Oxidative Stress in Bipolar Disorder

Victor Tang1 and Jun-Feng Wang1,2*  
1Department of Psychiatry, University of British Columbia, Vancouver, BC, Canada  
2Department of Pharmacology and Therapeutics, University of Manitoba, Winnipeg, MB, Canada

Abstract

Research on the complex and multifaceted nature of Bipolar Disorder (BD) pathophysiology has recently expanded to include oxidative stress. Several lines of evidence have reported higher reactive oxygen species production and results in increased oxidative damage in proteins, lipids, and nucleic acids. These findings have been observed in brains and peripheral samples of BD patients, as well as being reproduced in a number of animal model studies. Also discussed in this review is research highlighting antioxidant properties of existing mood stabilizing drugs, with consideration paid to novel therapeutic treatments for BD through the alleviation of oxidative stress. The maladaptive oxidative modifications of cellular macromolecules may be associated with impaired neuroplasticity and the development of functional abnormalities in the brain.

Keywords: Bipolar disorder; Lipid peroxidation; Protein oxidation; Antioxidant; Mood stabilizing drugs

Introduction

Bipolar Disorder (BD) is a severe psychiatric illness characterized by cycling episodes of mania and depression. Estimates frame the prevalence of BD as around 1-3% of the worldwide population. Mania is a state of excessive energy, associated with euphoria, impulsivity, lack of need for sleep, and psychosis. In contrast, depressive episodes are characterized by severely disoriented and non-reactive mood, anhedonia, fatigue, psychomotor retardation, disruption of sleep and eating routines, impaired concentration, and feelings of guilt and worthlessness. BD is a leading cause of disability among individuals with medical and psychiatric conditions, and is associated with substantial morbidity and mortality. Ramifications of this chronic disease include dependence on social welfare, criminal justice intervention, and serious economic strain to patients, their families, and the healthcare system. Moreover, the healthcare burden of this disorder is exacerbated by an increased risk of other medical conditions such as cardiovascular disease, obesity, and infectious diseases.

Advances in research on BD neurobiology have continued to elucidate the pathophysiology, genetic predispositions, and biochemical abnormalities of this disease. Abnormal neuroplasticity has recently been recognized in BD. Neuroplasticity is the ability to undergo and sustain change in the brain in order to maintain proper functioning. Neuroplasticity involves processes such as long-lasting modification of synapses and gene expression, neuronal growth and morphology, and the broader development of neural circuits in the brain. Dysregulation of intracellular cascades and associated gene regulation central to neuroplastic processes has been implicated in BD [1]. Evidence also suggests that monoaminergic, glutamatergic and neurotrophic signaling cascades are impaired [2]. Proteins involved in neuroplasticity, such as the Brain-Derived Neurotrophic Factor (BDNF) in synapse formation, differentiation, and neuronal survival, are abnormally expressed in BD [3]. Critical circuitry systems that regulate emotion, motivation, and memory are disturbed in BD, including the Prefrontal Cortex (PFC), cingulate, hippocampus and amygdala [4]. Taken together, the dysfunction found in neuroplasticity is likely to underlie deleterious effects on cell number, density, and morphology. Indeed, research has outlined significant structural abnormalities in BD, which may be related. Cellular abnormalities are observed in postmortem brain studies showing decreased pyramidal cells in the prefrontal cortex [5], while decreased non-pyramidal cell density was found in the anterior cingulate cortex [6]. Structurally, major findings from Magnetic Resonance Imaging (MRI) studies in BD patients have shown increased ventricular size, suggesting brain tissue volume decreases, and grey matter reductions in PFC, which plays a critical role in the control and direction of mood, cognition, and motor behavior [7].

Recently, oxidative stress has been suggested to play an important role in the pathophysiology of BD. Oxidative stress is defined as when the pro-oxidant levels in cells overwhelm the antioxidant capacity. Oxidative stress can occur through overproduction of free radicals or decrease in antioxidant defense systems or through both at once. These free radicals can cause substantial damage to macromolecules through the generation of adducts, destruction of unsaturated C-C bonds, and oxidation of disulfides [8]. More generally, they cause oxidative cell injury through damaging effects to proteins, lipids, and nucleic acids. However, when kept in controlled conditions by antioxidant systems; free radicals can serve important functions in physiological processes [9]. They are continuously produced in vivo by all tissues of the body, primarily during oxidative phosphorylation in the mitochondria. Oxygen-containing free radicals, or Reactive Oxygen Species (ROS), are typically produced as a byproduct of cellular respiration in the Electron Transport Chain (ETC). When electrons are prematurely released from complexes in the transport chain, they reduce molecular oxygen to superoxide radical, which subsequently produces various ROS through a series of biochemical cascades. A series of natural antioxidant defense systems normally eliminate ROS production and limit tissue damage: superoxide radical is dismutated by Superoxide Dismutase (SOD) to H$_2$O$_2$, which is further converted to H$_2$O by Catalase, Glutathione (GSH) plays a key role as a major antioxidant in supporting cellular redox balance and protecting brain cells against free radicals.

*Corresponding author: Jun-Feng Wang, Department of Pharmacology & Therapeutics, University of Manitoba, 406 - 753 McDermot Ave., Winnipeg, Manitoba R3E OE3, Canada. Tel. 204-975-7705; E-mail: wangj323@cc.umanitoba.ca

Received April 16, 2012; Accepted May 17, 2012; Published May 21, 2012

Citation: Tang V, Wang JF (2012) Oxidative Stress in Bipolar Disorder. Biochem Anal Biochem S2-002. doi:10.4172/2161-1009.S2-002

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oxidative damage by ROS. GSH is maintained by glutathione reductase that catalyzes the reduction of glutathione disulfide (GSSG) to GSH. Glutathione peroxidase (GSH-Px) catalyzes GSH reaction with $\text{H}_2\text{O}_2$ and converts to $\text{H}_2\text{O}$, and Glutathione S-transferase (GST) catalyzes the conjugation of oxidized products with GSH to form a non-toxic product.

Because the brain is only 2% of total body weight while metabolizing 20% of total body oxygen, it is particularly vulnerable to the production of ROS. As rising $\text{O}_2$ concentration is associated with an increased rate of electron leakage [10], the large amount of oxygen consumed by the brain may result in high superoxide formation. Furthermore, it has been widely argued that the brain is especially prone to oxidative damage, as attributable by the brain being(1) rich in peroxidizable fatty acids, (2) poor in catalase activity and having only moderate amounts of SOD and GSH-Px, and (3) rich in iron in several areas, which facilitates the production of highly reactive hydroxyl radicals [9,11]. In BD, accumulating evidence from a range of studies using post-mortem brain tissue and peripheral blood samples has demonstrated increased oxidative damage to cellular macromolecules. These studies have also demonstrated a decrease in antioxidant capacity. Relevant to current and future development of therapeutics, there have been numerous reports of mood-stabilizing drugs producing neuroprotective effects against oxidative damage and increased antioxidant expression and activity.

**Oxidative Stress in BD**

Direct evidence of increased oxidative stress has come from studies utilizing post-mortem brain tissue to measure markers of oxidative damage to cellular macromolecules. In 2009, Wang et al. [12] reported that 4-hydroxynonenal (4-HNE) protein adducts are significantly increased in postmortem Anterior Cingulate Cortex (ACC) of subjects diagnosed with either BD or schizophrenia when compared to non-psychiatric populations. These adducts are known to be a major product of lipid peroxidation, and are able to induce cytotoxic effects and disturb cellular function when they accumulate in excess. Therefore this finding suggests that oxidative damage to lipids and proteins is associated with BD and schizophrenia. It is very likely that ACC pathology is involved in BD symptomatology, as this structure has a vast array of connections to other frontal and limbic areas and contributes to processing of cognitive, emotional, and attention functions [13]. Interestingly, subjects without medication had higher levels of 4-HNE protein adducts compared to the total BD patient group, which may indicate an underlying marker of pathology in BD [12], although this remains to be validated in future studies with larger sample sizes. The authors also report a negative correlation between 4-HNE and pH, which may be indicative of a relationship between oxidative stress and mitochondrial dysfunction. More specifically, the mitochondrial dysfunction may lead to both an increase in ROS production and a shift of energy production from oxidative phosphorylation in the ETC towards glycolysis and lactic acid production.

It has also been found that levels of protein carbonylation and 3-nitrotyrosine (3-NT) are increased in prefrontal cortex tissue of BD subjects [14]. Protein carbonylation is an oxidative process where ROS react to form carbonyl groups into the side chains of amino acids in proteins. 3-NT is another marker for protein oxidation, and is a result of tyrosine residues being nitrated by the Reactive Nitrogen Species (RNS) peroxynitrite [9]. Interestingly, another study with hippocampal tissue of BD patients showed increases in neuronal nitric oxide synthase I, the enzyme that generates nitric oxide, the precursor to peroxynitrite [15]. Oxidation of proteins can lead to damaging effects through the formation of deleterious intermolecular aggregates and the alteration of protein function. It has also been reported that carbonylation and 3-NT levels were correlated with decreases in ETC complex I activity, suggesting that increases in oxidative dysfunction may be coming from mitochondrial dysfunction, specifically excessive leakage of electrons at complex I [14]. Indeed, protein oxidation from peroxynitrite can result in inhibition of complex I activity, as a study by Naoi et al. [16] demonstrated increases in 3-NT levels of complex I subunits compared to other ETC complexes in SH-SY5Y neuroblastoma cells incubated with peroxynitrite. Taken together, this evidence shows that oxidative damage to proteins is elevated in BD, and that it is likely to be interrelated with mitochondrial dysfunction.

Increased oxidative damage to nucleic acids has recently been observed in a post-mortem brain study of BD patients [17]. In the CA1, CA3, and dentate gyrus regions of the hippocampus, oxidative damage to nucleic acids was greater in BD subjects compared to controls. The hippocampus is best known for involvement in memory formation, but it is also connected with other limbic structures in order to regulate emotion. Moreover, hippocampus has been shown to be susceptible to damage from stress effects, as well as having volumetric abnormalities in BD patients [18]. RNA and DNA can be oxidized by ROS, with guanine residues most sensitive to attack compared to other bases. When these residues react with ROS, 8-oxo,7,8-dihydro guanosine (8-OHG) is produced in RNA and 8-oxo,7,8-dihydro-2-deoxy guanosine (8-OHdG) in DNA. In this study, oxidative damage was quantified through performance of immunohistochemistry with a monoclonal antibody recognizing both 8-OHG and 8-OHdG, and increases were found in BD subjects, along with major depression and schizophrenia subjects, when compared to controls. The study authors discuss the possibility of increased susceptibility to oxidative damage to RNA compared to DNA because of the fact that RNA are single stranded and without the protection from hydrogen bonds found in DNA, which is double stranded. Oxidatively modified RNA may cause downstream effects through errors in translation and altered or reduced protein expression. DNA fragmentation was also found to be increased in non-GABAergic neurons in BD [19], and it is suggested that this may confer vulnerability to oxidative stress and excytotoxicity [20]. Although these studies utilizing postmortem brain tissue from patients cannot rule out the effects of medication, substantial evidence from *in vivo* and *in vitro* studies suggests a neuroprotective role against oxidative stress in the major pharmacological treatments of BD. Thus it appears more likely that the significant differences found are due to an underlying pathology rather than a consequence of medication.

Deficiencies in brain antioxidants may also make significant contributions to increases in oxidative stress. Glutathione (GSH), the major free radical scavenger in the brain, was found to be lower in prefrontal regions of patients with BD, major depressive disorder and schizophrenia [21]. It has also been reported that gene expression of antioxidant enzymes such as SOD1, catalase, GSH-Px4, and GST is down-regulated in the hippocampus [22]. Fullerton and colleagues [23] also found that genes for SOD2 and GSH-Px3 are possible risk factors for BD. By targeting candidate genes in the oxidative stress pathway, the authors found that haplotypes within the SOD2 gene are associated with BD and interaction between SOD2 and GSH-Px3 at the haplotype and genotype level further increased this risk. Impairments in antioxidant defense systems in BD may indicate a decreased ability to neutralize ROS accumulation and infer susceptibility to oxidative stress.
Increases in peripheral biomarkers of oxidative stress have also been consistently reported in the literature. These were in biological samples of plasma, serum, white blood cells, red blood cells, or platelets from patients with BD. Studies have indicated significant increases in lipid peroxidation [24-26] and nitric oxide, a highly reactive oxidant molecule [24,27-29], along with DNA damage associated with oxidative stress [30], increased protein carbonyl content [31], and several antioxidant enzymes [25,27,32]. Some discrepancies in the literature exist surrounding SOD antioxidant levels being increased or decreased, which may reflect fluctuations in response to state dependent changes in oxidant levels [24,27,33]. Number of manic episodes was found to be correlated to serum nitric oxide levels [24], indicating that oxidative stress may be a trait marker of the disease, and that the chronic nature of the illness could lead to long term accumulation of oxidative stress, resulting in the poorer outcomes associated with length of disease and number of mood episodes [34]. Moreover, Andreazza et al. [30] found that increased DNA damage indicative of oxidative stress was associated with severity of mood symptoms. Importantly, increases in protein carbonylation were detectable in early stages of the illness, as reported in a recent population based study of young adults [35]. Research has also suggested that there may be particular oxidative imbalances present during active phases of BD, as Andreazza et al. [25] found increases in SOD activity only in patients during depressed or manic states, but not in euthymic episodes.

It has also been reported thatmania is associated with higher lipid peroxidation levels than any other phase of the illness [36]. In a recent study by Kapczinski et al. [31], such state dependent differences were examined in greater detail. Here they observed that manic and depressive episodes were distinguishable from control conditions by increased protein carbonylation and lipid peroxidation. 3-NT was elevated compared to controls only in depression [31]. Interestingly, a study by Yumru et al. [37] demonstrated that analyses of various markers of oxidative imbalance in BD patients had some ability to distinguish between bipolar subtype diagnoses. While peripheral samples suggest oxidative stress in the pathophysiology of BD, it is still unclear how blood biomarkers can specifically describe the state of oxidative stress in the brain. Thus, interpretation of these findings remains limited. Nonetheless, these studies have provided a valuable contribution by studying the relationship between oxidative stress and more particular and dynamic clinical factors such as illness duration, type of mood episode, and effects of medication. All together, amongst a wide range of patient populations, using different biological samples and experimental measures, these studies have generated a large body of literature to implicating oxidative stress in the pathophysiology of BD (Table 1).

Animal studies also provide evidence of oxidative stress in BD, although the interpretations are limited given the lack of a comprehensive model of the disease. Amphetamine induced hyperactivity is the most widely used animal model of mania, and as such is commonly used to study the manic component of BD. Studies have reported that repeated amphetamine administration was associated with increases in superoxide production [38], and protein and lipid oxidative damage in rat brain [39], with longer periods of exposure being positively correlated with oxidative stress. Another study by the same group found that the same model resulted in SOD and catalase activity decreases [40]. It has also been found that amphetamine treatment can induce increased peripheral and hippocampal DNA damage that is also correlated with increases in lipid peroxidation [41]. Tan et al. [42] also found that chronic amphetamine treatment increased 4-HNE protein adduction, particularly 4-HNE adduction with vesicular monoamine transporter 2 (VMAT2) in frontal cortex of rat brain. Monoamine neurotransmitters are involved in the etiology and pathology of bipolar disorder and other psychiatric diseases, and also contribute significantly to amphetamine-induced behavioral effects. Vesicular monoamine transporter 2 (VMAT2) is critical in packaging monoamine neurotransmitters. Similar results were found in an animal model of mania induced by ouabain, a Na+/K+-ATPase inhibitor, which mimics the hyperactivity and the sodium pump hypoaactivity found in BD [43]. Specifically, this model can demonstrate increases in TBARS, superoxide production, and protein carbonyl content [44-46]. Evidence for oxidative stress also exists in animal models of depression, although these usually take place in the context of studying major depressive disorder. Nonetheless, phenotypic expression of unipolar and bipolar depression is very similar, and animal models of depression are not distinguished for any particular type of depression. Lucca and colleagues [47] used a chronic mild stress model of depression on rats and found increases in superoxide generation in the hippocampus and TBARS in the cortex. Using the same model, Eren et al. [48] reported a decrease in antioxidants GSH-Px, glutathione, and vitamin C in rat cortex, and Eijchel-Cohen et al. [49] reported decreases in the antioxidant GST gene expression. Chronic stress models typically produce depressive symptoms in rats, such as anhedonia, weight loss, and decreased locomotors activity. Other studies using prolonged or chronic stress in animals have reported various increases in ROS and oxidative damage [50-52].

**Treatment of Mood Stabilizing Drugs**

Mood stabilizing drugs are medications for the treatment of BD and are useful for the treatment of acute manic episodes and for prophylaxis against future episodes. These drugs include lithium, often considered the “gold standard” mood stabilizing drug, anticonvulsant medications such as valproate, carbamazepine, lamotrigine, and atypical antipsychotics such as olanzapine. Despite the extended history of use in patients, as well as many years of research, the mechanism of action of these treatments is still unclear. A growing body of evidence shows that chronic treatment with mood stabilizing drugs is neuroprotective against oxidative stress both *in vitro* and *in vivo* [53] and this neuroprotective mechanism is currently undergoing extensive study. It has been reported that chronic treatment with the mood stabilizing drug lithium or valproate significantly inhibited oxidative damage to lipids and proteins induced by various insults in primary cultured rat cerebral cortical cells [54,55]. Treatment with lithium or valproate werealso shown to inhibit H$_2$O$_2$-induced and complex I inhibitor rotenone-induced cytochrome c release, caspase-3 activation, and cell death in human neuroblastoma cells and in murine hippocampal cells [56,57].

To further elucidate the mechanisms of action for these drugs, Cui et al. [57] showed that these drugs increase the levels of the antioxidant GSH and the expression of glutamate-cysteine ligase, the rate-limiting enzyme in GSH synthesis. Similar results were demonstrated with the mood stabilizing drugs lamotrigine and carbamazepine [57]. Lithium and valproate were also found to bolster antioxidant defenses through the regulation of GST. It was found that lithium increased mRNA expression of the M1,M3, M5, and A4 isoenzymes, while valproate increased levels of M1 and A4 but decreased A3 [58,59]. These findings were complimented by other reports showing lithium treatment in rats to increase brain SOD and GSH-Px activity [60] and also in an aggregate measure of total antioxidant activity [61]. Thus, current mood stabilizing treatments may exert their therapeutic effects at least in part by combating oxidative stress and strengthening antioxidant defense mechanisms.
Further evidence is provided by animal studies in vivo. Frey et al. [62] reported that chronic treatment with lithium and valproate inhibits amphetamine-induced hyperactivity and lipid peroxidation in rat brain. Inhibition of both hyperactive mania-like behaviour and oxidative stress suggests that this anti-oxidative stress effect may play an important role in the therapeutic effect of lithium and valproate. Similarly, lithium and valproate were able to modulate oxidative balance and prevent associated DNA damage in this animal model [41]. A study by Jornada et al. [44] showed that lithium and valproate reversed and prevented oxidative damage and production of superoxide in an animal model of mania using ouabain, as well as increasing catalase activity. These findings have therapeutic relevance because these drugs can also reverse the ouabain induced hyperactive behavior in an open field test [63]. It has also been found that when aluminum administration in rats causes ROS production and increased lipid peroxidation, these effects were decreased when followed by lithium supplementation [64]. The same study reported that changes in antioxidant enzymes by aluminum were normalized by lithium treatment. Most recently, chronic lithium treatment has been found to inhibit 4-HNE-protein adduction [42]. Repeated amphetamine stimulation significantly increased 4-HNE-VMAT2 adducts, while chronic lithium treatment reduced amphetamine-increased 4-HNE-VMAT2 adducts in rat frontal cortex. Taken together, these findings suggest that prevention of oxidative stress may contribute at least in part to the pharmacological action of mood stabilizing drugs for the treatment of BD.

Mood stabilizing therapy has also shown to decrease oxidative

| Reference                  | Patients/Controls | Sample            | Results                                      | Medication, drugs/alcohol                  | Mood Status       |
|----------------------------|-------------------|-------------------|----------------------------------------------|--------------------------------------------|-------------------|
| Oliveira et al. [15]       | 15/15             | Postmortem brain - hippocampus | ↑nNOS                                        | Lithium subjects higher in CA3 region than non-lithium, no effect from other medications | N/A               |
| Wang et al. [12]           | 15/15             | Postmortem brain - hippocampus | ↑nNOS                                        | Lithium subjects higher in CA3 region than non-lithium, no effect from other medications | N/A               |
| Andreazza et al. [14]      | 15/15             | Postmortem brain - PFC | ↑PCC, 3-NT                                   | No effect of medication on measures         | N/A               |
| Che et al. [17]            | 15/15             | Postmortem brain - hippocampus | ↑RNA oxidative damage                        | N/A                                        | N/A               |
| Gawryluk et al. [21]       | 15/15             | Postmortem brain - PFC | ↓GSH                                         | No effect from drug/alcohol abuse          | N/A               |
| Abdalla et al. [33]        | 20/58             | RBC               | ↓SOD                                         | No difference between lithium and lithium+antipsychotic | N/A               |
| Savas et al. [24]          | 44/21             | Plasma            | ↓NO                                         | N/A                                        | N/A               |
| Rajekar et al. [32]        | 10/31             | RBC               | ↓SOD, CAT, ↑TBARS                            | N/A                                        | N/A               |
| Ozcan et al. [66]          | 30/21             | RBC               | ↓CAT, GSH-Px, NO ↑MDA                        | After 3 months psychiatric inpatient clinic, GSH-Px increased | N/A               |
| Yanik et al. [28]          | 43/31             | Plasma            | ↓NO                                         | N/A                                        | N/A               |
| Savas et al. [24]          | 27/20             | Serum             | ↑SOD, ↑NO, SOD                               | SOD negatively correlated with number of manic episodes, correlation between NO and delusions | N/A               |
| Andreazza et al. [25]      | 85/32             | Serum             | ↑TBARS, SOD, SOD:GSH-Px+Cat                  | SOD, SOD:GSH-Px+Cat effects in manic or depressive phase | N/A               |
| Andreazza et al. [30]      | 32/32             | Serum             | ↑DNA damage                                  | Depressive/manic states correlate with damage | N/A               |
| Machado-Vieira et al. [65] | 30/30             | Serum             | ↑NO, SOD                                     | Currently euthymic. Manic episodes correlated with increasing NO | N/A               |
| Gergerlioglu et al. [29]   | 29/30             | Serum             | ↑NO, ↑SOD                                    | After 30 days of psychiatric hospital treatment, SOD decreased, no difference in NO | SOD negatively correlated with number of manic episodes, correlation between NO and delusions. |
| Kunz et al. [36]           | 83/32             | Serum             | ↑SOD, TBARS                                  | SOD with mania and depression, not euthymic. TBARS with all phases. | N/A               |
| Selek et al. [27]          | 30/30             | Serum             | ↑SOD, ↑NO                                    | Degree of normalization after 30 days hospitalization with drug treatments | No correlation with number of any type of episode |
| Yumru et al. [37]          | 94/41             | Plasma            | ↑Total oxidant, Total antioxidant, Oxidative stress (oxidant/antioxidant) | N/A                                        | N/A               |
| Kapczinski et al. [83]     | 80/60             | Serum             | ↑PCC, TBARS                                  | Depression and mania associated with PCC, TBARS, not euthymia. | N/A               |
| Magalhães et al. [71]      | 55/94             | Serum             | ↑PCC                                         | Current use of medication not correlated | N/A               |
| Banerjee et al. [26]       | 73/35             | Serum             | ↑TBARS                                       | TBARS lower in lithium treated vs. lithium naive | N/A               |

**Table 1:** Studies of oxidative stress in Bipolar Disorder patients. If available, findings of associated variables of mood stabilizing medications, alcohol and drugs of abuse, and different mood episodes with oxidative stress measures are presented. Brain samples: Anterior cingulate cortex (ACC), Prefrontal cortex (PFC). Peripheral samples: Red blood cells (RBC), Reactive Oxygen Species: Nitric Oxide (NO), Neuronal nitric oxide synthase (nNOS). Oxidative damage to proteins: Protein carbonyl content (PCC), 3-Nitrotyrosine (3-NT). Oxidative damage to lipids: Thiobarbituric acid reactive substances (TBARS), 4-hydroxynonenal (4-HNE). Antioxidants: Catalase (CAT), Glutathione (GSH), Glutathione peroxidase (GSH-Px), Superoxide dismutase (SOD).
stress markers in peripheral samples of human patients. Machado-Vieira et al. [65] reported that acute treatment with lithium was able to normalize SOD/CAT ratios and reduce lipid peroxidation levels. A longer study of treatment effects was conducted on 30 BD patients admitted to hospital, and showed that there was reduction of nitric oxide levels and increases in SOD activity between the 1st and 30th day of admission [27]. These results point to the possibility of oxidative stress alleviation with mood stabilizing treatments, although the study did note that SOD levels were not able to fully match control subject levels after 30 days, indicating a trait impairment in antioxidant capacity and possible predisposition to oxidative stress in BD. A similar study by Ozcan et al. [66] was conducted over an even longer period of 3 months, but with much smaller sample size, and observed a higher amount of GSH-Px activity in the post-treatment group compared to pre-treatment and control groups. Moreover, lithium alone or in combination with olanzapine appears to relieve lipid peroxidation and increase total antioxidant status [67]. In a study of healthy volunteers given therapeutic doses of lithium for 2–4 weeks, results showed decreases in hydrogen peroxide levels [68]. These studies are useful in providing evidence for the therapeutic effect of antioxidant properties in current mood stabilizing treatments. The sum of the evidence available shows that current mood stabilizing drug treatments have alleviation of oxidative stress has been supported by some human studies. In a double-blind randomized placebo-controlled trial, Berk et al. [69] used treatment with N-Acetyl Cysteine (NAC) in BD patients in their maintenance phase. NAC is an acetylated derivative of cysteine, which is the rate-limiting precursor in GSH production [70]. This study showed that NAC caused a significant improvement in assessments suggesting a neuroprotective effect against oxidative stress, and can modulate a variety of antioxidant pathways in the cell (Figure 1).

The possibility of novel drug treatments for BD through the alleviation of oxidative stress has been supported by some human studies. In a double-blind randomized placebo-controlled trial, Berk et al. [69] used treatment with N-Acetyl Cysteine (NAC) in BD patients in their maintenance phase. NAC is an acetylated derivative of cysteine, which is the rate-limiting precursor in GSH production [70]. This study showed that NAC caused a significant improvement in assessments suggesting a neuroprotective effect against oxidative stress, and can modulate a variety of antioxidant pathways in the cell (Figure 1).

**Figure 1:** Antioxidant mechanisms of mood stabilizers. Arrows represent increases or decreases in expression or activity due to stated mood-stabilizing drugs, numbers as reference citations. Superoxide radical (O$_2^-$), hydrogen peroxide (H$_2$O$_2$), hydroxyl radical (OH$^-$), nitric oxide (NO$^+$), peroxynitrite (ONOO$^-$), superoxide dismutase (SOD), catalase (Cat), glutathione (GSH), glutathione peroxide (GSH-Px), glutathione disulfide (GSSG), glutamyl-cysteine ligase (GCL), glutathione-S-transferase (GST). Mood stabilizers: Lithium (Li$^+$), valproate (VPA), olanzapine (OLZ), lamotrigine (LTG), Individualized psychiatric treatment in humans (Tx).

While the evidence for oxidative stress in BD continues to build, the implications of these findings to the overall pathophysiology of the illness still remain unclear. The significance of oxidative stress as a component of many disease processes in the central nervous system is increasingly recognized. Oxidative stress in the brain is seen in neurodegenerative disorders, Parkinson’s disease and Alzheimer’s disease. Many neurodegenerative disorders are strongly associated with increasing age [75], which attributes the precipitation of disease to gradual, long term accumulation of oxidative damage, finally causing neuronal loss in specific brain regions. However, onset of BD typically occurs in late adolescence or early adulthood [76]. BD is not as progressive as neurodegenerative disorders. Although less evidence suggests neuronal loss in BD, milder oxidative stress in this disorder may further modify various molecules such as synaptic proteins, enzymes and components of intracellular signaling. Oxidative modifications of these molecules that have been found in BD could affect neurotransmission and impair neuroplasticity in brain regions related to mood regulation.

Research on the neurobiology of BD has shifted focus towards more enduring cellular and molecular changes in BD. These changes in cellular structure, function, growth and development are...
characterized as mechanisms of neuroplasticity, and studies have accumulated to implicate dysfunction in these processes in BD [77]. Indeed, a recent genome-wide association study of BD revealed that most of the highly significant associations were implicated in signaling cascades of plasticity [78]. It has been proposed that abnormalities in neuroplasticity lead to maladaptive developments in neural circuits, affecting the information processing that mediates various facets of BD symptomatology [77,79]. The BDNF polymorphism gene (Val66Met) has been proposed to be associated with susceptibility to BD [80] and is said to alter the intracellular trafficking and activity-dependent secretion of BDNF [81]. This seems to be in line with other studies reporting BDNF decreases in serum of BD patients during manic and depressive phases [82]. Interestingly, Kapczinski et al. [83] showed that serum TBARS was negatively correlated with BDNF levels. Another study using peripheral markers showed that BD patients showed increased oxidative stress and decreased neuron-specific endolase, a neuronal glycolytic enzyme known to mediate neuroplastic pathways and cell survival [65], although the relationship between these variables were not analyzed. An animal study by Wu et al. [84] showed that oxidative stress is correlated with BDNF reductions, as well as reductions in cyclic AMP responsive element binding protein and synapsin I molecules that are involved in cellular plasticity cascades. Such research suggests that oxidative stress may play a role in the abnormal neuroplastic processes found in BD pathophysiology.

Increases in levels of ROS are likely due in large part to mitochondrial dysfunction. Imaging studies through magnetic resonance spectroscopy have demonstrated that the energy metabolism within the brain appears to be abnormal in BD patients, inferring mitochondrial dysfunction [85-87]. Concurrent with these studies has been genetic research highlighting increases in mitochondrial DNA deletions and mutations and altered expression of mitochondrial proteins [88,89]. While mitochondrial dysfunction has a myriad of maladaptive effects on the cell, of particular interest are findings of defects in the ETC, which is the primary site of electron leakage that occurs during oxidative phosphorylation. Along with DNA microarray analyses that have identified decreased expression of mRNAs that code for ETC complexes [90,91], postmortem brain tissue of BD patients also shows decreased levels of complex I subunit expression and impaired functionality [14]. Aberrant functioning in mitochondria is observed to increase production of ROS, and it is well understood that electrons are provided to form these free radicals when they are leaked from the ETC [92,93]. Mounting evidence suggests that mitochondria are directly involved in processes of synaptic plasticity. Mitochondria are enriched in presynaptic terminals, and play important roles in neurotransmission via energy production. In addition, mitochondria buffer the levels of calcium [94]. Calcium influx precedes neurotransmitter release and is necessary for vesicle docking and exocytosis. Mitochondrial dysfunction may be responsible for more pronounced calcium spikes and abnormal neurotransmitter release [95]. Deficiencies in ATP production, seen in BD, can impair the energy demanding processes of vesicle docking, fusion, and endocytosis.

Conclusions

As the neurobiology of BD continues to be elucidated, it has become clearer that oxidative stress likely plays an important role in the development of the illness. Studies of postmortem brain tissue have found evidence of oxidative damage to proteins, nucleic acids, and lipids, as well as decreased antioxidants, suggesting the presence of oxidative stress as a major component of BD pathophysiology. Other lines of evidence from animal model studies and peripheral blood studies of BD patients also converge to support this notion. Damage to the biochemical properties of cellular macromolecules is likely to underlie functional abnormalities in the brain, particularly the impairments in neuroplasticity that have been highlighted in the recent literature. Studies of pharmacological mechanisms of mood stabilizing treatment have also provided a large body of research supporting the therapeutic benefits of neuroprotection against oxidative stress. These findings contribute to our understanding of this complex and multidimensional disorder, and may reveal putative targets for novel drug development. Such efforts provide hope for discovering better treatments and outcomes for those with BD.

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