Modification of plasma glycosaminoglycans in long distance runners

M Contini, S Pacini, L Ibba-Manneschi, V Boddi, M Ruggiero, G Liguri, M Gulisano, C Catini

Background: It is well documented that exercise reduces the risk of thromboembolic disease, possibly by increasing the plasma concentration of anticoagulant-antithrombotic compounds.

Objectives: As plasma glycosaminoglycans (GAGs) play a role in the anticoagulant-antithrombotic potential of plasma, to examine the concentration and profile of these compounds in well trained, long distance runners and sedentary subjects.

Methods: Plasma GAGs were measured in 10 male, long distance runners and 10 sedentary counterparts before and after ergometric tests. GAGs were extracted, purified, and identified by electrophoretic and enzymatic methods, and measured as hexosamine.

Results: Plasma GAGs found in sedentary subjects were slow migrating heparan sulphates I and II, keratan sulphate I, and chondroitin 4–6-sulphate. Those found in trained athletes were slow migrating heparan sulphate I, chondroitin 4–6-sulphate (or keratan sulphate I), and fast migrating heparan sulphate. Total plasma concentrations of GAGs were higher in athletes than in sedentary subjects at rest. In sedentary subjects, plasma GAGs did not change after cycle ergometric exercise at 80% of their anaerobic threshold. However, the appearance of a novel band of heparan sulphate migrating faster than fast migrating heparan sulphate was observed in athletes after exercise.

Conclusions: Exercise changes the amount and profile of plasma GAGs; these changes may play a role in protecting subjects who practise aerobic sports against developing cardiovascular disease.

Methods

Subjects

Two groups of subjects were studied. The first was composed of 10 healthy men, aged 20–30, whose lifestyle could be defined as sedentary: none regularly engaged in strenuous exercise or hard physical work. The second was composed of 10 healthy, well trained, male, long distance runners, aged 18–34, with average training distance 90–130 km a week. Table 1 shows the physical characteristics of all the subjects.

Before being enrolled in the study, all subjects had a detailed medical examination, and only those showing no evidence of disease were included. They were asked not to take any drugs for at least three weeks before the day when blood was sampled or the day of the ergometry test, and not to change their usual dietary habits and level of physical activity. The dietary habits of the subjects did not differ significantly from those of the adult Italian population. All subjects gave written informed consent to the study according to the institutional review committee guidelines on human subjects.

Blood samples

For the evaluation of plasma GAGs, resting and fasting venous blood samples were taken from each sedentary subject and athlete, after they had avoided major physical activity for the preceding 24 hours. Each blood sample (9 ml), obtained by plastic syringe from the antecubital vein, was anticoagulated with 1 ml buffered sodium citrate (0.11 mol/l). Subjects were seated during blood withdrawal. Blood samples were taken at the beginning (phase A) and 1 hour after exercise (phase B).

Abbreviation: GAG, glycosaminoglycan
end (phase B) of the cycle ergometry test and in the recovery stage, when VCO₂ had returned to baseline (phase C).

Isolation and purification of GAGs from plasma
Platelet-poor plasma was obtained by centrifugation at 2500 g for 10 minutes at 20°C. The samples were lyophilised, weighted (dry weight), and subjected to lipid extraction with diethyl ether for 24 hours at 4°C. After rehydration of the samples, proteins were denatured and digested with Pronase E (Protease type XXV; Sigma Chemical Co, St Louis, Missouri, USA) for six hours at 20°C. The samples were lyophilised, further lyophilisation and lipid extraction with diethyl ether, for 10 minutes at 20°C. The samples were lyophilised, and the lipid extracts were dissolved in 0.1 M NaOH for 24 hours at 4°C. After rehydration, the samples were dialysed against running water for 48 hours, against deionised water, 80% ethanol.

Quantification and identification of plasma GAGs
GAGs were measured in terms of hexosamine content (mg/ml) using N-acetyl-d-glucosamine (Sigma) as standard. They were identified by the enzymatic/electrophoretic micro-method. The electrophoresis was carried out using 0.1 M barium acetate solution and cellulose acetate strips (Gelphore; Gelman Instruments, Wrightsville, Pennsylvania, USA), at 5 V/cm for 180 minutes. Electrophoretic patterns were visualised by staining with Alcian Blue 8GX 11 (Sigma), at 5 V/cm for 180 minutes. Then, each sample was dialysed against distilled water for 24 hours at 4°C. After further lyophilisation and lipid extraction with diethyl ether, the samples were solubilised, NaCl (0.03 M) was added, and GAGs were precipitated using cetylpyridinium chloride (Sigma), at 4°C for 24-48 hours. GAG–cetylpyridinium chloride complexes were removed by centrifugation at 1500 g for 20 minutes and washed three times with 80% ethanol saturated with sodium acetate and three times with 80% ethanol.

Statistical analysis
Plasma GAG concentrations were analysed by split-plot analysis of variance.

RESULTS
Table 2 gives total plasma GAG concentrations, measured in terms of hexosamine, in sedentary subjects and trained runners at rest.

The electrophoretic pattern of GAGs from sedentary subjects showed four bands, starting from the cathode (fig 1A). The first and second band were identified as slow migrating heparan sulphate I and II respectively, as they co-migrated with the appropriate standard and disappeared after treatment with nitrous acid (fig 1B). The third band disappeared after treatment with keratanase and was therefore identified as keratan sulphate I (fig 1C). The fourth band was identified as chondroitin 4-6-sulphate, as it co-migrated with the appropriate standard and disappeared after treatment with chondroitinase AC (fig 1D). The electrophoretic patterns were uniform, with few quantitative variations. As measured by densitometry, the first band, corresponding to the slow migrating heparan sulphate I, comprised, on average, 10% of the total GAGs (3.3 x 10⁻² mg/ml); the second band, identified as slow migrating heparan sulphate II, comprised 5% (1.6 x 10⁻² mg/ml); the third band, corresponding to keratan sulphate I, comprised 27% (9.1 x 10⁻³ mg/ml); and the fourth band, corresponding to chondroitin 4-6-sulphate, comprised 57% (18.9 x 10⁻³ mg/ml).

Plasma concentrations of GAGs in the trained athletes at rest were significantly higher (p<0.00005) than in the sedentary subjects. The electrophoretic pattern showed three bands which, starting from the cathode, were identified as first was an incremental ergometric test on an electromagnetically braked cycle ergometer (STS 3 Cardioline). After muscle adaptation and warm up, the initial work load of 50 W was increased by 30 W every two minutes, until exhaustion. During the test, heart rate was measured by continuous electrocardiogram recording. Standard techniques of open circuit spirometry were used to collect ventilatory and metabolic data. Respired gases were sampled through a low resistance breathing mask to measure oxygen uptake (V̇O₂), carbon dioxide output (VCO₂), expiratory ventilation (VE), tidal volume, and respiratory exchange ratio. These variables were continuously analysed and computed breath by breath during each exercise stage with a gas analyser (MMC4400TC; Sensor Medics, Yorba Linda, California, USA). This test allowed calculation, by the ventilatory method, of the anaerobic threshold and the maximum oxygen consumption (V̇O₂ MAX) for each subject (table 1). One week after the first test, all the subjects underwent a second ergometry test. This was performed for 12 minutes at a constant load equivalent to 80% of the anaerobic threshold assessed during the first test.

Table 1 Physical characteristics and ventilatory and metabolic variables of sedentary subjects (n = 10) and athletes (n = 10)

<table>
<thead>
<tr>
<th>Sedentary subjects</th>
<th>Athletes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>Median</td>
</tr>
<tr>
<td>Age (years)</td>
<td>24.0</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>63.0</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>171.3</td>
</tr>
<tr>
<td>BSA (m²)</td>
<td>1.73</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>21.3</td>
</tr>
<tr>
<td>V̇CO₂ (ml/min)</td>
<td>74.1</td>
</tr>
<tr>
<td>VE (l/min)</td>
<td>61.5</td>
</tr>
</tbody>
</table>

BSA, Body surface area, calculated as 0.007184 x height² x weight/height²; BMI, body mass index calculated as the Quetelet index as weight/height²; VO₂ MAX, maximum oxygen consumption; VE, expiratory ventilation; AT (%VO₂ MAX), percentage of VO₂ MAX corresponding to anaerobic threshold.

Table 2 Plasma glycosaminoglycans, measured in terms of hexosamine (mg/ml), in sedentary subjects and athletes, before (A) and after (B) the ergometric test, and when VCO₂ had returned to baseline (C)

<table>
<thead>
<tr>
<th>Sedentary subjects</th>
<th>Athletes</th>
</tr>
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<tbody>
<tr>
<td>Mean</td>
<td>Median</td>
</tr>
<tr>
<td>A</td>
<td>3.33</td>
</tr>
<tr>
<td>B</td>
<td>2.99</td>
</tr>
<tr>
<td>C</td>
<td>3.04</td>
</tr>
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Data are expressed as value x 10⁻². The means for the sedentary subjects are significantly lower than those for the athletes (p<0.00005).
slow migrating heparan sulphate I, chondroitin 4–6-sulphate (or keratan sulphate I), and fast migrating heparan sulphate (fig 2). The second band was identified as chondroitin 4–6-sulphate in six subjects, and as keratan sulphate I in four subjects. On average, the first band, slowly migrating heparan sulphate I, comprised 10% (4.9 $\pm$ 2.5 mg/ml) of total GAGs; the second band, chondroitin 4–6-sulphate comprised 45% (22.5 $\pm$ 2.5 mg/ml); the third band, fast migrating heparan sulphate, comprised 45% (21.9 $\pm$ 2.5 mg/ml).

After the second ergometry test, the total plasma GAG concentration did not change significantly in either sedentary subjects or athletes (table 2). However, athletes showed significant variations in the electrophoretic pattern, as shown by the appearance of a novel band in phases B and C, in all the athletes examined; this band migrated faster than fast migrating heparan sulphate (fig 3). As measured by densitometry, this band comprised on average 9% of total plasma GAGs (4.6 $\times$ 10$^{-5}$ mg/ml). Two of these bands were found in six subjects, whereas in the others only one preband was present. This preband behaved similarly to heparan sulphate when submitted to heparitinase and nitrous acid degradation (fig 3B–D).

DISCUSSION

“Beneficial” changes in the clotting state of the blood have previously been identified in athletes in aerobic sports such as marathon running and rowing. These changes may be similar to those that protect regularly exercising middle aged and older men against cardiovascular disease. We show here that long distance runners have different amounts and profile of plasma GAGs from sedentary healthy subjects. Firstly, the athletes had higher concentrations of total GAGs than the sedentary controls. Secondly, in the profile of electrophoretic migration, a band corresponding to fast migrating heparan sulphate was observed in athletes, but not in sedentary subjects. Moreover, after the cycle ergometry test, prebands migrated faster than fast migrating heparan sulphate bands in athletes, but not in sedentary subjects. As highly sulphated GAGs would precipitate in barium acetate, we assume that these fast migrating heparan sulphate species were low molecular mass compounds.

It has previously been shown that, low molecular mass heparan sulphates are powerful endogenous antithrombotic agents. Their effects are exerted primarily by inhibiting Factors Xa and IIa, acting as activators of antithrombin III. Interestingly, the profile of plasma GAGs in the athletes is similar to that of a novel antithrombotic drug, danaparoid sodium (Orgaran; Organon, Oss, the Netherlands); this is a low molecular mass heparinoid consisting of heparan sulphate, dermatan sulphate, and chondroitin sulphate, recently approved for prophylaxis of postoperative deep vein thrombosis in patients having hip replacement surgery.

The mechanism by which regular exercise increases the concentration of heparan sulphate and other GAGs remains
to be elucidated. Previous observations have suggested that a higher turnover rate of body cartilage matrix may account for the increase in hyaluronan and keratan sulphate in serum; this may correlate with the observation that exercise may occur at a higher level in athletes than in sedentary subjects. Other authors have suggested that intramuscular connective tissue is the main site of origin of plasma GAGs in athletes. However, the presence of low molecular mass compounds in athletes supports the hypothesis that a processive assembly of specific sequences in heparan sulphate may occur at a higher level in athletes than in sedentary subjects. Regular physical activity, in particular that typical of aerobic sports, may induce higher levels of expression of the genes involved in heparan sulphate biosynthesis leading to the generation of specific saccharide sequences.

In conclusion, we have shown that long distance runners have greater amounts and different composition of plasma GAGs compared with sedentary subjects; we postulate that these changes may be responsible for the variations in blood coagulation variables observed in athletes, which may reduce their risk of cardiovascular disease.

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REFERENCES

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Plasma glycosaminoglycans and exercise

Figure 3  Electrophoretic modification of plasma glycosaminoglycans after ergometric exercise in long distance runners. (A) Pattern before ergometry test; (B) modification after ergometry test (a new band on the right of the band corresponding to fast migrating heparan sulphate appeared); (C) this preband disappeared after treatment with nitrous acid, thus it was identified as heparan sulphate; (D) the sample was treated with keratanase before electrophoresis. DU, Arbitrary densitometric units.

This paper compares the plasma glycosaminoglycans of sedentary subjects and trained athletes. The two groups were characterised by clinical examination and ventilatory and metabolic variables. The glycosaminoglycans were identified by their electrophoretic mobility and digestion with specific enzymes or deamination with nitrous acid. The results obtained showed a clear difference in the plasma glycosaminoglycans of sedentary subjects and trained athletes, which may correlate with the observation that exercise reduces the risk of thrombosis. It is an interesting publication for those working in the field of the physiology of exercise and prevention of cardiovascular disease.

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