§ 90-18.4. Limitations on clinical pharmacist practitioners.

(a) Any pharmacist who is approved under the provisions of G.S. 90-18(c)(3a) to perform medical acts, tasks, and functions may use the title "clinical pharmacist practitioner". Any other person who uses the title in any form or holds himself or herself out to be a clinical pharmacist practitioner or to be so licensed shall be deemed to be in violation of this Article.

(b) Clinical pharmacist practitioners are authorized to implement predetermined drug therapy, which includes diagnosis and product selection by the patient's physician, modify prescribed drug dosages, dosage forms, and dosage schedules, and to order laboratory tests pursuant to a drug therapy management agreement that is physician, pharmacist, patient, and disease specific under the following conditions:

1. The North Carolina Medical Board and the North Carolina Board of Pharmacy have adopted rules developed by a joint subcommittee governing the approval of individual clinical pharmacist practitioners to practice drug therapy management with such limitations that the Boards determine to be in the best interest of patient health and safety.

2. The clinical pharmacist practitioner has current approval from both Boards.

3. The North Carolina Medical Board has assigned an identification number to the clinical pharmacist practitioner which is shown on written prescriptions written by the clinical pharmacist practitioner.

4. The drug therapy management agreement prohibits the substitution of a chemically dissimilar drug product by the pharmacist for the product prescribed by the physician without the explicit consent of the physician and includes a policy for periodic review by the physician of the drugs modified pursuant to the agreement or changed with the consent of the physician.

(c) Clinical pharmacist practitioners in hospitals and other health facilities that have an established pharmacy and therapeutics committee or similar group that determines the prescription drug formulary or other list of drugs to be utilized in the facility and determines procedures to be followed when considering a drug for inclusion on the formulary and procedures to acquire a nonformulary drug for a patient may order medications and tests under the following conditions:

1. The North Carolina Medical Board and the North Carolina Board of Pharmacy have adopted rules governing the approval of individual clinical pharmacist practitioners to order medications and tests with such limitations as the Boards determine to be in the best interest of patient health and safety.

2. The clinical pharmacist practitioner has current approval from both Boards.

3. The supervising physician has provided to the clinical pharmacist practitioner written instructions for ordering, changing, or substituting drugs, or ordering tests with provision for review of the order by the physician within a reasonable time, as determined by the Boards, after the medication or tests are ordered.

4. The hospital or health facility has adopted a written policy, approved by the medical staff after consultation with nursing administrators, concerning the ordering of medications and tests, including procedures for verification of the clinical pharmacist practitioner's orders by nurses and other facility employees and such other procedures that are in the best interest of patient health and safety.

5. Any drug therapy order written by a clinical pharmacist practitioner or order for medications or tests shall be deemed to have been authorized by the physician approved by the Boards as the supervisor of the clinical
pharmacist practitioner and the supervising physician shall be responsible for authorizing the prescription order.

(d) Any registered nurse or licensed practical nurse who receives a drug therapy order from a clinical pharmacist practitioner for medications or tests is authorized to perform that order in the same manner as if the order was received from a licensed physician. (1999-290, s. 3.)