Abstract
Mitogen-Activated Protein Kinase (MAPK) pathway is a signal transduction pathway that functions in a wide range of physiological and pathophysiological cellular events including cell proliferation, differentiation, apoptosis, migration, inflammation, metabolic disorders and diseases. In skeletal muscle, it plays an essential role in muscle fiber specialization, muscle mass maintenance, damage induced muscle regeneration and muscle diseases. This review provides an overview of MAPK pathway and its pathophysiological role in skeletal muscle diseases with a primary focus on muscular dystrophy and atrophy.

MAPK Pathway in Skeletal Muscle Diseases
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MAPK Pathway

MAPK pathway consists of at least 4 subfamilies that include extracellular signal-regulated kinase 1 and 2 (ERK 1/2), p38α/β/γ/δ MAPK, c-Jun NH2-terminal kinases 1, 2, and 3 (JNK 1/2/3), and ERK5 [1-4]. MAPKs are a family of protein phosphorylating enzymes [5] that regulate diverse aspects of cellular responses in physiology, immunology, neurobiology, and energy metabolism [6]. Cellular activities regulated by MAPKs include proliferation, differentiation, apoptosis, motility [7,8], stress responses, inflammation, and innate immunity [9-11]. MAPK functions through transcriptional activation and posttranslational modification on the downstream substrates. Specifically, MAPKs are involved in the production of antimicrobial factors, cytokines, chemokines, and other inflammatory mediatory factors [12]. In the central nervous system, MAPKs are required for proper neuronal axonal development [13]. MAPKs also play a critical part in energy metabolism through modulating lipid metabolism [14-16] and skeletal muscle growth and fiber type [6]. Given the fact that MAPKs play such a fundamental and integral role in a broad range of biological processes, interference of this pathway and their downstream effector proteins may have detrimental consequences leading to either metabolic disorder or diseases. This review mainly focuses on the functional role of MAPK pathway in skeletal muscle diseases.

MAPKs phosphorylate serine and threonine residues on their substrates [8] which may include protein kinases, phospholipids, transcription factors and even cytoskeletal proteins [17]. MAPKs are activated through a three-tiered phosphorylation relay that transmits signals from the cell surface into biochemical responses: MAPKs are phosphorylated and thus activated by upstream Mitogen-Activated Protein Kinase Kinases (MKKs) which are phosphorylated and activated by their upstream MKK kinases (MKKKs) [18,19]. Given the fact that each tier of phosphorylation consists of multiple kinases, one may question how an extracellular signal is decoded and transduced into a specific cellular response in a temporal and spatial manner. The complexity of regulation of the MAPKs provides a platform on which diverse signals converge and are precisely deciphered to generate a specific signal that fine-tunes the signaling network within a cell. On the other hand, an activated MAPK signal has to be curtailed in a timely manner so that cell will not overreact to a stimulus so to maintain metabolic homeostasis. One mechanism involves the dephosphorylation and thus inactivation of MAPKs by the MAPK phosphatases (MKPs) [5]. MKPs, also known as dual-specificity protein phosphatases (DUSPs), are a group of 10 catalytically active protein tyrosine phosphatases [9,20,21]. MKPs inactivate MAPKs through dephosphorylation of MAPKs on regulatory threonine and tyrosine residues. Though the substrates of the MKPs are to some extent overlapping, MKPs achieve their substrate specificity through binding affinity and sub-cellular localization [22].

MAPKs and Muscular Dystrophy
Muscular dystrophy (MD) is a group of over 30 genetic diseases that cause degeneration of skeletal muscles during voluntary movement and overall progressive weakness in muscle strength [23]. These diseases can be classified into nine major types including Duchenne, Becker, Myotonic, Congenital, Emery-Dreifuss, Facioscapulohumeral (FSHD), Limb-girdle,
Distal, and Oculopharyngeal. The most common and severe muscular dystrophy is Duchenne muscular dystrophy (DMD) which accounts for greater than 50% of all disease cases. DMD is a prevalent muscular dystrophy in humans, affecting 1:3,500 males in the United States [24]. Naturally occurring or spontaneous mutations in the DMD gene has been reported in Canine breeds such as Golden Retriever [25], German Short-Haired Pointer [26], and Cavalier King Charles[27]. Canine studies of X-linked DMD revealed that pups from an affected male were stunted on growth and showed progressive weakness and gait abnormalities which lead to muscle atrophy more over mine over they also exhibited fibrosis and contractures by 6 months of age. Smaller breeds are less affected than large crosses [28]. In feline DMD models, the disorder is referred to as hypertrophic feline muscular dystrophy (HFMD) due to a loss of dystrophin in skeletal and heart muscles, skeletal muscle hypertrophy occurs especially in the tongue muscles [29].

The role of MAPKs in dystrophy is not completely understood, and the inconclusive evidence from different diseases and experimental settings confounds a definitive role of MAPKs in the pathogenesis of muscular dystrophy [22]. Interestingly, a mechanistic link has been built between the dystrophin-glycoprotein complex (DGC), a structure that is dysfunctional in muscular dystrophy and the MAPK pathway. The DGC not only provides structural support to cells, but also bridges extracellular stimuli and intracellular reaction through specific physical interactions with intracellular proteins [30]. Some of these proteins, such as adapter protein Grb2, are involved in the MAPK signal transduction, which suggests that DGC may function to transduce signals through the MAPKs to maintain skeletal muscle fiber viability [22]. Our recent work on MAPK phosphatases (MKPs) has added another layer of regulation of MAPKs in Duchenne muscular dystrophy. MKP-1 is a nuclear phosphatase that dephosphorylates p38 MAPK, JNK, and ERK1/2 not only in skeletal muscle but also in the immune, metabolic, and nervous system as well [9,31-34]. In the cardiotoxin-induced skeletal muscle regeneration, knockout of MKP-1 impaired the recovery of the damaged muscle likely through the effect of p38 MAPK pathways on muscle stem cell function [33], since the precocious differentiation of muscle stem cells caused by enhanced p38 MAPK may slow down the injured muscle from returning to a normal muscleulature [33]. Interestingly, when another MKP family member MKP-5 was knocked out, muscle regeneration following damage was greatly improved [35]. When MKP-5 was knocked out in the mdx mouse, an animal model for Duchenne muscular dystrophy, the dystrophinopath was ameliorated and muscle functions were restored [35]. The obvious discrepancy between MKP-1 and MKP-5 knockout studies can be explained by the fact that MKP-5 is both nuclear and cytosolic, though its substrates include p38 MAPK, JNK and ERK1/2, it functions on distinct pools of MAPKs from MKP-1 due to their subcellular localization [35]. Together, these findings suggest that MAPKs may play pivotal roles in the pathogenesis of muscular dystrophy, however, the definitive role of each MAPK in the network of signaling pathways that contribute to the progression of the diseases remains to be further investigated.

**MAPKs and Muscle Atrophy**

Skeletal muscle atrophy is characterized by decrease in cell size through organelle, cytoplasm, and protein loss [36]. Biochemically, muscle atrophy is a result of negative balance between protein synthesis and degradation. This imbalance in protein turnover occurs during fasting, disuse of selected skeletal muscles, and a plethora of systemic diseases such as cancer, diabetes mellitus, AIDS, sepsis, and Cushing's Syndrome [37].

Traditionally, it was thought there are two types of muscular atrophy in skeletal muscles: neurogenic atrophy and muscle atrophy through ubiquitin-dependent proteolysis [38]. Healthy muscle maintains its size, structure, and function through innervation of motor neurons to skeletal muscle fibers, and through balance of protein accretion. Upon disruption of the nervous system, denervation occurs and neurogenic muscle atrophy ensues [38]. Such disorders are seen in amyotrophic lateral sclerosis (ALS), Guillain-Barré syndrome, polio, and polyneuropathy which ultimately leads to paralysis [38]. In contrast, ubiquitin-dependent proteolysis occurs during increased rate of protein degradation through the ubiquitin-proteasome pathway [39] and transcriptional adaptations called “atrophy program” [40]. Recently, emerging evidence indicates that there is another equally important protein breakdown machinery in skeletal muscle called autophagy-lysosome system. In mammals, there are three different types: macroautophagy, chaperone-mediated autophagy (CMA), and microautophagy [41], though the mechanistic aspect of each category remains yet to be further defined.

Due to the fact that muscle atrophy occurs in a large number of diseases and disorders in both humans and animals, it is a focal point of interest in basic research and clinical investigation. In agriculture and veterinary medicine, muscle atrophy caused by malnutrition is the culprit of inefficiently growing animals and alters meat quality. Atrophy, along with caloric restriction, induces cytoskeletal remodeling of muscle fibers which may contribute to lower quality less tender meat product [42]. Aging also proves to be a factor for muscle atrophy. In the hindlimbs of older dogs, a growing insensitivity to adenosine triphosphate (ATP) occurs and muscle fibers undergo fibrosis which weakens muscle [43].

Among the various pathways that participate in the pathogenesis of muscle atrophy, MAPKs play a critical role. There are several lines of evidence that support this notion. First, when MKP-1, a phosphatase that dephosphorylates and thus inactivate MAPKs, was overexpressed in skeletal muscle fibers, it induced profound muscle fiber atrophy possibly through the ubiquitin-proteasome pathways [44]. This atrophic effect may be conferred through ERK1/2 signaling pathway, yet it does not exclude the possibility of a combined effect of ERK, JNK, and p38 MAPK pathways [44]. Second, given the fact that MAPKs play integral roles in innate immunity [45], it is reasonable to speculate that MAPK-mediated cytokine secretion may serve as an indirect effect on skeletal muscle atrophy as seen in chronic inflammation diseases. Specifically, activation of JNK and p38 MAPKs are involved in cytokine-induced...
molecules. The ERK1/2 pathway, however, counteracts muscle wasting through enhanced protein synthesis by its control of ribosomal RNA gene expression [46]. Finally, diverse regimens of exercises have been developed to counter age-induced skeletal muscle atrophy in humans and animals. Analysis of exercise with aged animals has revealed that centrifugal activity, which includes weight training, may slow the weakening of muscles [47]. This type of exercise is not commonly employed in companion animal rehabilitation but water resistance activities may improve the quality of life in elderly animals [43]. Analysis of the MAPK activation during different exercise protocols revealed that ERK1/2 phosphorylation increases in nearly all forms of exercises, whereas p38 MAPK and JNK are activated in specific exercise protocols [48-51]. Together, these findings demonstrate that MAPK signal transduction pathway may serve as a potential therapeutic target to combat skeletal muscle atrophy in human and animals.

Future Perspectives

A wealth of data supports a critical role of the MAPKs in the pathogenesis of skeletal muscle diseases. MAPKs are activated and inactivated in a temporal and spatial manner through their upstream kinases and a family of phosphatases to achieve specific biological responses. Research in this area has begun to shed a light on the complexity and hierarchy of the MAPK regulation and how this signaling pathway is integrated into the intertwined signaling network system within a cell. Knockout animal models targeting specific MAPKs and their isoforms are useful tools to define the role of each individual MAPK in a specific muscle disease. Although mice are an excellent model to study the mechanistic insight of the MAPK in muscle diseases, other animal models are valuable especially in veterinary medicine and agriculture. Large animals may be more challenging for gene manipulation, yet the work in pigs has brought promise for the use of large animals to study human diseases. Animals with spontaneous mutations in genes of interest are another valuable tool to study muscle diseases. In all, research in the MAPK pathway will undoubtedly broaden our understanding of the etiology and pathogenesis of muscle disease and the molecular signatures that define the pathological process. All of these efforts are targeted to develop therapeutic means to improve the health of humans and animals. Given the fact that MAPK signaling pathway is ubiquitous in all cell types, drugs that inhibit MAPK signaling may have undesired effects. As such, identification of the downstream effectors of the MAPKs that convey MAPK effects in a specific disease will be an optimal choice.

References


