Guide to Fulfillment of Laboratory Results Comparability Requirements

Purpose

This document from the Diagnostic Accreditation Program (DAP) provides guidance for organizations on fulfilling the Laboratory Medicine Accreditation Standards.

If you have questions about items in the standards, please email them to laboratorymedicine@cpsbc.ca.

Terms and definitions

**comparability**

Agreement between patient results obtained for a measurand using different measurement procedures within a health care system.

**Note:** Results are considered to be comparable if the differences do not exceed a critical value established based on defined acceptance criteria.

**corrective action**

Action taken to eliminate the cause(s) of nonconformities.

**Note:** Corrective action is usually taken in response to nonconformity or an identified problem noted in audits or during the accreditation process.

**examination**

A set of operations having the object of determining the value or characteristics of a property.

Synonyms: analysis, assessment, investigation, measurement, study, test

**measurand**

Quantity intended to be measured.

Synonym: analyte

**procedure**

A documented, specified way to carry out an activity of a process.

**Note:** For the purposes of accreditation, procedures must always be documented, implemented and maintained.

**process**

A set of interrelated or interacting activities which transform inputs into outputs.
primary analyzer The identified method in a facility to which others will be compared for a particular examination. CLSI

sample A discrete portion of a body fluid, breath, hair or tissue taken for examination of one or more quantities or properties assumed to apply for the whole. One or more parts taken from a primary sample. ISO

Synonyms: specimen, primary sample, aliquot

user (of laboratory services) Physicians and others who order diagnostic examinations and/or receive diagnostic information and reports from laboratories. DAP

Synonyms: authorized requestor, ordering physician, clinical personnel

Introduction

The DAP accreditation process assesses the comparability of results at a regional or organizational level and at the individual facility level. Standards addressing comparability of laboratory results within a single facility in BC have been in place for many years. Incorporation of ISO 15189 requirements into the DAP laboratory medicine standards requires health authorities and organizations to ensure comparability of laboratory results across all sites within their jurisdiction. All laboratories must demonstrate comparability of results across their region or organization and within their individual facilities.

Objectives

The objectives of this document are to

- familiarize assessors with the concept of laboratory results comparability,
- introduce the approaches laboratories can use to ensure results comparability, and
- describe the assessment of comparability of laboratory results.

What is comparability of laboratory results?

The comparability of results provides assurance that the examination of a measurand is consistent within a laboratory system, even if different methods and instruments are used. There is no consensus procedure for demonstrating the comparability of laboratory results across different methodology and instrumentation. Laboratories must establish their own procedures to ensure the comparability of results in their individual system. This includes the frequency of comparability testing (at minimum every six months), the criteria for acceptability, consideration of the number of samples used for comparability validation, consideration of the range of clinical values, and the response for non-comparability of results. The procedures to ensure comparability of results must also specify the samples to be used. Samples used for verification of comparability can be individual patient samples, pooled patient samples, laboratory prepared material, proficiency testing material and QC material.

Summary reports prepared by proficiency testing vendors cannot be used for comparability testing. Proficiency testing data may be used if the laboratory has developed a mechanism to demonstrate comparability.
Manufacturer’s QC material can be used to demonstrate comparability of identical instrumentation and methodology. In instances where systems differ in instrumentation and/or methodology, QC material can be used to demonstrate comparability if:

- the numeric relationship between the means for QC results from the different measuring systems is known
- the result for native clinical samples have been verified to be comparable
- the performance of the measuring systems remains unchanged
- the above remain comparable following lot changes

It may be more practical to perform comparability monitoring using native patient samples than to conduct the validation required for QC results.

**Why must laboratories demonstrate comparability of results?**

Patients are often diagnosed and treated in multiple health care settings within a regional laboratory system. Within that laboratory system, there may be many instruments performing the same test using different methodology. Larger laboratories may have multiple instruments within one location, such as a primary analyzer, a backup analyzer, and point-of-care testing equipment. Even in small laboratories, the same examination will be performed in the laboratory using conventional instruments and near point-of-care testing equipment. These results may be reviewed by multiple clinicians at multiple patient care locations within a single laboratory system.

The diagnostic value of multiple laboratory results in multiple sites is optimized when there is agreement of those results. Conversely, multiple laboratory results from laboratories where comparability has not been achieved can result in confusion and distrust, and potentially compromise patient care. Clinicians should not have to consider when, where or how the examination was performed when interpreting results.

Non-comparability of laboratory results may occur because of the following:

- different methodologies
- differences in calibration or imprecision between procedures
- value assignment errors and variation of commutability between calibrator lots
- age/stage difference and time-dependent degradation of calibrators used simultaneously in different locations
- differences in commutability of calibrators with different procedures
- reagent degradation after calibration
- instrument drift/failure
- use of different reagent lots
- variation in shipping or storage conditions
- differences in instrument parameters (e.g. dilution rates, incubation times)
Laboratories may use a variety of processes to demonstrate comparability. In the above model, the organization consists of three sites, each having a primary analyzer, a backup analyzer and point-of-care testing. The laboratory must demonstrate comparability of results across the organization, and this will be assessed at both the regional assessment and at the individual facility assessments.

**Regional assessment of comparability**

At the regional assessment, laboratories must demonstrate comparability from site to site. This is most easily done by comparing Primary A to Primary B to Primary C.

**Facility assessment of comparability**

At the facility assessment, laboratories must demonstrate comparability within an individual site. This is usually done by comparing the primary analyzer to the backup analyzer to the point-of-care testing (e.g. Primary A to Backup A to POCT A).
Overall assessment of comparability

If the laboratory demonstrates site-to-site comparability and comparability within each site, then there should be comparability between all examinations at all sites within the laboratory system (e.g. Primary A = Backup B = POCT C).

DAP comparability requirements

There are six criteria used to evaluate the laboratory’s verification of the comparability of patient results within the region or organization. They are as follows:

<table>
<thead>
<tr>
<th>QUA3.1.1 M</th>
<th>There are procedures to establish the comparability of procedures, equipment and methods used, and establishing the comparability of results for patient samples. This is applicable to the same or different procedures, equipment, different sites or all of these.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interpretation:</td>
<td>A laboratory system must ensure comparability of results throughout its operation. This includes site-to-site comparability within a region or organization as well as comparability within a facility.</td>
</tr>
<tr>
<td>Evidence:</td>
<td>Specific procedures addressing comparability of results. The procedures should address regional and facility comparability.</td>
</tr>
<tr>
<td>Where:</td>
<td>Regional/organizational assessments</td>
</tr>
</tbody>
</table>
| QUA3.1.2 | M | Comparability encompasses the entire range of clinically relevant values.  

*Guidance: Freshly obtained patient samples are the preferred comparability material as commercial materials have had their matrices modified in ways that may significantly affect commutability with native clinical samples.*  

**Interpretation:** Laboratories must establish the types and numbers of samples used at which clinically significant points through the reportable range of the measurand.

**Evidence:** Specific records of the samples and results used to demonstrate comparability for each measurand.

**Where:** Regional/organizational and facility assessments

**Comment:** Ideally, all values throughout the reportable range would be checked during comparability testing, but that is often not possible. The laboratory must establish the number and value of points checked during comparability testing. The range of values tested will be specific to individual examinations with a range equivalent to the analytical measurement range or reference interval and include all clinical decision points.  

The preferred samples for comparability verification are individual patient samples. In the event that patient samples are unavailable, the following material can be used for comparability studies in order of preference: pooled patient samples, laboratory prepared (spiked) material, proficiency testing material and QC material.

| QUA3.1.3 | M | Acceptability criteria are defined for comparability of procedures, equipment and methods.  

**Interpretation:** Laboratories must establish the tolerance limits for comparability between sites and between instruments. See below.

**Evidence:** Specific records of the samples and results used to demonstrate comparability for each measurand.

**Where:** Regional/organizational and facility assessments

**Comment:** Laboratories must determine the limits of acceptable variability for results produced by different methods and analyzers for the same measurand. Acceptability criteria may need to be defined for each measurand. Evaluation of comparability may be based on published professional recommendations, based on goals set by external bodies or based on laboratory generated data.
| QUA3.1.4 | M | The laboratory documents, records and acts upon results from comparability. Any identified problems or deficiencies are addressed and records of action are retained.  
**Interpretation:** Laboratories should have documentation that demonstrates they have verified the comparability of results and have recorded any activity in response to any issues of comparability.  
**Evidence:** Specific records of the results of comparability for each measurand as well as any resulting activity.  
**Where:** Regional/organizational and facility assessments |
|---|---|---|
| QUA3.1.5 | M | The laboratory notifies users of any differences in comparability of results and discusses any implications for clinical practice when examination systems provide different reference intervals for the same measurand and when examination methods are changed.  
**Interpretation:** Laboratories should have a process to notify users of any differences in comparability of results.  
**Evidence:** Policy for notification of users in the event of differences in comparability of results and when available specific records of user notification of differences in results of comparability.  
**Where:** Regional/organizational and facility assessments |
QUA3.1.6 M Instruments and methods are checked against each other at least twice a year for comparability of results.

**Interpretation:** Laboratories should determine the risk for non-comparability of results. The higher the risk, the more frequent comparability of results should be checked. The minimum time frame for assessing comparability is every six months.

**Evidence:** Specific records demonstrating the frequency of the comparability of results assessment.

**Where:** Regional/organizational and facility assessments

**Comment:** There is no absolute answer to the question, “How often should verification of the comparability of results be performed?” Although the standards state that comparability verification should be performed at least twice a year, the laboratory should determine the risk for non-comparable results along with cost considerations and operational viability. Initially, a laboratory should perform comparability verification more frequently until confidence in comparability is established. Once the actual comparability performance has been established, the laboratory can evaluate the risk and reduce comparability testing appropriately. Laboratories should also consider those situations where special cause comparability verification is performed in addition to routinely scheduled comparability verification in instances of major maintenance, software update, clinician inquiry regarding results accuracy, or resolution of other problems.

**Examinations subject to comparability in DAP assessments**

As established by a multidiscipline comparability and measurement uncertainty guidance group, July 29, 2017.

**Anatomic pathology**

None at this time. For future consideration:

- immunohistochemistry

**Chemistry**

All measurands, except the following:

- body fluids
- trace metals
- urinalysis
- less frequent special chemistry testing
- results based on calculations from measured parameters when comparability already assessed

**Cytogenetics**

None at this time.
Cytology
None at this time.

Hematology
- CBC, differentials and any reportable results thereof
- reticulocytes
- coagulation testing including INR, PTT, D-dimer, fibrinogen, thrombin

Microbiology
- primary vs. backup methodology and instruments
- molecular testing
- serology
Automated vs. manual methods:
- antimicrobial susceptibility testing
- bacteria/yeast identification

Molecular diagnostics
None at this time.

Point-of-care testing
All measurands.

Transfusion medicine
- ABO/Rh typing
- antibody screening and identification
- crossmatch methods
- direct antiglobulin testing
For future consideration:
- reaction grading
- phenotyping
Example: Comparability of examination results at a facility

Consider the comparability of sodium results within a single laboratory site. To establish comparability of sodium results the region has determined the reportable range of the measurand (110-170 mmol/L).

The laboratory has determined the number of samples; in this case, three patient samples with results that span the reportable range as much as possible. Although the reportable range may be from 110-170 mmol/L, the laboratory is only able to find samples that span a smaller range.

The results for the patient samples are as follows:

<table>
<thead>
<tr>
<th>Instrument/Equipment</th>
<th>Sample 1</th>
<th>Sample 2</th>
<th>Sample 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary laboratory analyzer</td>
<td>120</td>
<td>138</td>
<td>152</td>
</tr>
<tr>
<td>Backup analyzer</td>
<td>125</td>
<td>141</td>
<td>150</td>
</tr>
<tr>
<td>Point-of-care equipment</td>
<td>123</td>
<td>145</td>
<td>154</td>
</tr>
</tbody>
</table>

QUA3.1.1 and QUA3.1.2 are assessed at the regional level and not used at the facility level.

**QUA3.1.3** Acceptability criteria are defined for comparability of procedures, equipment and methods.

**Interpretation:** Laboratories must establish the tolerance limits for comparability between sites and between instruments. See below.

The laboratory has also determined that results from the backup analyzer and point-of-care instrumentation must be within 5% of the primary laboratory analyzer to be considered comparable.

<table>
<thead>
<tr>
<th>Instrument/Equipment</th>
<th>Sample 1</th>
<th>Sample 2</th>
<th>Sample 3</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional laboratory analyzer</td>
<td>120</td>
<td>138</td>
<td>152</td>
<td>146–160</td>
</tr>
<tr>
<td>5% tolerance limit</td>
<td>114–126</td>
<td>131–145</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Backup analyzer</td>
<td>125</td>
<td>141</td>
<td>150</td>
<td>Pass</td>
</tr>
<tr>
<td>Point-of-care equipment</td>
<td>123</td>
<td>145</td>
<td>154</td>
<td>Pass</td>
</tr>
</tbody>
</table>
In this case, comparability has been demonstrated so there is no need for the laboratory to perform any tasks associated with QUA3.1.3 and QUA3.1.4 for this measurand.

**QUA3.1.6**

Instruments and methods are checked against each other at least twice a year for comparability of results.

The laboratory has records of comparability testing for each assessed measurand.

In this example, checking comparability has been performed across a suitable range with adequate samples. The backup analyzer and point-of-care testing is within the established tolerance limit compared to the primary laboratory analyzer with an acceptable frequency. Comparability of results has been demonstrated for this measurand at this site.

**References**

