Review Article

Lipid Peroxidation in Psychiatric Illness: Overview of Clinical Evidence

Yash B. Joshi and Domenico Praticò

Department of Pharmacology, Center for Translational Medicine, Temple University School of Medicine, MERB 947, 3500 North Broad Street, Philadelphia, PA 19140, USA

Correspondence should be addressed to Domenico Praticò; praticod@temple.edu

Received 18 March 2014; Accepted 8 April 2014; Published 27 April 2014

1. Introduction

Over the last century, few areas of clinical and biomedical inquiry have undergone as rapid a transformation as the neuroscience of psychiatric disorders. In the past several decades, increasingly sophisticated experimentation, investigational tools, and model systems have led to more nuanced approach to pharmacotherapy. Although different psychiatric disorders are currently thought to stem from unique abnormalities in neuronal biochemistry, circuitry, and/or brain architecture, emerging data indicates that oxidative stress is present and may play an active role in these psychiatric illnesses. In this paper, we will provide an appraisal of recent clinical findings on lipid peroxidation as it applies to the pathobiology of schizophrenia, bipolar disorder, major depressive disorder, and several other psychiatric disorders.

2. Lipid Peroxidation and the Central Nervous System

The degree of oxidative stress in the cellular milieu is a direct result of those processes that accelerate the production of reactive oxygen species and those that detoxify them (for a discussion on oxidative stress, see [1]). Reactive oxygen species such as superoxide anion and hydroxyl radicals are produced from a variety of cellular processes. These reactive oxygen species are neutralized by antioxidants such as vitamins C and E as well as enzymes such as superoxide dismutase (SOD), catalase, and glutathione peroxidase. Low levels of reactive oxygen species are utilized by redox sensors to modulate cell function, but high levels of reactive oxygen species damage and oxidize nucleic acids, carbohydrates, and lipids. The brain is approximately 2% of total body weight but utilizes 20% of total oxygen, allowing ample metabolic substrates for free radical generation [2]. In light of this, and given the fact that lipids are the major components of neuronal membranes as well as the myelin sheaths that help conduct neuronal signaling, their peroxidation and dysfunction can dramatically compromise brain function globally.

Thus far, brain lipid peroxidation is not able to be directly assessed in living subjects. In lieu of this, markers of global oxidative stress can approximate lipid peroxidation and include antioxidant status and the activity of antioxidant enzymes such as superoxide dismutase, catalase, and glutathione peroxidase peripherally [3]. More direct measures of lipid peroxidation in vivo include assessment of isoprostanes and malondialdehyde/thiobarbituric acid reactive species in a variety of biological fluids or alkanes such as pentane in
exhaled air [3, 4]. Although circuit level dysfunction and the dynamic pathophysiology of psychiatric disorders are poorly approximated by postmortem investigation of the brain, such tissue analysis can aid in localization of lipid peroxidation in regions known to be important to psychiatric disease.

3. Schizophrenia

Schizophrenia is characterized by the presence of positive symptoms such as delusions, hallucinations, disorganized thinking, and negative symptoms such as flat affect. In part due to the fact that the success of treating the positive symptoms of schizophrenia with dopamine receptor antagonists and that intoxication with substances that increase brain levels of dopamine such as amphetamine or cocaine can lead to episodes of psychosis, inappropriate activation of dopaminergic pathways was initially thought to underlie the disease. However, this dopamine hypothesis of schizophrenia is unable to account for several clinical and pharmacologic observations, and other pathologic mechanisms are likely to play a role (for a lengthier discussion, see [5]). Among these mechanisms, the role of toxic metabolites resulting in oxidative damage has been proposed since the mid-1950s [6]. In their 1954 manuscript, Hoffer, Osmond, and Smythies purported that a highly reactive "adrenochrome" was involved in the evolution of schizophrenia symptoms. Although this adrenochrome hypothesis was quickly shown to be inadequate, oxidative insults to the brain in schizophrenia have continued to be explored.

The largest body of evidence of lipid peroxidation in schizophrenia is from indirect markers such as activity of SOD, catalase, and glutathione peroxidase. Increased peripheral activity of SOD has been observed in patients with schizophrenia, correlating with positive symptoms (hallucinations, delusions, disorganized behavior), and is reduced upon administration of antipsychotic medications [7, 8]. However, several reports have suggested that increased SOD activity occurs late in the disease, with reduced SOD activity early and in younger patients, implying SOD function varies with disease progression [9, 10]. Recently, study of SOD polymorphism in patients with schizophrenia has shown an association between the Ala-9Val variant and poorer performance in neuropsychological assessment [11]. Catalase activity in schizophrenia has been reported as increased, unchanged, or reduced by different investigators, although it is likely that peripheral levels poorly approximate brain levels of catalase especially since catalase activity can be dramatically altered by antipsychotic medications [12–14]. Finally, glutathione peroxidase has been generally reported to be reduced peripherally as well as in postmortem brains. Overall, these findings imply that there is a prooxidative state in schizophrenia. Assessment of more direct markers of lipid peroxidation has generally supported the more indirect assessments—patients with schizophrenia have greater plasma and cerebrospinal levels of thiobarbituric acid reactive substances, increased exhaled pentane, and increased urinary isoprostane-8-epi-prostaglandin F2 alpha [15–18].

Clinical interventions attempting to mitigate the effects of lipid peroxidation have been attempted including administration of eicosapentaenoic acid, long-chain omega-3 fatty acid, vitamin antioxidant (primarily vitamins E and C), and N-acetyl cysteine [19–23]. However, no placebo-controlled study to date has demonstrated long-term symptom improvement following normalization of lipid peroxidation. While some have reported that antipsychotic medications used for schizophrenia have altered markers of oxidative stress, others have found no improvement [24–26]. Interestingly, electroconvulsive therapy in schizophrenia patients reduces plasma thiobarbituric acid reactive substances, which is only evident months after the initial therapy [13].

4. Bipolar Disorder

Bipolar disorder is a mood disorder characterized by periods of both mania and depression. While its underlying pathophysiology remains multifaceted and elusive, recent data have indicated that mitochondrial dysfunction and aberration in oxidation status are important components of bipolar disorder. Andreazza and colleagues have previously shown that thiobarbituric acid reactive substances are elevated in both episodes of mania and depression and are even elevated compared to controls when in remission [27, 28]. Beyond a general prooxidative state, oxidative damage appears to also be localized in bipolar disorder. Using postmortem brain samples, Andreazza and colleagues have noted increased carbonylation and nitration along with isoprostane-8-epi-prostaglandin F2 alpha levels in the prefrontal cortex of bipolar disorder patients [29]. Additionally, oxidation also appears to localize to dopamine transporter-positive and tyrosine-hydroxylase neurons in the prefrontal cortex [30]. In depressed or euthymic females patients with bipolar disorder, sleep cycle disruption positively correlates with measures of malondialdehyde levels, a relationship not found in age-matched healthy controls which suggests that lipid peroxidation follows disease course rather than occurring incidentally [31]. Catalase and glutathione peroxidase levels are also elevated in patients with bipolar disorder during depressive episodes [32]. Treatment with the mood-stabilizer lithium reduces thiobarbituric acid reactive substances in those patients presenting for an initial manic episode as well as in those patients with episodes of hypomania (bipolar disorder type II) [32, 33]. Lithium administration in bipolar disorder patients has been noted to increase the activity of the Na+K+ATPase, a cellular event which is independently associated with reduction in lipid peroxidation [34]. Correlation between peripherally collected markers of lipid peroxidation and magnetic resonance imaging of bipolar disorder patients also explains the majority of variance in white matter lesions on diffusion tensor imaging in their brains. Longitudinal prospective trials that reduce oxidative burden in patients in addition to treatment with mood stabilizers would be instructive to see how much of the disease burden can be attributable to lipid peroxidation in bipolar disorder.
5. Major Depressive Disorder

Major depressive disorder is characterized by depressed mood and symptoms that significantly impair normal function. The mainstay pharmacologic therapy in major depressive disorder involves usage of serotonergic and noradrenergic agents for symptomatic control, but several observations have led to consideration of lipid peroxidation targets in the disease. For example, individuals who are depressed as young adults show increased rate of cardiac morbidity and mortality later in life and meta-analyses have indicated that patients who are depressed after myocardial infarction have worse cardiac outcomes that non-depressed patients [35,36]. Since a major component of cardiovascular risk stems from lipid oxidation and atherosclerotic plaque progression, commentators have conjectured that elevated oxidative burden in major depressive disorder could account for this finding. In a recent study of community-dwelling elderly subjects, plasma levels of isoprostane-8-epi-prostaglandin F2 alpha were higher in those with depressive symptoms than in nondepressed controls [37]. These results have been observed in a younger cohort and other work has also indicated that increased urinary isoprostane-8-epi-prostaglandin F2 alpha is associated with depressed mood, especially in men [38,39].

Increased glutathione peroxidase and superoxide dismutase levels are found in depressed patients, particularly those with chronic disease and these elevations persist for months even after initiation of standard pharmacotherapy [40,41]. Thus, lipid peroxidation appears to be significantly perturbed in major depression, and its attenuation may be beneficial in reducing coincidental medical disease. As with the data in schizophrenia and bipolar disorders, longer-term studies must be carried out to better understand the role of lipid peroxidation in major depression.

6. Other Psychiatric Disorders

Evidence of lipid peroxidation has also been recently discovered in other psychiatric disorders. In a small study of adults with attention deficit hyperactivity disorder, serum malondialdehyde levels were elevated compared to nondiseased controls; however no correlation was found between symptom severity on neuropsychiatric battery and levels of malondialdehyde [42]. In children with attention deficit hyperactivity disorder, lipid peroxidation products are also found to be increased in urine when compared to controls [43]. By contrast, serum levels of malondialdehyde were found to correlate with behavioral assessment of severity of obsessive compulsive disorder [44]. Several reports have also described an association between other anxiety disorders and oxidative stress, including generalized anxiety disorder, panic disorder, and posttraumatic stress disorder [45–47]. Elevated markers of oxidation have also been found in brain regions of patients with autism spectrum disorder, including in temporal cortex and cerebellum [48–50]. Investigation of oxidation stress and lipid peroxidation in other psychiatric illness such as personality disorders and eating disorders, however, has not been extensively conducted.

7. Conclusion: Therapeutic and Diagnostic Implications

In summary, recent clinical data has revealed oxidative damage and lipid peroxidation is seen in several psychiatric disorders. Larger longitudinal studies must be conducted to see if measures of lipid peroxidation and oxidative stress can be used to determine risk of developing various psychiatric diseases and to see if long-term disease outcomes can be modified by interventions that mitigate reactive oxygen species. Assessment of oxidative damage and lipid peroxidation in patients is currently limited to using indirect peripheral assessment of brain lipid peroxidation or postmortem analysis of brain samples. Since lipid peroxidation appears to be present in several psychiatric disorders, measures of lipid peroxidation are unable to be used as specific biomarkers for screening or monitoring disease progression. Development of more sophisticated methods of detecting lipid peroxidation would undoubtedly be useful in this endeavor. Regardless, based on current evidence, further investigation of lipid peroxidation in psychiatric illness is likely to reveal clinically-relevant information and should be carried out.

Conflict of Interests

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

References

[1] R. M. Adibhatla and J. F. Hatcher, “Lipid oxidation and peroxidation in CNS Health and disease: from molecular mechanisms to therapeutic opportunities,” Antioxidants and Redox Signaling, vol. 12, no. 1, pp. 125–169, 2010.
[2] D. G. Smith, R. Cappaï, and K. J. Barnham, “The redox chemistry of the Alzheimer’s disease amyloid β peptide,” Biochimica et Biophysica Acta, vol. 1768, no. 8, pp. 1976–1990, 2007.
[3] C. M. Spickett, I. Wiswedel, W. Siems, K. Zarkovic, and N. Zarkovic, “Advances in methods for the determination of biologically relevant lipid peroxidation products,” Free Radical Research, vol. 44, no. 10, pp. 1172–1202, 2010.
[4] C. R. Wade and A. M. van Rij, “In vivo lipid peroxidation in man as measured by the respiratory excretion of ethane, pentane, and other low-molecular-weight hydrocarbons,” Analytical Biochemistry, vol. 150, no. 1, pp. 1–7, 1985.
[5] M. Toda and A. Abi-Dargham, “Dopamine hypothesis of schizophrenia: making sense of it all,” Current Psychiatry Reports, vol. 9, no. 4, pp. 329–336, 2007.
[6] A. Hoffer, H. Osmond, and J. Smythies, “Schizophrenia: new approach. II. Result of a year’s research,” The Journal of Mental Science, vol. 100, no. 418, pp. 29–45, 1954.
[7] X. Y. Zhang, Y. L. Tan, L. Y. Cao et al., “Antioxidant enzymes and lipid peroxidation in different forms of schizophrenia treated with typical and atypical antipsychotics,” Schizophrenia Research, vol. 81, no. 2–3, pp. 291–300, 2006.
[8] X. Y. Zhang, D. F. Zhou, L. Y. Cao, P. Y. Zhang, and G. Y. Wu, “Elevated blood superoxide dismutase in neuroleptic-free schizophrenia: association with positive symptoms,” Psychiatry Research, vol. 117, no. 1, pp. 83–88, 2003.
[9] S. Mukherjee, S. P. Mahadik, R. Scheffer, E. E. Correnti, and H. Kelkar, “Impaired antioxidant defense at the onset of psychosis,” Schizophrenia Research, vol. 19, no. 1, pp. 19–26, 1996.

[10] P. K. Ranjekar, A. Hinge, M. V. Hegde et al., “Decreased antioxidant enzymes and membrane essential polyunsaturated fatty acids in schizophrenic and bipolar mood disorder patients,” Psychiatry Research, vol. 121, no. 2, pp. 109–122, 2003.

[11] X. Y. Zhang, D. C. Chen, M. H. Xiu et al., “Cognitive function, plasma MnSOD activity, and MnSOD Ala-9Val Polymorphism in patients with schizophrenia and normal controls,” Schizophrenia Bulletin, vol. 40, no. 3, pp. 592–601, 2014.

[12] N. Srivastava, M. K. Barthwal, P. K. Dalal et al., “Nitrite content and antioxidant enzyme levels in the blood of schizophrenia patients,” Psychopharmacology, vol. 158, no. 2, pp. 140–145, 2001.

[13] S. Kartalci, A. B. Karabulut, A. C. Ozcan, E. Porgali, and S. Unal, “Acute and chronic effects of electroconvulsive treatment on oxidative parameters in schizophrenia patients,” Progress in Neuro-Psychopharmacology and Biological Psychiatry, vol. 35, no. 7, pp. 1689–1694, 2011.

[14] H. Herken, E. Uz, H. Özyurt, S. Söğüt, O. Virit, and O. Akyol, “Evidence that the activities of erythrocyte free radical scavenging enzymes and the products of lipid peroxidation are increased in different forms of schizophrenia,” Molecular Psychiatry, vol. 6, no. 1, pp. 66–73, 2001.

[15] M. Phillips, M. Sabas, and J. Greenberg, “Increased pentane and carbon disulfide in the breath of patients with schizophrenia,” Journal of Clinical Pathology, vol. 46, no. 9, pp. 861–864, 1993.

[16] R. G. McCready, E. MacDonald, D. Wiles, G. Campbell, and J. R. Paterson, “The Nithsdale schizophrenia surveys. XIV: plasma lipid peroxide and serum vitamin E levels in patients with and without tardive dyskinesia, and in normal subjects,” British Journal of Psychiatry, vol. 167, pp. 610–617, 1995.

[17] S. P. Mahadik, S. Mukherjee, R. Scheffer, E. E. Correnti, and J. S. Mahadik, “Elevated plasma lipid peroxides at the onset of nonaffective psychosis,” Biological Psychiatry, vol. 43, no. 9, pp. 674–679, 1998.

[18] A. Dietrich-Muszalska and B. Olas, “Isoprostanes as indicators of oxidative stress in schizophrenia,” World Journal of Biological Psychiatry, vol. 10, no. 1, pp. 27–33, 2009.

[19] P. Fusar-Poli and G. Berger, “Eicosapentaenoic acid interventions in schizophrenia: meta-analysis of randomized, placebo-controlled studies,” Journal of Clinical Psychopharmacology, vol. 32, no. 2, pp. 179–185, 2012.

[20] G. P. Amminger, M. R. Schäfer, K. Papageorgiou et al., “Long-chain ω-3 fatty acids for indicated prevention of psychotic disorders: a randomized, placebo-controlled trial,” Archives of General Psychiatry, vol. 67, no. 2, pp. 146–154, 2010.

[21] K. Soares-Weiser, N. Maayan, and J. McGrath, “Vitamin E for neuroleptic-induced tardive dyskinesia,” Cochrane Database of Systematic Reviews, vol. 16, no. 2, Article ID CD000209, 2011.

[22] G. N. Dakhale, S. D. Khanzode, S. S. Khanzode, and A. Saoji, “Supplementation of vitamin C with atypical antipsychotics reduces oxidative stress and improves the outcome of schizophrenia,” Psychopharmacology, vol. 182, no. 4, pp. 494–498, 2005.

[23] M. Berk, D. Copolov, O. Dean et al., “N-acetyl cysteine as a glutathione precursor for schizophrenia—a double-blind, randomized, placebo-controlled trial,” Biological Psychiatry, vol. 64, no. 5, pp. 361–368, 2008.

[24] J. K. Yao, R. Reddy, L. G. McElhinny, and D. P. van Kammen, “Effects of haloperidol on antioxidant defense system enzymes in schizophrenia,” Journal of Psychiatric Research, vol. 32, no. 6, pp. 385–391, 1998.

[25] A. Sarandol, S. Kiriçi, C. Akkaya, A. Altin, M. Demirci, and E. Sarandol, “Oxidative-antioxidantive systems and their relation with serum S100 B levels in patients with schizophrenia: effects of short term antipsychotic treatment,” Progress in Neuro-Psychopharmacology and Biological Psychiatry, vol. 21, no. 5, pp. 550–552, 2007.

[26] O. Virit, A. Altindag, M. Yumru et al., “A defect in the antioxidant defense system in schizophrenia,” Neuropsychobiology, vol. 60, no. 2, pp. 87–93, 2009.

[27] A. C. Andreazza, M. Kauer-Sant'Anna, B. N. Frey et al., “Oxidative stress markers in bipolar disorder: a meta-analysis,” Journal of Affective Disorders, vol. 111, no. 2–3, pp. 135–144, 2008.

[28] M. Kunz, C. S. Gama, A. C. Andreazza et al., “Elevated serum superoxide dismutase and thiobarbituric acid reactive substances in different phases of bipolar disorder and in schizophrenia,” Progress in Neuro-Psychopharmacology and Biological Psychiatry, vol. 32, no. 7, pp. 1677–1681, 2008.

[29] A. C. Andreazza, J. F. Wang, F. Salmasi, L. Shao, and L. T. Young, “Specific subcellular changes in oxidative stress in prefrontal cortex from patients with bipolar disorder,” Journal of Neuroscience, vol. 127, no. 4, pp. 552–561, 2013.

[30] H. K. Kim, A. C. Andreazza, P. Y. Yeung, C. Isaacs-Trepanier, and L. T. Young, “Oxidation and nitration in dopaminergic areas of the prefrontal cortex from patients with bipolar disorder and schizophrenia,” Journal of Psychiatry & Neuroscience, vol. 39, no. 1, pp. 130–155, 2014.

[31] L. E. Cudney, R. B. Sassi, G. A. Behr et al., “Alterations in circadian rhythms are associated with increased lipid peroxidation in females with bipolar disorder,” The International Journal of Neuropsychopharmacology, vol. 17, pp. 1–8, 2014.

[32] R. T. de Sousa, C. A. Zarate Jr., M. V. Zanetti et al., “Oxidative stress in early stage bipolar disorder and the association with response to lithium,” Journal of Psychiatric Research, vol. 50, pp. 36–41, 2014.

[33] R. Machado-Vieira, A. C. Andreazza, C. I. Viale et al., “Oxidative stress parameters in unmedicated and treated bipolar subjects during initial manic episode: a possible role for lithium antioxidant effects,” Neuroscience Letters, vol. 421, no. 1, pp. 33–36, 2007.

[34] U. Banerjee, A. Dasgupta, J. K. Rout, and O. P. Singh, “Effects of lithium therapy on Na+-K+-ATPase activity and lipid peroxidation in bipolar disorder,” Progress in Neuro-Psychopharmacology and Biological Psychiatry, vol. 37, no. 1, pp. 56–61, 2012.

[35] J. P. van Melle, P. de Jonge, T. A. Spijkerman et al., “Prognostic association of depression following myocardial infarction with mortality and cardiovascular events: a meta-analysis,” Psychosomatic Medicine, vol. 66, no. 6, pp. 814–822, 2004.

[36] R. Rugulies, “Depression as a predictor for coronary heart disease: a review and meta-analysis,” The American Journal of Preventive Medicine, vol. 23, no. 1, pp. 51–61, 2002.

[37] N. Dimopoulos, C. Piperi, V. Psarra, R. W. Lea, and A. Kalofoutis, “Increased plasma levels of 8-iso-PGF2α and IL-6 in an elderly population with depression,” Psychiatry Research, vol. 161, no. 1, pp. 59–66, 2008.

[38] S. Yager, M. J. Forlenza, and G. E. Miller, “Depression and oxidative damage to lipids,” Psychoneuroendocrinology, vol. 35, no. 9, pp. 1356–1362, 2010.

[39] Y. Milaneschi, M. Cesari, E. M. Simonsick et al., “Health ABC study,” PLoS ONE, vol. 8, no. 6, Article ID e65406, 2013.
[40] C. Stefanescu and A. Ciobica, “The relevance of oxidative stress status in first episode and recurrent depression,” Journal of Affective Disorders, vol. 143, no. 1–3, pp. 34–38, 2012.

[41] P. Galecki, J. Szemraj, M. Bieńkiewicz, A. Florkowski, and E. Galecka, “Lipid peroxidation and antioxidant protection in patients during acute depressive episodes and in remission after fluoxetine treatment,” Pharmacological Reports, vol. 61, no. 3, pp. 436–447, 2009.

[42] M. Bulut, S. Selek, Y. Bez et al., “Lipid peroxidation markers in adult attention deficit hyperactivity disorder: new findings for oxidative stress,” Journal of Psychiatric Research, vol. 209, no. 3, pp. 638–642, 2013.

[43] M. Kawatani, H. Tsukahara, and M. Mayumi, “Evaluation of oxidative stress status in children with pervasive developmental disorder and attention deficit hyperactivity disorder using urinary-specific biomarkers,” Redox Report, vol. 16, no. 1, pp. 45–46, 2011.

[44] S. Chakraborty, O. P. Singh, A. Dasgupta, N. Mandal, and H. N. Das, “Correlation between lipid peroxidation-induced TBARS level and disease severity in obsessive-compulsive disorder,” Progress in Neuro-Psychopharmacology and Biological Psychiatry, vol. 33, no. 2, pp. 363–366, 2009.

[45] M. Bulut, S. Selek, Y. Bez et al., “Reduced PON1 enzymatic activity and increased lipid hydroperoxide levels that point out oxidative stress in generalized anxiety disorder,” Journal of Affective Disorders, vol. 150, no. 3, pp. 829–833, 2013.

[46] H. Herken, O. Akyol, H. R. Yılmaz et al., “Nitric oxide, adenosine deaminase, xanthine oxidase and superoxide dismutase in patients with panic disorder: alterations by antidepressant treatment,” Human Psychopharmacology, vol. 21, no. 1, pp. 53–59, 2006.

[47] E. Tezcan, M. Atmaca, M. Kuloglu, and B. Ustundag, “Free radicals in patients with post-traumatic stress disorder,” European Archives of Psychiatry and Clinical Neuroscience, vol. 253, no. 2, pp. 89–91, 2003.

[48] S. Rose, S. Melnyk, O. Pavliv et al., “Evidence of oxidative damage and inflammation associated with low glutathione redox status in the autism brain,” Translational Psychiatry, vol. 10, Article ID 2e134, 2012.

[49] F. Gu, V. Chauhan, and A. Chauhan, “Impaired synthesis and antioxidant defense of glutathione in the cerebellum of autistic subjects: alterations in the activities and protein expression of glutathione-related enzymes,” Free Radical Biology and Medicine, vol. 65, pp. 488–496, 2013.

[50] A. Chauhan, T. Audhya, and V. Chauhan, “Brain region-specific glutathione redox imbalance in autism,” Neurochemical Research, vol. 37, no. 8, pp. 1681–1689, 2012.