Markers of Oxidative Stress and Neuroprogression in Depression Disorder

Magdaléna Vaváková,1 Zdenka Řuračková,1 and Jana Trebatická2

1Institute of Medical Chemistry, Biochemistry and Clinical Biochemistry, Faculty of Medicine, Comenius University, Sasinkova 2, 813 72 Bratislava, Slovakia
2Department of Pediatric Psychiatry, Medical Faculty, Comenius University and Child University Hospital, Faculty of Medicine, University Hospital, Limbová 1, 833 40 Bratislava, Slovakia

Correspondence should be addressed to Jana Trebatická; jana.trebaticka@fmed.uniba.sk

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Major depression is multifactorial disorder with high prevalence and alarming prognostic in the nearest 15 years. Several mechanisms of depression are known. Neurotransmitters imbalance and imbalance between neuroprogressive and neuroprotective factors are observed in major depression. Depression is accompanied by inflammatory responses of the organism and consequent elevation of proinflammatory cytokines and increased lipid peroxidation are described in literature. Neuropsychiatric disorders including major depression are also associated with telomerase shortening, oxidative changes in nucleotides, and polymorphisms in several genes connected to metabolism of reactive oxygen species. Mitochondrion dysfunction is directly associated with increasing levels of oxidative stress. Oxidative stress plays significant role in pathophysiology of major depression via actions of free radicals, nonradical molecules, and reactive oxygen and nitrogen species. Products of oxidative stress represent important parameters for measuring and predicting of depression status as well as for determining effectiveness of administrated antidepressants. Positive effect of micronutrients, vitamins, and antioxidants in depression treatment is also reviewed.

1. Introduction

Major depression is a mental disorder that occupies fourth position of worldwide disability list and is expected to become the second most frequent disease by 2030 [1]. Major depression not only decreases productivity and quality of life of patients but also represents a significant financial burden for health care [2, 3]. Depression is multifactorial disorder and its etiology includes genetics, environmental, psychological, and biological factors. Several molecular mechanisms play role in pathogenesis of depression. Depression is caused by abnormalities in metabolism of neurotransmitters (e.g., serotonin and dopamine) that is consequently affected by enzymes involved in their degradation (e.g., monoamine oxidase) or synthesis of their precursor tryptophan. To increase concentration of serotonin in the synaptic cleft is successful strategy for treatment of major depression via selective serotonin reuptake inhibitors (SSRI) antidepressants. SSRI represents the most frequently used antidepressants for treatment of major depression (88.5%) [2, 4–6]. Major depression is also associated with inflammatory processes and elevated levels of proinflammatory cytokines, decreased neurogenesis, and subsequent neuroprogression (pathological reorganization of the central nervous system), mitochondrial dysfunction, and dysfunction of hypothalamic-pituitary-adrenal axis. There are observed lowered antioxidant concentrations and increased oxidative stress levels as well [5, 7–9].

Oxidative stress is connected with many diseases including atherosclerosis, cardiovascular and neurodegenerative diseases, like Alzheimer’s disease and Parkinson’s disease, psychiatric disorders like Attention Deficit Hyperactivity disease and schizophrenia, or cancer, diabetes mellitus, and others. Oxidative stress is caused by imbalance between prooxidant and antioxidant in favor of prooxidant. Oxidative damage to cells and organs can be caused by free radicals (FR) (superoxide, hydroxyl radical) or nonradical molecules, like...
hydrogen peroxide, and their derivatives called reactive oxygen species (ROS) and reactive nitrogen species (RNS). They are expressed in the organism during standard metabolism. Mild oxidative stress and free radicals play an important role in the regulation of many processes in the organism, for example, during phagocytosis, apoptosis, egg fertilization, or activation of certain transcripational factors or in cell signaling pathways [10–13].

However, if ROS and RNS are produced in large amount and in the wrong place, they can cause oxidative modifications of lipids, proteins, and DNA [9, 14–16]. They can modify cell membranes and function of receptors and alter enzymes and genes activity. Oxidative stress contributes also to the aging [8, 11, 16–20]. Direct measurement of ROS concentration is difficult because of its extremely short half-lives. Obvious alternative option is to measure products of oxidative stress in proteins, lipids, or nucleic acids [12, 21].

To fight excessive production of ROS and RNS, the organism has built protective systems and mechanisms against their toxic effects. Protection is organized at three levels: (a) systems preventing FR formation, such as inhibitors of enzymes catalyzing FR formation. (b) When these primary protective systems are insufficient and FR and ROS are already formed, scavengers and trappers of FR come into action and eliminate high reactivity of ROS by turning them into nonradical and nontoxic metabolites. These compounds are called antioxidants and they prevent oxidation of biologically important molecules by FR or ROS. (c) If protection of the organism fails at this level, then repair systems recognize impaired molecules and decompose them, as it is in case of proteinases at oxidatively modified proteins, lipases at oxidatively damaged lipids, or DNA repair systems at modified DNA bases [13].

### 2. Neuroprogressive Processes and Hormones Activity in Depression

Balance between neuroprogressive and neuroprotective factors is established by grown and neurotrophic factors. Their expressions are also affected by stress (Table 1, Figure 1) [5].

Brain-derived neurotrophic factor (BDNF) is the most abundant neurotrophic factor in brain. Lowered concentration in serum is typical state for patient with depression and bipolar disorder. There is also correlation between changes in concentration of BDNF and severity of illness. Lowered concentration of BDNF increased level of oxidative stress which consequently affects expression, folding, and secretion of BDNF. These changes lowered neurogenesis of brain cells.

#### Table 1: Summarizing the main findings related to oxidative stress and major depression.

<table>
<thead>
<tr>
<th>Effect</th>
<th>Marker</th>
<th>Findings compared to controls</th>
<th>Subject</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neuroprogression</td>
<td>BDNF</td>
<td>↓</td>
<td>Human</td>
<td>[22–25]</td>
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<td>Signaling</td>
<td>5-HT</td>
<td>↓</td>
<td>Human</td>
<td>[26]</td>
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<td></td>
<td>NO</td>
<td>↑</td>
<td>Human</td>
<td>[27, 28]</td>
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<td>DNA damage</td>
<td>Telomerase</td>
<td>↓</td>
<td>Human</td>
<td>[29, 30]</td>
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<tr>
<td></td>
<td>8-OxoG</td>
<td>↑</td>
<td>Human</td>
<td>[31, 32]</td>
</tr>
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<td>Prooxidant enzymes</td>
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<td>↑</td>
<td>Human</td>
<td>[33–36]</td>
</tr>
<tr>
<td></td>
<td>MAO</td>
<td>↑</td>
<td>Human</td>
<td>[37]</td>
</tr>
<tr>
<td>Antioxidant enzymes</td>
<td>SOD</td>
<td>↑</td>
<td>Human</td>
<td>[38–40]</td>
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<tr>
<td></td>
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<td>↓</td>
<td>Human</td>
<td>[40–42]</td>
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<td>Catalase</td>
<td>↑</td>
<td>Human</td>
<td>[39]</td>
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<td>PON1</td>
<td>↓</td>
<td>Human</td>
<td>[43]</td>
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<td>↓</td>
<td>Human</td>
<td>[45]</td>
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<td>↓</td>
<td>Human</td>
<td>[46]</td>
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<td>↓</td>
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<td>[46–48]</td>
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<td>[38]</td>
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<td>TRYCATs</td>
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<td>[61–64]</td>
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<td>Human</td>
<td>[65]</td>
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<tr>
<td>Lipid peroxidation</td>
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<td>↑</td>
<td>Human</td>
<td>[39, 47, 66]</td>
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<tr>
<td></td>
<td>8-iso-P</td>
<td>↑</td>
<td>Human</td>
<td>[60, 67]</td>
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5-Hydroxytryptamine (5-HT), 8-oxoguanine (8-oxoG), 8-iso-prostaglandin F2 alpha (8-iso-P), brain-derived neurotrophic factor (BDNF), coenzyme Q10 (CoQ10), glutathione peroxidase (GPX), high density lipoprotein (HDL), immunoglobulin (Ig), monoamine oxidase (MAO), malondialdehyde (MDA), nuclear factor κ B (NFκB), paraoxonase 1 (PON1), superoxide dismutase (SOD), tryptophan catabolites (TRYCATs), and xanthine oxidase (XO).
An antidepressants usage increases concentration of BDNF and stimulates neurogenesis and regeneration of cells [5, 7, 22–25]. On the other hand, some studies fail to detect changes in concentration of BDNF in plasma between depressed patients and control group and between patients on antidepressants and medicine-free patients [68].

Probability of development of depression is increased with a lower level of insulin-like growth factor (IGF-I) in brain. An exact mechanism of how IGF-I influences depression is unknown. It is assumed that metabolic pathway of lithium efficacy in treatment of bipolar disorder is involved [69, 70].

Nuclear factor κ B (NFκB) is transcription factor that is also connected with development of depression. Proinflammatory cytokines (e.g., interleukin 6 and interleukin 8) are activated by NFκB. Activity of this transcription factor is affected by levels of ROS and glutamate. NFκB itself increases oxidative stress in the organism and causes inflammatory reaction which can lead to neuroprogression [5, 8, 70]. Increased levels of NFκB were observed in depressed patients [38].

An imbalance between production and release of neurotransmitters, like dopamine, serotonin (5-hydroxytryptamine), glutamate, or noradrenalin in brain, is also
associated with depression and neurodegenerative disease [4, 71].

Brain levels of neurotrophic factor 5-hydroxytryptamine (5-HT) affect second messenger signalization, expression of BDNF, and receptors for serotonin transporters. Lowered sensitivity of 5-HT receptors was detected in depressed patients and it is caused by inhibition of neurogenesis of 5-HT neurons [5, 26].

Secretion of melatonin affects a circadian cycle. Several depression symptoms, like restless sleep, early morning awakening, sleeplessness, daytime fatigue, and moodiness, are caused by interfering of day and night cycle. An antidepressant effect was recorded only after administration of agomelatine, a melatonin analogue. Improvement of patients with depression was markedly only in severe cases of depression. An antidepressant effect of agomelatine is presumably caused by synergic effect of melatonin (receptors MT1, MT2) and monoamine (receptors 5-HT2C) [4, 72–75].

Tryptophan catabolites (TRYCATs) are created by tryptophan breakdown by enzyme indoleamine 2,3-dioxygenase (IDO). Tryptophan, as a precursor of serotonin, is an important marker of depression. Its lowered level can lead to inflammatory reaction of the organism. IDO activity is negatively correlated with concentration of serotonin and tryptophan and is positively correlated with depression severity. Lowered levels of TRYCATs and tryptophan are also associated with melancholic depression. TRYCATs, like kynurenine, kynurenine acid, or xanthurenic acid, possess neuroprotective but also depressogenic, anxiogenic, and neurotoxic properties [4, 5, 61–64, 76].

Nitric oxide (NO), if it is produced in excess, is strong damaging-free radical. However, at low concentration NO is considered to be an important neurotransmitter connected to pathophysiology of depression, anxiety, epilepsy, and schizophrenia. NO is synthesized from L-arginine through constitutive and endothelial NO synthase (cNOS, eNOS) at physiological conditions. At some pathological conditions an inducible NO synthase (iNOS) forms high concentration of NO. However, at low concentration, NO acts as second messenger in the process of dopamine and noradrenergic releasing. NO affects sexual and aggressive behavior and via synthesis pathway also takes part in anxious behavior, inflammation, and depression elicited by interferon alpha (IFNα) [8, 17, 74, 77, 78]. Even temporary elevated levels of NO cause nitration and hypernitrosylation of amino acids and proteins. That leads to creation of highly reactive substances, for example, NO-tyrosine, NO-tryptophan, NO-arginine [79, 80], and NO-bovine serum albumin (NO-BSA) [81]. Indeed, Maes et al. [82] detected neoptepites and anchorage epitoites created by autoimmune response of IgM against palmitic and myristic acids and also elevated levels of IgM against NO-tyrosine, NO-aspartate, and NO-phenylalanine in serum of depressed patients [82]. Elevated concentration of NO was detected in patients with depression and in patients with suicidal thoughts compared to nonsuicidal patients [27, 28]. Antidepressant-like effect can be induced by inhibition of NO synthesis in brain. Inhibition of NOS can lead to increased effectiveness of serotonergic antidepressants and can be applied to patients suffering from drug resistant depression [8, 17]. Eight weeks of antidepressants administration leads to lowered concentration of NO [33].

3. Effect of DNA Damage on Depression

An excessive telomerase shortening is observed in several neuropsychiatric disorders like depression, schizophrenia, anxiety, and affective disorder, in consequence of lowered activity of telomerase. Length of telomeres is negatively correlated with longitude of nonmedicated depression and concentration of interleukin 6 (IL-6). IL-6 is considered to be a marker of inflammation. An exact mechanism of telomerase shortening and premature aging of the organism is unknown. Hypothesis said that oxidative stress and inflammation affect structure of replication fork in the vicinity of telomeres. Possibility is also that oxidative stress induced DNA damage preferably in telomeres in comparison to nontelomeric DNA [7, 8, 29, 30].

Oxidative changes in nucleotides are caused by ROS and RNS. 8-Oxodeoxyguanine (8-OxoG) is formed by oxidation of guanine and is diluted by urine. In urine of depressed patients significantly elevated levels of 8-OxoG are detected in comparison to healthy control [12, 17, 31, 32]. Another predisposition of depression can be a specific single nucleotide polymorphism (SNP). Change of nucleotide sequence can decrease activity of peptidases or influence several genes connected to metabolism of ROS and RNS. SNPs in genes for IL-1, IL-6, tumor necrosis factor alpha (TNFα), cyclooxygenase 2 (COX2), superoxide dismutase (SOD), and catalase is associated with depression [17, 61, 83]. SNPs in serotonergic genes can also alter antidepressants responses [84].

4. Oxidative Stress-Related Enzymes in Depression

Changes in activity of oxidative stress-related enzymes can be caused by depression and inflammation (Table 1, Figure 1). Several enzymes involved in ROS production showed elevated levels in depressed patients in comparison to nondepressed controls [7, 17].

Xanthine oxidase (XO) represents enzyme that catalyzes oxidation of xanthine. Products of this reaction are superoxide and hydrogen peroxide. Increased serum level of XO was observed in depressed patients and in thalamus brain area in postmortem depressed patients [5, 17, 33–36]. Oxidative deamination of monoamine neurotransmitters and metabolism of serotonin is affected by monoamine oxidase (MAO). Its byproducts (e.g., hydrogen peroxide) cause overproduction of ROS that can lead to neuron apoptosis and mitochondrial dysfunction [4, 7]. Elevated level of MAO was also detected in patients with depression and postpartum depression [37]. COX-2 inhibitors cause neuron inflammation that can consequently lead to worsening depression and increase probability of cardiovascular diseases. They can also damage mitochondrial, increase lipid peroxidation, and decrease antioxidant concentration [8, 9, 85]. Enzyme NADPH oxidase is associated mainly with production of superoxide by neutrophils.
during phagocytosis. In neurons NADPH oxidase positively correlate with processes of inflammation too. Inflammation-induced activation of microglial NADPH oxidase and iNOS has been reported to act synergistically to kill neurons through the formation of peroxynitrite. Peroxynitrite is a potent oxidant with biological reactivity similar to that of the hydroxyl radical. Expression of both enzymes is regulated by transcriptional factor NFκB [9, 86].

Activity of antioxidant enzymes was found controversial in patients with depression. In some studies increased activities were detected but, on the other hand, several studies published mixed or negative results of enzymes activity in depression in comparison to healthy control groups (Table 1, Figure 1) [38, 39, 41]. The contradiction can be caused by small study sample, heterogeneity of patients’ states, or variability in individual experiments [8].

Superoxide dismutase (SOD) with its cofactors (zinc and copper) catalyses break down from superoxide to oxygen and paradoxically to hydrogen peroxide which is at physiological conditions decomposed by catalase. An elevated level of SOD is observed as organism response to elevated concentrations of superoxide [7, 17]. In depressed patients increased and also decreased activity of SOD was shown [38–40]. Glutathione peroxidase (GPX) catalyzes reduction of peroxides, including hydrogen peroxide. GPX represents one of the main antioxidant enzymes in the organism [87]. In patients with affective disorder, depression and schizophrenia detected lowered activity of GPX in comparison to healthy control. Decreased activity causes accumulation of ROS and negatively correlates with severity of depression and its autonomous symptoms. GPX also protects the organism against cell death, DNA, and neurons damage [17, 40–42, 76]. Some studies on the other hand did not find significantly different level of GPX in depressed patients [38, 39]. Digestion of hydrogen peroxide to water and oxygen is catalyzed by enzyme catalase. Cell proliferation, apoptotic signals, and thrombocytes activation are influenced by catalase activity. Elevated activity of catalase was observed in patients in acute phase of depression and also in patients with bipolar disorder on lithium medication [39, 88, 89]. On the other hand, other works detected lowered activity of catalase in depression [43]. Oxidative stress and inflammation also affect activity of paraxenase 1 (PON1). PON1 is responsible for antioxidant properties of HDL and protects lipids against oxidative damage through lipoperoxidation. Together with HDL it possesses antiatherogenic properties [90]. Activity of PON1 is lowered in major depression and its activity also negatively correlates with smoking [44]. Other work did not find changes in PON1 activity in depressed women [40].

**5. Effect of Antioxidants, Micronutrients, and Vitamins in Depression**

Compounds capable of prevention or suppression of reaction between substrate and FR or ROS and RNS are defined as antioxidants. Antioxidants are able to be effective at very low concentrations and by ratio of 1:100 to amount of free radicals. Antioxidants in the organism are endogenous (e.g., glutathione, uric acid) and exogenous, like vitamin C, vitamin E, and polyphenols. An abnormal prefrontal level of GSH, the major cellular redox regulator and antioxidant in peripheral tissues and brains of patients with schizophrenia and bipolar disorder, was observed. Lowered concentration of antioxidants in the organism causes elevated production of ROS that leads to fatty acids membranes disruption and also to damage of proteins and DNA. Their concentration can be affected by diet (Table 1, Figure 1). Appraisal of antioxidant activity of natural substances is hindered by disagreement between results observed in vivo and in vitro [4, 7, 13, 61, 91].

Polyphenols are products of secondary metabolism of plants. They possess ability to absorb free radicals and to bond prooxidant ions of metals. They can increase activity of antioxidant enzyme or also increase expression of BDNF. There are observed anti-inflammatory effects of polyphenols. Increased cognitive abilities, mood improvement, and lowered probability of neurodegenerative disorders are accredited to diet rich in polyphenols. Among polyphenols are, for example, kurkumine, EGCG (epigallocatechin-3-gallat), resveratrol (presented in red wine, grapes, berries, and peanuts), chlorogenic acid (detected in apples, plums, and cherries), or EGB761 (detected in *Ginkgo biloba*) [6, 7, 13, 92, 93]. Kurkumine is contained in roots of *Curcuma longa*. It prevents lipid peroxidation and scavenges ROS and RNS. Apart from anti-inflammatory and antioxidant effect it is described also as having antiviral, antifungal, and antiamyloidogenic properties. It inhibits concentration decrease of 5-HT1A and BDNF; normalizes concentration of serotonin and dopamine, and inhibits activity of COX-1, COX-2, MAO-A, and MAO-B genes [7, 92, 94]. EGCG represents a main component of green tea (>20%). It possesses the ability of decreasing oxidative stress and TNFα production. Frequent green tea drinking can lower probability of depression occurrence and has potential to decrease symptoms of depression and stress and also increases cognitive abilities in consumers [7, 92]. Antidepressant-like effect was also confirmed in substance EGB761 found in *Ginkgo biloba* by affecting serotonin and dopamine concentrations [7]. Combined therapy with antidepressants and EGB extract increases sleep efficacy and reduces awakenings [95]. Polyphenolic extract from pine bark *Pinus pinaster*, Pycnogenol, improves symptoms of ADHD, normalizes level of neurotransmitters, and decreases oxidative stress markers [96–99].

Omega 3 polyunsaturated fatty acids (n-3 PUFA) can be found in sea food, fishes, and olive oil in high concentrations. Increased probability of depression and postpartum depression is associated with lowered food supply of n-3 PUFAs. In patients with depression, chronic fatigue and bipolar disorder are observed to have significantly decreased level of n-3 PUFA in comparison to healthy controls [61, 100–102]. Combined therapy of SSRI antidepressants with n-3 PUFA indicated to be more effective than antidepressants monotherapy [5, 103]. Consumption of at least one fish a week also decreases cognitive capability decline in old age approximately about 10–13% [104]. Other research, on the other hand, detected only small and nonsignificant effect of n-3 PUFA in depression [105, 106]. A number of accomplished suicides are also not significantly affected by n-3 PUFA consumption [107]. Among n-3 PUFAs are mainly eicosapentaenoic acid...
(EPA) and docosahexaenoic acid (DHA). DHA represents the main compound of phospholipid membrane in the brain. It inhibits apoptosis of neurons and ensures transfer of serotonin, norepinephrine, and dopamine. DHA also possesses antioxidant and antidepressant properties. EPA influences efficacy of immune system, decreases concentration of arachidonic acid (AA) in cell membranes, and lowers synthesis of prostaglandin E and cytokines like IL-1, IL-6, TNFα, and IFNγ. Consumption of EPA has significant antidepressive and anti-inflammatory effects. Omega 6 (n-6) PUFA, like AA, creates ROS by increasing proinflammatory signal molecules levels, for example, eikosanoids. n-6 PUFA promotes proinflammatory effects in the organism and increases production of IL-1, IL-6, and TNFα. It also directly induces brain tissue apoptosis [8, 18, 19, 108].

Coenzyme Q10 (CoQ10) represents an important antioxidant with anti-inflammatory and neuroprotective properties. It inhibits proinflammatory cytokines production and suppresses expression of NfκB. Lowered level of CoQ10 is associated with increased concentration of TNFα, ROS, and RNS and also mitochondrion dysfunction. Patients with drug resistant depression show lowered concentration of CoQ10 in comparison to healthy controls [5, 17, 45, 109, 110].

N-Acetyl cysteine (NAC) is an antioxidant that effectively reduces hydroxyl radicals. Intake of NAC increases concentration of cysteine in plasma and affects glutathione (GSH) levels. It also affects levels of glutamate and dopamine releasing. NAC has shown to possess anti-inflammatory properties and decreases levels of IL-1β, IL-6, and TNFα [7, 8]. Administration of NAC to patients with depression as part of bipolar disorder causes significant decrease of depression severity and improves antidepressants effectiveness [111, 112].

High density lipoprotein cholesterol (HDL) affects inflammatory and immune reaction of organism [113]. Lower level of HDL is associated with higher atherogenicity and risk for cardiovascular disease [90, 114]. Atherogenesis is frequently accompanied by depression in women [49]. Lowered serum level of HDL is detected also in depression, especially in patients with suicidal attempts [113] and with postpartum depression. There is also negative correlation between HDL concentration in serum and phobia, anxiety, and depression in women [115] and severity of postpartum depression [116]. On the other hand, no association between depression and level of HDL was found in elderly men [117]. Antidepressant treatment does not influence concentration of HDL in patients [113].

Depression is associated with deficiencies of some elements. For example, deficiency of magnesium causes mild inflammation. Its shortage also increases production of ROS and NO that can cause symptoms of depression especially in elderly people. In addition, intake of magnesium was associated with antidepressant-like effect in patients with depression [74, 118]. Besides magnesium deficiency, depression is usually accompanied also by iron shortage [41, 119]. Selenium is a main component of selenoproteins and amino acids, like selenomethionine and selenocysteine. Several selenoproteins, for example, GPX and thioredoxin reductase, possess antioxidant properties. They decrease oxidative damage of DNA and lower oxidative stress, increase expression of tumor suppressor p53, inactivate protein kinase C, and inhibit angiogenesis. Lowered level of selenium (62 μg/L) can be responsible for depressive mood, anxiety, and decline of cognitive function. Dietary intake of selenium can lower probability of postpartum depression and decrease depression severity. On the other hand, very high selenium status (110 μg/L) is also associated with depression [7, 50–53]. Deficiency of zinc is also associated with depression. It causes cognitive decay and humoral and cell immunity malfunction. Level of zinc also affects activity of antioxidant enzyme (Cu/Zn SOD) and enzymatic pathways of metalloenzymes dependent on zinc. Its concentration is important for homeostasis and signal transduction. It stimulates neurogenesis by increasing BDNF expression and by inhibition of glycohen synthase kinase 3 prevents neuroprogression. Deficiency of zinc increases oxidative stress and also concentrations of inflammatory cytokines. Zinc administration is efficient not only as prevention against symptoms of depression but also as effective antidepressant in postpartum depression and drug resistant depression. When combined with antidepressants usage, it increases efficacy of medication [5, 17, 54, 55, 120].

Deficiency of several nonenzymatic antioxidants is connected with worsening depression severity and anxiety. For example, significantly decreased level of vitamin E (α-tocoferol) was found in depressed patients in comparison to healthy controls [46]. Its concentration is negatively correlated with depression severity and also with levels of leukocytes and neutrophils. Antioxidant properties of vitamin E protect organism against lipid peroxidation. Diet rich in α-tocoferol also increases concentration of GSH and activity of glutathione reductase [7, 17, 121, 122]. In patients with depression lowered concentration of ascorbic acid (vitamin C) was also observed. Its intravenous administration not only amplifies antidepressants efficacy, but also act as antidepressant itself [7, 46–48]. Depression is associated with shortage of folate and vitamin B12 levels. Their deficiency affects metabolism of monoamines that can worsen depression severity in patients [123–126].

6. Effect of Mitochondrial Dysfunction on Depression

Production of ATP in mitochondrion ensures most of the energetic demand of the organism. This process is associated with excessive production of ROS and increasing of oxidative stress (Figure 1). Disrupted function of mitochondrion and an insufficient oxygen radical degradation increase concentration of ROS in the organism that consequently causes lipid, protein, and DNA damage. Depression is associated with decreased activity of enzymes involved in respiratory chain, decreased ATP production, and other mitochondrial dysfunctions caused by deletion in mitochondrial genome [5, 8, 61, 127, 128].

7. Correlation of Depression and Inflammation

Increased concentration of proinflammatory cytokines TNFα, IFN, and IL can lead to oxidative damage.
Overproduction of proinflammatory cytokines can be related to insufficient activity of antioxidant enzymes and low level of antioxidants. Overproduction can lead to pathological changes in brain that can escalate to cognitive dysfunction or neuropsychiatric disorders like schizophrenia, anxiety, depression, and depression states (lethargy, sleep disorders, weight loss, or anorexia). Levels of cytokines influence also neuroendocrine function and neurotransmitter metabolism. Imbalance of proinflammatory and anti-inflammatory cytokines can be a trigger of neurodegenerative disorders, for example, Alzheimer’s or Parkinson’s diseases. Inflammation processes in the organism are negatively correlated with concentration of antioxidants and antioxidant enzymes (Table 1, Figure 1) [5, 61, 129, 130]. In patients with depression and anxiety inflammatory reaction and mild chronic inflammation were observed to be caused by increased concentration of proinflammatory cytokines like TNFα, IL-1α, IL-1β, IL-4, IL-5, IL-6, IL-12, IFNγ, and C-reactive protein. Increased levels of inflammatory positive acute phase proteins, for example, haptoglobin, α1-antitrypsin, and ceruloplasmin, a α1-acid glycoprotein, were also detected. Antidepressants themselves possess anti-inflammatory effect by, for example, decreasing production of IFNγ and IL-12. In case of neurodegenerative disorders that are associated with chronic inflammation antidepressants act as preventives. In addition, inhibition of proinflammatory cytokines or their signal pathways increases effectiveness of antidepressants and decreases depression state in patients [4, 5, 7, 17, 41, 56–61, 127].

Depression is often accompanied by inflammation of digestive tract and increased number of gram-negative bacteria and Enterobacteria (e.g., Pseudomonas aeruginosa, Morganella morganii, Pseudomonas putida, and Citrobacter koseri). Their presence causes activation of immune response against lipopolysaccharides (LPS) of gram-negative bacteria especially by increasing immunoglobuline (Ig) A and IgM concentrations [5, 131, 132]. Immune reaction mediated by IgM towards phosphatidylinositol [133], NO-BSA [81], and NO-cysteiny l [134] is detected in serum of patients with chronic fatigue syndrome. IgM responses against oleic acid and phosphatidylinositol are also negatively correlated with severity of symptoms [134]. There also is recorded higher IgM response against NO modified amino acids, for example, NO-tyrosine, NO-arginine, and NO-tryptophan in serum of patients with chronic fatigue syndrome in comparison to healthy volunteers [79]. Major depression is also accompanied by elevated serum levels of IgM and autoimmune response against NO-tyrosine and oleic acid especially in chronically depressed patients. Antidepressants usage has no significant effect on IgM response [65].

8. Lipid Peroxidation in Depression

Increased lipid peroxidation is associated with depression. Lipid peroxidation is caused by increased concentration of proinflammatory cytokines that produce free radicals and increase metabolism of monoamines. Lipid peroxidation decreases life span of neurons, lowers expression of neurofilaments, reduces stability of membranes and activity of ion channels, affects neurotransmitter release, and by malondialdehyde (MDA) affects connection of 5-HT and membranes (Table 1, Figure 1). It belongs among important markers of cell damage caused by oxidative stress [7, 17, 135]. Decreased lipid peroxidation can be achieved by antidepressants and antioxidants (e.g., vitamin E) administration [39, 136].

Malondialdehyde (MDA) represents an end product of PUFA and AA peroxidation. MDA inhibits ligand-binding site of serotonin receptor and therefore affects metabolism of serotonin [5, 7, 17]. Increased concentration of MDA was detected in depressed patients in comparison to healthy control. Antidepressants administration, on the other hand, decreases MDA levels [39, 47, 66].

Product of AA peroxidation is 8-iso-prostaglandine F2α. 8-iso-prostaglandine F2α is important for stability and fluidity of cell membranes that influences consequential inflammatory response in the organism [17]. In patients with depression more than two times higher concentration of 8-iso-prostaglandine F2α are detected than in healthy controls. On the other hand, there does not seem to be a positive correlation between its concentration and depression severity [60, 67].

9. Conclusion

Major depression is associated with imbalance between several components, including neuroprogressive and neuroprotective factors (lowered concentration of BDNF [24], NFκB [38], and 5-HT [5]; elevated concentration of NO [27]), antioxidants and prooxidants (increased activity of XO [35] and catalase [39], decreased activity of GPX [40] and concentration of NAC [112] and HDL [113]), proinflammatory and anti-inflammatory molecules (increased levels of proinflammatory cytokines [57], autoimmune reactions against NO-modified proteins and decreased concentration of anti-inflammatory cytokines [65]), or production of reactive species and protective mechanism of the organism. Damaging changes can be observed in structure and utility of proteins (NO-modified proteins), lipids in cell membrane (increased levels of MDA [66] and 8-iso-prostaglandin F2α [67]), and nucleic acids (elevated levels of 8-OxoG [32]). Formation of new epitopes and IgM mediated immune response not only indicates nitrosatively and oxidatively damaged proteins and lipids but also triggers autoimmune and inflammatory response in pathophysiology of major depression [65, 82].

Besides affecting neurotransmitters concentration in brain by antidepressants it is possible to influence or even restore impaired balance by several other substances, including dietary intake (micronutrients, vitamins, omega-3 fatty acids, and antioxidants). On the other hand, effort to artificially decrease ROS and RNS, for example, usage of NOS inhibitor as a treatment drug for major depression, should be considered with caution [74].

Further research should provide answers to several unanswered questions. Effects of antidepressants on different aspect of oxidative and nitrosative damage in patients with psychiatric disorder is not fully understood. Is oxidative stress more harmful in early or chronic state of disease? Can
different markers of oxidative and nitrosative damage serve as valid predictor of psychiatric disorder? Can these markers distinguish between clinically different psychiatric disorders (e.g., major depression, anxiety, and schizophrenia)? Another problem is insecurity of results validation originating from blood and urine samples in comparison to values in brain or brain regions affected by disease [6]. Antioxidant effects of natural substances are also affected by discrepancies between results in vivo and in vitro [137]. Not only psychiatric disorders, but also several other diseases, for example, cardiovascular disorders, share common pathophysiological features with major depression [138]. Knowledge of connection between oxidative stress and depression is undoubtedly a step forward to better comprehension of pathophysiology of depression.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

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