Impact of Phlebotomy Tourniquet Use on Blood Lactate Levels in Acutely Ill Patients

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ABSTRACT

Objective: Lactate levels are increasingly used to guide resuscitation efforts. Some surgical literature suggests that tourniquet use during phlebotomy falsely elevates results, although studies in healthy volunteers have not demonstrated this. The purpose of this study was to determine in clinical practice whether tourniquet use during the drawing of a lactate results in significantly altered levels compared to the result of a level drawn without a tourniquet. Methods: A prospective cohort study was carried out on emergency department patients whose clinical presentation led a physician to order a lactate level. Written informed consent was obtained from patients or their proxies. Study lactates were obtained using a tourniquet during the draw sequence of other laboratory studies. Lactate levels for clinical use were drawn per hospital protocol with no tourniquet. The time of lactate measurements and patient demographic information were recorded. Lactate levels for each patient were compared with the Wilcoxon Rank-Sum Test.

Results: 40 patients were consented and enrolled. The median clinical lactate level was 1.9 (interquartile range 1.5-2.6), and the median study lactate level was 1.9 (interquartile range 1.4-2.7). There was no difference between paired lactate values (p = 0.95).

Conclusions: Tourniquet use appears to have no impact on measured lactate levels. Our findings suggest that current practices at many institutions regarding lactate collection are likely too stringent and should be changed.

RÉSUMÉ

Objectif: Le taux de lactate sert de plus en plus à guider les efforts de réanimation. D’après certains travaux de recherche en chirurgie, l’utilisation du garrot durant la ponction veineuse augmenterait erronément le taux de lactate, bien que certaines études réalisées chez des témoins en bonne santé ne soient pas arrivées à la même conclusion. L’étude décrite ici avait donc pour but de déterminer si l’utilisation du garrot en clinique durant les prélèvements de sang en vue de la mesure du taux de lactate avait pour effet de modifier considérablement le taux par rapport aux prélèvements de sang effectués sans garrot.

Méthode: Une étude de cohorte, prospective a été menée chez des patients traités au service des urgences dont le tableau clinique justifiait, selon le médecin, une mesure du taux de lactate. Les malades ou leur représentant ont donné par écrit leur consentement éclairé. Les prélèvements de sang faits pour la mesure du taux de lactate aux fins de l’étude ont été effectués avec garrot, en même temps que ceux faits pour d’autres examens de laboratoire. Les prélèvements de sang faits pour la mesure du taux de lactate aux fins cliniques ont été effectués sans garrot, selon le protocole en vigueur à l’hôpital. Ont été consignées l’heure de la mesure des taux de lactate ainsi que des données démographiques sur les patients. Il y a eu comparaison des taux de lactate de chaque patient selon le test de Wilcoxon.

Résultats: Quarante patients ont participé à l’étude après avoir donné leur consentement. Le taux médian de lactate, mesuré aux fins cliniques était de 1,9 (intervalle interquartile : 1,5-2,6) et le taux médian de lactate, mesuré aux fins de l’étude était de 1,9 (intervalle interquartile : 1,4-2,7). Aucun écart n’a donc été enregistré entre les valeurs appariées de lactate (p = 0,95).

Conclusions: Il semble donc que l’utilisation du garrot n’ait aucune incidence sur la mesure du taux de lactate. Les résultats de l’étude portent à croire que les pratiques en vigueur dans bon nombre d’établissements en ce qui concerne les prélèvements de sang pour la mesure du taux de lactate sont trop restrictives et qu’il faudrait les changer.

Keywords: blood lactate, tourniquet use, phlebotomy

INTRODUCTION

Lactate was first demonstrated to be present in the blood of acutely ill septic women in 1843.1,2 In the 1960s, lactate was noted to be a predictor of poor outcomes in hospitalized patients.3 Additional research has shown blood lactate to be one of the most useful...
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altered levels compared to the result of a level drawn
during the drawing of a lactate results in signi
determine in clinical practice whether tourniquet use
not been studied. The purpose of this study was to
tourniquet use on lactate levels in clinical practice has
be hypothesized that application of a tourniquet to
quets for hemostasis, lactate levels have been found to
increase; however, these studies also used tourniquet
times far in excess of what is applied for a blood
draw.

As a result of these studies as well as due to the
theoretical reasoning for tourniquet avoidance, many
hospitals have protocols in place indicating that lactate
levels must be drawn without a tourniquet. Some
manufacturers of lactate assays as well as the Surviving
Sepsis Campaign Bundle also recommend drawing
lactates without tourniquets. Phlebotomy without
the use of a tourniquet is technically more difficult as a
result of blood vessels’ smaller diameter and reduced
turgor. This increase in difficulty may lead to failure
to obtain a lactate level or a delay in the time necessary
to obtain this important test.

In healthy volunteers, tourniquet use has not been
found to increase blood lactate significantly with
tourniquet times up to 15 minutes, or with blood
pressure cuffs elevated to mean arterial pressure for a
duration of five minutes. However, the effect of
tourniquet use on lactate levels in clinical practice has
not been studied. The purpose of this study was to
determine in clinical practice whether tourniquet use
during the drawing of a lactate results in significantly
altered levels compared to the result of a level drawn
without a tourniquet.

MATERIALS AND METHODS

Study design

A prospective cohort study was undertaken of patients
presenting to the emergency department (ED) for
whom physicians ordered a lactate level as part of
clinical care. The reasons for physicians drawing clinical
lactates included evaluation of suspected sepsis or
ischemic bowel. Blood samples from these patients were
utilized for both the clinical lactate and the study
lactate. Participation in this study resulted in no addi-
tional cost, change in standard care, harm, or benefit to
the patients enrolled, as the attending physicians did
not have access to study results.

Study setting and population

This study was carried out at St. Luke’s University
Hospital in Bethlehem, Pennsylvania, a level 1 commu-
nity trauma center with an ED census of 60,000. A
convenience sample of adult patients (≥18 years) were
screened for eligibility over a one year period in
2013-2014, based on availability of research associates
to complete the informed consent process with the patients.
Patients were deemed eligible if the attending physician
ordered a lactate during their clinical care. Trauma
patients were excluded. Participants were enrolled after
written informed consent was obtained from the patient
or health care proxy. The study was approved by the
institutional review board at the study hospital.

Study protocol and measurements

Lactate levels for clinical use were drawn as per
standard protocol at the study hospital with no
tourniquet used during collection. Study lactates were
obtained using a tourniquet during the draw sequence
of other laboratory studies ordered at the discretion of
the treating provider. If consent was obtained prior to
establishment of intravenous access, the study lactate
was typically drawn last in the draw sequence of other
blood studies during intravenous catheter placement. If
the consent was obtained after blood studies had already
been drawn, the study lactate was typically drawn
during a second venipuncture for a second blood
culture. There were no circumstances in which the
patient underwent a blood draw solely for the purpose
of obtaining a study lactate.

Clinical lactate levels were reported in the electronic
laboratory result system as per hospital standard and
were available to physicians while patient care was
ongoing. Study lactate levels were stripped of patient
identifiers and coded for blinding purposes. Results of
these assays were mailed to the research investigators as
private correspondence and were not reported in the
medical record or used for patient care.
The hospital lactate assay was performed utilizing a Flex Reagent Cartridge in a Dimension Clinical Chemistry System analyzer with an analytical sensitivity of <0.3 mmol/L (Siemens Healthcare Diagnostics, NY).

The times of clinical lactate and study lactate draw were recorded, as were patient age and gender. These were entered into a standardized Microsoft Excel 2007 spreadsheet (Microsoft Corporation, Redmond, WA) by trained research associates.

**Data analysis**

Because the clinical and study lactates represent repeated measures in single patients, the two values were compared with Wilcoxon Signed-Rank test. This statistic was used because it was expected that the two lactates would be drawn by two methods at close to the same time. We did not use a T-test, because we had an a priori expectation that the lactate levels would display skew, with tail toward higher values clustering around the normal range. Linear correlation was used to assess overall relationship between clinical and study lactate. Demographic information was reported with standard descriptive statistics. Data were analyzed using VassarStats: Website for Statistical Computation (vassarstats.net, author Richard Lowry, PhD, Professor of Psychology Emeritus, Vassar College, Poughkeepsie, NY, © 1998-2013).

**RESULTS**

Forty patients were enrolled over the one year study period. The mean age of study participants was 72.7 years (standard of deviation 16.4 years), and 62.5% were male. Enrolled patients were typically ill and undergoing evaluation for infectious processes (sepsis) or intra-abdominal surgical disease (mesenteric ischemia or other surgical disease).

Clinical lactate and study lactate were drawn a median of six minutes apart (IQR 3-20 minutes, range 0-151 minutes). The lactate reference range for the laboratory performing the study was 0.5-2.2 mmol/L. For the patients enrolled in this study, the median clinical lactate level was 1.9 mmol/L (IQR 1.5-2.6, range 0.7-5.6). The median study lactate level was 1.9 mmol/L (IQR 1.4-2.7, range 0.8-5.7). 37.5% of clinical lactate levels fell above the reference standard (n = 15).

Using a two-tailed Wilcoxon Rank-Sum test, there was no statistical difference found between clinical and study lactate levels (p = 0.95, Figure 1). Considering only the 15 patients with clinical lactate levels above the reference standard, there were no differences in clinical and study lactate levels (p = 0.96). On simple linear correlation, the correlation coefficient between the clinical and study lactate was 0.97 (CI 0.94-0.98) with an r-square of 0.93 (Figure 2).

**DISCUSSION**

Accurate and timely measurement of lactate is important. Our findings suggest that current practices at many institutions regarding lactate collection are likely too stringent and should be changed. These practices appear to have been put in place based on extrapolation from
orthopedic and anesthesia research; however, studies of that nature involved higher tourniquet pressures and longer periods of time than that used in routine phlebotomy.

In routine phlebotomy, the standard of care is to apply a tourniquet that is tight enough to cause target vein dilation (i.e., tight enough to occlude venous flow), but not so tight as to cause arterial obstruction. If increasing pressures are responsible for increasing lactate in tourniquet patients, as some have suggested, it may be that the disposable tourniquets used in phlebotomy simply do not generate enough pressure to elevate lactate. If this is the case, situations in which phlebotomy is performed using a blood pressure cuff for target vein dilation may result in different blood lactate results depending on the level to which the cuff is inflated. Prior surgical studies generally involved increasing the tourniquet pressure to above the mean arterial pressure (MAP) in order to ensure a bloodless field during extremity surgery. These results may indicate that blood pressure cuff increase to above the MAP could also falsely increase lactate values in clinical practice; however, this has not been adequately studied.

In addition to lower pressure, phlebotomy tourniquets are usually applied for less time than those involved in surgical studies. In standard phlebotomy practice, tourniquet up-times are usually less than a minute, and the World Health Organization indicates a tourniquet up-time of two minutes or less is best practice. We did not record tourniquet up-time, but suspect that, since the study lactate was always the last blood drawn (so as not to interfere with necessary studies for patient management), study values represent lactates taken at the upper limit of tourniquet time. In a prior study of healthy volunteers, even 15 minutes of tourniquet up-time with a phlebotomy tourniquet did not alter lactate levels. Our study was not designed to find a specific tourniquet time beyond which blood lactate might be falsely elevated, but within the constraints of our methodology we found no evidence that lactate was affected by routine tourniquet application.

Our patient population was medically ill, and typically being evaluated for sepsis. We did not enroll trauma patients, although lactate levels are often used to guide resuscitative efforts during trauma. Elevated lactate levels in both trauma and sepsis are secondary to perfusion mismatch. As a result, we suspect that phlebotomy tourniquet use would not falsely elevate lactate levels in such patients, but our study was not adequate to determine impact of phlebotomy tourniquet use in all patients in whom a lactate might be obtained.

LIMITATIONS

Although our patients were all patients for whom a physician deemed a lactate level would be clinically useful, there are limitations introduced in any sample of this nature. Patients needed to be conscious and competent in order to consent to participate, or alternatively have a valid power of attorney accompanying them. Very ill patients are not always awake and alert, and clinicians do not usually approach the critically ill for research purposes due to the importance of timely resuscitation and treatment. As a result, the patients in whom this study might be most applicable (i.e., extremely ill septic patients) may not have been adequately represented in our study population. The population from which the lactate levels were measured was primarily made up of those who were not in extremis, but who were patients for whom it was judged by the attending physician involved that a lactate would be useful.

In addition, we did not attempt to standardize or record tourniquet up-time in study patients. It is possible that for some patients, for whom intravenous access was difficult to obtain, there were more prolonged tourniquet times in order to increase the success of phlebotomy. Since we did not standardize or record these times, we cannot evaluate the impact of time of tourniquet use on lactate levels.

We did not capture the clinical scenarios involved in the decision by treating physicians to order a lactate level. There may be differences in tourniquet effect based upon underlying patient comorbidities or the cause of an elevated lactate (i.e., “Type A,” secondary to perfusion mismatch, versus “Type B,” secondary to inability to clear the lactate).

Finally, our study was performed at a single institution with a small number of participants arising from convenience sampling, and thus may not be generalizable to other settings and populations. As a result of our methods, a selection bias for less sick patients existed, as our study could not result in delayed care for critically ill patients.

CONCLUSIONS

Tourniquet use appears to have no impact on measured lactate levels. Our findings suggest that current practices
at many institutions regarding lactate collection are likely too stringent and should be changed.

**Competing Interests:** None declared.

**REFERENCES**


