

Neocortical Development

Shirley A. Bayer, Ph.D.

*Department of Biology
Indiana University and
Purdue University at Indianapolis
Indianapolis, Indiana*

Joseph Altman, Ph.D.

*Department of Biological Sciences
Purdue University
West Lafayette, Indiana*

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AUTHORS' NOTE:

This book is out of print. Raven Press no longer exists. The copyright to this book is being transferred to us. We want to make this information available to all who are interested in neocortical development. In the files available for download, we add a few pages that review some of the relevant literature that has been published since 1991. Our observations in 1991 are relevant to the multitude of gene expression studies done since the mid-90s and up to the present day. Many of the hypotheses that we proposed in this book have been confirmed by these studies. We also re-examine cell migration in the neocortex; the labeling patterns in our autoradiograms are consistent with studies showing tangential migration of GABA-ergic cells from germinal zones in the basal ganglia into the developing neocortex. We invite your comments on our work, and would like to answer any questions that you want to ask about how your research correlates with our findings.

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Preface

The neocortex is the crown of the mammalian central nervous system both literally and figuratively. Because an animal can carry out many of its vital functions after structural or functional decortication, the neocortex is not essential for survival in the strict sense of the term. However, there is an abundance of direct and indirect evidence that suggests that the neocortex plays a crucial role in the control of higher perceptual processes, cognitive functions, and intelligent behavior. Ascending the phylogenetic scale, the six-layered neocortex steadily expands relative to the rest of the brain, a phenomenon referred to as progressive neencephalization. This expansion first becomes manifest as the smooth neocortical mantle (lissencephalic pallium) spreads over the rest of the forebrain and the midbrain. A later manifestation of progressive neencephalization, one especially pronounced in larger mammals, is the increasing convolution of the cortical surface. This brings about an increase in the ratio of nerve cell bodies and dendrites relative to the cortical afferents and efferents, reflecting gains in processing capability and computing power of the cortical gray matter. In parallel with progressive neencephalization, there is an increase in the number of different cytoarchitectonic subdivisions of the neocortex, particularly of those regions traditionally referred to as "association areas." It is widely assumed that the evolutionary growth of mental life that reaches its zenith in humans is attributable to the progressive expansion and elaboration of the neocortex.

Neuroanatomists have been studying the structural organization of the neocortex for centuries. This began with dissections aided by the naked eye and was followed by light microscopic examinations of regional differences in neocortical cytoarchitectonics. There is still an effort to unravel the gross and fine circuitry of the neocortex using chemical, biochemical, and physiological tracer techniques at light and electron microscopic levels of resolution. The investigation of the functional organization of the neocortex became possible with the introduction of electrical stimulation and recording techniques in the last century, and advances in electronics and computer techniques in this century are helping us to understand how information is conveyed to, and processed in, the neocortex. Neuropsychologists using ablation techniques, as well as electrical stimulation and recording procedures with implanted electrodes in conscious animals, have contributed their share to our current understanding of how the neocortex controls behavior. However, the investigation of the morphogenetic development of the neocortex was lagging behind the anatomical, physiological, and psychological studies of its structure and function. Gross descriptions of the developing neocortex in animals and humans were beginning to become available at the turn of the century. But the initial efforts of neuroembryologists to analyze the cytological processes underlying neural development tended to focus on the relatively simple caudal portion of the neural tube that gives rise to the spinal cord. This state of affairs began to change in the second half of this century. A steadily growing number of articles are now being published in various journals that deal with the development of the neocortex in a great variety of species and at different stages through the embryonic, fetal, and early postnatal periods. A few edited volumes have also appeared that contain valuable contributions by experts who deal with some selected aspect of neocortical development in one or another species. However, to our knowledge there is not a book currently available, written by a single team of investigators, that specifically deals with the development of the mammalian neocortex as a whole.

This book is the outcome of our own current investigations on neocortical development in which we relied on a limited number of techniques and concentrated on a single species, the rat. The lissencephalic cortex of the rat is far less complex structurally than the convoluted cortex of a carnivore or primate. This represents an advantage from the experimental point of view because

a study of the entire cortical mantle is possible. The major techniques and materials that we have used are limited to the following: (a) We have prepared a large collection (several hundred specimens) of high quality, methacrylate-embedded Nissl-stained sections that cut through the neocortex in the three cardinal planes. This material was used, first, to describe daily changes in the growth of the neocortex at the histological and cytological levels, and second, as normative material for comparisons with observations made in experimentally manipulated embryos; (b) A large collection (several hundred specimens) of embryos that received injections of [^3H]thymidine to label multiplying precursors of neurons and glia were prepared for autoradiography. Three variants of this technique were used. First, long-survival autoradiography after multiple injections was used to determine the time of origin of different classes of neurons in different regions of the developing cortex on successive embryonic days in neonates and adults. Second, short-survival autoradiography (2 hours) after single injections was used to locate the sites of cell proliferation and their magnitude as a function of embryonic age. Third, sequential-survival autoradiography (consecutive 24 hour intervals) after single injections was used to follow the dynamics of cell proliferation in the cortical germinal matrices; to trace the migration and temporary pause of cells through the subventricular and intermediate zones of the cortex; and to determine the time of arrival, final distribution, and settling of young neurons in the cortical gray matter. Finally, we used low-level x-irradiation as an ancillary technique to study changes in the cellular dynamics of the germinal matrices giving rise to neurons and glia, and to investigate certain aspects of neuronal differentiation. Because our techniques are best suited to study developmental change during the embryonic period at the histological level, developmental changes during the postnatal period at the cellular and subcellular levels, i.e., dendritic differentiation and synaptogenesis, have not been investigated.

*Shirley A. Bayer
Joseph Altman*

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Abbreviations

a, A	anterior	FM	foramen of Monro
abt	anterior basal telencephalic neuroepithelium	FR	motor area
ABT	anterior basal telencephalon (differentiating)	FR1	frontal cortex, area 1
AC	anterior commissure	FR2	frontal cortex, area 2
AD	anterodorsal thalamic nucleus	FR3	frontal cortex, area 3
AI	agranular insular cortex	fu	site of fusion of prosencephalon
AID	agranular insular cortex, dorsal part	GCC	genu of corpus callosum
AIP	agranular insular cortex, posterior part	GU	gustatory cortex
AIV	agranular insular cortex, ventral part	h	head of lateral cortical stream
AL	anterolateral	HC	hippocampal commissure
AM	anteromedial thalamic nucleus	hi	hippocampal neuroepithelium
ap	alar plate of neuroepithelium	HI, HP	hippocampus (differentiating)
APO	anterior primary olfactory cortex	HL	hindlimb motor area
AV	anteroventral thalamic nucleus	hy	hypothalamic neuroepithelium
bg	basal ganglia neuroepithelium	HY	hypothalamus (differentiating)
BG	basal ganglia (differentiating)	ib1	first inferior band
bp	basal plate of neuroepithelium	ib2	second inferior band
bt	basal telencephalic neuroepithelium	ibt	intermediate basal telencephalic neuroepithelium
BT	basal telencephalon (differentiating)	IBT	intermediate basal telencephalon (differentiating)
caz	callosal zone	ic	inferior collicular neuroepithelium
cc	cerebral cortical neuroepithelium	IC	internal capsule
CC	corpus callosum	ICP	insular cortical plate
ce	cerebellar neuroepithelium	IL	infralimbic area
CG	cingulate cortex	inr	infundibular recess
CG1	cingulate cortex, area 1	iz, IZ	intermediate zone
CG2	cingulate cortex, area 2	izl	lower intermediate zone
CG3	cingulate cortex, area 3	izm	middle intermediate zone
ch1, Ch1	channel 1 of developing cortex	izu, IZu	upper intermediate zone
ch2, Ch2	channel 2 of developing cortex	l, L	lateral
chp	choroid plexus rudiment	lcs, LCS	lateral cortical stream
CL	claustrum	LL	lateral limbic cortex
col	collapsed neuroepithelium	lo, h	low, heavy (labeling pattern in late cortical neuroepithelium)
CP	cortical plate	lo, l	low, light (labeling pattern in early cortical neuroepithelium)
CPa	cortical plate, anterior	LT	lower tier of the cortical plate (layers VI–V)
CPp	cortical plate, posterior	lv, LV	lateral ventricle
d, D	dorsal	m, M	medial
DC	diencephalon	mc	mitotic cells
DL	dorsolateral	MC	mesencephalon
DM	dorsomedial	ML	medial limbic cortex
DP	dorsal peduncular cortex	mn	zone of migrating neurons
E	embryonic day	MOC	motor cortex
ECL	lateral entorhinal cortex	mr	mammillary recess
ECM	medial entorhinal cortex	MS	mesencephalon
ez	ependymal zone	mSP	migrating subplate neurons
FI	fimbria	mz, MZ	mitotic zone of neuroepithelium
FL	forelimb area	NC	neocortex
		ne, NE	neuroepithelium (ventricular zone)
		OB	olfactory bulb

OC	orbital cortex	SE	septum (differentiating)
OC1B	occipital cortex, area 1 binocular part	SL	lateral somatosensory area
OC1M	occipital cortex, area 1 monocular part	SM	medial somatosensory area
OC2L	occipital cortex, area 2 lateral part	SP	subplate (layer VII)
OC2M	occipital cortex, area 2 medial part	SSC	somatosensory cortex
OCL	lateral occipital cortex	sSP	temporary positions of subplate neurons
OCM	medial occipital cortex	sv, SV	subventricular zone
ole	olfactory epithelium	sv1	lower subventricular zone
olp	olfactory placode	svu	upper subventricular zone
olv	olfactory ventricle	sz, SZ	synthetic zone of neuroepithelium
op	olfactory pit	sIV	sojourn zone of layer IV neurons
or	optic recess	sV	sojourn zone of layer V neurons
ov	optic vesicle	sVI	sojourn zone of layer VI neurons
p, P	posterior	TC	telencephalon
PAR	parietal cortex	te	tegmental neuroepithelium
PAR1	parietal cortex, area 1 (Somatosensory)	TE	temporal cortex (Auditory)
PAR1l	parietal cortex, area 1 lateral part (Somatosensory)	TE1	temporal cortex, area 1 (Auditory)
PAR1m	parietal cortex, area 1 medial part (Somatosensory)	TE2	temporal cortex, area 2 (Auditory)
PAR2	parietal cortex, area 2 (Somatosensory)	TE3	temporal cortex, area 3 (Auditory)
pbt	posterior basal telencephalic neuroepithelium	tf, TF	transitional field of neocortex
PBT	posterior basal telencephalon (differentiating)	th	thalamic neuroepithelium
PCP	primordial cortical plate	TH	thalamus (differentiating)
PI	pineal gland	tlv	telecephalic vesicles
PICP	piriform cortical plate	TT	tenia tecta
pl, PL	primordial plexiform layer	up, h	up, heavy (labeling pattern in late cortical neuroepithelium)
PM	posteromedial	up, l	up, light (labeling pattern in early cortical neuroepithelium)
PO	preoptic area	UT	upper tier of the cortical plate (layers IV–II)
POST	posterior	v, V	ventral
PPL	primordial plexiform layer	VB	ventrobasal thalamic nucleus
PPO	posterior primary olfactory cortex	VBL	ventrobasal thalamic nucleus, lateral
PR	perirhinal	VBM	ventrobasal thalamic nucleus, medial
prc	prosencephalic central canal	VC	visual cortex
prv	prosencephalic neuroepithelium	VL	ventrolateral
pt	pretectal neuroepithelium	VM	ventromedial
PT	pretectum	VM	ventromedial thalamic nucleus (Fig. 16–3)
r	reservoir of lateral cortical stream	VPm	ventroposteromedial thalamic nucleus (also called VM)
RC	rhombencephalon	vz, VZ	ventricular zone (neuroepithelium)
RF	rhinal fissure	VZa	anterior ventricular zone
Rp	Rathke's pouch	VZp	posterior ventricular zone
RS	retrosplenial cortex	v3	third ventricle
RSA	agranular retrosplenial cortex	v3d	third ventricle, dorsal
RSG	granular retrosplenial cortex	v3v	third ventricle, ventral
sb	superior bridge	WH, WM	white matter
sb1	first superior band	I	layer I of neocortex
sb2	second superior band	II	layer II of neocortex
sb3	third superior band	III	layer III of neocortex
sc	superior collicular neuroepithelium	IV	layer IV of neocortex
SSC	splenium of corpus callosum	V	layer V of neocortex
se	septal neuroepithelium	VI	layer VI of neocortex