Pathophysiological reasons for the failure of the cerebral perivascular drainage and progression of Alzheimer’s disease

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ABSTRACT:
The disorder of intracerebral perivascular drainage of amyloid beta (Aβ), as well as the drainage of various waste products related to a number of permanent biochemical reactions and physiological processes, leads to brain homeostasis disorders, breakdown of essential vital functions and rapid course of genetically programmed Alzheimer’s disease. As the brain does not have its own standard lymphatic system, this important function is mostly taken over by the perivascular system located within the basement membranes of capillaries, arterioles and arteries. The crucial driving force of this drainage, according to recent investigations, is conditioned by the internal force of arterial and arteriolar walls, the so-called vasomotion. With the direction opposite from the direction of the blood stream and pulse wave, this pulsating force is based on rhythmic intracellular oscillations of Ca2+ ion concentration in vascular smooth muscle cells. These oscillations are the fundament of the vasomotion phenomenon, and their disorder leads to perivascular drainage alterations and severe complications. The aim of this review is to present a detailed analysis of crucial events that are important for perivascular drainage system alterations.

KEYWORDS: Alzheimer’s disease, perivascular drainage alterations, lymphatic system, vasomotion phenomenon

SAŽETAK:
Patofiziološki uzroci poremećaja moždane perivaskularne drenaže i progresije Alzheimerove bolesti
Poremećaj intracerebralne perivaskularne drenaže amyloid beta (Aβ), kao i drenaže različitih otpadnih produkata, nastalih tijekom brojnih permanentnih biokemijskih reakcija i fizioloških procesa, vodi do poremećaja moždane homeostaze, raspada osnovnih vitalnih funkcija, te ubrzanog tijeka genetski programirane Alzheimerove bolesti. Obzirom na činjenicu kako mozk nema svoj standardni glimfatički sustav, ta je važna funkcija preuzeta sa strane perivaskularnog sustava lociranog unutar bazalnih membrana kapiara, arteriole i arterija. Presudna pokretačka snaga te drenaže, na osnovu najnovijih istraživanja, bazira se na urođenoj sili arterijalnih i arterioloarnih zidova, tzv. vazomociji. Smjerom suprotnim od smjera krvene struje i vala pulsa, ta pulzirajuca sila bazira se na ritmičkim intracelularnim oscilacijama koncentracije Ca2+ iona unutar vaskularnih glatkih mišićnih stanica. Te oscilacije su baza fenomena vazomocije, a njihov poremećaj vodi do alternacija perivaskularne drenaže i teških komplikacija. Namjera ove studije je da prikaže detaljnu analizu presudnih zbivanja koji su važni za alternacije perivaskularnog drenažnog sustava.

KLUČNE RIJEČI: Alzheimerova bolest, perivaskularni drenažni poremećaji, glimfatički sustav, fenomen vazomocije
**Introduction**

Due to the increasing rise of AD incidence and prevalence, Alzheimer’s disease is becoming one of the crucial problems of modern society. The world population is increasingly older, absolutely and relatively, and AD is closely connected with age and aging. This neurodegenerative disease with a chronic and progressive course is marked with a progressive decline of memory, disorientation, and general drop of cognitive functions. The costs for its prevention and therapy exponentially grow. By removing other nonconventional brain lymphatic drainage pathways (the lymphatic system) from the analytic process, which include drainage across the blood brain barrier (BBB), blood cerebrospinal fluid barrier (BCSFB), choroid plexus, arachnoid granulatious (AGs), paravascular drainage (connected with the cerebral vasculature surfaces), perineural drainage, lymphatic drainage through the orbits into the sinus sagitalis superior, and nasal cavity (lymph inflow along the olfactory tract), this study is exclusively oriented to the perivascular drainage analysis [1-6]. Without entering the extensive and complex analysis of the complete cerebral drainage system, the aim of this review is to explain the crucial elements in events connected with the functioning of the perivascular drainage pathway.

**Alterations of the perivascular drainage pathway from the brain**

Alzheimer’s disease, with its poligenetic but yet unknown etiology, is a severe, chronic, and lethal neurodegenerative phenomenon, in which the inadequate drainage of elevated concentrations of Aβ, of iron ions and waste products out from the brain, leads to intracerebral accumulation of these ingredients, with evidently progressive deteriorative effects on brain homeostasis, and through time, to the occurrence of a lethal end. This altered drainage is clearly connected with the aging process. Due to the permanent aging of the world population (senectual explosion), AD is becoming increasingly a crucial problem of the present-day human society. The two forms of this disease, the early form or EOAD (early onset AD-before the age of 65; APP gene-chr. 21q21.3; PSEN-1 gene-chr.14q24.2; PSEN-2 gene-chr.1q42.13; BACE-1 gene-chr.11q23.3), and LOAD (late onset of AD-65 and more; ADAM10 gene-chr.15q21.3; APOE gene-chr.19q13.32) have practically the same clinical picture with the progressive loss of memory, loss of speech, and dramatic failure of cognitive functions [7-9].

It is important to analyze some crucial events in Aβ and iron metabolism, which are in the centre of the AD pathophysiology. Bound with transferrin (Tf), two ferric ions (Fe³⁺), in the form of TfR/Tf/Fe³⁺ complex, enter from the blood into the endothelial cell membrane and into its embeded endosomes. In these structures, the transferrin receptor (TfR) is detached from the complex, and Fe³⁺ is reduced by ferric reductase located on the endosome membrane in Fe²⁺, and by the diveral metal transporter 1 (DMT1) and ferroportin (FPN1) transported into the neuropil. The detached TfR recycles back to the luminal cell membrane. Fe²⁺ in the neuropil is partially captured by DMT1 belonging to the neurons and microglia and becomes incorporated in their metabolism or stored as ferritin. The great part of Fe³⁺ is oxidized (loss of the electron) in Fe⁴⁺ by the locally present amylloid precursor protein (APP) and ceruloplasmin (CP) derived from the cell membranes. One part of Fe³⁺ bound with Tf/TfR enters into the adjoining cells, where it is used for their metabolic necessity or also stored as ferritin. A part of Fe⁴⁺ is attracted to the Aβ monomers and enters the oxidative/reductive cycle with the resulting Fenton reaction and *OH generation. On the other hand, Aβ, by its intracellular (proteasome and ubiquitin-proteasome route, lysosomal cathepsin enzymes, metalloendopeptidases) and extracellular (thrombin, matrix-metalloproteinasises 2,3 and 9) degradation, is exposed to its elimination and iron ions discharge. It is evident that in the case of Aβ overproduction typical for AD, in the neuropil occurs the rise of Aβ concentration and the accompanying rise of iron ions. Due to these events, the effective drainage of elevated concentration of both compounds, Aβ and iron ions, is necessary. This drainage is primarily obtained by the perivascular drainage pathway (Fig.1) [9].

**Elementary characteristics of the perivascular drainage pathway**

This study deals with the central parts of arterial, arteriolar and capillary walls, the basement membranes of vascular smooth muscle cells (VSMCs), the VSMCs, the ground substance of the interstitium, as well as with the local structures such as elastin and collagen fibers, cell receptors, and a lot of diverse stationary or moving dissolved molecules.

The perivascular drainage pathway, limited by the outer and inner basement membranes of vascular smooth muscle cells (VSMCs), comprised by central media location, actually is a tridimensional poroelastic net, along which pass collagen and elastic fibers. The average thickness of each VSMCs membrane which encircles every cell, is in the range of 20-200nm (1nm =1*10⁻⁹ m, nanometer). Its tridimensional net is predominantly composed of proteins i.e. laminins, collagen IV, nidogens, and heparan sulfate proteoglycans (HSPG). There are also expressed many other proteins like the insoluble fibronectin, fibulin 1 and 2, collagen type XVIII, thrombospondins1, and secreted protein acidic and rich in cysteine (SPARC). The fundamental element of this drainage pathway, crucial for the transport of molecules and particles, is the ground substance, an amorphous gel-like substance primarily composed of water, in which are embedded the mentioned fibrils and molecules. The starting point of this pathway is located in BBB capillaries, i.e. in their basement membranes (Fig. 2-4) [1-4].
**Driving forces for perivascular drainage**

The driving force for moving different particles and molecular complexes along this pathway in the direction opposite to the blood stream and pulse wave, is the recently in detail analyzed, but already previously discovered and not satisfactory explained, internal force of arterial and arteriolar walls, the so-called vasomotion. This pulsating force is based on rhythmic intracellular oscillations of $\mathrm{Ca}^{2+}$ ion concentrations in vascular smooth muscle cells. These oscillations cause the vasomotion phenomenon and their disturbance leads to perivascular drainage alterations and severe complications. 4,6,10,11).

**The crucial failures in the driving forces mechanism**

The crucial failures in the mechanisms responsible for regular perivascular drainage are primarily dependent on the elevated lipid peroxidation of VSMCs cell membranes, induced by free radicals. The regular contraction of these cells is important for the adequate generation of pressure gradient along the interior of arterial and arteriolar walls. The disruption of this adequate pressure gradient generation, primarily due to the VSMCs membranes lipid peroxidation, leads to the drop of cell contraction strength and decline of the vasomotion and drainage velocity.

Lipid peroxidation of VSMCs membranes is a permanent spontaneous process, but normally there are effective protective forces to combat with free radical effects and thus repair the alterations. In Alzheimer’s disease (AD), the oxidative stress is elevated. The generation of reactive oxygen species (ROS), especially free radicals with the crucial compound hydroxyl radical (*OH), is significantly elevated. The repair defensive forces are now not capable to combat and repair the failures in cell membranes structure 4,6,10,11).

**Lipid peroxidation**

Cell membranes, especially their main structural elements, polyunsaturated fatty acids (PUFAs), are enormously lean towards the destructive action of hydroxyl radical (*OH), one of the strongest member of free radicals, i.e. ROS. *OH is primarily generated during the Fenton reaction ($\mathrm{Fe}^{3+} + \mathrm{H}_2\mathrm{O}_2 = \mathrm{Fe}^{2+} + \cdot \mathrm{OH} + \cdot \mathrm{OH}$, which includes ferrous iron, hydrogen peroxide, hydroxyl ion, and ferric iron), extensively expanded practically in all tissues of AD patients. However, on account of the extremely great oxygen consumption, and the great protective forces of polyunsaturated fatty acids, the brain is especially prone to *OH and strong spontaneous lipid peroxidation. The initiation of this destructive process occurs by the random crash between the free *OH molecule and hydrogen atom (H) on polyunsaturated fatty acid (PUFA) methylene group (-CH$_2$-), between two double bonds, i.e. bis allylic hydrogen atom. Carbon-hydrogen (C-H) bond energy, due to the presence of the adjacent double bonds is very low and allows the H atoms to be readily abstracted by "OH. The consequence is the formation of the carbon centred radical on the carbon 2py orbital, the bonding of the abstracted hydrogen (electron in the 1s orbital) with the unpaired electron in the 2py orbital of *OH, and the formation of neutral H2O molecule, water (Fig. 5) 12-14).

The next step is the phospholipid carbon centred radical reaction with the molecular oxygen (O$_2$) and the formation of peroxy radical (LOO$^\cdot$). By reacting with the new PUFA, peroxy radical generates the new lipid radical (L$^\cdot$) and lipid hydroperoxide (LOOH). L$^\cdot$ reacts with molecular O$_2$ with the formation of new lipid peroxide (LOO$^\cdot$). This spontaneous reaction continues till the end, induced by the interplay of free radical scavengers, or by the mutual reaction of two *OH to form non-radical products (Fig. 5) 15).

It is also necessary to mention that a number of lipid peroxidation products and by-products is very destructive by inducing oxidative stress, oxidative damage and apoptosis. This is especially related to hydroperoxides and their aldehyde derivatives. Among aldehydes, most important is the 4-hydroxynonenal (HNE), the most effective product of lipid peroxidation, which induces evident cellular damage. Malonyl aldehyde (MDA) also produces protein damage 16).
Fig. 1. Schematic presentation of events connected with Aβ and Fe³⁺ elevated intracranial values

Oxidation, loss of electrons; reduction, gain of electrons; hydroxyl radical (·OH), highly reactive, aggressive and short lived; MsrA, peptide methionine sulfoxide reductase, repair enzyme for proteins inactivated by oxidation; MetS35, methionine one of nine essential amino acids; Fe³⁺, ferric iron; Fe²⁺, ferrous iron; H₂O₂, oxidizer, the simple peroxide; Aβ, amyloid beta peptide; Tf, transferin; for elevated iron ions concentration value, and its decrease, the crucial role has the perivascular drainage pathway; it is emphasized the process of Aβ intracellular (proteasome and ubiquitin-proteasome route, lysosomal cathepsin enzymes, thiolmetalloendopeptidases) and extracellular degradation (neprilysin, matrix-metalloproteinases 2, 3 and 9); small discrepancy in the production or clearance of Aβ unavoidably causes abnormal accumulation in AD.
Anterograde flow (paravascular drainage) AQP4 channel

Astrocyte processes of glia limitans

vessel pia cortical pia Virchow-Robin space Pial funnel

Paravascular space drainage ECS

Intercellular astrocytic endfeet cleft AQP4 channel

Aβ,waste

Basement membrane between VSMc – the pathway for Aβ and waste; located in the media; (IPAD).

inner basement membrane of SMCs (Without drainage)

lumen endothelial cells

ECS smooth muscle cells (SMCs, media, IPAD)

Blood flow from the heart
Anterograde flow
Blood cells and platelets O2, glucose, nutrients

Retrograde flow of waste and Aβ – perivascular drainage

Fig.2. Schematic presentation of the penetrate artery cross section

Clearly presented is the cross section of the cerebral parenchymal penetrating artery. By different colours are visible the essential artery layers including intra-mural peri-arterial drainage (IPAD) pathway and paravascular drainage pathway. By dark green colour are designated the astrocytic endfeets, and between them by black arrows intercellular astrocytic endfeet clefts. By small black rectangles, covered with bidirectional arrows, are presented the AQP4 channels, important for entrance in the artery wall and exit from it, of a lot of solutions and Aβ peptide. The big black arrow shows the most important layer in the artery wall, the media, and in its structures, located between vascular smooth muscle cells, the perivascular drainage pathway (black points in layers between the mentioned cells). Black points in the extracellular space (ECS) represent Aβ and waste. Between cortical and vessel pia there is Virchow-Robin space. On the outer side of the cortical pia, coloured by light blue colour, there is visible the paravascular drainage pathway. Cross section presents the situation on the middle level of the Virchow-Robin space. SMCs are in the relaxed phase of the vasomotion wave.

Figure 2. Schematic presentation of the penetrate artery cross section (from ref. 6)
Fig. 3. Schematic presentation of the Aβ, iron, and waste products drainage from the brain

CSF, cerebrospinal fluid; ECS, extracellular space; Aβ, amyloid beta; VRS, Virchow-Robin space; O₂, oxygen; C₆H₁₂O₆, glucose; CO₂, carbon dioxide ISF, interstitial fluid; SMC, smooth muscle cell; AQP4, aquaporin 4, channellocaled on astrocytic endfeet, important for Aβ drainage; SAS, subarachnoid space; AG, arachnoid granulation; Tf-Fe³⁺, transferin-ferric ion complex; SSS, superior sagital sinus; BBB, blood brain barrier; this figure represents the additional survey of Fig. 3. in the article GSR, 2019; 6:007-020 publishe online: March 31.2019. On the left side of the figure there is visible a rectangle limited by small red lines-it in  fact presents Fig. 5. The other rectangle including capillaries presents Fig. 4. BM at the capillary level is the result of the proximal integration of the outer and inner BM of VSMCs. HDL, high density lipoprote

Figure 3. Schematic presentation of the Aβ, iron, and waste products drainage from the brain (from ref. 5)
On the upper part of the figure there are visible two endothelial cells connected by the tight junction. Above the luminal side of both cells by the red arrow is denoted the capillary lumen, anterograde blood flow direction, some receptors (TIR, RAGE) and Aβ peptide molecules. Below the cells is visible the basement membrane which is composed of two layers, basal lamina and lamina reticularis; Hp, hephaestin. TIR receptor is localized on both luminal and abluminal membranes of brain capillaries; A part of HDL which enters into the EC is wrecked and a part continues to the L. Ret. ApoE can not be transported from the blood across the BBB into the neuropil. ApoE in the CNS is produced primarily by astrocytes. It is the principal lipid transporter in CSF. The extra and intracellular ApoE proteolysis has been discovered.
PUFA, polyunsaturated fatty acid; RAGE, receptor for advanced glycation end products; PKC, protein kinase C; NADPH oxidase, nicotinamide adenine dinucleotide phosphate-oxidase; ROS, reactive oxygen species; O2*, superoxide radical; H2O2, hydrogen peroxide; SOD, superoxide dismutase, strong antioxidant enzyme; catalase, enzyme, protects the cell from oxidative damage; GSH, glutathione peroxidase, enzyme, protection from oxidative damage; lipid peroxidation, oxidative degradation of lipids; Fe2+, ferrous iron; Fe3+, ferric iron; e.c.s., extracellular space.

Figure 5. Schematic presentation of important biochemical processes related to destructive free radical effects on essential molecular structures (from ref. 12)
The investigations show that the lipiddation (covalent binding of lipid group to a peptide chain) of APOEε4 lipoprotein is significantly weaker as related to APOEε2 and APOEε3. APOEε4-HDL particles are markedly smaller than the other two alleles. Otherwise, by aging, the ABCA1 (ATP-binding cassette transporter ABCA1, cholesterol efflux regulatory protein, CERP) expression and activity declines. APOE linked with the lipid (lipiddated form, HDL, LDL) conditions and increases its interaction with Aβ (Aβ, residues 12-28; APOE, N-terminal domain, residues 136-150) with the prompt transport of the whole complex by the effect of LRP1 (low density lipoprotein receptor-related protein1) and ABCB1 (P-glycoprotein, ATP-dependent translocaze ABCB1) receptor. ABCB1 is located on the luminal and abluminal side of the endothelial cells membranes. It is an active mediator for the Aβ peptide transport out from the brain. APOEε4 protein (lipiddated by ABCA1, but significantly weaker than other two alleles), in relation to the other two alleles, has an evidently weaker linking with Aβ, and the generated complex (Aβ/ApoEε4/HDL or LDL) transport is also weaker. The consequent result is stronger Aβ accumulation in the structures of the perivascular space (PVS, basement membranes of VSMCs), as well as its stronger accumulation (the result of the consequent decline of the perivascular drainage) in the brain parenchima. On both levels, its aggregation is stronger. The LRP1 expression drop in the aged has been emphasized earlier 17,18,19-22).

It is important to emphasize that ApoE cannot pass through the BBB. The central part of ApoE (CNS) is produced in the brain by astrocytes (mainly) and microglia. In the periphery, ApoE is produced primarily by the liver. The pools of ApoE in the CNS and periphery practically exist independently from one another (Fig. 4) 23,24).

On figure 4, well visible is the LRP1 effect on the Aβ transcytosis from the basal compartment of the EC to the cell luminal membrane, and its clearance into the blood. Internalization of the soluble Aβ from the lamina reticularis (capillary basement membrane) into the endothelial cell cytosol occurs by the interaction between LRP1 and ApoE. The effect of the latter, especially ε4 allele, in fact, blocks this interaction and diminishes the intracellular entrance of LRP1/Aβ complex. Bound with LRP1, Aβ transcytoses passes across the BBB, and by the help of PICALM (Phosphatidylinositol binding clathrin assembly protein) and Ras-related protein Rab5 (Rab5) as well as Ras-related protein Rab11 (Rab11), the small GTPases, completes the mentioned clearance route (Fig. 4) 25-27.

On figure 4 is shown that HDL particles, carried by apoprotein ApoEε4 (dominant allele in AD) incorporated molecules, arrive by blood in the BBB region where ApoEε4 recognizes the scavenger receptor class B type 1 (SR-B1) and binds with it. Now occurs the endocytosis and internalization of the formed complex HDL/ApoEε4/SR-B1, and its transcytosis. During this event follows the dissociation of ApoEε4 and its proteolytic degradation into a number of fragments, among them some very toxic for the cells. The proteolysis is mediated by the multiple intracellular proteolytic enzymes, among them especially the calpain proteolytic system (cp, calpain proteases). The obtained fragments are mainly toxic for a lot of cellular functions, for example the mitochondria oxidation (electron transport chain), cytoskeletal assembly, and stability. SR-B1 recycles and returns to the luminal membrane. Liberated HDL is partially wrecked and one part is used for the cell necessity. The other part continues its pathway to the abluminal membrane, and there is exocitosed into the capillary basement membrane. There it continues its binding with ApoEε4 (effect of ABCA1) produced by astrocytes, and exits from the basement membrane (BM) into the neuropil, or continues its movement along the perivascular pathway with the accepted Aβ. On Fig. 4 there is also presented the extracellular space with astrocytes which synthesise and secrete HDL and ApoE. There are also visible the Aβ monomers. A major part of generated HDL is served for the cells necessity, and the other part enters into the perivascular drainage pathway and leaves the brain (Fig. 4) 28,29.

**Fenton reaction and *OH generation**

Along with the strong extracellular intracerebral Aβ accumulation and aggregation, in AD, in the brain region, occurs simultaneous accumulation of transition metals, among them especially iron ions. It is considered that the main reason for this event lies in the decreased drainage of iron, especially along the perivascular pathway. Linked with one transferrin (Tf) molecule, two ferric iron ions (Fe³⁺) at the level of BBB enter from the neuropil into the capillary basement membrane (as well as Aβ and a number of other molecules and waste products), and continue their movement along the perivascular drainage pathway. In the case when in this pathway occurs an elevated concentration of Aβ monomers (AD), which readily bind Fe³⁺ ions on their MBD (metal binding domain-His 6,His 13,His 14) the monomers mutual contacts and binding become much easier and frequent with the elevated Fe³⁺ reduction (“electron hop”) in Fe²⁺ (MctS35 on one monomer by its oxidation dismises an electron to Fe³⁺), with the consequent accelerated Fenton reaction with *OH radical spreading. *OH radical overproduction primarily leads to the dangerous oxidative attacks (electron stealing) on the VSMCs cell membranes with their deterioration and fatty acids peroxidation. Earlier in the text, in the paragraph about lipid peroxidation, these events were explained in detail (Fig. 5) 7,12-16.

**Main receptors included in the perivascular drainage**

Basement membranes of cerebral capillaries are the starting points of the perivascular drainage pathway. Biological membranes on the luminal side of endothelial cells of BBB are structures in which are embeded a number of vital receptors.
Among them, it is necessary first to present the receptor for advanced glycation end products (RAGE) and explain its function. It is a transmembrane protein and belongs to the immunoglobulin superfamily. It can interact with multiple ligands such as advanced glycation end products (AGEs), high-mobility group protein (B1) (HMGB1), S-100 calcium-binding protein, Mac-1, phosphatidylserine, and amyloid-β-protein. After binding Aβ, RAGE, located on the apical endothelial cell side, induces its internalization and transcytosis from apical to the basolateral side, and its partial entrance into the brain. A portion of Aβ at the level of capillary basement membrane diverges from this route, and continues moving along this membrane and basement membranes of the perivascular pathway. In the case of elevated Aβ concentration or the decline of the perivascular drainage, Aβ local accumulation and aggregation can occur. This can also happen in the case of the decline of the Aβ drainage system, and generally, is strong. The expression of ABCA1 declines with aging and in AD 31).

LRP1 (low-density lipoprotein receptor-related protein-1) is a large endocytic receptor for many ligands located on the luminal and abluminal cell membrane of the BBB endothelial cells. It is involved in the lysosomal endocytosis and transcytosis of Aβ which enters into the cells from the capillary basement membranes. In these membranes Aβ arrives from the blood by RAGE activity. Bound with the angiopep 2, it can be transported from the blood into the brain, through the cell, as covalently conjugated drug. LRP1 expression levels decline in AD and in the course of aging. LRP1 is predominantly generated in the neurons and secreted into the interstitial fluid (ISF). It is abundantly expressed in neurons, glial cells and VSMCs. It has a great importance in maintaining brain homeostasis. After Aβ endocytosis, it efficiently recycles back to the cell surface 32,33).

LDLR (low-density lipoprotein receptor) and SR-B1 (scavenger receptor class B type 1) are surface receptors, located on the luminal side of the endothelial cell membrane. LDL and HDL particles are mainly synthesized in the liver and are carried through the blood into the brain, through the cell, as covalently conjugated drug. LDLR expression levels decline in AD and during aging 34,35). The receptor solute carrier family 40 member ferroportin1 (FPN1), is embedded in the abluminal side of the endothelial cell membrane. It transports Fe2+ out from the cell. Immediately after its exit, Fe2+ becomes oxidised by hephaestin (Hp) in Fe3+, and binds with transferin (Tf), locally present in the brain. Iron in the form of Fe3+ cannot be transported along the perivascular drainage pathway. In the case of the drainage drop, Fe3+ also accumulates, and connected with the elevated values of Aβ, is reduced in Fe2+ and enters the Fenton reaction. The *OH generation is the normal subsequent event. It is the only known iron exporter. Hphaestin is a transmembrane copper-dependent ferroxidase, functionally closely bound with the FPN1 receptor, responsible for the oxidation (electron loss) and transport of ferrous iron (Fe2+) from the cell interior across its abluminal cell membrane. The obtained Fe3+ binds promptly with the locally present transferrin (Tf)36).

The receptor P-glycoprotein1, ATP-binding cassette sub-family B member1 (ABCB1) is located on both sides of the cell membrane. Coordinated with LRP1, this receptor can transport intracellular Aβ into the vascular lumen. Its role on the abluminal side is a questionable, but it is possible that here, it blocks the entry of Aβ into the cell. In AD and during aging its expression declines 37,38).

The danger signal receptor (i.e., receptor for advanced glycation end products, RAGE) and explain its function. It is not exactly confirmed decrease in the sTfR concentration. (s, serum)40-42).

The transferin receptor (TfR) imports iron (Fe3+) into the cell by internalizing the Fe3+/transferin (Tf) complex through the receptor-mediated endocytosis. It is a transmembrane glycoprotein, located on the luminal and abluminal sides of capillary endothelial cells. According to some studies, it seems that during aging, there is a not exactly confirmed decrease in the sTfR concentration. (s, serum)40-42).

Events related to the scavenger receptor class B type 1(SR-B1) are explained in detail in the paragraph about the ApoE4 apolipoprotein lipidation (Fig.4).

The current prevention and therapy for AD

Unfortunately, it is evident that nowadays there still does not exist an effective prevention and therapy for AD and. The actual therapy for AD with cholinesterase blockers (donepezil, rivastigmine, galantamine) is not satisfactory, and a number of other current therapeutics have a problematic effect. Currently, there are also attempts with immunotherapy, therapy with free radical scavengers and chelators (for example curcumin). The therapy
with crosslinker breakers is also widely used. The correction of the fasting glucose level (nonenzymatic glycosylation of Aβ retardation) has an important preventive role. Generally speaking, presently the therapy of AD includes antioxidants, AGE breakers, RAGE blockers and antiglycation compounds. Nowadays, there are actual investigations about the effects of alagebrium (ALT-711), aminoguanidine, DPTC, tiamine, benfotiamine, and piridoxamine. Related to the extracellular Aβ degradation, the effects of nepriysine (metalloendopeptidase) and matrix-metallopeptinases 2, 3 and 9 inhibitors, are intensively investigated, as well as the functions of proteasomes and the ubiquitin-proteasome route. It is also necessary to mention vitamin C, vitamin E, superoxide dismutase (SOD), curcuma, Ginkgo biloba, and green tea (among the polyphenols found in green tea the most important is epigallocatechin-EGCG). In this paragraph regarding the current prevention and therapy it is necessary to emphasize the memantine as N-methyl-D-aspartate receptor (NMDAR) blocker. By this action it protects the brain cells from the harmful effects of, in AD, elevated concentrations of released glutamate from damaged brain cells. It is also important to emphasize the cholinesterase blockaders: Aricept (donepezil hydrochloride), Axelon (rivastigmine), and Raminyl (galantamine hydrochloride). Apart from acetylcholine blockers and NMDA receptor antagonists, there are no other generally accepted medications or supplements. Previous researches on antioxidants, AGE breakers, RAGE blockers and antiglycation compounds haven’t reached scientific evidence to be accepted as standard therapy for AD. It is justly to hope that the problem of the effective prevention and therapy of AD will be solved in the near future.

**Conclusion**

Due to the great importance of the Aβ accumulation and aggregation in the AD brain, it is evident and logical that a number of recent investigators and their institutions increasingly take part in the research of the etiology, genetics, and pathophysiology of this disease. These investigations are especially related to the failures of perivascular drainage and modes of its improvement.

**Conflict of Interest Statement**

The author states that the performance of this review entailed no issues representing a conflict interest.
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