Review on Advanced Glycation End Products and the Progress of Chronic Disorders

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Submission: February 12, 2018; Published: March 14, 2018

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Abstract

The study of AGEs represents one of the most promising areas of research today. This review covers the information and generation of AGEs in the field of disorders, which will be useful for biomedical professionals.

Keywords: Advanced glycation end products, chronic disorders, Amadori product

Introduction

Advanced glycation end products are a complex and heterogeneous group of compounds that is produce by non-enzymatic and spontaneous reactions of the carbonyl group of carbohydrates with proteins, nucleic acids or lipids through a series of reactions. In fact this process have two step:

i. The condensation of a sugar aldehyde with a free amino group of proteins (usually hydroxylsine or arginine residues) forming a non-stable Schiff base

ii. Formation of Amadori products from Schiff base.

This glycation process, is also known as the Millard action which was described in the early 1912’s, by Louis-Camille Maillard. The Schiff base and especially Amadori products may undergo oxidation, dehydration, and polymerization to rise other AGEs. The AGEs that formed under oxidative conditions called advanced glycoxidation end products [1].

Exogenous AGEs

Animal and human based studies have shown that along with endogenously production of AGEs in the body, exogenous AGEs can found naturally in many foods and tobacco. They are in the foods that cooked at high temperatures, especially animal-derived products that are rich in protein and fat. Indeed, some of the cooking methods including barbecuing, grilling, roasting, and frying may contribute to higher dietary AGEs [2]. Unfortunately, the differentiation of dietary AGEs type, amount and destructive mechanism with endogenously AGEs is not well known.

Most commonly AGEs

Pentosidine is a fluorescent glycoxidation product that is isolated by Sell and Monnier, as the common known AGEs. It can form by cross linking the glucose, fructose, or ascorbate to an arginine and a lysine residue [3].

Carboxymethyl-lysine (CML) also known as N (epsilon)-(carboxymethyl) lysine is a well-characterized and non-fluorescent AGEs that was first described by Ahmed in 1985. The CML formed by oxidation of fructosyl-lysine as an Amadori product or by reaction of glyoxal with the amino group of lysine [4]. It is the most prevalent marker for AGEs concentrations analysis in the foods.

Dicarbonyl compounds

Dicarbonyl compounds derive from oxidative degradation or autooxidation of Amadori products, which are leading to protein crosslinks. Methylglyoxal (MGO), glypxal (GO) and 3-deoxyglucosone (3-DG) are three well-known dicarbonyl compounds [5].

Receptors of AGEs

AGEs do not only act by altering the physicochemical properties of glycated proteins. Interestingly, many cells in the body such as endothelial cells and smooth muscle have receptors for binding AGEs (RAGEs). These receptors are not specific for AGEs, as other molecules such as S-100/calgranulins, amphotericin, β-amyloid peptides and β-sheet fibrils can bind them [6]. Excess accumulation of AGEs in the body may adversely affect these cells.

AGEs in disorders

AGEs affect nearly every type of cell and molecule in the body, which have pathogenic role in some conditions that causes intracellular damage and apoptosis. Recent studies has
been work done to elaborate on the treatment and prevention of related complications to some disorders including aging, coronary artery disease, kidney failure, Alzheimer’s disease, osteoarthritis, and diabetes and demonstrated that most of the complications especially in diabetes patients are due to the accumulation of AGEs [7]. Chronic disorders are significantly associated with oxidative stress in humans and animals, as decreased expression of antioxidative enzymes along with NADPH oxidase activation are contribute to high production of ROS in the tissues. Activation of nuclear factor (NF)-κB, and JNK pathways is accompanied by obesity, high-fat diet, and cellular stresses (ROS), which can increase inflammatory responses in body tissues. Some ligands including TNFR, IL-1R, and RAGE that are receptors for TNF-α, IL-1, and AGEs can also activate the JNK pathway during metabolic dysregulation [8]. The processes that are occurred in diseases are the principal causative factors for formation of AGEs.

Conclusion

The study of AGEs represents one of the most promising areas of research today. Long-term formation of AGEs affects all long-lived proteins, which they can only be removed from the body when the protein is removed. Therefore, many health professionals are calling for AGE levels to become a marker of overall health. Although, most of the AGEs have not yet been isolated or characterized, Pentosidine, CML, and some Dicarbonyl compounds such as MGO are used to identifying AGEs concentrations in the body. Methylglyoxal has been suggested to be a better marker for investigating the association between AGEs with adverse health outcomes.

References