

Evolutionary Mechanisms and Neural Adaptation: Selective Versus Constructive Strategies in the Development and Plasticity of the Nervous System

Ferdinando Rossi 5

Abstract The correct function of the nervous system requires complex neural 6 networks bearing precise connections. In principle, the high structural specificity 7 of neural circuits could be achieved by genetically-determined processes, selected 8 and refined during evolution. Highly conserved gene networks regulate some 9 crucial steps of neural development, such as the regionalization of the neural tube 10 and the initial phases of neurogenesis and synaptogenesis. A totally hardwired 11 nervous system may meet the requirements of adaptation and natural selection at 12 the population level. Nevertheless, it would be inadequate to allow individual 13 organisms to cope with rapid changes of environmental conditions. Neural adapta- 14 tion to external constraints can be partly achieved by introducing selective 15 mechanisms in neural development. Accordingly, neurons are generated in excess 16 and then partially eliminated to match the actual extension of innervation 17 territories. Such mechanisms, however, are restricted to a set of potentialities, 18 which must be predetermined in the ontogenetic program. On the other hand, 19 constructive mechanisms, in which external stimuli directly influence structural 20 modifications of neural circuits to produce adaptive responses, may allow individ- 21 ual organisms to cope with a wide variety of unprecedented situations. Thus, in the 22 last ontogenetic period as well as in the adult, when the organism actively interacts 23 with the external milieu, experience exerts a strong growth-promoting effect on 24 neural circuits and connections inducing the emergence of specific functional 25 properties. By this mechanism, which requires strict inhibitory control to prevent 26 aberrant growth and dysfunction, the nervous system exploits external stimuli to 27 create adaptive responses to unexpected situations. 28

3

F. Rossi (⊠)

Department of Neuroscience, Section of Physiology, University of Turin, Orbassano, Torino, Italy e-mail: ferdinando.rossi@unito.it



154 F. Rossi

29 1 Introduction

Over the last decades, substantial advancements have been obtained in the clucida-tion of the cellular and molecular interactions that regulate the development of the nervous system, govern its function and determine its plastic capabilities in adult-hood. These discoveries have led to the proposition of concepts and principles that relate, in a very peculiar manner, developmental neurobiology and neurophysiol-ogy to evolution. In addition to the obvious influence exerted by evolutionary processes on neural ontogenesis and on neurobiological mechanisms [57], this novel relationship stems from the understanding that both the construction of the nervous system and its operation are continuously scrutinized for their efficacy in enabling the organism to cope with environmental demands. Hence, the notion that neural development and plasticity represent the biological substrates of adaptation has led to propose that these processes are regulated by fundamental mechanisms that are shared with Darwinian evolution and, notably, the mechanisms of natural selection [8, 13].

This concept originated from the discovery that some fundamental ontogenetic phenomena, such as the formation of appropriate numbers of neurons or synapses in the brain, are subjected to environmental constraints, in a way that is reminiscent of the regulation of population size in living organisms. For instance, there is now general agreement that most neuron populations are initially generated in excess and attain their final numbers by a process of cell elimination, in which death or survival depend on the extension of innervation territories, the availability of target-derived trophic substances or the level of neuronal activity [27, 47]. Similar considerations are usually applied to synaptogenesis, where initially exuberant contacts are progressively withdrawn according to a set of restrictive parameters, including levels of activity, spatio-temporal patterns of synaptic activation or activity-dependent uptake of neurotrophic factors [27, 62].

This large body of evidence highlights the role of selective mechanisms in aspects of neural development and plasticity that are strictly related to adaptation. Nevertheless, a purely selective mechanism implies a range of pre-existing potentialities, which is restricted following confrontation with intervening demands. In other words, all the available options should be hardwired *ex ante* in the ontogenetic program responsible for constructing the organism. Now, is such a mechanism really compatible with adaptation? How can the variety of pre-existing potentialities be expanded at an adequate pace to match the speed of environmental change? Are the discarded options permanently lost or can they be rescued if they become again advantageous in the future?

A selective strategy is primarily designed to control adaptation at the population level. Hence, it is most efficient in regulating species evolution or, as we will discuss later, in defining the number of neurons belonging to a certain category. On the other hand, the main goal of neural adaptation is to allow individual organisms to cope with changing environmental conditions. A closer examination of neural development and plasticity in this perspective actually suggests that the nervous

85

103

Editor's Proof

system must be endowed with an intrinsic capability to construct neural circuits so 72 to create novel functional properties, beyond the original set of potentialities. As a 73 consequence, both selective and constructive mechanisms participate to determine 74 neural ontogenesis and plasticity. Constructive strategies, however, prevail over 75 selective ones when the individual nervous system has to face contextual environ- 76 mental demands.

Adaptive Mechanisms Can Be Either Predictive or Reactive

Biological modifications set up to cope with environmental changes occur according 79 to two main modes. On one side, the organism is able to predict the incoming 80 variation and builds up an anticipated response. On the other, the organism is unable 81 to foresee the external change and it can only react to novel conditions once 82 they have been established. Thus, predictive adaptation implies that the organism 83 is ready to face the novel environmental demand at the time when it materializes, whereas reactive adaptation will be only unfolded in a subsequent time.

At a first glance, predictive adaptation may appear more efficient in favouring 86 survival of the organism. Nonetheless, it can be only used in a restricted set of 87 situations. Actually, predictive mechanisms are only suitable to face extrinsic 88 changes that happen at a *constant* pace through a long period of time (essentially 89 forever). Organisms that spontaneously acquire predictive abilities are favoured 90 over their counterparts and, hence, these abilities become selected by evolutionary 91 mechanisms. Accordingly, predictive adaptation is usually sustained by highly 92 conserved gene networks, whose spatio-temporal patterns of activation correspond 93 to the time course or space distribution of the related environmental conditions. The 94 best example of this kind is the regulation of circadian and circannual functions [12, 95] 19]. These functions are operated by molecular cascades endowed with intrinsic 96 rhythms that match the duration of relevant environmental periods, to which they 97 become entrained by sensory information. As we will discuss here, predictive 98 mechanisms operate in some major ontogenetic processes, which are also governed 99 by highly conserved gene programs. For instance, the gene networks that direct the 100 building of the body (and neural) plan have clearly evolved to cope with consistent 101 environmental constraints, such as gravity, the sources of energy or relevant 102 sensory information (e.g. sunlight) or the mechanics of movement.

Albeit successful, predictive strategies take very long times to become 104 established and diffused. In addition, it is clear that the vast majority of environmental changes happen according to completely unpredictable frequencies and 106 locations. Such situations can be adequately faced only by means of reactive 107 processes, which allow individual organisms or populations to design and set up 108 novel responses. In these cases, evolutionary processes favour the emergence and 109 maintenance of certain abilities, but leave ample degrees of freedom in their actual 110 expression. Most homeostatic mechanisms work in this way. For instance, body 111 temperature is maintained by a series of evolutionary-selected interdependent 112



149

150

156 F. Rossi

devices, from thyroid hormones to horripilation, whose function is triggered and modulated by feedback loops that tune every response to the concomitant situation.

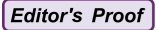
The vast majority of external conditions that may influence the function of the nervous system belong to the latter category. More, I would say that the main emerging property of the nervous system is to design novel strategies to solve unprecedented problems. Accordingly, neural cells and circuits must be endowed with the ability of reshaping connectivity so to generate new functional capabilities that are not part of the constitutive repertoire of the species. Acquiring new information or learning new skills are examples of this sort of morpho-functional modification that underlies neural adaptation. Hereafter, I will argue that these processes, that are crucial to regulate neural development and plasticity, cannot be solely explained in terms of selective mechanisms, but require constructive properties that allow the creative design of new adaptive strategies.

26 3 Neural Development and Evolutionary Mechanisms

127 In the perspective of this essay, neural development can be schematically subdivided in three main phases (Fig. 1): (1) neurulation refers to the formation 129 of the neural tube and its segmentation into discrete morphogenic regions; (2) 130 neurogenesis is the process by which neurons (and glia) are generated; (3) 131 synaptogenesis is the process by which neurons become connected to each other 132 into functional circuits. These phases comprise both addition (e.g. generation of 133 new neurons, formation of new synapses) and loss of elements (e.g. physiological 134 cell death, synaptic pruning). Therefore, the growth of the nervous system actually 135 results from the balance of concurrent expansive and regressive phenomena.

136 Neurulation is triggered by inductive signals issued by the notochord, a mesodermal structure lining the rostro-caudal axis of the embryo, which triggers 137 profound morphogenic rearrangement of the overlying ectoderm leading to the 138 formation of the neural tube [3, 27]. The latter is a highly polarised structure, which 139 soon becomes subdivided in discrete domains that acquire distinctive morphofunctional specification along the main spatial axes (Fig. 1) [3, 27]. The most important partition occurs along the rostro-caudal axis, where morphologically distinct segments appear, corresponding to the major subdivisions of the adult Central Nervous System (CNS). Within each of such segments, the dorso-ventral axis defines sensory or motor structures, whereas the medio-lateral axis defines the 146 relationship linking neural circuits to axial structures (the trunk) and distal appendages (the limbs). 147

The regionalization and spatial specification of the neuraxis are determined by the interplay between diffusible or contact signalling cues and the combinatorial expression of specific sets of transcription factors [3]. The whole process is regulated by gene networks, which direct the morphogenesis of the entire body plan. This gene program has been particularly successful during evolution: it has been inherited from invertebrates and it is highly conserved through the phyla of



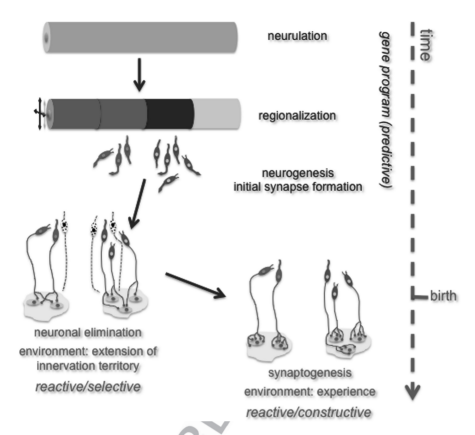
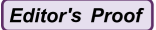


Fig. 1 Regulatory mechanisms of neural ontogenesis. The early phases of nervous system development are determined by the execution of a gene program that directs neurulation, the regionalization of the neural tube, the generation of nerve cells and the initial formation of synapses. While these processes are regulated in a predictive manner, later phases are accomplished according to reactive strategies, required to adapt ontogenetic processes to contextual environmental conditions. Surplus neurons are eliminated before birth by a selective mechanism depending on the extension of available innervation territories. On the other hand, synaptogenesis is carried out after birth, when the organism is interacting with the external world. Hence, synapse formation and reshaping are governed by experience-dependent constructive mechanisms

vertebrates [51]. The program assembles a structural scaffold, in which fundamental 154 morphogenic interactions are precisely regulated in space and time, securing the 155 coordinate development of intrinsic neural networks and their appropriate integration within the nascent organism. On this basic canvas, evolution creates diversity 157 by introducing domain-specific variations in the rate of growth and in the connection patterns. In this way, birds have a relatively large mesencephalon, whereas 159 mammals are characterized by a prominent telencephalon. Thus, neural morphogenesis is accomplished, in a predictive manner, by the intrinsic activity of specific 161 gene networks, whose success is determined a posteriori by natural selection.



207

158 F. Rossi

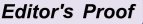
163 Neurogenesis, which is obviously interrelated with morphogenesis, comprises all the phenomena leading to the generation of neurons and glia from neural stem 164 and progenitor cells (Fig. 1) [27]. These cells proliferate in germinal structures 165 located at different levels along the neuraxis, become specified towards different 166 identities and migrate to specific locations, where they acquire mature phenotypes. 167 Then, the final size of each neuronal population can be refined through physiologi-168 cal cell death. The generation of phenotypic diversity is largely determined by 169 diffusible molecular cues or cell-to-cell interactions that regulate the expression of 170 particular combinations of transcription factors [14, 22, 37, 39]. Once cell fate choices have been taken, however, the differentiation into mature phenotypes is 172 achieved by the unfolding of type-specific gene programs, in an essentially cellautonomous manner. Hence, neuronal differentiation as well as the establishment of the basic framework of connectivity are also governed by predictive mechanisms 175 that determine a priori the capability of a given neuron to migrate into a certain position, orientate the navigation of its axon or recognize appropriate targets. 177

The situation is different when the regulation of neuron numbers is considered 178 179 (Fig. 1). The number of neurons generated for each category is determined by the interplay between intrinsic properties of neural progenitors and local regulatory 180 interactions that modulate the rhythm of proliferation, the relative proportion of 181 cells that initiate differentiation or continue to divide, and the duration of neuro-182 genic periods [7, 33]. All these mechanisms operate to regulate neuron numbers by 183 adjusting their production and, hence, work according to a predictive strategy. 184 Nevertheless, since the pioneering work of Rita Levi-Montalcini and Giuseppe 185 Levi [30], it is well known that most neuron populations are actually generated in 186 excess and the final amount of nerve cells that populate the mature nervous system 187 is achieved through the elimination of supernumerary elements [42]. Cell death or 188 survival depend on a set of parameters, including both intrinsic features of the 189 190 neurons (e.g. their level of activity) and environmental constraints (e.g. the extension of the target field or the availability of neurotrophic substances). This process 191 is suitable to match the size of each neuronal population to the amount of potential 192 synaptic partners or to the extension of innervation territories in the periphery. It 193 operates according to a selective mechanism that is most reminiscent of natural 194 195 selection: the juvenile neurons compete for limited quantities of available resources and their fate depends on their intrinsic ability to overcome their rivals [8, 47]. In 196 197 this case, however, the mechanism works following a reactive strategy, required to adjust neural development to individual fluctuations in the dimension of different 198 parts of the body. Accordingly, the size of most neuron populations can be signifi-199 200 cantly modified by experimental manipulations that increase or reduce the extension of the available innervation territory [27, 42, 44]. Therefore, the final number 201 of neurons belonging to each population derives from a dual mechanism, which 202 combines a predictive component, that determines the initial production of surplus 203 neurons, and a reactive component, that eliminates supernumerary elements in 204 205 response to contextual environmental conditions.

At a first glance, similar mechanisms may apply during synaptogenesis (Fig. 1). A well-established notion in developmental neurobiology is that synapses are

237

247



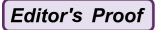
initially formed in excess and then partially withdrawn to shape the mature connec- 208 tivity [27, 47]. Since effective function of neural circuits depends both on the 209 number and on the specificity of synapses, the pruning process would be required 210 both to reduce the exuberant, supernumerary contacts and to remove aberrant, 211 wrong connections.

The initial formation of synapses is guided by recognition cues exposed on the 213 neuronal membrane, whose nature is determined by the intrinsic neurochemical 214 profile of the partner neurons [5, 61, 66]. Synaptic pruning is driven by activity- 215 dependent mechanisms that are directly influenced by the functional efficacy of the 216 developing circuitry [8, 47]. Thus, synaptogenesis also appears to depend on a dual 217 mechanism. Synapse formation is guided by molecular interactions determined by 218 the unfolding of neuronal-intrinsic gene programs that work in a predictive manner. 219 On the other hand, synaptic pruning is driven by an essentially reactive mechanism 220 that selects good connections on the basis of their functional meaningfulness. Again, 221 the latter phenomenon appears to follow some fundamental principles of natural 222 selection.

The analogy is partial at best. It is well established that a number of synapses are 224 withdrawn to shape appropriate spatial connection patterns on specific target 225 domains. Nonetheless, it is definitely clear that, when the number of contacts and/226 or their functional weight is considered, the final balance of the synaptogenic 227 process is a positive one: newly-formed synapses greatly outnumber the lost ones 228 [46, 49, 62]. This has been clearly demonstrated in a variety of experimental 229 models, including the autonomic nervous system [31, 48], the visual system [60], 230 or the cerebellar climbing fibres [23], just to cite a few ones. Even in the case of the 231 neuromuscular junction where mono-innervation of muscle fibres appears to be 232 solely achieved through the elimination of supernumerary axons, the winner 233 endplate undergoes a remarkable outgrowth to cover the entire postsynaptic surface 234 with additional junctional complexes and releasing sites [46, 55]. Therefore, the 235 reactive component of synaptogenesis is not a selective process, but rather operates 236 in a constructive manner.

This conclusion has profound implications in terms of structure-to-function 238 relationship during neural development. Indeed, while the initial formation of 239 synaptic contacts is essentially aimed at establishing a basic framework of neural 240 networks capable of initiating the interaction with the external world, the refine- 241 ment phase is aimed at modifying the structure of such networks to improve their 242 operational abilities. Thus, a fundamental circuit scaffold, assembled by executing 243 intrinsic gene programs, is confronted with experience and modified to achieve 244 adaptive function. The latter process involves the elimination of some unspecific 245 contacts, but it is primarily characterized by the strengthening of meaningful 246 connections with the addition of numerous new synapses.

This process of structural remodelling, which involves the simultaneous 248 outgrowth of both presynaptic axons and postsynaptic dendrites [44, 48], leads to 249 the emergence of novel functional properties, whose nature is influenced by the 250 specific features of the contextual environmental conditions. In other words, the 251 final structure of neural circuits is congruent with the actual experience: a particular 252



291

160 F. Rossi

interaction with the external world will always lead to an appropriate pattern of connectivity [53]. The essentially constructive nature of this process can be best 254 appreciated in extreme experimental conditions. For instance, severe manipulations 255 such as monocular deprivation or experimental squid during the critical periods of 256 visual system development induce extensive changes in the connectivity of the 257 subcortical and cortical visual system [60]. This peculiar structure, albeit strongly 258 divergent from that of the *normal* population, is clearly adaptive when the visual 259 experience of the relevant individuals is considered. Indeed, there is no reason to 260 leave half of the cortical territory to an eye that is not conveying any significant 261 sensory information. Similarly, there is no use to form binocular connections if the 262 two eyes are seeing different scenes. Yet, it is difficult to believe that such unusual 263 projection patterns result from the selection of pre-existing connections, rather than 264 being actively constructed by adapting the morpho-functional properties of the 265 circuit to real life experience. Similar considerations apply to other systems, such 266 as the peculiar tonotopic representation that can be induced in the auditory cortex 267 by exposure to auditory stimuli of specific frequencies [10]. 268

On the whole, the initial phases of nervous system development, which include neural morphogenesis, neuronal production and the establishment of basic connection patterns, are directed by the activity of species-specific gene networks that operate according to an essentially predictive strategy. These processes lead to assemble the fundamental framework of the nervous system, which then undergoes individual-specific morpho-functional adaptation according to reactive strategies. Neuron numbers are refined through a primarily selective process, whereas synaptic patterns are reshaped according to constructive mechanisms. The latter mechanisms have been likely evolved to exploit influences derived from contextual experience to favour the development of adaptive function.

279 4 Experience-Dependent Mechanisms, Neural Development 280 and the Emergence of Function

A major feature of the last phases of neural development is the appearance of 281 reactive processes that essentially shift adaptation from species to individuals. Such 282 processes, however, are accomplished during distinct ontogenetic phases, 283 characterized by strongly different conditions [27, 46]. Neurogenesis and physio-284 285 logical cell death primarily occur before birth and are influenced by somatic changes taking place within the same developing organism. On the other hand, 286 the bulk of synaptogenesis is carried out after birth, while the newborn organism is 287 actively interacting with the external world. The latter condition exerts a most 288 dramatic influence on the course and on the outcome of this process. 289

Higher vertebrates, notably mammals, are born with immature neural circuits, and this feature is most prominent in primates and humans [45, 57]. This implies that crucial phases of neural development occur while the organism is exposed to



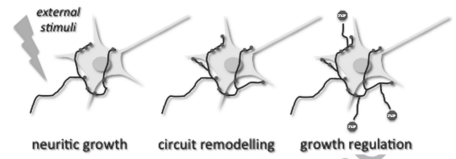
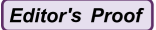


Fig. 2 External stimuli direct developmental synaptogenesis and adult circuit plasticity. External stimuli drive plastic modification of neural circuits by inducing neuritic remodelling and directing the formation of functionally meaningful contacts. The process is regulated by inhibitory cues present in the CNS microenvironment (represented by the STOP signals), required to prevent aberrant growth and dysfunction

the external environment rather than sheltered in an egg or in the uterus. The 293 newborn CNS, and particularly those structures that are more immature at birth 294 such as the neocortex, is subjected to a wide range of powerful stimuli, which 295 induce specific patterns of neuronal activation, stimulate neuritic extension and 296 influence the number and the distribution of newly-formed synapses [63]. This 297 ability of experience to stimulate neural growth is the crucial event that shifts the 298 nature of synaptogenesis from a selective process aimed at achieving synaptic 299 specificity to a constructive one capable of building new functionally meaningful 300 connections (Fig. 2).

Sensory deprivation experiments, such as dark rearing or exposure to un- 302 modulated acoustic stimulation [11, 60, 65], show how administration of meaningful 303 stimuli immediately activates neuronal growth mechanisms, associated with rapid 304 acquisition of new functional properties. All these examples of experience-dependent 305 structural remodelling are characterized by a clear prevalence of expansive phenomena, with the formation and strengthening of new synapses, over regressive events and 307 loss of contacts. Hence, experience drives neuronal growth to create adaptive function. The evolutionary advantage of this strategy is obvious: each individual organism 309 capable of exploiting contextual experience to generate appropriate novel responses 310 will be able to successfully cope with a wide range of unprecedented situations.

Once function is acquired, synaptogenic processes are greatly reduced if not 312 completely arrested [24]. This decline of neuronal growth properties, that marks the 313 end of developmental critical periods for the acquisition of experience-dependent 314 capabilities, has been attributed to a set of concurrent mechanisms. The remodelling 315 of neural circuits often leads to a substantial segregation of afferent axons, which 316 impinge upon private target domains, being individual dendrites, single neurons or 317 discrete anatomical modules. This process of input segregation would progressively 318 reduce the need and the opportunity for activity-dependent competitive interactions 319 that sustain synaptogenesis [62]. Hence, growth would be arrested when a stable 320 connection pattern is achieved and all partners had their share.



[26, 59].

347

361

162 F. Rossi

322 In spite of the attractive simplicity of this mechanism, the end of synaptogenic processes is actually coincident with profound modifications that occur in the 323 neurons themselves and in the surrounding microenvironment [53]. Within the 324 nerve cells, growth-associated gene programs are actively suppressed to favour 325 information processing and signalling function. Coincidentally, the maturation of 326 glia, namely myelination, and the deposition of the extracellular matrix are 327 accompanied by the appearance of a variety of growth-inhibitory molecules that 328 stabilize contacts and hamper further elongation of neuronal processes (Fig. 2). 329 These phenomena are precisely aimed at restricting growth properties of neural 330 circuits. As we will see in the next section, synaptogenic properties typical of 331 juvenile organisms can be restored in the mature CNS by specific manipulations that boost intrinsic neuronal growth properties or remove environmental inhibition. 333 The presence of such strict growth control mechanisms, which have been 334 progressively implemented during the evolution of vertebrates [17, 56], represents 335 an additional argument favouring the constructive nature of developmental 336 synaptogenesis. Indeed, a purely selective mechanism is self-limiting and does 337 not require additional regulatory devices to be terminated. On the contrary, a 338 constructive mechanism must be actively arrested, either by removing the sustain-339 ing stimuli or by dampening growth processes. Experience cannot be prevented or 340 abolished: the whole ontogenetic process is precisely aimed at making the nervous 341 system able to cope with external constraints. Therefore, when the development of 342 neural circuits adopted the constructive strategy driven by experience-dependent stimulation, a set of growth-inhibitory mechanisms evolved to stabilize meaningful connections and to restrain neuronal growth once function is achieved. Not surprisingly, the induction of such regulatory molecules is also triggered by experience

Constructive Mechanisms and Plasticity in the Adult

In spite of the clear decline of intrinsic neuronal growth potentialities, after the end of 349 350 canonical ontogenesis the nervous system retains a certain degree of ability to modify his structure and function in response to external stimuli or changes in the environ-351 ment. Adaptation in the mature nervous system, which is generally known as *plastic*-352 ity, shares some fundamental features and mechanisms with developmental processes. 353 The notion of plasticity in the adult CNS was established several decades ago with 354 355 the discovery of reactive synaptogenesis and synaptic turnover [9, 50]. Accordingly, for a long time the adaptive abilities of neural circuits were thought to be exclusively 356 sustained by changes of connectivity. Recently, however, the demonstration that 357 neurogenesis persists at least in some regions of the adult mammalian brain has 358 revealed that functional adaptation can be also carried out by integrating new 359 360 neurons in pre-existing circuits.

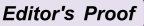
Compared to neural development, synaptogenic phenomena occurring in the adult nervous system are considerably restricted in space and time. They involve 362

Evolutionary Mechanisms and Neural Adaptation

372

385

386



both formation and withdrawal of synaptic contacts and, although they usually lead 363 to moderate changes of synaptic numbers, they obey to reactive mechanisms and 364 have a clear constructive character. One important difference with juvenile 365 synaptogenesis is the requirement of active participation [29, 64]. Synaptic 366 remodelling in immature organisms is usually triggered by the mere exposure to 367 external stimuli. In contrast, in adulthood plastic changes also require motivation and active participation of the involved organism. Hence, in mature individuals 369 adaptation is no more an automatic response to environmental conditions, but 370 requires an individual volition that determines the nature of the response and 371 influences its outcome.

Plasticity in the adult is strongly hampered by the presence of the abovementioned inhibitory mechanisms that terminate developmental synaptogenesis. 374 These mechanisms are partially counteracted by the growth promoting effect 375 exerted by external stimuli [18, 20, 54]. Accordingly, structural plasticity and 376 functional adaptation in the adult can be conspicuously enhanced by experimental 377 procedures that activate neuronal growth genes or neutralize inhibitory molecules 378 of the CNS microenvironment [53]. Nevertheless, whatever effective the simple 379 manipulation of the molecular devices that control neuritic growth is not sufficient 380 to induce adaptation. Endurable structural changes associated with significant 381 functional modifications can only be established if these procedures are combined 382 with specific environmental stimuli [43]. Hence, growth regulatory mechanisms exert a purely permissive role by setting the degree of plasticity of neural circuits, whereas environmental stimulation has a primarily instructive function in determining the shape of the connectivity that will be formed [53].

These features are consistent with a reactive mechanism that induces structural 387 remodelling of neural circuits to generate adaptive responses. As for developmental 388 synaptogenesis, the presence of multiple inhibitory mechanisms is required to 389 maintain constructive modifications within the limits of adaptive function. Indeed, 390 there are several examples showing that altered regulatory mechanisms and/or 391 unusual experience may induce unspecific growth associated with frank pathological 392 phenomena, such as seizures or dystonia [1, 6, 40]. A selective mechanism may fail 393 to generate an adaptive response if the required option is not available, but it should 394 be intrinsically unable to produce abnormal structures and aberrant function. Thus, 395 plasticity in the adult also follows a constructive strategy and, for this reason, it 396 must be subjected to inhibitory control.

Adult neurogenesis shares its major functional significance with adult plasticity. 398 In some CNS structures adaptation is not exclusively sustained by changes of 399 connectivity, but also involves the integration of newly generated neurons into 400 pre-existing circuits. As discussed above, developmental neurogenesis comprises a 401 predictive mechanism that generates excessive amounts of neurons, whose final 402 number is defined by a reactive mechanism that operates through selection. The 403 scenario of adult neurogenesis is very different. In both regions of mammalian brain 404 where new neurons are generated throughout life, the hippocampal dentate gyrus 405 and the olfactory system, the rate of neuronal generation is clearly influenced by 406 external stimuli and/or activity-dependent mechanisms [15, 35]. Thus, while the 407



412

419

420

421

422

423

424

425

426

427

428

429

430

164 F. Rossi

adult system retains the capacity for generating neurons, the course and outcome of the process are no more determined by an intrinsically-coded predictive mechanism, but regulated by extrinsic cues according to a reactive strategy.

Many of the newly generated neurons survive only for a short time, suggesting that survival may depend on selective mechanisms, as for developmental neurogenesis. However, the number and the specific features of the neurons that eventually become stably integrated in adult circuits depend on the activity of the involved network and on specific functional demands [2, 28, 32, 41]. In other words, integration of the newborn neuron is directly related to the function that is being established and not to the intrinsic receptive capacity of the system. Therefore, similar to synaptic remodelling, adult neurogenesis appears to work as a reactive device obeying to a primarily constructive strategy.

This conclusion is further supported by the observation that neurogenesis, or at least neurogenic attempts, may be induced in other regions of the CNS by strong stimulation or pathological conditions [4, 34, 52, 58]. In these instances, non-neurogenic structures react to extreme environmental constraints by redirecting the specification of local progenitors towards neuronal lineages. These phenomena of intraparenchymal neurogenesis are often abortive, because non-neurogenic regions fail to provide adequate conditions to support the differentiation and integration of new neurons. Hence, latent neurogenic potentialities may be diffused in many CNS regions, but actively repressed by local constraints. In any case, adult neurogenesis appears to be driven by environmental stimuli influencing the mature tissue, rather than local regulatory cues acting in a primary germinal structure.

Another feature that adult neurogenesis shares with adult plasticity is the pres-431 ence of strict inhibitory control. Intrinsic inhibitory control prevents adult neurons 432 from de-differentiating or re-entering the cell cycle [25]. In addition, environmental 433 cues regulate the proliferation of progenitors as well as the migration, differentiation and integration of newborn neurons [38]. Thus, successful incorporation of 435 new neurons in adult networks is restricted to precise phenotypes in defined circuits. 436 Furthermore, transplantation experiments show that the endogenous ability of the 437 adult CNS to accommodate donor neurons in functional circuits is limited to a few 438 types and locations [21, 36]. These inhibitory constraints also appear to be primarily aimed at preventing aberrant phenomena that may lead to maladaptive function or behaviour. However, these considerations indicate that adult neurogenesis also has 441 the main characters of a reactive/constructive process, in which experience-dependent growth is exploited to modify neural structures so to achieve adaption.

444 6 Conclusions

The initial phases of neural development are primarily regulated by predictive mechanisms that have been established by evolution. These processes, which are highly conserved throughout vertebrate phylogenesis, are designed to develop a nervous system that is suitable to control the main bodily functions of the organism

483

484

490



and is capable of interacting with the external world. The sensitivity of neural 449 circuits to external stimuli, however, profoundly influenced the strategy of neural 450 development. When coping with rather constant phenomena, such as the physio- 451 logical expansion or retraction of different body parts, suitable adaptation can be 452 obtained by merely selective mechanisms, which share some features with natural 453 selection. Hence, neurogenesis starts with the production of surplus neurons and 454 their final number is adjusted to match actual requirements, which may fluctuate 455 among individuals, but always remain within predictable ranges. A similar mechanism may also apply to synaptogenesis if the nervous system was designed to be 457 completely hardwired by intrinsic genetically-determined mechanisms.

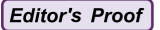
Quite surprisingly, however, the exposure of the immature nervous system to the 459 external environment dramatically changed the ontogenetic strategy. Now, the 460 ability of coping with a great variety of unpredictable environmental constraints 461 could not be adequately fulfilled by a selective process. Rather, the expanding 462 variety of situations favoured the emergence of an alternative mechanism, able to 463 create unprecedented structure and function to face unprecedented situations. Thus, 464 evolutionary pressure pushed developmental synaptogenesis, adult plasticity and 465 even adult neurogenesis to become reactive processes obeying to the rules of 466 constructive mechanisms. This constructive revolution of neural ontogenesis 467 induced the appearance of specific regulatory mechanisms, which evolved to 468 restrain the unchained growth driven by external stimuli within the limits of 469 adaptive function. These inhibitory cues first appeared in fish and amphibians 470 [56], but their importance consistently increased during later vertebrate evolution, 471 in parallel with the increasing complexity of CNS structure and function. Now, they 472 clearly fulfil the fundamental task of controlling potentially dangerous growth 473 properties that enable the nervous system of powerful plastic and adaptive 474 capabilities. However, they also bring with themselves some relevant side effects, 475 such as the loss of neural regeneration capabilities [16, 17]. In any case, constructive mechanisms, such as those directing adult plasticity and neurogenesis, repre- 477 sent a most successful phylogenetic invention that greatly increased the individual 478 ability to cope with increasingly wide ranges of environmental conditions.

Acknowledgements The scientific work of Ferdinando Rossi is supported by grants from 480 Ministero dell'Università e della Ricerca Scientifica e Tecnologica (MIUR-PRIN 2007 prog. 481 nr. 2007F7AJYJ), Compagnia di San Paolo (Neurotransplant Project 2008; GABAGEN Neuroscience project 2009), Regione Piemonte (Project A14/05; Ricerca Sanitaria Finalizzata, 2008, 2009), Ataxia UK; Fondazione Cavaliere del Lavoro Mario Magnetto of Turin.

References 485

1. Aigner L, Arber S, Kapfhammer JP, Laux T, Schneider C, Botteri F, Brenner H-R, Caroni P 486 (1995) Over expression of the neural growth associated protein GAP-43 induces nerve 487 sprouting in the adult nervous system of transgenic mice. Cell 83:269–278

2. Aimone JB, Deng W, Gage FH (2010) Adult neurogenesis: integrating theories and separating 489 functions. Trends Cogn Sci 14:325-337

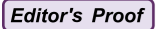


166 F. Rossi

3. Bronner-Fraser M, Hatten ME (2003) Neural induction and pattern formation. In: Squire LR,
 Bloom FE, McConnell SK, Roberts JL, Spitzer NC, Zigmond MJ (eds) Fundamental neuro science. Academic, New York/London

- 4. Brundin P, Winkler J, Masliah E (2008) Adult neurogenesis in neurodegenerative diseases.
 In: Gage FH, Kempermann G, Song H (eds) Adult neurogenesis. Cold Spring Harbor Labora-
- 496 tory Press, Cold Spring Harbor, pp 503–533
- 5. Burden SJ, Berg D, O'Leary DDM (2003) Target selection, topographic maps and synapse
 formation. In: Squire LR, Bloom FE, McConnell SK, Roberts JL, Spitzer NC, Zigmond MJ
 (eds) Fundamental neuroscience. Academic, New York/London
- 6. Byl N, Merzenich MM, Jenkins WM (1996) A primate genesis model of focal dystonia and
 repetitive strain injury. I. Learning-induced dedifferentiation of the representation of the hand
 in the primary somatosensory cortex in adult monkeys. Neurology 47:508–520
- 7. Caviness VS Jr, Nowakowski RS, Bhide PG (2009) Neocortical neurogenesis: morphogenetic
 gradients and beyond. Trends Neurosci 32:443–450
- 8. Changeux JP (1983) L'homme neuronal. Fayard, Paris
- 9. Cotman C, Sampedro N, Harris EW (1981) Synapse replacement in the nervous system of
 adult vertebrates. Physiol Rev 61:684–784
- 508 10. De Villers-Sidani E, Chang EF, Bao S, Merzenich MM (2007) Critical period window for 509 spectral tuning defined in the primary auditory cortex (A1) in the rat. J Neurosci 27:180–189
- 510 11. De Villers-Sidani E, Simpson KL, Lu Y-F, Lin RCS, Merzenich MM (2008) Manipulating
 511 critical period closure across different sectors of the primary auditory cortex. Nat Neurosci
 512 11:957–965
- 513 12. Dunlap JC, Loros LJ, De Coursey P (2004) Chronobiology: biological timekeeping. Sinauer,
 514 Sunderland
- 515 13. Edelman G (1987) Neural Darwinism. The theory of neuronal group selection. Basic Books,516 New York
- 517 14. Edlund T, Jessell TM (1999) Progression from extrinsic to intrinsic signaling in cell fate 518 specification: a view from the nervous system, Cell 96:211–224
- 519 15. Fabel K, Kempermann G (2008) Physical activity and the regulation of neurogenesis in the adult and aging brain. Neuromolecular Med 10:59–66
- 521 16. Fawcett JW, Rosser AE, Dunnett SB (2002) Brain damage, brain repair. Oxford University 522 Press, Oxford
- 523 17. Ferretti P, Zhang F, O'Neill P (2003) Changes in spinal cord regenerative ability through
- 523 17. Ferretti P, Zhang F, O Neill P (2005) Changes in spinar cord regenerative ability through 524 phylogenesis and development: lessons to be learnt. Dev Dyn 226:245–256
- 18. Foscarin S, Ponchione D, Pajaj E, Leto K, Gawlak M, Wilczynski GM, Rossi F, Carulli D
 (2011) Experience-dependent plasticity and modulation of growth regulatory molecules at
 central synapses. PLoS One 6:e16666
- 528 19. Foster RG, Kreitzman L (2004) Rhythms of life: the biological clocks that control the daily
 529 lives of every living thing, Profile Books, London
- 530 20. Gomez-Pinilla F, Ying, Z, Agoncillo T, Frostig R (2011) The influence of naturalistic experi-531 ence on plasticity markers in somatosensory cortex and hippocampus: effects of whisker use.
- 532 Brain Res (in press)
- 533 21. Grimaldi P, Carletti B, Rossi F (2005) Neuronal replacement and integration in the rewiring of
 534 cerebellar circuits. Brain Res Rev 49:330–342
- cerebellar circuits. Brain Res Rev 49:330–342
 535 22. Harris WA, Hartenstein V (2003) Cellular determination. In: Squire LR, Bloom FE,
- 536 McConnell SK, Roberts JL, Spitzer NC, Zigmond MJ (eds) Fundamental neuroscience.
 537 Academic, New York/London
- 538 23. Hashimoto K, Hichikawa R, Kitamura K, Watanade M, Kano M (2009) Translocation of a
 "winner" climbing fiber to the Purkinje cell dendrite and subsequent elimination of "losers"
 540 from the soma in developing cerebellum. Neuron 63:103–118
- 541 24. Hensch T (2004) Critical period regulation. Annu Rev Neurosci 27:549-579
- 542 25. Herrup K, Yang Y (2007) Cell cycle regulation in the postmitotic neuron: oxymoron or new
 543 biology? Nat Rev Neurosci 8:368–378

AU1



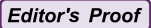
26.	Hockfield S, Kalb RG (1993) Activity-dependent structural changes during neuronal development. Curr Opin Neurobiol 3:87–92	544 548
27.	Jacobson M (1991) Developmental neurobiology. Plenum, New York/London	546
	Kempermann G, Fabel K, Ehninger D, Babu H, Leal-Galicia P, Garther A, Sa W (2010) Why	547
	and how physical activity promotes experience-induced brain plasticity. Front Neurosci 4:189	548
29	Keroughlian AS, Knudsen EI (2007) Adaptive auditory plasticity in developing and adult	549
۷).	animals. Prog Neurobiol 82:109–121	550
20	Levi-Montalcini R, Levi G (1944) Correlazioni nello sviluppo tra varie parti del sistema	
	nervoso. Pont Acad Sci 8:527–568	55°
31.	Lichtman JW (1977) The reorganization of synaptic connexions in the rat submandibular gnalgion during post-natal development. J Physiol 273:155–177	553 554
32	Lledo PM, Saghatelyan A (2005) Integrating new neurons into the adult olfactory bulb: joining	55
52.	the network, life-death decisions, and the effects of sensory experience. Trends Neurosci	556
	28:248–254	557
22		558
33.	Lukaszewicz A, Savatier P, Cortay V, Giroud P, Huissoud C, Berland M, Kennedy H, Dehay C	
	(2005) G1 phase regulation, area-specific cell cycle control, and cytoarchitectonics in the	559
2.4	primate cortex. Neuron 47:353–364	560
34.		56
	progenitors in the striatum. Neurodegener Dis 4:322–327	562
35.	Ma DK, Kim WR, Ming GL, Song H (2009) Activity-dependent extrinsic regulation of adult	563
	olfactory bulb and hippocampal neurogenesis. Ann NY Acad Sci 1170:664-673	564
36.	MacLaren RE, Pearson RA, MacNeil A, Douglas RH, Salt TH, Akimoto M, Swaroop A,	56
	Sowden JC, Ali RR (2006) Retinal repair by transplantation of photoreceptor precursors.	566
	Nature 444:203–207	567
37.	McConnell SK (1995) Strategies for the generation of neuronal diversity in the central nervous	568
	system. J Neurosci 15:6987–6998	569
38.	Ming GL, Song H (2005) Adult neurogenesis in the mammalian central nervous system. Annu	570
	Rev Neurosci 28:223–250	57
39.	Muotri A, Gage FH (2006) Generation of neuronal variability and complexity. Nature	572
	441:1087–1093	573
40.	Nudo RJ (2003) Retuning the misfiring brain. Proc Natl Acad Sci USA 100:7425–7427	574
41.	Oboti L, Savalli G, Giachino C, De Marchis S, Panzica GC, Fasolo A, Peretto P (2009)	575
	Integration and sensory experience-dependent survival of newly-generated neurons in the	576
	accessory olfactory bulb of female mice. Eur J Neurosci 29:679–692	57
42.	Oppenheim RW, Johnson JE (2003) Programmed cell death and neurotrophic factors.	578
	In: Squire LR, Bloom FE, McConnell SK, Roberts JL, Spitzer NC, Zigmond MJ (eds)	579
	Fundamental neuroscience. Academic, New York//London	580
43.	Pizzorusso T, Medini P, Berardi N, Chierzi S, Fawcett JW, Maffei L (2002) Reactivation of	58
	ocular dominance plasticity in the adult visual cortex. Science 298:1187–1189	582
44	Purves D (1988) Body and brain: a trophic theory of neural connections. Harvard University	583
	Press, Cambridge, MA/London	584
45	Purves D (1994) Neural activity and the growth of the brain. Cambridge University Press,	58
чЭ.	Cambridge	586
16		
40.	Purves D, Lichtman JW (1980) Synapse elimination in the developing nervous system.	587
47	Science 210:153–157	588
	Purves D, Lichtman JW (1985) Principles of neural development. Sinauer, Sunderland	589
48.	Purves D, Snider VD, Voyvodic JT (1988) Trophic regulation of nerve cell morphology and	590
40	innervation in the autonomic nervous system. Nature 336:123–128	59
49.	Purves D, White LE, Riddle D (1996) Is neural development Darwinian? Trends Neurosci	592
	19:460–464	593
50.	Raisman G, Field PM (1973) A quantitative investigation of the development of collateral	594
	reinnervation after partial deafferentation of the septal nuclei. Brain Res 50:241–264	598



168 F. Rossi

596 51. Reichert E (2009) Evolutionary conservation of mechanisms for neural regionalization, 597 proliferation and interconnection in brain development. Biol Lett 5:112–116

- 598 52. Robel S, Berninger B, Götz M (2011) The stem cell potential of glia: lessons from reactive gliosis. Nat Rev Neurosci 12:88–104
- 53. Rossi F, Gianola S, Corvetti L (2007) Regulation of intrinsic neuronal properties for axon
 growth and regeneration. Prog Neurobiol 81:1–28
- 602 54. Sale A, Berardi N, Maffei L (2009) Enrich the environment to empower the brain. Trends
- 603 Neurosci 32:233-239
- 55. Sanes JR, Lichtman JW (1999) Development of the vertebrate neuromuscular junction. Annu
 Rev Neurosci 22:389–442
- 606 56. Schweigreiter R (2008) The natural history of the myelin-derived nerve growth inhibitor Nogo-A. Neuron Glia Biol 4:83–89
- 608 57. Striedter GF (2004) Principles of brain evolution. Sinauer, Sunderland
- 58. Suh H, Weng D, Gage FH (2009) Signaling in adult neurogenesis. Annu Rev Cell Dev Biol25:153–175
- 611 59. Sur M, Frost DO, Hockfield S (1988) Expression of a surface-associated antigen on Y-Cells in 612 the cat lateral geniculate nucleus is regulated by visual experience. J Neurosci 8:874–882
- 613 60. Wiesel TN (1982) Postnatal development of the visual cortex and the influence of environ-614 ment. Nature 299:583–591
- 615 61. Williams ME, de Wit J, Ghosh A (2010) Molecular mechanisms of synaptic specificity in developing neural circuits. Neuron 68:9–18
- 617 62. Wong ROL, Lichtman JW (2003) Synapse elimination. In: Squire LR, Bloom FE, McConnell SK, Roberts JL, Spitzer NC, Zigmond MJ (eds) Fundamental neuroscience. Academic,
- 619 New York/London
- 620 63. Zheng D, Purves D (1995) Effects of increased neural activity on brain growth. Proc Natl Acad
 621 Sci USA 92:1802–1806
- 622 64. Zhou X, Merzenich MM (2009) Developmentally degraded cortical temporal processing 623 restored by training. Nat Neurosci 12:26–28
- 624 65. Zhou X, Ngarajan N, Mossop BJ, Merzenich MM (2008) Influences of un-modulated acoustic inputs on functional maturation and critical-period plasticity of the primary auditory cortex.
- 626 Neuroscience 154:390–396
- 627 66. Zipursky SL, Sanes JR (2010) Chemoaffinity revisited: dscams, protocadherins, and neural
 628 circuit assembly. Cell 143:343–353



Author Queries

Chapter No.: 10

Query Refs.	Details Required	Author's response
AU1	Please update the reference Gomez-	Brain Res
	Pinilla et al. (2011).	1388:39-47

