Analysis of the aerobic-anaerobic transition in elite cyclists during incremental exercise with the use of electromyography

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Abstract

Objectives—To investigate the validity and reliability of surface electromyography (EMG) as a new non-invasive determinant of the metabolic response to incremental exercise in elite cyclists. The relation between EMG activity and other more conventional methods for analysing the aerobic-anaerobic transition such as blood lactate measurements (lactate threshold (LT) and onset of blood lactate accumulation (OBLA)) and ventilatory parameters (ventilatory thresholds 1 and 2 (VT1, and VT2)) was studied.

Methods—Twenty eight elite road cyclists (age 24 (4) years; \( V_{\text{O}_2\text{MAX}} \) 69.9 (6.4) ml/kg/min; values mean (SD)) were selected as subjects. Each of them performed a ramp protocol (starting at 0 W, with increases of 5 W every 12 seconds) on a cycle ergometer (validity study). In addition, 15 of them performed the same test twice (reliability study). During the tests, data on gas exchange and blood lactate levels were collected to determine VT1, VT2, LT, and OBLA. The root mean squares of EMG signals (rms-EMG) were recorded from both the vastus lateralis and the rectus femoris at each intensity using surface electrodes.

Results—A two threshold response was detected in the rms-EMG recordings from both muscles in 90% of subjects, with two breakpoints, EMGT1, and EMGT2, at around 60–70% and 80–90% of \( V_{\text{O}_2\text{MAX}} \) respectively. The results of the reliability study showed no significant differences (p>0.05) between mean values of EMGT1 and EMGT2 obtained in both tests. Furthermore, no significant differences (p>0.05) existed between mean values of EMGT1 in the vastus lateralis and rectus femoris, and VT1 and LT (62.8 (14.5) and 69.0 (6.2) and 64.6 (6.4) and 68.7 (8.2)% of \( V_{\text{O}_2\text{MAX}} \) respectively), or between mean values of EMGT2 in the vastus lateralis and rectus femoris, and VT2 and OBLA (86.9 (9.0) and 88.0 (6.2) and 84.6 (6.5) and 87.7 (8.4)% of \( V_{\text{O}_2\text{MAX}} \) respectively).

Conclusion—rms-EMG may be a useful complementary non-invasive method for analysing the aerobic-anaerobic transition (ventilatory and lactate thresholds) in elite cyclists.

Keywords: electromyography; muscle; ventilatory threshold; lactate threshold; cycling; metabolic response

Surface electromyography (EMG) is an acceptable method for quantifying the total activity of working muscles and for estimating muscle fatigue non-invasively. An increase in EMG activity has been shown to reflect the recruitment of additional motor units and an increase in motor unit rate coding to compensate for the deficit in contractility resulting from impairment of fatigued motor units, as the strength of a muscle contraction increases. Along this line of thought, several studies have shown the existence of a non-linear increase in EMG during the aerobic-anaerobic transition phase in ergometer cycling. Indeed, an EMG threshold (EMGT) has been suggested to occur in the vastus lateralis, vastus medialis, rectus femoris, gastrocnemius, biceps femoris, and soleus of healthy not highly trained subjects during progressive tests on a cycle ergometer. The EMGT, in turn, would represent the point where an increased contribution from fast twitch motor units occurs to maintain the required energy supply for muscle contraction. In other studies, in contrast, a linear relation has been reported between EMG and exercise intensity in ergometer cycling.

In addition to these controversial findings, some questions still remain unanswered. First, only one study has assessed the test-retest reliability of the EMG response to progressive exercise. In this study, the EMG response was reported to be repeatable but linear (not threshold-like) during an incremental treadmill test. As reliability is an integral part of validity, no study has accurately determined the validity of the EMG method for analysing the aerobic-anaerobic transition. Furthermore, little research has been conducted using elite athletes (cyclists) as subjects, and the findings that have been reported seem somewhat controversial.

The aim of this study was to investigate the validity and reliability of EMG as a new non-invasive determinant of the metabolic response to incremental exercise in elite cyclists. We studied the relation between EMG activity and other more conventional methods of analysing the aerobic-anaerobic transition, such as blood lactate measurements (lactate threshold (LT) and onset of blood lactate accumulation (OBLA)) and ventilatory parameters (ventilatory thresholds 1 and 2 (VT1, and VT2)).

Methods

SUBJECTS

Twenty eight elite male cyclists (age 24 (4) years; height 177.1 (5.2) cm; body mass 67.2 (6.0) kg; all values mean (SD)) participated in

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this study. Sixteen subjects were professional road cyclists with a minimum competition experience of three years, and some of them had won several professional races. The other 12 were elite road cyclists (competition experience at least two years in the amateur category). Written informed consent was given before participation in the experiments, in accordance with the institutional human subjects guidelines (Complutense University of Madrid).

STUDY PROTOCOL

Before each exercise testing session, subjects were familiarised with the equipment and procedures used in this investigation. In addition, they were previously instructed to refrain from intense training during the day before testing. Fifteen subjects were randomly selected for the reliability study. Each of them performed two exercise tests on a bicycle ergometer (Ergometrics 900; Ergo-line, Barcelona, Spain) on different occasions and separated from each other by a period of no more than five days. Each of the two tests consisted of a ramp protocol until exhaustion, starting at 0 W. The power output was increased by 5 W every 12 seconds and pedalling cadence was kept constant at 70–80 rpm. The selection of a ramp-like protocol instead of a graded steady state test was chosen for two reasons: (a) in previous research conducted in our laboratory this type of protocol was used to analyse physiological responses—that is, ventilatory thresholds and lactate kinetics—in elite professional cyclists during incremental exercise; (b) previous research showing a non-linear increase in the EMG response to incremental exercise used exercise protocols comparable with that used in our investigation. For the validity study, each of 28 subjects performed a single bicycle ergometer test following the above protocol. Exercise tests were terminated (a) voluntarily by the subjects, (b) when pedalling cadence could not be kept at least at 70 rpm, or (c) when established criteria of test termination were met. Each test was performed under similar conditions (21–24°C and 45–55% relative humidity).

ANALYSIS OF EXPIRED GAS AND DETERMINATION OF VENTILATORY THRESHOLD

During the tests, gas exchange data were collected continuously using an automated breath by breath system (CPX; Medical Graphics, St Paul, Minnesota, USA). The measuring instruments were calibrated before each test and the necessary environmental adjustments made. VT1 was determined using the criteria of an increase in the ventilatory equivalent for oxygen (VE/VO₂) with no increase in the ventilatory equivalent for carbon dioxide (VE/VCO₂) and the departure from linearity of VE, whereas VT2 was determined by using the criteria of an increase in both VE/VO₂ and VE/VCO₂. Two independent observers detected VT1 and VT2, following the criteria previously described. If they did not agree, the opinion of a third investigator was included.

ANALYSIS OF BLOOD LACTATE

Blood samples (25 µl) for the measurement of blood lactate (YSI 23L; Yellow Springs Instruments, Yellow Springs, Ohio, USA) were taken from fingertips at rest, every two minutes during the test, and immediately after termination of exercise.

LT was determined by examining the "lactate concentration-power output (W)" relationship during the tests, using the methodology previously described by Weltman and co-workers. Thus the greatest power output that was not associated with a rise in lactate concentration above baseline was designated as the power output corresponding to LT. This always occurred just before the curvilinear increase in blood lactate observed with subsequent exercise intensities. A lactate increase of at least 0.2 mM (the error associated with the lactate analyser) was required for LT determination. OBLA, on the other hand, was defined as the power output corresponding to a blood lactate concentration of 4.0 mmol/l.

EMG

Electrode placement

Surface EMG recordings were taken from the vastus lateralis and rectus femoris (at sites respectively approximately one third and one half of the perpendicular distance from the superior border of the patella to the greater trochanter). Pairs of surface electrodes (Blue-sensor Medicotest Ag/AgCl electrodes; Rugmarken, Denmark) were attached to the skin with a 4 cm interelectrode distance. The electrodes were placed longitudinally with respect to the underlying muscle fibre arrangement. For those subjects who performed the test twice (reliability study), the skin was tattooed using ink in order to place the electrodes on the same site on the two tests. Two reference electrodes were placed over the anterior superior spine of the iliac crest. Before electrode application, the skin was shaved and abraded using sandpaper and cleaned with alcohol to minimise the source impedance. A saline EMG electrode gel was placed between the electrode and the underlying skin to enhance signal conductivity. The cables connected to the electrodes to measure myoelectrical activity were firmly attached with tape to minimise artefacts from leg movements.

EMG instrumentation and procedures

Myoelectrical activity was recorded using a ME3000P analyser (ME3000P; Mega Electronics Ltd, Kuopio, Finland). The measurement sensitivity of the instrument is ±1 µV and its range for bipolar EMG signals is ±5000 µV. The raw EMG signals were band-pass filtered between 20 and 480 Hz, amplified, and converted from analogue to digital at a sampling rate of 1 kHz. An EMG power spectral density was then computed for two second sampling periods, at fixed intervals throughout the tests, and the root mean square voltage (rms-EMG) contained in each two second
The degree of correlation between mean values of EMG$_{t1}$ and EMG$_{t2}$ reported with repeated tests.

In all 28 subjects (validity study), mean values of VT$_2$, LT (criterion methods), and EMG$_{t1}$ occurred on one hand, and those of VT$_1$, OBLA (criterion methods), and EMG$_{t2}$ on the other, were compared by using a one way repeated measures analysis of variance. In addition, total errors (Σ(Y−Y$'$)²/$N^{1/2}$ where Y is the criterion value and Y$'$ the predicted value) and Pearson’s correlation coefficients (criterion v predicted) were calculated to examine the error inherent in the EMG prediction technique. Further analysis of validity of the EMG method was accomplished by applying the procedures suggested by Bland and Altman. For this analysis, the mean differences (bias) and standard deviation (SD) of the differences between the mean values (in W) obtained with the two methods (EMG v criterion methods) were calculated. The data are presented graphically comparing the difference between the methods against their average value in W. The mean difference (bias) plus and minus two standard deviations is indicated on the graph. In this way, the bias and precision of the EMG technique could be calculated.

Significance was set at p<0.05 for all statistical analyses.

## Results

### MAXIMAL VALUES

Maximal values of $\dot{V}O_2$ and power output averaged 69.9 (6.4) ml/kg/min (range 60.0–82.3) and 432.8 (49.1) W (range 364–518) at the end of exercise.

### PATTERN OF EMG RESPONSE

A two threshold response was detected in 90% of subjects in both vastus lateralis and rectus femoris, and the two breakpoints EMG$_{t1}$ and EMG$_{t2}$ occurred around 60–70% and 80–90% of $\dot{V}O_2$MAX. In 10% of the cases, EMG$_{t3}$ could not be detected in either of the two muscles studied, whereas EMG$_{t2}$ was found in all 28 subjects.

Figure 1 shows an example of an EMG response in one subject.

### RELIABILITY

No significant differences existed between mean values of either EMG$_{t1}$ or EMG$_{t2}$ obtained in both tests (p<0.05) (Table 1).

Intraclass correlation coefficients (r) between repeated measurements were significant (p<0.05) and high (Table 1).

### VALIDITY

**Comparison between VT$_1$, LT, and EMG$_{t1}$**

Using the methodologies described above, VT$_1$ and LT could be detected in 100% of subjects. Average values of LT occurred at a blood lactate concentration of 1.9 (0.5) mM. Table 2 presents mean values of VT$_1$, LT, and EMG$_{t1}$, expressed in $\dot{V}O_2$ (ml/kg/min), % of $\dot{V}O_2$MAX, and W. No significant differences (p>0.05) were found between means.
Electromyographic analysis in elite cyclists

![EMG recording](image)

Figure 1 Example of electromyographic (EMG) recording in one subject. Each data point represents root mean square of an EMG signal (rms-EMG), recorded at two second intervals. The rms-EMG data against time were fitted mathematically to the corresponding straight lines by computerised multisegment linear regression. EMGT1 and EMGT2 are first and second EMG thresholds.

### Table 1 Reliability of Measurements of the Electromyographic Thresholds EMGT1 and EMGT2

<table>
<thead>
<tr>
<th></th>
<th>First test</th>
<th>Second test</th>
<th>Correlation (r)</th>
</tr>
</thead>
<tbody>
<tr>
<td>EMGT1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V0d (ml/kg/min)</td>
<td>45.5 (2.5)</td>
<td>43.9 (3.0)</td>
<td>0.76</td>
</tr>
<tr>
<td>% V0d MAX</td>
<td>68.8 (4.0)</td>
<td>63.3 (4.2)</td>
<td>0.82</td>
</tr>
<tr>
<td>W</td>
<td>252.7 (17.6)</td>
<td>238.2 (17.9)</td>
<td>0.83</td>
</tr>
<tr>
<td>EMGT2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V0d (ml/kg/min)</td>
<td>59.7 (1.4)</td>
<td>62.6 (1.4)</td>
<td>0.96</td>
</tr>
<tr>
<td>% V0d MAX</td>
<td>90.2 (1.3)</td>
<td>90.2 (2.3)</td>
<td>0.87</td>
</tr>
<tr>
<td>W</td>
<td>367.6 (15.4)</td>
<td>372.6 (14.7)</td>
<td>0.86</td>
</tr>
<tr>
<td>Rectus femoris</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V0d (ml/kg/min)</td>
<td>48.7 (2.5)</td>
<td>47.3 (1.7)</td>
<td>0.89</td>
</tr>
<tr>
<td>% V0d MAX</td>
<td>70.4 (3.0)</td>
<td>68.4 (2.9)</td>
<td>0.85</td>
</tr>
<tr>
<td>W</td>
<td>280.2 (22.4)</td>
<td>268.2 (20.8)</td>
<td>0.96</td>
</tr>
<tr>
<td>EMGT2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V0d (ml/kg/min)</td>
<td>59.6 (2.1)</td>
<td>59.8 (1.5)</td>
<td>0.73</td>
</tr>
<tr>
<td>% V0d MAX</td>
<td>89.9 (2.0)</td>
<td>86.2 (1.8)</td>
<td>0.75</td>
</tr>
<tr>
<td>W</td>
<td>374.2 (20.7)</td>
<td>360.8 (20.0)</td>
<td>0.87</td>
</tr>
</tbody>
</table>

All values are expressed as mean (SEM). No significant differences existed between means (p>0.05). All correlation coefficients were significant (p<0.05).

### Table 2 Comparison between Mean values of the First Electromyographic Threshold (EMG1), the First Ventilatory Threshold (VT1), and the Lactate Threshold (LT)

<table>
<thead>
<tr>
<th></th>
<th>EMG1</th>
<th>VT1</th>
<th>LT</th>
</tr>
</thead>
<tbody>
<tr>
<td>V0d (ml/kg/min)</td>
<td>43.9</td>
<td>47.0</td>
<td>45.1</td>
</tr>
<tr>
<td>% V0d MAX</td>
<td>62.8</td>
<td>69.0</td>
<td>64.6</td>
</tr>
<tr>
<td>W</td>
<td>240.3</td>
<td>270.8</td>
<td>257.8</td>
</tr>
</tbody>
</table>

All values are expressed as mean (SEM). No significant differences existed between means (p>0.05).

### Table 3 Comparison between Mean values of the Second Electromyographic Threshold (EMG2), the Second Ventilatory Threshold (VT2), and the Onset of Blood Lactate Accumulation (OBLA)

<table>
<thead>
<tr>
<th></th>
<th>EMG2</th>
<th>VT2</th>
<th>OBLA</th>
</tr>
</thead>
<tbody>
<tr>
<td>V0d (ml/kg/min)</td>
<td>61.5</td>
<td>59.7</td>
<td>59.0</td>
</tr>
<tr>
<td>% V0d MAX</td>
<td>86.9</td>
<td>88.0</td>
<td>84.6</td>
</tr>
<tr>
<td>W</td>
<td>371.1</td>
<td>367.5</td>
<td>352.8</td>
</tr>
</tbody>
</table>

All values are expressed as mean (SEM). No significant differences existed between means (p>0.05).

Comparison between VT2, OBLA, and EMG1. Using the methodologies described above, VT2, and OBLA could be detected in 100% of subjects. Table 3 presents mean values of VT2 and EMG2. In addition, for all comparisons, at least 90% of the individual values were within the limits of agreement.

### Table 4 Total Errors

<table>
<thead>
<tr>
<th></th>
<th>Vastus lateralis</th>
<th>Rectus femoris</th>
</tr>
</thead>
<tbody>
<tr>
<td>VT1</td>
<td>0.66</td>
<td>—</td>
</tr>
<tr>
<td>VT2</td>
<td>0.64</td>
<td>0.82</td>
</tr>
<tr>
<td>LT</td>
<td>0.64</td>
<td>0.80</td>
</tr>
<tr>
<td>OBLA</td>
<td>—</td>
<td>0.80</td>
</tr>
</tbody>
</table>

EMGT1 and EMGT2, first and second electromyographic thresholds; VT1 and VT2, ventilatory thresholds 1 and 2; LT, lactate threshold; OBLA, onset of blood lactate accumulation. No significant differences (p>0.05) existed between means.

### Table 5 Significant Correlation Coefficients (p<0.05)

<table>
<thead>
<tr>
<th></th>
<th>EMG1</th>
<th>EMG2</th>
</tr>
</thead>
<tbody>
<tr>
<td>VT1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>VT2</td>
<td>0.93</td>
<td></td>
</tr>
<tr>
<td>LT</td>
<td>0.96</td>
<td>0.83</td>
</tr>
<tr>
<td>OBLA</td>
<td>0.80</td>
<td>0.97</td>
</tr>
</tbody>
</table>

EMGT1 and EMGT2, first and second electromyographic thresholds; VT1 and VT2, ventilatory thresholds 1 and 2; LT, lactate threshold; OBLA, onset of blood lactate accumulation.

### Discussion

To our knowledge, this is the first report to determine the validity of the EMG method for analysing the aerobic-anaerobic transition phase in top level athletes (professional cyclists) during cycle ergometry using a ramp test. Previous studies on EMG response during such protocols have been conducted with subjects of considerably lower fitness levels (in most studies, mean VO2 MAX <65 ml/kg/min, and in all of them mean peak power output <360 W) than those selected for our investigation (VO2 MAX 69.9 (6.4) ml/kg/min; peak power output 432.8 (49.1) W). To date, no conclusion could be inferred from previous research.
The EMG method is both valid and reliable for analysing the aerobic-anaerobic transition. A second finding of our investigation was that the EMG activity of the muscles primarily involved in pedalling (vastus lateralis, rectus femoris) show a two threshold response with two distinct breakpoints, EMGT1 and EMGT2, occurring at an exercise intensity of 60–70% and 80–90% of VO\textsubscript{MAX} respectively. A novel finding of our study was that in elite cyclists the EMG activity of muscles that are primarily fatigued with increasing exercise intensity does not significantly affect muscle fatigue, as EMGT1 and EMGT2 occurred at similar intensities in the vastus lateralis and rectus femoris.

**METHODOLOGICAL LIMITATIONS**

Our study is not without potential limitations. Firstly, the literature on the usefulness of EMG in the determination of ventilatory or lactate thresholds is somewhat controversial. Differences in results could be attributed to difficulties in reliably following the myoelectrical activities in working muscles. In addition, during exertion the skin blood flow and temperature increase. This phenomenon could induce alterations in the electrical properties of the skin and affect the records of surface EMG. In this regard, previous research has shown no correlation between skin temperature and EMG measurements, excluding the possibility that EMG breakpoints are related to changes in electrical properties of the skin.

On the other hand, although we measured the myoelectrical activity of the muscles (vastus lateralis and rectus femoris) involved during the work (descending) period of pedalling only, no records were obtained for other muscles—for example, gastrocnemius, tibialis anterior—that are also involved during the rest (ascending) period of pedalling exercise. However, previous research has shown that a non-linear increase in EMG also occurs in the gastrocnemius muscle. In addition, previous EMG and biopsy studies suggest that the muscle most heavily involved during cycling is the vastus lateralis.

Furthermore, a single ramp increment (5 W every 12 seconds) was chosen in our protocol, whereas the results of a previous study using different ramp slopes (10, 20, 30 and 40 W/min) suggest that the metabolic state at which the EMG occurs in untrained subjects may differ during the different ramp exercises. Further research should be conducted with elite cyclists to determine the influence of varying ramp protocols on the EMG response. Finally, the results obtained during a progressive test in a cycle ergometer using a fixed cadence (70–80 rpm), as was used in this investigation, cannot be easily extrapolated to
Electromyographic analysis in elite cyclists

The EMG response to increasing exercise intensity is one of the key phenomena in exercise physiology. As exercise intensity increases, the force production of muscle fibres increases, leading to a greater recruitment of motor units. This process is marked by an increase in the amplitude of the myoelectric signal, or EMG, which can be recorded using surface electrodes. The breakpoint in this increase in EMG, often referred to as the EMG Transition (EMGT), is associated with the anaerobic threshold (AT) and is determined by the point at which the increase in EMG becomes linear.

The EMGT is a critical point in exercise physiology, as it marks the transition from aerobic to anaerobic metabolism. During exercise, muscle fibres are recruited in a specific order, with type I fibres being recruited first, followed by type IIa and then type IIb fibres. This recruitment pattern is determined by the exercise intensity and is associated with changes in muscle pH, with a decrease in pH indicating muscle fatigue.

In elite cyclists, the EMGT can be observed during exercise on a cycle ergometer. The EMGT is typically associated with a decrease in muscle pH, indicating the onset of anaerobic metabolism. This decrease in pH can be measured using a pH electrode inserted into the muscle, or estimated from changes in the blood lactate concentration.

The EMGT is also associated with changes in ventilatory and lactate thresholds. The ventilatory threshold (VT1) is the point at which ventilation begins to increase disproportionately with exercise intensity, while the lactate threshold (LT) is the point at which blood lactate concentration begins to increase. Both of these thresholds are associated with the EMGT, with the EMGT occurring at a percentage of VO2MAX that is lower than the VT1 and LT.

The EMGT is also associated with changes in muscle fibre recruitment. During exercise, muscle fibres are recruited in a specific order, with type I fibres being recruited first, followed by type IIa and then type IIb fibres. This recruitment pattern is determined by the exercise intensity and is associated with changes in muscle pH, with a decrease in pH indicating muscle fatigue.

In summary, the EMGT is a critical point in exercise physiology, marking the transition from aerobic to anaerobic metabolism. It is associated with changes in muscle pH, ventilatory and lactate thresholds, and muscle fibre recruitment. Understanding the EMGT is important for athletes and coaches, as it provides insights into the physiological mechanisms underlying exercise performance.
increase in lactate levels but also by other factors such as an increase in neural or hormonal activity.\(^4\)

**COINCIDENCE OF EMG AND VENTILATORY RESPONSES**

The correlation encountered between VT and EMG\(_T_2\), on the one hand, and VT\(_1\) and EMG\(_T_2\), on the other, could be due to a muscle derived signal to ventilation. Indeed, previous studies provide evidence for the existence of ergoreceptors or mechanoreceptors which respond to increases in the work performed per unit muscle.\(^20\) Moreover, Morikawa and co-workers\(^31\) found that a ventilatory response occurred in normal subjects with both active and passive leg exercise, but was absent with passive exercise in patients with thoracic spine transection. It could then be hypothesised that ventilation shows a first deflection point (VT\(_1\)) when muscle work is increased (at an exercise intensity corresponding to EMG\(_T_1\)) and a further increase (VT\(_2\)) when additional motor units are recruited to maintain power output—that is, recruitment of type IIb fibres at the EMG\(_T_2\).

On the other hand, the ventilatory thresholds observed during an incremental test could be elicited by enhancement of the neural activity that originates directly from the subthalamic motor region or indirectly via \(\alpha-\gamma\) coactivation of motoneurons innervating the muscle fibres of exercising limbs.\(^32\) This increase in neural activity may occur during incremental exercise in response to the need to progressively recruit additional motor units comprised of type IIa (possibly at the EMG\(_T_1\)) and IIb fibres (possibly at the EMG\(_T_2\)) respectively as the work rate is increased and individual fibres begin to fatigue.\(^35\) Indeed, numerous investigations have shown simultaneous increases in ventilation and EMG activity during incremental exercise.\(^3\)\(^4\)\(^5\)\(^32\)

The results of a recent study, however, do not support a link between motor unit recruitment and ventilation, as evidenced by the disassociation between the EMG, of the rectus femoris and the ventilatory threshold that existed in trained cyclists after glycogen depletion.\(^3\) A similar study should be conducted with professional cyclists in order to assess the effects of glycogen depletion on the rms-EMG response.

**ABSENCE OF EMG\(_T_2\) IN PREVIOUS RESEARCH**

The question remains to be answered of why a second EMG threshold (EMG\(_T_2\)) was found only in this study, in contrast with previous research. In this regard, a possible explanation could be that the significant increases in neural activity to the exercising muscles that occur at EMG\(_T_2\), were not detected in previous studies because EMG activity was mainly recorded from a single muscle—that is, the vastus lateralis. This hypothesis is supported by the findings of Green and Patla,\(^19\) which showed that many different muscle groups contribute to the completion of an incremental exercise task and that the average level of activation increases differentially among the muscle groups. It may be possible that only highly trained cyclists (such as those selected for our study) are able to effectively recruit a sufficient number of motor units (especially those with fast fibres) within individual muscles (vastus lateralis or rectus femoris) at near to maximum intensities during an incremental test such as to induce a second breakpoint in the EMG response to exercise. Indeed, EMG\(_T_2\) occurred at a power output as high as 370 W, which in turn elicited a \(V_{O_2}\) of about 60 ml/kg/min. Both values are well above the maximal levels of power output and \(V_{O_2}\) reported in most previous EMG studies (<350 W and <55 ml/kg/min respectively). Although two studies\(^7\)\(^8\) have been conducted in this area using highly trained cyclists with similar values of \(V_{O_2\text{MAX}}\) to those of our subjects (about 70 ml/kg/min), they attained considerably lower peak power outputs (<360 W) during incremental tests.

**SIMILAR RESPONSES IN VASTUS LATERALIS AND RECTUS FEMORIS**

Based on the results of previous research with non-elite athletes,\(^3\)\(^4\)\(^5\)\(^3\)\(^3\) a disparity in the EMG/exercise intensity relation would be expected to exist between the vastus lateralis and the rectus femoris. In effect, previous research has shown distinct responses in the two muscles, such as the occurrence of the EMG fatigue threshold at lower intensities in the rectus femoris.\(^34\) This disparity could be partly attributed to kinesiological differences, as the rectus femoris (unlike the vastii muscles) is a biarticular muscle involved in both leg extension and thigh flexion.\(^34\) On the other hand, possible differences in muscle fibre composition cannot be excluded, the rectus femoris being comprised of a greater percentage of fatigable fast twitch fibres.\(^3\)\(^4\)\(^3\) In this regard, few data are available on the muscle fibre composition of leg muscles in professional cyclists. In our investigation, however, EMG\(_T_1\) and EMG\(_T_2\) occurred at similar intensities in both muscles, suggesting a similar pattern of muscle fibre recruitment in the different leg muscles of top level cyclists as an adaptation to training and competition.

**CONCLUSION**

In elite cyclists the EMG activities of two leg muscles (vastus lateralis and rectus femoris) show similar patterns, with two distinct breakpoints, EMG\(_T_1\) and EMG\(_T_2\), occurring at an exercise intensity of 60–70% and 80–90% of \(V_{O_2\text{MAX}}\) respectively. The rms-EMG method seems to be both valid and reliable for analysing the aerobic-anerobic transition during cycle ergometer exercise.
Electromyographic analysis in elite cyclists


Take home message
During incremental cycle ergometer tests performed by top level cyclists, the relation between EMG activity (rms-EMG) of exercising muscles (vastus lateralis and rectus femoris) and exercise intensity exhibits two breakpoints, at around 60–70% and 80–90% of VO2MAX respectively. rms-EMG may be used as a complementary indicator of the aerobic-anaerobic transition in physiological evaluations of cyclists.
Analysis of the aerobic-anaerobic transition in elite cyclists during incremental exercise with the use of electromyography.

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