ilirubin AST NORMAL ? .. GGT ALT 3.5 Albumin mmo Lipid Profile 2.25 Cholesterol mm 1.0 Triglyceride m polesterol 1.5

Presented by:

Mike Kramer

Calibration Program Manager

Perry Johnson Laboratory Accreditation, Inc. 28-February-2020



This webinar is being recorded and will be available in it's entirely on the Perry Johnson Laboratory Accreditation Website.

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Also individual slides of this and previous presentation are available.

There is a space on your screen to ask questions. Please keep question related to today's topic. At the conclusion of the webinar, received questions will be reviewed and answered.

Duration of webinar is set for one hour.





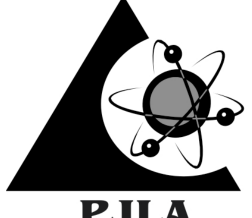
If you think about it, the report that a laboratory is providing it's customers is the final product. Things that can impact the final results may include having trained, personnel, appropriate procedures, appropriate facilities, traceability, equipment, and a handling process. All of which are addressed within ISOIEC 17025:2017





Reporting the Results

Calibration or test reports whom utilizes the symbol or references accreditation on calibration or test reports are considered accredited certificates or reports and come under the realm of Section 7.8 of ISO/IEC 17025:2017 and any applicable polices of the accreditation body. Refer to PJLA SOP 3 "Accreditation Symbol Procedure"





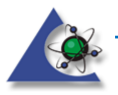
PJLA Calibration



This section covers all three area's of 17025 which includes testing, calibration, and sampling REPORTING THE RESULTS (7.8)

7.8.1 General – review and authorize prior to release

7.8.2 Common requirements for reports (test, calibration & sampling)
7.8.3 Specific requirements for test reports
7.8.4 Specific requirements for calibration certificates
7.8.5 Specific requirements for reporting sampling
7.8.6 Reporting statements of conformity
7.8.7 Reporting opinions and interpretations
7.8.8 Amendments to reports



7.8.1 General

7.8.1.1 The results shall be reviewed and authorized prior to release

By who?



6.2.5 The laboratory shall have procedure(s) and retain records for:e) authorization of personnel;

6.2.6 The laboratory shall authorize personnel to perform specific laboratory activities, including but not limited to, the following:

c) report, review and authorization of results.



7.8.1.2 The results shall be provided accurately, clearly, unambiguously and objectively, usually in a report (e.g. a test report or a calibration certificate or report of sampling), and shall include all the information agreed with the customer and necessary for the interpretation of the results and all information required by the method used. All issued reports shall be retained as technical records.

If something is unambiguous, there are no two ways to interpret it. When you do something objectively, you do it with an open mind, considering the facts rather than your personal feelings.



NOTE 1 For the purposes of this document, test reports and calibration certificates are sometimes referred to as test certificates and calibration reports, respectively.

NOTE 2 Reports can be issued as hard copies or by electronic means, provided that the requirements of this document are met



Even though the 2017 Standard is more in tune with today's electronic age, pager reports, records, and documents are still recognized.



7.8.1.3 When agreed with the customer, the results may be reported in a simplified way. Any information listed in 7.8.2 to 7.8.7 that is not reported to the customer shall be readily available.

--no longer specifies internal customers or written agreement with external customers;

From the 2005 Standard

In the case of tests or calibrations performed for internal customers, or in the case of a written agreement with the customer, the results may be reported in a simplified way;

-The lab should however maintain a record of this agreement PL-3 specifies written agreement in regards to uncertainty reporting;



7.8.2 Common requirements for reports (test, calibration or sampling)

- **7.8.2.1** Each report shall include at least the following information, **unless the laboratory has valid reasons** for not doing so, thereby minimizing any possibility of misunderstanding or misuse:
- a) a title (e.g. "Test Report", "Calibration Certificate" or "Report of Sampling");
- b) the name and address of the laboratory
- c) the location of performance of the laboratory activities, including when performed at a customer facility or at sites away from the laboratory's permanent facilities, or in associated temporary or mobile facilities;



On-site Calibration Services

We can visit your premises to calibrate your equipment to reduce your downtime

nd out more



Common requirements for reports (test, calibration or sampling

- d) unique identification that all its components are recognized as a portion of a complete report and a clear identification of the end;
- e) the name and **contact information** of the customer;
- 2005 specified name and address
- f) identification of the method used;
- Can be a lab procedure (SOP 123), or a Standard procedure such as ASTM , ASME or USP type method
- g) a description, unambiguous identification, and, when necessary, the condition of the item;
- -in other words what was tested or calibrated,



Common requirements for reports (test, calibration or sampling)

h) the date of receipt of the test or calibration item(s), and the date of sampling, where this is critical to the validity and application of the results;

-Shelf life may be critical if it sample may degrade over time (moisture content), therefore it would be important to record date of receipt.

- i) the date(s) of performance of the laboratory activity;
- j) the date of issue of the report; -new

k) reference to the sampling plan and sampling method used by the laboratory or other bodies where these are relevant to the validity or application of the results;



Common requirements for reports (test, calibration or sampling)

- l) a statement to the effect that the results relate only to the items tested, calibrated or sampled;
- -2005 specified where relevant
- Coordinate Measuring Machine with a serial number it would be obvious;
- If calibrating a dozen $\frac{1}{2}$ inch micrometers it would be relevant to state which results are for which micrometer;

m) the results with. where appropriate. the units of measurement;

Quantity	Unit	Abbreviation m kg	
Length	Meter		
Mass	Kilogram		
Time	Second	S	
Thermodynamic Temperature	Kelvin	К	
Amount of Substance	Mole	mol	
Electric Current	Ampere	A	



Common requirements for reports (test, calibration or sampling)

n) additions to, deviations, or exclusions from the method;

- o) identification of the person(s) authorizing the report;
- p) clear identification when results are from external providers.
- -this is new and would be from subcontractors

NOTE Including a statement specifying that the report shall not be reproduced except in full without approval of the laboratory can provide assurance that parts of a report are not taken out of context.

-cut and paste parts of a report



Common requirements for reports (test, calibration or sampling

7.8.2.2 The laboratory shall be responsible for all the information provided in the report, except when information is provided by the customer. Data provided by a customer shall be clearly identified. In addition, a disclaimer shall be put on the report when the information is supplied by the customer and can affect the validity of results. Where the laboratory has not been responsible for the sampling stage (e.g. the sample has been provided by the customer), it shall state in the report that the results apply to the sample as received

- Information provided by the customer may be needed to proceed with test or calibration if environmental criteria are not within predefined limits (customer location)



7.8.3 Specific requirements for test reports

- Specific requirements for test reports are in place without change from the 2005 Standard;

7.8.3.1 In addition to the requirements listed in <u>7.8.2</u>, test reports shall, where necessary for the interpretation of the test results, include the following:

a) information on specific test conditions, such as environmental conditions;

b) where relevant, a statement of conformity with requirements or specifications (see $\underline{7.8.6}$);



Specific requirements for test reports



c) where applicable, the measurement uncertainty presented in the same unit as that of the measured or in a term relative to the measured (e.g. percent) when:

- it is relevant to the validity or application of the test results;
- a customer's instruction so requires, or
- the measurement uncertainty affects conformity to a specification limit



Specific requirements for test reports

d) where appropriate, opinions and interpretations (see <u>7.8.7</u>);
e) additional information that may be required by specific methods, authorities, customers or groups of customers

7.8.3.2 Where the laboratory is responsible for the sampling activity, test reports shall meet the requirements listed in 7.8.5 where necessary for the interpretation of test results.

-This is when the laboratory is actually collecting the samples to be tested.



Sampling Clover Seed

Section 7.8 of ISO/IEC 17025:2017 Reporting of Results 7.8.4 Specific requirements for calibration certificates

7.8.4.1 In addition to the requirements listed in $\underline{7.8.2}$, calibration certificates shall include the following:

a) the measurement uncertainty of the measurement result presented in the same unit as that of the measurand or in a term relative to the measurand (e.g. percent);

NOTE According to ISO/IEC Guide 99, a measurement result is generally expressed as a single measured quantity value including unit of measurement and a measurement uncertainty

The 2005 Standard specifies "the uncertainty of measurement **and/or** a statement of compliance with an identified metrological specification or clauses thereof"



Specific requirements for calibration certificates calibration certificates shall include the following:

b) the conditions (e.g. environmental) under which the calibrations were made that have an influence on the measurement results;

c) a statement identifying how the measurements are metrologically traceable (see <u>Annex A</u>);

- Also see PL-2 "To SI through NIST"
- -2005 specified evidence and noted accreditation body symbol *NIST Commercial*

Lab





Specific requirements for calibration certificates

calibration certificates shall include the following:

d) the results before and after any adjustment or repair, if available;

e) where relevant, a statement of conformity with requirements or specifications (see $\underline{7.8.6}$);

- Pass/Fail In Tolerance/Out of Tolerance

f) where appropriate, opinions and interpretations (see $\underline{7.8.7}$).

7.8.4.2 Where the laboratory is responsible for the sampling activity, calibration certificates shall meet the requirements listed in $\underline{7.8.5}$ where necessary for the interpretation of calibration results

--sampling is primarily associated with testing labs.



Specific requirements for calibration certificates

Calibration certificates shall include the following:

7.8.4.3 A calibration certificate or calibration label shall not contain any recommendation on the calibration interval, except where this has been agreed with the customer.

--Not required to be on reports however if it is then it is required to be customer specified



Section 7.8 of ISO/IEC 17025:2017 Reporting of Results 7.8.5 Reporting sampling – specific requirements

- Where the laboratory is responsible for the sampling activity, in addition to the requirements listed in 7.8.2, reports shall include the following, where necessary for the interpretation of results
- a) the date of sampling;
- b) unique identification of the item or material sampled (including the name of the manufacturer, the model or type of designation and serial numbers, as appropriate);
- c) the location of sampling, including any diagrams, sketches or photographs;
- d) a reference to the sampling plan and sampling method;



Reporting sampling – specific requirements

e) details of any environmental conditions during sampling that affect the interpretation of the results

f) information required to evaluate measurement uncertainty for subsequent testing or calibration -this is new





Section 7.8 of ISO/IEC 17025:2017 Reporting of Results 7.8.6 Reporting statements of conformity

7.8.6.1 When a statement of conformity to a specification or standard is provided, the laboratory shall document the **decision rule** employed, taking into account the level of risk (such as false accept and false reject and statistical assumptions) associated with the decision rule employed, and apply the decision rule

As defined in ISO/IEC 17025:2017

decision rule - rule that describes how measurement uncertainty is accounted for when stating conformity with a specified requirement

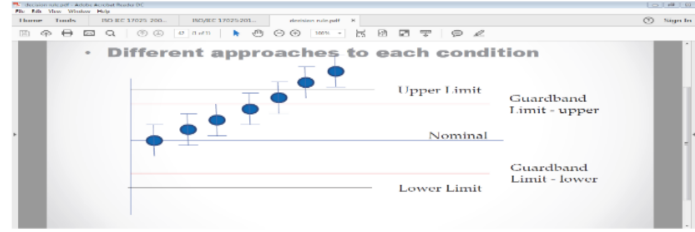


Reporting statements of conformity

7.8.6.2 The laboratory shall report on the statement of conformity, such that the statement clearly identifies:

- a) to which results the statement of conformity applies;
- b) which specifications, standards or parts thereof are met or not met;

c) the decision rule applied (unless it is inherent in the requested specification or standard



NOTE For further information, see ISO/IEC Guide 98-4

Reporting statements of conformity

If accepted by the customer as per the requirements specified in 7.1.3, the following decision rules can be documented:

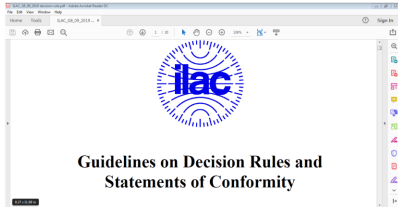
Accounting for the uncertainty will be taken to mean that at a 95% confidence level the measurement result plus and minus the expanded uncertainty (k=2) shall be totally within the specification limits

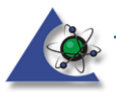




ILAC-G8:09/2019 Guidelines on Decision Rules and Statements of Conformity

- This guidance document has been prepared to assist laboratories in the use of decision rules when declaring statements of conformity to a specification or standard as required by ISO/IEC 17025:2017
- The revised ISO/IEC 17025:2017 recognizes that no single decision rule can address all statements of conformity across the diverse scope of testing and calibration





Reporting statements of conformity

There are testing methods that determine how the rules are applied. One good, common illustration is ASTM E18 for Rockwell Hardness where the testing and calibration decision rules take uncertainty into account effectively in the repeat testing and other "limits" as to the spread of the data etc. and the rules are defined in the method.

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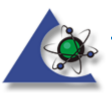


7.8.7 Reporting opinions and interpretations

7.8.7.1 When opinions and interpretations are expressed, the laboratory shall ensure that only personnel authorized for the expression of opinions and interpretations release the respective statement. The laboratory shall document the basis upon which the opinions and interpretations have been made.

-This is new

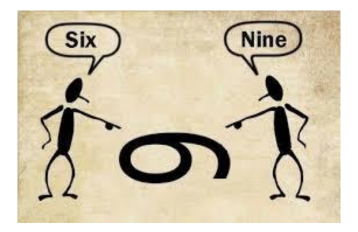
7.8.7.2 The opinions and interpretations expressed in reports shall be based on the results obtained from the tested or calibrated item and shall be clearly identified as such.



7.8.7 Reporting opinions and interpretations

7.8.7.3 When opinions and interpretations are directly communicated by dialogue with the customer, a record of the dialogue shall be retained

-This is a note in the 2005 Standard





Section 7.8 of ISO/IEC 17025:2017 Reporting of Results 7.8.8 Amendments to reports

7.8.8.1 When an issued report needs to be changed, amended or reissued, any change of information shall be clearly identified and, where appropriate, the reason for the change included in the report

- The concept of where appropriate the reason for the change included in the report is an addition

7.8.8.2 Amendments to a report after issue shall be made only in the form of a further document, or data transfer, which includes the statement "Amendment to Report, serial number... [or as otherwise identified]", or an equivalent form of wording

Such amendments shall meet all the requirements of this document



Amendments to reports

7.8.8.3 When it is necessary to issue a complete new report, this shall be uniquely identified and shall contain a reference to the original that it replaces

Certificate # 1234AR This report replaces certiticate #1234



Thank you!

This time is allocated for questions. You should have a space provided for submitting questions.





Save the Date

Next PJLA Webinar

Thursday March 19, 2020 1:00 PM EST

ISO/IEC 17025:2017 Section 7.8 Complaints

