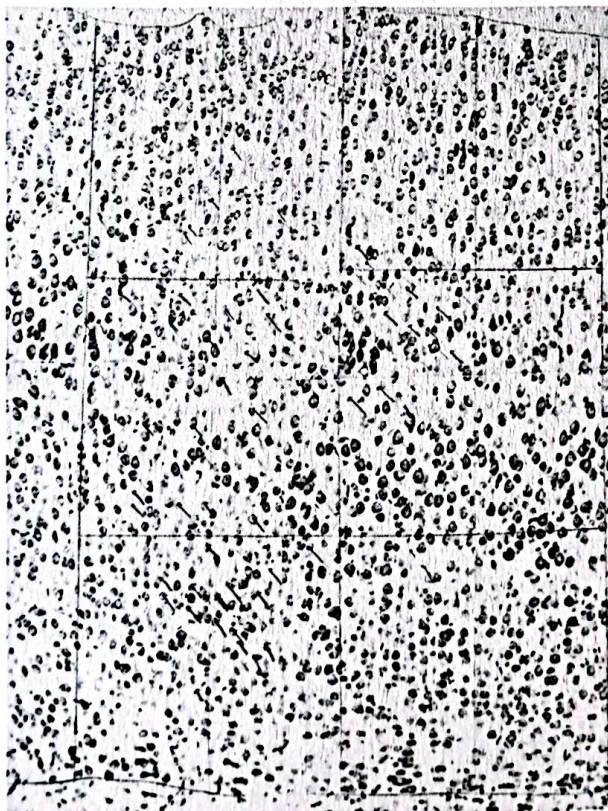


FIGURE 12. Montage of photographs illustrating neurons and glial cells in 1 mm^2 of cerebral cortex.

overlying Plexiglas. After all the cells in one sample were counted (there were approximately 600 to 725 nerve cells per square millimeter, depending upon the age of the animals), we replaced the marked piece of Plexiglas with a fresh one and made another cell count. The two completed sheets were then superimposed in order to declare a consensus on the counts.

The well-trained human eye connected to the discriminating brain has to decipher many variables in making a decision whether a cell should be counted or not. Not only are nerve cells of different shapes and sizes, but the several types of glial cells also display a variety of densities and sizes. The amount of overlap among all cell types is another consideration. When counting the neurons and glial cells, we do not wish to include the blood vessel lining cells, so they must be identified and excluded. No automatic scanner is yet sensitive enough to substitute for such judgments, even though many types of scanners are on the market.

Cells were counted on 6 micron ($6 \mu\text{m}$) sections beginning with 26-day-old male rats. With this thickness, we found we could not count the cortical cells prior to 26 days of age, because the cells had not grown enough branches to distinguish one cell from another (2). The density of cells per unit area was too great to allow us to see well-defined cell boundaries. After making counts on the 26-day-old animals' brains, we added additional age groups: 41, 108, and 650 days. All tissues were from our original longitudinal study with the Long-Evans rats.

At first glance, the results, shown in Figure 13, are consistent and revealing in one sense and very perplexing in another. Between 26 and 41 days of age, there was actually an increase in the number of neurons

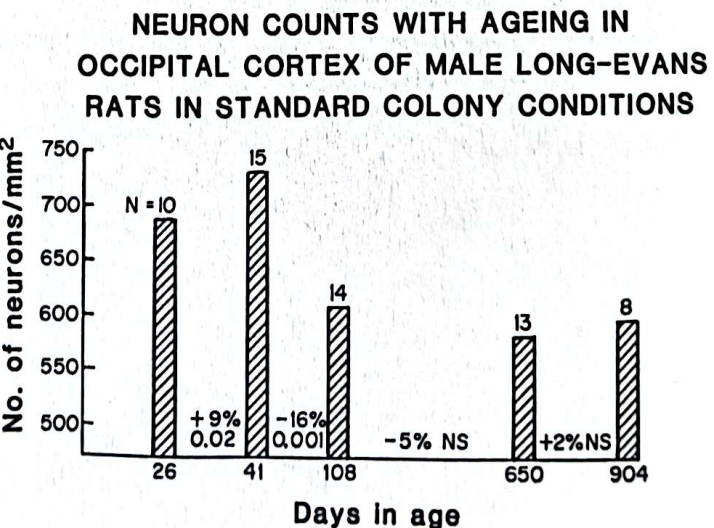


FIGURE 13. Neuron counts with aging in occipital cortex of male Long-Evans rats in standard colony conditions.

per unit area by 9% ($p < 0.02$). This increase can be explained by a decrease in total mass of cortex without a decrease in number of cells, which would reduce the connections between the cells and provide more cells to be counted per unit area. But between 41 and 108 days of age, a 16% ($p < 0.001$) decrease in neurons is seen. In this period of time, the cortex is also decreasing in thickness, and so this loss of cells could be a real one. The reason for these contradictory findings before 108 days of age is not absolutely clear.

If we had been comparing only the 26- and the 108-day counts, then a more satisfactory explanation could have been offered, since, as was pointed out, the greatest loss of neurons occurs during normal embryogenesis. A loss after birth could represent a continuation of the process that begins before birth, which could be explained as a tailoring of the nervous system to fit need and demand. Undoubtedly, as the animal is establishing a life pattern, the nervous system responds and molds its cell pattern and processes accordingly. Needless to say, more counts are necessary in the period between 26 and 108 days before any final conclusions can be made about neuron populations in the cerebral cortex during this postnatal period.

But between 108 days and 650 days of age, a more stable neuronal count is found with a nonsignificant loss of nerve cells of only 5%. This fact by itself is very encouraging to learn, that in a period of 542 days in the adult rat's life, no significant cortical neuron loss was measured in the occipital cortex. But still another remarkable finding was to emerge. The 650-day-old animals' cells were counted in 1977. In 1985, the counts on the 904-day-old animals' brains were completed (2, 3), and to our surprise, when all counts were tabulated, there was a nonsignificant difference between neuron counts at 650 and 904 days of age! The rat was not losing a significant number of neurons in the occipital cortex after early adulthood.

Table 1 shows a marked difference in the rate of neuron density changes between the upper layers and the lower layers of the occipital cortex between 26 and 650 days of age. For these counts, a grid overlying the photomicrographs was divided into two equal parts, upper and lower, between the beginning of layer II and the lower border of layer VI adjacent to the corpus callosum (see Figure 2 for a representation of the cortical layers). From 41 days to 108 days, the neurons in the lower layers decreased in density more markedly than those in the upper layers, a 24% decrease ($p < 0.001$) compared with a 9% decrease ($p < 0.01$). It is possible that the reduced physical activity experienced by the rats in their confined standard colony cages was influential in bringing about a

TABLE 1
Differences in Neuron Numbers in Upper and Lower Halves of the Rat Medial Occipital Cortex

AGE (DAYS)	NUMBER OF RATS	NUMBER NEURONS ($\bar{X} \pm S.D.$)	DIFFERENCE BETWEEN AGE GROUPS (%)	P
Upper half				
26	10	371 ± 27	(+) 5	NS
41	15	388 ± 33	(-) 9	0.01
108	14	352 ± 30	(-) 7	NS
650	13	327 ± 37		
Lower half				
26	10	309 ± 34		
41	15	338 ± 37	(+) 9	NS
108	14	258 ± 21	(-) 24	0.001
650	13	255 ± 28	(-) 1	NS

reduction in neuron numbers. (The lower layers of cells project out of the cortex to subcortical structures or to the spinal cord. If less motor activity is taking place, fewer cortical cells are needed to innervate the motor cells in the spinal cord.) For the other age comparisons, no significant differences were found in neuron density changes between upper and lower layers.

The cell count data clearly indicated that after 108 days of age, the male rats living in our standard colony conditions, receiving healthy, protein-rich diets, breathing relatively good laboratory air, and drinking tap water, did not lose a significant number of nerve cells. In other words, if the brain was given an adequate support system as it aged, the number of nerve cells in the cerebral cortex was quite stable. Other investigators have shown similar results in the somatosensory cortex: no significant loss of neurons during aging after adulthood has been reached (4).

We have already demonstrated that the cortical thickness decreases slowly with aging. However, according to this new knowledge, there is no significant loss of cortical neurons between 108 days and 904 days of age. It would seem, then, that the decrease in cortical thickness can be attributed to a loss of dendrites.

In addition to counting the numbers of neurons, we measured the area of the perikarya (cell bodies) and nuclei of neurons from area 18 of the occipital cortex in the 26-, 41-, 108-, and 650-day-old rats. We used a microfilm reader to make projections of the negatives of some of the pictures used for cell counts, and from each frame we made tracings on paper of three well-defined neurons (neurons in which we could clearly see a nucleus, nucleoli, and Nissl substance dispersed throughout the

perikaryon). The resulting total of 36 neurons per animal was measured with the aid of a planimeter. The total magnification from slide to drawing was 2000 \times .

Table 2 presents the data from the measured area of the perikarya and nuclei in area 18 from rats living in the standard colony conditions. Only the perikarya showed a significant decrease in area between postweaning and old age. As with the cell counts, the greatest decreases occurred in perikaryal area before 108 days of age, with a nonsignificant decrease during the next 542 days. It is important to stress that none of these animals was isolated; all were living at least two to three per small cage. Many other aging studies have dealt with animals housed alone and therefore in deprived sensory conditions; such studies have yielded smaller cell numbers and sizes than those reported here.

Glial cells are the second type of cell in the brain. Two kinds were quantified in our studies: oligodendrocytes (cells with few treelike processes) and astrocytes (star-shaped cells). The oligodendrocytes form myelin, the fatty sheath around nerve fibers which facilitates impulse conduction. Though oligodendrocytes are often found adjacent to the soma of the nerve cell or to capillaries, their function in these positions is not yet well understood.

Astrocytes are both structurally and functionally different from oligodendrocytes. Some of the processes of astrocytes terminate in expansions called *end feet*. These feet are found on blood vessels, on dendrites, or adjacent to the connective tissue membrane (pia mater) that surrounds the brain. One of their functions is to regulate metabolic and ionic conditions around blood vessels and dendrites; in connection with the covering around the brain, they serve a protective role.

TABLE 2

The Area of the Perikarya and Nuclei of Occipital-Cortical Neurons from 26 to 650 Days of Age

AGE (DAYS) ^a	AREA (IN. ²) $(\bar{X} \pm S.D.)^b$	DIFFERENCE (%)	p
<i>Perikarya</i>			
26	0.459 \pm 0.162		0.1
41	0.438 \pm 0.146	(-) 4.5	0.01
108	0.394 \pm 0.217	(-) 10.0	NS
650	0.385 \pm 0.235	(-) 2.3	
<i>Nuclei</i>			
26	0.210 \pm 0.145		NS
41	0.199 \pm 0.111	(-) 5.2	NS
108	0.189 \pm 0.104	(-) 5.0	NS
650	0.183 \pm 0.134	(-) 3.2	

^aN = 6 animals per age group with 36 cells per animal.
^bMagnification = 2000 \times .

The stain that we used for neuron counts (a composite of Luxol fast blue and cresylect violet) is an excellent one to bring out the types of glial cells separate from the neurons. Though in the end the glial cells were counted on the same pictures as the neurons, we first identified them using the microscope to take advantage of the very precise color differentiation and then marked them on the photographs. The glial cells were shades of blue and the neurons were pink. Those glial cells which had small, pale-blue nuclei were counted as astrocytes. The cells we identified as oligodendrocytes had smaller and darker-blue nuclei than the astrocytes. Those cells with blue nuclei in between the two clearly defined glial types we counted as glial cells but of unknown designation. Figure 14 presents the values of total glial counts from 26 to 904 days of age.

The greatest decrease in density in these glial cell types occurred between 650 and 904 days of age. There was an increase in glial cell counts between 26 and 41 days that followed a similar pattern to the neuron counts. The similarity lends credibility to the counts, because the neuron and glial cell counts were made at different times but on the same tissues. The exact reasons for the increase in glial and neuronal numbers during this time period are still a mystery.

In the older age groups, we found that both the neurons and the glial

Total Glial Numbers in Long-Evans Male Rat Occipital Cortex During Aging

N = 8-13 Animals/Age Group

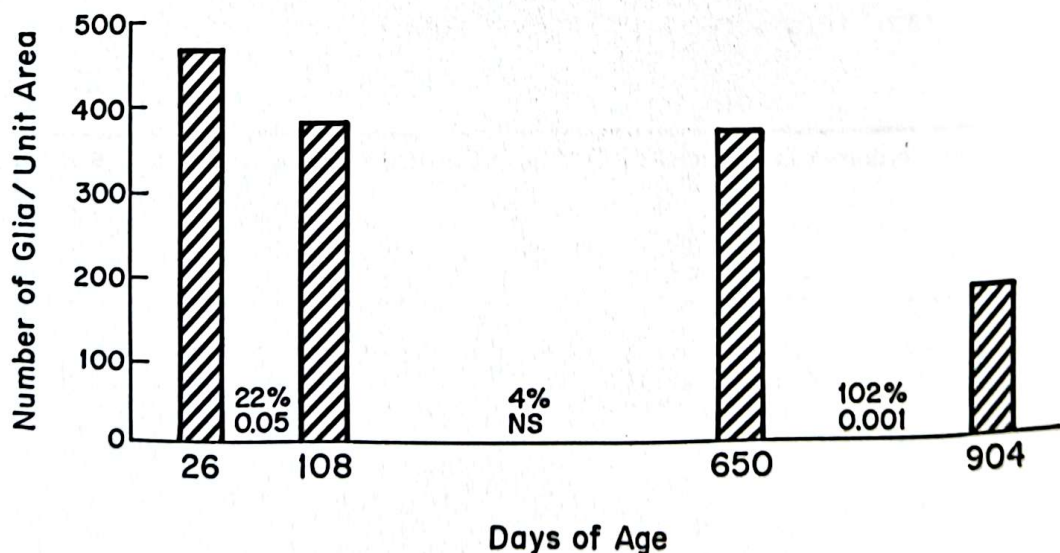


FIGURE 14. Total glial numbers in Long-Evans male rat occipital cortex sample during aging.

cells showed a density decrease by the same amount, 4 to 5%, from 108 to 650 days of age, and that both results were nonsignificant.

In summary, our results have shown a significant decrease in the density of neurons between 41 and 108 days of age, followed by a stability in number until 904 days of age. The density of astrocytes and oligodendrocytes decreased significantly from 26 to 108 days of age, yet no significant decrease occurred after this period until 650 days. Then a marked decrease took place between 650 and 904 days of age.

Having established methods for identifying neurons and glial cells in the cerebral cortex of the male rat, we next counted these types of cells in certain regions of the neurologically undamaged human male cortex. (We obtained the human brains from the Veterans Hospital in Martinez, California.) Glial cells reportedly increase in number as one ascends the phylogenetic scale. The hypothesis is that the more metabolically demanding nerve cells require more glial support cells. In other words, the number of neurons per glial cell or the neuron/glial ratio would be smaller in the more highly evolved brains.

We wondered whether the prefrontal lobe of the human brain was more highly developed and would have fewer neurons per glial cell than the inferior parietal lobe (Figure 15). The reason we decided to compare these two particular areas was that during evolution the prefrontal lobe of man has grown proportionately more than other lobes. But a neuroanatomist, Gerhard von Bonin, once proposed that the inferior parietal

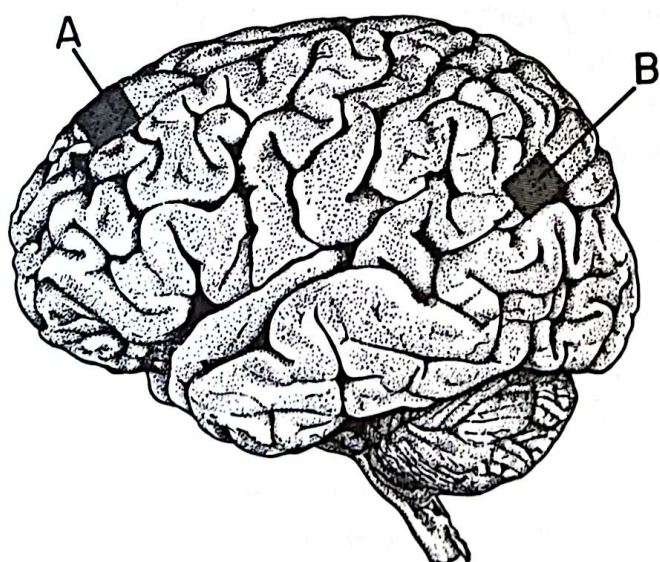


FIGURE 15. Lateral view of the human brain showing the position of the samples used for neuron and glial cell counts. (A) Superior prefrontal region (area 9); (B) Inferior parietal region (area 39).

cortex was also growing out of proportion to other cortical areas and was possibly "pushing" the frontal lobe forward during development. The inferior parietal region, like the prefrontal lobe, is an association cortex and does not receive primary sensory inputs from the external environment. These association cortices receive cortical input from the primary areas and other association areas. The inferior parietal cortex has frequently been called the "association area of association areas" because it receives information from several areas including the visual, auditory and somatosensory association cortices. Over the years, Professor von Bonin's comment had stayed in my mind.

We hypothesized that the region of the cortex that had more glial cells per neuron might prove to be the more highly evolved, or the more active, or both. With this hypothesis in mind, we began to collect cortical samples from neurologically undamaged male human brains from both the right and left prefrontal cortex (area 9) and the inferior parietal cortex (area 39) (see Figure 15). Sections of the tissue were prepared and stained in order to differentiate the glial cells from the neurons.

The results of the counts from 11 human male brains, ranging in age from 45 to 81 years, suggested that the prefrontal area had more glial cells per neuron than the inferior parietal lobe. These findings offer some support to our hypothesis that the prefrontal lobe is more highly evolved than the inferior parietal lobe.

We wanted to follow up our initial frontal and parietal human brain cell counts by studying an exceptional human brain, and we wanted to corroborate the results from our experiments on rats and those of others showing that glial cells increased with an enriched condition and did not increase with age in the controlled laboratory conditions (2, 3). Therefore, when we learned from an article in *Science* magazine in 1978 that no one had studied Albert Einstein's brain microscopically to compare it with "normal" male brains, we decided to request samples from Dr. Thomas Harvey, the pathologist who had obtained Einstein's brain at the time of autopsy in 1955. No one had attempted to make quantitative cell comparisons on Einstein's unique brain in over 25 years. After initially contacting Harvey in Weston, Missouri, we waited for three years before we received the small "cubes of brain" from the right and left hemispheres of Einstein's brain, including areas 9 and 39. The brain, when it arrived in Berkeley, was embedded in celloidin, a substance used to infiltrate tissue to harden it sufficiently to allow cutting micro-thin sections. Thus, the kinds of measurements we could attempt were very limited. Cell counts and cell size were among the few possibilities.

After completing the counts on Einstein's brain cells in a manner

similar to that used for the 11 "normal" male brains from individuals ranging in age from 45–81 years (Einstein was 76), we learned he did indeed have more glial cells per neuron than the normal males had in each of the four areas sampled (7). In fact, in the left area 39, the differences were statistically significant. Since glial cells have been reported by some to increase with aging, we also compared his brain with the three brains in his specific age group. Einstein's brain had more glial cells per neuron.

Our carefully controlled glial counts on aged rats informed us that the number of glial cells not only did not increase with aging, but in the very old animals, actually decreased. Thus, using the results obtained from our enriched rats as a general principle, the glial cell counts on Einstein's brain lent support to the findings of ours and others showing that the more active cortex possessed more glial cells per neuron (2, 3).

Not only did we have Einstein's samples to compare with our normal data base of human males, but, unexpectedly, we learned that we had been given two human female brains from the pathologists. Unaware that these female brains had entered the collection, we counted their cells along with those in the male brains. It was only after the counts were completed that we learned that the female brains had been included. The results showed that the neuron glial cell ratios in the human female inferior parietal cortex (area 39) were smaller than those of the males. Since the samples from the females were so few, only the suggested trends are mentioned and no statistics are offered.

But these apparent sex differences in neuron glial cell ratios in area 39 in human beings led us to attempt to substantiate the results by counting nerve and glial cells in area 39, the inferior parietal cortex, in male and female rats. More stringent controls (such as uniform environments, diets, and temperatures) can be utilized with rats than with human material. Would we also see neuron glial cell sex differences in the rat cerebral cortex under these conditions? McShane et al. have completed the counts in area 39 of 90-day-old male and female Long-Evans rats (8). The initial results suggest that the female does have more glial cells per neuron than the male. A replication experiment is essential to confirm these preliminary results.

In conclusion, our counts on cerebral cortical nerve and glial cells indicate that after 108 days of age, the cell populations remain quite constant until very old age. Before 108 days postnatally, the fluctuations in our cell counts cannot be explained satisfactorily at the present time. If the cortical cells are not decreasing significantly in number with aging, then the decrease in cortical thickness, which was demonstrated previ-

ously in our development and aging curves, must be due to depletion of cell size expressed primarily in the numerous dendritic ramifications. That there are sex differences in cell counts in one cortical area not only needs to be confirmed, but such counts need to be taken from many cortical regions as we attempt to understand the structures behind the functions of male and female cortices throughout a lifetime.

4

THE EFFECTS OF ENRICHMENT AND IMPOVERISHMENT

The History of an Idea

Knowing the basic patterns of development and aging and some of the factors that influence the forebrain structures, we can now begin to approach more directly the correlations among brain chemistry, brain structure, behavior, and environment utilizing various experimental conditions. The Krech-Rosenzweig-Bennett team had demonstrated that some aspects of the chemistry of the cerebral cortex were related to learning behavior. They showed that the enzyme acetylcholinesterase, which is associated with the breakdown of the neurotransmitter acetylcholine at the synapse, is more concentrated in the brains of maze-bright animals than in those of maze-dull animals. Upon joining the team at Berkeley in the late 1950s, I asked whether the distribution of acetylcholinesterase through the cortical layers was also different between these two strains of animals with different learning abilities. (See Figure 2 for a location of the cortical layers.) Not only did we have the maze-bright and maze-dull animals to examine, but two strains of rats were specifically bred to yield high and low concentrations of acetylcholinesterase in the cerebral cortex. We were curious to learn whether these strains as well as the maze-bright and maze-dull strains differed in distribution of this enzyme through the cortical layers.

We thought that a reasonable bridge between the chemistry that was being measured by Bennett, Krech, and Rosenzweig and the anatomy

that could be measured by my group would be a study of acetylcholinesterase concentrations in the anatomical layers of the cortex. Richard M. Diamond, a nuclear chemist-physicist, helped design a method utilizing radioactive copper (^{64}Cu) in place of the stable copper normally used in Gomori's thiocholine method to assay for acetylcholinesterase (1). In order to quantify the amount of acetylcholinesterase in the layers of the cortex, we took samples from rats with the different maze abilities and rats with the different enzyme concentrations. Cylinders of cortical tissue 2 mm in diameter were punched through the cortical surface down to the white matter. Frozen sections 25 μm thick were cut horizontally and were individually incubated with radioactive cupric thiocholine so that we could assess the acetylcholinesterase concentrations throughout the cortical layers, with the aid of a scintillation counter (2).

In the frontal lobe, the greatest differences in concentration of the esterase between the maze-bright and maze-dull strains were seen in the outer layers of the cortex—layers II, III, and IV—with the maze-bright having the higher enzyme concentration. In the somatosensory region, the greatest differences were also found in layers II, III, and IV. The finding that these layers, which are the last to develop embryologically and phylogenetically, reveal the greatest chemical change suggests that the cells in these layers are involved in the learning process.

Although this information was of immense interest, only later, by returning to the data after many years, could we put it together with other data to develop a complete story of the relationship between the chemistry of the layers and the accompanying cell counts. This chemical approach was no doubt fascinating, but I still wanted to learn more about the morphology of the cortex. How did brain cell structures differ?

Our laboratory spent the next few years counting cerebral cortical nerve cells and glial cells in the cerebral cortices of several rats of the maze-bright and maze-dull strains, as well as in high-acetylcholinesterase and low-acetylcholinesterase strains of rats. We found that the neuron-glial cell ratios between the strains were not significantly different. In counting cells we check only the nerve cell bodies and the nuclei of the glial cells. We did not measure the dendritic branches. Much of the cortex is occupied by the nerve cells' branches, which in turn are responsible for most of the acetylcholinesterase. Thus, as we look back, counting cell bodies was perhaps not the best indicator to determine areas of high or low enzyme concentrations in the cortex. After these studies were completed, we then turned our attention to the effects of different environmental conditions on brain anatomy.

As mentioned in Chapter 2, it was Donald Hebb at McGill University who began experiments dealing with the effects of stimulating experiences on behavior. He showed that rats living in enriched conditions were better learners than those which had not benefited from such experience. The exciting results from his report led to the quantitative study of the chemistry of the brains from enriched and impoverished rats which was initiated at Berkeley.

What is meant by an enriched or impoverished condition for rats? At Berkeley, an enriched environment contained many animals in a large cage with a variety of novel objects for the rats to explore, whereas the impoverished animals were caged singly and had neither the objects, nor the large living space, nor the companions. In essence, an enriched environment is one which introduces more stimulation to the body's surface receptors than does an impoverished one, whether it be for rats or human beings. For obvious reasons, the feral condition, the natural outdoor environment for the rat, could not be duplicated. The laboratory condition is sterile, controlled and protective by comparison, and even at the very best, not like living in the rats' natural habitat. Therefore, all types of laboratory environments have to be considered relative to the natural one. Nonetheless, the results from the experimental conditions in the laboratory can be validly compared to each other; one condition is more enriched than the other.

The experimental paradigm used for these initial environmental experiments was designed accordingly by Krech et al. in the Berkeley psychology laboratory (3). Twelve male animals were taken from their mothers in standard colony cages at 25 days of age and were housed together in a large cage ($64 \times 64 \times 46$ cm). Two different wooden "toys" from a set of seven were put into the cage each day. There was also a small wooden maze in the cage which the rats used as a nesting box. For 30 minutes each day, the rats were allowed to explore a Hebb-Williams maze with many partitions, in a box about a square meter in area. The pattern of barriers was changed daily. At about 50 days of age, formal maze training began in several other types of mazes, all providing different levels of problem solving. Glucose pellet rewards were given. The rats could normally take food and water ad libitum. Figure 16 illustrates the enriched, standard colony, and impoverished conditions we now use: the cages are built of sheet metal and metal mesh, and metal toys have replaced the wooden ones, which the rats used to chew upon.

Simultaneously, at 25 days of age, the littermate of each of the enriched rats was placed in the impoverished condition, consisting of pri-

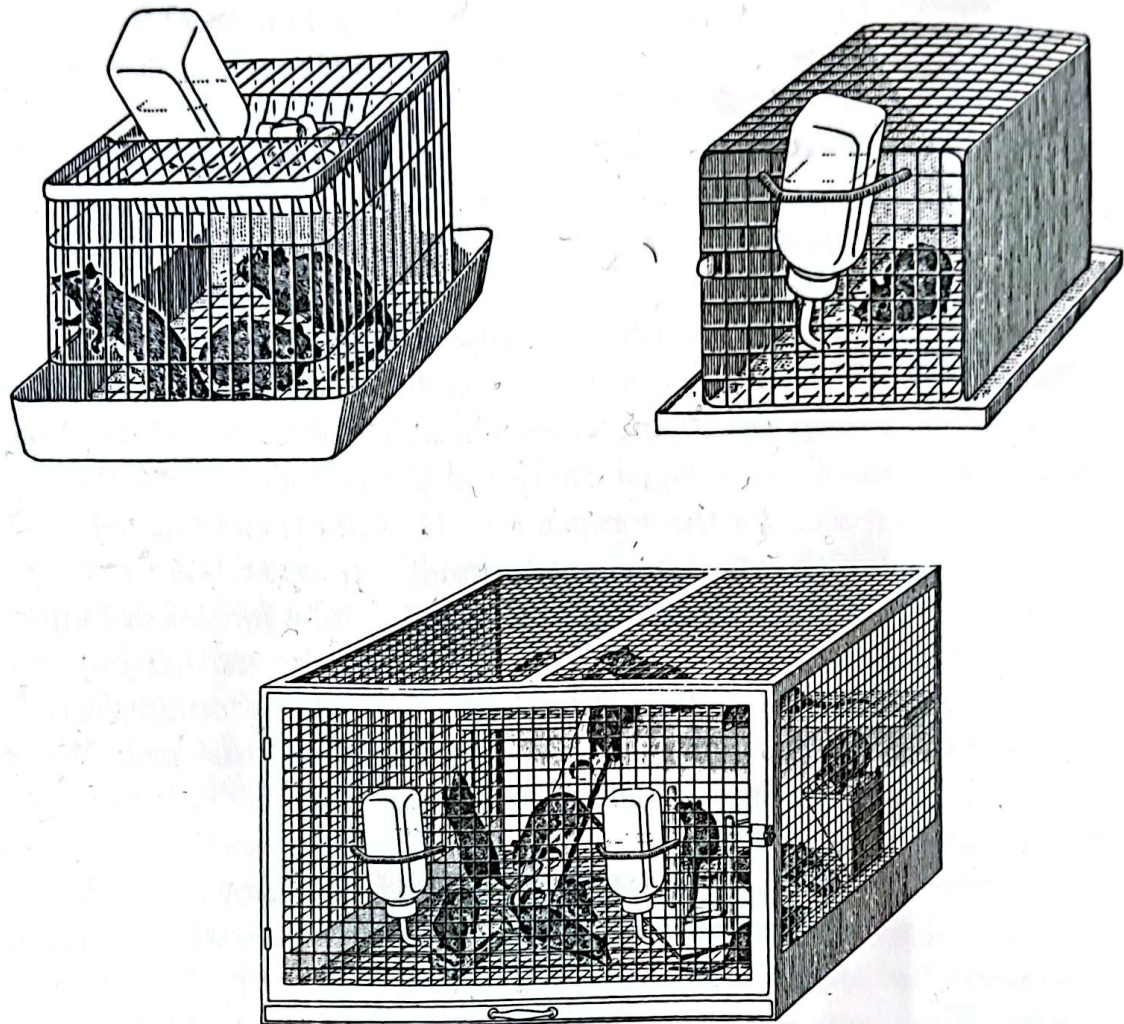


FIGURE 16. Illustration of environmental conditions: (below) enriched, a large cage with 12 rats plus toys; (upper left) standard colony, a small cage with 3 rats; (upper right) impoverished, a small cage with 1 rat.

vate cage ($28 \times 20 \times 20$ cm) with solid metal walls on three sides. (In later experiments the cages were all wire mesh or Lucite depending upon the experimental conditions, and the dimensions were slightly altered.) Food and water were available ad libitum, and every rat was given a glucose pellet each time its enriched littermate was rewarded with one. The impoverished animals did not see or touch another animal, nor were they handled during cleaning of the cages; but all animals in both groups were weighed at weekly intervals. After the animals had lived in their respective environmental conditions for the required period of time, all rats were coded so the anatomists were not aware of the previous experimental conditions, in order to prevent bias during the histological brain measurements.

The results indicated that either enriching or impoverishing the environmental complexity and training of rats caused measurable changes in

brain chemistry and brain weight (4). More specifically, some of the results demonstrated that the cerebral cortex of rats subjected to enriched or impoverished environmental conditions showed not only differences in the synaptic enzyme, acetylcholinesterase, but also the wet weight of samples removed for chemical analysis. It was the weight differences which concerned us as anatomists. Though small (8% in the samples from the visual cortex and 3% in the samples from the somatosensory cortex), the differences were consistent and could be replicated. The cortical weights from the enriched animals exceeded those from the nonenriched in 79% of the cases for the visual area and 64% for the somatosensory area.

Though cortical weight differences were found between the two experimental groups, no significant changes in the weights of the subcortical tissues were seen. More recently, others have found no environmental effect on total hindbrain weight (5, 6). Floeter and Greenough (7) did, however, clearly demonstrate neuronal differences in a hindbrain structure—the cerebellum—between the enriched and the nonenriched rats, but no brain weight changes were measured. Also, in the Berkeley experiments in the early 1960s, where the animals lived in their environmental conditions for 80 days, the body weights of the isolated or nonenriched rats were greater than those of the enriched by about 7%. The body weight gains were in the opposite direction from the cerebral-cortical gains. Here was an interesting paradox: the cerebral cortex weighed more in the enriched animals, yet their body weights were less. In other studies where we measured the dimensions of some subcortical structures, including the diencephalon, we learned that the areas of these regions were more positively correlated with body weight than were our cortical measures. The findings indicate the importance of quantifying differences in specific regions of the brain rather than taking whole brain measurements as many earlier studies did.

For example, total brain weight differences between the enriched and impoverished animals after 80 days in their respective conditions were only 1% ($p < 0.01$) with 175 pairs of rats (8). With a larger sample of animals, 200 pairs, ranging in age from 18 to 530 days, Walsh et al. (6) found a mean difference of 3%. Others have also shown significant differences in rat brain weights between the two groups (9, 5). These total brain weight differences were so small, in part, because the major percent changes were localized. Since many brain regions were unaffected by these environmental conditions, whole-brain weight measures masked the localized changes.

Within the cerebral cortex, the largest brain weight changes were

seen after 30 days of differential rearing (8). In male rats after 80 days, the occipital cortex showed a 6% difference in wet weight; at the same time, the somatosensory cortex showed only a 2% difference. This regional difference between the occipital cortex and the somatosensory will become of greater interest when cortical sex dissimilarities are reported in Chapter 6. Female rats were not used regularly in the early days of these experiments.

Wet- and dry-weight assessments were made in order to determine whether the difference in cortical weight was due to an increase or decrease in tissue fluid. By measuring both wet-weight and dry-weight brain samples from enriched and nonenriched rats, three separate investigations concluded that the experimental differences between the two conditions in rats were the same whether the tissues were wet or dry (10). Accumulation of fluid did not seem to be the factor causing the difference in brain weight.

Various investigators followed their brain weight measurements with macroscopic measurements of the cerebral cortex, such as cortical length and width. Reportedly, rat cerebral growth in width is essentially complete by 20 days after birth, but the length continues to grow at least to the age of 90 days (11, 12, 13). Altman et al. found that they could alter the length, but not the width, of the rat cortex by exposing the animals to 3 months of environmental complexity (12). Evidently, 3 months, or 90 days, was sufficient to show such a change, but 30 days proved an insufficient time to induce similar results (14). However, Walsh et al. chose a period in between, namely, 80 days, during which there was a highly significant change (2.5%) in length; whereas, with only 30 days, they found but a 1.2% difference in length of the cortex between enriched and nonenriched (13, 15). Others continued with this experimental approach and found that exposure to enriched conditions from 18 days to 530 days increased cerebral length (16, 17, 18, 19).

Walsh (15) pointed out that the length effect was limited to the anterior regions, and cortical weight effects were more marked at the posterior, and that they might therefore be largely independent of one another. This hypothesis would be consistent with results showing that weight and length effects appear to follow different temporal patterns depending upon the period of exposure to the experimental conditions (13, 18).

Prior to these experiments, the brain had generally been considered incapable of macroscopic physical changes as a consequence of alterations in experience. Here now was the opportunity to document weight and

length increase in the cerebral cortex with quantitative anatomical microscopic measures.

Anatomical Changes Within the Brain

When we were designing the first experiments on the Berkeley campus in the early 1960s to deal with the enriched and impoverished environments, it was not well established how long it would take to bring about measurable brain changes. Eighty days was the first time period we chose. Also in those early experiments, the rats were given maze training as part of their enriched conditions. This aspect of the enriched condition was dropped after 1965, when we found that it was not an important factor in producing the effects associated with enrichment.

In our first, and later replicated, enriched-impoverished anatomical experiment in the early 1960s, littermate male S₁ (maze-bright) rats from the psychology department colony were used (20). At 25 days of age one animal from each pair, chosen at random, was placed in the environmental complexity and training group (ECT). The littermate was assigned to the isolated-condition group (IC).

The simplest microscopic anatomical measurement we could make to compare the brains of the enriched and impoverished animals was our basic cortical thickness measurement. We measured it on transverse-cut sections 20 μm thick, utilizing subcortical landmarks to ensure uniformity in sampling from one rat to the next. Samples were taken from both the somatosensory cortex and the occipital cortex. For these early measurements of cortical thickness, we used an ocular micrometer, a small calibrated scale placed in the eyepiece of the microscope, to measure the distance from the beginning of layer II through layer VI on the brain slices (see Figure 2). We took differential counts of neurons and glial cells directly in the microscopic fields; in addition, we counted the numbers of blood vessels and classified them by size.

The results of these earliest experiments revealed significant differences in the thickness of the occipital cortex between rats living in enriched and impoverished environments for 80 days (see Table 3). In both the original and the replication experiments (Experiments I and II, respectively), the occipital cortex from the enriched rats was significantly thicker than that of the impoverished animals (6%, $p < 0.001$) (20). The results from both right and left hemispheres were combined to obtain these values.

TABLE 3

Thickness (μm) of Occipital Cortex from Enriched and Impoverished Rats (Excluding Layer I)

Experiment	N	ECT		IC		p		
		\bar{X}	S.E.	\bar{X}	S.E.	ECT/IC	ECT vs. IC	ECT > IC
I	11	1332	17	1271	22	1.048	< 0.001	7/11
II	9	1404	29	1298	27	1.082	< 0.001	9/9
I + II	20	1364	16	1284	17	1.062	< 0.001	16/20

ECT = Environmental complexity and training; IC = isolated condition.

Layer I was not included in these results because separate measurements of this layer alone in enriched and impoverished animals showed no significant differences. According to Eayrs and Goodhead (21), layer I shows very little change during the postnatal growth of the rat cerebral cortex. This fact alone is of interest when considering the dynamics of cortical development. Layer I is one of the early layers to form in the developing cortex; it receives terminal dendrites from cells in each of the underlying layers as well as axonal endings. These dendritic and axonal branches form a dense fiber plexus which constitutes layer I. Layer I apparently is not measurably altered in structure with our multisensory input: either our methods for determining morphological differences in this layer were not sophisticated enough or indeed no changes occur. The precise function of this layer, like that of many other cortical layers, is not known.

In the measurements of weight differences between the enriched and impoverished rats, the results for the somatosensory cortex had been smaller and less clear-cut than those for the occipital region. This finding was noted again in the present measurements of cortical thickness in the male S₁ rats. Although some differences in the somatosensory cortex were displayed between enriched and impoverished male rats, as seen in Table 4, the differences were not as striking as in the occipital cortex.

In an attempt to understand what cellular changes were responsible

TABLE 4

Thickness (μm) of Somatosensory Cortex from Enriched and Impoverished Rats (Excluding Layer I)

Experiment	N	ECT		IC		p		
		\bar{X}	S.E.	\bar{X}	S.E.	ECT/IC	ECT vs. IC	ECT > IC
I	10	1962	18	1825	22	1.075	< 0.001	10/10
II	8	1975	40	1985	38	0.995	NS	3/8
I + II	18	1968	20	1896	21	1.038	< 0.01	13/18

for the cortical thickness differences, cell counts were taken. In our first experiments, nerve cell and glial cell counts were made in each microscopic field, reading vertically from the pial surface of the cortex down to the underlying white matter. In the occipital cortex, the neurons per microscopic field were less numerous by 17% ($p < 0.01$) in the enriched animals than in the impoverished ones. The mean distribution of neurons in the rat occipital cortex from layers I through VI is shown in Figure 17 for the initial (I) and replication (II) experiments. The decreased number of neurons per microscopic field informed us that more intercellular material was present in the enriched animals than in the impoverished animals. It appeared that the nerve cells were not increasing in number but were getting larger by gaining more branches, so that their bodies were being spread farther apart and therefore the counts of nerve cells per microscopic field were fewer in the enriched animals. Since, at the present time, there is no evidence that the number of neurons increases in the cerebral cortex after birth, we conclude that any additional increase in cortical mass must be due in part to changes in the size of the nerve cells, including their dendritic branching.

The largest, most consistent differences between nerve cell counts in the enriched and the impoverished groups were apparent in the outer layers of the cortex, layers II, III, and IV (20). To be seen shortly, the greatest increase in the area of the neuronal soma also occurred in these

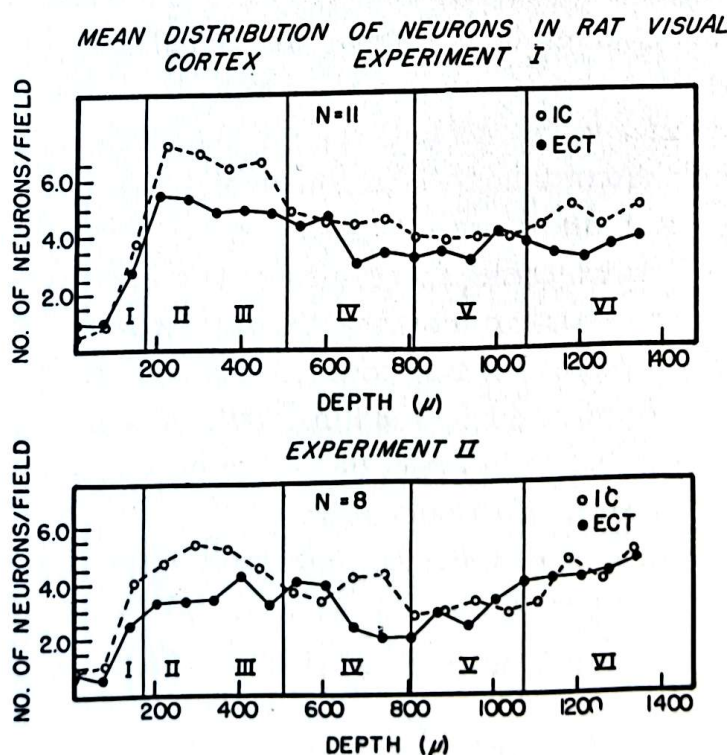


FIGURE 17. Mean distribution of neurons in rat occipital (visual) cortex showing the results in (I) the first and (II) the replication experiments.

layers. During embryological development, the lower layers of the cortex form first; each new group of cells has to migrate through the underlying layers to reach its outer position. In other words, an "inside-out" developmental pattern occurs. Our results indicating that the outer layers show the greatest plasticity suggest that those cells which are last to develop retain a greater ability to become modified in structure with an increase in new stimuli to the cortex.

It is clear from the above discussion that the animals living in enriched and impoverished conditions for 80 days, from 25 to 105 days of age, showed structural differences in the cerebral cortex. The enriched rats had thicker cortices, with the nerve cells spread farther apart, particularly in the outer layers. By counting the cortical cells per microscopic field, we did not record the total number of neurons and glial cells in a large cortical area but an average number in a single column of cells. With this method, no significant difference in number of neurons was noted between the enriched and impoverished animals. This was no surprise; we did not anticipate more neurons in the enriched rats because, to date, no cortical neurogenesis has been documented more than a few weeks after birth.

With our initial experiment in 1964 where we measured the area of the soma and the nucleus of the neuron by tracing their outlines from microscopic projected images, we found no significant differences between the size of the neurons from the enriched or the isolated rats. We were surprised with these results, for if we speculated that some of the thickness increases were due to a greater number of dendrites, then, in turn, we might expect a larger nerve cell body and nucleus to be supporting those dendrites. According to Glendinnen and Eayrs (22), the larger the neuron size, the more dendritic branching. All information suggested that we should be expecting larger neurons in our enriched rats.

For our later cell measurements, we turned to the negatives of the pictures previously used for cell counts. These negatives were projected through a microfilm reader; the final enlargement for the cells was 2000 \times . We expected these larger images to provide for a smaller margin of error in tracing the cell boundaries.

Since the photographs for cell counts were taken from the pial surface down to the underlying white matter, it was possible to divide the cortex (excluding layer I) into equal thirds in order to evaluate whether there was any difference in size between cells in the superficial layers and those in the deeper ones. The divisions were referred to as upper, middle, and lower. The results of the initial and replication experiments were that the upper division showed the greatest increases in both soma and nuclear

size. In the combined experiments, the increase in perikaryal size for the enriched rats in the upper division was 18% ($p < 0.001$), and in nuclear size it was 20% ($p < 0.01$). Though the upper division showed the greatest increases, each division indicated a significant increase in the size of the soma and nucleus in the enriched animals compared with the isolated animals. Thus, all layers of the cortex included in these arbitrary divisions were showing differences, with the outer cellular layers indicating the greatest changes.

In 1966 Altman proposed that the outer layers of the cortex play a special role in central nervous system function (23). They exhibit slow postnatal growth, and an increase in number of cells as we ascend the phylogenetic scale. Their strategic position makes it possible for these outer cells to become the modulating influences on the input to and output from the cerebral cortex. Our data also indicate that the outer layers appear to show a greater response to the additional stimuli entering the cortex in an experientially enriched condition than in an isolated one.

By refining our methods, we were able to demonstrate the predicted differences in soma and nuclear size. (We do want to point out again that all of the tissues were coded to prevent bias during measurement. Otherwise, subconscious influences might play a role.) In a stepwise fashion, we have demonstrated that the enriched animals have larger nerve cell bodies and nuclei in all three levels of the visual association cortex, with the outermost level showing the greatest differences.

In the initial studies, we found not only fewer neurons per microscopic field, but fewer glial cells as well. We found the density of glial cells per microscopic field to be less in the enriched than in the impoverished rats by 7% in the occipital cortex and 6% in the somatosensory cortex. These findings indicated an actual increase in glial cell number, because the glial cell-neuron ratio was greater in the enriched animals than in the impoverished ones, and the total number of neurons was presumably constant. The density of glial cells through the cortical layers was so small that we came up with no significant findings in comparing the glial populations per layer between experimental groups. Table 5 illustrates the cell counts per unit area of occipital cortex in two experiments.

For the more detailed counts, the cells were not differentiated directly with the aid of the microscope as previously, but instead, photographs were taken of microscopic fields extending from the pial surface through the six layers of the cortex. We had previously reported a higher glial cell-neuron ratio in the enriched brain because the decrease in glia per microscopic field was significantly less than the decrease in neurons. Even

TABLE 5
Differential Cell Counts in the Occipital Cortex from Enriched and Impoverished S₁ Male Rats (N = 17 Pairs)

CELLS	ECT X	IC X	ECT > IC	% DIFFERENCE	p
Neurons	485.8	500.6	6/17	-3.0	NS
Astrocytes	87.0	83.0	9/17	4.7	NS
Oligodendrocytes	75.2	62.5	13/17	20.4	<.02
Indeterminate glial cells	23.4	17.2	14/17	35.5	<.01
Total glial cells	185.6	162.8	13/17	14.0	<.01
G/N	0.385	0.332	12/17	15.9	<.02

though we found this relative increase in glial cells, we were not satisfied with the results. Using our new, more refined techniques, we found an absolute increase in glial cells in the enriched brains, giving a higher glial cell-neuron ratio of 16% overall, with the enriched animal having the larger ratio in 12 out of 17 pairs ($p < 0.02$).

We found that the number of oligodendrocytes (a type of glial cell) increased (20%, $p < 0.02$) and cells with characteristics of both oligodendrocytes and astrocytes were proliferating (36%, $p < 0.01$). Nothing definitive can be said regarding the astrocyte counts, because in the first experiment the enriched animals had more astrocytes than the impoverished (27%, $p < 0.01$) and in the replication experiment the enriched animals had fewer by 7% (NS). By combining the results of both experiments, we arrived at an astroglial density that was greater by 5%, but this difference failed to reach significance.

A glial cell increase due to behavioral manipulation was reported by Altman and Das in 1964 (24), using similar but not identical enriched conditions and using radioautography to measure cell proliferation. These investigators did not attempt to differentiate types of glial cells but attributed an increase in labeled cells largely to cells situated in the cortical radiation and corpus callosum rather than in the cortical gray. We counted cells only in the cortical gray. Evidently, in a more recent communication with Walsh, Altman now thinks the cells he was identifying as glial were postnatally derived undifferentiated precursors of both neurons and glial cells. At the present time a clear conclusion about glial cell proliferation and enrichment cannot be drawn from his data.

Studies on whether adult cortical neurons can proliferate in superenriched conditions are in progress in our laboratory (25). York is attempting to demonstrate neurogenesis using ^3H to quantify dividing cortical cells. She hopes to differentiate between dividing neurons and glial cells between adult enriched rats and rats in standard colony conditions by labeling cells with ^3H -thymidine. Only by counter-staining the nerve and

glial cells with a differential stain such as Luxol-Fast Blue-Cresylecht Violet will we really be certain that we are accurately separating the two populations of cells. This experiment has the potential to offer some clarification to the early experiment of Altman on whether the cells he identified were neurons or glial cells, though he was dealing with young animals and York (25) is using adult rats.

Since the early reports on glial cell counts, other investigators have also found increases in these cells with enriched conditions. In 1977, Szeligo and Leblond (26) confirmed our glial cell findings by reporting that 30 and 80 days of enrichment increased oligodendroglial densities by 27% and 33%, respectively, in comparison with those of social controls and isolated animals. (The glial cell counts of social controls and isolated animals did not differ significantly from each other.) Like us, they found that the differences in astrocyte counts were not as great as those in oligodendrocytes counts. Between animals experiencing their enriched and impoverished conditions for 80 days, the difference in astrocyte counts amounted to 13%. It has been reported that when neonates are handled in the first 10 days of life for a 15-minute period each day, the astrocytes increase by 12% but no significant changes are found in oligodendrocyte counts. But it is not clear what causes the astrocytes to be more susceptible to additional stimulation in the young and the oligodendrocytes to show a greater response in older animals. At one time, it was proposed that oligodendrocytes represented a later stage of astrocyte development; in other words, both came from the same stem cell but the astrocytes were younger than the oligodendrocytes. This theory has not been supported by additional experimentation.

Cummins and Walsh (27) continued to be intrigued with the role of glial cells and environmental input. They were able to demonstrate that the glial cell-neuron ratio was greater in the enriched brains by 8% after 90 days of rearing, thus supporting our original findings. However, in their experiments, it was the astrocytes that accounted for the increase in glial cell number. When such differences as these are obtained, it seems imperative for the investigators to collaborate and both count the same tissues to see where the differences lie. Since two groups—Szeligo and Leblond and Diamond et al. both showed that it was the oligodendrocytes that predominated following enrichment, it seems justified for now to accept these findings as the truer picture until proven otherwise. At least three groups of investigators have consistently reported an increase in glial cells in enriched conditions compared with impoverished. We must always consider when we compare only the enriched animals and the impoverished whether the enriched condition is only sparing or re-

taining the number of cells and not allowing them to decrease as they would with impoverishment. The ^3H -thymidine experiments being carried out with standard colony and enriched animals should help to clarify this possibility.

What is the advantage for the nerve cell in having more glial cells? Numerous functions have been attributed to glial cells. Some of these have already been mentioned in Chapter 3. Different types have different functions, but by addressing them collectively one can say, in general, that they form myelin in order to facilitate impulse conduction along the axon; they regulate ionic movement between nerve and glial cells and the vascular system; they give metabolic support to the nerve cell in a manner not yet completely understood; they give additional specificity to synaptic membranes because of their close proximity to the synaptic junction; they assist with protoplasmic movement in the axon; they help regulate vascular flow; they supply either energy or substrate for energy for axonal function; and they serve as a scaffolding for early neuronal migration. Any or all of these functions may be demanding more glial cells for increased neuronal function in the enriched animals.

Besides counting glial cells and neurons, we also made quantitative measurements of the blood vessels in the cerebral cortex. Our results indicated that the somatosensory areas had more blood vessels than the visual area, a finding in accord with another investigator, Craigie (28). Our own findings are reported in Table 6. That the enriched group showed fewer capillaries per microscopic field was in agreement with our

TABLE 6A

Blood Vessels per Field in the Occipital Cortex from Enriched and Impoverished Rats in Experiment I

Type of vessel	N	ECT		IC		ECT/IC	p	ECT > IC
		\bar{X}	S.E.	\bar{X}	S.E.			
Capillary	9	6.17	0.23	7.07	0.32	0.873	< 0.05	1/9
Over 5 μm	9	2.39	0.35	1.90	0.22	1.258	< 0.05	7/9

TABLE 6B

Blood Vessels per Field in the Somatosensory Cortex from Enriched and Impoverished Rats in Experiment I

Type of vessel	N	ECT		IC		ECT/IC	p	ECT > IC
		\bar{X}	S.E.	\bar{X}	S.E.			
Capillary	7	8.00	0.54	8.67	0.72	0.923	NS	2/7
Over 5 μm	8	2.89	0.36	2.83	0.29	1.021	NS	6/8

findings of a smaller number of neurons per field, indicating again that the difference in the thickness of the cortex was due in part to an increase in the dendritic branching between both the nerve cells and the capillaries. The enriched group showed an increase in the number of larger vessels, those over $5 \mu\text{m}$, presumably because this group required a greater blood supply (see Table 6). It is possible that all vessels increased in size with the enriched condition, thus pushing more vessels into the "above $5 \mu\text{m}$ " category. When the heart was examined after prolonged exercise, no new blood vessels were found, but it was shown that the existing vessels increased in diameter. It appears that a similar process occurs in the cerebral cortex.

The degree of vascularity corresponds directly to the degree of functional activity. In other words, a greater blood supply is necessary for higher metabolic activity. Thus, skeletal muscles need numerous blood vessels, in contrast to cartilage, which needs few. Dunning and Wolff (29) have reported that the vascularity of brain tissues varied with the number of synaptic structures and not with quantitative differences in nerve cell bodies. If an increase in cortical thickness is due to additional dendritic ramifications, which can account for more synapses, we would, in fact, expect an increased blood supply in the enriched brains.

After completing the nerve and glial cell counts and neuronal soma measurements, the next logical step in searching for anatomical changes due to modified environments led to the branches of the nerve cells, specifically dendritic and synaptic quantification. But before presenting these results, one study will be mentioned that provides beneficial background data for these more refined cellular measurements. We wanted to learn how much of a cortical area, such as area 18, was responding to our environmental conditions, the whole or only parts, by identifying and measuring areas including specifically active functional units called vertical columns.

Cortical cells are arranged in vertical columns with a width varying from 200 to $1000 \mu\text{m}$. The nerve cells in a particular vertical column are related to the same peripheral receptive field and are activated by the same peripheral stimulus. Though our lines drawn to measure cortical thickness did not trace individual vertical columns precisely, by examining each line separately we would have a better idea on whether our enriched environments were affecting specific columns.

Figure 18 illustrates that whether the animals had been in their environments at age 60 to 64 days or age 60 to 90 days; each separate line showed a significant difference due to enrichment (additional data from these age groups will be discussed in Chapter 5). These more specific

Measurements of Individual Lines in Areas 18 and 17

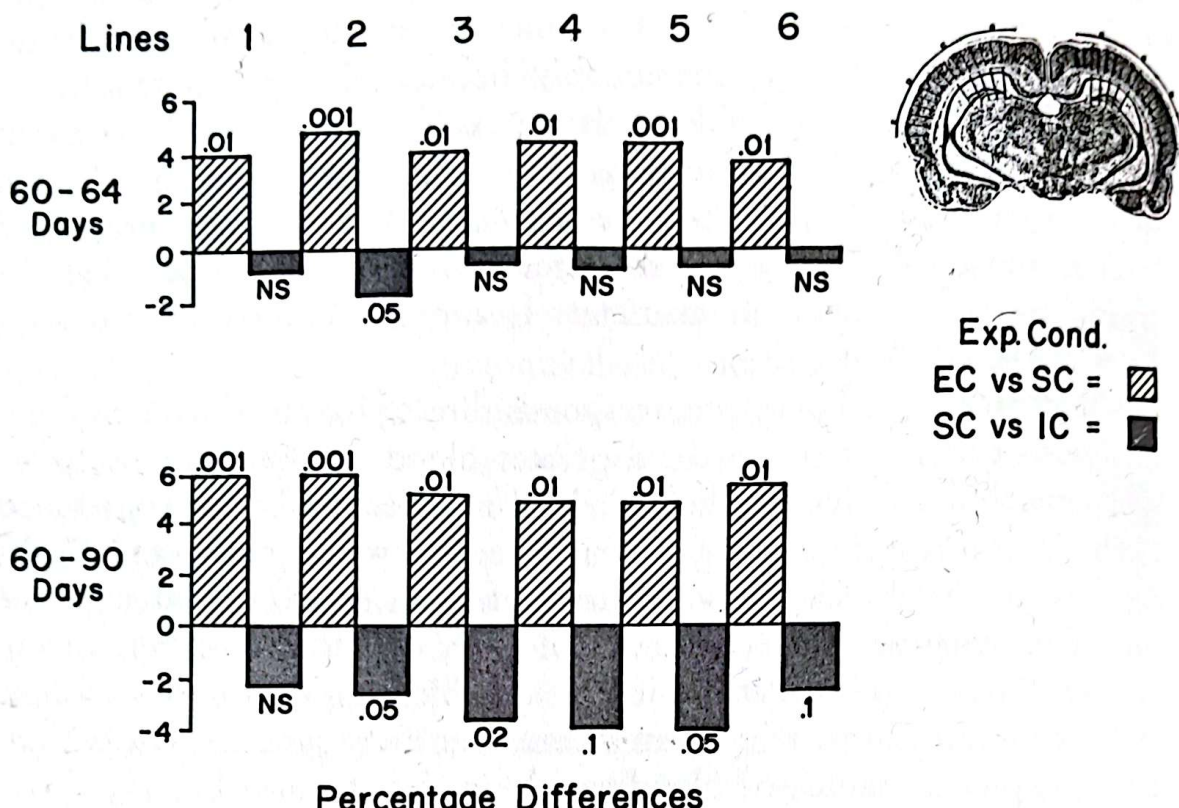


FIGURE 18. Cortical thickness percentage differences between individual lines drawn on the cortex to determine if separate "columns" were responding to the environmental conditions. EC = enriched, SC = standard, IC = impoverished.

data indicated that not just single columns of cells in the cortex were being affected by our environmental conditions: several millimeters of tissue were responding equally. This columnar information proved useful as we began to consider which nerve cell parts might be altered in their dimensions by the environment. Knowing that a large area of cortex was involved, we could now validly choose representative cortical cells from a broad region.

By this time we had thoroughly established that external environmental influences could alter the structure of the cerebral cortex, as shown by cortical thickness changes. Very early in our studies in 1964, we proposed that dendrites, the receptive extensions of the surface membrane of the nerve cell, were partially responsible for our cortical thickness changes. Small projecting receptive appendages on the dendrites called dendritic spines are known to be adaptable to severe deprivation, so these little projections seemed reasonable structures to quantify in our experiments. The functional junctions between the axon of one nerve cell and the

dendrite of another, a synapse, were an obvious next choice for measurements in brains from animals exposed to stimulating or deprived environments. By examining each of these structures systematically, we could gain a clearer understanding of how the environment affects the morphology of the cortical nerve cell.

Changes in the Structure of the Nerve Cell

DENDRITE BRANCHING. The first recorded results showing that dendrites were indeed being modified by enriched and impoverished environments were made in our laboratory by Holloway in 1966 (30). He demonstrated that the dendritic branches of stellate neurons (star-shaped nerve cells) in layer II of the occipital cortex (in area 18) were responding to the varied environmental conditions. He chose the stellate cells because they were more abundant in the outer layers of the cortex than in the lower layers. In the early description of our environmental results, we said that the outer cortical layers showed the greatest increases in neuronal soma size due to our form of enrichment. Holloway confirmed the prediction from our 1964 experiments; the changes in cortical thickness were due in part to changes in the dendritic branching or membrane arborization from the stellate nerve cells.

About 9 years after these results were obtained, the experiments of Greenough and Volkmar on small, medium, and large pyramidal neurons and on stellate neurons showed that enriched rats consistently had more higher-order dendritic branches than their impoverished littermates (31). In the pyramidal neurons, neurons whose cell bodies are shaped like pyramids, the branching differences occurred primarily in the basal dendrites, those extending from the base of the cell body (see Figure 19). Reportedly, these basal dendrites receive a rich input from nerve cells on the

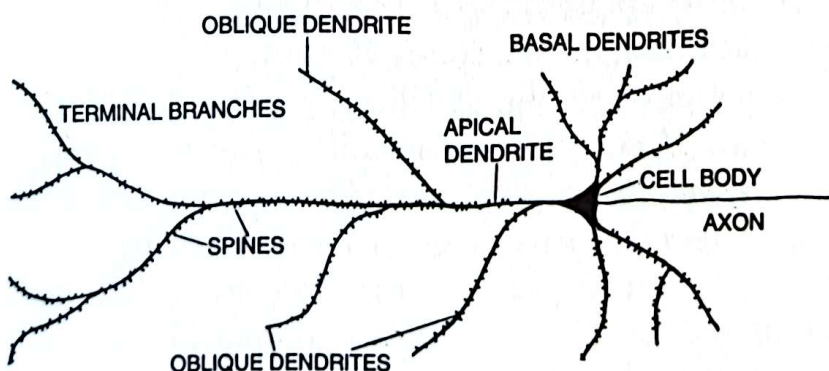


FIGURE 19. Pyramidal nerve cell showing basal dendrites, an apical dendrite, and terminal branches, all with spines.

same side of the cortex. That is, they do not primarily receive fibers from subcortical neurons or from nerve cells in the opposite cortex. It appears that they integrate information that has come into the same side of the cortex; this implies higher-order processing than would take place if the neurons were receiving information from subcortical areas, for example. No consistent differences in the length of branches were found either by Holloway or by Greenough and Volkmar.

The experiments of Holloway and of Greenough and Volkmar were carried out on neurons from the occipital cortices of young animals. It is possible that other regions of the cortex also change. In an attempt to find out, pyramidal cell dendrites were quantified in layers II and IV in the frontal cortex and layers IV and V in the temporal cortex (31). These investigators performed quite an elaborate analysis of the dendritic patterns. Their method was designed to give an accurate account of the number and length of dendritic branches and their location with respect to the cell body. Their type of analysis was reportedly more sensitive to the differences in higher-order dendrites than any previously used in the quantification of dendrites in the occipital cortex. The investigators found no systematic pattern of differences in dendrites from the frontal cortex between the enriched and the impoverished animals. However, in the temporal cortex, the effects of the two environmental conditions were clearly apparent in the higher-order branching of the basal dendrites. The enriched animals had significantly more basal branches on neurons in layer IV than their impoverished littermates. But the cells in layer V of the temporal cortex apparently had more basal branches in both the enriched and impoverished than in the standard colony group. The reason for this difference is not clear.

The investigations of dendritic patterns suggest that another factor to be considered in measuring dendritic branching is the variation of pattern between litters (31). These investigators found significant differences between litters in both the frontal and temporal cortices. Yet other investigators have noted the consistency of dendritic length between strains. It seems incongruous that litter patterns would vary yet strain patterns do not. Variations in dendritic responses were shown in the frontal and temporal cortical results. The fact that not all dendrites in the cortex respond equally to these experiential environmental conditions leads one to believe that the responses are specific to certain neuronal input patterns and not generally due to hormonal or nutritional factors, an observation made by Walsh as well (32).

If the dendrites could change in the young animals, we wished to learn whether they could do the same in the adult rats. A collaborative

effort between our laboratories and those of Uylings et al. (33) was begun in the mid 1970s to study adult animals, 112 to 142 days of age. The results revealed that the basal dendrites in pyramidal neurons in layers II and III of the enriched and standard colony rats showed lengthening in the terminal segments of the dendrites as well as an increase in the branching relative to the baseline animals at 112 days of age. The first-, second-, and third-order terminal segments all increased in length significantly in the enriched animals compared with the standard colony animals. These findings strongly indicate that the new branches were mostly formed at a considerable distance from the tips of the terminal segments. Nonetheless, here was evidence that the dendritic structure of 142-day-old adult rats could be changed by enriched conditions.

Next we examined older animals. The dendritic measurements were made on the same 600- to 630-day-old enriched animals that had lived for 30 days with young rats (age 60 to 90 days), an experiment to be covered more thoroughly in Chapter 5 (34, 35). The only statistically significant difference in dendritic pattern between the enriched and standard colony 600- to 630-day-old group in this study emerged in the analysis of the sixth-order dendrites. The first branches off the nerve cell body are the first-order dendrites, the next branches are the second-order, and so on. The sixth-order dendrites were significantly longer in the 630-day-old enriched rats than in the 630-day-old standard colony rats ($p < 0.03$). The mean length of the sixth-order segments from the enriched animals was 54 μm compared with a mean length of 29 μm for the sixth-order segments from the standard animals.

There are at least three possible explanations for the increased length observed in the enriched-environment animals: (1) a growth (elongation) of the sixth-order segments, (2) a retardation of the retraction process that occurs with age, and (3) a loss of segments which would result in greater internodal lengths and thereby artificially increase the remaining segment length. Neither the total number of terminal segments (34) nor the frequency of sixth-order segments (35) differed between the 630-day-old standard colony and enriched groups. Thus, it is unlikely that there was an artificial increase in length due to segment loss. Whether or not the increased length is due to a growth of the segment or a retardation of the retraction process cannot be clearly determined from these results. But recall that in the earlier study with Uylings et al., when a baseline group was available for comparison, the adult rat did show an increase due to enrichment and not just a maintenance of a status quo in dendritic length. Undoubtedly, for a final conclusion to be validated, another experiment will have to be done with a baseline group for the older animals.

to see whether the dendrites actually grew or maintained their original length. But whichever way this particular question is answered, the present study extended the findings of greater dendritic lengths in conditions of environmental stimulation from adult rats to middle-aged rats. It appears that the effect of the external environment on dendrites is still as important a consideration for the middle-aged rat as for the younger, adult one. It is possible that a longer exposure of the middle-aged animals to an enriched condition would result in a dendritic length change more similar to that observed in the adult animals.

Comparing our results on 630-day-old rats with those that Uylings et al. (33) had obtained in 142-day-old animals, we find that the lengths of segments of each order were markedly shorter in our study. Although our investigation used a different strain of rats, the differences in segment lengths were more apt to be due to the retraction of the dendritic length that occurred with age. Our belief that different strains of rats do not have measurable differences in dendritic lengths is supported by the similarity in dendritic lengths reported by Uylings et al. (33) and Lindsay and Scheibel (36).

Rats are not the only animals that experience brain changes in response to environmental conditions. Other species of rodents—mice and gerbils, for example—have shown occipital cortical weight and thickness changes as a consequence of exposure to various living conditions. Furthermore, the results from studies on monkeys have indicated that after 6 months of differential rearing, cells in the cerebellum, a more primitive hindbrain structure, also display differences (37). More specifically, the Purkinje cells in the archicerebellum, or old, vestibular cerebellum, displayed more extensive spiny branchlets in the enriched than in the nonenriched monkeys. The dendritic branchings from the smaller, excitatory granule cells of the cerebellum were not measurably affected. Perhaps of even greater interest, is the finding of changes in the old cerebellum, the nodulus and the paraflocculus. If one were to predict significant responses in a region such as the cerebellum, one might especially anticipate modifications in the neocerebellum, because of its rich associations with the cerebral cortex or neocortex, which have been shown to respond so readily to the enriched condition.

DENDRITIC SPINES. If the experiential environment can change the dendritic structure, it should be able also to modify the dendritic spines, the little appendages along the shafts of the dendrites. Spines were first described by Ramón y Cajal (38), who showed that they come in a variety of shapes. The classical spine structure with a thin stalk and a

bulbous terminal ending showed a remarkable constancy in size (39). Ramón y Cajal suggested that the spines served to increase the receptive area of the dendrite. More recently, in the late 1970s, several workers reported that 85% of the synapses on the dendrites are found on the spines (40, 41). There can be as many as five or six synapses on a single spine, indicating that information could be integrated by a tremendous amount within the spine before being passed on to the dendrite shaft.

The density of spines over a given length of dendritic surface can vary widely. Many investigators have illustrated that changes in the density can occur under several conditions; for example, with prenatal ionizing radiation (42), undernutrition (43), or alterations in the amount of light entering the eye (44, 45, 46, 47, 48). But we were mainly interested in reports of changes in spine density associated with variation in the complexity of the environment (48, 49, 50) and with the age of the animals (51, 52). In order to study spines, a collaborative effort between the Berkeley laboratories and the University of California at Irvine (49) was established. Spine density was measured on pyramidal cells in the occipital cortex of male S₁ rats. Four replication experiments were designed to show the effects of 30 days (between 25 and 55 days of age) of differential experience between enriched and impoverished conditions on pyramidal cell spines in 40 littermate pairs. The spines were counted using the rapid Golgi method, which clearly demonstrates their structure. Also the lateral width of the basal dendrites was measured. The soma of the pyramidal cells being measured were in cortical layers IV and V.

The results of the counts showed that the density of spines on the basal dendrites was 10% greater ($p < 0.01$) in the enriched than in the impoverished rats (see Figure 19). The oblique dendrites, which arise from the apical dendrite, and terminal dendrites showed barely significant spine changes in favor of the enriched over the impoverished (4% and 3%, respectively, $p < 0.05$). No significant environmental effects were measured in spine densities on the apical dendrites.

These were the first results to indicate dendritic spine changes in response to stimulating experiential environments, but unfortunately, we made no attempt to differentiate between types of spines; all visible spines were counted. Later we were interested in quantifying types of spines, because of their different aging patterns, but the emphasis was on the effects not of enrichment but of impoverishment. For these studies, Connor counted two types of dendritic spines in three different age groups (53). First, spines were counted on dendrites from animals living in standard colony conditions from age groups of 90, 400, and 630 days. Spines

with a "lollipop" configuration (a long stalk with a bulbous ending) decreased from 90 days to 400 days of age and then increased in density until 630 days of age. On the other hand, spines with a "nubbin" configuration (short, stubby, as wide at the top as at the bottom) were lowest in number in standard colony rats at 90 days of age and increased in density at every age period to a high at 630 days, the last age measured. Evidently, the two populations of spines have different aging patterns in the standard colony animal.

Next, Connor counted the two types of spines on dendrites from impoverished animals. The most consistent finding was a greater density of nubbin spines on all dendritic segments (oblique, apical, and basal) (see Figure 19) from the impoverished rats than from standard colony rats. Since both the impoverished environment and the standard colony aging process increased the number of the nubbin spines, it seems that this configuration might represent a degenerating spine or a spine remnant. Such an idea was proposed by Mervis (54) in his study of spines in the aging dog brain, where he found that the nubbin-type spine increased with age.

The lack of a long, thin stalk and a bulbous expansion that is characteristic of the nubbin spine could cause a significant reduction in the integration of afferent information passing into a given dendritic tree. It has been suggested that lollipop spines function to attenuate the post-synaptic potential, since the impulse would have to pass through the thin stalk to reach the main dendritic shaft (55). In addition, the thin stalk of the lollipop spine would effectively isolate the postsynaptic site from potential changes in the dendritic shaft (56). However, the exact significance of the nubbin-type spine is not apparent. Whether this type of spine is an actively degenerating spine, a spine remnant, or an immature spine (signifying an area of increased synaptic turnover) remains to be determined. But the fact remains that it is present more frequently in the aged and impoverished rat brain and consequently signifies some association with reduced stimulation of the nerve cells.

Besides measuring the effects of impoverished conditions and aging on the relative density of nubbin- and lollipop-type spines, Connor set out to determine whether spine density or type varied between superficial and deep layers of the cortex in impoverished animals. The fluctuations found in the density of spine types between neurons from different cortical layers suggest the need for treating the data from the various layers separately. For example, oblique dendrites in layer II had a greater density of nubbin-type spines on neurons from the impoverished animals than on those from the standard colony animals but this finding did not hold

in layer III. In other words, a reverse pattern existed between layers II and III. The same discrepancy occurred for the basal segments in layers II and III. It is possible that if the results from layers II and III had not been treated separately, the results would have canceled each other.

The importance of comparing the effects of environmental input separately for the different cortical layers was also evident in layers Va and Vb, where a distinction in the response pattern between the standard colony and the impoverished animals was observed. Neurons in layer Vb from isolated rats had more lollipop-type spines on oblique segments than those of standard colony rats. This finding, coupled with similar data on dendritic branching (57), indicated an increase in the receptive area of neurons in layer Vb from impoverished rats compared with the standard colony rats. The same finding was not true in layer Va. Lorente de Nò (58) observed that axons from layer Va remained in the cortex, whereas axons from layer Vb left the cortex. From this information, one can speculate that the isolated rats can exhibit only a direct input-output type of behavior, whereas the standard colony rats can integrate the information entering their cortex before producing an output. In other words, in the standard colony rats more neuronal connections are involved before the neuronal output takes place.

With the exception of the above example, the lollipop-type of spine was remarkably constant on the dendrites from neurons from rats exposed to the two experimental conditions, the standard colony and the impoverished. This type of spine was much more prevalent than the nubbin-type on all dendritic segments in all layers of the occipital cortex.

To indicate the reliability of the spine counts (i.e., the total spine density per micron of dendrite), our study was compared with the only other quantitative spine study in old rats, that of Feldman and Dowd (52). The results indicated an almost exact agreement for basal (0.5 vs. 0.56) and oblique (0.07 vs. 0.76) dendritic spine densities. The apical shaft spine density per micron for our study (0.77) was well within 1 standard deviation of the apical shaft spine density of 0.92 reported by Feldman and Dowd. It is amazing how consistent spine counts can be when taken in separate laboratories across the country. This consistency offers a great reassurance for quantitative morphological studies of this type.

If spines can be changed in their morphological configuration after 30 days of environmental alterations, one wonders in how short a period does a spine change its dimensions. Van Harreveld and Fikova (59) shed some light by comparing spines on the same dendrites of hippocampal granule cells after some had been stimulated and others had not. Two

minutes after 30 seconds of stimulation, noticeable spine enlargements were demonstrated. These investigators suggest that the swelling was caused by a release of glutamate from the intracellular compartment into the extracellular space, which caused an uptake of water and electrolytes. If such increases were maintained long enough for protein synthesis to occur, this process might possibly explain spine growth as a result of experience. On the other hand, what happens when the spine is not stimulated or is deprived? The spine might be completely reabsorbed into the membrane of the dendritic shaft, or the molecular configuration of the membrane may become permanently changed at the position of reabsorption. If the latter is the case, then regrowth of spines may more readily occur at a previous site.

Thus, it is clear that dendritic branches are altered as predicted by Cajal in 1895 and by us from our results in 1964. The dendritic branching patterns of stellate cells in the occipital cortex were different between the enriched and impoverished animals. In our studies of the effects of enrichment on dendrite branching in pyramidal cells from the occipital cortex (area 18), we found increases in the number of middle branches—primarily third- and fourth-order branches—of basal dendrites in young and adult rats; and in older rats we found increases in the length of the terminal branches.

Dendritic spine counts showed that on the basal dendrites of pyramidal cells from young enriched rats there were significantly more spines than on dendrites of the nonenriched. Although similar differences were also seen on oblique and terminal dendrites, they were not as great. Types of spines were not counted separately in the young animal brains, but they were on the older ones, and in these it was the "nubbin" types, those with a base as broad as the head, that were more prevalent in the impoverished animals than in the standard colony animals. The "lollipop" types, with a thin stalk and large head, were more frequent in the older enriched animals. The evidence is clear that down to the level of the spines, the nerve cell shows great structural adaptability to its environment at any age studied this far, at least two-thirds of the way through the lifetime of a laboratory rat.

While teaching a group of 13- to 15-year-old youngsters in Shanghai about our work on the brain, I called on one poised young lady who stood up and asked the following question: "If most of our creative work is done before 40 and yet wisdom does not come until after 60, how do we explain this with regard to nerve cells?" Obviously, I had no real answer, but I immediately drew on the knowledge from our data. If I were forced to speculate, I would say that results from enriched-

impoverished experiments tell us that it is the third- and fourth-order dendrites which increase in dimensions in the young rats. These are closer to the cell body and have a more limited expanse than the terminal dendrites and therefore may be dealing with more focused-creative activity. In the old rats, it is the terminal dendrites which show increases in length; they are broadly spaced and reach out widely. With such a massive extension, they sample more diffusely, making broad comparisons, and thus, allow for wiser judgment. The specific functions of the various levels of dendritic branches are not known at present; only further work will indicate whether there is a grain of truth in my speculation.

SYNAPSES. Since we had shown that both dendrites and spines change in response to environmental conditions and since both are sites of synaptic action, our next step was to study the effect of the environment on the dimensions of the synapse. It had been suggested that permanent changes occur in synapses as a result of stimulation and that these changes play some part in learning and memory (60). As we did in our earlier work when we were identifying ovarian hormonal effects on the synapse, we again chose to measure the length and number of postsynaptic thickenings, those structures which consist of a highly organized proteinaceous material immediately beneath the postsynaptic membrane.

Since we were ready to take a step in the direction of synaptic measurements when a medical doctor Møllgaard came to join our laboratory from the University of Copenhagen in Denmark, it seemed a reasonable project to offer to him. For the electron-microscopic study of synapses, we separated 12 newly weaned littermate pairs of 25-day-old male rats from the S₁ strain, placing one rat from each pair in the enriched condition and its littermate in the impoverished conditions. The rats were kept in the respective conditions until they were 55 days of age. Through all the anatomical procedures that followed, from sacrifice to completion of the measurement of synapses, the rats were studied as pairs. We sampled synapses in layer III of area 18 in the occipital cortex (61).

Measurements were made on the asymmetrical axodendritic synapses described by Colonnier (62). The great majority of dendritic spines are associated with asymmetrical synapses. The asymmetry of this type of synapse is characterized by the presence of postsynaptic densities of variable thickness that border on the postsynaptic membrane but not on the presynaptic membrane. Møllgaard measured only those junctions which reached an acceptable criterion of clarity in the photomicrographs. Curved thickenings were divided into smaller parts and the sum of the parts was taken. In some cases a large synapse had one or more discontinuities in the postsynaptic thickening; if the vesicles were uniformly dis-

tributed on the presynaptic side, the thickening was measured as one synapse in spite of the discontinuities. Serial sections and reconstructions have shown that such perforations are normally present in larger synapses (63). More recently, Greenough et al. (64) and Medosch and Diamond (65) reported that these discontinuous postsynaptic thickenings may represent more mature synapses, as discussed earlier. All of the synaptic measurements were done on coded prints so that the criteria of selection of synapses did not vary between the experimental groups.

In all, 2211 synapses were identified and measured, 1405 from the impoverished rats and 806 from the enriched animals. The number of synapses per print was 35% smaller for the enriched rats than for the impoverished in every pair. Though the enriched rats showed fewer asymmetrical synapses than the impoverished, the enriched synapses on the average were larger. After a cumulative frequency distribution of synaptic lengths was plotted for all enriched and impoverished rats, it was obvious that the enriched rats had significantly fewer small synapses than the impoverished.

The total area of enriched and impoverished synapses was calculated, on the basis of measured cross-section diameters and numbers of junctions; that is, we assumed that the junctions were roughly circular. We calculated the area for each size of cross section, and then multiplied by the number of cross sections per size. From these calculations, it appeared that the total area of the synaptic thickenings in the enriched exceeded that in the impoverished by 40%.

There was a 4.0% difference ($p < 0.05$) in cortical thickness between the enriched and impoverished rats in this experiment, which was close to the 5% mean found in four previous experiments on rats that had lived in their conditions in the period from 25 days to 55 days. A high positive correlation (0.79, $p < 0.001$) was found between cortical thickness differences and synaptic length differences. The correlation was not significant between the differences in cortical thickness and the number of synapses.

Scientists have long speculated that long-term memory might involve changes in synaptic size and number. Almost as soon as the neuron doctrine was enunciated near the end of the nineteenth century, Tanzi (66) speculated that learning might involve the growth of new neuronal terminals, and Ramon y Cajal (67) supported this hypothesis. Hebb (68) postulated that learning might occur through formation of new synaptic connections. The opposite possibility, that learning involves selective elimination of neurons or synaptic connections or both, has received less attention, although Ramon y Cajal (69) felt that in embryological devel-

opment there was a random selection among connections. In 1971 Dawkins (70) suggested that the relatively high continual rate of death of neurons did not occur at random but might be a mechanism of memory storage leaving those connections activated during memory consolidation. Other investigations (71) have considered whether learning and memory involve "growth just of bigger and better synapses that are already there, not of growth of new connections" (71). There were also those who doubted whether learning and memory involve any morphological changes; perhaps there are "only changes in physiological resistance and conductance . . . or various endogenous properties of neurons and glia" (72).

Our data demonstrated the presence of bigger synaptic junctions in the enriched than in the impoverished rats. The question which commonly arises in response to such findings is, Are bigger junctions more effective? It has been reported that the intensity of postsynaptic current increased with the local concentration of transmitter molecules and of available receptor molecules (73). If one hypothesizes that the amount of neurotransmitter liberated from the presynaptic terminal is proportional to the amount of membrane in which the receptors are localized, then the effectiveness of the synapse would increase as the size of the postsynaptic membrane increased. All of the morphological data collected on the enriched and impoverished rats support this hypothesis, as do the maze-testing data, which show that the enriched rats are better learners in many problem-solving situations.

Having found that the differences in synaptic measurements between the enriched and impoverished rats were very clear-cut, we had to ask ourselves, Was the difference in synaptic length mainly due to enrichment or impoverishment? In order to answer this question, another set of experiments (74) was designed to include not only the enriched and impoverished rats but the standard colony as well. We performed both initial and replication experiments, using littermate S₁ male rats that had lived in their conditions between 25 and 55 days of age, to replicate the setup of the original experiment we had done with Møllgaard. Measurements of the length of postsynaptic thickenings were taken from asymmetrical axodendritic synapses in layer IV of the dorsal medial occipital cortex (area 18). It should be noted that in the Møllgaard experiment it was synapses from layer III that were measured. After the publication of the results of the Møllgaard experiment (61), West and Greenough (75) reported that the enriched animal's synapses were longer by 10% in layer IV in cortices from four enriched-impoverished pairs. Since we were having some difficulty in replicating the large synaptic changes reported in

Møllgaard's results in layer III, we decided to increase the size of the sample used by West and Greenough and study layer IV synapses in our new experiment, as well as to include rats from the standard colony condition.

In the new experiment, we compared 675 synapses from enriched animals, 618 synapses from standard colony animals, and 680 synapses from the impoverished rats. We found an 8% difference ($p < 0.001$) in postsynaptic thickening length between the enriched and the impoverished rats that was primarily due to the effects of the impoverished condition. The number of synapses per unit area of neuropil was 15% more ($p < 0.01$) in the impoverished than in the enriched, and this difference was primarily due to impoverishment. The synapses in the impoverished brains were more in number but smaller—in essence a result similar to that found in the Møllgaard experiment but with a different emphasis where he noted fewer but larger synapses in the enriched compared to the impoverished.

A 7% ($p < 0.01$) cortical depth difference in area 18 was found between the enriched and impoverished animals used for synaptic measures; we could attribute it to either condition by comparing each with the standard colony rats. The effect of enrichment was about 4% ($p < 0.01$) and that of impoverishment about 2% (NS); but if the rats were older when entering their respective conditions, staying in them between the ages of 60 and 90 days, the effect of enrichment in area 18 was about 6% ($p < 0.001$) and that of impoverishment about 1%. The impact of enrichment is greater in the occipital cortex if the animals are older when placed in the enriched environment, while the effects of impoverishment are more significant earlier in life.

Thus, differential environmental experiences do significantly affect cortical thickness and synaptic length and number in the age groups measured in these experiments. In the Møllgaard experiment dealing with synaptic measurements, the differences between enriched and impoverished conditions were much greater than those found in our later experiments. In the Møllgaard experiment, layer III synapses were quantified, and in the more recent experiment it was the synapses in layer IV which were measured. In an earlier section, it was mentioned that the greatest differences in neuron dimensions between experimental groups were in the uppermost layers of the cortex. Is it possible that such large differences in synaptic dimensions as seen in the Møllgaard experiment were actually due to the presence of larger neurons found in layer III than those reported by West and Greenough and by us in layer IV? In the Møllgaard report, more synapses per picture were found in the impover-

ished rats than in the enriched ones. This finding was also encountered in the present investigation in layer IV, offering confirmation of some of the measures. Yet, Cummins (76), working with mice, has found a greater number of synapses in the enriched than in the impoverished. When we learned this, we asked to share photographs and compare measuring techniques and synapse identification criteria between our two laboratories. We could find no differences between our techniques, only in our results.

Aghajanian and Bloom (unpublished data), in a collaborative experiment with us in the late 1960s, found no synaptic changes in layer I between enriched and impoverished rats. We were not surprised by these findings, because layer I had not previously shown thickness changes in response to our environmental conditions. However, in 1972 West and Greenough showed synapses with greater lengths in layer I in enriched rats—thus adding ambiguities to these studies.

Like much of our work, our study of synapses requires additional data; ideally, we should compare synapses in all layers of the cortex, including layer I, in the same brains from animals exposed to these differential environments. Clearly, synapses are structurally modified by these environments, but we need much clearer and more detailed information on *how* they are modified before we can draw definitive conclusions. No one expected that the measurement of synapse numbers and dimensions would be simple—at least with regard to subtle environmental alterations. Severe environmental insults—such as reduction of visual input to the visual cortex—have been much easier to quantify. Synapses evidently change constantly, and to pick up the fine changes takes great patience and skill. With modern techniques using computer-integrated measuring devices, we should obtain better results in the future.

Other Brain Structures

Over the years, as we learned how readily the cerebral cortex responds to our external environmental paradigm, we wondered what other brain structures were similarly affected. Such structures as the corpus callosum, entorhinal cortex, hippocampus, amygdala, lateral geniculate body and cerebellum were examined in animals exposed to differential experiences.

Except for one area, lateral 10, of the frontal cortex, the increase in cortical thickness in our experiments was essentially the same in both hemispheres. Since the two sides were affected equally, we anticipated finding that enrichment causes an enlargement of the corpus callosum, the large fiber tract connecting the right and left cerebral cortices.

Though the tract has not been measured directly in our laboratory, there was one experiment that might indicate an increase in either size or number of callosal fibers or both. In the late 1960s, Leon Dorosz (77) set out to determine whether a transcallosal electrically induced response in the enriched cortex was different from that in the impoverished cortex. In his first experiment, he implanted electrodes directly into the cortices of the enriched and impoverished rats to record chronic activity at various times, only to learn that within a few days the enriched rats had chewed out each other's implants. (Group living and this kind of surgical intervention were not compatible.) He then performed acute preparations in which stimulation and recording were carried out directly from the brains just prior to the termination of the experiment. On one side of the cortex, he used a stimulating electrode, and on the other, he picked up the transcallosal response with a recording electrode. With this technique, he was able to demonstrate a significantly larger peak-to-peak amplitude in the occipital cortex from the enriched than from the impoverished rats after they had been living in their respective conditions for 80 days.

These results can be interpreted in one or both of the following ways: (1) the larger signals Dorosz was picking up resulted from the increased cellular dimensions in the occipital cortex, or (2) the fibers in the corpus callosum had increased in size or number. That they were due to an increase in callosal functioning, as suggested by Dorosz, was later substantiated by the work of Walsh et al. (78), which actually demonstrated a thicker callosum in the enriched rats than in the impoverished rats. Walsh et al. found that in the frontal region, but not the occipital, the lateral but not the medial, callosum was 14% thicker ($p < 0.01$) following environmental enrichment. With the hundreds of prepared tissues from rat brains in our experiments, this is one investigation we could easily duplicate.

All of our previous studies on enriched and impoverished environments have dealt only with the cerebral cortex and some of its related fiber tracts. But this does not mean that we have not tried to find morphological changes in response to the environment in other regions of the rat brain. None of the fifteen other areas of the brain which were sampled but the cerebellum gave a significant indication of a specific localization of an enrichment effect as measured by tissue weight or acetylcholinesterase.

Nonetheless, we continued to look for anatomical changes due to altered environments in various regions of the forebrain, because there was a chance that small alterations might be evident that were not noted

with previous wet weight or chemical measures. It seemed likely that other regions known to be involved in learning and memory should show differences, and so we obtained morphological measures for the entorhinal cortex and hippocampus as well as other forebrain regions, including the amygdala and the lateral geniculate body, the first three regions having been discussed previously with regard to growth patterns in Chapter 2.

Our attention was drawn in particular to the entorhinal cortex and the hippocampus because of their role in learning and memory. It was clear that the enriched condition facilitated learning in some way (for instance, the enriched animals were better maze learners), and so we expected that we might find some changes in these two forebrain areas. (See Figure 2 for identification of the areas and the manner in which their dimensions were determined.) We learned that the male entorhinal cortex exhibited a definite increase in thickness in response to the enriched conditions for two age groups, 60 to 64 and 60 to 90 days. However, in measuring the hippocampus on the same tissues used for both occipital and entorhinal cortical measurements, the environmental conditions did not significantly change the thickness of the hippocampus, for either the 4 or 30 day duration. In fact, totaling the results of our hippocampal thickness measures on as many as 231 pairs of male rats from enriched and nonenriched conditions, at no time were the differences significant. Since the entorhinal cortex showed changes in response to the environment and because it sends a major fiber tract into the hippocampus, we are still puzzled why changes of the magnitude found in the former did not create some measurable changes in the latter. We thought, perhaps, our measurements were not sensitive enough to detect the small changes in the hippocampus.

However, other investigators have noted some alterations by confining their measures to only the granular cell layer in the female hippocampal complex (79). They did find the anticipated increases resulting from the enriched environmental conditions, but only in the females. We looked for thickness differences in this layer in our oldest rats, the 904-day-old male rats, which are losing their asymmetry patterns to become more like the female pattern. Still we found no significant differences between male enriched and nonenriched rats.

These data suggest that there is a sex difference in the response of the hippocampal complex following differential experience. Since we do have brain sections from female rats exposed to either enriched or impoverished environments, we can measure the hippocampal complex in these brains to learn if our results concur with those of Juraska et al.

By providing greater detail on the changes in the male entorhinal cortex, we can see just how responsive it is in contrast to the male hippocampus. In our experiments S_1 males were maintained in enriched, standard colony, or impoverished conditions either between the ages of 60 and 64 or between 60 and 90 days of age. The entorhinal cortex showed a thickness 3% ($p < 0.01$) greater in the enriched rats than in the standard colony rats in as short a period as 4 days. The increase in thickness that occurred in this short period was of the same order of magnitude as the change that appeared during the 30-day period between 60 and 90 days of age. The entorhinal cortical change at 60 to 90 days was slightly less at 4% than the increase in thickness found in the medial occipital cortex at 5%. These results proved that the entorhinal cortex could show a definite structural change in response to our environmental conditions. Since the entorhinal cortex is known to be part of a memory pathway, this finding offers encouragement about the possibilities of improving one's memory through enriched conditions.

Scoville and Milner reported that the entorhinal cortex is related to memory in human beings and also stressed the importance of the anterior hippocampus and the hippocampal gyrus, which includes the entorhinal cortex, in retention of new experience (80). They claimed that if the medial temporal lobe was bilaterally removed in man, and if the removal of tissue extended far enough in the posterior direction to damage portions of the anterior hippocampus and hippocampal gyrus, a persistent impairment of recent memory would result. Stepien et al. (81) reported that monkeys that had undergone bilateral ablation of the hippocampus and hippocampal gyrus displayed a recent memory deficit in vision and auditory tests. These findings are supported by Drachman and Ammannya (82), who reported impairment in new-pattern discrimination in monkeys with lesions of the hippocampus and hippocampal gyrus. In the famous H. M. case so thoroughly studied by B. Milner in Montreal, bilateral removal of the temporal lobe caused him to lose recent memory permanently. And a study of eighty-five human brains from people with Alzheimer's disease, where clinical manifestations of recent-memory deficits are marked, revealed neurofibrillary degeneration within the neurons of the lateral entorhinal portion of the hippocampal gyrus, as well as in an area of the hippocampus (83). All of these reports made it seem imperative to understand the effects of an enriched or impoverished environment on the separate parts of the hippocampal complex.

In light of the close proximity of the amygdaloid nucleus to the hippocampus and entorhinal cortex in the temporal lobe, we continued our pursuit and examined the effects of our varied environments on this nu-

cleus (84). We undertook this study even though we knew that bilateral excision of the tip of the temporal lobe, which includes the amygdala (another term for the amygdaloid nucleus) but not the hippocampus and entorhinal cortex, did not harm memory function in any way, indicating that the amygdala is not primarily concerned with memory processing. Nonetheless, the amygdala has been associated with both active and passive avoidance learning (85). In addition, because it is known to mediate arousal, we thought that it might vary among our environmental groups since animals must be aroused to interact with their environment.

Studies have shown the amygdala to be associated with territorial behavior. Furthermore, Sherman et al. (86) and Webster (87) reported that preferential processing of spatial information in the right cerebral hemisphere in rats may be important for territoriality. The finding that both the right amygdala and the right visual cortex were larger than the left, may signify an interaction between the two areas in integrating territorial behavior. Thus, like the hippocampus, another older part of the forebrain, the amygdala does not vary in its dimensions as a consequence of changes in our environmental conditions, whereas the cerebral cortex readily does. A more plastic visual cortex could provide more variety in territorial behavior, while the amygdala's role is more stable.

In continuing our search for subcortical changes, we studied the lateral geniculate body within the visual pathway (88). This study was done on very young animals—28 days of age—that had lived either in a multifamily enriched condition (3 mothers with 3 pups each) or in a unifamily impoverished condition (1 mother with 3 pups) for 22 days. (The other subcortical measures were performed on animals at least 55 days of age.) Like the cerebral cortex in these young rats, the lateral geniculate body manifested a significant response to the environmental conditions: 20% fewer neurons per unit area in the multifamily enriched rats than in the unifamily impoverished rats ($p < 0.01$). These results indicated that the neuropil, or fibers between the neurons, had increased in either size or number to “push” the soma or cell bodies, of the neurons farther apart. Here, then, was one subcortical structure that showed measurable variation with the degree of enrichment in the environment. However, because the lateral geniculate was measured in animals who had not yet been weaned and the other subcortical areas—the hippocampus and amygdala—were examined in animals who had been weaned, it is possible that the plasticity of subcortical regions depends upon age, declining as the animal ages.

In our attempt to localize more precisely the regions in the rat brain which were responding to the enriched condition, we mentioned earlier

in this chapter that the cerebellum of rats in that condition did increase in weight. As Walsh mentioned (89), on theoretical grounds the cerebellum might be expected to play a role in response to enriched environments because studies have pointed to the necessity of active self-guided exploration and kinesthetic feedback for learning to take place (90). The cerebellum plays an important role in maintaining the position of the body in space, providing a rationale for the above behavioral patterns.

Active interaction with the toys was essential for the enrichment effect to occur in the cerebral cortex in our rats (91, 92). The animals climb, explore and balance on the ladders and wheels—all activities requiring cerebellar control. On the basis of what is known about the connections between the cerebral cortex and the cerebellar cortex, again one might predict cerebellar responses to the enriched condition. Floeter and Greenough (7) did find changes in the more primitive parts of the young cerebellum in enriched animals compared with the nonenriched. Between the overall cerebellar weight changes of Rosenzweig et al. (93) and these specific regional alterations reported by Floeter and Greenough, one is led to believe that even a hindbrain structure such as the cerebellum could show plasticity in young adult rats.

But we have the opportunity to study the cerebellum of the very old rats (904 days old) to determine whether an environmental response can be measured. The number of Purkinje cells and the thickness of the outer molecular layer, which is made up primarily of the dendrites from the Purkinje cells, are being measured in these old animals. Any proof that plasticity can exist in the aged cerebellum will be most comforting news to elderly people who are developing problems in balancing and coordination. Therapies designed to stimulate cerebellar function could be rationally encouraged.

Asymmetry

In our first enriched-impooverished morphological experiment carried out in 1964, on animals from the S₁ strain that had lived for 80 days in their conditions (from 25 to 105 days of age), we found that both hemispheres were affected equally by the environmental conditions (94). But having shown in our developmental studies that in the male Long-Evans rat the right hemisphere was significantly larger than the left in many areas, we wondered whether one hemisphere would be affected more than the other if we changed the age of the rats and the period of exposure to their respective environments. We carried out right-left compari-

sions on several groups of animals of the S₁ strain from which we had other enrichment data, including additional rats from the 25- to 105-day groups. The ages in days at commencement and termination of exposure to the environmental conditions were as follows: 25 and 55; 25 and 105; 60 and 64; and 60 and 90. A total of 267 rats from these various age groups were studied for these right-left comparisons.

Analyzing these data from the S₁ strain of rats for these left-right studies was useful in confirming other factors which had been found separately from combined right-left data. The statistically significant differences among the three conditions (enriched, standard, and impoverished) were confirmed for all regions measured. There was a statistically significant change in cortical thickness with age in all areas; from youngest to oldest, we found a general decrease in thickness, as was noted in Chapter 2 in connection with the development and aging study using the Long-Evans strain.

We learned that even with exposure to the different experimental conditions and for different periods of time, both hemispheres were still affected equally except for area 10 in the frontal cortex. Area 10, according to Krieg (95), is an association cortex; but according to Paxinos (96), it is also a motor cortex related to movement. We found that in medial area 10, the difference between the right hemispheres in the enriched and standard animals did not achieve statistical significance, although the right hemisphere differences between animals living in the standard and impoverished conditions did. It was the impoverished condition which decreased medial area 10 in the right hemisphere of these male rats. If one accepts the report of Paxinos, that the frontal cortex governs motor function then it is possible that without physical activity, as the rat lives in an impoverished condition, the right hemisphere decreases in dimensions. (It would be of interest to test whether the impoverished rats had changed a paw preference compared to the standard colony animals.) However, for the remainder of the cortex, one can conclude, in general, that with the multisensory environment provided by our type of enrichment, the hemispheres in the S₁ strain of rats are apparently affected equally.

These results do not lend support to Denenberg's report (97) on the effect of infantile stimulation and environment on brain lateralization in rats. He stated that the right brain was the repository for the interaction between environmental enrichment and handling in open-field activity. He concluded that lateralization of behavior occurred, and the effects of early experiences were asymmetrically distributed in the rat's brain. It is possible that lateralization of specific kinds of behavior occurs, but that

the neuronal protein changes which record an activity in the brain involve neuronal interplay between both hemispheres.

The results with the S₁ strain of male rats supported our previous findings with the Long-Evans strain showing that the right hemisphere was thicker than the left in the majority of comparisons. In the male S₁ rat, the right hemisphere was on the average thicker than the left in all areas except 18, where the left was thicker than the right, and area 2, where the hemispheres were equal. In areas medial 10, 4, 18, 17, 18a, and 39, the hemispheric differences were statistically significant. A particularly striking aspect of our findings was that the effects were seen in a large sample rather than only among a few individual animals.

In our studies we have been particularly interested in area 18 of the cerebral cortex, because this area is most frequently affected by the enriched condition. In our shortest duration of exposure to enrichment, from 60 to 64 days of age, it was area 18 which showed the only significant increase in cortical thickness over standard colony littermates. In addition, for all long durations in the environmental conditions, area 18 always showed a significant change, whereas other areas were not so consistent. As we search for a meaning to the susceptibility of this area to the environment, we noted that area 18 in the left hemisphere of the S₁ strain always showed a tendency to be thicker than in the right irrespective of the environmental conditions. This area is known to be a visual association area, but many subcortical regions project to this area. Examining our most recent data from crowded-enriched Long-Evans rats (36 enriched rats rather than 12), we find that area 18 shows an even greater difference in favor of the left hemisphere than is seen in the standard colony rats.

This latter fact becomes useful as we try to gain some understanding of the meaning of asymmetry by following only one area in rats that come from one strain but from different housing conditions. If we examine the male Long-Evans rats that have been raised in the Berkeley colony for over 50 years, we find very marked right-greater-than-left differences in area 18. If we look at right-left differences in Long-Evans rats from our local distributor, where the housing conditions are not identical to those at Berkeley, we find that area 18 in the left cortex is thicker than in the right. If we enrich these Long-Evans rats from the local distributor, we find that area 18 begins to show significant left-greater-than-right differences. But most important, if we now provide an environment that is both enriched and crowded, the left-greater-than-right asymmetry pattern in area 18 becomes very significant. These findings were present in both an initial and a replication experiment. It appears

that asymmetry patterns are subject to change in response to conditions as nonspecific, yet pervasive, as housing density.

Skull Size

Now that we had clearly demonstrated that the cerebral cortex varies in size in response to environmental variations, we wondered whether the intracranial capacity increases to accommodate an increased brain mass. We conducted experiments to study the skull dimensions of rats living in the different environmental conditions. The animals were exposed to their respective conditions—enriched or impoverished—for 77 days, from the age of 25 days to the age of 102 days. Roentgenograms for the initial and replication experiments were taken 8 days after the animals entered their respective conditions and again 69 days later (98). For our measurements, we used the methods established by Asling and Frank (99). The landmarks employed for the skull measurements on the Roentgenograms are shown in Figure 20. The results showed that the intracranial capacities of the two groups did not differ significantly. Evidently the two diverse environments did not alter internal cranial dimensions.

As was pointed out in Chapter 2, the cortex of the Long-Evans male laboratory rat reaches a peak in development sometime between 26 and 41 days of age and then begins to decline. We have one experiment (100) comparing the cortical thickness measurements of a baseline group at 112 days of age with those of animals from the standard colony and from the enriched group at 142 days of age. In this experiment we learned that growth of both the cerebral cortex and the dendrites did exceed the baseline as a result of enrichment. Here was evidence that enrichment actually caused the young adult cortex to grow and did not merely slow down or arrest a decrease in size. We can deduce from these data that an increased brain mass does accumulate with enrichment and as a consequence more intracranial space might be necessary. But our intracranial measurements show something different. It has been my fantasy that with further enrichment during early adulthood, an increase in cortical dimensions might produce the first cortical fold in the rat to accommodate more mass in a limited space. Cats, dogs, monkeys, and man all have folded cortices. Brain growth had to exceed intracranial dimensions to produce folds. There is no evidence of folds in the rat cerebral cortex. We need to find out more about the formation of the cranial cavity by examining the time factors involved in the formation of the sutures as related to brain development.

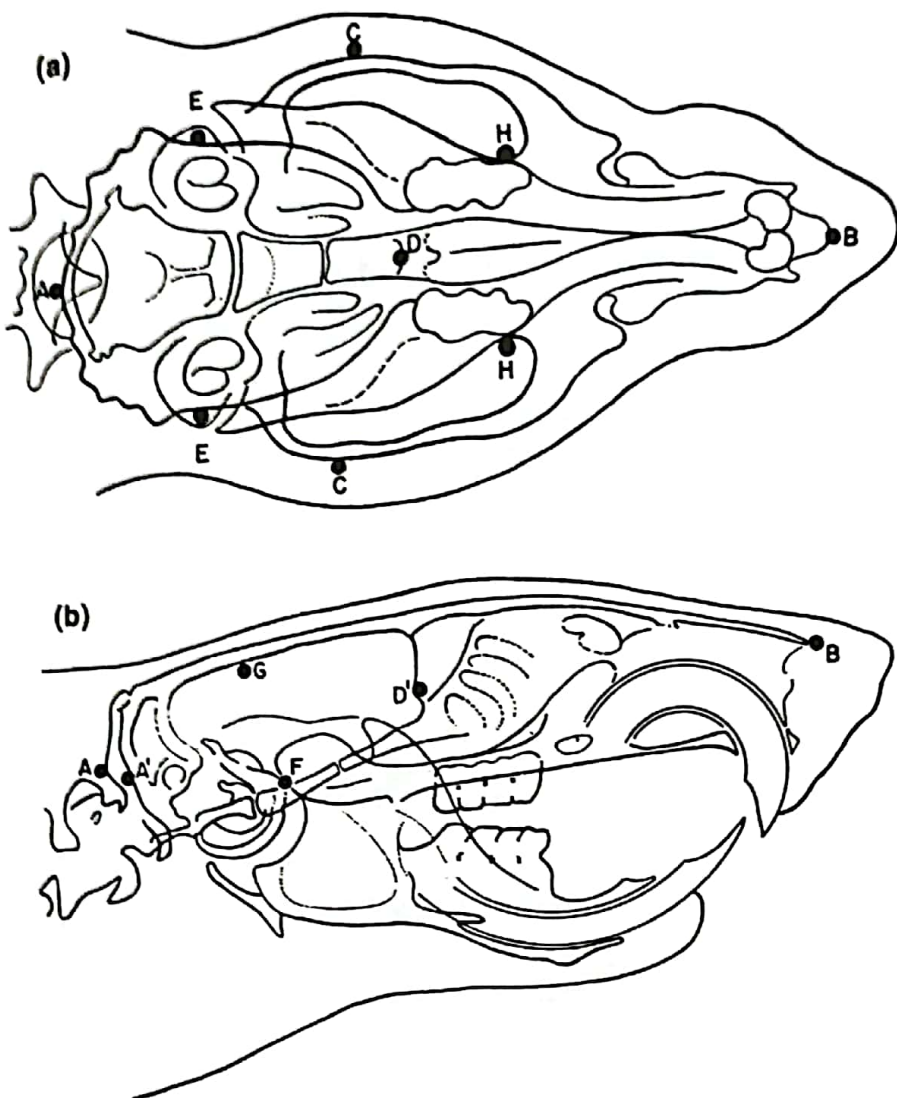


FIGURE 20. (a) Dorsal view of rat skull. (b) Lateral view of rat skull. Total skull length (A-B); maximum width (bizygomatic) (C-C); length of ventral neurocranium (A-D); external width of neurocranium (E-E); internal height of neurocranium (F-G); snout length (D-B); maxillary width (H-H); and lateral length of neurocranium (A'-D'). Neurocranium is that part of the skull which houses the brain.

According to Massler (101), the rat attains its maximum cranial width by 20 days of age, with the internasal and interfrontal sutures completed by 3 to 10 days and the sagittal suture completed by 20 days. The sutures of the calvarium, which are primarily responsible for cranial length, are completed by 40 days. From these data, it appears that the cranial size in our rats should have been well established approximately 15 days after the rats were put into the experimental conditions at 25 days of age.

However, Weidenreich (102) suggested that the problem cannot be solved merely on the basis of the conditions of the sutures. He pointed