Effects of aerobic and resistance exercise in murine models of rheumatoid arthritis and osteoarthritis – a systematic review

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Abstract

Introduction and Objective. Rheumatoid arthritis (RA) is an inflammatory and systemic autoimmune disease that affects peripheral joints leading to joint degradation, pain, deformities, decreased activities of daily living, and sedentary lifestyle, resulting in secondary sarcopenia. Osteoarthritis (OA), in turn, is a multifactorial disease associated with joint degeneration and impairment of the musculoskeletal system as a whole, affecting joint movement and stability. In both pathologies, there is loss of muscle mass and functional impairment, and physical exercise is a therapeutic alternative to minimize these consequences. The aim of the review is to demonstrate the effects of different physical exercise modalities on muscle mass loss in murine models of OA and RA.

Materials and method. The databases used were PUBMED, EMBASE, Cochrane Database, LILACS, and Google Scholar. The review included studies which induced experimental models of OA or RA in rats or mice; used any exercise modality as an intervention; and analyzed some quantitative muscle histomorphometric measures. The search strategy included all keywords on the topic identified in previous research and adapted for each database. Studies published in any language and in any year were included. An analysis of the studies was performed by two independent reviewers and data were extracted from the articles using tools developed by the reviewers.

Results. After completing the definitive search in the databases, 193 studies were found, of which only 2 were included in this review, one addressing the effects of resistance exercise in rats with RA, and another using aerobic exercise to treat rats with OA.

Conclusion. Treatment with different forms of physical exercise contributed to attenuation of muscle mass loss in animal models of OA and RA.

Key words

exercise therapy, animals models, inflammation, joints

INTRODUCTION

Rheumatoid arthritis (RA) is an autoimmune disease that causes inflammation and joint degradation, characterized by symmetric and chronic synovitis that mainly affects the peripheral joints. In addition to joint inflammation, RA most frequently triggers systemic inflammation that can lead to pain, with consequent decreased activities of daily living, sedentary lifestyle, and even joint deformities [1,2]. This systemic inflammation can also lead to secondary sarcopenia, which appears to be primarily driven by catabolic processes. When systemic inflammation persists and is associated with anorexia, asthenia, and/or physical inactivity, additional muscle loss pathways are activated, altering the balance of muscle tissue remodelling, and may activate different intracellular proteolysis pathways [3–5].

Osteoarthritis (OA), in turn, is a multifactorial disease that affects the musculoskeletal system as a whole, especially joint tissues (causing degeneration in meniscus, ligament, and synovium) and muscle tissues (leading to loss of periarticular muscle mass) [6]. Because the muscular system performs, among others, the functions of joint stability and strengthening [7], once compromised in OA, there is generalized fatigue, functional disability, increased risk of falls, as well as pain and stiffness, directly affecting the quality of life of sufferers [8].

Aimed at minimize the consequences of RA and OA, physical exercise has been used as a non-pharmacological treatment possibility, mainly because it is accessible and low-cost [9–11], and promotes benefits such as improved cardiorespiratory fitness and increased muscle mass, aerobic capacity, joint mobility, and physical function [12]. Overall, exercise promotes adaptive responses in skeletal muscle as a result of the cumulative effect of activating a series of molecular signalling pathways which, in turn, modulate protein synthesis and degradation to promote muscle hypertrophy while preventing atrophy [13].

Resistance exercise, which aims to provide resistance against muscle action through weights, is an important form of treatment for sarcopenia because it generates an increase
in the amount of type I muscle fibres and the cross-sectional area of type II fibres [14], promoting regulation of the rate of protein synthesis [15], and decreasing the levels of atrogin-1, MuRF-1, and myostatin; therefore, acting on the prevention of muscle atrophy [13]. Aerobic exercise, on the other hand, another modality of exercise which consists of long-duration, low-intensity resistance exercises [16], acts by improving pain, oedema, and joint function, which is limited in OA and RA [17,18].

However, although the benefits of physical exercise for the treatment of RA and OA and muscle atrophy are known, the studies are divergent as to the types of exercises and protocols used. Therefore, a systematic review compiling all this information is justified with the aim of future clinical direction. In view of this, the objective of the present study was to highlight the effects of different modalities and physical exercise protocols on muscle mass loss in murine models of OA and RA.

MATERIALS AND METHOD

Prior to conducting this review, a preliminary search in PROSPERO and PUBMED was conducted until 15 June 2021, aimed at finding other protocols/reviews with the same objective, and to rule out the possibility of duplication of studies.

Search strategy. This systematic review included studies published in any language and year in the following databases: PUBMED, EMBASE, Cochrane Database and LILACS, as well as Google Scholar as grey literature. The final search strategy was outlined in PUBMED and encompassed all keywords found in articles on the topic in the preliminary search. These were later adapted for each database (Appendix 1).

Inclusion and exclusion criteria. To be included in the review, the studies should adhere the following criteria: 1) use rats or mice of any gender, age or lineage, as samples induced to some experimental model of OA or RA; 2) use any modality of physical exercise as an intervention; 3) have analyzed at least one quantitative measure regarding muscle histomorphometry.

Articles were excluded if they involved the following: 1) the samples used were not rats or mice; 2) the sample was not submitted to some model of OA or RA, or had other associated pathology; 3) have analyzed at least one quantitative measure regarding muscle histomorphometry.

Article Selection. After the final search of the strategies in all databases, the retrieved citations were evaluated and duplicate studies excluded. Subsequently, two independent reviewers read the titles and abstracts to verify that they met the pre-established eligibility criteria. The studies initially included were read in full and again evaluated. Disagreements that arose between the reviewers were resolved by discussion, or by a third reviewer when necessary.

RESULTS

After the final search of the databases, a total of 193 studies were found, of which 32 articles were excluded because they were duplicates, 156 were excluded after reading the title and abstract, and 5 studies were read in full. After analyzing the eligibility criteria, only 2 articles were included in the present systematic review (Fig. 1).

Of the articles included, one was a RA model [19] and the other an OA model [20] – both written and published in English. In the article by Oliveira et al. [19], the resistance exercise (stair climbing with load) was used in a model of RA, and in the study by Assis et al. [20], aerobic exercise (running) associated or not with low-power LASER therapy, was used in a model of (Tab. 1).

DISCUSSION

The results obtained demonstrate that RA and OA cause morphological alterations in the muscle and in the expression of enzymes of the ubiquitin-proteosome system, consistent with muscle atrophy. Resistance and aerobic exercises provided regulation of myostatin, atrogin-1, MyoD, MuRF-
Table 1. Summary of included studies

| Author / Year | Country | Pathology | Sample | Affected joint / muscle analyzed | Intervention | Comparison | Outcome | Results |
|---------------|---------|-----------|--------|---------------------------------|--------------|------------|---------|---------|
| Oliveira et al., 2018 | Brazil | RA (Induced by methylated bovine albumin + CFA + Mycobacterium tuberculosis) | 26 Wistar rats (±126 g, 8 weeks old) | Tibiotarsal joint Gastrocnemius | Resistance exercise: – Stair climbing (1.1 x 0.18cm, 2cm between steps, 80° incline) with load. Training = 5 sets of 3 repetitions, with 120s interval between them, and progressive load of 0%, 25%, 50%, 75%, and 100% of body weight. If the animal failed, the load was reduced by 25%. Training was performed once, 6 hrs before euthanasia. | Control Group (CG): no intervention, n=7 | 1) Weight of gastrocnemius muscle 2) Cross-sectional area of gastrocnemius muscle 3) Expression of myostatin and atrogin-1 proteins 4) Expression of MuRF-1 enzyme | 1) Smaller in the GAR and GAR+EX groups vs. CG and EG (p<0.05) 2) Decreased in GAR and GAR+EX groups vs. CG and EG (p<0.05) 3) Increased in GAR group vs. EG and GAR+EX; and NSD in GAR+EX group (p<0.05) 4) All groups = NSD (p<0.05) |
| Assis et al., 2015 | Brazil | OA (Induced by sectioning the anterior cruciate ligament) | 50 Wistar rats (±150 g, 6 weeks old) | Femorotibial joint Quadriceps | Aerobic exercise: – Running, 16min/min, 3x/week, 50min/day, for 8 weeks. | Control Group (CG): no intervention, n=10 | 1) Cross-sectional area of quadriceps muscle 2) Quadricipus muscle fibre density 3) Expression of MuRF-1 enzyme and atrogin-1 protein | 1) Reduction in all groups vs. CG (p<0.001); and increase in groups OAT and OATL vs. OAC (p<0.001) 2) Increase in OAC, OAL, OAT, and OATL groups vs. CG (p<0.003); and decrease in OAT and OATL groups vs. OAC (p<0.02) 3) Increase in OAC vs. OAL (p<0.05); all other groups = NSD (p<0.05) |
1, and myogenin expression and prevented a decrease in the cross-sectional area of muscle fibres, thus preventing muscle atrophy in animal models of RA and OA.

In a study carried out in elderly people with rheumatoid arthritis, submitted to resistance and endurance exercise protocols, improvements in aerobic capacity, endurance and muscle strength were verified [21]. However, in individuals with knee osteoarthritis, submitted to resistance training, a reduction in pain was observed, including in the walking function [22]; however, these records point to gaps that still need to be filled in the literature regarding the forms of physical exercise, as well as the most suitable modality.

In both RA and OA, a large number of pro-inflammatory cytokines are produced, including TNF-α and IL-6 [23–25]. In the study by Adams et al. [26] it was observed that TNF-α induced MuRF-1 expression in skeletal muscle, leading to a reduction in muscle contractile strength by reducing myofibrillar sensitivity to calcium, which is an important factor in muscle weakness. Therefore, this is believed to be one of the reasons why skeletal muscle in individuals with RA exhibits impaired adaptation to exercise and injury repair [27]. Using an aerobic exercise protocol in mice with high levels of TNF-α, Mangner et al. [28] investigated whether exercise could reduce the loss of muscle mass in the diaphragm caused by TNF-α. They found that aerobic exercise promoted antioxidant action, minimizing TNF-α activity and preventing degradation activity through the ubiquitin-proteosome pathway. These findings are plausible for justifying the action of aerobic exercise in attenuating the loss of mass and muscle atrophy in animals with RA and OA.

Also using aerobic exercise, in the study by Assis et al. (Assis et al., 2015), rats were separated into 5 groups: a control group, osteoarthritis group, osteoarthritis group associated with laser, osteoarthritis and aerobic exercise group, and osteoarthritis group with laser associated with aerobic exercise. Three weeks after the induction of OA, running training was started on a treadmill at a speed of 16m/min, 50 minutes a day, 3 times a week for 8 weeks. It was observed that in the OA and OA groups with laser associated with aerobic exercise, there was no increase in MuRF-1 enzyme and Atrogin-1 protein expression. A similar study conducted by Martins et al. [29] in an experimental model of knee OA, moderate intensity aerobic training was applied on a treadmill, which started one week after OA induction, at a speed of 16m/min, 3 times a week, for 8 weeks and with a time progression from 30 minutes to 50 minutes in the fourth week. After the analysis of joint biomarkers and functional adaptations, aerobic training was found to reduce inflammatory marker levels of inflammatory markers (IL-1β, TNF, IL-10 and IL-6) and improve the functional performance of OA animals.

In turn, in another article included in this review, Oliveira et al. [19] opted for the use of resistance exercise and divided the rats into 4 groups: a control group (no intervention), RA group (induced RA), resistance exercise group (stair climbing with load) and RA group associated with exercise (induced RA and submitted to stair climbing exercise). In the results, the RA group showed increased levels of myostatin in the gastrocnemius muscle, while in the RA group associated with exercise this did not increase, demonstrating that resistance exercise promoted a protective effect in relation to muscle damage. Also using a RA model submitted to the resistance stair climbing exercise, Neves et al. [30] associated exercise with low-intensity LASER treatment. They concluded that the interventions (stair-climbing exercise and LASER), combined or not, caused positive effects on the modulation of the inflammatory process and leukocyte migration, both in the acute or chronic inflammatory process, indicating the benefits of the intervention at the beginning of the disease.

**CONCLUSION**

Based on the studies analyzed, it was possible to observe that physical exercise applied in murine models of RA and OA resulted in benefits to muscle tissue, which suggests an attenuation of the loss of muscle mass that is promoted by these inflammatory pathologies. However, the existence of several modalities and exercise protocols is observed, with varying intensity and parameters. This, added to the scarcity of experimental studies with protocols described in detail, which analyze the loss of muscle mass and the various ways in which exercise can interfere with this aspect, makes the research limited. In view of this, more quality primary studies on the subject are necessary so that future reviews can be more conclusive in answering the real effectiveness of each modality and protocol of exercises on the muscles in models of RA and OA.

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