Cognitive Improvement with Glutathione Supplement in Alzheimer’s Disease: A Way Forward

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Abstract. Alzheimer’s disease (AD) is a devastating neurodegenerative disorder affecting millions of people worldwide. The actual cause of AD is still unknown. Oxidative stress is believed to be an important player in AD. Glutathione (GSH) is a major antioxidant, and it is already known that GSH is depleted significantly in the hippocampal regions. Hence, there is a serious discussion to improve the brain GSH level by supplementation. This editorial highlights the need for GSH supplementation for the cognitive reserve of AD.

Alzheimer’s disease (AD) is a progressive neurodegenerative disorder manifested by cognitive deterioration, progressive impairment of activities of daily living, and behavioral disturbances. [1, 2]. Current conceptualizations of AD presume that the neurodegenerative changes occur well before the clinical manifestations of the disease becomes apparent [3]. With progressive neuronal degeneration, formation of neurofibrillary tangles and neurotic plaques gradually increases. As a result, it sets a threshold for the initiation of clinical symptoms of AD associated cognitive deficits which gradually worsen with time. The fundamental molecular etiology of neuronal loss resulting in cognitive decline in AD is still unknown. Although there are existing data to support amyloid [4], tau [5], oxidative stress [6, 7], membrane alternation [8], and soluble oligomeric amyloid-β (Aβ) [9, 10] hypotheses, research in the clinical setting has indicated that oxidative stress plays an important role in the pathogenesis of AD [11–14]. Oxidative stress is a general term used to describe the steady state level of oxidative damage in a cell, tissue, or organ, caused by the reactive oxygen species (ROS). ROS are chemically unstable and highly reactive, but their levels are kept relatively low by antioxidants (e.g., glutathione (GSH), vitamin E & C, etc.). These facts underline the importance of an effective brain antioxidant system during the life span. Antioxidant strategies for AD include...
systematic treatment with vitamin C and E [15, 16], but results indicated that there were no significant clinical protective effects of vitamin E and C in dementia [17, 18].

Glutathione is a potent antioxidant and detoxifying agent that is produced de novo in the cytoplasm of every cell of the human body. The majority of GSH in the human brain are used for radical scavenging and balancing ROS. The intracellular brain antioxidant GSH interconverts into reduced GSH and GSSG, maintaining the ratio in most cells around 500:1 [19, 20]. Transgenic mice (which develop Aβ plaques throughout the cortex) show a significant decrease in GSH levels compared to wild type mice [21]. Studies in AD patients have shown that GSH metabolism is altered and regulated differently in male and female patients [22].

A recent study has shown that the depletion of GSH level in mild cognitive impairment (MCI) and AD patients in the hippocampal regions is directly corrected with executive function using the Trail Making (B-A) test score [12]. Another study has demonstrated that intranasal administration of GSH elevates brain GSH levels in the brains of Parkinson’s disease (PD) patients [23]. However, to the best of our knowledge, no clinical trial yet has evaluated similar effects of GSH oral supplement and its subsequent bioavailability in the brain tissue. There is no GSH trial available for AD patients yet, where GSH level in the brain will be monitored non-invasively as well as behavioral studies in mild cognitive impairment (MCI) patients concurrently using various neuropsychological tests. Hence this editorial sensitizes the importance of possible GSH supplementation of MCI patients to prevent cognitive performance from deterioration using peripheral supplements.

GLUTATHIONE DETECTION IN PLASMA AND AUTOPSY STUDIES

GSH is a tripeptide consisting of the three amino acids namely: glutamate, cysteine, and glycine. GSH exists in two forms: reduced (GSH) and oxidized (GSSG) forms. The reduced form of GSH has two predominant conformations (extended and closed) as shown by recently conducted multi-center in vivo MR studies [24, 25]. GSH is reported to decrease with age and age-associated disorders as revealed from autopsy brain studies [26, 27]. The GSH concentrations in the normal healthy brain were reported to be 1.18 ± 0.09 mM and 0.89 ± 0.03 mM, respectively, from the white and the gray matter of parietal cortex regions (N = 5) [28]. Another autopsy study has reported GSH level decrease in the hippocampus (N = 25) and frontal cortex (N = 31) with age [27]. A clinical study reported significant decrease in plasma GSH levels in MCI (N = 34) and AD (N = 45) compared to age-matched control subjects (N = 28) [29]. The glutathione peroxidase (GPx) in plasma were measured in MCI (N = 25), AD patients (N = 63) and controls subjects (N = 53). Further, the activities of antioxidants in plasma were depleted in MCI and AD patients as compared to controls [30]. Autopsy studies involving AD, PD, and dementia with Lewy bodies (DLB) have reported that the GSH level in the cingulated cortex region of AD patients is reduced by 49% as compared to age-matched control subjects. Such a specific change of GSH depletion is not found in PD and DLB patients [29]. Another postmortem study in MCI, mild and severe AD brains has revealed depleted GSH levels in post-mitochondrial supernatant, mitochondrial, and synaptosomal fractions from frontal cortices, as compared to controls [31].

DETECTION OF GSH IN VIVO IN HEALTHY AND MCI/AD PATIENTS USING MR SPECTROSCOPY

Autopsy [27] as well as neuroimaging [13] studies have reported close association of the GSH distribution in human brain with gender, age, disease condition, and anatomical area. Glutathione can be detected in vivo selectively by the application of specialized pulse sequences [13, 32, 33]. GSH is distributed in various neuroanatomical regions of healthy subjects as well as patients with MCI and AD as reported using in vivo MR studies [13]. Accordingly, healthy young brains showed higher GSH content and a specific GSH distribution pattern in various brain regions. The mean GSH content and distribution varied between male and female healthy subjects when compared between left frontal cortex (LFC) and right frontal cortex (RFC) [13]. It was reported that the reduction of GSH in both hippocampus and frontal cortices correlated (p < 0.001) with decline in cognitive functions [12]. Receiver operator characteristics analyses showed that hippocampal GSH depletion distinguishes between MCI and elderly healthy controls with 87.5% sensitivity, 100% specificity, whereas cortical region GSH level differentiates MCI and AD with 91.7% sensitivity,
100% specificity [12]. The significant reduction in GSH selectively in the brain regions (frontal cortex and hippocampus) affected by AD pathology, but not in the cerebellum of AD patients [12]. The autopsy studies indicated significant depletion of GSH level in cingulated cortex regions by 49% in AD subject [29]. Moreover, this GSH depletion trend is found in MCI, and AD subjects from in vivo MR spectroscopic studies of frontal cortices and hippocampal regions [12].

In another MRS study involving MCI subjects, it was found that GSH level increases with MCI [34] compared to elderly control. This increase in GSH level was explained as an early compensatory or neuroprotective response [34]. However, in the same study, the GSH peak was identified using PRESS pulse sequence in a 3T scanner where GSH peak appeared in close proximity of high intense choline peaks. The probable anomaly of these results is due to the use of a PRESS sequence for detection of GSH. With this technique the GSH signal will be partly overlapped by intense signals arising from choline and creatine, thus complicating the accurate measurement of GSH. The MEGA-PRESS pulse sequence, on the other hand, is a very selective and confirmatory method to detect the GSH peaks. Hence, in any clinical trial involving GSH measurement, MRS with MEGA-PRESS should be the method of choice for AD clinical study with GSH [23].

VARIOUS TRIALS WITH GSH AND OTHER ANTIOXIDANTS

There are various clinical trials involving GSH to investigate GSH transportation across the intestinal epithelium [35], therapeutic effects of oral administration of glutathione in patients with non-alcoholic fatty liver disease [36], effect of oral GSH supplementation on changes in erythrocyte GSH level [37, 38]. The effect of GSH on anti-aging features (wrinkles in the skin) was also investigated [39].

It was reported that GSH can be transported across the intestinal epithelium in an orally administered GSH in animal model study and the initial uptake of GSH into cells is reported as a rapid process [35]. The ingested GSH has potent nutraceutical benefits for human health to improve oxidative stress and defense in human [35]. The therapeutic effects of oral administration of GSH in patients with non-alcoholic fatty liver disease in open label, single arm, multicenter trial [36] indicated a significant decrease in alanine aminotransferase levels as well as significant decreased of triglycerides, non-esterified fatty acids, and ferritin levels decreased [36]. To determine the effect of oral GSH supplementation on erythrocyte GSH concentrations, including total reduced glutathione (GSH), oxidized glutathione (GSSG), and their ratio in a randomized, double-blind, placebo-controlled clinical trial for 4 weeks on 40 healthy volunteers were conducted. Changes in erythrocyte GSH concentrations, including total reduced glutathione (GSH), oxidized glutathione (GSSG), and GSH: GSSG ratios were monitored. No significant changes in erythrocyte GSH concentrations were observed [37]. In a long-term study, the effect of oral GSH supplementation on body stores of GSH in healthy adults, randomized, double-blind, placebo-controlled trial GSH supplement for 6 months on 54 adults was conducted. GSH levels in blood increased after 1, 3, and 6 months versus baseline. At 6 months, mean GSH levels increased 30–35% in erythrocytes, plasma, and lymphocytes and 260% in buccal cells in the high-dose group. GSH levels increased 17% and 29% in blood and erythrocytes respectively, in the low-dose group. Natural killer cytotoxicity increased twofold in the high-dose group versus placebo [38]. This clearly demonstrates that study duration for oral GSH supplementation plays an important role.

The orally administered glutathione (500 mg per day for 4 weeks, on 60 healthy medical students) affects the skin melanin index, when compared with placebo in a randomized, double-blind, two-arm, placebo-controlled study. Mean reduction of melanin indices measured at six different sites, UV spots. It was observed that melanin indices decreased consistently at all six sites. Both glutathione and placebo were very well tolerated. Oral glutathione administration results in a lightening of skin color [40]. In another similar type of study, to evaluate the influences of both GSH at doses lower than 500 mg/d, on improving skin properties using randomized, double-blind, placebo-controlled, study was conducted [39]. Oral GSH was administered for 12 weeks in 60 healthy volunteers, and skin features including melanin index, wrinkles, and other relevant biophysical properties were measured. Blood samples were collected for safety monitoring. GSH showed a significant reduction in wrinkles compared with those taking placebo. A tendency toward increased skin elasticity was observed in GSH compared with placebo. There were no serious adverse effects throughout the study [39].
A phase IIb randomized double blind, placebo controlled trial of 45 H&Y stage 1–3 PD patients monitored the symptomatic relief effect of intranasal GSH supplement by administering 200 mg GSH daily for three months. The subjects were observed over a one-month washout period. 15 of these subjects participated in a sub-study in which MRS was performed at baseline and at the end of three-month GSH regime. All cohorts of this study, including placebo showed improvement during the intervention regime. However, as this study aimed at evaluating an appropriate trial design, conclusions regarding the superiority of GSH over placebo were not drawn [23].

To compare bioavailability, the effect on oxidative stress markers and the safety of a new sublingual form of GSH was studied with two commonly used dietary supplements: N-acetylcysteine (NAC) and oral GSH. In this randomized crossover study, sublingual GSH, Oral and NAC was administered for 3 weeks in 20 normal volunteers. Bioavailability, antioxidant efficacy, tolerance, reduced thiols, vitamin E, lipid status and adverse effects were monitored. Significant superiority of a new sublingual form of GSH compared to NAC, better effects of the sublingual form of GSH were also observed. All the dosage forms were very well tolerated and no adverse events were reported [41]. A nutraceutical formulation containing N-acetylcysteine among other compounds has shown some pro-cognitive benefits in AD and older adults, but the evidence for N-acetylcysteine alone is much weaker [42].

**FUTURE TRIAL INVOLVING GSH FOR COGNITIVE IMPROVEMENT IN AD**

To date, all GSH trials conducted address the bioavailability, tolerance level, dose dependence, and duration of the oral supplementation. Existing literature clearly indicates that oral supplementation for 500 mg/day indeed enters the blood stream and has enhanced the GSH level in the erythrocyte. To the best of our knowledge, there is no study/trial available in the clinical trial database or other cognitive performance studies. Therefore, there is an urgent need for the study of GSH supplementation involving MCI and AD patients. Further, MRS measurement of GSH in the hippocampus needs to be monitored on those participants at various time points. The neuropsychological battery of tests at various time points for cognitive profile evaluation should be part of the study. Finally, the GSH level in the hippocampus region and its correlation with the cognitive score will be a great measure for the identification of the impact of GSH supplementation both for MCI and AD compared to subjects treated with placebo.

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