INTRODUCTION

Previously results of analysis were dependent on analytical variation, but nowadays control and standardization of pre-analytical variability is a critical factor for achieving accuracy and precision in laboratory testing because about 32% to 75% laboratory errors are made in pre-analytical phase and majority of them are preventable. Venous blood sampling is usually performed using a tourniquet to help locate and define peripheral veins to achieve successful and safe venipuncture. During the last two decades significant improvements in blood sampling technique and equipments e.g. positive patient identification, vacuum tubes for blood sampling, improved needles for phlebotomy and operator safety are made, it is very important to give proper attention and careful supervision to prevent errors during phlebotomy procedure. Tourniquet is widely used for venipuncture by medical and laboratory staff but very few are aware of effect of tourniquet application on laboratory parameters. In addition definitive guide lines regarding how and when to use tourniquet for blood sampling are lacking. Tourniquet application should be minimized because it causes spurious and significant variation of several plasma analytes. For example prolonged venous stasis can cause 7% increase in prevalence of hypercholesterolemia and thus unnecessary medication can be introduced. Prolonged tourniquet application can lead to local hypoxia and thus acidosis, which can affect potassium measurement. With venous stasis, protein and protein bound constituents can increase upto 15%. Evidence from study in Verona, Italy have shown that by measuring plasma concentration of twelve common analytes, collected without and with stasis for one minute have clinically significant elevation for albumin (44.0±3.0, 45.5±3.4) mmol/L, calcium (2.32±0.13, 2.36±0.10) mmol/L, total cholesterol (5.07±0.83, 5.22±0.83) mmol/L and decrease in potassium (4.21±0.34, 4.09±0.27) mmol/L.

But other study indicates that blood gases, hematological parameters and serum electrolytes are not affected by application and removal of a tourniquet. So there are conflicting results available regarding the effect of venous stasis by tourniquet application and also there is no such study in this region. Therefore better of the two techniques will be
used in future.

**MATERIALS AND METHODS**

**Study Design**
The study was a Quasi experimental study.

**Setting**
The study was carried out in Pathology Laboratory, Quaid-e-Azam Medical College, Bahawalpur.

**Duration of Study**
From October 2011 to April 2012.

**Sampling Technique**
Non – probability, consecutive sampling.

**SAMPLE COLLECTION**

**Inclusion Criteria**
Blood samples were collected from all persons irrespective of age and gender who were willing to give 10ml of blood, visiting pathology laboratory, Quaid-e-Azam Medical College, Bahawalpur.

**Exclusion Criteria**
- Persons who refused to give 10ml of blood.
- Patients of disseminated intravascular coagulation.
- Subjects who were taking steroids and heparin for more than 4 weeks.
- Haemolysed and lipemic sample.

**DATA COLLECTION PROCEDURE**
The study was started after taking approval from Ethical committee of Bahawal Victoria Hospital. Written and informed consent was taken from all individuals who met the inclusion criteria. In persons with odd serial number, right arm was used as control and left arm as test arm, whereas, in person with even serial number right arm was used as test arm and left arm as control. The collection of diagnostic blood samples was accomplished by a single expert phlebotomist from median cubital vein in antecubital fossa after being seated for 15 minutes to eliminate the effects of standing posture with one difference that sample was withdrawn from the test arm after application of standardized external pressure of 60 mm Hg using a sphygmomanometer for one minute. The needles and vacuum tubes were of same lot.

All blood specimens were drawn by using a 20 G straight sterile needle directly into vacuum tubes with clot activator and acrylic gel separator.

**PROCESSING OF BLOOD SAMPLES**
All the tubes were left in upright position for 45 minutes at room temperature to allow complete blood clotting before centrifugation. Serum was separated after centrifugation at 1500 g for 10 minutes. No blood sample was discarded on the basis of unsatisfactory attempts, difficulty in locating venous access, missing veins and manifest haemolysis.

**LABORATORY TESTING**
The concentration of serum sodium, potassium, chloride was estimated on Ion Selective Electrode. Total serum protein and serum cholesterol and calcium were measured on fully automated chemistry analyzer (Selectra XL), by medical technologist with an experience of 5 years, according to the manufacturer's specifications and using proprietary reagents. Measurement bias was controlled by calibration of instruments and repeating each test twice within a single analytical session. The results were reported as the mean of paired measurements.

**DATA ANALYSIS**
Data was analyzed using statistical package for social sciences (SPSS). Frequency and percentage were calculated for sex while mean and standard deviation were calculated for sodium, potassium, chloride, calcium, protein and cholesterol. The mean change for these analytes was also calculated. T-test was used to compare the means and P value < 0.05 was taken as significant. Results were presented in the form of
In this study, blood samples were taken from 265 subjects and about half of the participants were female of different age group. The results of the study are shown in tables II.

The results of the study showed that the serum potassium, calcium, protein and cholesterol showed statistically significant difference in their mean concentration values while serum sodium and chloride measurements showed insignificant effect. The mean increase in serum potassium was 0.37 mmol/L (\(p \leq 0.0001\)), calcium showed an increase of 0.05 mmol/L (\(p=0.0064\)), total protein had increase of about 2.7 g/L (\(p=0.001\)) and there was increase of 0.25 mmol/L (\(p=0.02\)) in the measurement of cholesterol.

**DISCUSSION**

The clinical laboratory results are an essential part of the health care delivery. It has been estimated that 60 up to 70% of medical decisions and procedures, such as drug prescriptions, assessments prior to and in the course of further investigations or dialysis, are strongly dependent upon laboratory data.

The total testing process (TTP) is the total process from the ordering of a test to the interpretation of a test result. The TTP starts and ends with the patient, and can be subdivided into three distinctive phases: the pre-analytical step (before the analysis), the analytical step (the actual analysis) and the post-analytical step (after the analysis), as described in Figure 1.

Errors can occur in every step of the TTP. Of all errors in the TTP, approximately one fourth have consequences for the patient. These consequences include a delayed test result or new sample collection which result in increased workload on lab and financial burden on patient, but also life threatening and tragic consequences, such as unnecessary chemotherapy and coma.
It is evident that the majority of all errors in the TTP are of pre analytical in origin i.e. they occur before the sample arrives in the laboratory\(^2,9\). Previous reports suggested that about 46–68% of errors were made in preanalytical phase but advances in automation and instrument technology have improved the quality of test results and it is now stated that up to 93% of all errors are made in preanalytical phases\(^11\). Therefore, reduction of preanalytical errors is an important issue for everyone involved in the TTP.

Majority of the preanalytical errors are due to human mistakes\(^12\) and they are preventable in most of the cases\(^2,12\). The tourniquet application is an important controllable variable of pre analytical factor that can affect laboratory results. It is usual practice that tourniquet remains in place during blood collection so that continued venous dilation allows rapid sample collection which may result in its prolonged application. This application time is rarely regarded as a potential source of laboratory variability that may be lengthen due to difficult location of vein, selection of proper blood collection system, insertion of needle into the vein and collection of many tubes\(^13\).

The previous studies conducted on tourniquet application showed conflicting results. The study conducted by Broome et al on electrolytes showed no statistically significant difference when tourniquet was applied for two minutes\(^14\).

McMullan and his coworkers concluded that calcium and protein measurements showed negligible changes with one minute of tourniquet application\(^15\).

The results of other studies indicated that the mean change in concentration of analytes were not significant by the use of tourniquet\(^16,17\). However Renee et al showed an increase of about 6% in serum concentration of protein and 2.5% in calcium after one minute of tourniquet application\(^18\).

Venous stasis facilitates the exit of water and diffusible ions from the vessels. The resulting hemoconcentration raises the concentration of numerous blood analytes at the punctured site thus influencing the laboratory parameters interpretation.

The results of Lippi et al\(^1\) were consistent to this study and showed a mean increase in concentration of calcium was 0.04 mmol/L and mean increase of 0.15mmol/L in case of cholesterol with one difference that mean concentration of potassium showed decrease of 0.11mmol/L.

Another study that was performed by Oliveria\(^19\) showed mean increase of about 2g/L in case of protein, 0.03mmol/L in serum calcium level and 0.3mmol/L in serum potassium measurement that were similar to the results of this study. The possible mechanism may be that when vascular environment is subjected to hypoxia and stasis, local acidosis releases potassium from the cells that results in elevated potassium level.

Sonoli\(^20\) performed study on five routinely advised analytes and showed a mean increase of 0.18g/dl (\(p=0.006\)) in protein measurement after one minute.
of tourniquet use. Differences between the results may be due to difference in technique employed to apply tourniquet, biochemical characteristics of the measuring analyte. The studies were also conducted on different population with different instruments.

CONCLUSIONS
The venous stasis produced by tourniquet application of one minute can influence laboratory parameters. So every effort should be made to reduce its application time and medical and laboratory staff should be properly educated about the usage of tourniquet.

REFERENCES


“Don't find fault, find a remedy.”

Henry Ford