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Mitochondrial Lon protease in human disease and aging: Including an etiologic classification of Lon-related diseases and disorders

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Abstract

The Mitochondrial Lon protease, also called LonP1 is a product of the nuclear gene LONP1. Lon is a major regulator of mitochondrial metabolism and response to free radical damage, as well as an essential factor for the maintenance and repair of mitochondrial DNA. Lon is an ATPstimulated protease that cycles between being bound (at the inner surface of the inner mitochondrial membrane) to the mitochondrial genome, and being released into the mitochondrial matrix where it can degrade matrix proteins. At least three different roles or functions have been ascribed to Lon: 1) Proteolytic digestion of oxidized proteins and the turnover of specific essential mitochondrial enzymes such as aconitase, TFAM, and StAR; 2) Mitochondrial (mt)DNA-binding protein, involved in mtDNA replication and mitogenesis; and 3) Protein chaperone, interacting with the Hsp60-mtHsp70 complex. LONP1 orthologs have been studied in bacteria, yeast, flies, worms, and mammals, evincing the widespread importance of the gene, as well as its remarkable evolutionary conservation. In recent years, we have witnessed a significant increase in knowledge regarding Lon's involvement in physiological functions, as well as in an expanding array of human disorders, including cancer, neurodegeneration, heart disease, and stroke. In addition, Lon appears to have a significant role in the aging process. A number of mitochondrial diseases have now been identified whose mechanisms involve various degrees of Lon dysfunction. In this paper we review current knowledge of Lon's function, under normal conditions, and we propose a new classification of human diseases characterized by a either over-expression or decline or loss of function of Lon. Lon has also been implicated in human aging, and we review the data currently available as well as speculating about possible interactions of aging and disease. Finally, we also discuss Lon as potential therapeutic target in human disease.

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Keywords

Lon; Proteolysis; Mitochondria; Mitochondrial DNA; Aging; Cancer; Mitochondrial diseases; Cancer; Neurodegeneration

1. Introduction

Mitochondria are essential components of cellular metabolism, and rapid changes in the mitochondrial protein pool are required for adaptation to a variety of conditions, from increased demand for oxidative energy production to hypoxic adaptation. Mitochondrial homeostasis is crucial for cell survival, and mitochondrial dysfunction leads to apoptotic cell death in normal cells and to aberrant adaptation and selection of hypoxic phenotypes in pathologic conditions such as cancer.

The turnover of mitochondrial proteins was once believed to be a function of lysosomes which can carry out autophagy of the whole organelle [1]. Mitochondrial autophagy varies widely in different metabolic conditions, with an estimated median half-life for liver mitochondria of 1.83 days in control conditions, which can decrease to 1.16 days following 3 months of caloric restriction [2]. Lysosomal autophagy, however, cannot explain wide differences in the individual turnover rates of various mitochondrial proteins; for example, the two isoforms of mitochondrial acetyl-CoA acetyl-transferase (acetoacetyl-CoA thiolase, EC 2.3.1.9) A1 and A2 have half-lives of 24.5 h and 7.2 h respectively [3]. Similarly, the uncoupling proteins (UCPs) – a very important subset of the mitochondrial solute carrier family - have very different half-lives [4]. UCP2 has a remarkably short half-life of ~1 h in cells [5], while UCP1 has a much longer half-life, of ~1–4 days [6] and UCP3 has a short half-life of 0.5–4 h [4].

In order to maintain an active, balanced production of energy, mitochondria need to expeditiously remove and replace any modified or damaged proteins. This is absolutely crucial, due to the fact that oxidative phosphorylation (OX-PHOS) constantly induces oxidative damage to the respiratory subunits as well as to other essential enzymes involved in mitochondrial metabolism [7]. Mitochondrial proteases demonstrate substrate preferences, which suggest possible recognition based on protein modifications [100]. As the vast majority of mitochondrial proteins are encoded in the nucleus, an imbalance between the production of mitochondrial subunits – which is inducible – and mitochondrial degradation could lead to serious mitochondrial disorders of homeostasis and (finally) to cellular disease and death [8,9].

These findings suggest that mitochondria possess a complex proteolytic pathway which modulates the selective and precisely-timed degradation of internal proteins [10]. At least five different proteases are responsible for targeted protein quality control (UPRmt, the mitochondrial unfolded protein response): most prominently represented being the Lon protease and the other AAA + family proteases (ATPases associated with a variety of cellular activities) [11,116]. The mitochondrial ability to respond to organelle- specific stressors is essential for the degradation of unfolded proteins in the mitochondrial matrix [117].

The human Lon protease is highly conserved through multiple levels of evolution, and is the homolog of the bacterial La [12] and the yeast PIM1/Lon [13]. Lon expression is essential for survival in mammals, and homozygous deletion of *LONP1* causes early embryonic lethality [14]. In recent years, an explosion in our understanding of Lon function has occurred. As such, Lon dysfunctions/mutations have now been associated with multiple physiologic and pathologic conditions in humans, from aging to cancer and neurodegeneration [15]. This review attempts to provide a systematic approach to Lon involvement in health, disease, and aging, and also to provide an etiologic classification of Lon-related disorders.

2. Lon discovery, structure, and regulation

2.1. Lon discovery

The presence of an ATP stimulated proteolytic activity has been described in the matrix of mitochondria from a variety of mammalian organs and systems since 1982, including bovine adrenal cortex [16–18], rat liver [19,20] and human heart, brain, liver [21] and placenta [22].

The mitochondrial matrix protease responsible for this activity was initially purified by two different groups as a 600–650 kDa hexameric homoprotein complex (subunit mass 106 kDa) [23–25]. The human enzyme has a subunit size of 100 kDa in the mature form [26] and is widely expressed in multiple human tissues – with the highest levels in the most metabolically active organs – heart, brain, liver and skeletal muscle [21]. The nuclear gene encoding mitochondrial Lon is *LONP1* (*PRSS15*), situated on chromosome 19. There is also a peroxisomal form of the Lon protease, called LonP2; despite apparent similarities to mitochondrial LonP1 in proteolytic functions, the peroxisomal LonP2 is encoded by a completely different nuclear gene, and is regulated independently of LonP1 [27].

2.2. Lon protein structure

Lon's protein sequence was initially solved for the human protein. The cloned nuclear gene encoded a protein of 963 amino acids with a calculated molecular mass of 106 kDa [21], which is then processed to a mature 100 kDa mitochondrial matrix protein kDa [21]. There are three distinctive, highly conserved structural sequences that have been evolutionarily conserved: 1) a mitochondrial matrix targeting sequence [21], 2) a serine protease domain, [21] and 3) a Walker-type ATP binding motif (ATPase functional domain, AAA⁺ module) that is involved both with nucleotide binding and ATP hydrolysis. [28]

The complete crystal structure of Lon was resolved in *Thermococcus onnurineus* [29] and in the *Escherichia coli* [30] bacterium. The published 2.0-Å resolution crystal structure of *Thermococcus onnurineus* NA1 Lon (TonLon) reveals a three-tiered hexagonal cylinder with a large, central, unfolding and degradation chamber [29]. However, *E. coli* Lon analysis by size-exclusion chromatography (SEC) and multiangle laser light scattering (MALS) identified both hexamers (calculated MR 525 kDa) and dodecamers (calculated MR 1050 kDa) [29]. Formation of the dodecamer inhibits the proteolysis of large but not small protein substrates, suggesting that the repertoire of Lon substrates depends on its state of assembly [29]. The resolved structure of human mitochondrial Lon proteolytic domain is resembles

those of the previously determined Lon proteolytic domains from *Escherichia coli*, with six protomers in the asymmetric unit. The active site contains a 3_{10} helix attached to the N-terminal end of α -helix 2, which leads to the insertion of Asp852 into the active site [31]. The full –sequence human Lon protease is yet to be crystalized – but human Lon displays significant homology the bacterial protease La (*lon* gene) and the yeast form of Lon, which is called Pim-1 (*PIM1* gene) [13,32]. As such, the homology model of human Lon suggests a hexameric complex with an open-ring arrangement similar to that of yeast (Pim-1) Lon [33].

2.3. Lon regulation

The mitochondrial Lon protease is a one of the most important cellular stress-responsive proteins [34]. Acute stressors, such as heat shock [34], serum starvation, and oxidative stress [35,36] lead to Lon upregulation. However, regulation of Lon is biphasic, and chronic and severe stress conditions – such as aging [15], extensive hypoxia [35] and prolonged oxidative stress [37] cause Lon down-regulation, suggesting that Lon is a significant factor in aging and degenerative diseases.

The *LONP1* promoter region –623/+1 contains a Nuclear Respiratory Factor 2 (NRF-2) consensus binding site and is essential for response to reactive oxygen species [36]. Additional proximal NRF-2/GABP consensus sites were recently found to be essential for growth-related transcriptional activation of LONP1 [89]. The NRF-2 consensus binding site of *LONP1* should not be confused with the ARE or EPRE consensus binding sites on many other genes, for another well-known (and similarly named) transcription factor, the nuclear factor erythroid 2–related factor 2 (abbreviated as Nrf2) which plays a major role in cellular adaptation to various perturbations such as oxidative stress [38].

The ability of NRF2 to regulate Lon expression is a key factor in its ability to promote the expression of key components of the *mt*DNA transcription and replication machinery, as well as of genes encoding respiratory subunits [39]. Another very important transcriptional site is the region -2023/-1230 – containing a putative binding site for NF-kB [36]. The presence of an NF-kB binding site further consolidate Lon's role as a stress protein [34], as well as its involvement in oncogenesis [40] – a process which significantly involves NF-kB regulated pathways [41].

Lon expression is also significantly induced by Hypoxia-inducible factor 1 (HIF-1) [42]. The Lon gene promoter also contains of five potential HIF-1-binding sites within 0.6 kb of the transcription start site, and ChIP assays have demonstrated the binding of protein complexes containing HIF-1a and HIF-2a to the Lon promoter [42]. Another regulator of Lon expression is the Epidermal Growth Factor (EGF) [43]. EGF is one of the main genes involved in tumorigenic transformation, and disruption of mitochondrial homeostasis is an important characteristic of cancer. EGF regulation of Lon expression is mediated by two important oncogenic pathways: the Extracellular signal-regulated protein kinase (ERK) pathway and the Phosphoinositide 3-kinase (PI3K) pathway [42].

Lon is also regulated at a post-translational level. The NAD+ -dependent mitochondrial deacetylase sirtuin 3 (SIRT3) mediates Lon deacetylation, and SIRT3 silencing causes an increase of Lon protein levels but not of Lon mRNA [44].

3. Normal functions of Lon in healthy individuals

Lon can function as a protease, as a chaperone, and as a mtDNA–binding protein. These various functions are discussed below, and the spectrum of Lon functions described is also reviewed in Table 1.

3.1. Lon as a protease

The initial function discovered for mitochondrial Lon was its proteolytic activity. Our original work in human cell WI-38 VA-13 lung fibroblasts identified an ATP-stimulated serine protease localized in the mitochondrial matrix which is responsible for the degradation of oxidized mitochondrial aconitase [45]. The proteolytic activity is maintained even in the absence of ATP, but protein degradation is stimulated by to up to nine-times by ATP availability [25]. Importantly, ATP hydrolysis is required for this stimulatory effect and non-hydrolyzable ATP analogs are ineffective. Interestingly, Lon can degrade its substrates both in a folded and in an unfolded state – preferring the degradation of unassembled proteins [46]. In the case of properly folded, functionally competent mitochondrial proteins, the first cleavage occurs preferentially between hydrophobic amino acids located within highly charged environments, while subsequent cleavages proceed sequentially along the primary polypeptide sequence [46].

The list of Lon substrates is growing every year – and includes both specific and nonspecific targets. A very important function of Lon is to maintain mitochondrial homeostasis – by recognizing and degrading oxidatively modified proteins [45]. Mitochondrial proteins are constantly in contact with free radical species generated by the respiratory chain, and the oxidized proteins have to be quickly removed and replaced, before they accumulate and cause further toxicity [47]. In mammalian cells, Lon down-regulation by either genetic manipulation [48,49] or by physiologic conditions in intact organisms [37] (oxidative stress, aging) leads to accumulation of damaged proteins inside mitochondria. The proteolytic function is conserved from bacteria to yeast and to humans – as a yeast strain lacking a functional *Lon* gene also has a reduced ability to degrade mitochondrial matrix proteins, which accumulate as electron-dense inclusions [13]. As expected, the mutant yeast strain can be rescued by expressing bacterial (Lon) protease La [50].

The identified specific Lon targets are mitochondrial matrix proteins, involved either in mitochondrial bioenergetic processes, protection against oxidative damage, or in mitochondrial DNA maintenance. Some of the first targets identified were adrenodoxin reductase and cytochrome P-450 [16]. Another proposed protein substrate is SP-22 [51], a thioredoxin-dependent peroxide reductase which functions as a defense system against superoxide, hydrogen peroxide, and related reactive species in adrenocortical mitochondria [52].

We and others have published extensively on Lon protease's ability to selectively degrade oxidatively-modified aconitase [37,45,53]. The primary role of mitochondrial aconitase is to catalyze the stereo-specific isomerization of citrate to isocitrate, via cis-aconitate, in the mitochondrial matrix Krebs cycle [54]. Aconitase easily undergoes oxidative modification and its activity is widely used as a biomarker for oxidative stress. Aconitase inactivation in aging and certain oxidative stress-related disorders [37,53], and its accumulation, is a major pathologic event, which leads to cellular dysfunction and death [55].

Another important Lon substrate is the mitochondrial transcription factor A (TFAM) [56]. TFAM directly regulates the *mt*DNA copy number in mammals [57], and is one of the core mitochondrial nucleoid components, together with mitochondrial single-strand binding protein (mtSSB), polymerase gamma (POLG), and *mt*RNA polymerase (POLRMT). By controlling TFAM degradation, Lon regulates the DNA copy number and transcription [56]. TFAM is phosphorylated within its HMG box 1 (HMG1) by a cAMP-dependent protein kinase, and its phosphorylation impairs the ability of TFAM to bind DNA and to activate transcription. TFAM phosphorylation also promotes TFAM degradation, as Lon degrades only the phosphorylated TFAM [58].

Finally, Lon's proteolytic activity is also involved in hypoxic adaptation, through the Hypoxia-inducible factor 1 (HIF-1) pathway [42]. HIF-1 regulates COX4 subunit composition by increasing the transcription of both Lon and COX4-2. Lon overexpression then promotes the degradation of COX4-1. The change in COX4 subunit expression allows for the optimization of mitochondrial ATP production, O_2 consumption, and reactive oxygen species generation at different O_2 concentrations [42].

3.2. Lon as a chaperone

Co-immunoprecipitation experiments have identified 76 different mitochondrial proteins that bind Lon [59]. Some of these proteins are Hsp60, mtHsp70, Myosin-9/10, 78 kDa glucoseregulated protein (GRP78), [23] and protocadherin-18, and NADH: ubiquinone oxidoreductase core subunit S8 (NDUFS8). Lon contributes to the stability of Hsp60– mtHsp70 complex, and protects cell from apoptosis under environmental stress through interacting with and stabilizing Hsp60.[59]

3.3. Lon as a mtDNA-binding protein

Initial reports suggested Lon might be involved in controlling the number of mitochondria in cells, or the extent of the mitochondria reticulum. Lon over-expression and increased protease activity were reported in cells with an increased mitochondrial biogenesis, such as Zajdela hepatoma tumor [60]. We now know that human Lon can bind to mitochondrial promoters, and may control *mt*DNA transcription and translation by degrading multiple regulatory proteins involved in *mt*DNA maintenance (such as TFAM and POLG) [56,61]. Enzymatically active mouse Lon that hydrolyses ATP and degrades protein and peptide substrates in an ATP-stimulated manner also specifically binds to single-stranded but not to double-stranded DNA oligonucleotides [62]. Human Lon associates with *mt*DNA sequences containing at least 4 contiguous guanine residues - which can form G-quadruplexes (four-stranded intra- or inter- molecular structures with a tetrad arrangement of guanines) [63]. In

human fibroblasts and in colon cancer cells, Lon down-regulation is associated with loss of mitochondrial DNA [14,48]. However, this is not an universal finding – other groups have reported that Lon downregulation in the LS174T cells no significant difference in mtDNA copy number was observed [64] (Fig. 1)

4. Lon involvement in human diseases

As mentioned above, Lon expression is highest in the most metabolically active organs – brain, heart, liver and skeletal muscle [21]. Therefore, these organs seem to be commonly involved in conditions caused by Lon dysregulation (see Fig. 2). In addition, malignant transformation and oncogenic growth are highly dependent on adaptation to new sources of energy and to hypoxia – which explains the important role Lon plays in a variety of malignancies. As Lon is upregulated by acute oxidative stress, it is also involved in a number of conditions caused by the ingestion of toxins and drugs which cause damage via generation of reactive oxygen or nitrogen species such superoxide, hydrogen peroxide, and peroxynitrite, as in different tissues involved in detoxification such as the kidney and adipose tissues.

4.1. Lon involvement in neurological disorders

4.1.1. Lon and cerebral, ocular, dental, auricular, skeletal anomalies (CODAS) syndrome—Cerebral, ocular, dental, auricular, skeletal anomalies (CODAS) syndrome is a rare, multi-system developmental disorder originally described in 1991 [65]. Less than 20 cases have been reported to this day [65–67]. Two different groups reported in 2015 that the genetic abnormalities causing the CODAS Syndrome are compound heterozygous or homozygous mutations in *LONP1* (either missense or nonsense point mutations or small inframe deletions) [66,67]. The CODAS pattern of inheritance is autosomal recessive [66,67]. The amino acid substitutions cluster within the ATP-binding (AAA) and proteolytic domains of Lon [66,67]. Lymphoblastoid cell lines generated from CODAS patients have swollen mitochondria with electron-dense inclusions and abnormal inner-membrane morphology, aggregated *mt*DNA-encoded subunit II of cytochrome c oxidase; and altered respiratory capacity, leading to impaired mitochondrial energy production [67].

A number of children identified from the Old Order Amish community of Pennsylvania were homozygous for *LONP1* c.2161C>G [67]. The homozygous *LONP1* phenotype is very severe – confirming the very important role that *LONP1* regulation of mitochondrial activity has in human development [67]. All the affected children were very severely impacted by their disease – they either died of laryngeal obstruction in the first days of life or required tracheostomies to survive infancy. Another case was identified during pregnancy – and was a stillborn at 39 weeks of gestation due to severe abnormalities. Less than half of the children survived their early years – and had paretic, atrophic vocal cords, glottis narrowing, chronic sialorrhea, and swallowing dysfunction. They required a gastrostomy tube for nutrition.

The CODAS phenotype included broad skull and flattened midface, dysmorphic ears and noses, and with advancing age, short stature, scoliosis, *genu valgus*, and *pes valgus*. Metaphyseal dysplasia (most evident at hip joints), and both hypoplasia and delayed ossification of epiphyses were present. Scoliosis was diagnosed within the first few years of

life [67]. Teeth erupted late with cusp-tip extensions. Dense bilateral nuclear cataracts and hearing loss were present [67].

All the affected children had developmental and cognitive abnormalities – including hypotonia, developmental delay, and variable intellectual disability. Brain magnetic resonance imaging (MRI) studies were notable for revealing extensive brain atrophy, with prominent cortical *sulci* and ventriculomegaly as well as for subcortical hypomyelination, and a thin *corpus callosum* [67].

4.1.2. Lon and hereditary Parkinson disease—Loss-of-function mutations in the PARK6 gene coding for PTEN-induced kinase 1 (PINK1), which encodes a mitochondrially targeted serine/threonine kinase, cause an early-onset, autosomal recessive form of Parkinson's disease. PINK1 is constitutively degraded in healthy cells, but selectively accumulates on the surface of depolarized mitochondria, thereby initiating their autophagic degradation. Lon down-regulation causes dramatic accumulation of processed PINK1 species in the mitochondrial matrix. The processed forms of PINK1 that accumulate upon Lon inactivation are capable of activating the PINK1-Parkin pathway in vivo [68], and to target mitochondria for mitophagy [68].

4.1.3. Lon and Friedreich ataxia (FRDA)—Friedreich ataxia (FRDA) is a genetic neurodegenerative disease affecting the central and peripheral nervous systems, heart, skeleton, and endocrine pancreas [69]. The cause of the disease is a homozygous guanine-adenine (GAA) trinucleotide repeat expansion on chromosome 9q13 that causes a transcriptional defect of the frataxin gene, resulting in reduced expression of mitochondrial frataxin, an iron chaperone involved in the maturation of Fe-S cluster proteins [69]. In the FRDA mouse model, frataxin deficiency causes cardiodegeneration, deficiency of respiratory chain complexes I–III and aconitases, and mitochondrial iron accumulation [70]. The same animals show a clear progressive increase in Lon protein levels. Lon upregulation is also accompanied by an increase in proteolytic activity, and by decreased levels of mitochondrial Fe–S proteins. The effect of Lon upregulation on loss of mitochondrial Fe-S proteins during the progression of the disease suggests that Fe-S proteins may be targets of Lon in FRDA [71].

4.1.4. Lon and familial amyotrophic lateral sclerosis (ALS)—Amyotrophic lateral sclerosis (ALS) is a severe neurodegenerative disease characterized by progressive motor neuron death, causing progressive paralysis and lethal respiratory failure [72]. Mutations in copper-zinc superoxide dismutase (SOD1) represent the most common defect seen in the familial forms of ALS (fALS) [73]. Lon expression is down-regulated in the NSC34 cells (a motor neuron-like cell line) expressing mutant G93A-SOD1 gene [74]. Increasing evidence supports the concept that mitochondrial dysfunction and activation of apoptosis play critical roles in the fALS etiology. Lon down-regulation causes apoptosis in multiple human cell lines [48,59] – raising the question of a potential direct relation between Lon downregulation and motor neuron apoptosis in fALS [74].

4.1.5. Lon and Myoclonic epilepsy and ragged-red fibers (MERRF) syndrome— Myoclonic epilepsy and ragged-red fibers (MERRF) syndrome is a rare mitochondrial

disorder characterized by myoclonus, muscle weakness, cerebellar ataxia, heart conduction block, and dementia [75]. The increase in reactive oxygen or nitrogen species such superoxide, hydrogen peroxide, and peroxynitrite, caused by the respiratory chain dysfunction due to A8344G mutation in the tRNA^{Lys} mitochondrial gene causes damage proteins containing Fe–S clusters such as mitochondrial aconitase [76] – an enzyme predominantly degraded by Lon [45]. In MERRF cybrids containing >90% mtDNA with the A8344G mutation, aconitase activity is significantly decreased while Lon protein expression is highly up-regulated. However, the activity of Lon protease in the MERRF cybrids was significantly lower than that of the wild-type cybrids, suggesting that the up-regulation of Lon protease in the MERRF cells may be a compensation for the decrease of its enzymatic activity caused by oxidative damage.

4.1.6. Lon and myopathy, encephalopathy, lactic acidosis, stroke-like episodes syndrome (MELAS)—The myopathy, encephalopathy, lactic acidosis, stroke-like episodes syndrome (MELAS) is a mitochondrial disorder most frequently associated with an A to G transition at position 3243 of the mitochondrial tRNA(Leu(UUR)) gene [77]. MELAS patient-derived cells have increased mitochondrial Lon protein expression, which accumulates preferentially in MELAS cells containing numerous mitochondria with mutated genomes. Enzymatic assays revealed that –in contrast with the MERRF cybrids–Lon protease activity is also increased in MELAS cell lysates [78].

4.1.7. Lon and brain ischemia/stroke—Lon mRNA and protein levels are induced by hypoxia in cultured astrocytes. In the rat model of ischemic stroke (middle cerebral artery occlusion), both Lon mRNA and protein expression are induced by ischemia, especially in neurons [79]. Cerebral hypoxia/ischemia is known to induce severe endoplasmic reticulum (ER) stress. Attenuation of protein synthesis in response to ER stress occurs to lessen the load of protein entering the ER, and this pathway requires an activation of the ER-resident protein kinase, PERK. Lon induction by ER stress is inhibited in PERK (-/-) cells, confirming Lon's involvement in the ER stress response [79].

4.2. Lon involvement in cardiac disorders

4.2.1. Lon, cardiac ischemia and heart failure—In ischemic conditions, cardiomyocytes are subjected to increased generation of reactive oxygen and nitrogen species, mitochondrial dysfunction and apoptosis [80]. Lon is upregulated in hypoxia-induced cardiomyocytes, while Lon downregulation attenuates hypoxia-induced cardiomyocyte apoptosis through decreased oxidant generation. Lon overexpression stimulated oxidant production and induced apoptosis under normoxic conditions in cardiomyocytes [81] – in contrast with data in other human cells where Lon over-expression stimulates cell proliferation [60], while Lon downregulation causes senescence [47] and/or apoptosis [48,49].

Lon carbonylation and tyrosine nitration are prominent in a mouse model of pressure overload heart failure after transverse aortic constriction (TAC) [82]. The measurement of ATP-dependent proteolytic activity in isolated mitochondria revealed marked reductions in mitochondrial protease activity. Oxidative post-translational modifications (OPTM)

contribute to the proteolytic deficit of Lon and OPTM-mediated protease dysfunction and is closely associated with declines in electron transport chain (ETC) protein homeostasis, mitochondrial energetics, and left ventricular contractility. Infusion of mitoTEMPO, a mitochondria-targeted superoxide dismutase mimetic, improves mitochondrial respiration capacity and cardiac function at least partially through the improvement of Lon activity in heart failure [82].

4.3. Lon involvement in skeletal muscle pathology

4.3.1. Lon and muscle denervation—Muscle denervation is commonly seen in traumatic injuries. In the experimental model of sciatic nerve transection, Lon protease was down-regulated at day 30 after surgery, which correlated with a decreased number of functional mitochondria [83].

4.3.2. Lon and Hindlimb unloading (loss of gravitational pull in space travel)— Hindlimb unloading (HU) rodent models are frequently used to model the neuromuscular changes taking place in a real microgravity environment such as that experienced during spaceflights [84]. HU is characterized by muscle atrophy and dysfunction, and loss of mitochondria – with the loss of subsarcolemmal mitochondria being much greater than the loss of intermyofibrillar mitochondria [85]. Lon is down-regulated in HU exposed muscles, which might explain the decreased mitochondrial biogenesis, suggesting that Lon protease may play some role in mitochondrial adaptations to HU [85].

4.3.3. Lon and muscle regeneration—Skeletal muscles exhibit a high capacity to repair and regenerate after trauma or disease. Upon injury, satellite muscle cells can proliferate, and fuse with the existing myofibers or form new myofibers [86]. Myogenic differentiation is associated with an increase in mitochondrial biogenesis, and requires increased expression of mitochondrial biogenesis-related genes including Lon. These results suggest that mitochondrial biogenesis is activated early during muscle regeneration and that mitochondrial biogenesis plays a significant role in muscle regeneration. [87]

4.4. Lon involvement in steroid hormone synthesis (StAR protein degradation)

Steroidogenic acute regulatory protein (StAR) is required for steroid hormone production in the adrenal cortex and the gonads. StAR mediates the transport of cholesterol into the inner mitochondrial membranes where is converted to a steroid [88]. Lon protease is involved in StAR proteolysis [118], and gonadotropin administration to prepubertal rats stimulates ovarian follicular development and increases Lon expression. Furthermore, StAR expression stimulates *LONP1* gene transcription by acting through the NRF-2 pathways on the proximal promoter of the LON gene proteases [89] (which contains the NRF-2 responsive elements)-suggesting a novel regulatory loop through which Lon and StAR regulate each other's availability [90].

4.5. Lon involvement in hepatic disorders

4.5.1. Lon and 5-aminolevulinic acid synthase protein metabolism—5-

Aminolevulinic acid synthase (ALAS-1) is the first rate controlling enzyme that controls cellular heme biosynthesis in the liver and in bone marrow cells [91]. Negative regulation of

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ALAS-1 by the end product heme provides the foundation for heme treatment of acute porphyrias; these are a group of diseases caused by genetic defects in the heme biosynthesis pathway that are exacerbated by controlled up-regulation of ALAS-1. Heme and other metalloporphyrins decrease mitochondrial ALAS-1 protein levels by accelerating its proteolytic degradation by Lon protease. Conversely, down-regulation of Lon activity diminishes the negative effect of heme on mitochondrial ALAS-1 [92].

4.5.2. Lon and hepatic insulin resistance—Mitochondrial dysfunction is a cause of insulin resistance and type 2 diabetes [93]. In human liver cells, Lon downregulation causes impaired insulin signaling and increased levels of gluconeogenic enzymes, while Lon over-expression diminishes the insulin resistance induced by treatment with cholesterol and palmitate. In-vivo, Lon protein levels are significantly lower in the livers of diabetic db/db mice compared with their lean mice counterparts, suggesting that Lon down-regulation might be involved in the development of insulin resistance and type 2 diabetes [94].

4.6. Lon involvement in cancer

Mitochondrial dysfunction is a hallmark of cancer biology. Tumor mitochondrial metabolism is characterized by an abnormal ability to function in scarce oxygen conditions [95] by using glycolysis (the Warburg effect), and accumulation of mitochondrial DNA defects is present in both hereditary neoplasias and sporadic cancers [96]. Lon is highly expressed in aggressive tumors [97], which are characterized by extensive hypoxic areas caused by insufficient oxygen and nutrient supply which are the hallmarks of aggressive tumors [98]. Several key roles for Lon in cancer [99] include: 1) hypoxic survival, 2) invasion and metastasis, 3) resistance to apoptosis, and 4) treatment resistance.

4.6.1. Lon and malignant transformation—Lon is involved in Epidermal Growth Factor (EGF) and ErbB2/Her-2 induced tumorigenic transformation in both epidermal and mammary cancers. Irreversibly transformed mouse epidermal cells have higher Lon levels than the parent, non-transformed cells (JB6 P⁺). In addition the epidermal growth factor which induces tumorigenic transformation of the JB6 P⁺ cells also up-regulates Lon expression [43].

4.6.2. Lon and colon cancer—Lon silencing in RKO colon cancer cells, either by using either Lon shRNA or the triterpenoid Lon-inhibitor, 2-cyano-3,12-dioxooleana-1,9,-dien-28-oic acid, causes profound alterations in the mitochondrial proteome and function, and cell death. Lon-silenced cells display altered levels of mitochondrial proteins, low levels of mtDNA transcripts, and reduced levels of OX-PHOS complexes. Lon-deficient mitochondria have dilated cristae, vacuoles, and electron-dense inclusions [97]. Transgenic mice deficient in Lon protease are protected against the development of chemically-induced colon and skin tumors [14]. *LONP1* is upregulated in colon adenomas and in colon cancer specimens, and high *LONP1* expression correlated with decreased survival in colon cancer patients [14].

4.6.3. Lon and bladder cancer—Lon expression in bladder cancer tissues is significantly higher than in noncancerous tissues. Down-regulation of Lon in bladder cancer cells significantly blocked cancer cell proliferation and increased the sensitivity of bladder

cancer cells to chemotherapeutic agents by promoting apoptosis. {Granot, 2007 #244} High expression of Lon is related to the TNM (tumor, nodes, metastasis) stage, as well as histological grade of bladder cancer patients, and is an independent, negative prognostic factor for the overall survival of bladder cancer patients {Granot, 2007 #244}.

4.6.4. Lon and cervical cancer (HeLa cells)—Lon appears to play less of an essential role in the HeLa cells derived from a cervical cancer patient. In these cells, Lon depletion causes no significant alteration of mitochondrial activities and dynamics. However, the Lon depleted HeLa cells do show an increase in mitochondrial superoxide and hydrogen peroxide generation, and accumulate oxidized proteins, which also eventually results in inhibition of cytoplasmic 20 S proteasome [101].

4.6.5. Lon and leukemia—Adaptation of leukemia cell to hypoxia requires a successful transition from OX-PHOS to glycolysis. The THP-1 cells, which largely depend on OX-PHOS in mitochondria under normoxia, are able to accelerate their growth under hypoxia by changing metabolism from OX-PHOS to glycolysis . This transition is mediated by substituting COX 4 subunit (from COX 4-1 to COX 4-2) through Lon upregulation [102].

4.6.6. Lon and brain cancer (glioblastoma)—Our group showed that Lon controls metabolic adaptation to hypoxia and to starvation in glioma cells in direct response to HIF1- a activation [103]. Also, Lon induction is used by glioma cells to increase resistance to radiation and chemotherapy by direct repair of mtDNA [103].

4.6.7. Lon as a therapeutic target for cancer treatment—The wealth of data showing a crucial role for Lon in cancer growth, invasion and metastasis has also lead to the development of novel Lon inhibitors. The first published Lon inhibitors are coumarinic derivatives - small, non-peptidic molecules, with no reported ability to inhibit other proteases (such as the proteasome) and with potential to cross the blood-brain barrier - an important requirement for drugs targeted to central nervous system malignancies [104].

Triterpenoids such as 2-cyano-3,12-dioxooleana-1,9-dien-28-oic acid (CDDO) and its derivatives promote B-lymphoid cell apoptosis by inhibiting Lon's proteolytic activity [105]. Lon protein levels are substantially elevated in malignant lymphoma cells and Lon knockdown leads to lymphoma cell death, providing 'proof of concept' for the hypothesis that mitochondrial Lon can be a novel target for new anticancer drugs [105]. Similar activity of triterpenoids was found also in RKO human colon cancer cells, in HepG2 hepatocarcinoma cells and in MCF7 breast carcinoma cells [106].

Obtusilactone A and (–)-sesamin, compounds derived from *Cinnamomum kotoense* are potent Lon protease inhibitors, which interact with Ser855 and Lys898 residues in the active site of the Lon protease [107]. Lon is upregulated in non-small-cell lung cancer (NSCLC) cell lines, and downregulation of Lon with either Obtusilactone A or (–)-sesamin triggers caspase-3 mediated apoptosis [107].

4.7. Lon and the response to acute exposure to environmental toxins and to medications

4.7.1. Lon prevents renal cytotoxicity caused by ochratoxin A (OTA)—OTA is a naturally occurring mycotoxin and food contaminant produced by several species of Aspergillus and Penicillium. It is very commonly found in a variety of foods and represents a serious threat to human health. OTA has multiple toxicological effects, including nephrotoxicity, hepatotoxicity, genotoxicity, immunotoxicity, teratogenicity, mutagenicity and carcinogenicity (classified as a possible human carcinogen (group 2B) [108]. OTA exposure causes Lon down-regulation in kidney cells [109], and the Lon-deficient cells are more sensitive to OTA injury. Lon confers protection to OTA-induced renal cytotoxicity via 4 processes: defensing against OTA-induced oxidative stress in the mitochondria; regulating protein synthesis, modification and repair; maintaining the balance of carbohydrate metabolism; and assisting in mtDNA maintenance. [110]

4.7.2. Lon and HIV therapy (HAART) induced lipodystrophy—The combination of antiretroviral medication generally known as HAART (highly active antiretroviral therapy) has revolutionized HIV treatment, by allowing the infected patients to survive for a very long period of time. However, Nucleosidic inhibitors of the reverse transcriptase (NRTI), main components of HAART, display relevant toxicity for mitochondria and can provoke metabolic changes and body fat redistribution, resulting in lipodystrophy, a side effect significantly involving mitochondrial function. The *LONP1* gene is significantly upregulated in adipose tissue from lipodystrophy patients, and correlates with the high turnover of damaged proteins caused by increases in metabolic production of oxidants. However, increased Lon expression is not sufficient to eliminate all the oxidatively damaged proteins that accumulate and contribute to the development of lipodystrophy [111].

4.8. Proposed etiology of Lon involvement in human pathology and disease

Mitochondrial proteolytic mechanisms in mammals are complex, a feature required to maintain mitochondrial homeostasis. Potentially, any imbalance in the fragile equilibrium between components encoded by the mitochondrial genome and produced in the mitochondrial matrix, and components encoded by the nuclear genome and translocated from the cell cytoplasm can cause human pathology.

Though multiple proteolytic systems are described in mammalian mitochondria [10], the Lon protease seems to be the dominant enzyme involved in maintaining appropriate mitochondrial function, protecting the *mt*DNA against a variety of insults and controlling mitochondrial protein stoichiometry. Lon itself is a Janus-like protein, with tightly regulated levels, and with potential harm that can result from either downregulation or overexpression. In patients where either Lon expression or Lon function is significantly dysregulated, pathology inevitably seems to occur. Lon downregulation causes increased susceptibility to toxins, and is involved with a wide array of degenerative conditions, involving multiple organs. Lon upregulation promotes malignant transformation, increases the resistance of multiple malignancies to radiation and chemotherapy, and predicts worse outcomes in cancer patients.

In this paper we have reviewed existing knowledge of both the normal functions and the associated pathologies associated with Lon in humans. We would also like to propose a novel etiological classification of human disorders involving Lon. Our classification identifies four different categories of pathology, based on the presence or absence of a genetic mutation in the *LONP1* gene, Lon protein level, and Lon protease activity (see below, and also Table 2).

4.8.1. Proposed etiological classification of Lon-related disorders

Decreased Lon Expression and Loss of Function Caused by Mutations or Deletions in the Encoding Gene (*LONP1*):

Cerebral, ocular, dental, auricular, skeletal anomalies (CODAS) syndrome.

Decreased Lon Expression and Decreased Proteolytic Activity Causing/ Contributing to a Pathologic State (*no genetic defect in LONP1 gene*):

Early-onset, autosomal recessive form of Parkinson's disease.

Familial Amyotrophic Lateral Sclerosis (ALS).

-

Myopathy, encephalopathy, lactic acidosis, stroke-like episodes syndrome (MELAS).

—

Muscle denervation.

Hindlimb Unloading.

-

Insulin resistance and Type 2 diabetes.

-

Ochratoxin A Toxicity.

3.

1.

2.

Lon Overexpression and Increased Proteolytic Activity Causing/ Contributing to a Pathologic State (*no genetic defect in LONP1 gene*):

Friedreich ataxia (FRDA).

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Brain Ischemia/Stroke.

HAART Induced Lipodystrophy.

Multiple Malignancies.

Lon Overexpression but Decreased Lon Proteolytic Activity Causing/ Contributing to a Pathologic State (*no genetic defect in LONP1 gene*):

Myoclonic epilepsy and ragged-red fibers (MERRF) syndrome (increased expression but loss of proteolytic activity).

Cardiac Ischemia and Heart Failure (increased expression but loss of proteolytic activity).

5. Lon involvement in aging

4.

Levels of *Lon* mRNA transcripts were found to be about four-times lower in skeletal muscles from aged mice, than in young adult control animals, but this phenomenon was prevented by caloric restriction [112]. We next found that Lon protein levels, as well as total Lon proteolytic activity, were greatly reduced in the muscles of aged normal mice, and that this phenomenon was even more severe in mice burdened by a lifetime of chronic exposure to oxidative stress, caused by an approximate 50% reduction in manganese superoxide dismutase activity (manganese SOD heterozygote mice) [37]. Interesting, exercise can effectively minimize both the age-induced Lon decline and decreased mitochondrial biogenesis in aging muscle [113]. These results may actually suggest that declining Lon transcription, translation, and enzymatic activity in aging may result from a combination of both age and disuse, at least in skeletal muscles.

Age-related decline in Lon expression and/or its proteolytic activity might also be tissuespecific [114]. For example, a measurable decrease in Lon proteolytic activity occurs in parallel with the accumulation of damaged proteins in rat liver mitochondria isolated from aged animals [115]. However, Lon protease activity remains constant in the heart mitochondrial matrix isolated from the same aged rats, and the levels of expression of the Lon protease actually increased in the older animals in comparison with the younger ones [53]. This may, again, be evidence for a strong influence of tissue metabolic activity (mitochondrial respiration?) on Lon transcription, translation, and enzymatic activity, since heart muscle (unlike skeletal muscles) must be at least minimally active throughout life.

Clearly, the question of Lon regulation in aging remains largely unanswered and significant work will be needed to identify tissue-specific patterns of expression in aging, as well as the effects of exercise and other potential metabolic effectors. Finally, it will be very important

to discover the effects of interaction of age and disease on Lon transcription, translation, and enzymatic activity.

6. Conclusions

A number of human diseases have been identified whose mechanisms involve Lon dysfunction. These diseases and a proposed etiological classification are summarized in Table 2. Many of the same mechanisms that govern Lon-related pathology are also involved in the aging process, and in diminished resistance to multiple stressors including oxidative stress. Extensive further study of human diseases involving mitochondrial Lon, including mechanistic studies of enzymatic function and elucidation of substrate recognition, is required in light of the number and the variety of severe mitochondrial conditions of unknown etiology. New therapies focused on Lon inhibition as a potential target for cancer therapies are currently underway. However, their development has to be approached with careful attention to possible severe central nervous system and cardiac toxicities – that are potentially associated with chronic Lon down-regulation. Finally, the interaction of Lon regulation in aging, with the effects of many age-related pathologies will be a significant challenge for researchers in the field to tackle.

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Abbreviations

ALAS-1	5-Aminolevulinic acid synthase			
ALS	Amyotrophic Lateral Sclerosis			
CDDO	2-cyano-3,12-dioxooleana-1,9-dien-28-oic acid			
CODAS	Cerebral, ocular, Dental, Auricular, Skeletal Anomalies Syndron			
EGF	Epidermal Growth Factor			
ERK	Extracellular signal-regulated protein kinase			
ER	endoplasmic reticulum			
ETC	electron transport chain			
FRDA	Friedreich ataxia			
GRP78	78 kDa glucose-regulated protein			
HAART highly active antiretroviral therapy				
HIF-1	Hypoxia-inducible factor 1			

HU	Hindlimb unloading		
MELAS	Myopathy, Encephalopathy, Lactic Acidosis, Stroke-like Episodes Syndrome		
MERRF	Myoclonic Epilepsy and Ragged-red Fibers Syndrome		
mtSSB	mitochondrial single-strand binding protein		
NDUFS8	NADH:ubiquinone oxidoreductase core subunit S8		
NRF-2	Nuclear Respiratory Factor 2		
NRTI	Nucleosidic inhibitors of the reverse transcriptase		
NSCLC	non-small-cell lung cancer		
OPTM	Oxidative post-translational modifications		
OTA	Ochratoxin A		
OX-PHOS	Oxidative phosphorylation		
PI3K	Phosphoinositide 3-kinase		
PINK1	PTEN-induced kinase 1		
POLG	polymerase gamma		
POLRMT	<i>mt</i> RNA polymerase		
PERK	ER-resident protein kinase		
SOD	superoxide dismutase		
StAR	Steroidogenic acute regulatory protein		
TAC	transverse aortic constriction		
TFAM	mitochondrial transcription factor A		
TNM	tumor, nodes, metastasis stage		
UCP	Uncoupling proteins		

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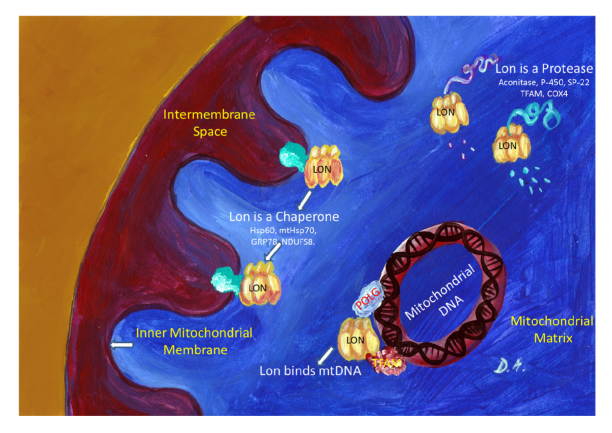
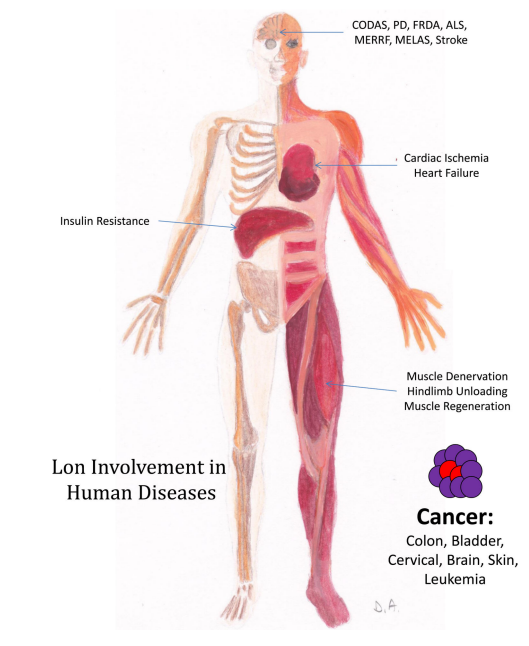


Fig. 1.

Schematic representation of Lon involvement in mitochondrial biology.



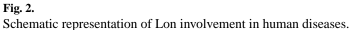


Table 1

Lon functions in mammalian cells.

Function	Partners/Substrates
Protease	adrenodoxin reductase [16]
	cytochrome P-450 [16] 99
	SP-22 [51]
	Aconitase [37,45,53]
	TM [56]
	StAR [89,90]
	COX4–1 [41]
	ALAS-1 [90]
	PINK1 [66]
	POLG [54]
Chaperone	Hsp60 [57]
	mtHsp70 [57]
	Myosin-9/10 [57]
	78 kDa glucose-regulated protein (GRP78) [57]
	protocadherin-18 [57]
	NADH:ubiquinone oxidoreductase core subunit S8 (NDUFS8) [57].
mtDNA binding	<i>mt</i> DNA sequences containing at least 4 contiguous guanine residues - G-quadruplexes (four-stranded intra- or inter- molecular structures with a tetrad arrangement of guanines) [60,61]

Table 2

Etiological classification of Lon-related human disorders.

Category	Disease	LONP1 Gene	Lon Expression	Lon Proteolytic Function
1) Decreased Lon Expression and Loss of Function Caused by Mutations or Deleons in the Encoding Gene	Cerebral, ocular, dental, auricular, skeletal anomalies (CODAS) syndrome [65–67]	mutated	decreased	decreased
2) Decreased Lon Expression and	Early-onset Parkinson's disease [68]	normal	decreased	decreased
Decreased Proteolytic Activity Causing/Contributing to a Pathologic State	Familial ALS [74]	normal	decreased	decreased
	MELAS [78]	normal	decreased	decreased
	Muscle denervation [83]	normal	decreased	decreased
	Hindlimb Unloading [85]	normal	decreased	decreased
	Insulin resistance and Type 2 diabetes 94]	normal	decreased	decreased
	Ochratoxin A Toxicity [109,110]	normal	decreased	decreased
3) Lon Overexpression and	Friedreich ataxia (FRDA) [71]	normal	increased	increased
Increased Proteolytic Activity Causing/Contributing to a Pathologic State	Brain Ischemia/Stroke [79]	normal	increased	increased
	HAART Induced Lipodystrophy [111]	normal	increased	increased
	Multiple Malignancies [99]	normal	increased	increased
4) Lon Overexpression but	MERRF [76]	normal	increased	decreased
Decreased Lon Proteolytic Activity Causing/Contributing to a Pathologic State	Cardiac Ischemia and Heart Failure [81]	normal	increased	decreased