A method-comparison study regarding the validity and reliability of the Lactate Plus analyzer

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ABSTRACT

Objectives: The aims of this study were to: (1) determine the validity and reliability of the Nova Biomedical Lactate Plus portable analyzer, and quantify any fixed or proportional bias; (2) determine the effect of any bias on the determination of the lactate threshold and (3) determine the effect that blood sampling methods have on validity and reliability.

Design: In this method comparison study we compared blood lactate concentration measured using the Lactate Plus portable analyzer to lactate concentration measured by a reference analyzer, the YSI 2300.

Setting: University campus in the USA.

Participants: Fifteen active men and women performed a discontinuous graded exercise test to volitional exhaustion on a motorised treadmill. Blood samples were taken via finger prick and collected in microcapillary tubes for analysis by the reference instrument at the end of each stage. Duplicate samples for the portable analyzer were either taken directly from the finger or from the micro capillary tubes.

Primary outcome measurements: Ordinary least products regressions were used to assess validity, reliability and bias in the portable analyzer. Lactate threshold was determined by visual inspection.

Results: Though measurements from both instruments were correlated (r=0.91), the differences between instruments had large variability (SD=1.45 mM/l) when blood was sampled directly from finger. This variability was reduced by ~95% when both instruments measured blood collected in the capillary tubes. As the proportional and fixed bias between instruments was small, there was no difference in estimates of the lactate threshold between instruments. Reliability for the portable instrument was strong (r=0.99, p<0.05) with no proportional bias (slope=1.02) and small fixed bias (~0.19 mM/l).

Conclusions: The Lactate Plus analyzer provides accurate and reproducible measurements of blood lactate concentration that can be used to estimate workloads corresponding to blood lactate transitions or any absolute lactate concentrations.

ARTICLE SUMMARY

INTRODUCTION

Not only is blood lactate accumulation a common measure in the physiological assessment of endurance athletes, but is also an important clinical measure.¹–⁴ Portable lactate analyzers have advantages over bench top models including: (1) their ability to rapidly sample blood lactate concentration ([lactate]), in or outside the laboratory;
(2) they require a much smaller sample of blood (0.5–0.7 μl) than many bench top analyzers (25–75 μl) and
(3) they can be purchased and operated at a lower cost
than many bench top models.

Several studies have attempted to evaluate the validity
and reliability of these portable analyzers.3–10 While the
majority of studies report that the [lactate] measured
using portable analyzers is similar to those of various
bench top models, the mean difference between the ref-
ERENCE and portable analyzers can be as much as 1.0 mM/l. This can represent nearly 10% of the full
range of values in some populations.11 This level of dis-
agreement could be explained by the presence of sys-
Tematic measurement error. Systematic measurement
error can result in a proportional bias, where one instru-
ment produces values that are different from those of
another instrument by an amount that is proportional to
the level of the measured variable, and/or a fixed bias,
where one instrument gives values that are different
from those of another instrument by a constant
amount.12 13 Thus, similar mean values between lactate
analyzers could occur while the portable analyzer pro-
duces low values at lower [lactate] and high values at
higher [lactate] or vice versa. Previous studies have pri-
marily relied on Bland-Altman analysis to determine the
presence of any fixed bias. However, this approach does
not allow the independent determination of bias, and
thus has limited utility in assessing the presence of sys-
tematic measurement error. Therefore, while most data
appear to show a substantial proportional and/or fixed
bias, the presence and degree of bias in portable lactate
analyzers remains unresolved.3 4 6–10 Furthermore,
because previous studies have not directly examined
these biases it is unclear if they are large enough to
affect estimates of various lactate parameters, such as pH
or lactate threshold (LT).

Blood sampling techniques may also affect measure-
ment accuracy and reliability. Previous studies have
either used intravenous blood drawn directly into a
syringe,5 7 9 or capillary blood from a finger stick drawn
into capillary tubes and then mixed as would be done in
the laboratory.5 10 Portable analyzers, however, are
designed to sample blood directly from a puncture for
ease of use in the field. When using a finger stick to
draw blood it is not uncommon to require ‘milking’ of
the finger to get an adequate sample. This may dilute
the lactate concentration by increasing interstitial fluid
in the sample. It would seem important to understand
and quantify the effect of differing blood-sampling pro-
cedures on the accuracy and reliability of these portable
analyzers.

Given the questions that remain regarding the validity
and reliability of portable lactate analyzers the specific
aims of the present study were: (1) to determine the val-
idity and reliability of the Lactate Plus analyzer (Nova
Biomedical), and quantify any fixed and/or propor-
tional bias and (2) determine the effect that blood
sampling methods have on validity and reliability.

METHODS
Fifteen young (20–36 years; mean=24.5 years) men and
women (6 women) participated in the study. All subjects
reported at least 90 min of moderate to vigorous physical
activity each week. All individuals read and signed an
informed consent. The Institutional Review Boards at
Wheaton College and the Northern Illinois University
approved this study. All procedures conformed to the
Declaration of Helsinki.

Instruments
To determine the validity of the Lactate Plus analyzer we
used the YSI 2300 Stat Plus Glucose and Lactate analyzer
from Yellow Springs Instruments (Yellow Springs, Ohio,
USA) as our reference instrument. This bench top
laboratory analyzer uses a membrane-bound enzyme
electrochemical methodology. L-Lactate oxidase is immo-
bilised in a thin membrane placed over an electrochem-
ical probe. The enzyme catalyses the conversion of
L-lactate to pyruvate and hydrogen peroxide, the latter
then being oxidised at the platinum anode to measure
lactate concentration in whole blood or plasma. A new
membrane was used for each data collection session.

The analyzer was initially calibrated using 5, 15 and
30 mM/l solutions. In addition, an automated quality
control was performed in triplicate every 45 min using a
5 mM/l solution. Blood samples were collected from a
finger stick into two heparinised capillary tubes. Blood
was then mixed in a micro centrifuge tube. Two 25 μl
samples were sequentially aspirated and measured by
the analyzer.

The Lactate Plus analyzer uses an electrochemical lactate
oxidase biosensor to measure lactate concentration in a
0.7 μl sample. Following the manufacturer’s instructions we
used low (1.0–1.6 mM/l) and high (4.0–5.4 mM/l) quality
control solutions to ensure the lactate analyzer was oper-
ating properly at the beginning of each data collection
session. For the first nine participants three blood
samples were taken directly from the finger between each
stage of the graded exercise test (GXT). All samples were
taken in this order: (1) portable directly from finger,
(2) capillary tubes for the YSI 2300 from the finger and
(3) a second sample directly from finger using the port-
able analyzer. To assess the effect of blood sampling tech-
niques on the accuracy of the portable analyzer blood
was drawn from the finger into capillary tubes and allo-
cated to both the YSI 2300 and portable analyzer for the
last six participants.

Graded exercise
Participants performed a discontinuous GXT on a
motorised treadmill (Quinton TM65). Each stage lasted
2 min with a 1 min blood sampling period between
stages. The finger was prepared for sampling just prior
to the end of each exercise stage. During the 1 min
blood collection period participants straddled the tread-
mill belt while blood samples were taken from a finger.
After 1 min the participants resumed exercise at a
higher speed or grade. The initial speed was 1.55 m/s and 0% grade. The speed was increased by either 0.50 or 0.67 m/s for each stage until the participant’s heart rate was at least 80% of their age-predicted maximum (220 age). After this point the speed remained constant while grade was increased 2.5% for each stage. Exercise was continued until volitional exhaustion.

Data analysis
Two methods were used to assess validity. First, a Bland-Altman plot was constructed to allow the reader to more directly compare our data with those of previous studies since this is the approach typically used. However, because fixed and proportional biases cannot be determined independently from these plots, ordinary least products regression analysis was used. Validity was determined from the correlation coefficient in combination with the presence and degree of bias. The degree of fixed bias was determined from the y-intercept 95% CIs. If the CI for the intercept includes the value of zero, then there is no fixed bias. Proportional bias was determined from the 95% CI for the slope. If the CI for the slope includes the value of 1.0, then there is no proportional bias. Ordinary least products regression gives different slopes and y- intercepts than does least squares regression because error is assumed in both portable and bench top analyzers.12 13

LT was defined as the point at which blood [lactate] began to increase in a non-linear fashion.14 15 The threshold was estimated by plotting [lactate] against GXT stage. These graphs were visually inspected to determine the lines of best fit by the two evaluators. The following guidelines were used to help guide the evaluators: (1) at least three data points were included in each line, (2) both lines contained unique data points and (3) lines were chosen that produced the highest R² with the smallest CIs. Once the lines were chosen the equations for each line were set equal to one another and solved for the point of intersection (figure 1). The values from each evaluator were averaged.16 These equations were also used to calculate the stage that corresponded to an absolute blood [lactate] of 2.5 and 4.0 mM/l. A t test for paired data was used to compare means between analyzers. A p value of<0.05 was considered statistically significant.

Reliability was determined using ordinary least products regression to quantify the relationship between sequential measurements for both instruments.

RESULTS
Validity
Lactate values during graded exercise ranged from 1.2 to 16.4 mM/l. When both portable and bench top blood samples are each taken directly from the finger the mean difference between [lactate] measured by the portable and bench top analyzers was small across the full range of lactate values as depicted in figure 2. While the mean difference between the two instruments was near zero, differences between the instruments had a large variability (SD=1.45 mM/l). Even though there can be large differences between values measured by the portable and bench top analyzers, the paired measurements were highly correlated as shown in figure 3A. Least-product regression indicated a small fixed bias (y-intercept=-0.28 mM/l) between [lactate] measured with the portable and bench top analyzers. There was no evidence of a proportional bias (95% CI 0.94 to 1.15). When the same mixed blood sample was used by both analyzers, the fixed bias was reduced to −0.056 mM/l, while a small proportional bias was evident (slope=1.08) as shown in figure 3B.

Regardless of a blood sampling approach there was excellent agreement between estimates of the LT based on
lactate values from the portable analyzer compared with those from the bench top analyzer ($r=0.97$). Moreover, there was neither a proportional bias (95% CI for slope: 0.910 to 1.098), nor a fixed bias (95% CI for y-intercept: −0.396 to 0.325) in estimates of the LT from the portable analyzer. Given the lack of bias it is not surprising there was no difference between blood [La] at the LT (2.88NOVA±0.53 vs 3.15YSI±0.46 mM/l; $p=0.32$). In addition the stages corresponding to absolute blood lactate values of 2.5 mM/l (2.99NOVA vs 2.92YSI) and 4.0 mM/l (4.64NOVA vs 4.61YSI) were not different between portable and bench top values ($p=0.86$ for both).

**Reliability**

The relationship between duplicate measurements of [lactate] by the bench top analyzer was very strong ($r=0.99$, $p<0.05$). Ordinary least products regression indicated no proportional bias (slope=0.99), and a small fixed bias (0.059 mM/l; figure 4). Ordinary least products regression revealed a small proportional (slope=1.20) and fixed bias (−0.54 mM/l; figure 3A) when the two duplicate blood samples for the portable analyzer were taken directly from the fingers. Thus, the reading from the second sample was typically lower than that from the first. However, when two duplicate measurements were taken from the same mixed blood sample, there was no proportional bias (slope=1.02) and the fixed bias was reduced to −0.19 mM/l).

A total of 242 blood samples were taken using the portable analyzer. Twenty-seven of these attempts resulted in error messages (E-4—insufficient sample). Thus, about 1-in-10 measurement attempts resulted in errors.

**DISCUSSION**

There were three new findings in our study: (1) The very small proportional bias indicates that the Lactate Plus analyzer is a highly linear instrument, (2) multiple blood samples directly from the finger increases measurement error and (3) the small proportional and fixed bias in the portable analyzer does not affect the ability to determine the LT.

We chose to use ordinary least products regression to characterise the relation between the Lactate Plus analyzer and our reference analyzer. Most studies have employed a combination of Bland-Altman plots and least squares regression to determine the degree of agreement between various portable analyzers and a corresponding reference analyzer.3–10 The mean difference between analyzers, as determined through Bland-Altman plots, is determined by the interaction of any fixed and proportional bias. Therefore, the mean difference between methods does not solely reflect the accuracy or fixed bias of the device, but in some cases, the presence of a proportional bias or loss of linearity. The use of least squares regression to characterise the level of proportional bias, as reflected in the slope of the linear relation, is skewed because all error is assigned to the dependent variable, in this case the portable analyzer.
Numerous studies have compared blood lactate measured with various portable analyzers to several different bench top analyzers. All have reported that these portable analyzers produce similar lactate values compared with their bench top counterparts with average differences ranging from -0.8 to 1.0 mM/L. However, differences of almost 1.0 mM/L can significantly impact the use of absolute [lactate] to characterise training intensity or efficacy. Weltman et al. reported that women who trained at an intensity corresponding to the 2.5 and the 4.0 mM/L absolute lactate concentrations. These lactate concentrations were chosen because they have both sport and clinical significance. The strong correlation coefficient and small biases suggest that the Lactate Plus analyzer can be used to accurately determine exercise intensities based on any blood lactate parameter.

Determination of the LT by visual inspection has come under scrutiny. To reduce subjectivity our approach to visual inspection is guided by several principles similar to those used by others. Several methods of assessing the LT have been proposed that purport to be more objective. However, many of these methods are known to be significantly affected by data outliers and/or missing data. Therefore, the choice of any analytical approach has a subjective component. While our approach likely produces LT values that are different from other approaches, it produced values consistent with other studies that employed similar approaches to LT estimation.

Duplicate sample readings from the Lactate Plus analyzer were strongly related, however there was a small fixed bias, indicating that the values from the second sample were consistently lower than the values from the first sample. In addition, there was a very small proportional bias. Both of these biases may be explained by using separate samples collected directly from the finger. The milking of the finger to obtain a blood sample can cause the dilution of the blood sample by interstitial fluid. The manufacturer warns the user against vigorous squeezing of the finger to obtain a blood drop. The use of a vasodilating agent would likely increase the lactate content of the blood sample.

The use of least products regression to compare methods avoids both of these issues, allowing independent and more accurate determination of any fixed or proportional bias. The use of absolute [lactate] to characterise training intensity or efficacy. Weltman et al. reported that women who trained at an intensity corresponding to about 2.5 mM/L showed greater improvement in blood lactate parameters, but less of an improvement in VO2 max than did women training at their LT. If true, then an error in the measurement of blood lactate concentration could lead to suboptimal improvements in either lactate parameters or VO2 max. Of the two studies that have tested the Lactate Plus analyzer, only Tanner et al. reported the absolute difference between this portable analyzer and a reference analyzer (-0.8 mM/L). Our data show a much smaller difference between the Lactate Plus and the YSI bench top analyzers (fixed bias=-0.056 mM/L). Though not specifically assessed, it does appear that Tanner’s reported difference between the hand held and reference analyzer is significantly influenced by a proportional bias (figures 4 and 5 from reference 8). The fact that our data shows little proportional bias (figure 3) may account for the greater agreement between analyzers that we observed. It is possible that if Tanner had been able to independently determine the proportional and fixed biases, their analysis may have revealed a small bias similar to ours. Differences in reference instruments would not likely explain the greater measurement error reported by Tanner, given that their instrument undergoes a three-point and two-point calibrations check every few hours, similar to our reference instrument.

Given that we found a very small proportional bias the estimation of the LT from [lactate] measured by the Lactate Plus analyzer agreed very well with those determined from [lactate] measured by the reference analyzer. Moreover, given the small fixed bias, it was not surprising that the lactate values from the portable analyzer provided similar estimates of the workload corresponding to the 2.5 and the 4.0 mM/L absolute lactate concentrations. These lactate concentrations were chosen because they have both sport and clinical significance. The strong correlation coefficient and small biases suggest that the Lactate Plus analyzer can be used to accurately determine exercise intensities based on any blood lactate parameter.

Figure 5: Ordinary least products regression analysis of the relation between sequential estimates of blood lactate concentration by the Lactate Plus portable lactate analyzer. (A) When separate samples for each analyzer were collected directly from finger. (B) When a common sample of blood was used by both analyzers. Regression equation and CIs for slope (B) and y-intercept (A) are presented.
Validity and reliability of a portable lactate meter

cream may resolve this issue. When we used the same mixed blood sample as the reference analyzer, the proportional bias was eliminated, while the fixed bias was reduced by approximately 65%.

We also found that the portable analyzer was unable to analyse the blood sample 11% of the time, presumably from an insufficient sample volume. This was surprising given that the Lactate Plus lactate analyzer provides an audible signal to indicate when the test strip has a sufficient volume of blood for analysis. Our experience has shown that anticipating the filling of the test strip can result in both the audible signal and an error. However, even when great care is taken, one can still get an audible full signal and the error message.

Ridenour et al. advocated for a switch from fetal blood sampling to lactate analysis. However, their data showed that the variability in blood [lactate] accounted for only 46% of the variability in pH. This could be owing to the significant proportional bias that is apparent in their data (see ref 1, figures 1 and 3). However, our analysis shows a fixed and proportional bias that are less than half reported by previous studies relying on Bland-Altman plots and simple comparison of means. This suggests the modest correlation between fetal [lactate] and blood pH is best attributed to the independent regulation of blood lactate and pH rather than unreliable measurement of [lactate].

We did not compare the Lactate Plus lactate analyzer with known standards. This limits the precision with which we can quantify the accuracy of the portable analyzer. However, our reference instrument was calibrated using three known lactate standards across a supraphysiological range. Our analysis assumes measurement error in both the portable and reference instrument. Thus it is likely that by comparing the Lactate Plus lactate analyzer directly to known lactate standards, our fixed bias would be reduced.

While some studies have used blood collected from trained athletes to compare portable lactate analyzers to bench top models, several do not. This seems quite appropriate given that the importance of accurate lactate measurement extends well beyond the athletic field. Our subjects were healthy and physically active, but not highly trained. This is unlikely to account for any difference between previous studies and ours given that we can find no reason to speculate that either lactate analyzer would more accurately measure [lactate] in one population compared with another.

Similarly, the choice of graded exercise protocol can affect LT determination. Thus, our use of a personalised, discontinuous GXT likely produced LT values different from some other protocols. However, this would have no affect on our ability to accomplish the aims of our study, specifically to compare estimates of LT between lactate measurements produced by the portable and reference analyzers.

In summary, the Lactate Plus analyzer is a valid and reliable instrument across a wide range of blood lactate concentrations. Any proportional or fixed bias in blood lactate concentration is nearly indistinguishable from zero. Therefore, the portable analyzer can be used to determine exercise intensities based on absolute or relative blood lactate concentrations. Sampling procedures can have a significant effect on the reliability of the portable analyzer, and the portable analyzer is prone to technical issues in nearly 1 of 10 samples.

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REFERENCES


