Swimming training potentiates the recovery of femoral neck strength in young diabetic rats under insulin therapy

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OBJECTIVE: To test whether swimming training benefits femoral neck strength in young diabetic rats under insulin therapy.

METHODS: A total of 60 male Wistar rats (age: 40 days) were divided equally into the following six groups: control sedentary, control exercise, diabetic sedentary, diabetic exercise, diabetic sedentary plus insulin and diabetic exercise plus insulin. Diabetes was induced with a unique intraperitoneal injection (60 mg/kg body weight) of streptozotocin. Seven days after the injection and after 12 hours of fasting, the animals with blood glucose levels $\geq$ 300 mg/dL were considered diabetic. Seven days after the induction of diabetes, the animals in the exercise groups were subjected to progressive swimming training (final week: 90 min/day; 5 days/week; 5% load) for eight weeks. The animals in the insulin groups received a daily dose of insulin (2-4 U/day) for the same period.

RESULTS: Severe streptozotocin-induced diabetes reduced the structural properties of the femoral neck (trabecular bone volume, trabecular thickness and collagen fiber content). The femoral neck mechanical properties (maximum load and tenacity) were also impaired in the diabetic rats. Insulin therapy partially reversed the damage induced by diabetes on the structural properties of the bone and mitigated the reductions in the mechanical properties of the bone. The combination of therapies further increased the femoral neck trabecular bone volume ($\geq$ 30%), trabecular thickness ($\geq$ 24%), collagen type I ($\geq$ 19%) and type III ($\geq$ 13%) fiber contents, maximum load ($\geq$ 25%) and tenacity ($\geq$ 14%).

CONCLUSIONS: Eight weeks of swimming training potentiates the recovery of femoral neck strength in young rats with severe streptozotocin-induced diabetes under insulin therapy.

KEYWORDS: Diabetes; Exercise Training; Bone Fragility; Collagen Fibers.

INTRODUCTION

The risk of high hip fracture in people with type 1 diabetes mellitus (T1DM) is thought to be caused by poorer bone quality, which is related to impaired bone structure and tissue integrity (1,2,3). Long periods of hyperglycemia lead to nonenzymatic glycation (4) in bone tissue along with augmented cross-links in the bone collagen matrix, thus reducing bone formation (1,5) and increasing bone brittleness and fragility (6,7). Poor bone quality has been reported in adolescents with T1DM (8) and in a young rat model of streptozotocin (STZ)-induced diabetes; the characteristics of poor bone quality included impaired structure and mechanical properties of the femoral neck and impaired resistance to fracture (9,10). Fractures of the femoral neck account for over 50% of the hip fractures that occur in populations with elevated bone fragility (i.e., osteoporosis) (11).

Insulin therapy is essential for glycemic control and is linked with steady bone mineral density (BMD) as well as marked decreases (i.e., 38%) in bone resorption in patients with T1DM (12). In animal models, insulin reversed the reduction in BMD caused by STZ (13) and mitigated the deleterious effects of STZ on the structure and function of the femoral neck in young rats experimentally exposed to severe hyperglycemia (9).

Regular exercise is recommended to counteract bone fragility because of the mechanical stimulation and benefits to bone health that are associated with exercise (14).
Therefore, swimming is an attractive therapeutic option for people with high bone fragility, such as those with severe T1DM, because of its safety, especially in relation to falls and excessive loads. Despite the lower impact of swimming exercise and the consequent osteogenic benefits compared to those from impact exercises (15), swimming has been shown to significantly improve femoral structural and mechanical properties in osteoporotic rats (16,17).

Therefore, in this study, we employed a rat model of STZ-induced diabetes to test whether swimming training improves the strength of the femoral neck in young rats under insulin therapy.

## METHODS

### Animals

Forty-day-old male Wistar rats were allocated into one of the following six experimental groups (n=10 each group): control sedentary (CS), control exercise (CE), diabetic sedentary (DS), diabetic exercise (DE), diabetic sedentary + insulin (DSI), and exercise + insulin (DEI). The rats were housed in an environment with controlled humidity (60-70%), a controlled light-dark cycle (12/12 hours) and controlled temperature (~22°C) and had free access to water and commercial chow. All experimental procedures were approved by the Ethics Committee on Animal Use of the Universidade Federal de Viçosa, Brazil (Protocol n. 51/2011), and the study was performed in accordance with international ethical standards (18).

### Induction of diabetes and blood glucose monitoring

After fasting for 12 hours, the animals from the diabetic groups received a single intraperitoneal injection (60 mg/kg of body weight (BW)) of STZ (Sigma-Aldrich, USA) diluted in 1.0 ml of a buffer solution (sodium citrate - 0.1 M, pH 4.5). The control rats were injected with the same dose of only the buffer. To confirm the induction of diabetes, one week after STZ injection, the blood glucose (BG) levels of the animals were measured (One Touch Ultra - Johnson & Johnson, Mexico) at rest after fasting for 12 hours. The rats that exhibited fasting hyperglycemia (i.e., BG levels ≥300 mg/dL) were considered diabetic. BG was monitored weekly on Monday mornings at 68 and 15 hours after the training sessions and insulin injections performed on Friday, respectively.

### Swimming training and insulin therapy

After 7 days of hyperglycemia, the animals in the DE, DEI and CE groups started the training program in which they swam 5 days/wk (Monday to Friday) for 8 weeks. This training program was adapted from a program discussed in a previous study (19). Briefly, in the 1st week, the rats swam for 10 min/day, and the exercise duration was then increased by 10 min/day with no additional load (Table 1). By the 2nd week, the duration of the exercise was increased again by 10 min/day until it reached 90 min with no additional load. Such conditions were maintained during the 3rd week. From the 4th week on, the swimming duration of 90 min/day was maintained, and a progressive load of 1% of BW was added to the tails of the animals weekly until the load reached 5% on the 8th week (Table 1).

The rats from the DSI and DEI groups received human insulin at a dose of 2-4 U/day/rat (i.e., 1U per 60 g of BW) for the same 8-week period. The insulin therapy started with a daily dose of 2 units (subcutaneous) at 6 p.m., 6 hours after the training session. To simulate severe T1DM, the insulin dosage was adjusted gradually to maintain an average BG level of ~300 mg/dL.

### Sample collection

Forty-eight hours after the end of the 8-week swimming program, the animals were euthanized and their right and left femurs were removed and dissected from the connective tissue. The left femur was fixed in neutral buffered formalin (10%) and used for histomorphometry. The right femur was immersed in saline solution and stored at -20°C until its use in the femoral neck mechanical testing.

### Femoral neck histomorphometry

The left femur was decalcified and dehydrated, and longitudinal sections (5 μ-thick) were cut from the midneck region using a microtome (Leica 2065, Germany) and were affixed to histology slides as described previously (10). Slides stained with hematoxylin and eosin (H&E) were used to determine the trabecular bone volume per total bone volume (BV/TV), while the slides stained with Sirius Red were used to assess the collagen content. Representative photomicrographs of femoral neck collagen type I (red and yellow colors) and type III (green color) as well as the trabeculae and bone marrow of the animals in the experimental groups are shown in Figure 1.

The BV/TV was measured as described previously (10). Briefly, 5 images (200x) in distinct fields from each rat were digitalized using a photomicroscope (Olympus Biological CX31, Japan) equipped with the analySIS® getIT software (Olympus Soft Imaging Solutions GmbH). The images were analyzed using Image-Pro Plus software, version 4.5.0.29 (Media Cybernetics, USA). The percentage of trabecular bone calculated from the images was used to express BV/TV. The percentage of bone marrow calculated from the images was

### Table 1 - Swimming training schedule.

| Week | Load (% BW) | Monday (Time - min) | Tuesday (Time - min) | Wednesday (Time - min) | Thursday (Time - min) | Friday (Time - min) |
|------|-------------|---------------------|----------------------|-----------------------|----------------------|---------------------|
| 1    | 0           | 10                  | 20                   | 30                    | 40                   | 50                  |
| 2    | 0           | 60                  | 70                   | 80                    | 90                   | 90                  |
| 3    | 0           | 90                  | 90                   | 90                    | 90                   | 90                  |
| 4    | 1           | 90                  | 90                   | 90                    | 90                   | 90                  |
| 5    | 2           | 90                  | 90                   | 90                    | 90                   | 90                  |
| 6    | 3           | 90                  | 90                   | 90                    | 90                   | 90                  |
| 7    | 4           | 90                  | 90                   | 90                    | 90                   | 90                  |
| 8    | 5           | 90                  | 90                   | 90                    | 90                   | 90                  |

BW: body weight.
used to express trabecular spacing. Trabecular thickness was determined by measuring the width of all trabeculae in the images (5 images per rat). Because the diabetic rats exhibited reduced BV/TV, we used the 5 thinnest trabeculae of each rat for comparison purposes.

The collagen content was assessed as described previously (20). In brief, we digitalized 5 images (200x) in distinct fields from each rat using a microscope (Olympus AX-70) with polarized light equipped with SPOT Basic Software, version 3.5.9 (Diagnostic Instruments, USA). The images were analyzed using Image-Pro Plus software (version 4.5.0.29 - Media Cybernetics). The collagen type I and type III fibers were counted and expressed as percentages.

Measurements of femoral neck mechanical properties

The mechanical properties of the femoral neck were measured in the right femurs as described previously (10). In brief, to measure the strength of the femoral neck, we placed the femur axially in a supportive holder mounted on a computer-controlled universal testing machine (EMIC, DL 3000, Brazil) equipped with a 2000-N load cell. The loading force was applied downward to the top of the femoral head at a constant speed (0.5 mm/min) using a loading cup until the femoral neck fractured. Data were recorded at a frequency of 60 Hz and converted into a load-displacement curve from which we determined the maximum load, stiffness, tenacity, yield point energy and postyield energy. Since diabetes reduced the BW of the rats, the femoral neck mechanical properties were normalized by BW.

Statistical analysis

A two-way repeated-measures analysis of variance (ANOVA) and Tukey’s post hoc test were used to compare the BW and BG data. To compare the structural and mechanical data, we used the two-way ANOVA followed by Tukey’s post hoc test.

RESULTS

Blood glucose and body weight

Diabetes augmented BG levels to ~500 mg/dL from week 2 until week 8 (Figure 2A). Insulin treatment reduced the BG levels to ~320 mg/dL in both the sedentary and trained
diabetic rats from week 4. Exercise training did not affect BG levels in the control or diabetic rats, and the combination of treatments had no effect on BG levels. Independent of the treatments, the control rats presented a higher BW than the diabetic rats from week 3 onward (Figure 2B). Insulin therapy partially restored the BWs of the diabetic rats, and the difference in BW between the diabetic insulin-treated rats and nontreated rats was significant from week 7 on. Exercise training had no effect on BW in the control or diabetic rats, and the combination of treatments did not affect BW.

Femoral neck structural properties

The BV/TV was diminished in the diabetic rats (−45%; Figure 3A). However, insulin therapy restored the BV/TV to the levels of the control rats (−62% increase). The swimming program increased the BV/TV in control rats (−33%), but in the diabetic rats that were not treated with insulin, this effect was not statistically significant. Nevertheless, the combination of therapies further increased the femoral neck BV/TV (−30%). Diabetes reduced (−28%) and increased (−37%) the trabecular thickness (Figure 3B) and spacing (Figure 3C), respectively. In contrast, insulin therapy increased (−28%) and reduced (−21%) the trabecular thickness and spacing, respectively. The swimming training program increased the trabecular thickness (−24%) and diminished the trabecular spacing (−27%) in the femoral neck of the control rats. However, in the diabetic rats that were not treated with insulin, the effects of exercise on these parameters were not
significantly different. When the therapies were combined, the trabecular thickness and spacing in the diabetic rats were further increased (~24%) and reduced (~21%), respectively.

Diabetes reduced the total collagen content (~39%), which was partially restored by insulin therapy (~53% increase; Figure 4 A) and completely restored when swimming training was combined with insulin therapy (~13% increase).

Collagen type I was diminished in the diabetic rats (~43%) (Figure 4 B); however, it was almost completely restored by insulin therapy alone (~76% increase), and the level was further increased by the combination of insulin and exercise training (~19%). Collagen type III was also diminished in the diabetic rats (~45%) (Figure 4 C); nevertheless, it was partially restored by insulin therapy alone (~40% increase), and the level was further increased by the combination of insulin therapy and swimming training (~23%). Swimming training alone did not significantly affect these collagen parameters.

**Femoral neck mechanical properties**

The maximum load was significantly reduced in the diabetic rats (~35%), and it was partially restored by insulin therapy (~28% increase) (Table 2). The exercise training increased the maximum load in the control rats (~26%) and had no effect on the diabetic rats that were not receiving insulin therapy. The combination of treatments, however, further increased (~25%) the femoral neck maximum load in the diabetic rats. Diabetes status did not affect femoral neck stiffness. Insulin therapy, either alone or in combination with swimming training, had no effect on bone stiffness. Likewise, no effect of swimming training alone was observed. The femoral neck tenacity was significantly reduced in the diabetic rats (~43%) and was partially restored by insulin therapy (~47% increase). The femoral neck tenacity was further increased by the combination of insulin and exercise training (~14%). Swimming training alone increased the femoral neck tenacity (~35%) in only the control rats. The yield point energy was not significantly reduced in the diabetic rats. However, insulin therapy alone enhanced the yield point energy (~56%) and further increased this parameter when combined with swimming training (~13%). Swimming training alone increased the yield point energy in only the control rats (~16%). The postyield energy was not significantly different among the groups and was not significantly affected by diabetes status, insulin therapy or swimming training.

**DISCUSSION**

In this study, severe diabetes markedly reduced the BV/TV in the femoral neck and subsequently resulted in negative changes in trabecular thickness (i.e., diminished) and spacing (i.e., augmented). We also observed reduced collagen fibers in the diabetic rats. The diminished secretion and/or action of insulin, which is associated with the formation of advanced glycation end-products (AGEs) induced by oxidative stress and hyperglycemia, debilitates new bone formation by decreasing osteoblast activity and increasing osteoclast activity, thus reducing bone mass (21,22). In a rat model of diabetes, active osteoblasts may revert into inactive bone-lining cells, and a reduction in the proliferation of preosteoblastic cells has been proposed (23). The formation of AGEs also inhibits type I collagen synthesis (24) and the attachment of osteoblasts to the collagen matrix (22). Thus, since the collagen type III network plays a pivotal role in bone strength as it facilitates musculoskeletal exchange (25), the reduction in collagen content found in the femoral neck of the diabetic rats also contributed to the reduced trabecular BV/TV and impaired mechanical properties (see below).

Insulin therapy restored the femoral neck BV/TV in the diabetic rats, which was also reflected in the recovery of

![Figure 4](https://example.com/figure4.png)

*Figure 4 - Femoral neck total collagen (A), collagen type I (B) and type III (C) fibers. Data are expressed as the means ± SEMs of 5 images per rat of the 10 rats in each group. CS: control sedentary; CE: control exercise; DS: diabetic sedentary; DE: diabetic exercise; DSI: diabetic sedentary plus insulin; DEI: diabetic exercise plus insulin. *#§Significant difference between groups, p<0.05; *vs. CS, *vs. CE, *vs. DS, *vs. DE, *vs. DSI.
trabecular thickness and space. Likewise, insulin therapy restored the collagen content in the femoral neck of the diabetic rats. Insulin exerts an anabolic effect on bone and regulates bone turnover (26). Regarding collagen fibers, insulin acts directly on osteoblasts to increase bone collagen synthesis (27). Indirectly, by reducing BG levels, insulin decreases nonenzymatic glycation and, hence, the formation and accumulation of AGEs; insulin also decreases the formation of cross-links in the bone collagen matrix, thereby reducing its consequent mineralization (4).

While swimming training alone did not counteract the severe damage associated with T1DM, our results showed that combining insulin therapy with swimming training for 8 weeks further improved the recovery of the BV/TV and collagen fiber content (i.e., a major increase in type III collagen fibers) in the femoral neck of diabetic rats. This finding indicates that swimming training provided supplementary osteogenic stimuli in addition to the effects of insulin. Nonweight-bearing exercise, such as swimming, positively influences the skeleton indirectly by promoting increases in muscle mass and strength, which stimulates mechanoreceptors to trigger biological signals in the bone (12) and alter hormonal levels, leading to bone deposition (28). The collagen content in the femoral neck of nondiabetic rats has also been shown to be increased by swimming training (10). It has been suggested that the collagen fibers of the femoral neck at the point of muscle insertion are augmented as a result of forces exerted by the strong musculature (29).

Along with worsened structural properties, the maximum load and tenacity of the femoral neck were impaired by severe diabetes. The maximum load is a measure of bone strength that is underpinned by bone mass and collagen. Therefore, the reduced femoral neck BV/TV and collagen content resulted in decreased trabecular bone resistance to fracture in the diabetic rats. The lower tenacity that was observed in the T1DM rats is in line with the reduced femoral neck collagen content in these rats, as bone tenacity is determined primarily by the unmineralized matrix (30). In fact, reduced bone collagen content is suggested to result in bone loss and diminished bone strength in T1DM (4), resulting in fragile bones. Moreover, the increased cross-links in the bone collagen matrix lead to premature increases in bone brittleness (6,7). However, other factors, such as bone microarchitecture and microdamage accumulation, may negatively affect bone biomechanical integrity in diabetic rats (31).

Treatment with insulin improved both femoral neck maximum load and tenacity in diabetic rats. Although swimming training alone did not efficiently counterbalance the T1DM-associated deteriorations, the combination of swimming training and insulin therapy further improved the femoral neck strength, as an additional increase in the maximum load and tenacity was found in the diabetic rats. Such benefits in these mechanical properties were associated with restored femoral neck structural properties resulting from insulin and the combination of insulin and swimming exercise, as mentioned above. These findings reinforce the usefulness of swimming training as an osteogenic stimulus, especially in populations under treatment for bone fragility. Nevertheless, swimming training may improve bone strength by positively affecting other factors, such as trabecular bone microarchitecture, thus benefitting the structural efficiency of the trabecular bone (16,17).

Despite the differences in bone loading patterns between rats and humans, the effects of swimming training on diabetic rat trabecular bone may be of clinical relevance for humans because of the impaired femoral neck properties and high risk for fracture in patients with T1DM. Moreover, the results obtained here represent a scenario of severe T1DM (i.e., BG >300 gm/dl) and may not be generalizable to patients under insulin therapy, although a clinical study reported reduced bone mass in young subjects with moderate T1DM who had good metabolic control (32).

Finally, this study has limitations. First, serum and urinary bone biochemical markers were not assessed, and the bone mass in diabetic rats may have been influenced by urinary calcium and magnesium excretion, as well as intestinal calcium absorption (33). Second, insulin concentrations were not assessed. Nevertheless, the measurement of BG levels is a good indicator of insulin deficiency.

In conclusion, these results indicate that eight weeks of swimming training potentiates the recovery of femoral neck strength in young rats with severe STZ-induced diabetes under insulin therapy.

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### AUTHOR CONTRIBUTIONS

Gomes GJ was responsible for the intellectual and scientific content of the study, the technical procedures that were followed, and the acquisition and interpretation of the data. Del Carlo RJ was responsible for the intellectual and scientific content of the study and the critical revision of the manuscript. da Silva MF, Cunha DNQ, da Silva E and Silva KA were responsible for the acquisition and interpretation of the data and the technical procedures that were followed. Carneiro-Junior MA and Primola-Gomes TN were responsible for the intellectual and scientific content of the study and the interpretation of the data. Natali AJ was responsible for the design and intellectual and scientific content of the study, for writing and critically revising the manuscript and for securing funding. All authors approved the final version of the manuscript.

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**Table 2 - Femoral neck mechanical properties.**

|               | CS        | CE        | DS        | DE        | DSI       | DEI       |
|---------------|-----------|-----------|-----------|-----------|-----------|-----------|
| Maximum load (N) | 0.391 ± 0.031 | 0.492 ± 0.029* | 0.252 ± 0.030* | 0.281 ± 0.029* | 0.323 ± 0.030* | 0.404 ± 0.031* |
| Stiffness (N/mm)     | 0.441 ± 0.060 | 0.611 ± 0.089* | 0.510 ± 0.061 | 0.532 ± 0.041 | 0.382 ± 0.051 | 0.471 ± 0.039 |
| Tenacity (μm)       | 0.260 ± 0.030 | 0.352 ± 0.031* | 0.150 ± 0.030* | 0.181 ± 0.029* | 0.225 ± 0.028* | 0.252 ± 0.030* |
| Yield point energy (μJ) | 0.057 ± 0.009 | 0.087 ± 0.008* | 0.048 ± 0.009 | 0.052 ± 0.009* | 0.075 ± 0.008* | 0.085 ± 0.008* |
| Postyield energy (μJ) | 0.239 ± 0.043 | 0.297 ± 0.040 | 0.201 ± 0.043 | 0.224 ± 0.040 | 0.180 ± 0.043 | 0.185 ± 0.040 |

Data are expressed as the means ± SEMs of 10 animals in each group. CS: control sedentary; CE: control exercise; DS: diabetic sedentary; DE: diabetic exercise; DSI: diabetic sedentary plus insulin; DEI: diabetic exercise plus insulin. *Significant difference between groups, p<0.05; vs. CS, †vs. CE, ‡vs. DS, §vs. DE. ¡vs. DSI.
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