Multiple Pathways to the Same End: Mechanisms of Myonuclear Apoptosis in Sarcopenia of Aging

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Sarcopenia, the age-related decline in muscle mass and function, represents a significant health issue due to the high prevalence of frailty and disability associated with this condition. Nevertheless, the cellular mechanisms responsible for the loss of muscle mass in old age are still largely unknown. An altered regulation of myocyte apoptosis has recently emerged as a possible contributor to the pathogenesis of sarcopenia. Studies in animal models have shown that the severity of skeletal muscle apoptosis increases over the course of aging and correlates with the degree of muscle mass and strength decline. Several apoptotic pathways are operative in aged muscles, with the mitochondria- and TNF-α–mediated pathways likely being the most relevant to sarcopenia. However, despite the growing number of studies on the subject, a definite mechanistic link between myocyte apoptosis and age-related muscle atrophy has not yet been established. Furthermore, the evidence on the role played by apoptosis in human sarcopenia is still sparse. Clearly, further research is required to better define the involvement of myocyte apoptosis in the pathogenesis of muscle loss at advanced age. This knowledge will likely help in the design of more effective therapeutic strategies to preserve muscle mass into old age, thus fostering independence of the elderly population and reducing the socioeconomic burden associated with sarcopenia.

KEYWORDS: aging, sarcopenia, myonuclear apoptosis, mitochondria, tumor necrosis factor-alpha, caspases, endonuclease G, apoptosis inducing factor

INTRODUCTION

The age-related loss of muscle mass and function, referred to as sarcopenia, is a highly prevalent condition among older persons, affecting 10–25% of the population under the age of 70 years, and over 40% of the individuals aged 80 years or older[1,2]. This age-dependent muscle loss has been reported
SIGNALING PATHWAYS OF APOPTOTIC CELL DEATH

Apoptosis, a process of programmed cell death, is an evolutionarily conserved, tightly regulated, systematic set of events resulting in cellular self-destruction without inflammation or damage to the surrounding tissue[21]. Apoptosis plays a fundamental role in the development and maintenance of tissue homeostasis. Proteolytic enzymes, known as caspases, perform the dismantling of the cell and are normally present as inactive zymogens (procaspases)[22]. Upon appropriate stimuli, initiator caspases (i.e., caspase-8, caspase-9, caspase-12) are activated, leading to the activation of effector caspases (i.e., caspase-3, caspase-6, caspase-7) responsible for the cellular degradation and DNA fragmentation via a caspase-activated DNase (CAD)[23]. Activation of caspases can occur through extrinsic and intrinsic pathways (Fig. 1)[22]. The extrinsic apoptotic signaling is initiated by death receptors located on the cell surface, such as the tumor necrosis factor receptor (TNF-R) and the Fas receptor[24]. Activation of these receptors by binding of death-stimulating ligands results in the recruitment of adaptor proteins, such as Fas-asssociated death domain (FADD), TNF-receptor-associated death domain (TRADD), and TNF-receptor-associated factors (TRAFs), forming the death-inducing signaling complex (DISC), which engages and activates procaspase-8. Active caspase-8 subsequently activates caspase-3[24]. Intrinsic pathways of caspase activation include those triggered by the endoplasmic reticulum (ER) and the mitochondrion[22]. Under stress conditions, such as calcium dyshomeostasis, the ER-specific procaspase-12 can be activated by m-calpain, leading to caspase-3 activation[25]. However, the mitochondria-mediated intrinsic signaling pathway of apoptosis is currently considered the most relevant means of programmed cell death and has been the subject of intense scientific focus[26].

Notably, mitochondria can induce apoptosis via both caspase-mediated and caspase-independent pathways[26]. The former is initiated by the release of cytochrome c into the cytoplasm, where it complexes with apoptotic protease-activating factor-1 (Apaf-1), dATP, and procaspase-9, forming the apoptosome[26]. Within this complex, procaspase-9 becomes activated, subsequently engaging caspase-3[26]. In addition, mitochondria can release the second mitochondria-derived activator of caspases/direct inhibitor of apoptosis-binding protein with low pi (Smac/DIABLO) and heat requirement A2 protein (Omi/HtrA2), both of which block the activity of the inhibitor of apoptosis proteins (IAPs)[27]. Hence, inhibition of caspase activity is removed and caspase-mediated proteolysis proceeds.
Aside from their role in cytochrome c–mediated induction of cell death, mitochondria also participate in a pathway of apoptosis that does not involve caspase activation. Endonuclease G (EndoG) and apoptosis inducing factor (AIF) are two mediators that, upon release from mitochondria, translocate to the nucleus and perform DNA fragmentation independent of caspases[27,28]. Specifically, AIF binds to DNA to induce chromatin condensation[28]. AIF can also interact with cyclophilin A, forming a DNase responsible for large-scale DNA fragmentation[29]. EndoG, a mitochondrion-specific nuclease, is thought to participate directly in oligonucleosomal DNA fragmentation[30].

Upstream of the release of mitochondria-specific apoptotic mediators is the process of mitochondrial outer membrane permeabilization (MOMP)[31], whose intimate mechanisms are currently under intense investigation. The integrity of the outer mitochondrial membrane (OMM) is regulated by the Bcl-2 family
of proteins that includes pro- (e.g., Bax, Bak, Bad, Bim, Bid, Puma, Noxa) and antiapoptotic (e.g., Bcl-2, Bcl-X<sub>L</sub>, Bcl-w, Mcl-1, A1) members[32]. The balance between these mediators (e.g., Bax-to-Bcl2 ratio) is considered a fundamental control point for the cell fate by regulating OMM stability. Under normal conditions, Bcl-2 and Bcl-X<sub>L</sub> block Bax and/or Bak[33]. In apoptotic conditions, Bax translocates from the cytosol to the mitochondria, oligomerizes, and inserts into the OMM in concentrated foci[34]. This process results in the formation of a pore through which apoptotic factors stored in the intermembrane compartment are released[34]. Similarly, Bak, which is constitutively anchored to the OMM, can form homo-oligomeric complexes upon apoptotic stimulation[34]. Bak/Bak hetero-oligomers have also been described[35]. The proapoptotic factors Bid and Bim participate to Bax and Bak activation, through neutralizing Bcl-2 and Bcl-X<sub>L</sub> activity[36]. In addition, Bid allows cross-talk between intrinsic and extrinsic apoptotic pathways, after truncation by caspase-8[37].

MOMP is also thought to occur via opening of the mitochondrial permeability transition pore (mPTP)[38]. The exact nature of the mPTP is still unclear. However, three putative components have been suggested: a voltage-dependent anion channel (VDAC) located in the OMM, the adenine nucleotide translocase (ANT) in the inner mitochondrial membrane (IMM), and cyclophilin D (CyPD) in the matrix[39]. Upon certain stimuli, such as oxidative stress or calcium overload, ANT interacts with VDAC, generating an unselective pore spanning the IMM and OMM[40]. This interaction is stabilized by CyPD. Opening of the mPTP causes a sudden increase in membrane permeability to solutes with molecular weight up to 1,500 Da. This results in dissipation of membrane potential, mitochondrial swelling, and rupture of the OMM, with subsequent release of mitochondrial apoptotic mediators[38]. Interestingly, cross-talk between the mPTP and Bcl-2 proteins has been described, with Bid and Bax promoting mPTP opening[41,42], while the antiapoptotic Bcl-2 and Bcl-X<sub>L</sub> possess an inhibitory effect[43].

THE INVOLVEMENT OF APOPTOSIS IN THE PATHOGENESIS OF SARCOPENIA

A wealth of experimental animal data indicates that the apoptotic program is activated in the aged skeletal muscle, likely contributing to the pathogenesis of sarcopenia (Fig. 1)[44].

Given the central role played by mitochondria in the induction and regulation of programmed cell death, intensive investigation has focused on mitochondria-driven myonuclear apoptosis[45].

Alteration in the expression of Bcl-2 family proteins has been reported in skeletal muscles of aged experimental animals[18,46,47,48,49]. Specifically, a proapoptotic shift in the expression pattern of Bcl-2 proteins (i.e., increased Bax and decreased Bcl-2 levels) has been observed in muscles of aged rodents[47,48,49]. However, other studies did not detect such an adaptation[18,46]. Strikingly, lower levels of Bax and Bcl-2 mRNA were found in the plantaris muscle of old and senescent rats, resulting in a reduced, rather than increased, Bax-to-Bcl-2 ratio in very old animals compared to younger controls[50]. Interestingly, the age-related pattern of Bcl-2 and Bax expression may be muscle-type specific[51]. In fact, Bax content was found to be increased in the fast-twitch extensor digitorum longus (EDL) of old rats, whereas no changes were observed in the slow-twitch soleus. In contrast, both muscles displayed an increased Bcl-2 expression at advanced age. The elevation of Bcl-2 detected in aged muscles may be interpreted as a compensatory attempt to limit myonuclear loss. Alternatively, the increased serine-phosphorylation and subsequent inactivation of Bcl-2 recently reported in old mice[52] might abolish the antiapoptotic activity of this molecule despite elevated expression[53].

Besides alterations in the Bcl-2 rheostat, increased susceptibility towards opening of the mPTP has also been demonstrated in aged skeletal muscles[54,55,56]. Interestingly, elevated mitochondrial levels of CyPD relative to ANT and VDAC were detected in the gastrocnemius muscle of very old rats[18]. This finding underlines the major role postulated for CyPD in the formation of the mPTP, and suggests a higher susceptibility to mPTP opening at an older age. Indeed, it was demonstrated that mPTP formation and apoptotic cell death could still occur in the absence of either ANT[57] or VDAC[58]. In contrast,
overexpression of CyPD induced mitochondrial swelling and spontaneous cardiac apoptosis, whereas mice lacking cardiac CyPD were protected from ischemia/reperfusion-induced cell death[59].

Once MOMP has occurred, the release of apoptogenic factors stored in the mitochondrial intermembrane space ensues, initiating the series of events that culminate in cell death[31]. Elevated cytosolic levels of cytochrome c have been detected in the skeletal muscle of old rats[46]. Accordingly, levels of Apaf-1[13,46,60], active caspase-9 content[52,54,60], and caspase-9 proteolytic activity[46] were also found to be higher in aged muscles compared to young controls. However, other studies did not observe increases in cytosolic cytochrome c and/or caspase-9 expression in skeletal muscles of old rats[12,13,18,49]. Surprisingly, it was observed that gene expression levels of Apaf-1 were even reduced in the plantaris of old and senescent rats relative to younger controls[50].

These observations have cast doubts about the involvement of mitochondrial caspase-dependent apoptosis in the pathogenesis of sarcopenia. In contrast, several studies have shown that caspase-independent apoptotic pathways are activated in old age[12,18,50,61]. It was found that translocation of mitochondrial EndoG to the nucleus was increased in the soleus muscle of old rats. In addition, an age-related increase in both cytosolic and nuclear levels of AIF and EndoG was observed in the rat gastrocnemius muscle[18]. Moreover, AIF gene expression progressively increased during aging in the rat plantaris muscle, and correlated with the progression of sarcopenia[50]. Collectively, these findings support a role for the mitochondrial caspase-independent pathway of apoptosis in the pathophysiology of sarcopenia. Indeed, this apoptotic pathway might be particularly relevant in sarcopenia, since it may allow for the elimination of superfluous/damaged nuclei without the demolition of the entire muscle fiber by caspases[62].

Besides mitochondria-driven apoptosis, other pathways of myonuclear apoptosis have been found to be operative at advanced age. In particular, the death receptor–mediated pathway, triggered by TNF-α, is thought to play a significant role in age-related muscle loss[14,17,19,20], owing to its ability to promote muscle protein wasting[63,64] and myonuclear apoptosis[14,17,19,20,65]. Interestingly, the transduction of TNF-α signaling during aging may be fiber-type specific[14]. In fact, TNF-α–mediated apoptosis was enhanced at old age in the rat superficial vastus lateralis muscle (predominantly comprised of type II fibers), whereas increased signaling through the NF-κB pathway was observed in the soleus muscle, characterized by type I fiber dominance. Notably, NF-κB may promote the expression of antiapoptotic genes such as cIAP-1, cIAP-2, and FLICE-like inhibitory protein long form (FLIPL)[66,67]. This phenomenon could at least partly explain the lower susceptibility to age-related atrophy displayed by slow-twitch muscles compared to those with fast-twitch properties[68]. In support of these findings, Marzetti et al.[19] recently detected an age-related increase in myocyte apoptosis in the rat extensor digitorum longus (EDL, type II) muscle, whereas the extent of DNA fragmentation was unaffected by age in the soleus. Furthermore, the death receptor pathway of apoptosis was activated in the EDL, but not in the soleus of aged rats. Interestingly, 4-week treadmill exercise training reversed the age-related elevation of DNA fragmentation and TNF-α apoptotic signaling in the EDL muscle. Importantly, mitigation in the extent of apoptosis was accompanied by improved exercise tolerance and grip strength, suggesting a strong association between the severity of myocyte apoptosis and physical performance. In another very recent study, the age-dependent increase in TNF-α apoptotic signaling was paralleled by decreased expression of the antiapoptotic interleukin 15 (IL-15) and its specific receptor subunit α (IL-15Rα) in the rat gastrocnemius muscle[20]. In contrast, lifelong calorie restriction (CR), an intervention known to attenuate apoptosis in the skeletal muscle of old rodents[69], preserved IL-15 signaling and muscle mass into very old age, while preventing the activation of the death receptor–mediated apoptosis[20]. These findings suggest that the loss of the anabolic drive by IL-15 may unbridge the proapoptotic actions of TNF-α in skeletal muscle, thus contributing to the development of sarcopenia and loss of physical function in old age.

Similar to the TNF-α/IL-15 axis dysfunction, an altered balance between insulin-like growth factor-1 (IGF-1) and TNF-α may also contribute to the proapoptotic environment taking place in the aged muscle. IGF-1, either systemic or muscle derived, is a potent stimulator of muscle development, hypertrophy, and
maintenance[70]. Importantly, IGF-1 is also able to counteract the proapoptotic actions of TNF-α in skeletal muscle. Indeed, incubation of C2 myoblasts with IGF-1 decreases the incidence of TNF-α–induced apoptosis[71]. Levels of IGF-1 are reduced with advanced age, as a consequence of impaired growth hormone (GH) production[72]. In addition, muscle production of mechano growth factor (MGF), a local splice variant of IGF-1, is decreased in old rats in response to mechanical overload[73]. Furthermore, both the density and affinity of the IGF type 1 receptor are reduced in the aged muscle[74]. Downstream of IGF-1 is the protein kinase B (PKB), also known as Akt. Akt plays a central role in muscle anabolism by stimulating glucose uptake, glycogen synthesis, and protein synthesis[75]. Akt also inhibits myonuclear apoptosis and muscle protein degradation by inactivating Bad[76] and FoxO transcription factors[77], respectively. Insulin-stimulated Akt activation is significantly decreased in advanced age in the rat skeletal muscle[78], as a result of TNF-α–induced insulin resistance[79]. Interestingly, Wu et al.[80] recently reported an age-related decrease in Akt kinase activity in the rat skeletal muscle, which was associated with elevated levels of Akt S-nitrosylation. Noteworthy, amelioration of Akt activity following acetaminophen administration was accompanied by attenuation of myonuclear apoptosis and increases in myocyte cross-sectional area and contractile protein expression.

Other apoptotic pathways that may be operative in the aged muscle include those initiated by caspase-2 and -12; however, their contribution to sarcopenia has yet to be established. Braga et al.[52] recently found an increased caspase-2 expression in the skeletal muscle of old mice. This caspase is thought to cooperate with MOMP during stress-induced apoptosis[81]. However, it is still unclear whether caspase-2 is an initiator or an effector caspase and how it is activated.

Finally, recent evidence suggests that the ER-induced apoptosis may participate to age-related muscle atrophy. In fact, increased caspase-12 expression was detected in the skeletal muscle of old rats[13,60,82], suggesting a role for the ER stress response in sarcopenia.

THE INVOLVEMENT OF MYOCYTE APOPTOSIS IN HUMAN SARCOPENIA

Although studies in animal models appear to support a key role for apoptosis in age-related muscle atrophy, evidence in humans is still lacking. To date, only three reports have been published examining the occurrence of myocyte apoptosis in older persons[11,16,83]. However, none of these studies has examined either specific biochemical pathways of apoptosis or functional implications. Interestingly, Whitman et al.[16] found that the increase in apoptotic myonuclei in older persons was not accompanied by changes in caspase-3 and -7 activity. This may indicate that the caspase-independent pathway of myonuclear apoptosis might be selectively activated during aging, as reported by Marzetti et al.[18], in aged rats. In support of this hypothesis, Park et al.[84] recently found increased expression of AIF in the semi-tendinosus muscle of middle-aged men relative to younger controls, whereas no age-related changes in caspase-3 levels were detected. Clearly, further research is required in order to understand the signaling transduction pathways of myonuclear apoptosis in older human subjects, as well as their contribution to sarcopenia and physical function decline.

CONCLUSIONS

Growing evidence indicates that an age-related deregulation of apoptosis may contribute to the onset and progression of sarcopenia. However, several critical issues are still unsolved. First, a definite mechanistic link between myonuclear apoptosis and muscle loss has yet to be established. Furthermore, the relative contribution of the various apoptotic pathways to sarcopenia is still unresolved. Likewise, it is unclear whether specific signaling transduction pathways are selectively activated during the progression of muscle wasting. Finally, the role of apoptosis in human sarcopenia warrants further investigation. In particular, a deeper understanding is necessary regarding the relationship between the severity of myonuclear apoptosis, muscle atrophy, and physical function decline in older persons. By answering
these fundamental research questions, novel biomarkers of sarcopenia will be provided that may be utilized for the development of new therapeutic remedies for sarcopenia and functional decline in old age.

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