

QSE 6: Equipment Management

Policy	The laboratory keeps detailed records on each piece of equipment that is critical in the processes across the laboratory's path of workflow.	
Purpose	This policy provides direction for the processes and procedures to effectively manage the laboratory's equipment.	
Responsibility	The Laboratory Director is responsible for laboratory acquisition decisions, implementing an instrument verification and maintenance program, and documentation review.	
	The Purchasing Department is responsible for the actual purchasing process associated with instrument purchases.	
	The Laboratory Supervisor is responsible for instrument justification and comparative analyses.	
	The Lead Technologists and Quality Manager are responsible for the validation, maintenance and repair once the instruments are on-site.	
Equipment Selection, Acquisition, Installation, and Inventory	The laboratory acquires and installs the equipment needed for producing quality results. An inventory of all laboratory equipment is maintained.	
Validation Studies	Validation studies are performed and documented as required on laboratory instrumentation to include accuracy, precision, linearity, reportable ranges, reportable reference ranges, sensitivity (as needed) and specificity (as needed).	
Method Comparison	When more than one method or instrument is used to conduct the same test, the methods or instruments are compared semi-annually.	
Carryover	Studies are performed to show non-FDA approved instruments or test systems do not have any carryover.	
Maintenance, Calibration, Use, Troubleshooting, Service, Repair, and Records	Each instrument in use has a separate manual that includes all instructions and documentation generated during the life of the instrument indefinitely or as otherwise directed.	
Retiring Instruments	The laboratory retires instruments that are no longer in use and has procedures for storage of retired instruments and their records.	
Supporting Documents	 The following processes support this policy: Equipment Acquisition, Installation, Identification, and Inventory Instrument Validation Studies Method Comparisons Carryover Studies Instrument and Equipment Preventative/Corrective Maintenance Troubleshooting and Corrective Actions Retiring Instruments 	



Process for Equipment Selection, Acquisition, Installation, Identification, and Inventory

What Happens	Who's Responsible	Procedures
Decision is made to purchase new equipment or replace equipment in use	 Laboratory Director Laboratory Supervisor Lead Technologist 	 Laboratory Testing Turn Around Times Equipment Inventory Specific Equipment Service and Maintenance Records
Research is performed for available instrumentation	 Laboratory Director Laboratory Supervisor Lead Technologist 	 Requesting Bids from Suppliers Test-Specific Procedures Equipment Selection
Equipment justification is performed	Laboratory DirectorLaboratory SupervisorFinance Department	Finance Department Policies and Procedures
Equipment purchase request is completed and approved	Laboratory SupervisorPurchasing Department	 Completing a Purchase Request Form Equipment Acquisition
New equipment is installed by manufacturer or approved representative	 Laboratory Supervisor Lead Technologist Quality Manager Manufacturer/ Representative 	 Manufacturers' Equipment Manuals Installing New Laboratory Equipment
Equipment is assigned a unique identifier	Laboratory SupervisorLead Technologist	Equipment Identification
Equipment is included in an inventory	Laboratory SupervisorLead Technologist	Equipment Inventory
All instructions and documentation related to an instrument are kept in an equipment manual specific to that piece of equipment	 Laboratory Supervisor Lead Technologist Quality Manager Technicians/Technologists 	 Documenting Equipment- Related Activities Equipment-Related Records



Process for Instrument Validation Studies

What Happens	Who's Responsible	Procedures
New instrument validation studies are performed, documented, and evaluated for approval in- house	Laboratory DirectorLead TechnologistsQuality Manager	 Instrument-Specific Validation Procedures Laboratory Section- Specific Validation
	Technicians/Technologists	Procedures
Annual or as needed validation studies are performed, documented, and evaluated for approval	 Laboratory Director Lead Technologists Quality Manager Technicians/Technologists 	 Instrument-Specific Validation Procedures Laboratory Section- Specific Validation Procedures
When equipment is removed from the laboratory or is serviced/repaired, performance is validated before use in patient testing	 Laboratory Director Lead Technologists Quality Manager Technicians/Technologists 	 Instrument-Specific Validation Procedures Laboratory Section- Specific Validation Procedures



Process for Method Comparisons

What Happens	Who's Responsible	Procedures
Two laboratory instruments or methods are used in testing of the same analytes	 Lead Technologist Quality Manager Technicians/Technologists 	 Performing Comparison Studies Test-Specific SOPs Instrument-Specific User Manuals
Both manual and automated methods are used to result a laboratory analyte A manual method is used to confirm an analyzer- generated result	 Lead Technologist Quality Manager Technicians/Technologists Lead Technologist Quality Manager Technicians/Technologists 	 Performing Comparison Studies Test-Specific SOPs Instrument-Specific User Manuals Performing Comparison Studies Test-Specific SOPs Instrument-Specific User Manuals
An instrument/method comparison study is performed semi-annually	 Lead Technologist Quality Manager Technicians/Technologists 	 Performing Comparison Studies Test-Specific SOPs Instrument-Specific User Manuals



Process for Carryover Studies

What Happens	Who's Responsible	Procedures
New instrument carryover studies are performed, documented, and evaluated for approval	 Laboratory Director Lead Technologists Quality Manager Technicians/Technologists 	 Instrument-Specific Carryover Study Procedures
Additional instrument carryover studies are performed, documented, and evaluated for approval as needed	 Laboratory Director Lead Technologists Quality Manager Technicians/Technologists 	Instrument-Specific Carryover Study Procedures



Process for Instrument and Equipment Preventive/Corrective Maintenance

What Happens	Who's Responsible	Procedures
Instrument and/or equipment maintenance is performed on a daily, weekly, monthly, quarterly, semi-annual, or annual schedule	 Lead Technologist Technologists/ Technicians 	 Equipment-Specific Procedures Manufacturer's Equipment Manual Maintenance Schedule
Maintenance, service, or repair is performed on instruments or equipment at times other than scheduled	 Lead Technologist Technologists/ Technicians Instrument Manufacturer Service Personnel Facility Engineering Department Service Personnel 	 Equipment-Specific Procedures Manufacturer's Equipment Manual Emergency Service Requests Requesting Service or Repair from Facility Engineering Department
All maintenance, service, and repair activities are documented and maintained in the equipment manual	 Lead Technologist Technologists/Technicians 	 Maintenance, Service, and Repair Records Equipment-Specific Procedures
Documentation is reviewed at least monthly	Quality ManagerLaboratory Director	 Equipment-Specific Procedures Maintenance, Service, and Repair Records Maintenance Schedule Review and Storage of Laboratory Records



Process for Calibration

What Happens	Who's Responsible	Procedures
Calibration schedule is developed based on manufacturer's recommendations	 Laboratory Supervisor Lead Technologist Quality Manager 	 Manufacturer's Equipment Manual Calibration Procedure Instrument/Equipment- Specific Procedures
Calibration is performed and documented at regularly scheduled intervals or as needed for troubleshooting	 Lead Technologist Quality Manager Technicians/Technologists 	 Manufacturer's Equipment Manual Calibration Procedure Instrument/Equipment- Specific Procedures
Documentation is reviewed and filed	Quality ManagerLaboratory Director	 Manufacturer's Equipment Manual Calibration Procedure Instrument/Equipment Specific Procedures- Review and Storage of Laboratory Records



Process for Troubleshooting and Corrective Actions

What Happens	Who's Responsible	Procedures
Trouble-shooting and training schemes are developed	 Laboratory Supervisor Quality Manager Lead Technologist Technologists/Technicians 	 Instrument/Laboratory Section-Specific Procedures Manufacturer's Manuals
Problem is identified and documented	 Quality Manager Lead Technologist Technologists/Technicians 	 Instrument/Laboratory Section-Specific Procedures Manufacturer's Manuals Documenting Problems and Corrective Actions
Corrective Action is performed and documented	 Technologist/Technicians Lead Technologist Quality Manager 	 Instrument/Laboratory Section-Specific Procedures Manufacturer's Manuals Documenting Problems and Corrective Actions
Corrective Action documentation is reviewed and filed	Laboratory DirectorQuality Manager	 Documenting Problems and Corrective Actions Review and Storage of Laboratory Records



Process for Retiring Instruments

What Happens	Who's Responsible	Procedures
New instrument validation studies are completed	Lead TechnologistQuality Manager	 Technical SOPs Instrument-Specific Validation Procedures
Retired instrument is cleaned and decontaminated for storage	Quality ManagerTechnicians/Technologists	 Retiring Laboratory Instruments/Equipment Equipment Decontamination
Instrument is packed and removed for storage	 Facility Maintenance Department Facility Engineering Department Quality Manager 	Retiring Laboratory Instruments/Equipment
Instrument records are stored indefinitely (or as otherwise directed) after retirement	Laboratory SupervisorLaboratory DirectorQuality Manager	 Retiring and Archiving Laboratory Records Retiring Laboratory Instruments/Equipment