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## **Documentation and Error Correction for Laboratory Records**

#### **Purpose**

This procedure describes how to document laboratory work for written records, as well as how to document error correction on records.

#### **Policy**

- All work performed is recorded legibly. Work is recorded in such a manner that another individual, competent in the same field, may interpret the work solely from the description written without additional explanation.
- Observations and calculations are to be clearly and permanently recorded at the time they are made (concurrent documentation).
  - Work that is not documented concurrently must be identified as a late entry or as an addendum with current date/time.
- Entries contain a date, and initials/signature of person creating the record.
- Entries are made in indelible blue or black ink. Space not used (e.g., no testing performed on that day) is indicated as specified in the legend on the form. If no legend exists, annotate the form to include the abbreviation used. Example: N/A = Not Applicable, or NT = Not Tested
- Corrections are made by drawing a single line through the incorrect entry, entering the correct information; initial and date the change.
  - The original entry must not be obliterated and should remain legible.
- Records or documents must be kept for the appropriate timeframe per the policy *Retention of Laboratory Records and Materials*. Records within the retention guideline must not be discarded without explanation.
- Examples of unacceptable documentation or error correction include:
  - Ditto marks, arrows, or continuation lines instead of entering data
  - A stamp in lieu of a handwritten signature
  - White-out or any similar product, blacking out with marker
  - Erased entry
  - Write over
- Use of scrap paper, Post-it notes or any similar product that would intentionally record raw data on non-official records is prohibited.
- Use form legend when using abbreviations on records.

### Scope

This procedure applies to all Laboratory staff, supervisors, and managers related to the management of all laboratory records.

# Special safety precautions

Refer to the safety manual for general safety requirements.

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# **Documentation and Error Correction for Laboratory Records,**Continued

#### **Definitions**

- Electronic Record An electronic record is information recorded in a form that only a computer can process. Electronic records include numeric, graphic and textual information.
- Form A form is a document with a fixed arrangement of captioned spaces designed for entering and extracting prescribed information. Forms become a record once filled out.
- Non-record Non-records are copies of memoranda or letters sent to an office or an employee for information only and for whose filing or maintenance no one in the office is responsible.
- Quality records Quality records include the following: laboratory operating procedures, preventive maintenance charts, quality control (QC) charts, internal audit reports, management reviews, corrective and preventive actions.
- Records Records are materials created or received by a department and that are preserved as evidence of the activities of the department or for its information value. Records include reports, correspondence, diaries, quality records and technical records.
- Technical records Technical records are accumulations of data and information which result from carrying out tests or calibrations and which indicate whether specified quality or process parameters are achieved. Technical records include forms, worksheets, control graphs, inspection reports, and test reports.

#### **Documentation**

Follow these steps for documentation.

Step	Action
1	Where documents require the entry of data, enter the data clearly
	and legibly with indelible blue or black ink.
2	<ul> <li>Make data entries or complete records at the time each action is taken or document as a late entry. Temperatures and other observations that may change over time may not be entered as a late entry.</li> <li>If there are missing entries at the time of record review, the empty field should be circled. Multiple missing entries should be investigated and corrective or preventive action taken. If necessary, make photocopies of missing entries to be used for corrective or preventive action.</li> </ul>

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# **Documentation and Error Correction for Laboratory Records, Continued**

# Documentation, continued

Step	Action
3	Do not leave spaces for handwritten entries blank. Space not used
	(e.g., no testing performed on that day) is indicated as specified in
	the legend on the form. If no legend exists, annotate the form to
	include the abbreviation used. Example: $N/A = Not$ Applicable, or
	NT = Not Tested

# Omission of **Documentation**

Follow these steps correcting the omission of documentation.

Step	Action
1	Omission of documentation can be corrected as late entry or an
	addendum to the document with current date and/or time as
	indicated. It is considered falsification of records to go back and
	complete or to fill-in the blanks.
2	Late entry is allowed only when there is supporting information to
	validate the entry.
3	Late entry is allowed within the designated time frame and is at the
	discretion of department administrator.
4	Operating departments are encouraged to have frequent
	monitoring to assure that documentation is complete and timely.
	Examples include shift-to-shift review or daily peer review.

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# **Documentation and Error Correction for Laboratory Records, Continued**

## Error correction

Follow these steps for error correction.

Step	Action		
1	Verify the written error before making the correction.		
	The person making the correction must be knowledgeable of		
	the incident and authorized by training to make the change.		
	All changes that would change the intent or substance of the		
	original documentation must have supervisor approval.		
2	Make corrections of daily temperatures or preventive		
	maintenance on the Action Log or in the space marked		
	accordingly.		
	• Draw one line across the incorrect data, place an asterisk (*)		
	next to the incorrect entry, and document the data correction		
	on the form appropriately.		
3	Make clear corrections of QC records on the form when possible.		
	If the correction cannot be made clearly, then draw one line		
	across the incorrect data and place an asterisk (*) next to the		
	incorrect data entry.		
	Document the correction and any additional details on the form		
	appropriately.		
4	Make any correction on the manual records only by drawing a line		
	through the original entry, so as not to obliterate that record, and		
	then writing the corrected entry with an initial and date of the		
	person making the correction.		

# Non-controlled documents

The following non-controlled documents support this procedure:

- J. Ezzelle, I. R. Rodriguez-Chavez, J. M. Darden, M. Stirewalt, N. Kunwar, R. Hitchcock, T. Walter, and M. P. D'Souza. *Guidelines on Good Clinical Laboratory Practice*, <a href="http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2213906/?tool=pubmed">http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2213906/?tool=pubmed</a>, downloaded July 5, 2010.
- Good Documentation Practice, Wikipedia, https://en.wikipedia.org/wiki/Good\_documentation\_practice, downloaded May 30, 2017.

### Signature Manifest

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### **Doc & Error Correction for Lab Rec**

## **Initial Approval**

Name/Signature	Title	Date	Meaning/Reason
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Fred Ung (K057175)	SCPMG LABORATORY QCD	09 May 2017, 09:53:39 AM	Approved

### **Final Approval**

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### **Set Effective Date**

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