Neurological and psychiatric disorders as a neuroglial failure

ALEXEY VERKHRATSKY1,2,3
VLADIMIR PARPURA4,5
1 Faculty of Life Sciences, The University of Manchester, Manchester, M13 9PT, UK;
2 Achucarro Center for Neuroscience, IKERBASQUE, Basque Foundation for Science, 48011 Bilbao, Spain;
3 University of Nizhny Novgorod, Nizhny Novgorod 603022, Russia;
4 Department of Neurobiology, Center for Glial Biology in Medicine, Atomic Force Microscopy & Nanotechnology Laboratories, Civitan International Research Center, Evelyn F. McKnight Brain Institute, University of Alabama, Birmingham, AL 35294, USA;
5 Department of Biotechnology, University of Rijeka, 51000 Rijeka, Croatia

Correspondence:
Prof. Alexei Verkhratsky
The University of Manchester,
Oxford Road, Manchester, M13 9PT, UK.
E-mail: Alexej.Verkhratsky@manchester.ac.uk
or
Prof. Vladimir Parpura,
Department of Neurobiology,
1719 6th Avenue South, CIRC 429,
University of Alabama,
Birmingham, AL 35294;
E-mail address: vlad@uab.edu

Key words: neuroglia; astrocyte; oligodendrocyte; microglia; reactive gliosis; NG2 glia; neurological diseases; neurodegeneration; psychiatric diseases

Received March 14, 2104.

Abstract

Neuroglia are a diverse non-neuronal population of cells in the central and peripheral nervous system. These cells have a variety of functions that can all be summed up as the maintenance of homeostasis of the nervous system. It is the loss of homeostasis that represents the culprit of all disorders. Thus, neuroglia can be envisioned as the pivotal element in all neural disorders, be that neurological or psychiatric. In this review, we discuss the role of glia in homeostasis and defence of the nervous system as well as changes in the morpho-functional characteristics of these cells in various disorders.

Prelude: neurological and psychiatric disorders as a homeostatic failure

Last century witnessed a remarkable progress in medicine that made most of the somatic diseases cureable; antibiotics conquered infections, advances in immunology and surgery allowed organ transplantation, while oncology developed treatments for many types of cancer. These successes, however, are in stark contrast with the status of medicinal options in neurology, and especially in disorders of the central nervous system (CNS). Indeed, mechanical trauma of the spinal cord invariably results in paralysis, the best cure for stroke is represented by cooling of the brain (known already to ancient Egyptians), and neurodegenerative diseases inexorably proceed to dementia (Alzheimer’s disease), or trigger rapid death (motor neurone disease, also referred to as amyotrophic lateral sclerosis). Similarly hopeless is the realm of psychiatric and neurodevelopmental diseases, as neither cure nor preventive care exists for major psychiatric disorders, such as schizophrenia and major depression, or for neurodevelopmental diseases represented, for example, by heterogeneous autistic spectrum disorders. Modern drugs acting on the CNS are generally agonists or antagonists of major types of neurotransmitter receptors or neurotransmitter metabolic pathways that try to modify (by inhibition or activation) the chemical transmission that underlies synaptic connectivity within neuronal networks. These agents have little spatial specificity, being indiscriminate to the receptors of its relevant kind throughout the nervous system and peripheral organs, and their action is rather generic, being manifested either in stimulation or slackening of nervous activity. When it comes to specific brain and chronic CNS disorders, the therapeutic options are simply non-existing.

The limited cure reflects a fundamental problem: the cellular pathobiology of neurological disorders is ill defined and cell-based therapy has
been developed on the widespread assumption of neurons being the central element in both physiology and pathophysiology, with synapses and neurotransmitter receptors being the chief regulatory pathways in neuronal networks. This neuron-centric dogma is almost universal, being central for the philosophy of experimental and clinical neurology.

This assumption of the dominant role of neurons and neuronal networks in the initiation and progression of neurological disorders, however, is at odds with the general logic of disease nature. Indeed, every disease can be defined as a homeostatic failure in which various exogenous factors (physical, chemical or genetic) interfere with living tissues and infringe their ability to maintain homeostasis, which is the fundamental requirement of life. In other words, disease can be defined as a homeostatic failure and the depth of the failure determines the compatibility with life. According to this logic, the mechanisms of neurological and psychiatric diseases should be sought in homeostatic systems of the nervous system, which are represented by neuroglia, the long-time neglected pariah of neurobiology.

**Neuroglia: the general overview**

The term Neuroglia (or Nerevenkitt; the closest translation from Greek and German is “the neural putty”; the concept and the name were introduced by Rudolf Virchow in 1856-58 (1-3)) defines a highly heterogeneous population of cells responsible for the homeostasis and defence of the nervous system. The homeostatic and defensive roles are the systemic and most fundamental functions of neuroglial cells. The neuroglia comprise cells of ectodermal (i.e., neural) and mesodermal (myeloid) origins (4); generally, neuroglia are sub-classified into peripheral glia and CNS (the brain and the spinal cord) glia (Fig. 1). The glia of the peripheral nervous system incorporate satellite glial cells that localise in sensory and sympathetic ganglia; the numerous and highly heterogeneous enteric glia; the olfactory ensheathing cells and Schwann cells that support and myelinate peripheral axons, and cover neuromuscular junctions. The neuroglia of the CNS are subdivided into macroglia and microglia. The macroglia comprises the astrocytes and cells of oligodendroglial lineage that are further subdivided into oligodendrocytes and NG2 glia. The astrocytes or astroglia (αστρον κυτος; astroon, star and kyton, a hollow vessel, latter cell or the star-shaped cell, the term introduced by Michael von Lehnossek, (5)), encompass protoplasmic and fibrous astrocytes of grey and white matter respectively, the radial glia of the developing CNS, the close relatives of which in the adult CNS are represented by the retinal Müller glia and cerebellar Bergmann glia, velate astrocytes of the cerebellum, interlaminar and polarised astrocytes of the primate cortex, tanyocytes (found in the periventricular organs, the hypophysis/pituitary gland, and the raphe part of the spinal cord), pituicytes in the neuro-hypophysis, and perivascular and marginal astrocytes. Astroglia also include several types of cells that line the ventricles or the subretinal space, namely ependymocytes, choroid plexus cells and retinal pigment epithelial cells. Oligodendrocytes (identified and named so by Pío del Río-Hortega (6)) are myelinating cells in the white and grey matter of the CNS, whereas NG2 cells (discovered by William Stallcup and colleagues (7)) belong to the oligodendrocyte precursor lineage and may also contribute to the homeostasis of the CNS.

The non-neural subpopulation of neuroglia known as microglia are the cells of myeloid origin that represent the main defensive and innate immune system of the CNS. The microglial cells were discovered and characterised by Pío del Río-Hortega in the early 20th century (8, 9). Microglial cells originate from progenitors that derive from primitive c-kit+ erythromyeloid precursors, which come from the extra-embryonic yolk sac (10). These progenitors migrate into developing CNS early in embryogenesis (about embryonic day 10 in mice (11)). After entering the nervous tissue, microglial precursors undergo a substantial transformation and acquire an idiosyncratic morphology, characterised by small cell bodies and several thin and motile processes, and physiology, characterised by the expression of numerous receptors to neurotransmitters and neurohormones concomitant with an expression of “immuno-competent” receptors (e.g., Toll-like receptors and receptors for chemokines and cytokines (12)). Besides being the principle elements of CNS defence, microglial cells play an important role in the development of the nervous system being, for example, pivotal for synaptic pruning, phagocytosis of redundant neurons and shaping synaptic network properties (13).

**Neuroglia: the central element of CNS homeostasis and defence**

The preservation of homeostasis of the nervous system is the main function of neuroglia, which functions include the housekeeping of the neural tissue by astrocytes, maintenance of interneuronal “connectome” by oligoden droglia-dependent axonal myelination and providing defensive homeostasis. Astrocytes perform virtually every conceivable homeostatic function (for recent reviews and extensive references lists see (4, 14, 15)). For example, astroglia are fundamental for structural integrity of the CNS, dividing the grey matter into individual gliovascular units that couple brain parenchyma to the local circulation. Astrocytes control the emergence of the blood-brain barrier (by regulating the expression of tight junctions between endothelial cells) and represent its neural side; similarly, astroglia is central for the formation of the cerebrospinal fluid (CSF)-brain barrier. Through astroglial endfeet covering 99% of CNS capillary walls astrocytes participate in regulated transport of various matter through these barriers and contribute to the regulation of local blood flow. Astrocytes, by virtue of multiple
plasmalemmal transporters and channels, as well as by numerous astroglia-specific enzymes, control CNS homeostasis of ions and neurotransmitters, most notably glutamate, γ-aminobutyric acid (GABA) and ATP/adenosine, or their precursors, in particular supplying neurons with glutamine, which is a precursor for both glutamate and GABA. It is important to emphasise that glutamine supply is critical for neurotransmission, because neurons are devoid of enzymes for de novo synthesis of glutamate (and hence GABA for which glutamate is a precursor). Astrocytes provide for water transport, metabolism, synaptogenesis and the removal of reactive oxygen species. Astrocytes also contribute to systemic homeostasis, being involved in central chemosensation, circadian rhythm and regulation of sleep (16). Oligodendrocytes provide CNS myelination; they are involved in a complex bidirectional communication with axons and contribute to periaxonal ion and transmitter homeostasis, to axonal metabolic support, and are able to dynamically influence the action potential propagation (17-19).

Another fundamental function of neuroglia is the formation of brain defence system. First, neuroglial cells protect nervous tissue through their homeostatic mechanisms, which are, for example, fundamental for containing excitotoxic damage (by removing excess of glutamate and buffering extracellular K+), supporting brain metabolism in conditions of ischaemia through mobilising glycogen, and supplying neurons with energy substrates as well as secreting numerous trophic and neuroprotective factors (20, 21). Furthermore, insults to the brain trigger evolutionarily conserved glial response, generally defined as reactive gliosis, which includes reactive astrogliosis, proliferative response of NG2 cells and the activation of microglia (12, 22-25). The gliotic response is, fundamentally, a defensive reaction responsible for neuroprotection, isolating injured area through the formation of glial scar, removing pathogens, dying cells and cellular debris, and remodelling the nervous tissue after the resolution of pathology.

Reactive astrogliosis, which is activated in most of the pathological processes in the CNS is manifested by a complex biochemical remodelling of astrocytes, their hypertrophy and proliferation and up-regulation of intermediate filaments, i.e., cytoskeletal proteins glial fibrillary acidic protein (GFAP), vimentin and nestin (23, 26). A substantial increase in GFAP immunoreactivity is regarded as a specific marker for astrogliotic response. Importantly, reactive astrogliosis is a controlled, multi-stage and diverse process, which may involve heterogeneous cell populations with distinct neuroprotective capabilities. Furthermore, the manifestation of astrogial reactivity is context-specific and different pathways are associated with distinct reactive astrogial phenotypes (27, 28). Inhibition of reactive astrogliosis generally reduces neuronal viability and worsens the outcome of neurological pathology (24). Finally, reactive astrocytes are instrumental for post-pathology neural tissue regeneration and repair, contributing, for example, to the rewiring of neuronal networks, lesion-induced neurogenesis and reconstitution of blood-brain barrier (26, 29).

Broadly, reactive astrogliosis is classified into isomorphic (i.e. preserving morphology) or anisomorphic (i.e., changing the morphology). In isomorphic gliosis, astroglial hypertrophy, physiological and biochemical changes proceed without altering normal astroglial morphological domain organisation, which is endowed by minute overlap between individual cells at their very periphery (30). Isomorphic astrogliosis is neuroprotective, fully reversible and it facilitates neurtile outgrowth and synaptogenesis. In anisomorphic gliosis, astrocytes became hypertrophic and start to proliferate; glial territorial domains are disrupted, astrocytic processes intermingle and finally a permanent glial scar is formed. The glial scar effectively seals the damaged area and prevents axonal growth, because of chondroitin and keratin secreted by reactive astrocytes (24, 31).

The NG2 glia also respond to various types of CNS pathology by increased proliferation and morphological changes. The processes of NG2 cells in the affected regions become shorter and thicker; this is also accompanied by a substantial increase in NG2 (i.e., chondroitin sulphate proteoglycan 4) expression. The reactive NG2 cells can also proliferate and, at least in spinal cord, the NG2 cells can generate oligodendrocytes that may remyelinate pathologically affected axons. In the spinal cord, activated NG2 cells generate new oligodendrocytes that remyelinate the demyelinated lesions (32, 33). Arguably, NG2 cells may also contribute to scar formation by secreting chondroitin sulphate proteoglycan 4.

Another important component of neuroglial defence is represented by microglia. Insults to the nervous tissue initiate the activation of microglia, which is a multi-stage controlled process progressing through different stages and phenotypes with a variety of morphological, biochemical, functional and immunological changes and producing a variety of phenotypically distinct types of activated microglia. Responses of microglia to pathology are multi-faceted. For example, localised insults trigger rapid converging of microglial processes to the site of injury (34, 35). Stronger and more persistent lesions induce morphological remodelling; microglial somata enlarge and processes become fewer and thicker. Microglial cells alter their expression of various enzymes and receptors, and begin to secrete immune response molecules. At even stronger insults, microglial cells enter proliferative stage, become motile, acquire macrophage-like morphology, migrate and accumulate around the sites of damage and finally transform into phagocytes (12, 25, 36).

Neuroglial reactivity is a central element of CNS response to damage and chronic pathologies. Contributions of activated neuroglia can, however, be not only neuroprotective, but also deleterious. This reflects an intrinsic...
dichotomy of every homeostatic system. The very same molecular cascades that underlie neuroprotection can also contribute to neurotoxicity. Overstimulation of astrocytes can induce the excessive release of glutamate through various release pathways and this release can add to the glutamate toxicity and eventually neuronal death. Abnormal water transport through astroglial aquaporins is a leading mechanism of cellular oedema, whereas deficient astroglial K+ buffering contributes to spreading depression. Similarly, over-activation of microglia results in the release of neurotoxic factors and phagocytosis that can exacerbate neuronal damage.

**General pathology of neuroglia: Reactivity versus atrophy and asthenia**

Neuroglial reactivity in neurological disorders could be considered as dedicated defensive response. At the same time, an opposite process, a loss of function, atrophy or asthenia of glial cells can be directly involved in pathological progression. Evidence for the loss of function of neuroglial cells that accompany different neurological conditions begun to accumulate in recent years. Astrocytes, for example, show signs of morphological atrophy at the early stages of several neurodegenerative conditions (37). In diverse neurotoxic impairments of the CNS, astrocytes lose their ability to control extracellular glutamate, which may be a leading mechanism for ensuing excitotoxicity and neuronal death. Similarly, atrophy or death of astroglia is observed in a variety of neuropsychiatric disorders. In demyelinating conditions, oligodendrocytes fail in remyelination, whereas a loss of function of microglia is involved in neurodevelopmental diseases and is observed in neurodegeneration and in tumorous growth in the nervous system (see (21, 38) and references therein).

**Specific gliopathology in neurological and psychiatric diseases**

**Genetic astroglialopathy: Alexander disease**

The inherited gliopathy, associated with sporadic mutations in the GFAP encoding gene, was described in 1949 by Stewart Alexander (39). Here, the impaired function of astroglia affects brain development and results in severe deficit of white matter manifested by profound leukodystrophy. Histopathologically, Alexander disease is associated with an appearance of cytoplasmic inclusions in astroglial cells known as Rosenthal fibres; these corkscrew-shaped inclusions are formed by GFAP and stress proteins. The pathogenesis of Alexander disease remains unknown and the prognosis is pessimistic with most of the patients dying in early childhood or in adolescence (40).

**Neurodevelopmental disorders**

The glial impairment in neurodevelopmental disorders such as autistic spectrum disorders (ASD) begun to be considered only very recently (see (41) for details and references). Many forms of ASD reflect abnormal formation of neuronal networks and disbalanced neurotransmission. These could result from environmental factors (e.g., exposure to heavy metals or other toxins), oxidative stress, hormonal impairments or early neuroinflammation in combination with genetic predisposition. Astrocytes are the main source for reactive oxygen species scavengers such as glutathione and ascorbic acid, and hence astroglial weakness can lie at the core of oxidative damage to nervous tissue. Astrocytes are also involved in the regulation of neurogenesis and neuronal migration in early postnatal period and hence astroglial weakness can contribute to the malformation of neuronal networks. Astroglia are critical for synaptogenesis (42), and hence for shaping the synapticly connected neuronal networks. Astroglia-derived cholesterol is one of the critical elements of the synaptogenesis and abnormalities of cholesterol metabolism have been detected in ASD (43); these abnormalities may reflect impaired astroglial function and could be linked to oxytocin-mediated signalling pathways acting through oxytocin receptors expressed in astroglia. Finally, ASD is associated with neuro-immune alterations such as an increase in the levels of numerous cytokines (44), which are mainly secreted by microglia. Microglial cells and astrocytes are also implicated in the pathogenesis of Rett syndrome (45, 46) and trichotillomania (47).

**Toxic encephalopathies**

Astrocytes play a primary role in neurotoxic damage to the brain. Astroglial cells are primary targets for heavy metal induced brain damage in Minamata disease (poisoning by methylmercury), and in manganese, lead or aluminium toxic encephalopathies (21, 48-50). In all these toxicities, astroglial cells accumulate heavy metals through astroglial-specific transporters, which in turn affect the plasmalemmal glutamate transporters. Decrease in the activity of the latter results in chronic elevation of extracellular glutamate with ensuing glutamate neurotoxicity and neuronal death. Similarly, astrocytes appear as a primary target in hepatic encephalopathy, which accompanies liver failure. Here, the brain is being poisoned by ammonia, concentration of which markedly increases in the blood and in the CNS following liver insufficiency; the symptoms of ammonia toxicity include confusion, memory deficits, lethargy, somnolence and, in the terminal stages, coma. Astrocytes are chiefly responsible for ammonia detoxification; ammonia is metabolised by glutamine synthetase, astroglia-specific enzyme catalising the condensation of glutamate and ammonia to form glutamine (51). Increased activity of glutamine synthetase in response to elevated ammonia concentration overloads astrocytes with glutamate, impairs K+ buffering and inhibits glutamate transporters. All these result in osmotic shock and cell swelling, brain oedema and glutamate excitotoxicity (52, 53).

**Ischaemia and stroke**

In ischaemic damages to the CNS, neuroglial cells are intimately involved into pathological progression, con-
tributing to both neuroprotection and neurotoxicity (54-56). Normal astrocytes are substantially more resistant to hypoxia than neurones and oligodendrocytes, and hence they can survive in conditions of limited oxygen supply that is characteristic for ischaemic penumbra. Here, astroglial performance is critical for neuroprotection, through removal of glutamate, K⁺ buffering, release of reactive oxygen species scavengers and supplying stressed neurones with lactate. Removal of astrocytes greatly increases neuronal vulnerability in experiments in vitro (57). Such astroglia-dependent neuroprotection is critical for containing the spread of neuronal death through penumbra, which in turn defines post-ischaemic neurological deficit. Astrocytes, however, could exert fundamentally different effects, mediating neurotoxicity, especially in conditions of severe and prolonged ischaemia. The astroglia-mediated neurotoxicity can be mediated through the release (instead of removal) of glutamate via, e.g., the reversal of glutamate transporters or glutamate diffusion through astroglial hemichannels. Astroglial cells can increase extracellular acidosis as a by-product of anaerobic glycolysis; this could be seen in experimental conditions whereby an increase in glucose levels exacerbated the ischemic neuronal damage. Finally, astrocytes can mediate neuronal death through propagating aberrant astroglial Ca²⁺ waves causing distal (to the infarction core) release of glutamate and other neurotoxic factors (58).

Neuropsychiatric diseases

The causes, nature and pathogenesis of neuropsychiatric diseases remain generally enigmatic, albeit there is a recent shift towards the role for disharmony of neurotransmission and in particular deficient glutamatergic mechanisms that include altered glutamate homeostasis and possible endogenous inhibition of N-methyl-D-aspartate (NMDA) glutamate receptors (59, 60). These alterations may certainly be centered on neuroglia which is indispensable for gluta-

tamate turnover, catabolism and synthesis. Morphological studies have confirmed neuroglial alterations such as reduced density and atrophic changes in astroglia and oligo-
dendroglia to be prominent in all three major psychiatric disorders, that is in schizophrenia, bipolar disorder and major depressive disorder; incidentally, no signs of apparent neuroglial reactivity were identified (61). The pathological remodelling of astroglial biochemistry may also be relevant for the progression of schizophrenia. Astrocytes are the main producers of kynurenic acid (through astrogial-specific enzyme kynurenine aminotransferase II, or KAT II (62)); kynurenic acid acts as an endogenous inhibitor of NMDA receptors, and the levels of kynurenic acid are increased in the cortex and in the CSF of patients with schizophrenia (63). Finally, astroglia and kynurenic acid may be a critical link between Toxoplasma gondii infection and an increased risk for schizophrenia. It appeared that T. gondii selectively infects astrocytes, which results in an increased production of kynurenic acid; this may account for the increased risk of schizophrenia (64).

Epilepsy

The pathological cellular substrate of epilepsy is represented by a synchronous slow depolarisation of neurones within an epileptic focus, known as a paroxysmal depolarization shift, which in turn is mediated by the activation of ionotropic glutamate receptors. Epilepsy, in its various forms, is usually associated with prominent reactive astrogliaosis, which often underlies the formation of glial scar. Astroglial reactivity develops at the early stages of the epileptic disorders (which has been observed in both human post-mortem tissues and in animal models) and proceeds in anisomorphic fashion so that reactive astrocytes in epileptic tissue lose their domain organisation (65). Specific feature of astroglial reactivity in epilepsy is represented by (i) an increased expression of ionotropic and metabotropic glutamate receptors, (ii) aberrant calcium signalling; (iii) a decreased presence of inward rectifier K⁺ channels and aquaporins and (iv) a decreased expression and activity of glutamate transporters and glutaminate synthetase. All these changes result in aberrant K⁺ buffering and deregulated glutamate/GABA homeostasis, which may affect neuronal excitability and contribute to the generation of seizures (66-69).

Neurodegenerative diseases

Neuroglia play much more important role in neurodegeneration than has been previously thought, and likely it does play the leading role in some (if not in all) forms of neurodegenerative diseases. Sporadic neurodegenerative process (in contrast to acute neurodegeneration that is a consequence of trauma or ischemic attack), occurs almost exclusively in the CNS of humans; Alzheimer’s disease (AD), Huntington disease (HD), Parkinson disease, motor neuron disease (MND)/amyotrophic lateral sclerosis (ALS) or other forms of dementia do not affect animals. This specificity to humans remains an unsolved conundrum that represents a substantial obstacle to experimental studies of these diseases. In the recent decade, numerous animal models of neurodegenerative diseases, that transgenically insert disease-associated human genes into mice, have been developed (70-73). It has to be remembered, however, that all these models, although reproducing certain parts of human pathologies and often showing disease-specific histopathology, remain imperfect replicas of the naturally occurring diseases.

In MND/ALS (also known as Lou Gehrig’s disease in the US in memory of a baseball player who suffered and died from this pathology) astrocytes are the first cells to undergo pathological remodelling. In a mouse model of MND/ALS (which expresses a human mutant superoxide dismutase 1, or SOD1, associated with a familial form of the disease) astroglial cells in the spinal cord undergo degeneration and become atrophic; these cells have deficient plasmalemmal glutamate transporters and hence cannot contain the excitotoxic build-up of extracellular glutamate (74) that is arguably the leading cause for con-
sequent neuronal death. Furthermore, the MND/ALS progression could be mimicked by astroglia-specific genetic deletion of glutamate transporter 1 in mice (of which excitatory amino acid transporter 2, EAAT2, is a human analogue) (75), whereas selective silencing of the SOD1 mutant gene in astrocytes markedly delayed the progression of MND/ALS (76, 77).

Impairment of the astrocytic ability to clear extracellular glutamate appears as a key pathogenetic mechanism for Wernicke’s encephalopathy that represents an organic substrate for Korsakoff’s psychosis (78, 79). In this disorder, the expression of astroglial-specific glutamate transporters EAAT1/EAAP2 in humans is decreased by 60-70% of the physiological norm. A similar decrease in glutamate transporters was observed in astrocytes from the beriberi (thiamine deficiency) rat model (80, 81). Here, the failure of astroglial glutamate uptake causes profound neurotoxicity, rapid neuronal death with consequent psychotic abnormalities, cognitive deficiency and death.

In AD, which is arguably one of the most common cases of dementia in the developed world, all types of neuroglia are affected and are most likely, linked to pathological progression. AD is characterised by conspicuous atrophy of brain tissue and histopathological hallmarks in the form of the extracellular deposits of β-amyloid protein, known as senile plaques, and intra-neuronal accumulation of abnormal tau-protein filaments, known as neuronal tangles (82, 83). Astrocytes in AD show two types of apparently opposing changes: the relatively early and region-specific atrophy and, at the later stages of the disease characterised by the formation of senile plaques, region-specific reactivity (37, 84, 85). Morphological atrophy, detected as a decrease in astroglial profiles positive to astroglia-specific proteins GFAP, S100β and glutamine synthetase, has been observed in entorhinal and prefrontal cortices, and the hippocampus of several AD animal models (86-91); it also seems to exist in post-mortem human tissues (Rodriguez & Verkhratsky own observations). The early dystrophy of astroglial cells can be pathologically relevant because reduced astroglial synaptic coverage could impair the synaptic strength and synaptic maintenance. Moreover, this reduced astroglial coverage may also influence β-amyloidogenesis. The latter is apparently regulated by glutamatergic transmission; in particular, the activation of synaptic NMDA receptors favours non-amyloidogenic processing of amyloid precursor protein, whereas the stimulation of extra-synaptic NMDA receptors stimulates β-amyloid production (92). Reduced astroglial perisynaptic coverage facilitates glutamate spillover from the synaptic cleft and hence may increase activation status of extra-synaptic NMDA receptors and thus favours β-amyloid production.

At the later stages of the AD, the appearance of senile plaques presents a signal for reactive astrogliosis, and, indeed, an accumulation of reactive hypertrophic astrocytes around β-amyloid deposits have been detected in post-
mortem human tissues as well as in AD animal models (85, 87). The reactive astrocytes show an increased expression of GFAP and S100β, along with a reduced expression of glutamine synthetase, which indicates an impairment of glutamate homeostatic function (93). In addition, reactive astrocytes localised in senile plaques display aberrant Ca²⁺ signalling (94). Astrogial reactivity is region-specific, and it is absent in entorhinal and prefrontal cortices of AD mouse model (89, 90), which may contribute to a higher vulnerability of these brain portions to AD-like pathology.

Progression of AD also affects oligodendroglia and myelination; oligodendrocytes show atrophic changes (95) and reduced densities (96) in AD-affected tissues, which may result in a decreased myelination in the CNS. AD pathology also affects microglia; in the early, preplaque stages, a substantial increase in the microglial densities was observed in AD mice; at the later stages, activated microglia is associated with senile plaques (97, 98). Importantly, however, the activated microglia in AD brains show a loss of function, manifested in the impairment of phagocytosis (99).

Astroglia are also affected in non-AD type dementias, such as fronto-temporal dementia, Pick’s disease, fronto-temporal lobar degeneration, thalamic dementia, or -associated dementia in which both astroglial atrophy and reactive astrogliosis have been identified (100, 101). Primary astrogial pathology, represented by both gliotic and dystrophic changes, is observed in thalamic dementia, in which the loss of astrogial homeostatic functions induces widespread neuronal loss, hippocampal sclerosis and white matter lesions (102). Loss of function (due to mutations) of astroglia-specific protein NPC-1, which appears to function as a transporter in the endosomal-lysosomal system, contributes to the Niemann-Pick disease type C (103). In Huntington’s (HD) disease decrease in astroglial glutamate transporters and possibly in the production of ascorbic acid may contribute to neurotoxicity (104). It should be noted that HD causes preferential loss of a subset of neurones in the brain, although the huntingtin protein is expressed broadly in various neural cell types. Recently, it has been demonstrated that full-length mutant huntingtin expression perturbs astrocyte glutamate transporter release. Hence, BACHD astrocytes show augmented exocytotic glutamate release with unaltered Ca²⁺ dynamics. These astrocytes have a biochemical footprint that would lead to increased availability of cytosolic glutamate, i.e., augmented de novo glutamate synthesis due to an increase in the level of the astrocyte specific mitochondrial enzyme, pyruvate carboxylase. This work identified a new mechanism in astrocytes that could lead to increased levels of extracellular glutamate in HD and thus may contribute to excitotoxicity in this devastating disease (105). Similarly, a loss of astrogial-dependant neuroprotection may contribute to the demise of dopaminergic neurones in Parkinson’s disease (106).

Conclusions: Translational outlook

Neurological and psychiatric disorders have been almost entirely considered from the neurone-centric point of view, with neurons being the principal, if not the sole, cellular element of disorderly process. However, it is neuronoglia, but not neurones that control the nervous system homeostasis, the dysregulation of which is the common denominator in all diseases. Recently, it is becoming clear that neuroglial cells play an active role in pathophysiological processes and that understanding the underlying mechanisms shall provide novel targets for much needed therapeutic intervention.

Acknowledgements: We thank Manoj K. Gottipati for comments on a previous version of this manuscript. VP research is supported by the Eunice Kennedy Shriver National Institute of Child Health and Human Development (HD078678), AV was supported by the Alzheimer’s Research Trust (UK), by European Commission, by IKERBASQUE and by a research grant of Nizny Novgorod State University.

REFERENCES

1. KETTENMANN H, VERKHRATSKY A 2008 Neuroglia: the 150 years after. Trends Neurosci 31: 653-659


8. DEL RÍO-HORTEGA P 1919 Poder fagocitario y movilidad de la microglia. Bol de la Soc esp de Biol 9: 154


analysis reveals that adult microglia derive from primitive macrophages. *Science* 330: 841-845

12. KETTENMANN H, HANISCH UK, NODA M, VERKHARSKY A 2011 Physiology of microglia. *Physiol Rev* 91: 461-553


32. TRIPATHI RB, RIVERS LE, YOUNG KM, JAMEN F, RICHARDSON WD 2010 NG2 glia generate new oligodendrocytes but few astrocytes in a murine experimental autoimmune encephalomyelitis model of demyelinating disease. *J Neurosci* 30: 16383-16390


42. PFRIEGER FW 2010 Role of glial cells in the formation and maintenance of synapses. *Brain Res Rev* 63: 39-46


44. WEI H, MORI S, HUA K, LI X 2012 Alteration of brain volume in IL-6 overexpressing mice related to autism. *Int J Dev Neurosci* 30: 554-559

45. MAEZAWA I, JIN LW 2010 Rett syndrome microglia damage dendrites and synapses by the elevated release of glutamate. *J Neurosci* 30: 5546-5556

46. MAEZAWA I, SWANBERG S, HARVEY D, LASALLE JM, JIN LW 2009 Rett syndrome astrocytes are abnormal and spread MeCP2 deficiency through gap junctions. *J Neurosci* 29: 5051-5061


60. STEINER J, BOGERTS B, SARNYAI Z, WALTER M, GOS T, BERNSTEIN H G, MYINT A M 2012 Bridging the gap between the immune and glutamate hypotheses of schizophrenia and major depression: Potential role of glial NMDA receptor modulators and impaired blood-brain barrier integrity. World J Biol Psychiatry 13: 482-492


63. ALEXANDER K S, WU H Q, SCHWARCZ R, BRUNO J P 2012 Acute elevations of brain kynurenic acid impair cognitive flexibility: normalization by the α7 positive modulator galantamine. Psychopharmacology (Berl) 220: 627-637

64. SCHWARCZ R, HUNTER C A 2007 Toxoplasma gondii and schizophrenia: linkage through astrocyte-derived kynurenic acid? Schizophrenia Bull 33: 652-653


73. SKAPER S D, GUPTI P 2010 Transgenic mouse models of Parkinson’s disease and Huntington’s disease. CNS Neurol Drug Targets 9: 455-470


80. HAZELLE S A 2009 Astrocytes are a major target in thiamine deficiency and Wernicke’s encephalopathy. Neurochem Int 55:129-135


88. RODRIGUEZ J J, TERZIEVA S, OLABARRIA M, LANZA R G, VERKHATSKY A 2013 Enriched environment and physical
activity reverse astrogliodegeneration in the hippocampus of AD transgenic mice. Cell Death Dis 4: e678.


