Depletion of Glutathione during Oxidative Stress and Efficacy of N-Acetyl Cysteine: An Old Drug with New Approaches
Manjula Ramen T*
Sri Ramachandra Medical College and Research Institute, Sri Ramachandra University, Chennai, India

Abstract
Glutathione, a non-protein thiol is most abundant in cells and glutathione-glutathione disulfide is the major redox pair in animal cells. Its synthesis is from two main enzymes gamma glutamyl cysteiny1 synthetase and GSH synthetase. Also cysteine availability as N-acetyl cysteine and GSH feedback mechanism determines the reduced glutathione status of the cell. Additionally parenteral NAC, cysteine, methionine are effective precursors of cysteine for tissue GSH synthesis. Adequate protein in the diet is crucial for tissue GSH synthesis. The role of NAC as an antioxidant, in gene expression, and regulation of cellular events must be emphasized. GSH deficiency contributes to oxidative stress and it progresses in diseased conditions such as HIV, AIDS, cancer, diabetes, also metabolic syndrome. New knowledge of the nutritional regulation of GSH metabolites is critical to improve health and treat these diseases.

Keywords: N-Acetyl L Cysteine; Oxidative stress; Antioxidant; Cell regulator

Introduction
Glutathione is a tripeptide that maintains intracellular levels of reduced glutathione. Its redox action makes it an important biological antioxidant. Gamma-glutamyl-cysteinyl-glycine is synthesized from glutamate, together with cysteine and glycine through the gamma glutamyl cycle. As a key modulator of cell functions, the most abundant non protein thiol (GSH) has important roles in cellular defense against cellular oxidant aggression, redox regulation of protein thiols and maintaining redox homeostasis that is critical for proper functioning of cellular processes including apoptosis. The shift in the cellular GSH-GSSG decides the fate of the cell.

Glutamate is a precursor of two gamma-glutamyl compounds of major biochemical importance glutamine and glutathione. Gamma amino butyrate, a neurotransmitter and glutatione are synthesized from glutamate. Glutamine synthetase reaction takes place in a number of mammalian tissues eg, liver, brain, kidney, muscle, intestine [1]. It is the central compound in nitrogen metabolism. It functions in the uptake, storage and formation of ammonia, the homeostatic control of amino acid balance, the synthesis of the purine and pyrimidine moieties of nucleic acids, ATP, and other nucleotides and coenzyme formation and formation of amino sugars.

The synthesis of glutathione involves the uptake and release of amino acids from gamma glutamyl linkage, in a cyclical process termed, the gamma glutamyl cycle (Figure 1). The gamma glutamyl cycle has properties that fulfill the requirements of an amino acid transport system.

The presence of gamma glutamyl cyclotransferase, transpeptidase and gamma glycylsynthetase leads to catalytic events of synthesis and breakdown of glutathione and coupled uptake and release of free amino acids from gamma glutamyl linkages through gamma glutamyl cycle.

The gamma glutamyl cycle may play a role in the transport or secretory functions of many tissues and the ubiquitous occurrence of glutathione reflects its central role in amino acid transport [2].

The transport of amino acid involves a number of steps such as binding of the amino acid to the site on the cell membrane, its carrier mediated translocation, intracellular release of the amino acid from the carrier and reactivation of the carrier in an energy requiring process utilizing glutathione.

*Corresponding author: Manjula Ramen T, Sri Ramachandra Medical College and Research Institute, Sri Ramachandra University, Chennai, India, Tel: 91-44-45928500; E-mail: manjarumtk@yahoo.co.in

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Excess production of free radicals such as superoxide (O2), hydrogen peroxide (H2O2) and hypochlorous acid (HOCl) by activated phagocytes are produced as a consequence of the underlying disease process.

Oxidation-reduction based regulation [5,6] of signal transduction and gene expression is emerging as a fundamental regulatory mechanism in cell biology. Electron flow through side chain functional CH2SH of conserved cysteinyl residues in proteins account for their redox signaling properties. Because in most intracellular proteins, thiol groups are strongly buffered against oxidation by the highly reduced environment inside the cell.

The list of redox sensitive signal transduction pathways are steadily growing and current information suggests manipulation of the cell redox state may prove to be a strategy for the management of AIDS and some forms of cancer. The endogenous thioredoxin and glutathione system are of central importance in redox signaling and N-acetyl L-cysteine (NAC) and alpha lipoic acid are known for clinical use in their efficacy to modulate cellular redox status. Lipoate captures the metabolic power of cell to continuously regenerate its reductive dithiol form.

N-Acetyl L-Cysteine (NAC)

NAC the preacetylysed form of simple amino acid cysteine, a synthetic precursor of cysteine and reduced glutathione has been in clinical use for more than 40 years. It is a powerful intracellular antioxidant, antitoxin improves immunity and is found naturally in foods.

Structure

NAC is a thiol compound which has a chemical formula C5H9NO3S and a molecular weight of 163.2 [7] NAC gets deacetylated to cysteine in the liver [8] In addition to increase in free and total NAC, additional increase in non-protein and protein SH groups, and small molecular weight protein bound thiols are found in the human plasma.

Both L-Cysteine and glutathione has been explored for their usage as supplements and NAC shows increased bioavailability. Glutathione has limited therapeutic usage as it is being rapidly hydrolysed by the intestines.

NAC may have a direct chelating effect on lead as seen by lowered serum lead levels. It prevents lead toxicity and reduces oxidative sequelae of lead exposure. NAC crosses cell membranes and is rapidly consumed in producing intracellular glutathione. By reducing extracellular cystine to cysteine, it acts as a source of SH groups and it can stimulate glutathione synthesis enhance glutathione-S-transferase activity promote detoxification and act directly on reactive oxidant radicals [9].

NAC corrects the reduction in glutathione concentration and results in significant preservation of fluidity of membranes and of the activities of catalase, mitochondrial superoxide dismutase and different forms of glutathione peroxidase in biliary obstructed rats [10]. NAC is a powerful scavenger of hypochlorous acid and is capable of reducing hydroxyl radicals and hydrogen peroxide. SH groups are essential for defence against reactive oxygen species [11]. NAC can also prevent apoptosis caused by oxidative stress and promote cell survival by activating signal regulating pathways [12].

NAC on oxidative damage in urolithiasis

Hyperoxaluria is one of the major risk factors in human idiopathic calcium oxalate stone disease and its experimental induction was studied in rats. Exposure to oxalate has been shown to be toxic to renal epithelial cells with increased urinary excretion of marker enzymes [13]. Renal cell damage is associated with lipid peroxide production primarily caused by hyperoxaluria and the development of tissue injury is relative to its oxidant – antioxidant status of the cells [14]. Increased urinary calcium levels found in hyperoxaluric animals was controlled using methionine [15]. Phyllanthusniruri normalizes elevated urinary calcium levels found in stone formers [16] due to antioxidant action of NAC. Jonassen [17] proposed high concentrations of oxalate promotes stone formation in 2 ways.
1) By providing urinary conditions favorable to the formation of calcium oxalate crystal

2) By inducing renal injury that generates cellular debris and promotes crystal nucleation and attachment. NAC decreased urinary oxalate in hyperoxaluric animals which coincided with cysteine administration to ethylene glycol treated rats [18] with the decrease in urinary oxalate due to an adduct formation with cysteine and its conversion to oxalate. A similar mechanism may be proposed for NAC both being structural identities. Increased activities of urinary marker enzymes gamma glutamyl transferase, alkaline phosphatase, lactate dehydrogenase and pyrophosphatase were seen in hyperoxaluric rats indicative of cell injury [19,20] mainly proximal tubular epithelium damage. Histologic changes in kidney tissue showed cellular injury in hyperoxaluria induced rats with bleb formation but NAC pretreated rats showed improvement. (Proceedings NHFR Conference2004)

**NAC on oxidative damage in diabetic rats**

NAC is a potent antioxidant due to its ability to stimulate reduced glutathione (GSH) synthesis therefore maintaining intracellular levels. NAC can be used as an antioxidant agent in alloxa- induced diabetic rats showing modulatory action on oxidative stress biomarkers [21].

Pubmed Therapeutic goods administration of Australia and Food Administration suggests NAC use in psychiatric disorders schizophrenia and bipolar disorder. Enhanced oxidative stress pathogenesis of several disorders like ischemic diseases of the heart, brain, neurogenerative disorders and cancer and also in human immunodeficiency virus (HIV) infection [22-26] as impaired antioxidant disturbance and particularly disturbed glutathione metabolism is seen in HIV patients [25-28]. Also impaired responsiveness and enhanced apoptosis may possibly involve increased intracellular hydroxyl radicals [10] tumor necrosis factor (TNF)-alpha related mechanisms [29,30]. Consumption of GSH is high during HIV infection and disease progression. GSH is known to play a major role in regulation of T-cell immune function. The clinical significance of the HIV associated glutathione disturbances is reflected as low thiol levels in both plasma [31] and CD4+ T cells [32] and is strongly associated with decreased survival of HIV infected patients. GSH depletion can enhance oxidative stress and also may increase excitotoxic molecules that can initiate cell death. GSH status is diminished in Lou Gehrig’s disease (ALS)/Parkinson’s and Alzheimer’s disease. Hence can propose potential therapeutic approaches for preventing these diseases [33].

**References**