
Abbott Value Assignment Sheets

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UNDERSTANDING ABBOTT VALUE ASSIGNMENT SHEETS: A GUIDE FOR CLINICAL LABORATORIES

In the fast-paced world of clinical diagnostics, precision is not just a goal—it is an absolute necessity. Every test result that leaves a laboratory carries the weight of clinical decisions, from routine health screenings to critical disease management. Behind the scenes, ensuring that every instrument and reagent performs as expected is a complex process involving rigorous quality control protocols. Central to this process in laboratories utilizing Abbott diagnostics are the documents known as **Abbott Value Assignment Sheets**. These sheets serve as the authoritative reference for the target values and acceptable ranges of control materials, playing a pivotal role in maintaining the accuracy and reliability of diagnostic testing. This article delves into the purpose, structure, and importance of these sheets, providing a natural and comprehensive look at how they support laboratory quality assurance and, ultimately, better patient outcomes.

WHAT ARE ABBOTT VALUE ASSIGNMENT SHEETS?

At their core, Abbott Value Assignment Sheets are technical documents that accompany Abbott quality control materials, calibrators, and certain reagents. They provide the assigned

target values—often referred to as the mean—and the acceptable range, usually expressed as a standard deviation or a confidence interval, for each analyte measured. For example, a sheet for an Abbott LiquidQC control material will list the expected concentrations for a panel of analytes such as glucose, creatinine, sodium, potassium, cardiac troponin, or thyroid hormones, specific to a particular instrument platform and reagent lot.

These sheets are not generic; they are lot-specific and instrument-specific. This means that each batch of control material produced by Abbott undergoes a rigorous process of value assignment, and the resulting values are printed on the sheet that accompanies that specific lot. Laboratories rely on these sheets to set up their internal quality control procedures, ensuring that the controls they run daily yield results within the expected range. When a control result falls outside this range, it signals a potential issue with the reagent, calibrator, instrument, or technique, prompting corrective action before patient samples are reported.

Why Value Assignment Sheets Matter in

Quality Control

The foundation of any robust quality control (QC) program is the ability to distinguish between normal analytical variation and true error. Abbott Value Assignment Sheets provide the benchmark against which all QC results are compared. Without these assigned values, laboratories would be operating without a reliable reference, essentially flying blind. The sheets enable laboratories to implement statistical QC rules—such as Westgard rules—by providing the necessary parameters to calculate means, standard deviations, and control limits.

Furthermore, these sheets support compliance with regulatory standards. Accreditation bodies such as the College of American Pathologists (CAP), the Joint Commission, and ISO 15189 require laboratories to have documented procedures for verifying that control materials perform as expected. Using the values from the Abbott Value Assignment Sheets as the target standard is a recognized and accepted practice for meeting these requirements. This ensures that the laboratory is not only following best practices but also maintaining its accreditation status.

THE ANATOMY OF AN ABBOTT VALUE ASSIGNMENT SHEET

While the specific layout may vary slightly depending on the product and instrument platform, most Abbott Value Assignment Sheets share common elements. Understanding these components helps laboratory professionals interpret the information correctly and apply it in their QC workflows.

1. **Product Identification:** The sheet clearly identifies the control material or calibrator by name, catalog number, and lot number. This ensures that the correct sheet is used for the specific material in use.
2. **Instrument Platform:** The values are specific to the Abbott instrument model, such as the Architect i2000, Alinity i, or Alinity c series. Using a sheet for the wrong platform can lead to incorrect target values and QC errors.
3. **Analyte List:** Each analyte tested is listed with its name, unit of measure, and assigned value (mean). This may include multiple levels of control, such as Level 1, Level 2, and Level 3, each with its own target.
4. **Acceptable Range:** The sheet provides the acceptable range, typically expressed as a lower and upper limit. This range is derived from statistical analysis of replicate measurements performed during the value assignment process.
5. **Standard Deviation (SD) and %CV:** Often, the sheet includes the standard deviation and percentage coefficient of variation (%CV) associated with the assigned value. These metrics give laboratories insight into the expected variability of the control material.
6. **Lot Numbers of Reagents and Calibrators:** Some sheets list the lot numbers of the reagents and calibrators that were used during the value assignment. This helps laboratories trace any potential interactions between different lots.

7. Date of Assignment and

Expiration: The date the values were assigned and the expiration date of the control material are clearly marked, ensuring that the sheet is used only while the material is valid.

THE PROCESS OF VALUE ASSIGNMENT: HOW ABBOTT ENSURES RELIABILITY

The values printed on an Abbott Value Assignment Sheet are not arbitrary; they are the result of a meticulous and standardized process designed to ensure accuracy and reproducibility across different laboratories and instrument systems. Understanding this process reinforces confidence in the sheets.

Abbott typically performs value assignment using a multi-site, multi-instrument study. This involves running the control material on multiple instruments of the same model across different geographical locations. By pooling data from numerous instruments, Abbott can calculate a robust mean and standard deviation that account for both within-instrument and between-instrument variability. This statistical approach ensures that the assigned values are representative of what a well-calibrated instrument in a typical laboratory should produce.

Reference methods and traceability to higher-order standards are also key. For many analytes, Abbott calibrators and controls are traceable to reference materials from organizations like the National Institute of Standards and Technology (NIST) or the International Federation of Clinical Chemistry and Laboratory Medicine (IFCC). This traceability chain ensures that the values on the sheet are not just arbitrary

numbers but are anchored to an international standard of measurement. This is essential for the comparability of results across different laboratories and over time.

PRACTICAL USE OF ABBOTT VALUE ASSIGNMENT SHEETS IN THE LABORATORY

For laboratory technicians and clinical scientists, the Abbott Value Assignment Sheet is a daily working document. Here is how it is typically used in practice:

Setting Up Quality Control Protocols

When a new lot of control material is received, the laboratory enters the assigned values and ranges from the sheet into the laboratory information system (LIS) or the instrument software. This establishes the target mean and control limits for that specific lot. The laboratory then begins running the control material daily, plotting results on a Levey-Jennings chart or similar QC graph. Each result is compared to the assigned mean and range from the sheet. This process is a fundamental requirement of ISO 15189 and CAP accreditation standards.

Investigating QC Failures

If a control result falls outside the acceptable range on the sheet, the laboratory immediately initiates a troubleshooting workflow. Common

steps include checking the expiration date of the control material, verifying proper storage and handling, inspecting reagent integrity, reviewing calibration status, and running a new aliquot of control. The sheet serves as the authoritative reference for determining whether a result is actually out of range or whether the sheet itself might be misinterpreted. In rare cases, if a systemic issue is suspected, the laboratory may contact Abbott technical support, referencing the lot number from the sheet for further investigation.

Comparing Across Platforms and Sites

For laboratories that operate multiple Abbott instruments or have multiple testing sites, the Value Assignment Sheets help ensure consistency. When the same control material lot is used across different instruments, the assigned values should theoretically be the same. If significant discrepancies appear between instruments, it indicates a calibration or performance issue that needs attention. This cross-site comparability is a major advantage of using lot-specific, instrument-specific value assignment sheets.

COMMON CHALLENGES AND BEST PRACTICES FOR USING VALUE ASSIGNMENT SHEETS

While Abbott Value Assignment Sheets are designed to be clear and reliable, laboratory professionals may encounter certain challenges. Awareness of these issues and adherence to best practices

can prevent errors and ensure consistent quality control.

- **Lot-to-Lot Variability:** Each new lot of control material may have slightly different assigned values. It is critical to update the QC software with the new values every time a new lot is introduced. Failure to do so is a common source of QC errors.
- **Reconstitution and Handling Errors:** For lyophilized (freeze-dried) controls, improper reconstitution—such as using the wrong volume of diluent or insufficient mixing—can alter the actual concentration, leading to results that do not match the sheet. Always follow the instructions provided with the control material and referenced on the sheet.
- **Storage Conditions:** Storing controls at incorrect temperatures or beyond their expiration date can degrade the analytes. The sheet specifies valid storage conditions, and these must be strictly followed.
- **Misinterpretation of Ranges:** Some users may mistakenly use the range on the sheet as fixed limits, rather than understanding that the range is a statistical estimate. In a robust QC program, the laboratory may calculate its own cumulative mean and range over time, which may differ slightly from the manufacturer's assigned values. This is acceptable when done correctly and documented.
- **Cross-Platform Use:** Using a Value Assignment Sheet intended for one instrument platform on another platform is a serious error.

Always verify that the sheet matches the specific instrument model in use.

BEST PRACTICES FOR LABORATORY STAFF

1. Always verify that the lot number on the control material vial matches the lot number on the Value Assignment Sheet.
2. Enter the assigned values and ranges into the QC system immediately upon receipt of a new lot.
3. Store the sheets in a designated binder or digital folder for easy reference during troubleshooting.
4. Train all laboratory personnel on how to read and interpret the sheets, including understanding units, SD, and range limits.
5. Participate in proficiency testing programs such as CAP surveys to verify that your laboratory's results align with peer groups using similar methods and materials.

THE ROLE OF ABBOTT VALUE ASSIGNMENT SHEETS IN PATIENT SAFETY AND REGULATORY COMPLIANCE

Beyond the laboratory bench, Abbott Value Assignment Sheets have a direct impact on patient care. When a laboratory follows the values on these sheets correctly, the QC process acts as a safety net, catching potential errors before they affect patient results. For example, if a control result for troponin—a critical cardiac marker—falls outside the expected range, the laboratory can delay reporting patient troponin results until the issue is

resolved. This prevents potentially misleading results that could affect clinical decisions such as ruling in or ruling out myocardial infarction.

From a regulatory perspective, these sheets are often reviewed during laboratory inspections. Surveyors from CAP, the Joint Commission, or other accrediting bodies may ask to see the Value Assignment Sheets for the control materials currently in use. They will verify that the laboratory has correctly entered the values, that the controls are within their expiration date, and that the laboratory has documented procedures for handling out-of-range results. Having well-organized and properly used sheets demonstrates a commitment to quality and can facilitate a smoother inspection process.

FUTURE TRENDS IN VALUE ASSIGNMENT: DIGITALIZATION AND AUTOMATION

As clinical laboratories continue to evolve toward greater automation and digital integration, the traditional paper-based Abbott Value Assignment Sheet is also changing. Many laboratories now use laboratory information systems (LIS) that allow for the direct import of value assignment data via barcode scanning or electronic data transfer. This reduces the risk of manual data entry errors and streamlines the QC setup process.

Abbott has been moving toward providing these sheets in electronic formats, accessible through portals that integrate with instrument software. In the future, we may see real-time updates to assigned values based on ongoing QC data aggregation across a network of instruments, providing even more robust and dynamic reference ranges. However, the fundamental

principle will remain the same: providing evidence-based reference for ensuring laboratory professionals with a trusted, the accuracy of diagnostic testing.