Original Article

Identification of types and frequency of pre-analytical errors in hematology laboratory at a tertiary hospital of Nepal

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**Keywords:**
Hematology; Preanalytical error; Quality control; Samples;

**ABSTRACT**

**Background:** In the laboratory, errors can occur at any stage of sample processing; pre-analytical, analytical, and post-analytical. Since the pre-analytical phase is the most common source of laboratory errors, the goal of this study is to identify the types and frequency of pre-analytical errors in the hematology laboratory.

**Materials and Methods:** This is a cross-sectional descriptive study done at Nepal Medical College Teaching hospital for a duration of nine months (January 2020 to September 2020). All blood samples received at the hematology laboratory were included whereas biochemistry and special tests blood samples were excluded. Samples were checked for misidentification (incorrectly labeled vials/vials without labels/incorrectly filled forms), incorrect samples (wrong choice of vials), clotted samples, inadequate samples, diluted samples, hemolyzed samples. The errors that occurred in these samples (both inpatient and outpatient) were noted down and measures were taken accordingly before analyzing the sample.

**Results:** The total number of samples received was 15,337. Pre-analytical errors were seen in 857 samples (5.5%). Inadequate samples (25%) were the most common error followed by incorrect samples (20%), hemolyzed samples (20%), misidentification (14%), clotted samples (12%), and diluted samples (9%). Complete blood count test was most affected. Samples from the inpatient department were most affected.

**Conclusion:** The preanalytical error rate in the hematology unit was 5.5% with an inadequate sample being the commonest error. Most of the errors were seen in the test requested for a complete blood count. Samples from the inpatient department showed the most errors.

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**INTRODUCTION**

The clinical laboratory is a rapidly changing field that has a very strong impact directly on patient’s treatment. Even with the advancement and automation in the clinical laboratory, the zero error rate in the quality report has not been achieved to date. Quality assurance in the laboratory is a must to ensure laboratory users of reliable test results with a high degree of precision and accuracy.¹ Total quality in laboratory medicine should be defined as the guarantee that each activity throughout the total testing process is correctly performed, providing valuable medical decision-making and effective patient care.²
Laboratory errors can often have serious adverse consequences. Lack of standardized procedures for sample collection accounts for most of the errors encountered within the total testing process. They can also have clinical consequences as well as a significant impact on patient care, especially those related to specialized tests as these are often considered as "diagnostic". In the laboratory, errors can occur at any stage of sample processing: pre-analytical, analytical, and post-analytical stages. However, evidence shows most of the laboratory errors occur during the pre-analytical stage. A pre-analytical error is defined as a rejected specimen (blood or urine sample), which cannot be successfully tested as it does not meet the acceptability criteria of the laboratory or if the sample is not received. The receipt and processing of specimens are one of the main steps in the pre-analytical stage. Errors in this stage could be due to mislabeling, incorrect test entry, and entering the wrong location. Most of these errors are preventable.

The pre-analytical phase is the stage of greatest risk and is most vulnerable but pre-analytical errors may go undetected until post-analytical validation and interpretation which impacts the good quality report of a laboratory. Recently the test performances have improved and new parameters have been introduced, as well as internal and external quality controls have been used for the monitoring of accuracy to minimize the error rates.

The quality of the report will not be accepted until the laboratory minimizes the pre and post-analytical errors. Proper and timely recognition of these loopholes in quality control will help lead to the correct therapeutic strategy and better patient care. Since pre-analytical errors are not the sole responsibility of the lab and its staff, the management and other staff involved with blood collection should be aware of this problem to minimize the pre-analytical errors hence helping in generating a quality report. Since the pre-analytical phase is the most common source of laboratory errors, the goal of this descriptive study is to identify the types and frequency of pre-analytical errors at Nepal Medical College, Teaching hospital.

MATERIALS AND METHODS

This is a cross-sectional descriptive study done at Nepal Medical College, Teaching Hospital (NMCTH) for a duration of nine months (January 2020 to September 2020). All blood samples received at the hematology unit of the laboratory (samples for complete blood count, coagulation studies, peripheral blood smear, and malarial parasites) during the study period were included whereas biochemistry and special tests blood samples were excluded. Before submitting them to the respective unit, they were checked for mismatching which included: misidentification (incorrect labeling of vials/ vials without labels/ incorrectly filled forms), incorrect samples (wrong choice of vials), clotted samples, inadequate samples, diluted samples, and hemolyzed samples. The types of error, the place from where the sample was sent (outpatient or inpatient) as well as in which test samples the error has occurred was noted down and measures were taken accordingly before analyzing the samples in automated and semi-automated machines. The data thus obtained was entered in Excel Microsoft and analyzed using SPSS version 16.0. Ethical clearance was taken from the Institutional Review Committee of NMCTH.

RESULTS

The total number of samples received in the hematology unit was 15,337 samples in nine months duration. Pre-analytical errors were seen in 857 samples. Hence, the frequency of pre-analytical errors was 5.5%. Out of 15,337 samples, error-free samples were 14,480 (94.5%). The different types of errors identified in the hematology unit are tabulated in table 1. Out of total errors, inadequate samples (219 samples; 25%) were the most common followed by incorrect samples and hemolyzed samples (169 and 172 samples; 20% each). The less common errors in descending order were misidentification (117 samples; 14%), clotted samples (105 samples; 12%), and the least common was diluted samples (75 samples; 9%). Table 2 shows different sources of errors. The samples for complete blood count accounted for most of the sources of errors (n=553; 64%) followed by coagulation studies (n=179; 21%), peripheral blood smear (n=111; 13%) and test for malarial parasite in blood smear (n=14; 2%).

The rejected samples were traced whether they were from the in-patient department (IPD; ward samples) or out-patient department (OPD). Out of 15,337 samples, most of the samples were from OPD (8351 samples; 54%) followed by IPD (6986 samples; 46%). It was found out that out of 857 rejected samples, 809 samples (94%) were from IPD whereas 48 samples (6%) were from OPD. The inadequate sample was the most common error in both IPD as well as OPD. Among IPD, hemolyzed samples (n=168, 2.40%), incorrect sample (n=158, 2.26%), clotted sample (n=103, 1.47%), misidentification (n=102, 1.46%) and diluted samples (n=75, 1.07%) were the samples with error. Among OPD cases, errors were seen as misidentification (n=15, 0.17%), incorrect samples (n=11, 0.13%), hemolyzed samples (n=4, 0.04%) and clotted samples (n=2, 0.02%).

<table>
<thead>
<tr>
<th>Table 1: Types of errors in the hematology laboratory</th>
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<td>Types of error</td>
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<tr>
<td>Misidentification</td>
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<td>Incorrect samples</td>
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<td>Inadequate samples</td>
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<td>Clotted samples</td>
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<tr>
<td>Diluted samples</td>
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<td>Hemolyzed samples</td>
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Total number of samples: 15,337
Total number of samples with error: 857 (5.5%)
Pre-analytical errors in hematology laboratory

<table>
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<th>Table 2: Different sources and percentage of each pre-analytical error</th>
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<td>Sources of error</td>
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<tr>
<td>Complete blood count</td>
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<td>Coagulation test</td>
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<td>Peripheral blood smear</td>
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<td>Malarial parasite test in blood smear</td>
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<td>Total</td>
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Diluted samples were not received in OPD samples. The distribution of pre-analytical errors among IPD and OPD is tabulated in table 3.

**DISCUSSION**

The increase in the prevalence of medical errors represents a disturbing trend; hospital-based errors are the eighth leading cause of death in the United States. Clinical laboratories have long focused their attention on quality control methods and quality assessment programs dealing with analytical aspects of testing. The phases before the sample reaches the laboratory (preanalytical) and the phase after the sample is analyzed (post-analytical) are equally important. However, the preanalytical phase is challenged with many shortcomings like improper filling up of request forms with illegible handwriting, improper blood collection by the staff, and improper mixing up of blood with anticoagulant, etc. The total number of samples received in hematology was 15,337 during nine months duration. Out of these samples, 857 samples were rejected with an error rate of 5.5% which is higher than other studies. Following are different rates of error found in different studies: Sakyi et al (4.7%), Chawla et al (1.9%), Uperti et al (1%), Arul et al (0.43%), and Rajalakshmi et al (0.3%). The higher rate of an error occurring in the present study may be because only the hematology unit of whole laboratory samples was taken into account in this study. Since the error rate is significantly higher than the error rates of other studies, there seems a need to educate the staff dealing with phlebotomy. A similar study by HarsimranKaur et al concluded that preanalytical errors were frequent in laboratories and can be corrected by regular analysis of the variables involved. Rectification of these types of errors could be done by regular education of the staff.

Out of total errors, an inadequate sample was the most common error followed by incorrect samples and hemolyzed samples. The inadequate sample was most often error even in a study by Singla et al and Arul et al. Appropriate knowledge regarding the important role of proper labeling of the samples should be made known to all the staff involved in the collection as it is the most common form of error seen in this study. Since samples from the hematology unit were only taken into account, most of the errors were found out to be occurring in the samples collected for complete blood counts followed by coagulation studies.

The present study showed a vast difference between the error rates in between samples from OPD and IPD. Most of the errors occurred from samples collected from IPD (94%) which is significantly higher than the samples collected from the OPD. This variation may be due to the collection of blood samples by laboratory technicians if the samples were collected from OPD whereas the inpatient samples were collected by the ward staffs hence, there seems a dire need to arrange for the workshop and training program for all those involved in sample collection to reduce the preanalytical errors which are completely human dependent. The main areas of training should be focused on phlebotomy methods as well as about the adequacy of the sample, how to not let the sample be hemolyzed, and proper filling of the requisition forms. This trend of higher error rates among the samples of inpatient departments is seen in other studies as well. Kadic et al stated that the proportion of inpatient rejected samples was 8.7 fold higher than in the outpatient samples.

Rajalakshmi et al stated that adequate training, regular maintenance of records of errors, and periodic auditing will result in effective reduction of such errors with improvement in the overall performance of laboratory works. Ying et al emphasized applying a training system between various departments whereas Lippi et al gave much importance to standardization and monitoring preanalytical variables which will be associated with the most efficient and well-organized laboratories resulting in reduced operational costs and increased revenues. Tadesse et al, Wiwanikit et al, Plebani et al, and Boon et al gave importance to the close communication between clinicians and laboratory personnel to improve laboratory quality in general. Hence, a training session and interdepartmental communication should be organized to reduce the error rates.

**CONCLUSIONS**

The preanalytical error rate in the hematology unit was 5.5% with an inadequate sample being the commonest error. Most of the errors were seen in the test requested for a complete blood count. The error rate was very high in
samples collected from IPD. To minimize this error, training session and interdepartmental communication is a need of time. Since laboratory work is a product of teamwork, the main areas of training should be focused on phlebotomy methods as well as about the adequacy of the sample, how to not let the sample be hemolyzed, and proper filling of the requisition forms.

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Conflict of interest: None

REFERENCES


