(A) Policy Statement

Laboratory must comply with all College of American Pathologists terms of accreditation in order to maintain two year accreditation on a continuous basis.

(B) Purpose of Policy

To help ensure UTMC laboratory maintains the high standard for which it received CAP accreditation.

(C) Procedure

The following requirements must be met to maintain continuous two year accreditation:

1. Provide an inspection team of a size and composition similar to that required for your inspection and perform at least one inspection during the two-year accreditation cycle, if asked to do so by Commissioner.

2. Submit an application for re-inspection in sufficient time that the re-inspection may be accomplished prior to your anniversary date.

3. Successfully participate in the CAP surveys or a CAP-approved alternative proficiency testing program.

4. Notify the commission of changes in location, ownership, directorship, location, insolvency, or bankruptcy of the laboratory. Notification must occur prior to the change(s); or, in the case of unexpected changes, no later than two working days afterwards.

5. Notify the Commission when there is a change in the test menu. Notification must occur prior to starting new patient testing.

6. Cooperation with CAP when the laboratory is subject to a CAP investigation or inspection.

7. Submit a completed Self-Evaluation Verification Form in the interim year.

8. Have a written procedure for employees to communicate concerns about quality and safety to management to investigate employee complaints. Incorporate corrective or preventive actions into the laboratory Quality Management plan.

9. Authorize the CAP to release its inspection and proficiency testing data to the appropriate regulatory oversight agencies such as CMS, Joint Commission, UNOS, or state/provincial agencies.

10. Submit only documentation and other materials to CAP that have been de-identified of all protected health information (PHI) in accordance with the requirements of the Health Insurance Portability and
Accountability Act of 1996 and its implement regulations, see 45 C.F.R. 164.514(b) unless the laboratory must submit PHI to CAP in order to respond to a deficiency or patient complaint.

11. Refrain from copying or distributing the CAP Checklists or any content thereof except for use by inspectors in conduction a CAP inspection and by the laboratory in preparing for such an inspection.

12. Adherence to the Terms of Use for the CAP Certification Mark of accreditation.

13. CAP must be notified if any of the following occurs:
   a. Investigation of the laboratory by a government entity or other oversight agency, or adverse media attention related to laboratory learns of an investigation or adverse media attention. This is to include and complaint investigations conducted or warning letters issued by any oversight agency (i.e. CMS, State Department of Health, The Joint Commission, FDA, OSHA).
   b. CAP must be notified as soon as laboratory finds itself to be subject to validation inspection.
   c. Discovery of actions by laboratory personnel that violate national state or local regulations.

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Approved by:

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Date

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Chief Executive Officer-UTMC

Date

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Policies Superseded by This Policy: None