Tribulus terrestris extracts alleviate muscle damage and promote anaerobic performance of trained male boxers and its mechanisms: Roles of androgen, IGF-1, and IGF binding protein-3

Yiming Ma, Zhicheng Guo, Xiaohui Wang *

School of Kinesiology, Shanghai University of Sport, Shanghai 200438, China

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Abstract

Purpose: To investigate the effects of Tribulus terrestris (TT) extracts on muscle mass, muscle damage, and anaerobic performances of trained male boxers and its mechanisms: roles of plasma androgen, insulin growth factor 1 (IGF-1), and IGF-1 binding protein-3 (IGFBP-3).

Methods: Fifteen male boxers were divided into exercise group (E, n = 7) and exercise plus TT group (E + TT, n = 8). The 2 groups both undertook 3-week high-intensity and 3-week high-volume trainings separated by a 4-week rest. TT extracts (1250 mg/day) were orally administered by boxers in E + TT group. TT extract compositions were detected by UHPLC–Q-TOF/MS. Before and at the end of the 2 trainings, muscle mass, anaerobic performance, and blood indicators were explored.

Results: Compared with E group, decreases of plasma CK (1591.5 ± 909.6 U/L vs. 2719.9 ± 832.5 U/L) and IGFBP-3 (3075.5 ± 1072.5 ng/mL vs. 3950.8 ± 479.3 ng/mL) as well as increases of mean power (MP , 459.4 ± 122.3 W vs. 434.6 ± 69.5 W) and MP/body weight (MP/BW, 7.5 ± 0.9 W/kg vs. 7.1 ± 1.1 W/kg) were detected in E + TT group after a high-intensity training. For high-volume training, reduction of IGFBP-3 (2946.4 ± 974.1 ng/mL vs. 3632.7 ± 470.1 ng/mL) and increases of MP (508.7 ± 103.2 W vs. 477.8 ± 49.9 W) and MP/BW (8.2 ± 0.3 W/kg vs. 7.5 ± 0.9 W/kg) were detected in E + TT group, compared with E group. Muscle mass, blood levels of testosterone, dihydrotestosterone (DHT), and IGF-1 were not significantly changed between the 2 groups.

Conclusion: Taking 1250 mg capsules containing TT extracts did not change muscle mass and plasma levels of testosterone, DHT, and IGF-1 but significantly alleviated muscle damage and promoted anaerobic performance of trained male boxers, which may be related to the decrease of plasma IGFBP-3 rather than androgen in plasma.

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Keywords: IGF binding protein-3; Insulin growth factor 1 (IGF-1); Muscle damage; Performance; Testosterone; Tribulus terrestris

1. Introduction

Tribulus terrestris (TT) is a famous traditional Chinese medicine that has been widely used in many countries for thousands of years. TT revealed many compounds including steroidal saponins, flavonoids, alkaloids, and amino acids. TT saponins are considered the most important active components that possess a broad range of biological effects such as relieving sexual dysfunction and improving erectile function in rabbits and males, protecting myocardium against ischemia/reperfusion injury and treating hypertension and coronary heart disease. TT, claimed to be a testosterone booster, is a popular nutritional supplement in athletes and physically active men for enhancing gain in muscle mass, strength, and performance. Supplement of TT extracts increased serum testosterone levels on male rats, primates, rabbits and castrated rats. Our previous studies demonstrated that TT extracts improved exercise performance of rats with high-intensity endurance training and overload training by increasing plasma level of testosterone. However, different views still exist. TT has no significant influence on serum testosterone concentrations, strength, lean body mass, and exercise performance in elite rugby league players, resistance-trained males, and normal females as well as intact and castrated rats. Although there is little strong
evidence to prove that TT truly has the effects of testosterone booster and muscles anabolism promoter,7 TT extracts are still used constantly by many athletes. More clinical trials should be carried out to get a clear conclusion about the effectiveness of TT extracts.

Skeletal muscle is a highly dynamic tissue that responds to endogenous and external growth factor stimuli, among which insulin growth factor 1 (IGF-1) is one of the primary regulators affecting muscle growth, damage, repair, and regeneration. IGF-1 reduced aged-related wasting of skeletal muscle,15 and resistance training which is the most useful treatment for the loss of muscle mass and strength in elderly people upregulated the expression of IGF-1.16 IGF-1 injected soleus muscles of C57BL6 mice resulted on average 19% larger than the contralateral muscles and produced 16% more force.17 Local upregulation of IGF-1 has been also observed during muscle repair and regeneration in a variety of animal models of muscle damage.18 The action of IGF-1 is modulated by high-affinity binding proteins known as IGF binding proteins (IGFIBPs), and until now 7 of IGFIBPs (IGFBP1–7) have been found, among which IGFBP-3 is the most abundant in blood and tissue fluid combined with more than 80% IGF-1 normally.19 As the most important inhibitor of IGF-1 activity, the changes of IGFBP-3 level may have an indirect effect on muscle mass, muscle damage, and performance.

2. Materials and methods

2.1. Subjects

Fifteen male boxers (national second-level athletes, 2–3 years of training) were recruited from boxer team of Shanghai University of Sport Affiliated School of Sports in China. The boxers were randomly divided into exercise (E, n = 7) group and exercise plus TT (E + TT, n = 8) group, and 2 subjects in the E + TT group quit the experiment for leaving the school after 6 weeks. The baseline parameters of the participants are shown in Table 1. The experiment was approved and supervised by the Ethics Committee of Shanghai University of Sport (No. 2014002). An informed consent form was signed by the boxer who was equal or older than 18 years or by the guardian of the boxer who was younger than 18 years.

Table 1
The baseline parameters of the participants (mean ± SD).

| Parameter                              | E (n = 7) | E + TT (n = 8) |
|----------------------------------------|-----------|---------------|
| Age (year)                             | 16.6 ± 1.9| 16.1 ± 1.8    |
| Height (cm)                            | 172.7 ± 4.0| 174.0 ± 8.1  |
| Weight (kg)                            | 64.1 ± 6.6| 62.8 ± 15.2   |
| Body fat percentage (%)                | 9.6 ± 3.2 | 9.8 ± 2.4    |
| Maximum strength (1RM of barbell bench press, kg) | 72.0 ± 2.0| 71.0 ± 2.5    |
| Aerobic endurance (10,000 m race, min) | 41.8 ± 2.5| 42.2 ± 2.5    |
| Anaerobic endurance (peak power/body weight, W/kg) | 8.6 ± 1.3| 8.3 ± 1.1       |
| Anaerobic endurance (mean power/body weight, W/kg) | 7.2 ± 0.8| 6.7 ± 0.7       |

Abbreviations: E = exercise; E + TT = exercise plus TT; TT = Tribulus terrestris; RM = repetition maximum.

2.2. Administration and composition determination of TT extracts

The capsules of TT extracts (TT saponin > 40%) were purchased from Pronova Biocare company of Sweden. Two TT capsules a day (1250 mg, recommended dosage) were orally administered by male boxers of E + TT group every morning during 3-week high-intensity training and 3-week high-volume training, while placebo capsules of starch were taken by E group boxers. The capsules were administered in a double-blind fashion.

The compositions of TT extracts were detected by ultra-high performance liquid chromatography–quadrupole-time of flight mass spectrometry (UHPLC–Q-TOF/MS; Agilent Technologies, Santa Clara, CA, USA). Briefly, the powder of TT extracts from a capsule was dissolved in 70% alcohol and the supernatant was obtained to identify the compositions of TT extracts after ultrasonic extraction and centrifugation. An Agilent 1290 infinity UHPLC with binary pump, auto-sampler, thermostatted column compartment coupled with 6538 Q-TOF/MS system was used for the study on MS characterization of TT extracts. The mobile phase consisted of water containing 0.1% formic acid (A) and acetonitrile containing 0.1% formic acid (B) with the following gradient: 0–1.0 min, 5% B; 1.1–6.0 min linearly increased B from 5% to 20%; 6.1–9.0 min, linearly increased B from 20% to 50%; 9.1–13.0 min, linearly increased B from 50% to 95%; 13.1–18.0 min, 95%. Dual Agilent jet stream electrospray ion source was used and ran at both positive and negative modes. The temperature of gas was set as 350°C and the flow rate was 11 L/min. The nebulizer was 45 psi and the capillary voltage was set at 4000 V for positive mode and 3000 V for negative mode.

2.3. Exercise protocol

All athletes received similar 3-week high-intensity training and 3-week high-volume training separated by a 4-week rest. Besides special technical training, the main part of the high-intensity training was strength training including maximum strength training (twice a week, on Tuesday and Friday) and speed strength training (twice a week, on Monday and Thursday). For high-volume training, the boxers undertook endurance training (10,000 m race every day and low-to-moderate intensity rope skipping twice a week, on Tuesday and Friday), and special technical training and speed strength training similar to high-intensity training. Table 2 shows the training protocol of the boxer with high-intensity training and high-volume training.

2.4. Blood index assays

Fasting blood samples were collected before and at 40 h after the last training session to avoid the potential acute influence of the training on the levels of humoral factors. Blood levels of creatine kinase (CK), blood urea nitrogen (BUN), and hemoglobin (Hb) were detected by colorimetry (Nanjing Jiancheng Bio-engineering Institute, Nanjing, China). Plasma testosterone was determined using chemiluminescence immunoassay, while plasma DHT (ALPCO, New Hampshire, NH, USA), IGF-1
| High-intensity training                                      | High-volume training                                      | Note                                                                 |
|------------------------------------------------------------|-----------------------------------------------------------|----------------------------------------------------------------------|
| Monday  1. Barbell bench press, horizontal press, and power clean (40%–50% of 1RM, 6–8 sets × 12–15 rep); 2. Boxing (HR: about 190 bpm, 2 min × 8 times); 3. Punching sandbag (HR: about 190 bpm, 2.5 min × 4 times); 4. 3000 m race (HR: about 170 bpm); 5. Simulated actual combat (2.5 min × 10 times). | 1. 10,000 m race (less than 42 min); 2. Barbell bench press, horizontal press and power clean (40%–50% of 1RM, 6–8 sets × 12–15 rep); 3. Boxing (HR: about 190 bpm, 2 min × 8 times); 4. Punching sandbag (HR: about 190 bpm, 2.5 min × 4 times). | For training used barbell: 1st week: 40% of 1RM, 6 sets × 12; 2nd week: 40% of 1RM, 8 sets × 15; 3rd week: 50% of 1RM, 8 sets × 15. |
| Tuesday  1. Barbell squat and bench press (85% of 1RM, 6–8 sets × 2–3 rep); 2. Boxing (HR: about 190 bpm, 2 min × 8 times); 3. Punching sandbag (HR: about 190 bpm, 3 min × 5 times); 4. Rope skipping (HR: about 190 bpm, 2 min × 4 times); 5. 600 m race (HR: about 190 bpm, 2 min × 4 times). | 1. 10,000 m race (less than 42 min); 2. Rope skipping (HR: about 150 bpm, 30 min); 3. Boxing (HR: about 190 bpm, 2 min × 8 times); 4. Punching sandbag (HR: about 190 bpm, 3 min × 5 times); 5. Simulated actual combat (2.5 min × 10 times). | For training used barbell: 1st week: 6 sets × 2; 2nd week: 8 sets × 2; 3rd week: 8 sets × 3. |
| Wednesday 1. Rope skipping (HR: about 190 bpm, 2 min × 4 times); 2. 10,000 m race (less than 42 min); 3. Simulated actual combat (2.5 min × 10 times). | 1. 10,000 m race (less than 42 min); 2. Simulated actual combat (2.5 min × 10 times). | For training used barbell: |
| Thursday  1. Barbell bench press, horizontal press and power clean (50%–75% of 1RM, 6–8 sets × 8–10 rep); 2. Boxing (HR: about 190 bpm, 2 min × 4 times); 3. Punching sandbag (HR: about 190 bpm, 3 min × 5 times); 4. Football (60 min). | 1. 10,000 m race (less than 42 min); 2. Barbell bench press, horizontal press and power clean (50%–75% of 1RM, 6–8 sets × 8–10 rep); 3. Football (60 min). | For training used barbell: 1st week: 50% of 1RM, 6 sets × 8; 2nd week: 60% of 1RM, 8 sets × 10; 3rd week: 75% of 1RM, 6 sets × 8. |
| Friday  1. Barbell squat and bench press (85% of 1RM, 6–8 sets × 2–3 rep); 2. Boxing (HR: about 190 bpm, 2 min × 8 times); 3. Punching sandbag (HR: about 190 bpm, 3 min × 5 times); 4. Simulated actual combat (2.5 min × 10 times). | 1. 10,000 m race (less than 42 min); 2. Rope skipping (HR: about 150 bpm, 30 min); 3. Boxing (HR: about 190 bpm, 2 min × 8 times); 4. Punching sandbag (HR: about 190 bpm, 3 min × 5 times). | For training used barbell: 1st week: 6 sets × 2; 2nd week: 8 sets × 2; 3rd week: 8 sets × 3. |
| Saturday Same as Wednesday | Same as Wednesday | |

Abbreviations: bpm = beats per minute; HR = heart rate; rep = repetition; RM = repetition maximum.

2.5. Detection of muscle mass and fat mass

Muscle mass and fat mass of male boxers in the 2 groups were determined by dual-energy X-ray absorptiometry (DEXA) (Lunar Prodigy; GE, Madison, WI, USA). The test included a complete body scan of the boxers, in supine position, with the apparatus always regulated and operated by a technically trained professional. Lean soft tissue mass including total, appendicular, and trunk as well as fat mass and body fat percentage were obtained. The mass of total skeletal muscles was calculated by the formula: 1.115 × mass of appendicular lean soft tissues (kg) − 1.135.  

2.6. Determination of anaerobic performance

No. 894E Wingate anaerobic power bicycle (Monark Company, Vansbro, Sweden) was used to test anaerobic performance before and after the 2 trainings. After the rest period followed by a 5 min warm-up, the subject began to pedal maximally at the signal “go”. When the subject’s cadence reached 100 rpm at no resistance, the electromagnet released the weight pan and the 30 s test began. Computer software (Monark Anaerobic Test Software Version 3.0.1) was used to calculate PP, PP/BW, MP, MP/BW, and fatigue index (PP − Pmin)/PP × 100% throughout the 30 s test.

2.7. Statistical analysis

All values were expressed as mean ± SD and statistical significance was set as p < 0.05. Data were analyzed using SPSS Version 20.0 for windows (IBM Corp., Armonk, NY, USA) and entered into a two-way repeated measure analysis of variance (ANOVA) model with group (E vs. TT) as the between-subject effect and experimental condition (high-intensity and high-volume trainings) as the within-subject effect. The correlations of the change of IGFBP-3 with the changes of CK, MP, and MP/BW in E + TT boxers were evaluated by the correlation of Pearson’s χ² test.

3. Results

3.1. TT extract components by UHPLC–Q-TOF/MS

Alcohol-soluble components of TT extracts were separated by UHPLC (or UPLC), and their accurate molecular weights and molecular formula were identified through the information of positive ion and negative ion determined by Q-TOF/MS (Fig. 1). Then, names of the components of TT extracts were obtained by
Referring to literatures about TT extracts. As shown in Table 3, 22 constituents were identified from the TT extracts and the most abundant constituents were 6 kinds of TT saponins (25(R)-Spirostan-3,6,12-trione/25(R)-Spirostan-4-ene-3,12-dione, TT saponin A, gitogenin, TT saponin, tigogein, and diosgenin) which covered about 58.86% of the total peak area. Amino acids (valine and phenylalanine) and flavonoids (rutin, kaempferol-3-glucoside, quercetin, tricin, and kaempferol) comprised about 20.67% and 10.24% of the total peak area, respectively.

3.2. Effect of TT extracts on muscle mass and muscle damage

No significant post-training change was found in total muscle mass, total fat mass, and the percentages of total muscle and fat between the main effects for groups, experimental condition, or their interaction ($p > 0.05$) (Table 4).

For the change in CK after high-intensity training, the ANOVA showed significant main factors of group ($F(1, 11) = 5.53, p = 0.038$) and experimental condition ($F(1, 11) = 53.33, p < 0.001$). In addition, the interaction between the above 2 main effects indicated that the CK in the E + TT group following high-intensity training was significantly lower than that in the E group ($F(1, 11) = 5.00, p = 0.047$) while there was no significant difference between groups after high-volume training. Finally there was no significant post-training change in BUN between the main effects for groups, experimental condition, or their interaction ($p > 0.05$) (Table 5).

![Total ion chromatogram from UHPLC–Q-TOF/MS of Tribulus terrestris extracts in positive (A) and negative (B) modes. ESI = electrospray ionization; TIC = total ion chromatogram.](image-url)

### Table 3

Q-TOF/MS analysis results for components of TT extracts.

| No. | Name                              | RT (min) | Formula              | M ± X | Expected m/z | Experimental m/z | Error (ppm) | Area (%) |
|-----|-----------------------------------|----------|----------------------|-------|--------------|------------------|-------------|----------|
| 1   | Rutin                             | 7.727    | C_{12}H_{16}O_{16}   | M + H | 611.1624     | 611.1607         | −2.87       | 3.659    |
| 2   | Kaempferol-3-glucoside            | 7.931    | C_{21}H_{20}O_{11}   | M + H | 449.1092     | 449.1078         | −3.04       | 1.542    |
| 3   | Quercetin                         | 7.010    | C_{15}H_{10}O_{7}    | M + H | 303.0503     | 303.0499         | −1.24       | 1.674    |
| 4   | Valine                            | 0.790    | C_{5}H_{9}NO_{2}      | M + H | 116.0713     | 116.0706         | −0.73       | 18.438   |
| 5   | Valine acid                       | 7.206    | C_{10}H_{12}O_{4}    | M + H | 169.0481     | 169.0495         | 0.82        | 1.805    |
| 6   | TT amide                          | 9.064    | C_{15}H_{14}O_{3}    | M + H | 328.1211     | 328.1179         | −0.34       | 4.977    |
| 7   | Tricin                            | 10.010   | C_{15}H_{14}O_{3}    | M + H | 331.0821     | 331.0912         | −2.77       | 1.957    |
| 8   | 25(R)-Spirostan-3,6,12-trione/25(R)-Spirostan-4-ene-3,12-dione | 13.458 | C_{27}H_{38}O_{4} | M + H | 427.2881 | 427.2843 | −1.46 | 36.807 |
| 9   | Diosgenin                         | 11.836   | C_{15}H_{10}O_{5}    | M + H | 417.3369     | 417.3363         | −0.14       | 0.431    |
| 10  | Tigogenin                         | 12.533   | C_{15}H_{10}O_{5}    | M + H | 417.3369     | 417.3363         | −0.14       | 0.431    |
| 11  | TT saponin A                      | 13.630   | C_{15}H_{10}O_{5}    | M + H | 431.3196     | 431.3156         | −3.10       | 18.584   |
| 12  | Gitogenin                         | 13.950   | C_{15}H_{10}O_{5}    | M + H | 433.3325     | 433.3312         | −0.30       | 1.946    |
| 13  | Palmitic acid monoglyceride       | 14.477   | C_{15}H_{34}O_{4}    | M + H | 331.2847/353.2666 | 331.2843/353.2662 | −0.47/−1.17 | 0.070 |
| 14  | Phenyllalamine                     | 5.731    | C_{6}H_{8}O_{4}      | M + COOH | 210.0767 | 210.0772 | 0.67 | 2.228 |
| 15  | N-trans caffeoyltyramine          | 8.511    | C_{9}H_{10}O_{4}     | M + H | 298.1076     | 298.1085         | 0.31        | 1.325    |
| 16  | TT amide                          | 9.098    | C_{15}H_{10}O_{5}    | M + H | 326.1026     | 326.1179         | 0.50        | 1.613    |
| 17  | TT saponin                        | 9.253    | C_{15}H_{10}O_{5}    | M + H | 593.1295     | 593.1301         | 0.08        | 0.831    |
| 18  | Kaempferol                        | 9.375    | C_{15}H_{10}O_{5}    | M + H | 285.0405     | 285.0405         | −0.15       | 0.398    |
| 19  | Physcion                      | 10.044   | C_{15}H_{10}O_{5}    | M + COOH | 329.0667 | 329.0652 | −0.21 | 0.374 |
| 20  | 7-Hydroxy-4′-methoxyisoflavone    | 10.582   | C_{9}H_{8}O_{4}     | M + H | 267.0655     | 267.0663         | 0.35        | 0.201    |
| 21  | Emodin                            | 12.000   | C_{15}H_{10}O_{5}    | M + H | 269.0454     | 269.0455         | 0.40        | 0.071    |
| 22  | Kaempferol                        | 14.006   | C_{15}H_{10}O_{5}    | M + COOH | 367.2488 | 367.2490 | 0.60 | 0.809 |

Abbreviations: m/z = mass-to-charge ratio; M = molecule; RT = retention time; TT = Tribulus terrestris; X = uncertain material.
3.3. Effect of TT extracts on anaerobic performance

There was no significant post-training change in PP, PP/BW, and fatigue index between the main effects for groups, experimental condition, or their interaction (p > 0.05). For the change in MP after high-intensity training, the ANOVA yielded significant main effect of MP (F(1, 11) = 31.81, p < 0.001), but group effect of MP (F(1, 11) = 0.07, p = 0.799). However, a significant group × MP interaction (F(1, 11) = 7.79, p = 0.018) was found, and post hoc test revealed a significant decrease in MP for E group (p < 0.001) with no difference for E + TT group (p = 0.078). After high-volume training, there was also an interaction of group × MP (F(1, 11) = 4.81, p = 0.050) but not group (F(1, 11) = 0.06, p = 0.819). Post hoc test showed a decrease in MP for E group (p = 0.017) with no difference for E + TT group (p = 0.702) (Table 6). These results indicated that the decrease of MP after the 2 trainings was modulated by TT supplement (p < 0.05).

Similar results were found in MP/BW between groups. After high-intensity training, the ANOVA showed significant main effect of MP/BW (F(1, 11) = 30.78, p < 0.001) and interaction of group × MP/BW (F(1, 11) = 6.64, p = 0.026). Although there was no group discrepancy of MP/BW (F(1, 11) = 0.17, p = 0.685), post hoc test revealed a significant decrease in MP/BW for E group (p < 0.001) with no difference for E + TT group (p = 0.068). After high-volume training, significant main effects of MP/BW (F(1, 11) = 6.97, p = 0.023) and group × MP/BW interaction (F(1, 11) = 7.82, p = 0.017) were found. Although there is no group difference in MP/BW (F(1, 11) = 0.03, p = 0.867), post hoc test revealed a significant decrease in MP/BW for E group (p = 0.002) with no difference for E + TT group (p = 0.917) (Table 6). These results indicated that the decrease of MP/BW after the 2 trainings was modulated by TT supplement (p < 0.05).

3.4. Effect of TT extracts on plasma levels of testosterone, DHT, IGF-1, and IGFBP-3

No significant post-training change was found in testosterone (T), DHT, and IGF-1 between the main effects for groups, experimental condition or their interaction (p > 0.05). For the change in plasma IGFBP-3 after high-intensity training, the ANOVA showed significant group × IGFBP-3 interaction (F(1, 11) = 14.73, p = 0.003), but not group (F(1, 11) = 0.57, p = 0.467). Post hoc test revealed a decrease in IGFBP-3 for E + TT group (p = 0.005) with no difference in E group (p = 0.091). For high-volume training, significant main effects of IGFBP-3 (F(1, 11) = 5.07, p = 0.046) and group × IGFBP-3 interaction (F(1, 11) = 5.78, p = 0.035) but not group (F(1, 11) = 0.35, p = 0.568) were found. Post hoc test revealed

![Table 6](image)

|                      | Baseline        | E + TT          | High-intensity training | E + TT          | High-volume training | E + TT          |
|----------------------|-----------------|-----------------|-------------------------|-----------------|----------------------|-----------------|
| PP (W)               | 667.1 ± 112.6   | 616.1 ± 135.7   | 572.9 ± 84.8            | 551.2 ± 136.5   | 603.1 ± 54.3         | 632 ± 147.5     |
| PP/BW (W/kg)         | 10.8 ± 1.2      | 10.2 ± 1.4      | 9.3 ± 1.1               | 9.1 ± 1.1       | 9.5 ± 0.9            | 10.2 ± 0.9      |
| MP (W)               | 550.6 ± 75.5    | 498.7 ± 112.8   | 434.6 ± 69.5*           | 459.4 ± 122.3*  | 477.8 ± 49.9*        | 508.7 ± 103.2*  |
| MP/BW (W/kg)         | 8.9 ± 0.7       | 8.2 ± 0.9       | 7.1 ± 1.1*              | 7.5 ± 0.9*      | 7.5 ± 0.9*           | 8.2 ± 0.3*      |
| Fatigue index (%)    | 32.9 ± 7.5      | 35.1 ± 5.5      | 45.5 ± 12.9             | 36.9 ± 8.7      | 42.4 ± 12.5          | 35.7 ± 13.8     |

* p < 0.05, compared with the baseline values in the same group; † p < 0.05, compared with E group after the same training.

Abbreviations: BW = body weight; E = exercise; E + TT = exercise plus TT; MP = mean power; PP = peak power; TT = Tribulus terrestris.
a significant decrease in IGFBP-3 for E + TT group (p = 0.009) with no difference for E group (p = 0.912), indicating that the significant main factor of IGFBP-3 was modulated by TT supplement (p < 0.05) (Table 7).

3.5. No correlation between the change of IGFBP-3 with the changes of CK, MP, and MP/BW

The correlation coefficients of the change of IGFBP-3 with the changes of CK, MP, and MP/BW in E + TT groups were not statistically significant after the 2 trainings (data not shown).

3.6. No role of TT extracts on RBC and Hb

There was no statistical difference in RBC and Hb between E and E + TT groups after high-intensity or high-volume training (data not shown), which was similar to our previous results in rats² and report of Milasius et al.²¹

4. Discussion

4.1. Bioactive components of the TT extracts

Q-TOF mass spectrometry can provide high resolution and accurate mass measurement of both the precursor and product ions, thus it has become more and more popular for identification of drug components and metabolites.²² Twenty-two constituents were identified from the TT extracts, among which the most abundant constituents were TT saponins (25(R)-Spirostan-3,6,12-trione/25(R)-Spirostan-4-ene-3,12-dione and TT saponin A). The compositions and quantitative contents of TT extracts are not stable, depending on geographical region, climate²³ and part of herb, which may partly explain the divergent results of TT extracts from different studies.

4.2. TT extracts mitigated muscle damage and increased anaerobic performance while unchanged muscle mass

The levels of serum or plasma CK and BUN are commonly used to judge the severity of muscle damage and muscle protein catabolism, respectively.²⁴,²⁵ In the present study, mitigating muscle damage was found in male boxers who took TT extracts during high-intensity training. Decreasing contents of CK induced by TT extracts in rats were also found by another study.²⁶

TT extracts have been regarded as natural substances that can be added in diets in order to improve exercise performance and increase lean body mass.²⁷ Our previous studies demonstrated the increased effect of TT extracts on rat performance.⁹,¹⁰ In the present study, similar results were found in male boxers, that TT capsules (1250 mg, about 20 mg/kg a day for 3 weeks) significantly increased anaerobic performance (absolute and relative mean power). The promoting effect of TT on anaerobic performance was also demonstrated by Milasius et al.,²¹ who reported that anaerobic alactic muscular power and single muscular contraction power were significantly increased among youth men after consuming TT capsules (1875 mg, about 25 mg/kg a day) for 20 days. In contrast to the above data, other studies did not confirm the increased performance effect of TT on athletes, such as on resistance-trained men²¹ and rugby players.¹¹ In addition, no significant discrepancy of muscle mass and fat mass with or without TT extracts was demonstrated during training, similar to the results of our previous work in rats⁹ and other researchers,¹¹,¹² suggesting that the increased anaerobic performance by TT extracts was not mediated by increasing muscle mass.

4.3. TT extracts play no role on plasma testosterone and DHT

Testosterone is an important androgenic anabolic hormone for its long-term anabolic actions²⁸ and a close connection between testosterone and hypertrophy-type training was demonstrated.²⁹ So far, the results of TT extracts on blood androgens among humans and animals were both contradictory, and limited animal studies displayed a significant increase in blood testosterone levels after TT administration.⁴,⁵ but this effect was not found in humans except that TT was part of a combined supplement administration.³ The present study also demonstrated that TT extracts did not affect the blood testosterone and DHT of trained male boxers, indicating that plasma androgens were irrelevant to the enhanced effect of TT extracts on anaerobic performance.

Nutritional supplements recommended for competitive athletes to enhance their performance may be contaminated intentionally by androgenic-anabolic steroids,³⁰ which may lead to inadvertent doping in competitive sports. However, taking TT without any contaminations did not cause positive anti-doping tests.¹⁴ We believe that 2 things should be done for using TT safety and effectively among athletes and physically active men. One is to purify and separate TT bioactive components and the other is to avoid contamination of TT by androgenic-anabolic steroids.
4.4. Changes of plasma IGF-1 and IGFBP-3: possible mechanisms for TT extracts?

IGF-1 has great effect on muscle hypertrophy, muscle repair, and alleviating muscle damage. Over-expressed IGF-1 exhibited dramatically enlarged skeletal muscles in a number of animal models; in contrast, decreased IGF-1 in circulation and skeletal muscle was found to be dramatically reduced in muscle mass. Recently, a potential role of IGF-1 in protecting unloaded skeletal muscles from damage and accelerating muscle repair and regeneration was reported. In addition, the change of IGFBP-3 is considered a crucial aspect in modulating IGF-1 bioactivity. Serum concentrations of total IGF-1 decreased and IGFBP-3 increased were found after 16 weeks of resistance training in young women and in endurance-trained elite athletes at the end of exercise. However, increases, decreases, and no changes of circulating total IGF-1 and IGFBP-3 after both acute and chronic exercises have been reported because of the heterogeneity in subject characteristics, physical activity (type, intensity, and duration), and training state. In the present study, supplement of TT extracts decreased the plasma level of IGFBP-3 in male boxers after the 2 trainings. But there was no statistical significant correlation of the change of IGFBP-3 with the changes of CK, MP, and MBP in E + TT boxers, which may be related to the small sample size (n = 6). These results suggested that the effects of TT in trained male boxers may be mediated by decreasing IGFBP-3, ultimately increasing the bioactivity of IGF-1.

5. Conclusion

Taking 1250 mg capsules containing TT extracts did not change muscle mass and plasma levels of testosterone, DHT, and IGF-1, but significantly alleviated muscle damage and promoted anaerobic performance of trained male boxers, which may be associated with the decrease of IGFBP-3 rather than androgen in plasma.

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Authors’ contributions

YM carried out ELISA and DEXA studies, participated in statistical analysis, and drafted the manuscript; ZG carried out Wingate test and UHPLC–Q-TOF/MS analysis; XW conceived of the study, participated in its design and coordination, and helped to draft the manuscript. All authors have read and approved the final version of the manuscript, and agree with the order of presentation of the authors.

Competing interests

The authors declare that they have no competing interests.

References

1. Melnyk JP, Marcone MF. Aphrodisiacs from plant and animal sources—a review of current scientific literature. Food Res Int 2011;44:840–50.
2. Do J, Choi S, Choi J, Hyan JS. Effects and mechanism of action of a Tribulus terrestris extract on penile erection. Korean J Urol 2013;54:183–8.
3. Qureshi A, Naughton DP, Petroczi A. A systematic review on the herbal extract Tribulus terrestris and the roots of its putative aphrodisiac and performance enhancing effect. J Diet Suppl 2014;11:64–79.
4. El-Tantawy WH, Temraz A, El-Gindi OD. Free serum testosterone level in male rats treated with Tribulus alatus extracts. Int Braz J Urol 2007;33:554–8.
5. Singh N, Sair V, Gupta YK. Evaluation of the aphrodisiac activity of Tribulus terrestris Linn. in sexually sluggish male albino rats. J Pharmacol Pharmacother 2012;3:43–7.
6. Tyagi M, Aswar UM, Mohan V, Bodhankar SL, Zambare GN, Thakurdessai PA. Study of furostenol glycoside fraction of Tribulus terrestris on male sexual function in rats. Pharam Biol 2008;46:191–8.
7. Gauthaman K, Adaikan PG, Prasad RN. Aphrodisiac properties of Tribulus terrestris extract (protodioscin) in normal and castrated rats. Life Sci 2002;71:1385–96.
8. Gauthaman K, Ganesan AP. The hormonal effects of Tribulus terrestris and its role in the management of male erectile dysfunction—an evaluation using primates, rabbit and rat. Phytomedicine 2008;15:44–54.
9. Yin L, Wang XF, Cao XZ, Wang XH. The effects of Tribulus terrestris on the time of exhaustion in rats with high intensity training and its mechanism. J Shanghai Univ Sport 2013;37(5):73–7. [in Chinese].
10. Wang XH, Sun JY, Qu J, You SZ, Yang WJ. Effects of Tribulus terrestris on exercise ability, endocrine and immune functions of over-trained rats. J Shanghai Univ Sport 2010;34(1):46–9. [in Chinese].
11. Rogerson S, Riches CJ, Jennings C, Weatherby RP, Meir RA, Marshall-Gradisnik SM. The effect of five weeks of Tribulus terrestris supplementation on muscle strength and body composition during pre-season training in elite rugby league players. J Strength Cond Res 2007;21:348–53.
12. Antonio J, Uelmen J, Rodriguez R, Earnest C. The effects of Tribulus terrestris on body composition and exercise performance in resistance-trained males. Int J Sport Nutr Exerc Metab 2000;10:208–15.
13. Martino-Andrade AJ, Morais RN, Spercoski KM, Rossi SC, Vechi MF, Golin M, et al. Effects of Tribulus terrestris on endocrine sensitive organs in male and female Wistar rats. J Ethnopharmacol 2010;127:165–70.
14. Saudan C, Baume N, Emery C, Strahm E, Saugy M. Short term impact of Tribulus terrestris intake on doping control analysis of endogenous steroids. Forensic Sci Int 2008;178:e7–10.
15. McMahon CD, Chai R, Radley-Crab HG, Watson T, Matthews KG, Sheard PW, et al. Lifelong exercise and locally produced insulin-like growth factor-1 (IGF-1) have a modest influence on reducing age-related muscle wasting in mice. Scand J Med Sci Sports 2014;24:e422–35.
16. Luo L, Lu AM, Wang Y, Hong A, Chen Y, Hu J, et al. Chronic resistance training activates autophagy and reduces apoptosis of muscle cells by modulating IGF-1 and its receptors, Akt/mTOR and Akt/FOXO3a signaling in aged rats. Exp Gerontol 2013;48:427–36.
17. Ye F, Mathur S, Liu M, Borst SE, Walter GA, Sweeney HL, et al. Overexpression of insulin-like growth factor-1 attenuates skeletal muscle damage and accelerates muscle regeneration and functional recovery after disuse. Exp Physiol 2013;98:1038–52.
18. Barton ER, Morris L, Musaro A, Rosenthal N, Sweeney HL. Muscle-specific expression of insulin-like growth factor 1 counters muscle wasting in mdx mice. J Cell Biol 2002;157:137–48.
19. Alem E, Elshayeb A, Elhabachi N, Mansour AR, Gowily A, Hela A. Serum IGFBP-3 is a more effective predictor than IGF-1 and IGF-2 for the development of hepatocellular carcinoma in patients with chronic HCV infection. Oncol Lett 2012;3:704–12.
20. Kim J, Shen W, Gallagher D, Jones Jr A, Wang Z, Wang J, et al. Total-body skeletal muscle mass: estimation by dual-energy X-ray absorptiometry in children and adolescents. Am J Clin Nutr 2006;84:1014–20.
21. Milasius K, Dadeliene R, Skernevicius J. The influence of the Tribulus terrestris extract on the parameters of the functional preparedness and athletes’ organism homeostasis. Fiziol Zh 2009;55:89–96.
22. Wu JL, Leung EL, Zhou H, Liu L, Li N. Metabolite analysis of toosendanin by an ultra-high performance liquid chromatography-quadrupole-time of flight mass spectrometry technique. Molecules 2013;18:12144–53.
23. Dinchev D, Janda B, Evstatieva L, Oleszek W, Aslani MR, Kostova I. Distribution of steroidal saponins in Tribulus terrestris from different geographical regions. Phytochemistry 2008;69:176–86.
24. Clarkson PM, Kearns AK, Rouzier P, Rubin R, Thompson PD. Serum creatine kinase levels and renal function measures in exertional muscle damage. Med Sci Sports Exerc 2006;38:623–7.
25. Karabulut M, Sherk VD, Bemben DA, Bemben MG. Inflammation marker, damage marker and anabolic hormone responses to resistance training with vascular restriction in older males. Clin Physiol Funct Imaging 2013;33: 393–9.
26. Zhang S, Li H, Xu H, Yang SJ. Effect of gross saponins of Tribulus terrestris on cardiocytes impaired by adriamycin. Yao Xue Xue Bao 2010;45:31–6. [in Chinese].
27. Kreider RB, Wilborn CD, Taylor L, Campbell B, Almada AL, Collins R, et al. ISSN exercise & sport nutrition review: research & recommendations. J Int Soc Sports Nutr 2010;7:7. doi:10.1186/1550-2783-7-7
28. Wood RL, Stanton SJ. Testosterone and sport: current perspectives. Horm Behav 2012;61:147–55.
29. Schoenfeld BJ. Postexercise hypertrophic adaptations: a reexamination of the hormone hypothesis and its applicability to resistance training program design. J Strength Cond Res 2013;27:1720–30.
30. Cavalcanti Gde A, Leal FD, Garrido BC, Padilha MC, de Aquino Neto FR. Detection of designer steroid methylstenbolone in “nutritional supplement” using gas chromatography and tandem mass spectrometry: elucidation of its urinary metabolites. Steroids 2013;78:228–33.
31. Frystyk J. Exercise and the growth hormone-insulin-like growth factor axis. Med Sci Sports Exerc 2010;42:58–66.
32. Frost RA, Lang CH. Regulation of insulin-like growth factor-1 in skeletal muscle and muscle cells. Minerva Endocrinol 2003;28:53–73.
33. Arikawa AY, Kurzer MS, Thomas W, Schmitz KH. No effect of exercise on insulin-like growth factor-1, insulin, and glucose in young women participating in a 16-week randomized controlled trial. Cancer Epidemiol Biomarkers Prev 2010;19:2987–90.
34. Berg U, Enqvist JK, Mattsson CM, Carlsson-Skwirut C, Sundberg CJ, Ekblom B, et al. Lack of sex differences in the IGF-IGFBP response to ultra endurance exercise. Scand J Med Sci Sports 2008;18:706–14.
35. Copeland JL, Heggie L. IGF-I and IGFBP-3 during continuous and interval exercise. Int J Sports Med 2008;29:182–7.
36. Gatti R, De Palo EF, Antonelli G, Spinella P. IGF-I/IGFBP system: metabolism outline and physical exercise. J Endocrinol Invest 2012;35:699–707.