Review

Epigenetic Factors in Late-Onset Alzheimer’s disease: MTHFR and CTH Gene Polymorphisms, Metabolic Trans-sulfuration and Methylation Pathways, and B vitamins

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Abstract: DNA methylation and other epigenetic factors are important in the pathogenesis of late-onset Alzheimer’s disease (LOAD). Methylene tetrahydrofolate reductase (MTHFR) gene mutations occur in most elderly patients with memory loss. MTHFR is critical for production of S-adenosyl-L-methionine (SAM), the principal methyl donor. A common mutation (1364T/T) of the cystathionine-γ-lyase (CTH) gene affects the enzyme that converts cystathionine to cysteine in the trans-sulfuration pathway causing plasma elevation of total homocysteine (tHcy) or hyperhomocysteinemia – a strong and independent risk factor for cognitive loss and AD. Other causes of hyperhomocysteinemia include aging, nutritional factors, and deficiencies of B vitamins. We emphasize the importance of supplementing vitamin B12 (methylcobalamin), vitamin B6 (pyridoxine), and SAM to patients in early stages of LOAD.

Keywords: Alzheimer’s disease; CTH gene; DNA methylation; epigenetics; epigenome-wide association study; methylome; MTHFR gene; nutrition; S-adenosylmethionine; vitamin B complex

1. Introduction

Most genetic research on late-onset Alzheimer’s disease (LOAD) has focused on genome-wide association studies (GWAS) that in general have provided low effect size results, with the exception of apolipoprotein E (ApoE) [1,2]. Studies of monozygotic twins with Alzheimer’s disease (AD) showed discordance in onset and progression indicating a role for non-genetic factors in disease pathogenesis [3]. For these reasons, in the last few years genetic research turned to epigenetic modifications using epigenome-wide association studies (EWAS) [4,5].

Bonasio et al [6] defined epigenetics as “the study of molecular signatures that provide a memory of previously experienced stimuli, without irreversible changes in the genetic information.” Therefore, epigenetic refers to potentially heritable and non-heritable modifications in gene expression induced by environmental factors without changes in DNA base sequences [1,2]. These epigenetic processes include DNA methylation, histone modification and expression of long non-coding RNAs and non-coding microRNAs (miRNAs) that primarily repress target messenger RNAs (mRNAs) [1]. In AD, the miRNA-125b is overexpressed enhancing neuronal apoptosis and tau phosphorylation by activation of cyclin-dependent kinase 5 (CDK5) and p35/25. Forkhead box Q1 (FOXQ1) is the direct target gene of miR-125b [7]. The miR-125b has been found to be overexpressed
and circulating in patients with cardiovascular diseases and cancer [8]. Epigenetics has been extensively used in oncology, but epigenetic markers have been demonstrated to be also important regulatory factors of brain function [9], particularly in AD and other neurodegenerative diseases, as well as in aging. Experimental anti-aging epigenetic interventions attempt to reverse age-related changes in DNA methylation [10].

This review focuses on DNA methylation dynamics and other epigenetic changes, including the role of methylenetetrahydrofolate reductase (MTHFR) gene polymorphisms and its metabolic pathways particularly in aging and LOAD pathology [11], as well as polymorphisms of the cysteathionine-gamma(γ)-lyase (CTH) gene [12] the enzyme that converts cystathionine to cysteine in the trans-sulfuration pathway and is responsible for plasma elevation of total homocysteine (tHcy). Also, we review relevant nutritional factors including folate, vitamin B₁₂, and vitamin Be status, as well as hyperhomocysteinemia—an independent vascular risk factor linked to coronary disease, stroke, dementia, as well as cognitive impairment in the elderly and LOAD. Hcy is important in oxidative stress contributing to the decrease of S-adenosyl-L-methionine (SAM) levels, which induce demethylation of DNA resulting in overexpression of genes involved in AD pathology such as presenilin (PSEN1) and beta-secretase (BACE1), the β-site APP-cleaving enzyme that increases hypomethylation and Aβ42 deposition [9].

2. DNA Methylation Studies

5-cytosine methylation and DNA methyltransferases. Methylation at the 5-position of the cytosine base (5mC) is considered a critical phase of epigenetic regulation [1] and 5mC mutations introduced into the germline produce severe developmental restriction [13] and finally a lethal phenotype [14]. Cytosine base methylation occurs mainly at cytosine-phosphate-guanine (CpG) dinucleotides [1]. Gene regulation is achieved by 5mC silencing gene expression via high-density CpG areas, known as CpG islands, which remain largely unmethylated [9]. In humans, genomic DNA methylation of cytosine results from the addition of a methyl group from SAM to the cytosine, catalyzed by DNA methyltransferases (DNMT1, DNMT3A, and DNMT3B) [9]. In addition to 5mC, hydroxymethylation at the 5-position of the cytosine base (5hmC) derived from the oxidation of methylated cytosines by ten-eleven translocation (TET) enzymes is another epigenetic regulatory mechanism, which is particularly abundant in the brain [9].

In humans, DNA methyltransferases are involved in tumor transformation and progression resulting in genome-wide hypomethylation of tumor cells and silencing of tumor-suppressor genes [15]; also, DNMT3A mutations have been associated with poor prognosis in acute myeloid leukemia [17]. DNMT1 mutations occur in hereditary sensory and autonomic neuropathy type 1 (HSAN1) [9]. In mice, DNMT1 mutations induce global hypomethylation along with cortical and hippocampal neuronal dysfunction causing neurodegeneration with severe deficits in learning, memory, and behavior [16]. Hypomethylated excitatory neurons have postnatal maturation defects including abnormal dendritic arborization and impaired neuronal excitability [16].

Grossi et al [17] used artificial neural network analysis to illustrate how low cobalamin; low folate and high Hcy are linked to AD. Low PSEN1 methylation was linked to low folate levels and low promoter methylation of BACE1 and DNMT genes. High levels of folate-vitamin B₁₂ and low Hcy promoted methylation of genes required for DNA methylation reactions (DNMT1, DNMT3A, DNMT3B, and MTHFR) [18].

DNA methylation in Alzheimer's disease. Early studies of DNA methylation in LOAD from peripheral blood lymphocytes [19,20], brain biopsies and autopsy material [21-29], demonstrated variable results of cytosine methylation at CpG dinucleotides. Wang et al [30] studied postmortem pre-frontal cortex tissue and peripheral lymphocytes of AD patients and showed that specific loci in MTHFR gene promoter regions were hypermethylated compared to healthy controls. Ellison et al [31] using gas chromatography/mass spectrometry found abnormal levels of 5mC and 5hmC in the superior and middle temporal gyri, hippocampus and parahippocampal gyrus in early stages of AD, as well as in frontotemporal lobe degeneration and Lewy body dementia; these global values returned to control levels as the disease progressed suggesting that methylation changes occur in
early stages of neurodegenerative dementias. Chouliaras et al [32] confirmed the presence of significant decreases in levels of 5mC and 5hmC in the hippocampus of AD patients compared with negative controls. Levels of 5mC were inversely proportional to the deposition of neurofibrillary tangles in the same hippocampal cells. Hernández et al [33] studied DNA methylation patterns of cortical pyramidal layers in 32 brains of patients with LOAD demonstrating hypermethylation of synaptic genes and genes related to oxidative-stress including HOXA3, GSTP1, CXXC1-3 and BIN1.

One of the major problems of initial methylation studies was the small sample size. This was solved by De Jager et al [4] utilizing one of the largest clinicopathological studies to date, the Religious Orders Study, with 708 brains to assess the methylation state of the brain’s DNA correlated with AD pathology. Almost half million CpGs were interrogated including CpGs in the ABCA7 and BIN1 regions. The authors also identified genes whose RNA expression was altered in AD including ANKI, CDH23, DIP2A, RHBDL2, RPL13, SERPINF1 and SERPINF2. A companion study by Lunnon et al [5] found robust association between differences in methylation, mRNA levels, and tau-based Braak staging. Dysregulation of DNA methylation occurred earlier in brain areas affected at onset by AD and appeared to have stronger effects (28.7%) than the combination of ApoE and other risk genes (13.9%) identified by GWAS [1,2], indicating the importance of epigenetic changes in AD. Additional studies by Yu et al [34] confirmed the association of DNA methylation in SORL1, ABCA7, HLA-DRB5, SLC24A4, and BIN1 genes with pathological diagnosis of AD including both Aβ load and tau tangle density. RNA expression of transcripts of SORL1 and ABCA7 was associated with tau tangle density, and the expression of BIN1 was associated with Aβ load [34]. Moreover, Lunnon et al [5] found hypermethylation of the ankyrin 1 (ANK1) gene in the entorhinal cortex, superior temporal gyrus and prefrontal cortex in LOAD. These findings confirm that AD involves significant disruption of DNA methylation. Epigenetic age-associated alterations of DNA methylation have also been reported in animal models of AD, in particular global DNA hypomethylation in the J20 model and DNA hypermethylation in the triple transgenic 3xTg-AD model [35].

3. Trans-sulfuration metabolic pathways and remethylation defects

The metabolism of sulfur-containing amino acids in the trans-sulfuration pathway involves the transfer of the sulfur atom of methionine to serine to produce cysteine (Figure 1). Methionine first reacts with ATP to form S-adenosyl-L-methionine (SAM), then S-adenosyl-homocysteine (SAH) and finally, homocysteine. Plasma elevation of total homocysteine (tHcy) or hyperhomocysteinemia may result from congenital deficiency of cystathionine β-synthase (CBS) leading to homocystinuria, or more frequently from polymorphisms of the cystathionine gamma(γ)-lyase (CTH) gene (OMIM *607657; EC 4.4.1.1.) in chromosome 1 (1p31.1) [36]. CTH is the enzyme that converts cystathionine to cysteine, the last step in the trans-sulfuration pathway. Wang et al [12] demonstrated that a single nucleotide polymorphism (SNP), namely c.1364G>T in exon 12 of the CTH gene causes cystathioninuria and elevation of tHcy. In Caucasian subjects homozygous for the CTH 1364T/T SNP the elevation of tHcy reached effects sizes similar to those caused by the 677C>T MTHFR polymorphism [12].

Closely related to the trans-sulfuration pathway are the remethylation defects resulting from the failure to convert homocysteine to the amino acid methionine (Figure 1). This pathway requires the integrity of the gene encoding methylenetetrahydrofolate reductase (MTHFR) required for the interaction of folate and cobalamin (vitamin B12). Folate provides the methyl group required for the remethylation pathway (Figure 1) to finally produce SAM, the main methyl donor for epigenetic processes.

The human MTHFR gene (OMIM *607093; EC 1.5.1.20) is localized in chromosome 1 (1p36.3) and it encodes for 5,10-methylenetetrahydrofolate reductase (MTHFR) [37]. This enzyme catalyzes the conversion of 5,10-methylenetetrahydrofolate to 5-methyltetrahydrofolate, a co-substrate with vitamin B12 for the remethylation of homocysteine to methionine [11]. Mutations of this gene occur in 10-15% of the population and the resulting MTHFR deficiency affects the production of methionine and SAM. Linnebank et al [38] demonstrated a decrease of SAM in the cerebrospinal fluid (CSF) of patients with LOAD, mainly among ApoE ε4 carriers.
MTHFR gene polymorphisms cause enzyme thermolability and involve C-to-T substitution at nucleotide 667 and A-to-C at nucleotide 1298; these MTHFR mutations have been associated with homocystinuria, neural tube defects, preeclampsia, cleft lip and cleft palate, cerebrovascular disease, and psychiatric disorders including susceptibility to depression and schizophrenia [39,40]. Population-based international studies showed no increased risk of dementia in subjects with MTHFR polymorphisms [41,42]. In Japan, Nishiyama et al [43] found a slight association of the MTHFR-C667T polymorphism with senile cognitive decline in men but not with AD. In Australia, a causal link between high tHcy and incident dementia was demonstrated [44] but the study lacked power to determine an effect of the MTHFR-C667T genotype. In contrast, in the normal elderly population of the Rotterdam Study de Lau et al [45] observed that the MTHFR-C667T genotype was associated with elevated tHcy but not with cognitive loss or white matter lesions. In a small patient population in Tunisia [46], the MTHFR-A1298C mutation was associated with susceptibility to AD. As mentioned earlier, Román [11] found a very high frequency (above 90%) of MTHFR gene mutations in an elderly population attending a memory clinic in the USA, with diagnoses ranging from mild cognitive impairment (MCI) to LOAD; about 65% had single mutations; the MTHFR-C667T mutation was found in 58.5% of the patients and 41.5% had the MTHFR-A1298C mutation whereas 20% were compound heterozygous for both mutations [11]. MTHFR and epigenetic drift. In 2005, a multinational study of identical twins by Fraga et al [47] first demonstrated that whereas DNA methylation and histone acetylation in young identical twins are indistinguishable, older identical twins showed substantial differences; epigenetic changes were up to four times greater than those of young twin pairs. The authors concluded that this “epigenetic drift” was associated with aging [47]. Epigenetic drift of identical twins with aging also occurs among a large number of animal species [48] following a non-Mendelian pattern. In identical twins with AD, the prognosis and onset of AD can differ by more than ten years [3,49-53]; young identical twin pairs are essentially indistinguishable in their epigenetic markings while older identical twin pairs show substantial variations. Breitner et al [50,53] suggested that twins with a history of systemic infection developed AD at an earlier onset than their identical twin. Epigenetic drift can be caused by lifestyle, diet, infections, folate status, homocysteine status, or toxic exposure [51]. Wang et al [52] demonstrated that the MTHFR gene promoter in the brain displayed high interindividual variance in DNA methylation among twins. The methylation level of MTHFR and APOE in individuals 30 years of age apart decreased by 10.6%, whereas in patients with AD the methylation level increased by 6.8%. The epigenetic drift increases with age particularly in genes that play pivotal roles in removing β-amyloid such as PSEN1 and APOE and among methylation genes such as MTHFR and DNMT1 [9,54].
**Figure 1**: Homocysteine metabolism: B12=cobalamin. B6=pyridoxine. MTH=methylenetetrahydrofolate. MTHFR=methylenetetrahydrofolate reductase. SAM=S-adenosylmethionine. SAH=S-adenosylhomocysteine. 5-Me THF=5-methyltetrahydrofolate. (From Spence, J.D.; Yi, Q.; Hankey, G.J. B vitamins in stroke prevention: Time to reconsider. *Lancet Neurol.* 2017, 16, 750–760.

4. Homocysteine: A risk factor for cognitive loss and dementia

Hcy is a sulfur-containing amino acid produced in the *trans-sulfuration pathway* (Figure 1) from the reaction of methionine with ATP to form SAM, then SAH and finally homocysteine. Homocystinuria due to congenital deficiency of the CBS gene causes hyperhomocysteinemia. Polymorphisms of the CTH and MTHFR genes are common genetic causes of hyperhomocysteinemia [36,37]. The *remethylation pathway* (Figure 1) involves reactions enzymatically mediated by MTHFR requiring as co-substrates the B-group vitamins folic acid (vitamin B₉) and cobalamin (vitamin B₁₂) for the remethylation of homocysteine to methionine. Pyridoxine (vitamin B₆) is required by CBS for the conversion of homocysteine to cysteine (Figure 1).

Elevation of plasma or serum tHcy (hyperhomocysteinemia) is an independent vascular risk factor linked to coronary disease, peripheral vascular disease, stroke and small-vessel cerebrovascular disease [55]. More importantly, elevated tHcy is considered a risk factor for dementia and cognitive decline in the elderly, particularly in association with low levels of folate and cobalamin [56]. A number of studies in cognitively normal elderly subjects, demonstrated that baseline tHcy is a strong and independent predictor of cognitive decline after observation periods ranging from 3 years (USA, n=321 men [57] and Sydney, Australia, n=889 [58]); 4 years (France, n=1241) [59]; 5 years (Wales, United Kingdom, n=32) [60]; 6 years (Norway, n=2,189) [61]; 7 years (Finland n=274) [62], up to 10 years (United Kingdom, n=691) [63]. In the Finland cohort [62], the MRI study demonstrated the association of higher baseline vitamin B₁₂ and holotranscobalamin levels with a decreased rate of total brain volume loss during 8 years of the study period [64]. Increased tHcy levels were associated with faster rates of total brain volume loss and with progression of white matter hyperintensities among participants with hypertension (systolic blood pressure > 140 mm Hg) [64].

Regarding the risk of AD associated to elevated tHcy, in the Framingham Study, Seshadri and colleagues [65] demonstrated in elderly subjects (mean age, 76 years) that raised tHcy above 14 μmol/L nearly doubled the risk of LOAD over a period of 8 years. Similar findings were corroborated.
in two large Finnish [62,64,66] and Australian [67] cohorts. In 2008, Smith [68] performed a comprehensive review of cross-sectional and prospective studies involving >46,000 subjects and confirmed the association between elevated tHcy and cognitive deficit or dementia.

According to a recent international consensus statement [69], moderately raised homocysteine (>11 μmol/L) increases the relative risk of dementia in the elderly 1.15 to 2.5 fold, and the Population Attributable risk from 4.3 to 31% [69]. From the Public Health viewpoint, homocysteine-lowering treatment with B vitamins that markedly slows down the rate of brain atrophy and cognitive decline in the elderly offers the possibility that, in addition to folic acid fortification, mandatory methylcobalamin supplementation should also be considered for the prevention of LOAD [44,62,68,69].

Elevation of tHcy is caused by numerous factors including advancing age, diet, supplementation of B-vitamins, obstructive sleep apnea, smoking, Helicobacter pylori infection, and renal failure, among others [55,56]. As indicated earlier, both CBS gene polymorphisms and the C667T and the A1298C S4-NPs in the MTHFR gene decrease the activity of the MTHFR enzyme leading to hyperhomocysteinemia.

### Table 1. Harmful effects of homocysteine on vascular function and cognition (Modified from Smith & Refsum [56])

<table>
<thead>
<tr>
<th>Proposed mechanisms</th>
<th>Vascular Mechanisms</th>
<th>Neuronal Mechanisms</th>
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<td></td>
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<tr>
<td>1</td>
<td>Impairs endothelial function reducing inducible NO synthase</td>
<td>Direct activation of NMDA receptor causes excitotoxic neuronal death</td>
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<tr>
<td>2</td>
<td>NO-mediated endothelial dysfunction in brain vasculature</td>
<td>Homocysteic acid and cysteine sulfenic acid activate NMDA receptor causing neuronal death by excitotoxicity</td>
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<td>3</td>
<td>Causes a leaky blood-brain barrier</td>
<td>Oxidative stress induced by generating superoxide and reactive oxygen species</td>
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<td>4</td>
<td>Induces thrombosis</td>
<td>Decreased activity of antioxidant enzymes</td>
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<td>5</td>
<td>Cerebrovascular ischemia leading to neuronal death and tau tangle deposition</td>
<td>Formation and deposition of β-amyloid</td>
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<td>6</td>
<td>Affects lipid metabolism increasing cholesterol synthesis</td>
<td>Potentiates neurotoxic effects of β-amyloid by itself or via homocysteic acid</td>
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<td>7</td>
<td>Reduces synthesis of apolipoprotein 1</td>
<td>Activates tau kinases, such as Cdk5, causing tau tangle deposition</td>
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<tr>
<td>8</td>
<td>Causes cerebral amyloid angiopathy</td>
<td>Triggers the cell cycle in neurons, leading to tangle formation and cell death</td>
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<td></td>
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<td>Causes DNA damage, limits DNA repair, leading to apoptosis</td>
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<td></td>
<td>Increases SAH inhibiting methylation reactions, such as DNA cytosine methylation in promoters for amyloid genes, causing epigenetic effects</td>
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<td></td>
<td>Inhibits PP2A activity leading to tau tangle deposition</td>
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<td></td>
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<td>Inhibits methylation of phosphatidylethanolamine</td>
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<td>Stimulates endoplasmic reticulum stress response leading to amyloid formation</td>
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<td></td>
<td>Activates the immune system</td>
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<td></td>
<td></td>
<td>Decreases SAM-dependent synthesis of catecholamines and other neurotransmitters</td>
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</table>
Smith and Refsum [56] reviewed the proposed mechanisms responsible for the harmful cognitive effects of hyperhomocysteinemia (Table 1). These include impaired endothelial function with reduced inducible nitric oxide synthase; augmented oxidative stress and decreased activity of key antioxidant enzymes; raised generation of the superoxide anion; alterations of lipid metabolism with increased cholesterol synthesis and reduced synthesis of apolipoprotein 1; and, carotid stenosis and induction of thrombosis [55,56].

Minagawa et al [70] found that elevated Hcy inhibits the dimerization of ApoE3 and reduces ApoE3-mediated high-density lipoprotein (HDL) concentrations involved in degradation of soluble Aβ within microglia. ApoE4 was not affected; in patients with hyperhomocysteinemia the CSF levels of ApoE3 dimers were significantly lower than in controls. Minagawa et al [70] suggested that the effects of elevated Hcy on ApoE3 contribute to the pathogenesis of AD.

Hyperhomocysteinemia induces a decrease in the SAM-dependent synthesis of catecholamines including dopamine, norepinephrine, and epinephrine, as well as non-catecholamine neurotransmitters such as melatonin and serotonin (5-HT) that contribute to development of depression [71]. Moreover, elevated tHcy produces two neurotoxic products, homocysteic acid (HCA) and cysteine sulfonic acid (CSA), which are agonists of the N-methyl-D-aspartate (NMDA) glutamate receptor, with neurotoxic effects on dopaminergic neurons derived from excessive Ca++ influx and reactive oxygen generation [72]. The beneficial effects of B-group vitamins on elevated tHcy will be reviewed next.

5. Folate metabolism

Vitamin B9 or folic acid (from the Latin folium, leaf) is abundantly found in green leafy vegetables. Folate is vital for cell development and growth given its role in numerous biochemical one-carbon (methyl-group, -CH3) reactions, many of them critical for cognition. The Nun Study [73] first provided epidemiological and neuropathological data demonstrating that limited lifetime consumption of salads with low blood folate levels increased the risk of cognitive decline and dementia. Also, the severity of the atrophy in the neocortex and of the Alzheimer disease lesions were strongly correlated with low serum folate levels; none of 18 other nutrients, lipoproteins, or nutritional markers measured in the study correlated with the atrophy [73]. Further studies confirmed that normal cognitive scores were highly associated with elevated blood folate despite the neuropathological evidence of LOAD brain lesions [74].

The primary methyl-group donor for DNA methylation reactions is 5-methyl-tetrahydrofolate (CH3THF) required for the transformation of homocysteine into methionine mediated by methionine synthase with cobalamin (vitamin B12) as a co-substrate (Figure 1), leading to the synthesis of SAM.

Also, CH3THF is critical in the de novo purine synthesis to convert dUMP (deoxyuridylate) into dTMP (thymidylate) for DNA and RNA synthesis, DNA repair or replication. Several forms of cancer are associated with epigenetic differential methylation causing disturbances in nucleotide synthesis; for instance, hypermethylation may inhibit tumor suppressors. Folate, therefore, is an important epigenetic determinant in gene expression, DNA stability, DNA integrity and mutagenesis. Abnormal folate status is an important factor in neural tube defects, cardiovascular and cerebrovascular diseases, cleft lip and palate, neurodegenerative diseases, schizophrenia, and depression [75–77].

Low folate levels are associated with short telomeres due to DNA damage in the telomeric region. Telomere length is epigenetically regulated by DNA methylation and directly influenced by folate status, a process independent of DNA damage due to uracil incorporation. Shorter telomeres occur with age, infection, stress, and chronic diseases including LOAD [78]. Paul et al [79] observed that decreased plasma folate concentration to <11.6 μmol/L was correlated with a decrease in mean telomere length. In this population, carriers of homozygous MTHFR C677T gene mutation showed decreased levels of plasma folate [80]. Decreased serum folate induces anomalous integration of uracil in place of thymidine in DNA [81], a mechanism corrected by folic acid supplementation.

Troesch et al [82] summarized the importance of reduced SAM-dependent methylation reactions, due to genetic factors along with reduction of folate, vitamin B9 and vitamin B12 levels, for the
development of LOAD. The resulting elevation of Hcy levels and the reduced capacity to synthesize, methylate and repair DNA, along with the impaired modulation of neurotransmission, appears to favor the development of AD particularly when combined with increased oxidative stress, particularly in ApoE ε4 carriers [83].

6. Vitamin B12 Deficiency and β-amyloid deposition

Smith, Warren and Refsum [84] have recently provided a comprehensive review of vitamin B12. Only bacteria can biosynthesize vitamin B12; in humans B12 from the diet is a cofactor for the enzymes methionine synthase and L-methyl-malonyl-CoA mutase. B12 deficiency results in build-up of homocysteine and lack of interaction with folate that is trapped as CH3-THF leading to depletion of tetrahydrofolates used in thymidylate and purine synthesis blocking DNA for the production of red cells in the bone marrow. B12 deficiency impedes cellular proliferation and protein synthesis and thereby causes development of megaloblastic anemia [84]. In 1920, pernicious anemia was successfully treated by adding liver to the diet. In 1955, Dorothy Hodgkin used crystallography to first identify the molecular structure of cyanocobalamin or vitamin B12 from the deep-red cyanide-containing pigment isolated from liver tissue. Pernicious anemia was the first disease identified as caused by vitamin B12 deficiency [84].

Stabler [85] reviewed the clinical manifestations of vitamin B12 deficiency. In addition to megaloblastic anemia, acidemia from elevation of serum methylmalonic acid (MMA), and methylmalonic aciduria, the neurological manifestations of pernicious anemia include memory loss and cognitive decline, visual disturbances from optic nerve neuropathy, burning and painful sensations in hands and feet from peripheral neuropathy, and spinal cord involvement with subacute combined degeneration resulting in loss of proprioception from dorsal column involvement and pyramidal tract symptoms such as paralysis and incontinence.

Dietary sources of B12 include liver, meat, fish, shellfish, and dairy products; vegans are prone to B12 deficiency [84,85]. Vitamin B12 deficiency occurs from inborn metabolic errors, alterations of B12-binding proteins including haptocorrin (HC) found in saliva, intrinsic factor (IF) produced by parietal cells in the stomach (pernicious anemia is associated with anti-parietal-cell and anti-IF auto-antibodies), and transcobalamin (TC) which binds B12 to facilitate uptake by the cells [84]. According to Stabler [85], measurement of total serum B12 levels is unsatisfactory because it reflects B12 that is bound to either HC or TC and up to 60% of bound materials are cobalamin analogues (corrinoids). Therefore, “normal” total serum B12 levels can mask deficiency if serum contains relatively large amounts of cobalamin analogues [84]. Levels below 200 pg/mL usually indicate biochemical B12 insufficiency. Serum B12 <350 pg/mL along with tHcy >14 μmol/L indicate metabolic B12 deficiency [84,85]. For this reason, holotranscobalamin, MMA and tHcy levels should be included in the evaluation of a patient suspected of having B12 deficiency [86]. Other causes of B12 deficiency include atrophic body gastritis, malabsorption of vitamin B12, gastrectomy, gastric bypass or other bariatric surgery, inflammatory bowel disease, tropical sprue, use of metformin, anticonvulsants, drugs to block stomach acid, and vegetarian diets low in meat and dairy products. Hemodialysis patients, nitrous oxide inhalation, and cholinesterase inhibitors in LOAD patients [87] also increase the risk of vitamin B12 deficiency. Epidemiological studies have shown that prevalence of vitamin B12 deficiency increases with age [88,89], due to decreased saliva (dry eyes-dry mouth) and gastric atrophy with deficits respectively of haptocorrin and intrinsic factor. Andrès et al [90] have emphasized that as many as 20% of elderly people may have unrecognized B12 deficiency due to food-cobalamin malabsorption plus insufficient dietary intake. According to Spence [91], metabolic B12 deficiency occurs in 30% of vascular patients older than 71 years, increasing to as many as 40% in patients above age 80 years; these patients usually have plasma levels of tHcy >14 μmol/L resulting from B12 deficiency. Inadequate supply of B12 and folic acid is not only a strong and independent vascular risk factor but it is also responsible for cognitive impairment and memory complaints in the elderly promoting the development of LOAD [92]. Animal experimental data confirms the importance of B-vitamin deprivation in the expression of AD [93].
Despite the negative results of meta-analyses reviewing results from inadequately controlled clinical trials [94], solid positive results such as those of the OPTIMA trial [95–97] indicate that supplementation of B12, pyridoxine, and folic acid in subjects with MCI and hyperhomocysteinemia decreases tHcy resulting in improved episodic memory and global cognition and, most importantly, halting the progression of the brain atrophy in areas affected by AD [97]. Current recommendation is to provide oral supplementation of methylcobalamin 1000 μg/d, folic acid 800 μg/d and pyridoxine 100 mg/d.

7. Conclusions

It is well established that the damaging effects of deficiencies of folate and cobalamin and the resulting elevation of tHcy contribute to the development of LOAD [69]. The numerous detrimental effects of elevated tHcy include, among others, endothelial and cerebrovascular damage of large-vessels as well as small-vessel disease [98]; activation of tau kinases; inhibition of methylation reactions; epigenetic effects on the β-amyloid pathway; reduced protein phosphatase-2A; and, impaired formation of phosphatidylcholine. Adequate supply of B-vitamins in the elderly, particularly in subjects with MTHFR and CTH gene mutations, appears to be critical to prevent the development of cognitive decline and to halt the progression of LOAD.

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