Peripheral muscle fatigue is a common experience in daily life. Every individual at some point in their life has realized the inability to maintain muscle contraction, a phenomenon known as fatigue. Interestingly, neurological patients with peripheral sequelae such as spastic muscle contraction are able to remain in a pattern of muscle contraction for prolonged periods. The effects of laser therapy are already recognized in muscle contraction to delay skeletal muscle fatigue, prolong physical activity, and reduce delayed onset muscle soreness. However, the effects of photobiomodulation on neurological patients with muscular spasticity are still not well established. The present literature review seeks to recognize articles about the application of laser irradiation, also known as photobiomodulation, to patients with muscle fatigue and/or spastic palsy. To perform a literature review, we used the systematic review methodology for the literature search. The following keywords were searched: (skeletal muscle fatigue) AND (spastic patients) AND (low-level laser therapy OR low intensity laser therapy OR low energy laser therapy OR LLLT OR LILT OR LELT OR infrared laser OR IR laser OR diode laser), and these were used for search on the following databases: PubMed, Embase, Web of Science, BIREME, Scopus, and SciELO. Besides that, a literature review concerning on muscle physiology, fatigue, and LLLT was made. No language filter was applied, and altogether, 689 papers were identified. A group of 3 physiotherapists and 1 pharmaceutical scientist performed the literature review, and every exclusion was confirmed by at least two reviewers. After inclusion and exclusion criteria, 128 studies were included in this review. Conclusion, the LLLT can contribute to the recovery of spastic patients and muscles in fatigue. However, the real effect of laser photobiomodulation on muscle spasticity remains to be established. Only a much reduced number of clinical trials have been performed with a small number of participants. There is a lack of clinical trials from different research groups that could help to understand and elucidate the effects of laser in prolonged muscle contraction in spastic palsy.

1. Introduction

Looking into the past, more specifically since the 1960s, it was consistently seen that low-level laser therapy (LLLT) could present positive effects on the muscle-skeletal system and related diseases such as joint inflammation, sports injuries, muscle fatigue, and lower back and neck pain among other conditions [1]. Photobiomodulation has been covering new areas of study, and with them, there is a range of possibilities for spastic patients, to reduce or eliminate muscle fatigue in some cases, especially in patients with traumatic brain injury, for example [2].

Previously, photomodulation was called a "low-level laser therapy" or LLLT, including light from light-emitting diodes (LEDs), lasers, and other light sources with wavelengths ranging from visible to infrared. It has also become an
increasingly mainstream treatment for muscle fatigue, especially in the areas of physical medicine and rehabilitation, in patients with a type of degenerative disease [3].

It is well known that there are dose responses to simultaneous applications of several different wavelengths in some cases, for example, to kill cancer cells and promote significantly faster healing in chronic and diabetic wounds and sports injuries, reduction of pain in arthritic joints and back and neck, reduction of inflammation by increasing the number of fibroblasts and myofibroblast, induction of the accumulation of collagen, and decrease of fatigue in the muscles in spastic patients [4]. Additionally, recent studies indicate that photobiomodulation can also become promising in clinical cases of patients suffering from myocardial infarction, stroke, brain injuries caused by trauma, or degenerative and spinal cord injury, with the last three triggering injuries in motor neurons and ending up directly affecting the muscles, causing spasms and finally fatigue and pain [5].

The wavelengths for photobiomodulation are more regularly used in the 500–700 nm range for treating superficial healing, while wavelengths between 800 and 1000 nm are used for deeper tissues and skeletal muscle fatigue [4].

This review focuses on the efficacy of photobiomodulation in muscle fatigue in spastic patients, its mechanisms of action, and how its efficacy may be increased by using multiple wavelengths. The review examines research related to the induction of muscle fatigue signaling pathways and the elimination of pain by the application of photobiomodulation and their mechanisms of action. Photobiomodulation application is making clear headway toward the development of a reliable and successful set of irradiation parameters for inducing significantly faster and more complete healing of illness related to muscles that does not respond to conventional treatment [3]. However, further work is required to optimize its therapeutic value to determine the integrative nature of fatigue in spastic patients.

2. Methodology

The following bibliographic databases were searched in MEDLINE via PubMed, Embase, Web of Science, BIREME, Scopus, and SciELO. The search strategy was (skeletal muscle fatigue) AND (spastic patients) AND (low-level laser therapy OR low intensity laser therapy OR low energy laser therapy OR LLLT OR LILT OR LELT OR infrared laser OR IR laser OR diode laser). No language filter was applied, and altogether, 689 papers were identified. A group of 3 physiotherapists and 01 pharmaceutical scientist performed the literature review, and each exclusion was confirmed by at least two reviewers. After inclusion and exclusion criteria, 128 studies were included in this review.

Reviewers independently identified titles and abstracts relevant to applying LLLT to patients suffering from muscle fatigue and/or spastic. Full texts of the published articles, unpublished articles, and unpublished data of completely finished and analyzed studies were included. The reference list of full-text articles was also reviewed. Figure 1 illustrates the selection process for including studies.

3. Homeostasis

In the human body, one of the most plastic and dynamic tissues is the skeletal muscle. There is about 40% of the human total body weight in humans that is made up of skeletal muscle and is composed of 50–75% of all body proteins and accounts for 30–50% of the whole-body protein turnover. The composition of the muscle is mainly of water (75%), protein (20%), and other substances including inorganic salts, minerals, fats, and carbohydrates (5%) [6].

The muscle mass depends on the conditional nutrition, hormonal condition, and the balance between physical activity/exercise and illness or injury. In addition, the balance between protein synthesis and degradation is extremely sensitive to factors that deregulate the different cellular compartments (structural, contractile, and regulatory); for this reason, it has received significant scientific attention because of their important contribution to mobility, contractility capacity, and functioning [7].

Skeletal muscle contributes significantly to multiple bodily functions, since mechanical function for converting chemical energy into mechanical energy to generate force and power, walking motion, or stand, besides allowing participation in social and sportive settings, maintains a possibility of functional independence. From a metabolic point of view, several skeletal muscle functions include a contribution to basal energy metabolism, like storage for important substances such as amino acids and carbohydrates used by other tissues such as the skin, brain, and heart, and the production of heat for the maintenance of core temperature, and it is responsible for the consumption of the majority of oxygen and energy during physical activity and exercise [8].

Furthermore, amino acid release from muscle contributes to the maintenance of blood glucose levels during conditions...
of starvation, in that case, a person with a reduced muscle mass impairs the body’s ability to respond to stress and chronic illness [9].

The architecture of skeletal muscle is characterized by myofi bers or muscle cells and associated connective tissue. The structure of muscle is characterized by the number and size of individual muscle fi bers, but any pathological infi ltration by fat and connective tissue may alter this relationship [10, 11].

Observing the cellular level, the muscle fi bers are multinucleated and postmitotic, and some part of each nucleus within a muscle fi ber controls the type of protein synthesized in that specific region of the cell; these regions are known as nuclear domains and have a highly regulated, but not uniform, size [12]. Protein expression found in the adjacent domains of a single fi ber appears to be coordinated in such a way that the type of protein, in this case, the myosin produced, is almost the same across the length of the fi ber [13].

In the human body, there are several cell storage sites with stem cell potential; knowing this, it is important to note that skeletal muscle has cells of this nature, which are known as satellite cells and are located between the sarcolemma and the basal lamina and contribute to muscle growth, repair, and regeneration [12, 14]. When these cells are activated by myogenic factors, satellite cells proliferate and differentiate into new muscle fi bers and repair the local injury.

The functional unit of the muscle is divided into compatibilities and is separated by a dense connective tissue called a sarcolemma. The outermost layer that contemplates the bundles of muscular fi bers is called epimysio. The perimeter consists of the grouping of fi bers within and surrounded by another layer of connective tissue. In addition, there is a complex of several proteins associated with the sarcolemma and are usually connected to the myofilament of actin. A single-muscle fi ber has approximate dimensions of 100 µm in diameter and 1 cm in length [15].

The contractile units of skeletal muscle are composed of grouped and organized myofilaments, called sarcomeres. These myofilaments are actin and myosin proteins (70–80% of the total proteins present). Myosin is the main driver of the muscle fi ber and is present in a total of eleven sarcomere myosin genes, but only a few can be described in humans, and at least two of these genes are expressed in cardiac muscle. There are many other proteins present in the sarcomere and sarcoplasm, which act as the coupling of the processes of excitation and contraction, release of energy, structuring of the cytoskeleton, and generation of strength and power [16]. Among them, in particular, we can mention the regulatory proteins that form the calcium-dependent troponin complex, which are troponins C, I, and T. Tropomyosin plays an essential role in the activation process that leads to slippage and the creation of force, which is directly associated with the actin fi lament [17].

In addition to the proteins mentioned above, there are other proteins that contribute to the physiological and mechanical properties of muscle, titin, and nebulin [18]. The titin is able to bind to the Z disk of the sarcomere and to the myosin, to provide stability and alignment to the thick filament, and this is only possible thanks to its large width.

Nebulin proteins, on the other hand, are integrated with other proteins in the thin fi laments that contain actin, in which they help to maintain the sarcomere integrity [19] and influence the cell stiffness and tension that proportionally influence the formation of myofibrils and signaling local cell. In addition, in vitro studies suggest that titin may assist in muscle contraction and strength generation [20]. In the sarcomere, disk Z has several proteins; among them, the one that fi xes the actin myofilament is called actinin. There are other proteins that also connect the Z disk to the sarcolemma and the extracellular matrix, called desmin [20].

Sarcoplasm is composed of several cellular elements, among them, a transverse tubular system (T tubule), the sarcoplasmic reticulum, and a mitochondrial network; based on each type of fi ber, the amount of these elements can be estimated. Detailing the cellular elements above, the T tubule system is nothing more than an invagination of the sarcolemma and its important role in driving the nerve action potential into the cell is of utmost importance for local homeostasis [21]. This network of tubules is in contact with the exterior of the cell and ensures that the excitation can spread uniformly throughout the fi ber and the change in the calcium concentration was detected by the protein Dysferlin located in the membrane of the T tubule system [22].

The concept that mitochondria, in isolation, provide the necessary energy for muscle fi bers, according to some studies, is not applicable. It was seen that there is the formation of a three-dimensional network around the cell and generates the necessary energy for muscle action when oxygen is available [23]. In this way, some mitochondria remain close to the sarcolemma to decrease the time of diffusion of oxygen through the capillaries; thus, during aerobic exercise, when the oxygen demand increases, other mitochondria, located in the intermyofibrillar space, are recruited to supply the new demand [24].

It is well known that several neuromuscular pathologies, the practice of physical exercises, and even aging directly influence the structure and function of myofilaments and the surroundings. For example, training with resistance exercises induces the production of mitochondria through the activation of the coactivator-1a of the c-receptor, activated by the peroxisome proliferator [25]. The number and size of mitochondria increase with resistance training program and type of aerobic exercise. On the other hand, observing muscle aging, the sarcoplasmic reticulum shows a fragmented aspect and this impairs calcium release and muscle activation, so we can understand why there is muscle weakness in this population, specifically [26].

Due to the heterogeneity of human skeletal muscle in the biochemical, mechanical, and metabolic phenotypes of the individual fi bers, it is possible to highlight the presence of different muscle fi bers with different properties with each one specifically imposed by its respective motor neuron [27]. Through this, several physiological properties of muscle allow the plasticity capacity of the muscle according to various metabolic and mechanical demands, such as the architecture of capillary supply networks, which varies according to the type of fi ber [28].
The three basic ways of obtaining energy for muscle fiber are the stocks of ATP and CP, anaerobic glycolysis, and oxidative phosphorylation. This obtaining in the form of ATP is of extreme importance, because according to the metabolic pathway, the production of energy varies and this must supply the energy requirement of the muscle during the period and the intensity of the activity or exercise [26].

Muscle contraction is related to the extension of the development of the sarcoplasmic reticulum, so that its contraction speed is directly proportional to this development, looking from another angle, or the oxidative capacity also directly influences this speed, but it is more observed in fatigue. The most used classifications for muscle in adults are three types of fibers: type I (slow, oxidative, and fatigue resistant), IIA (fast, oxidative, and intermediate metabolic properties), and IIX (fastest, glycolytic, and fatigable) [26, 27].

In the muscle, an activity that uses the stocks of ATP and CP assists high-intensity short-duration (few seconds) activities because the reserves in muscle fibers are just few. Looking into anaerobic glycolysis, it produces ATP quickly to sustain muscle actions for a couple of minutes but the end products (H⁺ and lactate) are associated with muscle fatigue and consequently damage the muscle. Eventually, the energy obtained by oxidative phosphorylation is carried out within the mitochondrial network, which is able to supply the energy demand for an intense activity or activity lasting minutes and even hours [28–30].

Obtaining the raw material for energy production can be done through carbohydrates and free plasma fatty acids and muscle triglycerides, in addition to the use of amino acid metabolism, but the percentage obtained is small compared to the energy production total [31, 32].

4. Muscle Fatigue

Muscle fatigue is usually defined as a loss of strength or power in response to contractile activity. These can occur as a result of injury or physiological or pathological adaptations [33].

In order to understand the nature of skeletal muscle fatigue, we must take into account external factors such as injuries and disuse that somehow influence the fatigue mechanism and the internal factors, which include cell signaling, vascular function, neurophysiology, bioenergetics, and molecular mechanics [34, 35].

Observing the loss of maximum strength or capacity to generate energy during a muscle contraction, this would be a physiological response to all muscle contractions; however, in the shortest, this cannot occur [36].

The contractions can be sustained or repetitive, and the intensity of these contractions can vary from submaximal to maximum, and from a mechanical point of view, they can be concentric, eccentric, and isometric [37, 38]. That way, according to researchers, the magnitude of loss of force or power in response to fatigue in the muscle can occur in a continuous way or only for a few moments [39].

Following this line of thought, quantifying fatigue as a single continuous contraction or a series of contractions with submaximal tension, further studies are needed. However, the study of fatigue recovery proves to be valuable, in particular, for detecting and understanding low-frequency fatigue (LFF) associated with loss of strength due to low activation frequencies and failure of the excitation and contraction coupling (ECC) of skeletal muscle [40–43].

In general, the causes of fatigue vary widely, depending on the model studied, the experimental conditions, and the tasks imposed on the muscle. In in vitro studies, it is typical to manipulate the conditions to directly determine the cause(s) of the contractile failure and fatigue is quickly observed as the drop in strength or power of a single cell or group studied. In situ studies also usually use this approach, to mimic physiological conditions and compare them by proceeding in more basic ways compared to the in vitro study.

In vivo, usually, the mechanisms of muscle fatigue magnitude vary according to the animal model and the task imposed, this is called “task specificity,” and to demonstrate these mechanisms, it is common to use approaches that can cover and evaluate directly and indirectly the location(s) of failure in muscle strength and power [44–47].

5. Fatigue in Cells

Studies using a material that mimics fine muscle fibers to observe motility in vitro have provided insights into how actin (thin filament) interacting with myosin (thick filaments) is capable of generating strength, speed, and power [48–50]. From these studies, it was possible to observe that the main stages related to fatigue are the transition from the state of low strength to the state of high strength of the actin-myosin (AM) cross-bridge and the step of dissociation of adenosine diphosphate (ADP) from the cross-bridge AM. The first stage would be associated with the release of inorganic phosphate (Pi) and hydrogen (H⁺) ions; this stage is being considered as limiting because the rate of force development and the force inhibition site immediately begin to accumulate these ions and fatigue starts during the exercise, being of high or low intensity [45, 51]. It has been known for years that a high rate of ATP utilization accelerates the reaction of creatine kinase (CK) and this can be seen in the last step, the release of ADP, in which the loaded contractions limit the speed of the AM cross-bridge cycle and therefore the speed of the fiber [52].

Due to the high turnover of ATP and the increase in anaerobic metabolism, as a result of the intense contractile activity, [H⁺] also increases, causing the decrease in intracellular pH in fast glycolytic fibers (FG) from 7.0 to 6.2 and for the end resulting in the decline of cellular phosphocreatine (Pi) [45, 53]. However, other studies indicate that the increases of Pi decrease the isometric strength and fiber stiffness proportionally, indicating that the cross-bridge strength remains constant and that the strength decline is directly connected with the transition period from a bonding state of the fibers and these are also affected by the concentration of Pi and H⁺ [52, 54–56].

Adversely, studies show that a low pH in the presence of Ca²⁺ decreases the strength and this may be related to the hypothesis that a high [H⁺] limits the amount of high-strength AM cross-bridges, inhibiting the direct velocity
constant for the transition from the weak to the strong link state of the cross-bridge AM [57–59].

The Westerblad and Allen studies demonstrated that when the fast fibers of the mouse flexor digitorum brevis (FDB) are stimulated in vitro, the initial decline in strength occurs before any detectable change in intracellular \([Ca^{2+}]\) [60, 61]. The hypothesis of some authors was that the early loss of strength was mediated by increased \(\text{p}i\), which was confirmed by the observation that fibers isolated from CK knockout mice, which when stimulated, did not show an increase in \(\text{p}i\) or loss of strength during the initial phase of muscle contraction [62, 63].

In the stimulus phase, with the high contraction rate, it was observed that the initial drop in strength would be mediated by the combined effects of the increase in \(\text{p}i\) and \([H^+]\) and that the effects of these ions on the myofilaments have been shown to involve additives that involve a direct inhibition of strength and reduced sensitivity to \(Ca^{2+}\) and that at the beginning of fatigue, the amplitude of intracellular \(Ca^{2+}\) is high, causing local sensitivity to be impaired and preventing fatigue from ceasing [62–64].

However, a subsequent decline in the amount of \(Ca^{2+}\) that had been released by the sarcoplasmic reticulum (RS) and was impairing local sensitivity causes muscle fatigue as the muscle needs to recover from this uncoordinated imbalance, resulting in more loss of strength during peak saturation and less close to physiological temperatures compared to muscle temperatures that did not contract [50, 65].

It is important to note that studies have shown that the \([Ca^{2+}]\), necessary to elicit half the maximum strength in the muscle of mice, was significantly increased by 30 mM \(\text{p}i\) and that the effect was two times greater on slow fibers and was performed at temperatures of 30°C and 15°C [66]. The mechanism of sensitivity to \(Ca^{2+}\) is not yet fully understood; however, at a pH of approximately 6.2, it is partially mediated by the competitive inhibition of \(Ca^{2+}\) binding to troponin-C [67, 68].

One factor that contributes to the certain change in the ratio of \([Ca^{2+]}\) and strength is the decline in coarse filament cooperative, which is a direct result of the reduced number of high-strength cross-bridges [69–71]. In addition to the high \([H^+]\), the micromolar increases in cell ADP, due to intense contractile activity, depress the fiber speed, but the strength increases; this is probably due to the increase in the amount and type of cross-bridges in the high-strength peaks, these being cross-bridges AM-ADP-Pi, and cross-bridges AM-ADP [72, 73]. The most important role for increases in cellular ADP in eliciting fatigue seems to be related to the inhibition of the \(Ca^{2+}\) pump present in the sarcoplasmic reticulum, causing direct effects on the AM cross-bridge.

Looking at the effects of low pH in conjunction with high \(\text{p}i\) and high [ADP], on slow fibers and fast fibers, it is not entirely clear how it can influence fatigue [74–77]. However, recent studies have shown that the effects of low pH and high ADP and \(\text{p}i\) have been investigated at the molecular level using the in vitro motility assay to assess the slip speed of actin filaments without the regulatory proteins tropomyosin and troponin in myosin [78, 79].

To confirm that using single-fiber studies, the high \(\text{p}i\) had no effect on the sliding speed, while at pH 6.4, the speed of the actin filament showed a 36% reduction. The increase in ADP from a low resting value of 0.02 mM to a fatigue level of 0.3 mM caused an 18% drop in filament speed and with phosphorylation of the N-terminal region of the myosin light chain demonstrating that the inhibition increased to 34% [79, 80].

Thus, both high \([H^+]\) and ADP seem to decrease the sliding speed of the filament without load, increasing the affinity of ADP for the myosin head, thus decreasing the rate of ADP release. Besides that, the additional inhibition of speed caused by phosphorylation of the N-terminal region of the myosin regulatory light chain can be explained by the high affinity for ADP observed in phosphorylated myosin compared to nonphosphorylated myosin.

This suggests that, although high ADP can reduce the maximum shortening speed without load, ADP may have little or no effect on fiber speeds during peak power generation at the high intensity of force. However, it is worth mentioning that phosphorylation in the N-terminal region of the myosin light chain reduces the inhibitory effects of conditions similar to fatigue on the decrease in speed by only 28% [81–83].

6. Excitation-Contraction Coupling Failure

The first step for excitation-contraction (ECC) coupling to occur is the generation and propagation of the sarcolemma (SL) action potential and the proper functioning of the neuromuscular junction in nonfatigued muscle cells. It is well known that the SL membrane potential at rest is approximately +80 mv and, during activation, the peak action potential approaches +20 mv. The duration of the action potential is short, varying from 1 to 1.5 milliseconds (ms), and the membrane is capable of responding to stimulation frequencies even greater than 150 Hz [84–87].

With the development of fatigue induced by high rates of muscle contraction, the surface membrane potential, both at rest and in action, demonstrates characteristic changes, as the resting potential becomes depolarized in 10 to 20 mv and the height of the peak of the action potential decreases proportionately; however, the duration of the action potential becomes prolonged [84, 85].

The great lack of information about how and when these changes about the extent to which the tubular membrane T is changed and how these changes can be mitigated during exercise is still a challenge, but the hypothesis would be that the changes observed in the T tube should probably be greater compared to the changes observed in SL [86, 87].

The depolarization of the SL and T tube membranes interferes in two processes: first, they interfere with the generation and propagation of the action potential due to reduced activation and slower inactivation kinetics of the voltage-dependent Na+ channels. Second, they interfere with the inactivation of the intramembranous T tubular protein, more specifically the voltage sensor, called the 1,4-dihydropyridine receptor (DHPR), obtaining the reduction in the peak of the action potential as a direct result of the
inactivation induced by the depolarization of the voltage-regulated Na⁺ channels and, now, by the reduced electrochemical gradient and inducing the influx of Na⁺ in the T system, so that the extracellular [Na⁺] decreases and intracellular [Na⁺] increases [88–90].

For inactivation of the Na⁺ channel and DHPR, depolarization is necessary if the cell has an intracellular [K⁺] reduction and an extracellular [K⁺] increase. Studies have shown that small increases in [K⁺] increase the subtetan strength, while large increases reach a critical level where it causes tetany and an abrupt decrease in the contraction force. Besides that, the malfunction of the Na⁺ K⁺ pump may result in unregulated pH-free energy of ATPase hydrolysis by increasing ADP/Pi intracellular and extracellular [91, 92].

In order for the processes to work correctly, the following are required: maintaining the ideal rate of activation of motor activity, diversified recruitment of motor units carried out in the CNS and β-adrenergic stimuli, stimuli for the Na⁺ K⁺ pump to function, and an increase in [Cl⁻] to control the influx of K⁺ to prevent it from reaching a critical level, resulting in fatigue [93–95].

Checking the amount of K⁺ used by muscle cells during contraction and the amount of efflux released by the sarcolemma during action potentials, we can conclude that the account is far from closing. Thus, studies suggest that the depolarization observed in fatigued muscle cells is the result of the effects of high K⁺ contractions, both intracellular and extracellular, and of the increase in their conductance. The increase in K⁺ conductance can cause the activation of K⁺ channels, which depend on ATP or dependent on Ca²⁺, and the activation of these channels can directly influence the local pH, making it more basic, creating a cascading effect that would contribute to fatigue by cell depolarization or an attempt to return homeostasis [96–98].

Studies suggest that the infusion of nonspecific antioxidant N-acetylcysteine prolongs fatigue time and reduces the decline in Na⁺ K⁺ pump activity by approximately half, but the mechanisms of pump inactivation are not yet known; however, data suggest that it can be related with reactive oxygen species (ROS) [99].

Studies suggest that the main evidence that disturbances in eccentric contraction (ECC) contribute to fatigue is the low transient Ca²⁺ amplitude and that the greatest decline in Ca²⁺ release tends to occur late in fatigue. From this, we can observe that changes in the amplitude of Ca²⁺ also influence the strength, because when inducing metabolic factors signaled by the concentration of Ca²⁺ in CPB, fatigue becomes early and is characterized by an increase in ATP turnover during the initial stage of contractile activity and consequently by an increase in pi and H⁺. Thus, so that there is a release of Ca²⁺ by SL, it has become quantitatively more important in the process of developing fatigue independently if the Ca²⁺ ions act on the AM cross-bridge [97, 100].

Furthermore, there is strong evidence that Ca²⁺ plays an important role in cell signaling that triggers fatigue; however, the nature and how it occurs are still unknown. Studies suggest that in ECC, fatigue can be compensated by activating the Ca²⁺/ATPase pumps present in the T tubes, in which they can move the Ca²⁺ from the cytoplasm to the extracellular space; from that, it is expected that local factors can activate mechanisms to correct this deregulation, such as the activation of the local [Ca²⁺] pumps due to the modification of the local pH, and thus inhibit the activity of ATPase. However, [Ca²⁺] can interfere with DHPR and thus decrease the likelihood of ATPase inactivation, consequently changing the threshold of the action potential in the lumen of the T tube to increasingly positive values, making it deeply difficult for the muscle to leave the stage of contraction for relaxation [101–103].

Once knowing that recovery from fatigue usually occurs within minutes to an hour, we must understand that some studies theorize that part of the fatigue problem is actually related to cell signaling, more precisely, in the direct inhibition of the Ca²⁺ channel released by SL, called RyR1, because the content of intracellular Ca²⁺ depends on the activity of this channel, both for entry and removal, mainly in fast fibers. The permeability of the channel is directly regulated by Ca²⁺ already released in the SR, so that the ion itself, with the smallest fluctuation in concentration, regulates the RyR1 channel [104, 105].

Beyond that, considering that the intensity of the force is intracellular [Ca²⁺] related and that in turn depends on the release of Ca²⁺ and the self-regulation of some channels for removal or influx, it is important to note that the decrease in [Ca²⁺], even if it is physiological, tends to prolong the relaxation period, the opposite being also possible, and the increase in [Ca²⁺] tends to prolong the contractile period of the fibers, even when the reduction of the Ca²⁺ of the SR during the contractile activity remains due to the binding of increased Ca²⁺ intracellular fast fiber proteins and the SR pump, thus consequently removing more slowly. In summary, the cell signaling that controls the period and the place where Ca²⁺ must be maintained during the contraction time and the relaxation time after tetanus contraction in the muscle are essential to understand muscle fatigue [106, 107].

Comparatively, the mechanism for the development of muscle fatigue also depends on the amount of ATP recruited by the muscle, from which it will then be degraded into ADP + Pi, as well as the extent of phosphocreatine depletion. Studies suggest that muscle fibers have the ability to sustain ADP decline without opening Ca²⁺ channels and flooding the muscle with these ions, so that action potentials with 0.5 mM cell decline did not activate Ca²⁺ release. However, when injected with Mg²⁺ ion in the declining action potential, an inhibition of Ca²⁺ channels was observed, due to the competition of intracellular Mg²⁺ by ATP, resulting in the fall of free ATP, which was possible to be released after intense exercise [108, 109].

7. Fatigue in Organs

Fatigue occurs in whole muscles when there is a loss of peak strength, speed, and power, in which the time course of change in function depends on the intensity of activation and the composition of the type of muscle fiber [92, 109]. The human body has some muscles made up of slow muscle fibers, so they are called slow muscles, such as the sole of the calf. These fibers are composed mainly of slow type I or slow
oxidative fibers, that is, they generate ATP mainly from oxidative phosphorylation and are kept tonically active and are extremely resistant to fatigue. On the other hand, muscles made up of fast fibers usually have greater capacity to obtain ATP from glycolysis and other energy sources, are not kept tonically active, and are not resistant to fatigue [79, 96].

Going deeper into the types of fibers present in the human body. There are two types of fast fibers: IIa or fast-oxidizing glycolytic fibers and IIx or glycolytic fibers. In addition, the hybrid fast fibers that contain myosin are IIa and IIx. The hybrid fiber IIx has the highest maximum speed and glycolytic capacity, while the hybrid fiber IIa has properties between fast IIx and slow type I fibers. Additionally, there is a third fast type IIb fiber, which is the fastest of all the fibers, but is expressed, in extremely inactive muscles [108, 110].

Studies have shown that understanding the isometric contraction properties before and after fatigue has helped to identify cell fatigue sites. For example, postfatigue muscle contractions show reduced muscle tension and prolonged contraction and relaxation times. The duration of the contraction depends on the detachment rate of the AM cross-bridge and the duration of the transient intracellular Ca^{2+}; therefore, the prolongation of fatigue-induced contraction is an indication that the Ca^{2+}/ATPase reuptake by the SR may have been inhibited; note as well that the duration of relaxation after a peak of contraction in tetany is limited by the activity of the Ca^{2+}/ATPase/SR pump [50, 105].

According to studies that analyzed the initial phase of relaxation of a muscle in tetany, the lengths of the sarcomere remain constant and it is believed that the relaxation rate depends directly on the detachment rate of the AM cross-bridge, so that this phase showed a longer fatigue time compared to the second relaxation phase and should probably be limited by the activity of the Ca^{2+}/ATPase/SR pump [23].

Another point to be considered would be that the development of fatigue would be a force-frequency relationship. If the strength continuously decreases and relatively less frequently compared to the high ones, a solution would be that the transient Ca^{2+} induces in the prolongation of fatigue with low injections, however, continuous so that the muscular strength in fatigued muscles also decreases [78].

Along with the slowing of relaxation, α-motoneuron firing rates decline with fatigue in vivo. This response is consistent with studies that observe that the optimal stimulation frequency for peak force showed a decline in muscle with fatigue [39].

Some studies show that the aim should be the muscle contractile process and should include disturbances in cross-bridge AM interactions caused by increases in pH and Pi and alterations in SR function or in the Ca^{2+}/ATPase pump, besides the changes that can occur in cross-bridge AM. Therefore, the force, velocity, and power all are reduced. In addition, the hydrolysis of ATP produces an increase in free Mg^{2+} and inhibition of the RyR1 and Ca^{2+} release [106].

It is well known that in cases of prolonged exercise, fatigue is inevitable, as this is correlated with muscle glycogen depletion and hypoglycemia. Currently, studies are needed to explain this mandatory carbohydrate oxidation; it is hypothesized that the decrease in muscle glycogen and the increased dependence on fat oxidation end up limiting the ability of mitochondria to provide important components for oxidative phosphorylation such as NADH, before the muscle suffers; in addition, it was also observed that glycogen has a greater affinity with DHPR and RyR1, thus making the alternatives to meet the need for the muscle obsolete [102, 111].

A reduced uptake of SR Ca^{2+} without alteration in ATPase activity suggests a decoupling of the transport or leaking vesicle, through which Ca^{2+} escapes into the intracellular medium and changes the concentration and local pH. Another organelle that can be damaged due to prolonged physical exercise would be the mitochondria, in which studies indicate that it may present some edema, but there is no evidence that it can impair its function; however, it was observed that the fibers showed some wear after prolonged exercise [105, 109].

The oxidative phosphorylation that occurs in the mitochondria produces ATP and consumes O_2; this process is also capable of generating ROS, such as hydrogen peroxide (H_2O_2) and superoxide anion (O_2^-). The rate of production of ROS increases according to the intensity of work and the rate of respiration in skeletal muscle, and it is commonly recognized that ROS play a role in muscle fatigue by the oxidation of critical cellular proteins, such as the Na^+ K^+ pump, myofibrils, DHPR, and RyR1. The oxidation of ROS at critical RyR1 cysteine residues and/or myofibril serine groups can assist in the development of low-frequency fatigue and inhibit the release of Ca^{2+} from SR and the sensitivity of Ca^{2+} in myofibrils [91, 103, 112].

It is believed that there is a mechanism that involves a structural alteration of the Ca^{2+} SR release channel and/or associated proteins, since the amplitude of the Ca^{2+} transient is decreased for all stimulus frequencies, so that the concentration ratio of Ca^{2+} and strength is increasingly observed, and it can be concluded that Ca^{2+} mediators probably involve calmodulin (CaM), calcium-activated proteases, or ROS [64, 89, 104].

RyR1 is part of a multiprotein complex that includes proteins involved in phosphorylation/dephosphorylation, linked to different types of cell signaling, such as cAMP-dependent protein kinase (PKA), Ca-CaM-dependent protein kinase (CaMKII), phosphodiesterase-4-D-3 (PDE4D3), protein phosphatase 1 (PP1), the Ca^{2+} binding protein CaM, and modulation of Ca^{2+} activation of the channel [23, 105, 113].

It is well established that CaM plays a dual role in the regulation of the Ca^{2+} in SR release channels. CaM binding activates or inhibits the opening of the Ca^{2+} channel in the cytoplasm whose concentration is low and high. In a nonfatigued muscle fiber, this process probably contributes to the cyclic activation and inactivation of the release channel. There is no evidence linking Ca^{2+} with CaM to an altered RyR1 function. However, there is a possibility that the elevated cytoplasmic Ca^{2+} associated with fatigue leads to an altered CaM binding, so that the channel becomes more difficult to activate. The increase in cytoplasmic Ca^{2+} can lead to prolonged elevation of CaMKII and cause excessive phosphorylation of DHPR and/or RyR1. Looking from another point, prolonged β-adrenergic activity with exercise can also
lead to an increase in PKA activity and result in hyperphosphorylation of RyR1. In addition, the inhibition of the PP1 and/or PDE4D3 protein can also lead to excess phosphorylation of RyR1, which can directly reduce the probability of its opening or mediate the dissociation of the stabilizing protein from the FKBP12 channel [7, 48, 106].

The exact mechanisms and relative importance of RyR1 and the Ca\(^{2+}\)/ATPase/SR pump are not completely understood and represent an important area for future research [78, 102, 107].

8. Photobiomodulation

Low-level laser therapy (LLLT) or laser therapy has been used for more than 40 years. The idea came up in 1960 after the invention of the laser, making it a widespread treatment with a variety of clinical applications. Scholars of the time decided to use different tools and models to be used according to their functions, the expressions like "photobiostimulation" and "biostimulation" are often related to the stimulation effect that the LLLT was used for. However, a few years later, it was possible to verify that the LLLT also has an inhibitory effect; from that, it was established to coin the term "biomodulation" [108, 110].

Therapeutic treatments are based on three principles: first, minimize inflammation, edema, and chronic disorders of the joints, brain, skin, etc; second, promote wound healing in superficial and deep tissues, etc; and third, treating neurological disorders and pain [108, 111].

Recent studies indicate that the most used wavelengths for photobiomodulation therapy (PBM) are infrared (IR) and 700 nm to near infrared (NIR), in which they have shown more beneficial impacts than light-red in many medical conditions [109, 111].

In general, the laser therapy involves portions of the electromagnetic spectrum (390–1600 nm and 1013–1015 Hz), which is red until NIR, and they are absorbed according to the specific application for each biological tissue and the appropriate wavelengths [4, 112].

Unlike high-power "hard" lasers, LLLT provides low energy, just enough to induce a response in body tissue. In addition, it has a wavelength-dependent shape capable of altering cell function in the absence of significant heating. Thus, LLLT is also called "soft" laser therapy or cold laser, as it has low energy without thermal effects [114, 115].

Studies show that the wide range of laser therapy includes effects at the molecular, cellular, and tissue levels and the ways in which LLLT works can vary according to different application factors [116]. For the LLLT, to produce a photobiological effect, it is necessary for the photon absorption of the laser radiation to occur; the photons are captured by the initial photoreceptor molecules, which can be endogenous or exogenous chromophores. The energy absorbed from a photon can be transferred to another molecule, which can then cause a chemical reaction without changing the temperature in the surrounding tissue and consequently triggers local biochemical reactions without any discomfort [109, 117].

Several studies have suggested that mitochondria are the cellular component most sensitive to visible light and NIR; this stimulus results in increased ATP production, increased deoxyribonucleic acid (DNA) synthesis, ROS modulation, and nitric oxygen species (NOS) and in the induction of transcription factors. In addition, PBM at red and NIR wavelengths stimulates an increase in intracellular ionic Ca\(^{2+}\). However, recent studies emphasize that blue (420 nm) and green (540 nm) lights are more effective in increasing intracellular Ca\(^{2+}\). Researchers also suggest the use of blue or green light for better interaction with light-dependent ion channels, which allows light to control electrical excitability, intracellular acidity, and calcium influx, among other processes. The most likely ion channel is the rhodopsin of the light-gated channel, because the spectrum of action of the rhodopsin family exhibits peaks in the blue-green spectral region. However, the mechanism of laser-tissue interaction has not yet been fully described [118–120].

At the cellular and molecular levels, there are still open arguments and responses about the effectiveness of lasers in producing the desired responses [121]. Photobiomodulation is a form of phototherapy, which is designed to apply light with specific wavelengths from red to NIR with output powers of up to 500 mW. One of the great advantages of LLLT is the use of photon energy at low levels to alter biological activity without thermal reactions, since there is little increase in the temperature of the irradiated tissue, in addition to being nontoxic and nonallergic, and due to the ease of application that its study has been spreading increasingly [122].

Studies indicate that clinical treatment with LLLT in various intensities has a stimulating effect on cellular processes [123]. Recently, it has been reported that at low levels of red or infrared light, LLLT can prevent cell apoptosis; stimulate mitochondrial activity; increase in cell recruitment, proliferation, and renewal; and modulate cell metabolites. In addition, it has also been suggested that LLLT may promote changes in the cell’s redox state, playing an important role in ion homeostasis and consequently in cellular activity and induce photobiostimulation processes. Besides that, preexposure of PBM has a protective effect against many external agents, such as hydrogen peroxide (H\(_2\)O\(_2\)) and UV radiation [124, 125].

9. Discussion

Studies suggested that the implementation of low-level laser therapy (LLLT) may cause biomodulatory, biochemical, biophysical, and bioenergetic effects. That is why we can use it in the repair of fatigued muscles, muscular disorders, and improved muscle performance in specific muscles already treated with LLLT. According to [126], the experimental study with 15 volunteers including female and male stroke patients who presented with poststroke spasticity, the LLLT (diode laser, 100 mW, 808 nm, beam spot area 0.0314 cm\(^2\), 127.39 J/cm\(^2\)/point, 40 s) was applied mainly in areas where spasticity was present like the rectus femoris muscle and to the vastus medialis muscle. After LLLT, intervention was observed to contribute to the increase in the recruitment of muscle fibers, reduction in the scale for pain intensity
because of its anti-inflammatory effect in stroke patients with spasticity, time delay to the fatigue onset, improvement of muscle performance, and increase in peak torque. The study also observed that LLLT provided major breakthroughs in the treatment of muscular disorders and prevention of muscle fatigue.

The literature shows that positive results were found with a single application of LLLT on tissue regeneration and decreased neurological injury but there are not enough information about laser therapy related to spastic muscle and it shows increased resistance to passive stretching that can be explained by changes in muscle tissue properties and early onset of muscle fatigue. In addition, the authors found that the application of laser photobiomodulation in spastic muscle prior to isometric exercise contributes to the decrease in blood lactate concentration after exercise possibly because laser radiation increases microcirculation, increases metabolite removal, and prevents local ischemia. Besides, it seems that the vascular effects of PBM may be due to the nitric oxide release, leading to smooth muscle relaxation and increased peripheral microcirculation. Considering the bioenergetic effects, the LLLT application is related to mitochondrial function and light interacts with the mitochondria and promotes cellular changes that also may contribute to delay muscle fatigue [127].

Impairment of basic motor functions, such as muscle weakness in limbs affected by spasticity, leads to peripheral fatigue and impaired functionality. According to [128], the clinical use of PBM has provided major advances in the treatment of muscular disorders and prevention of muscle fatigue. Their study were structured with 10 sessions of PBMT (laser 100 mW, 808 nm, 159.24 J/cm²/point, 5 J/point); PBM active or placebo was associated with exoskeleton-assisted functional treatment. A double-blind, placebo-controlled sequential clinical trial was conducted with 12 healthy volunteers and 15 poststroke patients who presented upper limb spasticity. Ultimately, it suggests that the application of PBM may contribute to an increased range of elbow motion and muscle fiber recruitment, increases in muscle strength, and, hence, increases in signal conduction on spastic muscle fibers in spastic patients.

10. Conclusion

It could be concluded that LLLT may contribute to an increased range of muscle motion and fiber recruitment and increased strength to increase signal conduction on spastic muscle fibers in spastic patients. However, further studies are needed to understand how fatigued fibers behave in these patients, since clinical studies used different laser wavelengths and numerous illumination parameters, which influence the determination of different biological parameters.

In addition, the literature is not explicit about the application of PBM in spastic patients but that this possibility could reduce pain and fatigue regarding biochemical and neuromotor mechanisms that may represent a new therapeutic use for neurological patients.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

References


O. Friedrich, M. B. Reid, G. Van den Berghe et al., The Sick and the Weak: Neuropathies/Myopathies in the Critically Ill, Physiological reviews, 2015.


