Plasma cortisol concentration in Thoroughbred horses during and after standardized exercise tests on a treadmill and effect of conditioning on basal cortisol values

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Summary
The behaviour of cortisol concentration in plasma of clinically normal horses performing two different exercise tests on a treadmill was investigated to study the possible value of measuring cortisol for performance diagnosis. In addition, the horses were exercised with different conditioning programmes and their effect on basal plasma cortisol concentration was measured. Finally, the repeatability of the cortisol values measured before and after exercise were examined. Horses were always exercised on a treadmill. The multiple step exercise test consisted of five gallop workouts lasting five minutes’ each. The velocity in the first step was 6.0 m/s. Each consecutive step velocity was increased by 0.5 m/s. The two-speed exercise test consisted of two runs. In the first run horses were galloped over 1,200 m at a constant velocity of 10, 10.5 or 11 m/s. Thereafter horses were walked for 30 minutes. The second run over 1,200 m was conducted at speeds of 13, 13.5 or 14 m/s. Each conditioning programme examined consisted of eleven repetitions of exercise of 5, 15 or 25 minutes’ duration at a velocity at which, mathematically, horses had a blood lactate concentration of 2.5 or 4 mmol/l. During the multiple step test the mean cortisol concentration in plasma showed a tendency to increase (p = 0.07). The mean peak of the cortisol concentration in plasma was measured 10 minutes after the test. Thereafter mean cortisol concentration continuously decreased and the day after the test values were similar to those before the test. After the first run of the two-speed test mean plasma cortisol concentration tended to increase (p = 0.08), while significant changes were registered after the second run (p < 0.01): It decreased up to the fifth minute after exercise (p < 0.05) and increased thereafter reaching a plateau between the 15th and 45th minute after exercise (p < 0.05). The morning after the test mean plasma cortisol concentration had returned to the values measured the morning before the test. None of the conditioning programmes had an effect on the mean plasma concentration of cortisol in the horses at rest. The mean coefficient of variation of plasma cortisol concentration in five horses before exercise was 34.1% and 36.4% after exercise.

The large individual variability of plasma cortisol concentrations before and after exercise does not allow a good repeatability of results, and enforces the need for multiple blood sampling during exercise. The multiple step exercise test as well as the two-speed exercise test induced increases of the plasma cortisol concentration after exercise. The changes measured supply a basis for studies on treadmills on the value of monitoring the plasma cortisol concentration of sport horses for performance diagnosis.

Keywords: horse, exercise, cortisol, blood plasma, lactate

Konzentration von Plasmakortisol bei Pferden während und nach Laufband-Belastungstests und Einfluß von Training auf den Plasmakortisolgehalt bei Pferden unter Ruhebedingungen


Der festgestellte Variationskoeffizient ist hoch. Dies bedeutet eine schlechte Wiederholbarkeit für die Kortisolmessungen und betont die Bedeutung der mehrfachen Blutentnahme bei Belastung. Die Veränderungen der Kortisolkonzentration im Blut bei den Belastungstests dienen als Referenz für weitere Studien, um den Nutzen der Messung dieser Variablen für die Leistungsdagnostik zu prüfen.

Schlüsselwörter: Pferd, Belastung, Kortisol, Blutplasma, Laktat
Introduction

Standardized exercise tests are used for performance diagnosis of horses (Persson 1983). Variables normally determined in this test are blood lactate concentration, heart rate and maximal oxygen consumption capacity. Other biochemical variables like cortisol could be measured too to possibly provide additional information on performance capacity. Physical stress influences the cortisol response of the horse (Thornton 1985). Glucocorticoids are anti-inflammatory, increase glycogen deposition, stimulate lipolysis and enhance hepatic gluconeogenesis (Keelie et al. 1982). The significance for performance diagnosis of the changes in cortisol concentration with horses during and after exercise is not clear. Literature data is controversial. Garcia and Beech (1986) and Church et al. (1987) did not find training-induced changes of plasma cortisol after exercise, while Müller et al. (1990), Freestone et al. (1991) found smaller increases and a faster return to base levels after exercise following training and Grosskopf et al. (1983) observed a similar pattern of cortisol concentration in horses with a better performance after endurance competition compared to pre-training levels and to horses with lower performance. Wilson et al. (1991) reported on a decrease of baseline cortisol concentration in thoroughbred horses in the initial stage of the training for racing and that horses fit for racing had lower baseline cortisol concentration than when in early race training. But, Persson et al. (1980) found that standardbred horses with poor performance had also lower basal cortisol concentration and the increase of cortisol after ACTH application was smaller than in horses performing at an expected level, and Baker et al. (1982) could not find differences in the cortisol concentration in plasma between resting thoroughbred horses performing at the expected level and not performing well. Finally, Golland et al. (1996) demonstrated that standardbred horses with clinical signs of experimentally induced overtraining had a lower increase of cortisol in plasma in response to exercise as compared with values before overtraining and with a control group. They also reported that basal plasma cortisol concentration and the via ACTH injection induced plasma cortisol reaction were not different between overtrained and control horses.

The behaviour of cortisol concentration in plasma of clinically normal horses performing two different exercise tests on a treadmill was investigated to study the possible value of measuring cortisol for performance diagnosis. In addition, the horses were exercised with different conditioning programmes and their effect on basal plasma cortisol concentration was measured. Finally, the repeatability of the cortisol values measured before and after exercise was examined.

Materials and methods

Horses

Five conditioned thoroughbred horses 3.4±0.5 (SD) years old, weighing 467±29 (SD) kg trained for a period of two months were used for the studies. Horses were kept in 3 x 3 m boxes at night, at day they were turned out on pasture. They were fed a concentrate of 4.5 kg daily (Reform-Mix; Hoeveler Spezialfutterwerke, Germany), and 5 kg of silage (in a relation of 2:1 grass to cornsilage). Hay, straw and water were always available. All the exercise tests and workouts were performed on a high-speed treadmill (Mustang®, Kagra AG, Fahrwangen, Switzerland) with horses wearing a heart rate meter (Polar Electro OY, Finland). The horses were always warmed up following the same routine: 5 minutes walk at 1.6 m/s and 5 minutes trot at 3.4 m/s without incline. After exercise they walked during the whole blood sampling period at 1.6 m/s without incline.

Design of studies

Study to determine the effect of standardized exercise tests

Horses were submitted to two different test protocols: a two-speed exercise test and a multiple step exercise test. During and after the multiple step test blood samples for measuring cortisol were collected at 7:00 a.m., before exercise but after warm up, within 10 seconds after each step and in the 5th, 10th, 15th, 30th, 45th and 60th minute, and at 7:00 a.m. on the morning after the test.

To examine the effects of the two-speed exercise test blood samples for measuring cortisol were collected at 7:00 a.m., before exercise but after warm up, in the 1st, 5th, 10th and 15th minute after both runs and in the 30th, 45th and 60th minute and at 7:00 a.m. on the morning after the second run of each test. Blood samples for lactate determination were taken in the 1st, 3rd, 5th minute after both runs of a test and additionally in the 7th, 10th and 12th minute after the second run of each test.

Study to determine effect of conditioning programmes

In a randomized 5x6 latin square cross over study design (five horses x 6 conditioning programmes), horses were exercised at v2.5 or v4 during 5, 15 or 25 minutes for 11 times with a one day rest between two consecutive workouts (v2.5 or v4; velocity at which, mathematically, a lactate concentration of 2.5 or 4 mmol/l blood is determined, when it is run under defined conditions). Before each conditioning programme horses performed a standardized exercise test to determine their individual v2.5 and v4. To examine the effect of the conditioning programme on cortisol concentration in plasma, blood samples were always taken from the horses at 7:00 a.m., before and after each conditioning programme. Horses had about one week without standardized exercise between conditioning programmes. During this period horses were walked and trotted on the treadmill every second day.

Study to determine coefficient of variation of cortisol

Blood was taken from five of the horses before exercise at 7:00 a.m. and within 10 seconds after the end of exercise. This was repeated four times for each horse on days 1, 7, 13 and 19 of a single conditioning programme. Duration and intensity of the exercise were the same for a single horse, but differed between horses (Table 1). The duration of exercise was 5, 15 or 25 minu-
Plasma cortisol concentration in Thoroughbred horses during and after standardized exercise tests on a treadmill

tests, and horses were run at $v_{2.5}$ or $v_4$. The $v_{2.5}$ and $v_4$ of each horse was determined with a standardized multiple step exercise test two days before the beginning of the conditioning period.

Tab. 1: Duration and velocity run ($v_{2.5}$ or $v_4$) by horses on a treadmill during four exercise sessions/horse and mean (± SD) heart rate during, blood lactate concentration after exercise and plasma cortisol concentration before and after exercise.

<table>
<thead>
<tr>
<th>Horse</th>
<th>Duration (minutes)</th>
<th>$v_{2.5}$ (m/s)</th>
<th>$v_4$ (m/s)</th>
<th>Heart rate (beats/min)</th>
<th>Blood lactate (mmol/l)</th>
<th>Cortisol (nmol/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5</td>
<td>7.5</td>
<td>-</td>
<td>160±10</td>
<td>2.74±0.59</td>
<td>248±89</td>
</tr>
<tr>
<td>2</td>
<td>5</td>
<td>-</td>
<td>6.9</td>
<td>154±14</td>
<td>5.34±1.40</td>
<td>306±72</td>
</tr>
<tr>
<td>3</td>
<td>25</td>
<td>-</td>
<td>7.6</td>
<td>164±3</td>
<td>3.14±0.53</td>
<td>271±74</td>
</tr>
<tr>
<td>4</td>
<td>15</td>
<td>-</td>
<td>7.4</td>
<td>173±9</td>
<td>3.55±0.27</td>
<td>287±78</td>
</tr>
<tr>
<td>5</td>
<td>15</td>
<td>-</td>
<td>8.6</td>
<td>177±1</td>
<td>2.56±0.43</td>
<td>433±245</td>
</tr>
</tbody>
</table>

Standardized exercise tests

Multiple step exercise test
It consisted of five gallop workouts of five minutes’ duration each. Between two consecutive steps there was a resting period of 60 s. The velocity in the first step was 6.0 m/s. Each consecutive step was increased by 0.5 m/s. $v_{2.5}$ and $v_4$ were determined from the individual blood lactate concentration-running speed relationship by exponential regression equation (Galloux 1991).

Two-speed exercise test
This test consisted of two runs. In the first run horses were galloped over 1,200 m at a constant velocity of 10, 10.5 or 11 m/s. Thereafter horses were walked for 30 minutes. The second run over 1,200 m was conducted at speeds of 13, 13.5 or 14 m/s (Table 2). The speed at which horses ran depended on their $v_4$ values determined with the multiple step test.

Tab. 2: Running speed of horses in the 1st and 2nd run over 1,200 m of the two-speed exercise test on a treadmill, maximal blood lactate concentration after each run and mean heart rate during exercise.

<table>
<thead>
<tr>
<th>Horse</th>
<th>1st run</th>
<th>2nd run</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Velocity (m/s)</td>
<td>Maximal blood lactate (mmol/l)</td>
</tr>
<tr>
<td>1</td>
<td>11.0</td>
<td>3.80</td>
</tr>
<tr>
<td>2</td>
<td>11.0</td>
<td>6.15</td>
</tr>
<tr>
<td>3</td>
<td>10.0</td>
<td>4.25</td>
</tr>
<tr>
<td>4</td>
<td>10.0</td>
<td>5.75</td>
</tr>
<tr>
<td>5</td>
<td>10.5</td>
<td>3.95</td>
</tr>
<tr>
<td>6</td>
<td>10.0</td>
<td>8.30</td>
</tr>
<tr>
<td>7</td>
<td>10.5</td>
<td>4.75</td>
</tr>
</tbody>
</table>
test. Horses with lower \( v_4 \) values galloped at 10 m/s in the first run and at 13 m/s in the second run, and horses with the higher \( v_4 \) values galloped at 11 m/s in the first and 14 m/s in the second run.

**Blood sample handling**

For the cortisol and lactate measurement blood was collected from a jugular vein into evacuated Na-heparinate containing tubes. Within 30 minutes samples were centrifuged at 6,000 \( \times \) g for 10 minutes, and the plasma was transferred to plastic vials and kept stored at \(-20^\circ\text{C}\) until analysis, which was normally done within two months.

For the lactate determination 20 \( \mu\text{l} \) of blood were transferred as soon as possible after collection into vials with 200 \( \mu\text{l} \) ice-cold 0.6 n perchloric acid. Samples were centrifuged for 5 minutes by 12,000 \( \times \) g and the supernatant was transferred to empty vials and stored at 4\(^\circ\text{C}\) until analysis, normally within two days.

**Analysis**

**Cortisol**

Plasma concentration of cortisol was analyzed in duplicate with an immunoenzymatic assay (Boehringer Mannheim Immuno-diagnostics no. 649945). The coefficient of variation between days was 8%.

**Lactate**

The blood lactate analysis was done with an EPOS 5060 analyzer (Eppendorf-Netheler-Hinz) using an enzymatic test kit (Behring, OSUA 40). The coefficient of variation for this enzymatic method from day to day was 7.5% at a lactate concentration of 2.15 mmol/l (Precipath\textsuperscript{®} S Boehringer Mannheim, Nr. 125202) and 3.7% at 4.4 mmol/l (SIGMA lactate, Nr. 82610).

**Statistics**

Data shown are means of duplicate analyses for each sample. Data are presented as mean ± standard deviation. The coefficient of variation of the variables before and after exercise was calculated dividing the standard deviation through the mean and multiplying by 100.

To determine whether an exercise test had an effect an analysis of variance for repeated measures was applied. Effects of conditioning programmes on variables were examined with analysis of variance for repeated measures. If significant effects were found, Fishers’ test was used post-hoc. \( p<0.05 \) was used as a level to denote significant differences.

**Results**

During the multiple step test there was a tendency for the cortisol concentration in plasma to increase \((p = 0.07; \text{Figure 1})\). The peak cortisol concentration in plasma was measured 10 minutes after the test \((\text{Figure 2})\). Thereafter the cortisol concentration decreased continuously \((p < 0.01)\). On the morning after the test the mean cortisol concentration was back to the values measured on the morning before the test \((\text{Figure 2})\).

![Fig. 1: Cortisol in plasma of horses before and during multiple step exercise test (mean ± standard deviation; \(n = 5\))](image1.png)

**Kortisol im Plasma von Pferden vor und während eines Mehrstufen-Belastungstests (Mittelwert ± Standardabweichung; \(n = 5\))**

![Fig. 2: Cortisol in plasma of horses after multiple step exercise test (mean ± standard deviation; \(n = 5\))](image2.png)

**Kortisol im Plasma von Pferden nach einem Mehrstufen-Belastungstest (Mittelwert ± Standardabweichung; \(n = 5\))**

After the first run of the two-speed test plasma cortisol concentration tended to increase \((p = 0.08)\). After the second run the mean plasma cortisol concentration changed significantly \((p < 0.01)\). Initially it decreased in the 5th minute after the run and thereafter it increased reaching a plateau between the 15th and 45th minute after the run \((p < 0.05)\) respectively compared with values measured before the 2nd run. The mean cortisol concentration in plasma measured at 7:00 a.m. on the morning after the test was not different from the value measured at 7:00 a.m. on the testing day \((\text{Figure 3})\).

None of the conditioning programmes had an effect on the plasma concentration of cortisol at 7:00 a.m. \((\text{Table 3})\).
The coefficient of variation of cortisol before exercise of single horses ranged from 24% to 57%. After exercise the values were between 22% and 60%. After exercise, in all horses plasma cortisol concentration was higher as compared with values before exercise taken at 7:00 a.m. (Table 1).

Discussion and conclusions

The two-speed and especially the multiple-step exercise tests are used routinely under field conditions by our group to estimate endurance of sport horses by calculating their \( v_4 \) from the blood lactate-running speed relation established with the tests (\( v_4 = \) velocity run under defined conditions at which, mathematically, a blood lactate concentration of 4 mmol/l is determined; von Wittke et al. 1994; Werkmann et al. 1996; Guhl et al. 1996 a+b; Lindner 1998). It has been shown that these two types of exercise tests increase plasma cortisol concentration in thoroughbred and standardbred horses when run on a track (Lindner 1992; Lindner et al. 2000). Both types of exercise tests induced increases of the plasma cortisol concentration after exercise on the treadmill too. However, changes produced by the two-speed test were smaller than those by the multiple-step test. The different magnitude of reaction of the cortisol concentration in plasma of horses seems to indicate that duration of exercise induces a larger increase than speed of exercise. But, this assumption may be valid for treadmill conditions only, because speed of the second run of the two-speed test and therefore exercise stress may not increase sufficiently to examine a horse with high endurance. It has been stated that measurable increases of plasma or serum cortisol concentration in horses after exercise only occur if exercise load is sufficiently stressful (Thornton 1985). In man this depends on the fitness of an athlete (Kjaer 1996), and it may also hold for horses (Snow and McKenzie 1977; Müller et al. 1990; Freestone et al. 1991). From the results of the few studies available with sufficient sampling times after exercise it can be deduced that the peak cortisol concentration will normalize within the first 60 minutes after exercise (Flisinska-Bojanowska et al. 1974; Colborn et al. 1991). This is substantiated by the results of this study.

A singular behaviour of the plasma cortisol concentration was observed after the 2nd run of the two-speed exercise test: first there was a pronounced decline of the mean plasma cortisol concentration in the 5th minute and thereafter only, the mean cortisol concentration increased and showed a plateau between the 15th and 45th minute after exercise. The plasma cortisol concentration of four out of five horses reacted this way, and in the other horse the drop was protracted by five minutes followed by an increase as well. Between the 1st and the 2nd run of the test 30 minutes of time elapsed. It may be that the cortisol secretion after an exercise stimulus is delayed within a certain amount of time if another exercise stimulated the cortisol secretion before. This behaviour certainly documents the importance of consecutive blood sampling in short intervals after exercise.

### Table 3: Cortisol concentration in plasma (nmol/l) of horses at 7:00 a.m. before and after different conditioning programmes (n=5; mean ± standard deviation)

<table>
<thead>
<tr>
<th>Time of blood sampling</th>
<th>( v_{2.5} ) 5 minutes</th>
<th>( v_{2.5} ) 15 minutes</th>
<th>( v_{2.5} ) 25 minutes</th>
<th>( v_4 ) 5 minutes</th>
<th>( v_4 ) 15 minutes</th>
<th>( v_4 ) 25 minutes</th>
</tr>
</thead>
<tbody>
<tr>
<td>before</td>
<td>280±80</td>
<td>295±72</td>
<td>296±89</td>
<td>297±94</td>
<td>257±89</td>
<td>242±52</td>
</tr>
<tr>
<td>after</td>
<td>219±69</td>
<td>265±92</td>
<td>304±112</td>
<td>262±63</td>
<td>251±80</td>
<td>219±51</td>
</tr>
</tbody>
</table>

Fig. 3: Cortisol in plasma of horses before and after the 1st and 2nd run of a two-speed exercise test (mean ± standard deviation; n = 5)

Kortisol im Plasma von Pferden vor und nach dem ersten und zweiten Lauf eines Zweigeschwindigkeiten-Belastungstests (Mittelwert ± Standardabweichung; n = 5)
There was no measurable effect of the examined conditioning programmes on mean plasma cortisol levels in resting horses (7:00 a.m. samples). It may be that exercise workload of the conditioning programmes applied was not large enough for the thoroughly prepared horses to induce adaptations which would have been reflected in the cortisol concentration in plasma. This is the most reasonable explanation for the result because none of the other physiological and biochemical variables measured in these horses demonstrated an effect of the conditioning programmes (Ferlazzo et al. 1996; Werkmann et al. 1996). Only the blood lactate concentration after exercise at \( v_2 \), during 25 minutes’ duration decreased during the conditioning period (Werkmann et al. 1996). It is also possible, that the cortisol concentration in plasma of resting horses does not allow their performance capacity to be determined, and an adaptation would only have been demonstrable evaluating the response to standardized exercise. Another approach could have been to measure the response to the application of ACTH (James et al. 1970; Rossdale et al. 1982; Linden et al. 1991a; Golland et al. 1996). However, this method may not be readily applicable to sound sport horses, and may be more appropriate for the diagnosis in horses not performing at the expected level.

The rather high variability of the plasma cortisol concentration within a horse before and after exercise observed in this study may also account for the lack of significant effects. Linden et al. (1990) calculated a coefficient of variation for plasma cortisol concentration in five horses during five consecutive days between 21% and 64%, and Baker et al. (1982) reported on the coefficient of variation of one horse between days being 20%. The reason for this variability is most likely that the plasma cortisol concentration in horses exhibits an ultradian rhythm superimposed upon the circadian rhythm, and that the individual rhythms are variable within the same horse (Evans et al. 1977). Reproducibility of plasma cortisol measurements in experiments can be expected to be poor, and the value of measuring a single cortisol concentration for performance diagnosis questionable. This is the opinion of many authors (Evans et al. 1977; Thornton 1985; Wilson et al. 1991). However, in most studies only one blood sample is taken in resting horses (Baker et al. 1982; Linden et al. 1990), and after exercise (James et al. 1970; Grosskopf et al. 1983; Snow et al. 1983; Linden et al. 1991a; Desmecht et al. 1996), or more than 15 minutes are within two blood samples (Rossdale et al. 1982; Garcia and Beech 1986; Church et al. 1987; Freestone et al. 1991).

In conclusion, the large individual variability of plasma cortisol concentrations before and after exercise does not allow a good repeatability of results. Exercising conditioned horses repeatedly for up to 25 minutes’ duration at \( v_2 \) and \( v_4 \) during a period of three weeks does not affect plasma cortisol concentrations sampled in horses at 7:00 a.m.. The multiple step exercise test as well as the two-speed exercise test induced increases of the plasma cortisol concentration after exercise. The range of changes measured may supply a basis for future studies on treadmills on the value of monitoring the plasma cortisol concentration of sport horses for performance diagnosis.

Acknowledgements

We are very grateful to the Verein zur Förderung der Forschung im Pferdesport, Wissenschaftliche Gesellschaft der Schwarzwal-Tierklinik, Höveler Spezialfutterwerke and Horst Dieter Beyer for the material and financial support, to Dr. Bidlingmaier of the Institut für Klinische Biochemie of the University of Bonn for allowing us to run all the lactate analysis in his laboratory. This work was supported by a grant (60%) from MURST, Italy.

References


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