The Relationship Between Lactate and Ventilatory Thresholds in Runners: Validity and Reliability of Exercise Test Performance Parameters

Víctor Cerezuela-Espejo, Javier Courel-Ibáñez, Ricardo Morán-Navarro, Alejandro Martínez-Cava and Jesús G. Pallarés*

Human Performance and Sports Science Laboratory, Faculty of Sport Sciences, University of Murcia, Murcia, Spain

The aims of this study were (1) to establish the best fit between ventilatory and lactate exercise performance parameters in running and (2) to explore novel alternatives to estimate the maximal aerobic speed (MAS) in well-trained runners. Twenty-two trained male athletes ($\dot{V}O_{2max} 60.2 \pm 4.3$ ml·kg·min$^{-1}$) completed three maximal graded exercise tests (GXT): (1) a preliminary GXT to determine individuals’ MAS; (2) two experimental GXT individually adjusted by MAS to record the speed associated to the main aerobic–anaerobic transition events measured by indirect calorimetry and capillary blood lactate (CBL). Athletes also performed several 30 min constant running tests to determine the maximal lactate steady state (MLSS). Reliability analysis revealed low CV (<3.1%), low bias (<0.5 km·h$^{-1}$), and high correlation (ICC > 0.91) for all determinations except V-Slope (ICC = 0.84). Validity analysis showed that LT, LT+1.0, and LT+3.0 mMol·L$^{-1}$ were solid predictors of VT$_1$ (<0.3 km·h$^{-1}$; bias = 1.2; ICC = 0.90; $p = 0.57$), MLSS (<0.2 km·h$^{-1}$; bias = 1.2; ICC = 0.84; $p = 0.74$), and VT$_2$ (<0.1 km·h$^{-1}$; bias = 1.3; ICC = 0.82; $p = 0.919$), respectively. MLSS was identified as a different physiological event and a midpoint between VT$_1$ (bias = −2.0 km·h$^{-1}$) and VT$_2$ (bias = 2.3 km·h$^{-1}$). MAS was accurately estimated (SEM ± 0.3 km·h$^{-1}$) from peak velocity ($V_{peak}$) attained during GXT with the equation: $MAS_{EST}$ (km·h$^{-1}$) = $V_{peak}$ (km·h$^{-1}$) * 0.8348 + 2.308. Current individualized GXT protocol based on individuals’ MAS was solid to determine both maximal and submaximal physiological parameters. Lactate threshold tests can be a valid and reliable alternative to VT and MLSS to identify the workloads at the transition from aerobic to anaerobic metabolism in well-trained runners. In contrast with traditional assumption, the MLSS constituted a midpoint physiological event between VT$_1$ and VT$_2$ in runners. The $V_{peak}$ stands out as a powerful predictor of MAS.

Keywords: blood lactate, ventilation threshold, maximal aerobic speed, $V_{O2max}$, endurance, maximal lactate steady state
INTRODUCTION

Numerous studies have embraced the question of how training programs based on individualized physiological parameters may increase cardiorespiratory performance in endurance sports like running or cycling. Evidence suggests establishing exercise workloads or intensities based on the individual physiological events (i.e., setting training zones) allows athletes to minimize injury and fatigue risks, but above all to enhance individual adaptations and respond to the training plan (Scharhag-Rosenberger et al., 2012; Mann et al., 2014; Wolpern et al., 2015). A recent review (Stöggl and Sperlich, 2015) addressed the fact that similar training intensity distribution shows different efficacy and adaptations depending on the competitive stage, the endurance discipline, and the athlete’s performance levels. Thus, the more individualized and accurate the thresholds and training zones, the more precise the exercise prescription and the greater the athletes’ adaptation and performance enhancement (Garcia-Pallarés et al., 2009; Wolpern et al., 2015). From a competition point of view, exercise test performance parameters are useful to track cardiopulmonary and specific adaptations to the entire season training plan, and to explain performance (Lucia et al., 2000; Esteve-Lanao et al., 2007; Garcia-Pallarés et al., 2009, 2010).

Physiological variables such as maximal oxygen uptake (VO$_{2max}$), submaximal metabolic inflection points like the pulmonary ventilation thresholds (VT) and lactate thresholds (LT), the maximal aerobic speed (MAS: the speed associated with VO$_{2max}$), or the peak velocity (V$_{peak}$: the highest speed attained at the end of the test) are regular variables used by coaches and scientists to estimate and monitor running performance during training and competition events (Farrell et al., 1979; di Prampero et al., 1986; Stratton et al., 2009; McLaughlin et al., 2010). For the evaluation of these parameters in runners, it is common to use graded exercise tests (GXTs) on the treadmill, consisting of a series of stages lasting 1–5 min. Differences in the duration of each stage and the load increments can alter the cardiorespiratory and metabolic response, and therefore the measurement (Bentley et al., 2007; Julio et al., 2017). As suggested by pioneering studies (Buchfuhrer et al., 1983; Lukaski et al., 1989), recent investigations (Midgley et al., 2007) and reviews (Julio et al., 2017), traditional longer GXTs (i.e., 20–30 min) to determine LT including increments each 3–5 min would prevent the athlete from achieving their MAS due to accumulative fatigue, dehydration, muscle acidosis, and cardiovascular drift. This is critical because MAS is a pertinent and widespread criterion to set training intensities for endurance disciplines (Billat and Koralsztein, 1996; Jones and Carter, 2000). An interesting approach carried out with cyclists revealed that shorter protocols (12–14 min) including 1-min stages are valid both, to estimate submaximal metabolic inflection points (VT and LT), and to identify true values for VO$_{2max}$ and MAS in cyclists (Lucia et al., 1999, 2000; Gaskill et al., 2001; Midgley et al., 2007; Pallarés et al., 2016). However, the validity and reliability of GXT with 1-min stages protocol in runners needs to be fully verified.

Physiological response to exercise in endurance sports is commonly assessed though measurements based on ventilatory and lactate methods. However, the relationship between the two methods is not yet clear (Pallarés et al., 2016). Recent findings support the idea that a training model based on ventilatory thresholds (VT$_1$ and VT$_2$) could be very effective to set individual exercise intensity in endurance sports given that it takes into account individual metabolic responses (Wolpern et al., 2015). One of the most accurate systems to obtain these ventilatory responses is on the basis of gas exchange parameters using indirect calorimetry (Lucia et al., 2000; Gaskill et al., 2001; Pallarés et al., 2016). In VT$_1$, the VO$_2$ and carbon dioxide production (VCO$_2$) increase proportionally, while HCO$_3$– acts to buffer lactic acid concentration in blood (Wasserman et al., 1973; Del Coso et al., 2009); this intensity is ideal for high-volume low-intensity exercise (Stöggl and Sperlich, 2014). In turn, in VT$_2$, the blood lactate accumulation boosts and rises considerably and the system collapses due to the homeostatic compromise and metabolic acidosis (Wasserman et al., 1973; Jones et al., 2007); this intensity sets a critical limit for high-intensity interval training (Stöggl and Sperlich, 2014). However, gas exchange systems require the use of expensive equipment and laboratory conditions which most teams, coaches, and athletes are not equipped with or cannot afford.

A further method to set individual exercise intensity is based on capillary blood lactate (CBL) measurements (Beneke et al., 2011). A number of authors have defined a list of CBL parameters associated with specific exercise intensities such as LT (Wasserman et al., 1973), maximal lactate steady state (MLSS, Beneke and van Duvillard, 1996), OBLA (onset of blood lactate accumulation, Sjödin and Jacobs, 1981), or the D$_{MAX}$ (Cheng et al., 1992). An accurate detection of MLSS is particularly important due to it being considered the highest intensity in which glycogen stores are the main exercise limiting factor (Coyle et al., 1986) and constitutes a prominent part of aerobic training in world-class athletes (Garcia-Pallarés et al., 2009, 2010). Although CBL methods are commonly used for coaches to set individual training workloads, the relationship between lactate-based parameters and VTs load intensities is still an open debate. In cyclists, it seems clear that workloads at the VT$_1$ are very related to LT (Lucia et al., 1999; Amann et al., 2006; Pallarés et al., 2016). However, the estimation of VT$_2$ from lactate methods generates some controversy. Traditionally, VT$_2$ intensities have been associated with MLSS (Svedahl and MacIntosh, 2003). In contrast with this assumption, recent evidence in cyclists demonstrates that MLSS encompasses a different metabolic pathway and limiting factor than VT, and constitutes a midpoint between VT$_1$ and VT$_2$ (Pallarés et al., 2016; Peinado et al., 2016). The determination of VT$_2$ is essential due to represents a turn point at which metabolic acidosis cannot be buffered by ventilation (Lucia et al., 2000) and sets a critical limit for high-intensity training (Stöggl and Sperlich, 2014). One previous study conducted with cyclists has attempted to clarify the relationship between VTs and CBL methods, and reported a high reliability and validity of the following relationships: (1) VT$_1$ and LT, (2) VT$_2$ and LT+2 mMol·L$^{-1}$, and (3) MLSS and LT+0.5 mMol·L$^{-1}$ (Pallarés et al., 2016). To the best of our knowledge, there are no previous studies examining these relationships in runners. This is an important gap considering the existing differences between cycling and running, such as the
more impaired ventilation in cycling and the higher muscle mass involved, greater muscle pump efficiency, and the implication of eccentric muscle actions in running (Bijker et al., 2002; Millet et al., 2009). Given that these differences may alter the physiological response to exercise, prescribing training plans for runners based on cyclists’ reference values could be imprecise. Thus, the relationships between CBL and VTs intensities in runners need to be fully clarified.

In addition to the aerobic–anaerobic transition, another ventilation parameter to predict running performance is the MAS (McLaughlin et al., 2010), considered as the minimum speed at which V\textsubscript{O}\text{2 max} is reached (Lacour et al., 1991). As a rule of thumb, high intensity training in endurance athletes is established at 90–105% of the MAS (Stöggel and Sperlich, 2015). Given its importance for training plans and workload distribution, coaches and researchers have invested effort in designing maximal field tests to estimate the MAS and predict V\textsubscript{O}\text{2 max} in endurance athletes to establish the aerobic performance limits (Léger and Boucher, 1980; Berthon et al., 1997). However, these tests have important limitations: (1) the equations proposed to estimate the MAS from field tests are not based on accurate measurements such as gas exchange systems using indirect calorimetry (Lucia et al., 2000; Gaskill et al., 2001), and (2) maximal efforts criteria were not tested to ensure reaching values of V\textsubscript{O}\text{2 max} (ACSM, 2013). As stated above, a valid alternative to these field tests is to determine the MAS through GXT with 1-min increments using gas exchange systems. These short protocols allow the athletes to reach their maximal cardiac output, and therefore make possible obtaining a true V\textsubscript{peak} value (Pallares et al., 2016; Julio et al., 2017). Given that both MAS and V\textsubscript{peak} correspond to very similar intensities (LaCOUR et al., 1991), the calculation of an estimated MAS (MAS\textsubscript{EST}) from the V\textsubscript{peak} when gas exchange systems are not available, seems promising. However, this hypothesis is still to be proven.

Therefore, the aims of this study were (1) to establish the best fit between ventilatory and lactate exercise performance parameters in running (and (2) to explore novel alternatives to estimate essential running performance indicators such as the MAS from similar intensity parameters like the V\textsubscript{peak} when gas exchange systems are not available.

**METHODS**

**Participants**

Twenty-two trained male athletes (runners and triathletes) volunteered to participate in this study (age 25.9 ± 8.0 years, body mass 68.2 ± 6.1 kg, height 174.8 ± 5.8 cm, body fat 11.4 ± 1.9%, VO\textsubscript{2 max} 60.2 ± 4.3 ml·kg·min\(^{-1}\), endurance training experience 7.1 ± 4.0 years). All participants were competing at regional and national level races and following a regular training load of 4–6 days per week, 1–2 h per day. Measurements were obtained during the pre-competitive season. All participants underwent a complete medical examination (including ECG) that showed all were in good health. No physical limitations or musculoskeletal injuries that could affect testing procedures were reported. None of the subjects were taking drugs, medications, or dietary supplements known to influence physical performance.

The Bioethics Commission of the University of Murcia approved the study, which was carried out according to the declaration of Helsinki. Subjects were verbally informed about the experimental procedures and possible risk and benefits. Written informed consent was obtained from all subjects.

**Experimental Design**

Participants visited the lab 5–7 times separated by 2–7 days. All participants had at least 6 months of familiarization with the testing procedures used in this investigation. On the first day, participants completed a preliminary GXT with 1-min increments (GXT\textsubscript{PRE}) to determine individuals’ MAS and V\textsubscript{peak}, including 48–72 h rest before the next session. In the following two sessions, separated by 48 h, athletes performed two identical experimental GXT (GXT\textsubscript{EXP} 1 and GXT\textsubscript{EXP} 2). For these two GXT\textsubscript{EXP} protocols, initial running speed and workload increments were individually set according to participants’ V\textsubscript{peak} previously determined in the GXT\textsubscript{PRE}.

The GXT\textsubscript{EXP} started with a 5-min warm-up at 13 km·h\(^{-1}\) less than each athlete’s V\textsubscript{peak} followed, without a break, by a GXT 1-min (i.e., increments of 1 km·h\(^{-1}\)·min\(^{-1}\)) until exhaustion. Lastly, athletes came back to the lab two to three more times to perform a 30 min submaximal constant running test to determine the speed associated with the MLSS (Beneke, 2003). To maintain physical performance during the investigation period (2–3 weeks) participants followed an individual training protocol consisting in: running sessions (runners) or swimming, cycling, and running sessions (triathletes) of 90 min at individual VT1 intensity interspersed with efforts of 5–7 min at 90–95% of VT2 intensity each 20 min. Training sessions were repeated each 48 h with 24 h rest before each evaluation to ensure a full recovery.

**Individualized Maximal Treadmill GXT Protocol**

All the running trials were performed on the same treadmill (HP Cosmos Pulsar, HP Cosmos Sports and Medical GMBH, Nußdorf Traunstein, Germany) with an incline of 1.0% (Jones and Doust, 1996). Evaluations were performed under similar environmental conditions (21–24°C and 45–55% relative humidity) at the same time of day (16:00 to 19:00 h) to minimize the circadian rhythm effects (Mora-Rodriguez et al., 2015). Air ventilation was controlled with a fan positioned 1.5 m from the subject’s chest at a wind velocity of 2.55 m·s\(^{-1}\).

The GXT\textsubscript{PRE} under medical supervision to fulfill three objectives: (1) discard cardiovascular diseases, (2) to minimize the bias of progressive learning on test reliability, and (3) to determine the athletes’ MAS and V\textsubscript{peak} subsequently used to set up the individualized GXT\textsubscript{EXP} workload (i.e., treadmill speed). Participants’ HR was monitored by standard 12 lead ECG (Quark T12, Cosmed, Italy), ventilatory performance (V\textsubscript{O}2, V\textsubscript{O}\text{2 max}, and VE) was recorded on a breath-by-breath basis using a metabolic cart averaging data every 5 s (MetaLyzer 3B-R3, Cortex Biophysik GmbH, Leipzig, Germany) and the rate of perceived exertion (RPE) was assessed using the 6–20 Borg Scale (Borg, 1998) every 2 min. The MAS was determined from metabolic cart measurements as the first running velocity where V\textsubscript{O}2 max was reached (Billat and Koralsztein, 1996). The V\textsubscript{peak
was automatically obtained from the treadmill software using the Kuipers et al.’s formula (Kuipers et al., 2003): \( V_{\text{peak}} = V_{\text{complete}} + \left( \text{Inc} \; \times \; t/T \right), \) in which \( V_{\text{complete}} \) is the running velocity of the last complete stage, \( \text{Inc} \) is the speed increment (i.e., 1 km·h\(^{-1}\)), \( t \) is the number of seconds sustained during the incomplete stage and \( T \) is the number of seconds required to complete a stage (i.e., 60 s).

The two GTX\(_{\text{EXP}}\) were individually set up according to the MAS previously determined in the preliminary test (GXT\(_{\text{PRE}}\)), as follows: starting with a 5-min warm-up at 13 km·h\(^{-1}\) less than each athlete’s \( V_{\text{peak}} \), followed, without a break, by a GTX 1-min (i.e., increments of 1 km·h\(^{-1}\)·min\(^{-1}\)) until exhaustion. Ventilatory parameters and RPE were assessed as aforementioned in the GXT\(_{\text{PRE}}\). The HR was continuously monitored (V800, Polar, Finland). Capillary blood lactate samples from the finger were collected (Lactate Pro, Arkray, Japan) every 2 min (i.e., each 2 km·h\(^{-1}\) increments). The design of this particular protocol and its duration (min–max) were deliberate, given that: (1) It allows a clear detection of ventilatory thresholds (VT\(_1\) and VT\(_2\)) by indirect calorimetry (Lucía et al., 1999, 2000; Pallarés et al., 2016); (2) It is effective in determining a true \( V_{\text{O2max}} \) (Midgley et al., 2007); (3) The protocol duration was short enough (12–14 min) to avoid the local acidosis and HR rise (cardiac drift) to obtain a true maximum cardiovascular performance (Dawson et al., 2005); (4) The short duration allows athletes to achieve a true MAS and \( V_{\text{peak}} \) (Julio et al., 2017); and (5) By the end of the test, seven to nine capillary blood samples can be collected from each participant before exhaustion, which enable the plotting of a complete lactate curve. In particular, fingerprint blood samples were collected by a specialist placed beside the treadmill without any pause during the participants’ running test (i.e., in movement) to make the process less invasive and ensure a constant effort during the GXT protocols.

Maximal effort criteria (ACSM, 2013) were considered to verify the outcomes, from which participants must reach at least three from the list: (i) failure of HR to increase with further increases in exercise intensity; (ii) a plateau in \( V_{\text{O2}} \) (or failure to increase \( V_{\text{O2}} \) by 150 mL·min\(^{-1}\)) with increased workload; (iii) a respiratory exchange ratio (RER) \( \geq 1.10; \) CBL \( > 8 \text{ mmol·L}^{-1}; \) (iv) a rating of perceived exertion (RPE) \( > 17 \) on the 6–20 scale. If verified, physiological parameters were determined, and the individuals’ treadmill speed at each of the physiological parameters studied were considered for subsequent analysis. Blood lactate analyzer and indirect calorimetry devices were calibrated before each test according to the manufacturer’s instructions.

**Determination of MLSS**

Several 30 min constant workloads on a treadmill were performed to identify the highest workload (km·h\(^{-1}\)) at which CBL increased \( > 1 \text{ mmol·L}^{-1} \) between the 10th and 30th min of exercise (Beneke, 2003). After 7 days from the second GTX\(_{\text{EXP}}\), all participants performed the first MLSS trial at the individual workload associated to their 85% of VT\(_2\), based on previous studies (Llodio et al., 2016; Pallarés et al., 2016). Depending on the results of the first MLSS-test, successive trials with a 48-h rest between sessions were increased or decreased 0.5 km·h\(^{-1}\) until MLSS criteria was fulfilled (Pallarés et al., 2016).

**Determination of Ventilation Parameters**

VT\(_1\) was determined using the criteria of an increase in both ventilatory equivalent of oxygen (\( V_{\text{E/VO2}} \)) and end-tidal pressure of oxygen (\( P_{\text{ETO2}} \)) with no concomitant increase in ventilatory equivalent of carbon dioxide (\( V_{\text{E/VO2}} \)). VT\(_2\) was determined using the criteria of an increase in both the \( V_{\text{E/VO2}} \) and \( V_{\text{E/VCO2}} \) and a decrease in \( P_{\text{ETCO2}} \) (Lucía et al., 2000; Figures 1A,B). V-Slope load was identified in that intensity of exercise which, in a plot of the minute production of CO\(_2\) over the minute utilization of oxygen (\( \dot{V}O_2 \)), shows an increase in the slope above 1.0 (Wasserman et al., 1973; Gaskill et al., 2001). The \( V_{\text{O2max}} \) was defined as the highest plateau (two successive maximal within 150 mL·min\(^{-1}\)), averaging the data every 5 s) reached. MAS was defined as the minimum speed at which maximum oxygen uptake \( V_{\text{O2max}} \) is reached (Lacour et al., 1991). \( V_{\text{peak}} \) was taken from the highest velocity reached during this GXT protocol and calculated according to the Kuipers et al. (2003).

**Determination of Lactate Parameters**

LT was determined by examining the CBL speed relationship (\( [\text{Lact}]_{\text{blood}} / \text{km·h}^{-1} \)) during the GXT as the highest speed not associated with a rise in CBL above baseline (Weltman et al., 1990). Baseline CBL was the average during the initial stages with values 0.8 mmol·L\(^{-1} \) above rest state. This always occurred just before the curvilinear increase in blood lactate observed at subsequent exercise intensities (Coyle et al., 1983; Lucía et al., 2000). Lactate Threshold + 1.0 mmol·L\(^{-1} \) (LT + 1.0) represents the speed which causes an increase of 1 mmol·L\(^{-1} \) above baseline measurements (Coyle et al., 1983). Following this criterion, five LT-based events were established as previously described (Pallarés et al., 2016): LT + 0.5, LT + 1, LT + 1.5, LT + 2.0, LT + 2.5, and LT + 3.0 mmol·L\(^{-1} \). \( D_{\text{MAX}} \) method was determined by plotting the lactate response to exercise intensity in a third-order polynomial regression curve. \( D_{\text{MAX}} \) was defined as the point on the regression curve that yields the maximal distance to the straight line formed by the two end points of the curve (Cheng et al., 1992). Onset of blood lactate accumulation (\( OBLA_{\text{4mm}} \)) was defined as the exercise intensity identified by interpolation across the 4 mmol·L\(^{-1} \) point in the plot of \( [\text{Lact}]_{\text{blood}} \) during incremental exercise (Sjödin and Jacobs, 1981). Two independent observers detected all ventilatory and LT following the criteria previously described. If they did not agree, the opinion of a third investigator was sought (Lucía et al., 1999; Figure 1C).

**Statistical Analyses**

Standard statistical methods were used for the calculation of means, standard deviations (SD), and 95% confidence interval. The reliability of ventilation and lactate parameters was analyzed comparing the consistence among trials (i.e., GXT\(_{\text{EXP}}\) 1 vs. GXT\(_{\text{EXP}}\) 2) by calculating the coefficient of variation (CV), intraclass correlation coefficient (ICC), and Bland–Altman plots. Linear regression analysis was employed to estimate a theoretical MAS (\( \text{MAS}_{\text{EST}} \)) from the average of the \( V_{\text{peak}} \) achieved at the end of the two GXT\(_{\text{EXP}}\) trials. Validity analysis of the ventilatory thresholds (VT\(_1\)
RESULTS

All participants reached at least two of the criteria for achievement of maximal efforts during all the GXT-tests, therefore maximal ventilation and cardiovascular performance was verified. The initial speed ranged from 6 to 10 km·h⁻¹ and no fatigue was detected following the warm-up (i.e., all participants maintained a RER < 0.85 and CBL under the baseline). The \( V_{\text{peak}} \) reached during the GXT \( \text{PRE} \) ranged from 18 to 22 km·h⁻¹. Linear regression analysis (Figure 2) revealed a very strong association between \( MA_{\text{S}} \) and \( V_{\text{peak}} \) \((p < 0.01; r = 0.954; \text{SEM} = 0.3 \text{ km·h}^{-1})\) and yielded the equation:

\[
MA_{\text{S}} = V_{\text{peak}} \times 0.8348 + 2.308
\]

Intra-subject reliability between GXT \( \text{EXP} \) trials (Table 1) revealed low CV (<3.1%), low bias (<0.5 km·h⁻¹), and high correlation (ICC > 0.91) for all determinations except V-Slope (ICC = 0.84).

Table 2 shows the validity analysis comparing VT₁, MLSS, VT₂, and MAS workloads against the rest of the parameters. The strongest associations (ICC > 0.82, \( p > 0.57 \)) were: VT₁ with LT (−0.3 ± 1.2 km·h⁻¹), MLSS with LT+1.0 (−0.2 ± 1.2 km·h⁻¹), VT₂ with LT+3.0 (<0.1 ± 1.3 km·h⁻¹), and MAS with \( MA_{\text{S}} \) (<0.1 ± 0.4 km·h⁻¹).

Table 3 shows the 95% confidence interval for main physiological parameters under study.

DISCUSSION

The main findings of the current study were that (1) LT obtained during a 12–14 min, 1 km·h⁻¹ per minute GXT is a valid method to determine the main physiological parameters of the aerobic–anaerobic transition, (2) LT, LT+1 and LT+3.0 are

and VT₂), MLSS, and MAS against the other parameters was conducted over the means obtained in the trials by ANOVA, ICC, and Bland-Altman bias. Analyses were performed using GraphPad Prism 6.0 (GraphPad Software, Inc., CA, USA) and SPSS software version 19.0 (IBM Corp., Armonk, NY, USA).

\[
MA_{\text{S}} = V_{\text{peak}} \times 0.8348 + 2.308
\]
solid predictors of VT1, MLSS, and VT2, respectively, (3) the MLSS was identified as a midpoint between VT1 and VT2, and (4) an estimated maximal aerobic speed (MASEST) can be accurately obtained (error ± 0.3 km h⁻¹) from the fastest speed achieved during the current GXT (Vpeak). This study adds to the existing literature by providing a valid alternative test based on blood lactate to obtain performance workloads without the need of using indirect calorimetry (less affordable technology). In addition, we contribute with an accurate method to estimate the MAS, which is one of the most used indicators to set training intensities in running. To our knowledge, this is the first report examining the validity and reliability of such an extensive battery of tests and parameters to determine critical workloads in runners.

The high reliability values found in physiological measurements between the two GXT EXP treadmill trials concurs with those previously reported in cycle ergometer (Pallarés et al., 2016). In addition, our results allow us to discourage using V-Slope when other parameters are available. Although the causes that might explain these effects are very difficult to isolate and quantify, it is arguable that an individualized workload adjustment approach accounted for these increments (García-Pallarés et al., 2009; Wolpern et al., 2015). In the current GXT with 1-min increments, athletes started at 13 km h⁻¹ below their Vpeak previously determined during the GXT PRE session. By doing this, it is guaranteed that the athlete is running at the optimal intensity to end up at their maximum workload after 12–14 min avoiding cardiac drift, local acidosis, and allowing a clear detection of ventilatory and LT. Additionally, getting maximal values of VO₂max. Considering this information, individual GXT protocols based on athletes’ maximal speed should be developed to enhance the consistency of data during physiological evaluations.

A number of studies have investigated the relationship between ventilatory threshold and blood lactate concentration in endurance athletes. Authors agreed that workloads at the first ventilatory threshold (i.e., VT1) are strongly related to the workload at which lactate starts to increase above resting values (LT; Wasserman et al., 1973; Lucía et al., 2000; Pallarés et al., 2016). Our findings corroborate this association between VT1 and LT but showing a greater external workload in runners (VT1 = 59–65% of MAS) compared to cyclists [VT1 ∼ 51.5% of maximal aerobic power (MAP)]. These findings suggest that running describes a great relative external workload associated with the VT1 response. Therefore, smaller errors in detecting ventilatory thresholds may have a greater negative impact on the running performance compared to other disciplines like cycling, for instance, misguided training prescription, undesirable physical adaptations, and a greater probability of the appearance of the interference phenomenon during concurrent training (García-Pallarés and Izquierdo, 2011). In turn, there is no clear agreement about which LT better reflects VT2 intensities. A previous experiment in cyclists (Pallarés et al., 2016) determined a high correlation between VT2 and LT+2 mMol L⁻¹, followed by the OBLA4mM, which established high intensities at ∼80% of their maximal aerobic power (MAP). Interestingly enough, we identified a greater CBL in runners during the transition phase, locating the VT2 at LT+3 mMol L⁻¹ intensities, setting the high intensity limit at 84–87% of the MAS. The existing physiological differences between running and cyclists may explain these disparities. Millet et al. (2009) reviewed the literature and identified a list of potential distinguishing factors between running and cycling physiological demands. The authors pointed out differences on ventilatory responses to exercise in terms of exercise-induced arterial hypoxaemia, O₂ diffusion capacity, ventilatory fatigue, and pulmonary mechanics. Moreover, other factors like running/cycling economy (higher delta efficiency in running), muscle recruitment patterns (greater muscle mass involved and eccentric phase activity in running), and ventilation impairment (higher in cycling) may account for these differences.

The MLSS constitutes another essential physiological event in endurance performance, as it is the maximal workload that can be maintained without elevations in blood lactate concentration (MLSS). Previous authors have proposed that MLSS workload coincides with the one for VT2 (Smekal et al., 2012). In contrast to this assumption, recent investigations in cyclists elucidated that MLSS may correspond to a lower exercise intensity of VT2 and matches better with the midpoint between both ventilatory thresholds (Pallarés et al., 2016; Peinado et al., 2016). In support of this theory, our findings revealed that MLSS intensity (72–74% of MAS) constitutes a transition between VT1 (59–65% of MAS) and VT2 (84–87% of MAS). Moreover, MLSS was highly associated with LT+1 mMol L⁻¹. In cyclists, MLSS has been associated with LT+0.5 mMol L⁻¹ (Pallarés et al., 2016). These increments on CBL at the same relative intensity might indicate

### TABLE 1 | Reliability of lactate and ventilatory tests.

|                  | CV   | ICC  | Bias (SD) | LoA 95% |
|------------------|------|------|-----------|---------|
| VT1              | 2.08 | 0.98 | 0.22 (0.47) | −0.71; 1.15 |
| VT2              | 1.92 | 0.95 | 0.13 (0.63) | −1.11; 1.37 |
| MAS              | 2.20 | 0.91 | 0.36 (0.64) | −0.9; 1.62 |
| LT               | 1.99 | 0.98 | 0.09 (0.43) | −0.76; 0.94 |
| VT2+0.5          | 1.23 | 0.96 | 0.07 (0.87) | −1.84; 1.78 |
| VT2+1.0          | 3.49 | 0.96 | 0.07 (0.79) | −1.48; 1.62 |
| VT2+1.5          | 3.08 | 0.96 | 0.10 (0.76) | −1.39; 1.59 |
| VT2+2.0          | 2.99 | 0.97 | 0.07 (0.69) | −1.29; 1.43 |
| VT2+2.5          | 2.53 | 0.96 | 0.11 (0.73) | −1.33; 1.56 |
| VT2+3.0          | 2.46 | 0.96 | 0.10 (0.77) | −1.41; 1.61 |
| V-Slope          | 2.58 | 0.84 | 0.31 (1.11) | −1.87; 2.49 |
| DMAX             | 2.12 | 0.94 | 0.27 (1.09) | −1.87; 2.41 |
| OBLA4mM          | 3.08 | 0.96 | 0.48 (0.97) | −1.43; 2.39 |
| Vpeak            | 2.79 | 0.94 | 0.12 (0.42) | −0.71; 0.96 |

CV: Coefficient of variation; ICC: Intraclass coefficient; and Bland-Altman results. Vpeak, the fastest velocity achieved at the end of the graded exercise testing protocol; MAS: Maximal aerobic speed; VT1, First ventilatory threshold; MLSS: Maximal lactate steady state; VT2, Second ventilatory threshold; LT, Lactate threshold; LT+0.5, +1.0, +1.5, +2.0, +2.5, +3.0. Concentrations above lactate threshold. DMAX, Maximum distance between the slope of a polynomial and the line connecting both ends; OBLA4mM, Onset blood lactate accumulation 4 mM; LoA, 95% limit of agreement.
| Speed (km·h⁻¹) | Blood lactate (mMol·L⁻¹) | VT₁ | MLSS | VT₂ | MAS |
|----------------|--------------------------|-----|------|-----|------|
| M ± SD         | M ± SD                   | Student t | ICC | Bland altman | p | Bias (SD) | LoA 95% | p | Bias (SD) | LoA 95% | p | Bias (SD) | LoA 95% |
| VT₁            | 11.5 ± 1.8               | 2.2 ± 0.7 | <0.01 | 0.91 | −2.0 (0.9) | −3.8 | −0.3 | <0.01 | 0.94 | −2.3 (0.7) | −3.6 | −1.0 |
| MLSS           | 13.5 ± 1.1               | 3.3 ± 1.1 | <0.01 | 0.92 | −4.3 (0.9) | −6.1 | −2.6 | <0.01 | 0.95 | −5.0 (0.6) | −6.2 | −3.9 | <0.01 | 0.91 | −3.1 (1.8) | −6.7 | 0.5 |
| VT₂            | 15.8 ± 1.4               | 5.7 ± 1.9 | <0.01 | 0.86 | −7.6 (2.0) | −11.6 | −3.7 | <0.01 | 0.95 | −5.0 (0.6) | −6.2 | −3.9 | <0.01 | 0.83 | 4.1 (1.4) | 1.4 | 6.9 |
| MAS            | 18.1 ± 1.3               | 10.6 ± 3.1 | <0.01 | 0.57 | 0.90 | −0.3 (1.2) | −2.7 | 2.1 | <0.01 | 0.82 | 1.8 (1.3) | −0.9 | 4.4 | <0.01 | 0.84 | 3.0 (1.4) | 0.3 | 5.8 |
| LT             | 11.8 ± 1.8               | 2.3 ± 0.4 | 0.05 | 0.90 | −0.3 (1.2) | −2.7 | 2.1 | <0.01 | 0.82 | 1.8 (1.3) | −0.9 | 4.4 | <0.01 | 0.84 | 3.0 (1.4) | 0.3 | 5.8 |
| LT+0.5         | 12.7 ± 2.1               | 2.8 ± 0.4 | 0.03 | 0.93 | −1.3 (1.1) | −3.5 | 0.9 | <0.01 | 0.82 | 0.7 (1.4) | −2.0 | 3.5 | <0.01 | 0.84 | 3.0 (1.4) | 0.3 | 5.8 |
| LT+1.0         | 13.6 ± 2.0               | 3.2 ± 0.4 | <0.01 | 0.93 | −2.2 (1.1) | −4.4 | −0.1 | <0.01 | 0.84 | 2.1 (1.4) | −0.7 | 4.9 | <0.01 | 0.77 | 4.8 (1.5) | 2.0 | 7.7 |
| LT+1.5         | 14.3 ± 1.9               | 3.8 ± 0.4 | <0.01 | 0.93 | −2.8 (1.1) | −5.0 | −0.7 | <0.01 | 0.84 | 1.5 (1.3) | −1.1 | 4.1 | <0.01 | 0.78 | 4.1 (1.4) | 1.4 | 7.0 |
| LT+2.0         | 14.8 ± 1.9               | 4.3 ± 0.4 | <0.01 | 0.92 | −3.4 (1.1) | −5.6 | −1.3 | <0.01 | 0.84 | 0.9 (1.3) | −1.7 | 3.5 | <0.01 | 0.78 | 3.6 (1.4) | 0.9 | 6.4 |
| LT+2.5         | 15.3 ± 1.9               | 4.8 ± 0.4 | <0.01 | 0.91 | −3.9 (1.1) | −6.1 | −1.8 | <0.01 | 0.85 | −1.8 (1.1) | −4.1 | 0.5 | <0.01 | 0.78 | 3.1 (1.4) | 0.4 | 5.9 |
| LT+3.0         | 15.8 ± 1.8               | 5.3 ± 0.4 | <0.01 | 0.91 | −4.3 (1.1) | −6.5 | −0.3 | <0.01 | 0.85 | −2.3 (1.2) | −4.6 | −0.1 | <0.01 | 0.78 | 2.7 (1.4) | −0.1 | 5.4 |
| V-Slope        | 14.6 ± 1.4               | 4.4 ± 2.1 | <0.01 | 0.91 | −3.1 (1.5) | −6.1 | −0.2 | <0.01 | 0.85 | −1.1 (1.2) | −3.6 | 1.4 | <0.01 | 0.57 | 3.9 (1.5) | 1.0 | 6.8 |
| DMAX           | 14.1 ± 1.6               | 3.7 ± 1.0 | <0.01 | 0.96 | −2.6 (0.8) | −4.2 | −1.1 | <0.01 | 0.94 | −0.6 (0.8) | −2.1 | 0.9 | <0.01 | 0.92 | 4.4 (0.9) | 2.6 | 6.2 |
| OBLA₉₄ mmol·L⁻¹| 14.5 ± 2.1               | 4.0 ± 0.1 | <0.01 | 0.93 | −3.0 (1.1) | −5.2 | −0.9 | <0.01 | 0.91 | 1.7 (1.0) | −0.3 | 3.7 | <0.01 | 0.78 | 3.9 (1.5) | 1.0 | 7.0 |
| Vpeak          | 19.3 ± 1.3               | 12.0 ± 3.9 | <0.01 | 0.93 | −8.5 (2.1) | −12.7 | −4.4 | <0.01 | 0.96 | −5.8 (0.6) | −7.0 | −4.7 | <0.01 | 0.92 | −0.9 (0.5) | −1.9 | 0.2 |
| MASest         | 18.5 ± 1.1               | 7.6 ± 2.0 | <0.01 | 0.91 | −7.6 (2.0) | −11.6 | −3.7 | <0.01 | 0.91 | −3.1 (1.9) | −6.9 | 0.7 | <0.01 | 0.98 | <0.1 (0.4) | −0.9 | 0.9 |

Comparison of running speeds at VT₁, MLSS, VT₂, and MAS against the rest of parameters. VT₁, First ventilatory threshold; MLSS, Maximal lactate steady state; VT₂, Secondary ventilatory threshold; MAS, Maximal aerobic speed; LT, Lactate threshold; LT+0.5, +1.0, +1.5, +2.0, +2.5, +3.0. Concentrations above lactate threshold, DMAX, Distance between the slope of a polynomial and the line connecting both ends, OBLA₉₄ mmol·L⁻¹, Onset blood lactate accumulation 4 mMol·L⁻¹, Vpeak, Fastest velocity achieved at the end of the graded exercise testing protocol; MASest, Theoretical MAS estimated from Vpeak. The strongest relationships are highlighted in bold. LoA, 95% limit of agreement.
a greater energy cost in running at MLSS workload, which may imply earlier fatigue and lower performance by accelerating glycogen depletion (Coyle et al., 1986).

A main contribution of the current study is to provide an estimated MAS (MAS<sub>EST</sub>) from the maximal speed achieved (V<sub>peak</sub>) at the end of the GXT protocol with a minimal error of ±0.3 km·h<sup>-1</sup>. Main physiological events (VT and MLSS) are related to a given percentage of MAS (<sup>2</sup>V<sub>T</sub>, or <sup>2</sup>LT) or MAS, Maximal aerobic speed; HRR, Heart rate reserve; RPE, Rate of perceived exertion.

### REFERENCES

ACSM (2013). ACSM’s Guidelines for Exercise Testing and Prescription, 9th Edn. Philadelphia, PA: Lippincott Williams and Wilkins.

Aman, M., Subudhi, A. W., and Foster, C. (2006). Predictive validity of ventilatory and lactate thresholds for cycling time trial performance. Scand. J. Med. Sci. Sports 16, 27–34. doi: 10.1111/j.1600-0838.2004.00424.x

Beneke, R. (2003). Methodological aspects of maximal lactate steady state-implications for performance testing. Eur. J. Appl. Physiol. 89, 95–99. doi: 10.1007/s00421-002-0783-1

Beneke, R., Leithäuser, R. M., and Ochentel, O. (1996). Determination of MAS using a cheaper and affordable method such as CBL. Furthermore, the current GXT individualized protocol (i.e., starting at 13 km·h<sup>-1</sup> below the athlete’s MAS with increments of 1 km·h<sup>-1</sup>/min until) appears to be a promising method to determine training zones in well-trained runners. What is now required is to test the effectiveness of training plans according to the current 5-zone proposal (Table 4). In addition, future investigations should examine the validity of this protocol in amateur and female runners to enhance its applicability within the endurance sport community.

### ETHICS STATEMENT

All procedures performed in this study involving human participants were in accordance with the ethical standards of the institutional Human Research Ethics Committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. The study was approved by the Bioethics Commission of the University of Murcia. Written informed consent was obtained from all subjects prior to participation.

### AUTHOR CONTRIBUTIONS

JP and VC-E: Conception and design of the experiments; JP, VC-E, RM-N, and AM-C: Pre-testing, experimental preparation, and data collection; VC-E, JC-I, and JP: data analysis. The first draft of the manuscript was written by VC-E, JC-I, and JP. All co-authors edited and proofread the manuscript and approved the final version.

### ACKNOWLEDGMENTS

The authors wish to thank the subjects for their invaluable contribution to the study.

---

TABLE 3 | 95% confidence interval values for main physiological events.

| Intensity | Zone | MAS (%) | V<sub>peak</sub> (%) | HR<sub>Max</sub> (%) | HRR (%) | RPE<sub>6–20</sub> |
|-----------|------|---------|------------------|------------------|---------|-------------|
| VT<sub>1</sub> | 59–65 | 77–81 | 68–74 | 10–12 |
| MLSS | 72–74 | 85–87 | 79–83 | 12–13 |
| VT<sub>2</sub> | 84–87 | 91–93 | 81–98 | 15–16 |
| MAS | 100 | 98–100 | 98–100 | 18–20 |

VT<sub>1</sub>, First ventilatory threshold; MLSS, Maximal lactate steady state; VT<sub>2</sub>, Second ventilatory threshold; MAS, Maximal aerobic speed; HRR, Heart rate reserve; RPE, Rate of perceived exertion.

TABLE 4 | Personal author’s approach for exercise prescription (training zones).

| Intensity | Zone | MAS (%) | V<sub>peak</sub> (%) | HR<sub>Max</sub> (%) | HRR (%) | RPE<sub>6–20</sub> |
|-----------|------|---------|------------------|------------------|---------|-------------|
| 70–90% VT<sub>1</sub> or LT | R0 | 43–56 | 40–52 | 55–70 | 50–64 | 8–10 |
| 90–110% VT<sub>1</sub> or LT | R1 | 57–68 | 53–64 | 71–83 | 65–77 | 11–12 |
| 95–105% MLSS or LT+1.0 | R2 | 69–79 | 65–75 | 84–88 | 78–84 | 13–14 |
| 95–105% VT<sub>2</sub> or LT+3.0 | R3 | 80–93 | 76–89 | 89–94 | 85–93 | 15–16 |
| 95–105% VO<sub>2</sub>max | R3 | 94–105 | 90–100 | >95 | >94 | >17 |

V<sub>peak</sub>, the fastest velocity achieved at the end of the graded exercise testing protocol; VT<sub>1</sub>, First ventilatory threshold; MLSS, Maximal lactate steady state; VT<sub>2</sub>, Second ventilatory threshold; MAS, Maximal aerobic speed; HR<sub>Max</sub>, Maximal heart rate; HRR, Heart rate reserve; RPE, Rate of perceived exertion.
Jones, A. M., Wilkerson, D. P., DiMenna, F., Fullford, J., and Poole, D. C. (2007). Muscle metabolic responses to exercise above and below the “critical power” assessed using 31P-MRS. Am. J. Physiol. Regul. Integr. Comp. Physiol. 294, R853–R893. doi: 10.1152/ajpregu.00731.2007

Julio, U. F., Panissa, V. L. G., Shvivien, S. A., and Franchini, E. (2017). Effect of protocol manipulation for determining maximal aerobic power on a treadmill and cycle ergometer: a brief review. Strength Cond. J. 5, 58–71. doi: 10.1519/JSC.0000000000000350

Kuipers, H., Rietjens, G., Verstappen, F., Schoenmakers, H., and Hofman, G. (2003). Effects of stage duration in incremental running tests on physiological variables. Int. J. Sports Med. 24, 486–491. doi: 10.1055/s-2003-42020

Jacour, L. R., Padilla-Maguncalaya, S., Chatard, J. C., Arsac, L., and Barthélémy, J. C. (1991). Assessment of running velocity at maximal oxygen uptake. Eur. J. Appl. Physiol. Occup. Physiol. 62(2), 77–82. doi: 10.1007/BF00626760

Léger, L., and Boucher, R. (1980). An indirect continuous running multistage field test: the Université de Montréal track test. Can. J. Appl. Sport Sci. 5, 77–84.

Llodi, I., Gorostiaga, E., García-Tabar, I., Granados, C., and Sánchez-Medina, L. (2016). Estimation of the maximal lactate steady state in endurance runners. Int. J. Sports Med. 37, 539–546. doi: 10.1007/s00421-010-22653

Lucía, A., Hoyos, J., Pérez, M., and Chicharro, J. L. (2000). Heart rate and performance parameters in elite cyclists: a longitudinal study. Med. Sci. Sports Exerc. 32, 1777–1782. doi: 10.1097/00005768-200010001-00018

Lucía, A., Sánchez, O., Carvajal, A., and Chicharro, J. L. (1999). Analysis of the aerobic-anaerobic transition in elite cyclists during incremental exercise with the use of electromyography. Br. J. Sports Med. 33, 178–185. doi: 10.1136/BJSMM.33.3.178

Lukasci, H. C., Bolonchuk, W. W., and Klevay, L. M. (1989). Comparison of metabolic responses and oxygen cost during maximal exercise using three treadmill protocols. J. Sports Med. Phys. Fitness 29, 223–229. doi: 10.1519/JSMS.0000000000000193

Mann, T. N., Lambert, R. P., and Lambert, M. I. (2014). High responders and low responders: factors associated with individual variation in response to standardized training. Sports Med. 44, 1113–1124. doi: 10.1007/s00421-014-0197-3

McLaughlin, J. E., Howley, E. T., Bassett, D. R., Thompson, D. L., and Fitzhugh, E. C. (2010). Test of the classic model for predicting endurance running performance. Med. Sci. Sports Exerc. 42, 991–997. doi: 10.1249/MSS.0b013e3181c0669d

Midgley, A. W., McNaughton, L. R., and Carroll, S. (2007). Time at V02max during intermittent treadmill running: test protocol dependent or methodological artefact? Int. J. Sports Med. 28, 934–939. doi: 10.1007/s00421-007-94972

Millet, G. P., Vleck, V. E., and Bentley, D. J. (2009). Physiological differences between cycling and running: lessons from triathletes. Sports Med. 39, 179–206. doi: 10.1007/s00421-007-9303-0

Mora-Rodríguez, R., Pallarés, J. G., López-Gullón, J. M., López-Samanes, A., Fernández-Elias, V., and Ortega, J. F. (2016). Improvements on neuromuscular performance with caffeine ingestion depend on the time-of-day. J. Sci. Med. Sport 18, 338–342. doi: 10.1016/j.jsams.2014.04.010

Pallarés, J. G., Moran-Navarro, R., Fernando Ortega, J., Emilio Fernandez-Elias, V., and Mora-Rodríguez, R. (2016). Validity and reliability of ventilatory and blood lactate thresholds in well-trained cyclists. PLoS ONE 11:e0163389. doi: 10.1371/journal.pone.0163389

Peinado, A. B., Filho, D. P., Díaz, V., Benito, J. P., Álvarez-Sánchez, M., Zapico, A. G., and Calderón, F. J. (2016). The midpoint between ventilatory thresholds approaches maximal lactate steady state intensity in amateur cyclists. Biol. Sport 33, 373–380. doi: 10.5604/20831862.1221812

Scharhag-Rosenberger, F., Walitzeck, S., Kindermann, W., and Meyer, T. (2012). Differences in adaptations to 1 year of aerobic endurance training: individual patterns of nonresponse. Scand. J. Med. Sci. Sports 22, 113–118. doi: 10.1111/j.1600-0838.2010.01139.x

Sjödin, B., and Jacobs, I. (1981). Onset of blood lactate accumulation and marathon running performance. Int. J. Sports Med. 2, 23–26. doi: 10.1055/s-2008-1034579

Smoral, G. von, Duvillard, S. P., Pokan, R., Hofmann, P., Braun, W. A., Arciero, P. J., et al. (2012). Blood lactate concentration at the maximal lactate steady state is not dependent on endurance capacity in healthy recreationally trained individuals. Eur. J. Appl. Physiol. 112, 3079–3086. doi: 10.1007/s00421-011-2283-7
Stöggl, T., and Sperlich, B. (2014). Polarized training has greater impact on key endurance variables than threshold, high intensity, or high volume training. *Front. Physiol.* 5:33. doi: 10.3389/fphys.2014.00033

Stöggl, T., and Sperlich, B. (2015). The training intensity distribution among well-trained and elite endurance athletes. *Front. Physiol.* 6:295. doi: 10.3389/fphys.2015.00295

Stratton, E., O’Brien, B. J., Harvey, J., Blitvich, J., McNicol, A. I., Janissen, D., et al. (2009). Treadmill velocity best predicts 5000-m run performance. *Int. J. Sports Med.* 30, 40–45. doi: 10.1055/s-2008-1038761

Svedahl, K., and Maclntosh, B. R. (2003). Anaerobic threshold: the concept and methods of measurement. *Can. J. Appl. Physiol.* 28, 299–323. doi: 10.1139/h03-023

Wasserman, K., Whipp, B. J., Koyl, S. N., and Beaver, W. L. (1973). Anaerobic threshold and respiratory gas exchange during exercise. *J. Appl. Physiol.* 35, 236–243. doi: 10.1152/jappl.1973.35.2.236

Weltman, A., Snead, D., Stein, P., Seip, R., Schurrer, R., Rutt, R., and Weltman, J. (1990). Reliability and validity of a continuous incremental treadmill protocol for the determination of lactate threshold, fixed blood lactate concentrations, and VO$_2$max. *Int. J. Sports Med.* 11, 26–32. doi: 10.1055/s-2007-1024757

Wolpern, A. E., Burgos, D. J., Janot, J. M., and Dalleck, L. C. (2015). Is a threshold-based model a superior method to the relative percent concept for establishing individual exercise intensity? A randomized controlled trial. *BMC Sports Sci. Med. Rehabil.* 7:16. doi: 10.1186/s13102-015-0011-z

**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2018 Cerezuela-Espejo, Courel-Ibáñez, Morán-Navarro, Martínez-Cava and Pallarés. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.