Review

Hypoxia-Induced Degenerative Protein Modifications Associated with Aging and Age-Associated Disorders

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Abstract: Aging is an inevitable time-dependent decline of various physiological functions that finally leads to death. Progressive protein damage and aggregation have been proposed as the root cause of imbalance in regulatory processes and risk factors for aging and neurodegenerative diseases. Oxygen is a modulator of aging. The oxygen-deprived conditions (hypoxia) lead to oxidative stress, cellular damage and protein modifications. Despite unambiguous evidence of the critical role of spontaneous non-enzymatic Degenerative Protein Modifications (DPMs) such as oxidation, glycation, carbonylation, carbamylation, and deamidation, that impart deleterious structural and functional protein alterations during aging and age-associated disorders, the mechanism that mediates these modifications is poorly understood. This review summarizes up-to-date information and recent developments that correlate DPMs, aging, hypoxia, and age-associated neurodegenerative diseases. Despite numerous advances in the study of the molecular hallmark of aging, hypoxia, and degenerative protein modifications during aging and age-associated pathologies, a major challenge remains there to dissect the relative contribution of different DPMs in aging (either natural or hypoxia-induced) and age-associated neurodegeneration.

Key words: protein aggregation, hypoxia, aging, neurodegenerative disease, dementia, proteostasis, proteomics

1. Introduction

Aging is nature’s most complex phenomena, where an inevitable time-dependent progressive functional decline of numerous physiological functions finally leads to death. These changes, either programmed or due to protein damage, lead to an imbalance in a regulatory system including hormones, repair, immune and neuroendocrine. Many theories of aging have been proposed [1]. The theory of longevity believes switching “on” and “off” of certain genes [2, 3]. The endocrine theory considers hormones control biological clocks [1].

While the immune theory recommends the programmed decline of immunity over time [4]. Of numerous damage theories, i.e. wear and tear, cross-linking and free radical theory; the majority of them are centered on protein damages through cross-linking, oxidative modifications by reactive oxygen species (ROS) or reactive nitrogen species (RNS) [1]. In short, aging is a consequence of two independent biological processes, one programmed loss of functionality and other damage related changes [1-3]. The damage-related changes include protein aggregation, structural modifications, loss of resistance to stress, and failure to repair the damages. According to an aging

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theory proposed by Harman in 1956 [5], the alterations in biological function with time is due to the accumulation of damages caused by free radicals.

It is not clear how the loss of physiological integrity with cumulative dysfunction occurs in cells, and which factors are responsible for aging and unique aspects increase susceptibility to diseases, respectively. Basically, proteins, carbohydrates, lipids, and nucleic acids are four fundamental molecules without which cells and organisms cannot be living, hence they are named “molecules of life”. Aging of these molecules cannot be ignored since they are susceptible to free radical attack and actively involved in tissue aging. Recently, Lopez-Otin et al., [6] reviewed the recent literature and proposed nine tentative hallmarks of aging including genomic instability, epigenetic alterations, loss of proteostasis, telomere attrition, deregulated nutrient sensing, mitochondrial dysfunction, cellular senescence, stem cell exhaustion, and altered intercellular communication (Fig. 1). In all nine hallmarks of aging, proteins remains as a key component e.g. proteins stabilize genome through dynamic modification of chromatin architecture [7], epigenetic alterations are regulated by histones [8], proteostasis and mitochondrial functions are governed by proteins, and DNA-damages are repaired by proteins, and damaged DNA are finally translated into proteins. Thus, the protein damage could be crucial in aging and pathogenesis.

Proteins are the building blocks of life as they are not only the structural constituents of the living organisms but also a final functional molecule governing most of the biological functions. The proteins undergo alterations by spontaneous non-enzymatic Degenerative Protein Modifications (DPMs) including oxidation, deamidation, carbamylation, carboxylation, glycation etc. The DPMs change protein charge state, hydrophobicity and three-dimensional structure that influence functional activities and induce aggregation [9-15]. These protein modifications and accumulation of modified proteins are allied to aging and the development of age-associated pathologies like neurodegenerative diseases [10, 16]. DPMs like spontaneous protein deamination characterized by the modification of glutaminyl and asparaginyl residues were hypothesized as a molecular timer of biological events including protein turn over, development and aging [17-20]. Protein deamination progressively disrupts structural integrity of the protein and alter their biological activity [12, 13, 21]. Other
DPMs including glycation, advanced glycation end products, oxidation, carbonylation, carbamylation, etc., impart deleterious structural and functional changes in proteins and impair their normal function [12, 13, 22, 23].

Hypoxia, a condition where oxygen supply to tissue is inadequate, induces free radical generation leading to oxidative protein modifications and tissue damage [24-27]. Oxygen supply also acts as a modulator of aging processes [28]. The cerebrovascular disorders and hypoxia-ischemia injuries in the brain are projected as a primary cause of protein pathologies that leads to cognitive impairment and dementia [29, 30]. In short, hypoxia-ischemia injury in the brain persuades DPMs that can lead to aging, age-associated diseases and neurodegeneration. The mechanisms that initiate and promote DPMs remain poorly understood, partly due to technical challenges. However, mass spectrometry not only has the potential to identify and quantify proteins but also identify DPMs quantitatively (Fig. 2). Therefore, the main purpose of this review was to present the recent advances in DPMs research in aging, hypoxia, and neurodegenerative diseases. Given the complexity of DPMs and their role in aging and age-related neurodegenerative diseases, we highlighted current discoveries of DPMs and their impacts on various cellular processes. We attempted to identify the linkage between hypoxia-induced protein modifications and dysfunctions, the aging process, and age-related neurodegenerative diseases.

Figure 2. Advanced novel proteomic approaches to elucidate degenerative protein modifications (DPMs) and post-translational modifications (PTMs) of protein in aging, hypoxia and neurodegenerative diseases or other biological samples. (VaD: vascular dementia, CSF: cerebrospinal fluid, CI: cerebral ischemia, AD: Alzheimer disease, PD: Parkinson disease)
2. Hypoxia

2.1. Correlation between hypoxia, aging and neurodegenerative diseases

The regular function of virtually all tissues and organs depends on adequate blood flow and oxygen supply. Any condition that results in insufficient blood flow leads to acute or chronic hypoxia and numerous complications including diminished metabolic processes, compromised neuronal signaling [31], cellular damages and functional impairments in the affected tissue[32]. Based on the oxygen level in tissue, hypoxia is classified as physioxia (5% oxygen), physiological hypoxia (2%), pathological hypoxia (1%), radiobiological hypoxia (0.4%) and anoxia (0%) [33]. The short-term exposure to hypoxia, ischemia, hyperoxia, hyperthermia, and hypothermia may enhance the adaptive mechanism by overcoming stress, and remains beneficial [34, 35]. In mild hypoxia, the oxygen sensing system stimulates gene expression including hypoxia-inducible transcription factor-1α (HIF-1α), which in turn controls the expression of survival genes to protect the tissue from damages[36]. It’s important to note that the arterial blood oxygen concentration remains at 9.5% (95000 ppm) and the cellular damage begins to occur when it drops below 5000 ppm oxygen [37].

Patients with obstructive sleep apnea, Cheyne-stokes respiration, and nocturnal hypotension experience chronic intermittent hypoxia (CIH) more often than continuous hypoxia [38]. CIH induces modification of the proteins involved in transcriptional activation, signaling pathways, transmitter synthesis and cardioprotection [39]. The decrease in air pressure and air density with increase in altitude (during aviation) result in hypobaric hypoxia, which causes neurodegeneration and memory impairment in animal models [40, 41]. Chronic cerebral ischemia induced by cerebral hyperperfusion in a bilateral carotid artery stenosis mouse model damaged brain tissue, intercellular communication and caused impairment of working memory and neurovascular unit dysfunction observed in early Alzheimer's disease [42, 43]. Biswal and colleagues [44] noted increased neurodegeneration with altered mitochondrial morphology and aggregation of lipofuscin granules in the hippocampus region. These authors also found an elevated level of pro-inflammatory S100A9 protein and reduced expression of synaptosome associated protein 25 (SNAP25), total Tau, superoxide dismutase 2 (SOD2), and apolipoprotein E (ApoE) in young rats when exposed to chronic global hypoxia. These findings correlate hypoxia and neurodegeneration with aging in the hippocampus. Further, neurodegeneration and dendritic atrophy in the hippocampus acts as a contributing factor for spatial memory impairment upon chronic exposure to hypobaric hypoxia [41]. The magnitude of deleterious effects increases with a decrease in oxygen concentration.

A continuous uninterrupted enough supply of oxygen to the brain is crucial to maintain its proper metabolic functions and to avoid tissue damage. However, diseases including cardiorespiratory (heart attack, lung injury, peripheral vascular disease, hypertension, asthma, bronchiecstasis, bronchitis etc.), carotid stenosis and cardiovascular disorders (coronary artery disease, heart attack, heart valve disease etc.) results in persistent systemic hypoxia. In hypoxia, the brain is not completely deprived of oxygen, but yet it causes chronic decreased cerebral oxygenation, generation of free radicals, oxidative stress, protein oxidation, altered neurotransmitter synthesis, cellular apoptosis, neurodegeneration, and memory impairment [45]. In hypoxia, the metabolic support for neuronal signaling gets compromised in many regions of the brain leading to brain injury, neurological and the memory dysfunction [46, 47]. While comparing time-dependent and region-specific changes in the cortex, hippocampus, and striatum upon hypoxia, Maiti et al., [48] found an increase in oxidative stress, free radicals, nitric oxide level, lipid peroxidation, and these authors concluded that the hippocampus is more susceptible to hypoxia than the cortex.

Hypoxia/reoxygenation (H/R) such as induced by ischemia/reperfusion results in neuronal injury mediated by the glutamate/N-methyl-D-aspartate (NMDA)/Ca++/nitric oxide (NO) and free radical pathway [49]. The increased neuronal damage with time was noted when primary cultures of rat cortical neurons and glia were exposed to H/R [50]. In hypoxic-ischemic condition, Kim et al [51] noted neuronal death due to the accumulation of Ca++ and Na+. Variation in oxygen supply to brain tissue in H/R triggers cellular and molecular alterations including protein turnover, protein aggregation, impairment in neural plasticity, perturbed calcium homeostasis, neuronal survival, neuroinflammation etc. which eventually affect brain functions leading to aging and neurological disorders. However, it’s somehow difficult to differentiate normal brain aging and hypoxia-induced aging, because in many cases normal brain aging switches to pathological aging with drastic deterioration in cognitive abilities and motor skills [52].

In hypoxia and under disrupted redox equilibrium due to excessive accumulation or depletion of ROS, the cellular signaling pathways are influenced leading to cellular dysfunction and development of various diseases [53]. Under physiological state, a balance between ROS generation and clearance is regulated by antioxidative defense mechanisms. The major antioxidant enzymes involved are Cu/Zn-superoxide dismutase (Cu/Zn-SOD, SOD1) in the cytosol, manganese superoxide dismutase (Mn-SOD, SOD2) in the mitochondrial matrix, catalase,
glutathione peroxidase (GPx), and glutathione reductase (GR). However, when ROS overproduction overcomes intrinsic antioxidant capacity, then the oxidative stress occurs, which damages the biomolecules of cells[54]. The oxidative stress usually results from either excessive ROS production, mitochondrial dysfunction, impaired antioxidant system, or a combination of these factors[54].

Exposing oxygen-sensing pheochromocytoma (PC12) cells, human embryonic kidney cells (HEK-293 in which stably expressed the human cardiac L-type Ca^{2+} channel α1C subunit), the human neuroblastoma (SH-SY5Y), cultures of cortical astrocytes and central neurons to hypoxia (1-2.5% oxygen) for a period of 6-48 h resulted in increased production of amyloid peptides that affected cell calcium channel currents and impaired calcium signaling [55, 56]. As reviewed by Peers, [56] disrupted oxygen supply to central nervous system promote the formation of ROS, alters mitochondrial energy metabolism, expression of various proteins of calcium homeostasis; causes neuronal death and the onset of dementia. In short, hypoxia induces production of amyloids alters signaling mechanism, causes mitochondrial dysfunction, and neuronal death. On the other hand, these alternations are the root cause of neurodegeneration.

Under the anoxic condition, the brain is completely deprived of oxygen due to sudden cardiac arrest, choking, strangulation, which may lead to major acute damage. Apparently, a few seconds of oxygen deprivation won’t cause major damage, but the total circulatory arrest results in loss of ATP production and dysfunction of membrane ATP–dependent Na-K pumps, affect the release of glutamate that further induces excitotoxic injury through NMDA receptors [57, 58]. Activation of the NMDA receptor elevates intracellular calcium, which further increases the free radical level [59, 60]. The elevated level of free radicals is known to cause more damage through lipid peroxidation, protein oxidation, and DNA fragmentation, all of which contribute to cell death [61]. The documented literature clearly indicates the deleterious effects of hypoxia in brain damage, signaling, cellular and metabolic processes. Strokes including both ischemic and hemorrhagic, deprive oxygen in the brain and lead to brain cells death. Desmond et al., [62] found the 4-fold increased risk of incident dementia among ischemic stroke patients who were initially non-demented relative to clinically stroke-free elderly control subjects.

Oxygen is the ultimate electron acceptor in the electron transport chain [48]. The oxygen level in cells remains low in hypoxic conditions, which leads accumulation of electrons. These accumulated electrons attack the ground state of available oxygen to form superoxide anion (O_2^-) and chain reaction, leading to the formation of H_2O_2 and hydroxyl radicals (OH^-). Importantly, hypoxia-induced oxidative stress not only impairs mitochondrial function, neuronal damage/apoptosis through the nitric oxide synthase (NOS) pathway, [63, 64]; but also causes damage to lipids, proteins, and DNA [65]. According to recent literature, cerebral hypoxia, ROS, mitochondrial dysfunction associated with accumulation of amyloid plaque, development of intra-neuronal neurofibrillary tangles, hyperphosphorylated tau, loss of synaptic integrity and neuronal death in the cortical region of brain are the main risk factors and causative agents of dementia [66, 67].

Mitochondria are important regulators of longevity, and a direct correlation exists between mitochondrial ROS production and lifespan [68, 69]. Mitochondrial DNA (mtDNA) is a major target for age-associated mutations due to the oxidative environment in mitochondria and lack of efficient histone protective mechanism [6]. The accumulated oxidative stress and ROS lead to oxidation of macromolecules. These oxidized macromolecules are no longer degraded or repaired due to defects in proteostasis [14, 70]. Under hypoxic conditions, protein oxidation and aggregation of lipofuscin in neurons were noted [71]. These events are a hallmark of aging [72]. Hypoxia induces ROS and protein modifications through RNS, primary radical species (•OH, O_2•−, CO_2•−, NO•), nonradical species (H_2O_2, HOCl, O_3, ONO_2•−, ONOCO_2•−, CO, N_2O_2, NO_2, 1O_2), and free radicals (•C, RS•, RSO•, RSSR•, R•, RO•, ROO•)[73]. In addition, proteins can also be modified by highly reactive aldehydes and ketones produced during ROS-mediated oxidation of lipids[74]. Hypoxia alters mitochondrial biogenesis and dynamics[76], and thus leads to mitochondrial dysfunction which is one of the key hallmarks of aging process and numerous age-related pathologies.[75] In summary, technological advances including proteomics, genomics, lipidomics, glycomics, transcriptomics etc. suggest that hypoxia alters mitochondrial biogenesis [76], causes DPMs [77], which leads to aging and age-related neurodegenerative diseases. It’s clear that mitochondria retain a central role in complex balance of cellular processes that contribute to aging, age-associated disorder. The key challenge remains to clinically translate this knowledge into model system and develop therapeutics.

2.2. Protein aggregation in hypoxia, aging and neurodegenerative disorders

Amyloidosis is a hallmark of neurodegenerative diseases where different amyloidogenic species cause a variety of neuropathic diseases such as Alzheimer’s disease (AD), Parkinson’s disease (PD), Huntington disease (HD), Prion diseases including Creutzfeldt-Jakob disease, Lewy body
The protein aggregates are often correlated with diseases, e.g., AD is characterized by aggregates of the tau protein and β-amyloid, PD by alpha-synuclein, and HD by aggregates of the huntingtin protein (Htt). On the contrary, the protein aggregates are not only pathogenic but also exert a protective effect on the stressed striatal neurons [79, 80]. According to Letiman et al. [79], the process of forming an aggregate is protective, isolating and segregating the problematic proteins. Arrasate et al. [81] found that the inclusion body (IB) formation reduced intracellular level of diffuse protein huntingtin (Htt) and prolonged survival which suggest IB formation protect neurons by decreasing the levels of toxic diffuse forms of Htt.

Similar findings of neuroprotective role of aggregated protein were noted in a mouse model of spinocerebellar ataxia in which the polyglutamine-rich forms of the ataxin-1 protein are expressed [82].

Proteins need to be folded into their stable three-dimensional structure for being functional. However, due to the folding errors, mutations, DPMs, failure of the proteostasis network, pathological conditions, and unfavorable conditions like elevated temperature, extreme pH, high pressure and agitation, protein get unfolded/misfolded and aggregation. Aberrant protein aggregation remains a common feature of neurodegenerative diseases, where mis-assembly of Aβ1-42 has been linked to AD. The relative increases in Aβ1–42 levels promote aggregation of Aβ into toxic species. However, aggregation-mediated Aβ1-42 toxicity was reduced in C. elegans when aging was slowed by decreased insulin/insulin growth factor–1–like signaling pathway (IIS) [83, 84]. Thus, modulation of IIS pathway could be a promising approach for the development of AD therapy. It’s important to note that in the nematode C. elegans, flies, mice and in human, IIS pathway regulates stress resistance, aging and determines the lifespan [83, 85].

Using C. elegans aging model, Kaufman et al., [86] found an aggregation of mitochondrial proteins and failure of mitochondrial protein homeostasis as a characteristic feature of hypoxia, while mitochondrial dysfunction and decreased protein turnover is linked to aging [87]. Recently, Adav et al. [88] evaluated the quantitative profile of Alzheimer disease (AD) brain mitochondrial proteins using both isobaric tags for relative and absolute quantitation (iTRAQ) and label-free quantitative technique and found altered mitochondrial proteins. These authors found destabilization of the junction between the membrane and matrix arm of mitochondria in AD. As evaluated by Kaufman and Crowder, [89] an increase in the detergent-soluble proteins, which are believed to be misfolded and aggregated, were the consequences of the hypoxic conditions. Thus, protein aggregation is common feature of aging and hypoxia and neurodegenerative diseases.

During hypoxia and aging, proteostasis pathway is compromised due to protein aggregation. The proteins of these aggregates have been found associated with neurodegenerative diseases, which indicates that the protein aggregation is an inherent part of aging and neurodegenerative diseases and consequences of hypoxia [90, 91]. Not only aggregation of protein but the protein or peptide-based drug (i.e., small peptides with 7, 6, 5, and 4 residues) is of significant concern [92]. In protein drugs, domain swapping technique is adopted to avoid aggregation, but swapping an aggregation-prone segment from amyloidogenic protein to a non-amyloidogenic homolog triggers amyloid formation [93]. Thus, the short aggregation-prone region of protein/peptide sequence and structural specificity can induce protein aggregation. However, the exact cause of protein aggregation, and the mechanistic link between the protein and tissue degeneration are not yet fully understood.

The protein misfolding and aggregation can induce cellular dysfunctions, cell death, and organelle failure that are the major pathological findings in post-ischemic brain, where impaired autophagy was thought to be the main cause of abnormal proteostasis and protein aggregation [94]. The most prominent structural motif of the functional protein in its native conformation is an alpha helix [95]. The beta sheet conformation also exists in many functional native proteins, but the transition from alpha-helix to beta-sheet is the main feature of amyloids and protein aggregation [95]. In physiological flow conditions, the shear flow induces protein aggregation and amyloid formation [96]. The loosely packed proteins in unfolded state expose their hydrophobic core which may interact with the cellular environment and undergo self-aggregation, while partially folded proteins act as precursors in the protein aggregation process [97].

While investigating the main cause of protein aggregation, the physicochemical analysis revealed that these aggregates differ in size, their proteomic composition and cellular location [98, 99]. Age-related and oxidative stress-induced protein aggregates were having compact conformation and protein carboxylation was a cause of compact aggregation [100]. Based on the solubility, aggregates were classified into soluble and insoluble, where soluble form can be more easily unfolded, while insoluble aggregates accumulate and impair cellular function [101]. When the proteomic composition of soluble and insoluble aggregates of human brain tissue was profiled, it revealed proteins like S100A9, ferritin, hemoglobin subunits, S100-A8, S100-B, collagens, mitochondrial creatine kinase (U-type), β-tubulin and laminin exclusively in compact aggregates and most importantly they were deamidated [23]. The
site-specific deamidation revealed deamidation of more than one residue of S100A9, which may introduce a negative charge altering Ca\(^{2+}\) binding capability and enhancing the capacity of the protein to form pathological aggregates in the brain. Using different aging models including C. elegans, murine bone marrow, and spleen cells, DPMs were found as a cause of protein aggregation [90, 91]. In conclusion, recent literature suggests that the ROS generated during hypoxia and aging, increases non-enzymatic DPMs such as carbonylation, oxidation, glycation, deamidation, citrullination, and lipoxidation. These protein modifications alter the protein charge and hydrophobicity, cause protein misfolding and aggregation, and their deposition in brain tissues could be key features of degenerative disease and aging. Despite plentiful evidences of the formation of aggregates and their effects, the exact mechanism(s) of protein aggregation and/or triggering nuclei remains unclear.

### 2.3. Hypoxia-induced DPMs and its role in aging and neurodegeneration

Hypoxia is known to induce broad changes in gene expression. In addition to affecting gene expression, hypoxia also alters the functions of proteins via posttranslational protein modifications. The clinical and experimental studies concluded that cerebrovascular disease and hypoxic-ischemic brain injury are the primary causes of cognitive impairment and dementia [29]. The progressive cycle of hypoxic-ischemic brain injury induces protein misfolding, aggregation and DPMs, leading to cognitive decline and dementia[15]. Further, the changes and posttranslational modification of proteins in response to chronic sustained and intermittent forms of hypoxia have been reviewed by Kumar and Klein [102]. Zanelli et al [103] tested the impact of hypoxia-induced protein nitration on the distribution of nitrotyrosine-like immunoreactivity in the hippocampus of the guinea pig fetus. Hypoxia-induced modifications were documented and reviewed including Slug SUMOylation [104], acetylation [105], glycosylation[106]. Apart from these protein modifications, this review article focuses on hypoxia-induced non-enzymatic post-translational degenerative protein modifications.

#### 2.3.1. Protein Carbonylation

At physiological concentration, ROS is a cellular requirement since it is involved in signaling pathways and in the regulation of numerous cellular activities like cytokine secretion, growth, differentiation and gene expressions, and defense [107]. A cellular balance between ROS production and clearance exits where antioxidant enzymes like Cu/Zn-superoxide dismutase (Cu/Zn-SOD, SOD1), manganese superoxide dismutase (Mn-SOD, SOD2), catalase, glutathione peroxidase (GPx), and glutathione reductase (GR) play a major role[54]. However, when ROS overproduction overcomes intrinsic antioxidant capacity, then the oxidative stress occurs. Thus, beyond the physiological concentration, ROS and secondary by-products of oxidative stress impose detrimental biological damage through protein oxidation. Cysteine and methionine are particularly prone to oxidative attack by ROS. The direct oxidative attack on lysine, arginine, proline or threonine leads to the formation of 2-pyrrolidone from proline, glutamic semialdehyde from arginine and proline, α-aminoacidic semialdehyde from lysine, and 2-amino-3-ketobutyric acid from threonine residues [108, 109]. The ketone carbonyls were also observed from histidine, proline, threonine, and tryptophan residue modification. The secondary reaction of cysteine, histidine, or lysine residues with reactive carbonyl compounds also leads to the formation of protein carbonyl derivatives, aldehydes, and ketones. Aldehydes produced from hydroperoxidation of lipid undergo Schiff-base formation with lysine residues through Michael addition and produces a lipid acyl group containing a free carbonyl that is capable of the secondary Schiff-base formation with an adjacent amine or cyclization. The degradation products of lipid oxidation can also bind mostly to cysteine, lysine, and histidine through Michael addition or Schiff base formation [110, 111].

The accumulation of carbonylated proteins with aging also corresponds to the alterations in hypoxic conditions, but the mechanism that causes accumulation of these proteins, though interesting, but not established. As reviewed by Solaini et al [112], hypoxia modulates mitochondrial oxidative metabolism. Upon exposure of yeast cells to hypoxia, abnormal protein carbonylation and protein tyrosine nitration were noted [113]. An elevated level of protein oxidation has been noted in hypoxic chronic obstructive pulmonary disease (COPD) patients compared with COPD patients who do not exhibit hypoxia [114]. Constant accumulation of oxidized proteins also takes place during aging and the possible causes of accumulation could be i) an increase in the rate of oxidizing species, ii) decrease in scavenging molecules, iii) impaired repair mechanism or iv) decreased clearance of oxidized proteins.

The clearance of oxidized protein is carried out by the proteasomal system. The partial inactivation of the proteasomal system results in the failure of oxidatively damaged protein repair or clearance of oxidized proteins during aging [115]. In mammalian cells, oxidized proteins are removed by the 20S proteasome [116]. The proteasome activity was inhibited or decreased in the episodes of repetitive and intermittent hypoxia in
Sprague-Dawley rat brain [117], cerebral ischemia-reperfusion [118] and in aging [119]. In addition to aging, there are several human diseases associated with carbonylated proteins such as AD, amyotrophic lateral sclerosis (ALS), acute respiratory distress syndrome, cataractogenesis, chronic lung disease, dementia with Lewy bodies, diabetes, ischemia-reperfusion, pre-eclampsia [120-123]. Thus, hypoxia induces ROS that leads to protein oxidation causing numerous diseases including neurodegenerative diseases and its shared feature of aging.

To evaluate the correlation between accumulated carbonylated proteins with age, Aguilaniu and colleagues [124] successfully retained higher carbonylated proteins in mother cells compared to daughter cells in the yeast model and confirmed that the accumulation of carbonylated proteins is a characteristic of aging processes. These authors successfully maintained carbonylated protein content in *Saccharomyces cerevisiae* but having such experimental studies in a higher organism such as an animal species is required. The support for the oxidative-stress hypothesis of aging, which postulates that aging is associated with the molecular damage caused by ROS came from the studies by Sohal et al., [125] who found a higher rate of ROS in the mitochondria of older rodents and flies than younger animals.

Figure 3. Possible routes of AGE formation (Adopted from Kikuchi et al [208]). AGEs get generated through decomposition of Amadori products, from glycolysis intermediates like glycolaldehyde, a Schiff base fragmentation product such as glycolaldehyde, fragmentation of Amadori products including methylglyoxal and 3-deoxyglucosone, and the autoxidation of glucose to glyoxal. (GO imidazolone, glyoxal imidazolone; MGO imidazolone, methylglyoxal imidazolone).  

2.3.2. Protein Glycation

Protein glycosylation is an enzyme-catalysed process that attaches glycans to proteins, lipids, and other organic molecules in a site-specific manner. While, protein glycation is a non-enzymatic reaction between reducing sugar and the free amino group of a target protein. Protein glycation is a consequence of the elevated production of
dicarbonyl species such as glyoxal, methylglyoxal, and 3-deoxyglucosone that non-enzymatically react with protein amino groups under conditions of hyperglycemia and oxidative stress. French scientist Louis Camille Maillard, observed a browning reaction when glycine heated with glucose, hence it is called Maillard reactions. It is not a single reaction, but proceed by a series of reactions [126] where the process begins with the reaction of the carbonyl group of glucose with the amino group of a protein to form a Schiff base, followed by conversion into stable Amadori product, which gets further modified to form advanced glycation end-products (AGEs). AGEs formation occurs through decomposition of Amadori products, glycolysis, Schiff base fragmentation product and many more routes as depicted in Fig. 3. Glycation site includes amino groups of protein, particularly, lysines, arginines and N-terminus of amino acids. In addition to protein glycation, DNA glycation also produces AGE, which acts as a potential carcinogen [127]. AGEs are also formed when food is processed at elevated temperatures, and thus, deep-fried, roasted, grilled food act as a source of AGEs; but their importance was largely ignored due to their poor absorption [128]. In mice model, consumption of AGE-rich diets has been linked to conditions such as atherosclerosis and kidney disease[129]. The food-derived AGEs induce protein cross-linking and intracellular oxidant stress similar to their endogenous counterparts [130]. The restriction of AGE-containing diet reduces oxidative stress, AGE accumulation and tissue damage resulting in extension of lifespan in mice [131]. The findings from animal and human studies suggest that chronic diseases and aging can be delayed by avoiding dietary AGEs. In other words, restriction of AGE-rich diet could be a novel therapeutic target to prevent age associated disorders.

AGEs induce damage to cells and extracellular matrix (ECM), thereby contributing to aging and age-related diseases. The buildup and accumulation of AGEs alters the structure and function of proteins, thus affecting several of the hallmarks of aging and responsible for the development of many age-related morbidities [132]. The mechanisms involved are: (i) accumulation of AGEs within the ECM, cross-linking between AGEs and ECM triggering a decrease in tissue elasticity, (ii) modification of proteins, which further results in a loss of the original cellular function, and (iii) interaction of AGEs with receptor for advanced glycation end-products (RAGE), which leads to activation of inflammatory signaling pathways, ROS generation, and apoptosis [133, 134]. The connection between aging, age-related disorders and AGEs remain difficult to unravel due to 1) wide-diverse sources for AGEs, 2) difficulties in detecting accumulation of AGEs, 3) technical limitation in detection and quantitation of AGEs, 4) a lack of models that reiterate the pathologies resulting from the accumulation of AGEs [132]. Moreover, there is no enzyme to remove glycated products from the human body. However, AGEs studies in short lifespans model organisms may help to find answers to their mechanistic role in aging.

Hypoxia condition escalates inside a solid tumor mass due to insufficient oxygen supply. Hypoxia-driven AGE accumulation and RAGE activation is well documented in published literature [135]. Hypoxic conditions elevates expression of RAGE [136] and contribute to protein modifications and, thus influence aging and neurodegeneration processes. Its well-known fact that the cells of hypoxic tumor adopt anaerobic glycolytic process instead of mitochondrial aerobic respiration and these hypoxic cells induce accumulation of di-carbonyls, a precursor of AGE [137]. The methylglyoxal (MG), an intermediate compound generated during glycolysis also act as a precursor molecule for AGE[138]. According to Chang et al [139], a rapid generation of AGE after hypoxia act as a precursor in endothelial cells that further activates RAGE-mediated signaling. These oxygen deficient cells actively participate in tumor growth and metastasis through activating several signaling events[140]. The contribution of AGE in enhancing proliferation of breast cancer progression[141], a key role in prostate cancer[142] has been shown. The mechanism of hypoxia-driven glycation and role of protein glycation in various cancers have been reviewed [143]. Detailed understanding of the AGE mediated cancer onset could open avenues in cancer therapeutics.

Apart from cancer, the roles of AGEs have been linked to various diseases including diabetes, cardiovascular disease and neurodegenerative disorders [144]. RAGE-AGE interaction mediates myocardial injury after ischemia attack[145]. The hyperglycemia-induced AGE activation followed by retinal neovascularization was studied by Shin et al. [146]. Glycation process is a toxic cascade reaction in which different oxidative products with high oxidative potential than the parent compound are produced, and their quantity determines its destructiveness [147]. In fact, the toxicity of glycation results in loss of protein function through cross-linking, aggregation and deposition, and production of reactive species, which are an analogy of hypoxia.

### 2.3.3. Protein Carbamylation

Carbamylation is an irreversible, non-enzymatic spontaneous reaction of primary amino groups or a free sulfhydryl group of proteins with an isocyanate as reported in the 1960s by Stark et al., [148]. The spontaneous decomposition of urea into ammonium and cyanate generate isocyanic acid, which is a reactive
species. The reactive cyanate generated through thiocyanate metabolism and neutrophil-derived myeloperoxidase catalyzes the oxidation of thiocyanate at sites of inflammation and atherosclerotic plaque. Urea is present abundantly in the human body and can decompose spontaneously forming cyanic acid and cytanate, where cyanic acid is in rapid equilibrium with isocyanic acid [148]. When carbamylation occurs on lysine residue then it generates ε-carbamyl-lysine, also called as “homocitrulline” and it remains challenging to differentiate it from citrulline since just with one additional methylene group homocitrulline residue generated by carbamylation becomes structurally similar to the citrulline residue formed by pyrroldid arginine deiminase activity [149]. Again, a mass shift of +43 Da (carbamylation) needs to be distinguished from +42 Da (trimethylation or acetylation) to avoid artificial identification of specific modification. Carbamylation can also occur at the guandine moiety of arginine and a reduced thiol of cysteine.

Biomolecules such as albumin, low-density lipoprotein (LDL), collagen, and many more, undergo carbamylation in both pathological and physiological conditions that alters these biomolecules structurally and functionally. The accumulation of carbamylated proteins is considered as a hallmark of aging. The carbamylation of crystallin is well studied [150]. The carbamylated erythropoietin (CEPO) is attracting widespread interest due to its neuroprotective effects without erythropoiesis [151]. While other carbamylated proteins namely, carbamylated-haemoglobin and carbamylated-LDL have been implicated in hypoxia and atherosclerosis, respectively [152]. The carbamylation of the erythropoietin (EPO) causes a drastic decrease in its signaling ability and loss in erythropoietic activity. The carbamylation of the erythropoietin is linked to hypoxia in patients with end-stage renal disease [153]. It’s important to mention that in hypoxia, hypoxia-inducible factor 1α (HIF-1α) activates erythropoietin. The distinct receptors for EPO and carbamylated EPO exist. According to Brines et al [154] homodimeric EPO receptor binds only to EPO and regulates its erythropoietic function, while the heterodimeric receptor for carbamylated EPO regulates tissue protection. Thus, carbamylation of EPO results in decreased erythropoiesis and further hypoxia due to the altered synthesis of RBCs and HBs.

### 2.3.4. Protein deamidation

Deamidation is a non-enzymatic post-translational modification in which amide functional group in the side chain of amino acid asparagine (Asn) is converted into an aspartic acid or iso-aspartic acid, and glutamine (Gln) into glutamic acid. At neutral pH, deamidation introduces a negative charge at the reaction site and lead to biological and structural alterations in peptide and protein. At physiological pH, deamidation is a two-step processes wherein a first step-the peptide bond nitrogen of the N+1 amino acid attacks the carbonyl carbon of the asparagine or aspartate side chain leading to the formation of a five-membered ring structure named as a succinimide or cyclic imide; and in a second step-the succinimide is rapidly hydrolyzed at either the alpha or beta carbonyl group to yield iso-aspartate (beta-aspartate) and aspartate in a ratio of approximately 3:1 [17, 155]. At low pH i.e. pH≤2, direct hydrolysis of the side chain amide generates aspartase as a sole product. At alkaline pH, an elevated rate of succinimide formation has been documented [156] and presumed that the elevated rate is due to greater deprotonation of the peptide bond nitrogen.

Under physiological conditions, L-aspartyl (L-Asp) and L-asparaginyl residues in proteins undergo spontaneously nonenzymatic deamidation leading to the formation of an abnormal isoaspartyl residue. The enzyme protein-L-isoaspartate (D-aspartate) O-methyltransferase (PIMT) recognizes and repairs the abnormal L-isoaspartyl residues in proteins. Since PIMT is an enzyme with methyltransferase activity, Yan et al [157] believe the possibility of regulation of mammalian sterile 20-like kinase (Mst1) activity by PIMT through altering either deamidation or methylation of Mst1. Mst1 regulates apoptosis and tumor suppression in mammals and play a key role in heart disease since its activation causes cardiomyocyte apoptosis and dilated cardiomyopathy. Under hypoxic conditions and UV radiations, Mst1 increases protein deamidation [158]. Thus, it will be fascinating to investigate the role of deamidated Mst1 and impact of PIMT under hypoxic conditions. The possibility of interaction of PIMT with Mst1 and conformational changes of Mst1 can’t be ignored. These studies will highlight the critical role of deamidated Mst1 and PIMT. Further it will shed light on the impact of conformational changes on the formation of the Mst1/Hippo signalling complex with other proteins, such as Rassf1, hWW45, and Lats, which have been shown to play essential roles in the regulation of cardiomyocyte apoptosis and heart failure [159, 160].

Asparagine deamidation was elevated in hypoxic RBCs relative to normoxia and such phenomena are described as a function of human aging, and RBC senescence [161]. Hypoxic stress also causes chromatin modification including a pool of histone modifications which plays a role in gene regulatory switches [162]. Recent studies implicated the role of protein deamidation in regulating signal transduction in innate immune responses which were reviewed by Zhao et al., [163].
Despite the accumulating evidences, research on the direct impact of DMPs in disease pathology and progression remain limited and still needs further exploration. Proteomics technology remains a state-of-art technology for quantifying DPMs in order to understand their associated biological functions. However, to study the functional alteration that DMPs causes requires a larger toolbox with combination of MS and other more diverse methodology that can highlight the impact of DPMs and help to go beyond DPMs, protein identification and quantification. Future studies are required to focus on the specific DPMs as a therapeutic target and to develop and validate drug targets and effectiveness.

3. Role of degenerative protein modifications in aging and age-associated disorders

More than 40 human neurodegenerative diseases have been characterized by proteins misfolding, aggregation, and deposition, including the aggregation of α-synuclein in PD [164], tau or beta-amyloid in AD [165, 166], and huntingtin in HD [167] are the examples. Thus, the etiologies of different neurodegenerative disease differ widely but the pathological signature remains protein aggregation. In neurodegenerative diseases including AD, PD, HD, amyotrophic lateral sclerosis (ALS) and prion diseases, protein aggregation remains as a common feature. It’s been believed that the DMP may facilitate protein aggregation. Oxidative modification of protein α-synuclein via dopamine adducts facilitated protein aggregation [168]. Hence, we reviewed the role of DMPs in aging and age-associated disorders as below

3.1. Protein Carbonylation

Several theories of dementia and AD are proposed since they were reported a century ago. The initial hypothesis proposed that ischemic cerebral vascular disease or strokes are the main cause of age-related neurodegenerative diseases [169]. The presence of hyperphosphorylated microtubule-associated tau protein, intracellular neurofibrillary tangles and extracellular amyloid deposits derived from amyloid precursor proteins in the cerebral cortex and hippocampus are the major hallmarks of neurodegenerative diseases. The injection of Aβ (1-42) into the nucleus basalis of the rat, showed a congophilic deposit and microglial and astrocyte activation which steered a strong inflammatory reaction and induced nitric oxide synthase expression [170]. Aβ is found to be neurotoxic and its neurotoxicity is mediated through its potential to induce free radical mediated oxidative stress, including protein oxidation and lipid peroxidation [171, 172]. In a well-established HD mouse model (R6/2 transgenic mice), α-enolase, γ-enolase, creatine kinase, aconitase, voltage-dependent anion channel 1, and Hsp90 were identified as the main carbonylated proteins that potentially contribute to the impairment of energy metabolism and the pathogenesis of HD [173]. In human brain striatum, Sorolla et al [174] found increased carbonyl levels of glial fibrillary acidic protein, aconitase, γ-enolase (neuron-specific enolase), and creatine kinase B (the brain-specific isoform) in HD patients. Similarly, oxidative and nitrative protein modifications in PD have been documented [175].

Under insufficient oxygen, brain induces oxidative stress, which is indexed by an elevated level of lipid peroxidation, protein oxidation, and neuronal dysfunction or death [121]. Protein carbonyls are the results of a direct free radical attack on the protein backbone or from the products of glycation, glycoxidation, and lipid peroxidation reactions with proteins [176]. Oxidative protein modifications diminish specific protein functions and causes cell death [176, 177]. Mis-regulation of programmed cell death is implicated in aging and age-related neurodegenerative diseases. Protein oxidation is indexed by the presence of protein carbonyls. Protein carbonylation in neurofibrillary tangles, neuronal cell bodies, dendrites and glial nuclei in hippocampal sections have been linked to AD [178]. Further, protein carbonylation in synaptic and non-synaptic mitochondria in the frontal cortex have also been correlated with the AD [179]. Smith et al [180] noted significantly elevated levels of protein oxidation in AD frontal lobe. While Lyras et al [181] noted increased protein carbonyls in the frontal, temporal and occipital lobes and hippocampus in AD.

Boyd-Kimball and colleague [182] identified significantly oxidized numerous proteins which lost their functions due to conformational changes. An altered structural conformation induced protein aggregation, neurofibrillary tangles pathology, loss of synapse and neuronal communication which are associated with the AD. Restated, protein oxidation plays a central role in aging, in the pathogenesis of age-related neurodegenerative diseases, where the possible link could be hypoxia-induced ROS that causes protein oxidation leading to neurodegenerative diseases. Further, an increase in lipid peroxidation, protein oxidation, and accumulation of oxidized biomolecules in neurons exposed to hypoxia may be interpreted as hypoxia accelerate the aging process. In short, hypoxia-induced protein oxidation causes protein aggregation, alter structural conformation and leads to loss of protein function, which is a common feature of aging and age-related disorders.

3.2. Protein Glycation and its role in aging and age-associated diseases
It’s well known that the proteins of the extracellular matrix and vascular basement membrane are long-lived. The stiffness of the tissues which are composed of long-lived proteins including skeletal muscle, tendons, joints, bone, heart, arteries, lung, skin, and lens have been correlated with aging [183]. However, these long-lived proteins are susceptible to glycation [184]. The collagen is the most abundant structural protein in skin, tendon, bone, articular cartilage and vascular system; which provides functional properties to different tissues such as renal basement membrane, cardiovascular and retinal capillaries. Hence, the glycation of fibrous and non-fibrous collagen and their correlation with aging remain the subject of research. The glycation of collagen, cross-link between collagen and AGEs in the aorta, carotid and conduit arteries enhances their stiffness with age [185]. Thus, glycation is directly associated with aging. AGE-modified collagen reduces cell migration by impairing mechanism of cell adhesion and proteolytic degradation of collagen by matrix metalloproteinase, [186] while in another study, AGE-modified collagen impaired the potential of chondrocytes of the articular cartilage [187]. Thus, glycation promote collagen cross-linking, increase in tissue stiffness, reduced skin elasticity, modify matrix proteins, and impair several repair processes contributing to vessel rigidity and hypertension, which are an integral part of the aging process. Thus, AGEs could be good therapeutic target to slow down aging process.

Protein glycation is associated with several neurodegenerative disorders, including AD, PD and HD. AGEs are reported in senile plaques, NFTs, and cerebral amyloid angiopathy from AD brains [188]. In established yeast, mammalian cell and fly models of HD, protein glycation impairs Htt clearance thereby promoting its intracellular accumulation, enhancing its aggregation and pathogenicity [189]. Glycation exacerbates the accumulation, aggregation and toxicity of Aβ and α-synuclein [190]. The protein like tau, Aβ, α-synuclein, and prions were found to be glycated and the extent of glycation is correlated with the pathologies of the patients [191]. AGEs modification triggers the abnormal deposition and accumulation of these modified proteins, and hence it can be hypothesized that glycation contribute to the development of neurodegenerative disease. Thus, drugs targeting the glycation precursors, or promoting the clearance of glycated proteins may be beneficial for AD, PD and HD patients. The targeted studies on characterization of the molecular mechanisms responsible for AGEs mediated neurotoxicity are required to discover the possible link and novel strategies for the prevention and development of therapeutics to treat neurodegenerative diseases.

Glycation of DNA alters the structure of DNA macromolecule, and it leads to depurination, strand breaks, and mutations such as insertions, deletions, and transposition [192]. Glycation of DNA gives rise to characteristic nucleotide adduct, some of which were found to increase in oxidative stress. Methylglyoxal (MG) is extremely reactive glycating agent that reacts with free amino group of nucleic acids resulting in the formation of DNA–AGEs. MG reacts with guanine residues in DNA to form a tricyclic compound [193] and its mutagenicity at G:C base pairs has been reported [194]. Again, AGE product, carboxymethyllysine, accumulates in nuclear proteins like histone causing extensive DNA strand cleavage. Thus, DNA-AGEs leads to the loss of genomic integrity during aging and age-related complications. However, recently Richarme and colleques [195] found that parkinsonism-associated protein DJ-1 and its bacterial homologs Hsp31, YhbO, and YajL could repair methylglyoxal- and glyoxal-glycated nucleotides and nucleic acids.

Receptor for advanced glycation end products (RAGE) when binds to AGE, it initiates a further signaling mechanism [196]. RAGE also acts as a receptor for Damage-Associated Molecular Pattern (DAMP) molecule, high mobility group box 1 (HMGB1), the prototypical DAMP, and S100 proteins [196, 197]. Hypoxic conditions elevates expression of RAGE [136] and contribute to protein modifications and, thus impact aging and neurodegeneration processes. AGE-RAGE complex initiates aberrant signaling pathways initiating the onset of diseases such as diabetes, AD, atherosclerosis, heart failure, and various cancers like oral, breast, gastric, colorectal, pancreatic, intestinal and others as reviewed by Khan et al., [198]. AGE-RAGE complex provokes oxidative stress that further induces proliferative, inflammatory, thrombotic and fibrotic reactions [199]. Its well-known fact that the cells of hypoxic tumor adopt anaerobic glycolytic process instead of mitochondrial aerobic respiration and these hypoxic cells induce accumulation of di-carbonyls, a precursor of AGE [137].

Glycation process is a toxic cascade reaction in which different oxidative products with high oxidative potential than the parent compound are produced, and their quantity determines its destructiveness [147]. In fact, the toxicity of glycation results in loss of protein function through cross-linking, aggregation and deposition, and production of reactive species, which are an analogy of hypoxia. The cross-link between adjacent protein and rearrangement of Amodori product remain the cause of protein aggregation and AGEs production, and the consequence of the accumulation of AGEs is plaques and tangles [200]. Accumulated AGEs have also been discovered in senile plaques, primitive plaques, classic plaques and some glial cells of AD brain [201]. It was found that the expression of amyloid precursor protein was elevated in vitro and in...
 vivo by AGEs, indicating AGEs plays a role in AD pathogenesis [191, 202]. The dimeric form of Apolipoprotein E (ApoE) binds to AGEs and promote the aggregate formation in AD brain by binding to plaque components [203], which may explain how ApoE contributes to increased risk of the AD. In rats, tau protein was glycated at its tubulin binding site [204]. This glycated tau protein induced oxidative stress and hyperphosphorylation of tau, which further impairs memory via RAGE-mediated GSK-3 activation. Thus, AGEs play a major role in AD pathogenesis, amyloid formation, protein aggregation, and participates in NFTs formation. AGEs have been intensively studied in diabetes mellitus [205, 206], cardiac dysfunctions [207], visual disorder, nephropathy, vascular disorders [208] and diabetic atherosclerosis.

![Figure 4](image-url)

**Figure 4. Separation of Asn and Gln deamidation peptides from trypsin-digested human brain tissue using LERIC-MS/MS.** (Adopted from Serra et al.[233]) A) Extracted ion chromatograms of peptide N#GFDQCDYGWLSASVR showing separated triad of Asn deamidated peptides B) Extracted ion chromatograms of VDKGVPLAGTN#GETTTQGLDGLSER peptide showing separation of carbon-13 peak of nondeamidated peptide (Asn-C13), Asp (asparyl isomer) and isoaspartyl isomer (isoAsn). C) Extracted ion chromatograms of GVVPLAGTNGETTTQGLDGLSER nondeamidated peptide. D) Extracted ion chromatograms of two deamidated proteoforms with asn and Gln deamidated residues in peptide GVVPLAGTN#GETTTQGLDGLSER and GVVPLAGTNGETTTQ#GLDGLSER where Gln-Asn-C13 is a carbon-13 peak of the nondeamidated peptide; α-Glu is α-glutamyl isomer; Asp is Asp aspartyl isomer; γ-Glu is γ-glutamyl isomer; and isoAsp reamin isoaspartyl isomer. #indicates site of modification.

### 3.3 Protein Carbamylation

Carbamylation of the protein causes changes in protein structure and charge, which alter the molecular activity of enzymes, cofactors, hormones, low-density lipoproteins, antibodies, receptors, and transport proteins [209]. Carbamylation induces a conformational change in crystallins, it alters collagen type I structure [210].
diminish the activity of insulin and erythropoietin [153],
and also induces atherosclerotic plaques formation [211].
The carbamylation of a few residues per α-chain of the protein destabilizes its structure leading to decreased thermal stability, fibrillogenesis impairment and altered sensitivity to proteases [210, 212]. Earlier, the quantitative increase in protein carbamylation has been correlated with kidney disease, atherosclerosis, or coronary artery disease [213, 214].

Using murine, bovine, and human species from younger to older ages, an increase in the carbamylated proteins in the skin with time regardless of lifespan have been observed [215]. The type I collagen and elastin of skin was highly carbamylated in older subjects than in younger, and authors projected protein carbamylation as a hallmark of the aging process [215]. Thus, the discovery of carbamylated skin protein has established a link between carbamylation and aging. The carbamylation of brain protein by cyanate [216] caused a decline in learning ability [217], which associate protein carbamylation with neurodegeneration. The carbamylated low-density lipoprotein (cLDL) induced endothelial cell injury, enhanced expression of cell adhesion molecules, and proliferation of vascular smooth muscle cell, which are identical with atherosclerosis [218, 219]. From inflammation to aging, many diseases have been correlated with protein carbamylation and their pathology [215, 220, 221].

3.3. Protein deamidation and its role in aging and age-associated diseases

According to comprehensive research by Robinson and colleagues [222-226], though, each deamidation site has programmed rate of deamidation, but it also depends on the primary sequence, neighboring amino acids, three-dimensional structure, pH, temperature, ionic strength, and other solution properties. The deamidated Asn and Gln serves as molecular clocks which time biological processes including protein turnover, organismic development, and aging [17, 19, 227, 228]. Deamidation process occurs in vivo but it also occurs spontaneously in vitro during proteomic sample preparation and introduces artifact. To overcome artefactual identification, Sze and colleagues [229-231] proposed an improved protocol at acidic pH to minimize artefactual non-enzymatic deamidation during sample preparation. Notably, Asn deamidation proceeds via formation of succinimide ring intermediate which gets hydrolyzed to n-Asp and isoAsp [155]. It’s difficult to separate, distinguish and identify n-Asp and isoAsp deamidation products due to their identical mass and charge. The techniques like 2-dimensional reverse phase liquid chromatography coupled with electrostatic repulsion-hydrophilic interaction chromatography (RPLC-ERLIC)-coupled mass spectrometry [232] and long-length electrostatic repulsion-hydrophilic interaction chromatography (LERLIC) tandem mass spectrometry have been optimized for distinguishable separation as well as identification and quantification of deamidation products [233]. The separation of Asn and Gln deamidation peptides by LERIC-MS/MS is shown in Fig 4. An altered brain protein function and aggregation are characteristics of neurodegenerative diseases, however, which specific protein modification recruit plaque formation is not conclusive. But Sze and colleagues proposed that the protein deamidation alters the charge of the protein, influence protein folding, structure, and stability; and initiate protein aggregation in neurodegenerative diseases like dementia [12, 13, 21, 23].

Recently, the critical role of isoDGR motif has also been evaluated by Dutta and Sze et al., [234], the deamidation of extracellular matrix (ECM) proteins in vascular bed and atherosclerotic plaque promotes atherosclerosis by enhancing monocyte-macrophage binding to the ECM proteins in the blood vessel.

The effects of age-dependent protein deamidation were extensively studied in eye lens proteins as a model system due to their exceedingly slow turnover during the patient lifetime. To retain life-long transparency in the absence of protein turnover, eye lens crystalline maintains its longevity by retaining long-term native structure, stability and refolding of non-native proteins by chaperones. However, the disruptive protein modifications, particularly deamidation and accumulation of deamidated proteins remain the main concern. Despite the features that cause age-dependent changes in lens crystallines are still not fully understood, it was found that the aggregate size of crystalline increased with age and was composed of deamidated alpha- and beta- crystalline proteins [235]. Age-related cataract has been characterized by the protein aggregation and DPMs, many reports suggests that lens protein aggregation and deterioration are correlated to protein deamidation and oxidation [236-239]. Age-dependent increase in deamidated histone H1 in the brain of mice and rats over 20-days old to 300-days old [240] also supports the role of deamidation in aging. The proportion of deamidated H1 was 8-fold higher in mice of 10 months when compared with 20 days old animals. It is proposed that the deamidation of linker histone would add negative charge to nucleosome that could influence the binding of DNA, chromatin structure and stability.

4. Future perspectives

The main goal of biomedical research is to understand physiological and pathological processes to develop
preventive measure and therapeutics to maintain good health throughout life and improve human healthspan. However, maintaining good health become very challenging in the modernized demographic scenario, partly, due to increased oxidative stress that induces various pathologies including ischemic heart disease, cancer, stroke, and diabetes; and neurological disorders. These chronic diseases are the cause of physiological and structural deterioration. With advancing age, several cellular, structural and functional changes occur, where, alterations arisen at pre-mature period are beneficial, but those that occur during the post-maturational period of life are mostly deteriorative, except a few. These deteriorative cellular damages keep accumulating, remain vulnerable to challenges, and lead to a decline in cognitive functions. Therefore, this review attempted to summarize up-to-date information on different aspects of aging, the effects of hypoxia, DPMs during aging and age-associated neurodegenerative diseases. An attempt has been made to find the link between DPMs and aging, as well as the impact of DPMs on age-related pathologies. Hypoxia and disrupted oxygen supply generate oxidative stress, enhance ROS, cellular apoptosis, abnormal mitochondrial metabolism, influence lipid peroxidation, neurodegeneration and memory impairment, and most importantly increased DPMs. The deposition and accumulation of these modified proteins act as pathogenic factors. These protein damages may not be repairable due to defective repair mechanism. This leads to the aggregation of proteins and their accumulation with age which has been linked to aging and age-associated disorders.

Due to the role and relationship between Aβ in protein aggregation and AD, the so-called "secretases" that produces Aβ have been targeted for development of therapeutics. A new generation of small molecule β-secretase (BACE) inhibitors are being evaluated in clinical trials [241]. The difficulties in direct γ-secretase inhibition, directed the focus towards modifying γ-secretase substrate specificity[242]. Emerging strategies to avoid or clear protein aggregates includes protein reduction using as β-secretase inhibitors/modulators[241], γ-secretase modulators [242], chemotherapy [243], proteasome inhibitors [243], liver transplantation mediated gene therapy [244] etc. Therapeutics used to modulate amyloid deposition and age-associated disorders are reviewed by Yiannopoulou and Papageorgiou [245]. The preclinical studies in mice highlights possible novel immunotherapeutic strategies like DNA epitope vaccine, antibodies against the β-secretase cleavage site of the APP and mucosal vaccination as an effective AD therapy. Removal of hyperphosphorylated tau by immunotherapy inhibits retrograde neurodegeneration [246]. Cholinesterase inhibitors like donepezil (Pfizer, New York, NY, USA), rivastigmine (Novartis, Basel, Switzerland) and galantamine (Janssen, Beerse, Belgium) [247] are being used to delay the decline in cognitive function. Antidepressant drugs like mirtazapine, venlafaxine and duloxetine, combined selective noradrenaline and serotonin inhibitors (SNRIs), and bupropion are being used to treat depression in dementia. However, a further research that focuses on the root cause of the protein aggregation, factors that trigger and sustain aggregate formation is needed.

Neurodegenerative disorders are global public health priorities and require urgent action to address at healthcare level. Elucidating the molecular mechanism and investigating main initiating factors will be a key in developing therapeutic interventions. Administration of N-acetyl cysteine to animals exposed to hypoxia exhibited a decrease in the free radical generation and improvement in memory functions [248], while acetyl-L-Carnitine treatment improved the cognitive capabilities [249]. The implications of such compounds as anti-aging and anti-senescence needs further detailed investigations. AGE-RAGE mediated pathologies are gaining significant importance but the mechanism of different AGE-RAGE mediated signaling pathways need to be explored using proteomic approaches, and we believe such studies will provide new insights into aging and AGE-mediated pathogenesis. Further, such studies in an animal model may discover a direct link between protein modification and their role in aging and age-associated diseases. AGEs are involved in aging, diabetes, cancer, as well as neurodegenerative diseases. Therefore, designing selective inhibitors for AGE could help to treat several diseases. To develop such therapies, it’s essential to correlate Hif1α signaling pathway in hypoxia and other diseases to find out a specific target. The structural modification and aggregation of the protein is a core issue in disease development and need focused detailed research. An advanced mass spectrometry-based proteomic technology can be used to profile global effects since its highly sensitive and powerful tool for global proteome profiling and PTMs/DPMs characterization. Aging and age-associated neurodegenerative diseases are multifactorial, hence adopting unbiased, global, the discovery-driven proteomic approach can emphasize global quantitative proteome with novel protein modifications. Based on this discovery data as a steppingstone, targeted proteomic approach on the DPMs might provide promising insight and assist to elucidate pathologies and develop novel therapeutics.
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Conflicts of interest

The authors declare no conflict of interest.

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