INTRODUCTION

Skeletal muscle is the largest tissue in the animal body, accounting for 40%–50% of body weight and playing an essential role in motion. One of the most distinctive features of skeletal muscle is its plasticity, as it can increase or decrease its mass (hypertrophy or atrophy, respectively) in response to environmental factors such as growth, aging, and exercise. Muscle remodeling is caused by increasing or decreasing the cross-sectional area of muscle fibers (myofibers), as an increase or decrease in the number of myofibers in myofibers directly leads to muscle hypertrophy or atrophy, respectively. The extracellular growth factors that induce muscle hypertrophy or atrophy, such as myostatin (growth differentiation factor-8, GDF-8; McPherron, Lawler, & Lee, 1997) and insulin-like growth factor-1 (Barton-Davis, Shoturma, Musaro, Rosenthal, & Sweeney, 1998) have received considerable attention. However, little attention has been paid to the mechanisms underlying the assembly and structural maintenance of myofibrils. In this review, to better understand these fundamental issues regarding hypertrophy and atrophy in skeletal muscle, we focus on the thick filament and its main component, myosin. First, we describe the structure of myosin, a main component of the thick filament, in thick filament formation and the dynamics of myosin in skeletal muscle cells. Changes in the number of myofibrils in myofibers can cause muscle hypertrophy or atrophy. Therefore, it is important to understand the fundamental mechanisms by which myofibers control myofibril formation at the molecular level to develop approaches that effectively enhance muscle growth in animals.
approximately 0.1 mm in diameter, which varies in length, and is filled with numerous myofibrils consisting of tandemly aligned sarcomeres. The sarcomere is the smallest basic contractile unit and is the most regular macromolecular protein complex (Figure 1). Thus, skeletal muscle fibers are entirely dedicated to generating force.

The sarcomere has a highly ordered semi-crystal structure consisting of four main components: Z-bands, thin filaments, thick filaments, and connectin/titin. The size of each sarcomere component is highly regulated, and each component is formed of a substantial number of myofibrillar proteins and their associated proteins (Henderson, Gomez, Novak, Mi-Mi, & Gregorio, 2017; Sweeney & Hammers, 2018). The Z-band is a complex protein network which coordinates the contraction generated by individual sarcomeres. Its major components are sarcomeric-α-actinin, (s-α-actinin), α-actin, T-cap, C-terminus of nebulin, and N-terminus of connectin/titin (Gregorio, Granzier, Sorimachi, & Labeit, 1999). The thickness of the Z-band is 30–120 nm depending on the type of muscle fiber (Engel & Franzini-Armstrong, 2004). The thin filament consists of α-actin, tropomyosin, the tropinin complex, tropomodulin, and nebulin. Although the length of the thin filament is known to vary depending on developmental stage (Ohtsuki, 1979), Gokhin et al. recently revealed that the thin filament is longer in slow type myofibers than in fast type myofibers (Gokhin et al., 2012). The bipolar thick filament is formed of approximately 300 myosin molecules and their associated proteins (Craig & Offer, 1976; Luther, Munro, & Squire, 1981). It has a strictly uniform dimension of 1.6 µm in length and 15 nm in diameter, which is independent of the type of muscle fiber (Huxley, 1963). The thick filament is located at the center of the sarcomere as the giant elastic protein connectin/titin spans half sarcomere along the thick filaments, linking the Z-band and the M-lines (Labeit & Kolmerer, 1995; Maruyama, 1976; Wang, McClure, & Tu, 1979). Connectin/titin provides the sarcomere not only with passive tension (Horowits, Kempner, Bisher, & Podolsky, 1986; Trombitas et al., 2000) but also acts as a scaffold for associated proteins, which are implicated in connectin/titin-based signaling pathways (Miller et al., 2003; Ojima et al., 2010).

1.2 | Myosin: One of the main myofibrillar proteins in skeletal muscles

Myosin is the most abundant myofibrillar protein, accounting for more than 40% of the myofibrillar proteins in skeletal muscle (Yates & Greaser, 1983). Myosins are a large superfamily of motor proteins which play key roles in the fundamental cellular functions of nonmuscle and muscle cells, such as locomotion, cytokinesis, and contraction (Engel & Franzini-Armstrong, 2004; Sellers, 2000). The sarcomeric myosins specifically expressed in skeletal and cardiac muscle cells to generate contractile force are categorized as either conventional myosins or class II myosins.

A single myosin molecule consists of two myosin heavy chains (MYHs), which interact with two pairs of myosin regulatory light chains and myosin essential light chains to form a hexamer (Weeds & Lowey, 1971). Sarcomeric myosin is generally divided into the head domain and the rod domain (Figure 2). The head domain [subfragment-1 (S1)] is located at the N-terminus of myosin and is the motor domain that generates force by ATP hydrolysis when myosin interacts with actin-containing thin filaments. Thus, S1 determines the biochemical and physiological functions of myosin, such as ATPase activity and actin-filament velocity (Resnicow, Deacon, Warrick, Spudich, & Leinwand, 2010). The rod domain is proteolytically cleaved into subfragment-2 (S2) and light meromyosin (LMM). S2 has

**FIGURE 1** Structure of myofibrils in the myofiber. Skeletal muscle tissue consists of bundles of myofibers. Each myofiber contains millions of myofibrils that are comprised of longitudinally aligned sarcomeres. The sarcomere is the minimum contractile unit, which consists of Z-bands, thin filaments, thick filaments and connectin/titin. Z-bands are indicated by “Z”. A transmission electron micrograph of the sarcomere structure is shown at the bottom of panel. Connectin/titin is omitted in a model of "Myofibril". The figure was created using BioRender
Sarcomeric myosin has the remarkable ability to form highly organized bipolar thick filaments in myofibrils. Extensive biochemical studies have demonstrated that thick filament formation is mediated by the distribution of charge along LMM (Atkinson & Stewart, 1991; Sohn et al., 1997). Myosin forms filaments in an antiparallel fashion at the center of the thick filament, while myosin forms filaments in a parallel way in the rest of the thick filament. Consequently, a bipolar thick filament is formed, leaving a central bare zone in the middle.

Which myosin domain is critical for forming the thick filament in skeletal muscle cells? To answer this question, we evaluated whether the thick filament is formed by exogenously expressed MYH deletion variants in cultured skeletal muscle cells. S1 is dispensable for thick filament formation (Ojima, Oe, et al., 2015b), consistent with the findings of other studies which showed that the thick filament is formed without S1 in Drosophila melanogaster and cultured muscle cells (Cripps, Suggs, & Bernstein, 1999; Thompson, Buvoli, Buvoli, & Leinwand, 2012). S2-LMM is perfectly incorporated into the entire thick filament, while the expression of S2 alone does not form any structure with S2 diffusing into the cytoplasm. Furthermore, LMM is incorporated into the restricted region of the thick filament, in the vicinity of the central bare zone. These results demonstrate that the LMM domain is not sufficient for thick filament formation in cultured muscle cells although it is essential for forming myosin filaments or paracrystals in vitro (Sohn et al., 1997). Interestingly, S2-LMM with N-terminal or C-terminal deletions are assembled into the M-line neighboring region but not into the entire thick filament. These results suggest that the S2 N-terminus region and the LMM C-terminus region (also known as the tail piece) are necessary for incorporation of myosin into the entire thick filament or for thick filament stabilization (Ojima, Oe, et al., 2015b). However, the molecular structure of the thick filament has not yet been completely determined in vertebrates (Irving, 2017).

In nonmuscle COS7 cells, exogenously expressed MYH forms a filamentous structure or aggregates (Moncman, Rindt, Robbins, & Winkelman, 1993; Ojima, Oe, et al., 2015b; Vikstrom et al., 1997), however, the authentic thick filament does not form. Therefore, it is likely that the myogenic intracellular environment, including unknown myocytosolic factors and/or other myofibrillar proteins such as connectin/titin (Myhre, Hills, Prill, Wohlgemuth, & Pilgrim, 2014; Tonino et al., 2017) is required to generate properly assembled-thick filaments.

1.4 | Replacement of myosin in the thick filament

How often are myosin molecules exchanged in the thick filaments? To address this question, we used fluorescence recovery after photobleaching (FRAP), which determines the dynamics of fluorescently labeled-molecules in living cells (Wang et al., 2005). FRAP experiments revealed that green fluorescence protein (GFP)-tagged MYH molecules are actively and continuously substituted in the thick filaments of cultured myotubes (Ojima, Ichimura, Yasukawa, Wakamatsu, & Nishimura, 2015a). The simultaneous uptake and release of myosin was observed in myofibrils while the half-life of GFP-MYH replacing myosin in the thick filament was estimated to be 3–4 hr (Ojima, Ichimura, et al., 2015a). Given that the half-life of myosin replacement is 3 hr, more than 90% of the myosin molecules in a single thick filament will be replaced within 12 hr. Myosin exchange is maintained in every myofibril of every myotube as long as myotubes are alive since the pioneering works using radioactive isotopes demonstrated that the half-life of the MYH protein turnover rate, i.e., from synthesis to degradation, is approximately 6 days in vivo and in vitro (McManus & Mueller, 1966; Zak, Martin, Prior, & Rabinowitz, 1977; Rubinstein, Chi, & Holtzer, 1976).

The exchange rate of myofibrillar proteins in other sarcomere components has also been determined by other research groups. The half-lives of thin filament components (α-actin and tropomyosin) and Z-band components (s-α-actinin) have been reported as minutes (Littlefield, Almenar-Queralt, & Fowler, 2001; Wang, Fan, Dube, Sanger, & Sanger, 2014; Wang et al., 2005), whereas the half-life of the giant molecule connectin/titin is approximately 2 hr (da Silva Lopes, Pietas, Radke, & Gotthardt, 2011). Likewise, the half-lives of thick filament associated-proteins, such as myosin binding protein C1 and myomesin, are approximately 2 hr (Ojima, Ichimura, et al., 2015a). The different exchange rates of each myofibrillar protein may, in part, reflect their molecular size, since the rate of protein diffusion is dependent on molecular mass (Papadopoulos, Jurgens, & Gros, 2000). In fact, connectin/titin (~3 MDa) and myosin (~500 KDa) are relatively large molecules compared to α-actin (~42 KDa), tropomyosin dimers (~66 KDa), and s-α-actinin dimers (~200 KDa). Besides molecular size, the structural complexity of the thick filament may account for the slow myosin exchange rate in skeletal muscle cells. In nonmuscle cells, approximately 30 nonmuscle

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FIGURE 2 Schematic model of sarcomeric myosin. Sarcomeric myosin is formed by a pair of myosin heavy chains, essential light chains, and regulatory light chains. The N-terminus of myosin is S1, referred to as the head, which contains nucleotide and actin binding sites. The rest of myosin is the rod, which is further divided into two parts, S2 and LMM, which form an α-helical coiled-coil structure.
myosin molecules form a bipolar filament. FRAP experiments revealed that the half-life of nonmuscle myosin exchange is less than one minute, indicating that myosin substitution is more rapid in nonmuscle myosin filaments than in the thick filaments of myofibrils (Hu et al., 2017). The molecular masses of these two distinct myosins (nonmuscle and sarcomeric myosins) are almost identical, however, the number of myosin molecules is 10 times higher in a single thick filament than in a single nonmuscle filament. Furthermore, thick filaments contain myosin-associated proteins including myosin binding protein C, myomesin, and connectin/titin, which help to pack the filament together. Therefore, the thick filament in striated muscles is larger, more complicated and has a higher-order structure than the nonmuscle bipolar myosin filaments (Dasbiswas, Hu, Schnorrer, Safran, & Bershadsky, 2018). The intricate structure of thick filaments may reduce the exchange rate of myosin molecules in skeletal muscle cells.

1.5 | Sources of myosin for replacement in the thick filament

As myosin molecules are continuously replaced in the thick filament of myofibrils, what are the sources of these replacement myosin molecules? The primary source is de novo synthesized myosin molecules, since newly synthesized myosin molecules are immediately incorporated into thick filaments (Isaacs & Fulton, 1987). In fact, blocking myosin biosynthesis using a translation inhibitor significantly slows the substitution rate of myosin in myofibrils (Ojima, Ichimura, et al., 2015a). The secondary source is cytosolic myosin that is not incorporated into the thick filament. Reducing the level of cytosolic myosin using Streptolysin-O, which perforates the plasma membrane to release cytosolic components such as myosin, leads to a lower myosin exchange rate (Ojima et al., 2017). These results also demonstrate that myosin allocated for replacement is present in the myocytosol, although myosin forms aggregates or filaments in physiological ionic conditions in vitro (Perry, 1955). In other words, the myosin exchange rate in thick filaments declines when myosin supply sources, including de novo synthesized myosin and pooled cytosolic myosin, are depleted.

1.6 | Modulation of the myosin replacement rate by myosin chaperones

Heat shock proteins (HSP) are a family of molecular chaperones which assist in protein folding and remodeling, and are rapidly upregulated in response to cellular stressors. HSP90 is strongly implicated in thick filament organization (Smith, Carland, Guo, & Bernstein, 2014), with mutations or disrupted chaperone activity causing the disassembly of thick filaments in myofibrils (Du, Li, Bian, & Zhong, 2008; He, Liu, Tian, & Du, 2015). We investigated the effects of HSP90 on myosin substitution in thick filaments. HSP90 overexpression increased the rate of myosin replacement, while inhibiting its chaperone activity reduced myosin substitution in myofibrils, indicating that HSP90 chaperone activity is closely associated with myosin replacement (Ojima et al., 2018). As HSP90 was originally found to function as a chaperone to ensure the correct folding and conformational structure of myosin S1 (Srikakulam, Liu, & Winkelmann, 2008; Srikakulam & Winkelmann, 2004), its overexpression may increase the rate of myosin folding following translation, increasing the amount of myosin available for exchange. Upregulated MYH gene expression is also observed in myotubes overexpressing HSP90, alongside an increase in myosin content, however, the molecular mechanisms underlying the upregulation of MYH expression by HSP90 remain unclear.

What are the conditions in muscle cells overexpressing HSP90? The induction of HSP expression is an important response of skeletal muscle cells to physical exercise-induced stress (Harris, Mitchell, Sood, Webb, & Venema, 2008; Locke, Noble, & Atkinson, 1990). Myofibrillar protein synthesis is also elevated in exercised skeletal muscle, supplying newly synthesized myofibrillar proteins to replace those damaged by exercise-induced stress and to build new myofibrils in hypertrophy (Camera et al., 2015; Di Donato et al., 2014). Considering these findings, HSP90 overexpression may partly mimic the intracellular environment of exercised muscle, leading to the upregulation of myosin transcription, translation, and content in muscle cells. As a result, the myosin replacement rate may increase rapidly in myotubes overexpressing HSP90.

Uncoordinated mutant number 45 (UNC45) also functions as a chaperone for myosin (Barral, Hutagalung, Brinker, Hartl, & Epstein, 2002; Lee, Melkani, & Bernstein, 2014; Liu, Srikakulam, & Winkelmann, 2008; Srikakulam et al., 2008). UNC-45 chain formation provides multiple HSP90 and myosin binding sites, with the correct spacing for myosin heads on thick filaments (Gazda et al., 2013). Although myofibril organization is impaired in Caenorhabditis elegans when UNC45 is overexpressed or knocked down (Landsverk et al., 2007), the myosin replacement rate is not affected in myotubes overexpressing UNC45b (Ojima et al., 2018).

1.7 | Proposed schematic model of myosin dynamics

Our proposed model of myosin dynamics in myofibers is illustrated in Figure 3. Newly synthesized myosin is immediately incorporated into the thick filament following translation (Isaacs & Fulton, 1987) while myosin is simultaneously released from the thick filament (Ojima, Ichimura, et al., 2015a). Myosin is pooled in the myocytosol, providing another source of replacement myosin. Although myosin forms aggregate under physiological ionic conditions in vitro (Perry, 1955), myosin is protected from aggregate formation in the myocytosol by an unknown mechanism. HSP90 induces the expression of MYH and subsequently increases the myosin content of the myocytosol, leading to rapid myosin replacement in thick filaments. Although damaged myosin is degraded by the ubiquitin-proteasome system (Cohen et al., 2009; Flieltz et al., 2007), it is not yet known how and where damaged myosin is selected. The mechanisms of the myosin degradation system in this model should be investigated in future studies.

2 | CONCLUSION

Based on our current and previous experiments, we have described the role and the dynamics of myosin in the formation and
maintenance of thick filament structure. Despite the collection of skeletal muscle data by multiple researchers, the dynamic control of myofibrillar proteins such as myosin is still poorly understood. How are hundreds of myofibrillar and myofibrillar-associated proteins assembled into the highly ordered sarcomere structure? How do myofibers regulate, maintain, and degrade the myofibril structure at the molecular level? Importantly, these processes proceed without impeding muscle contraction. While sarcomeres appear to have a hard, rigid structure when observed using electron microscopy, each myofibrillar protein is continuously replaced. This is a crucial field for future research in animal science and life science in order to understand the fundamental regulation of plasticity and remodeling in the skeletal muscle of animals.

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