61-02-01-03. Pharmaceutical compounding standards. The minimum standards and technical equipment to be considered as adequate shall include:

1. Definitions.
   a. "Active chemical or ingredient" refers to chemicals, substances, or other components of articles intended for use in the diagnostics, cure, mitigation, treatment, or prevention of diseases.
   b. "Aseptic processing" is the method of preparing pharmaceutical and medical products that involves the separate sterilization of the product and of the package, the transfer of the product into the container and closure of the container under ISO class 5 or superior conditions, and using procedures designed to preclude contamination of drugs, packaging, equipment, or supplies by micro-organisms during the process.
   c. "Beyond-use date" refers to the date placed on preparation label that is intended to indicate to the patient or caregiver a time beyond which the contents of the preparation are not recommended to be used. The beyond-use date is determined from the date and time compounding of the preparation is completed.
   d. "Component" is any ingredient used in the compounding of a drug product, including any that are used in its preparation, but may not appear on the labeling of such a product.
   e. "Compounded sterile preparation" (CSP) will include all of the following:
      (1) Preparations prepared according to the manufacturer's labeled instructions and other manipulations when manufacturing sterile products that expose the original contents to potential contamination.
      (2) Preparations containing nonsterile ingredients or employing nonsterile components or devices that must be sterilized before administration.
      (3) Biologics, diagnostics, drugs, nutrients, and radiopharmaceuticals that possess either of the above two characteristics, and which include baths and soaks for live organs and tissues, implants, inhalations, injections, powders for injection, irrigations, metered sprays, and ophthalmic preparations.
   f. "Compounder or compounding personnel" is the pharmacist or other licensed or registered health care professional responsible for preparing the compounded preparations.
   g. "Compounding" is the preparation, mixing, assembling, packaging, and labeling of a drug or device in accordance to a licensed practitioner's prescription or medication order. Compounding does not include tablet splitting, reconstitution of oral or topical products as intended by the manufacturer, or repackaging of nonsterile dosage forms for redistribution, dispensing, or administration. Compounding includes:
      (1) Preparation of drugs or devices in anticipation of prescription drug orders based on routine, regularly observed prescribing patterns.
      (2) The addition of one or more ingredients to a commercial product as a result of a licensed practitioner's prescription drug order.
      (3) Preparation of drugs or devices for the purposes of, or as an incident to, research, teaching, or chemical analysis.
      (4) Categories of compounding.
         (a) Nonsterile simple. Should be conducted according to USP chapter 795.
         (b) Sterile compounds. Risk levels of compounded sterile preparations. Risk levels are assigned according to the corresponding probability of contaminating a preparation with microbial organisms, spores, and endotoxins, or chemical and physical contamination such as foreign chemicals and physical matter.
Preparations should be compounded according to USP chapter 797 based on the appropriate risk level.
(c) Radiopharmaceuticals. See article 61-05.
(d) Veterinary pharmaceuticals. Standards for veterinary pharmaceuticals are consistent with all parts of section 61-02-01-03.

h. "Compounding supervisor" is a person who supervises and is responsible for the compounding and dispensing of a nonsterile or sterile preparation. This may be the pharmacist on duty or the pharmacist-in-charge.

i. "Critical site" is a location that includes any component or fluid pathway surfaces (such as injection ports) or openings (such as opened ampules or needle hubs) exposed and at risk of direct contact with air, moisture, or touch contamination.

j. "Direct and contiguous compounding area" refers to the specific area where a compound is prepared.

k. "Disinfection" is the process by which the total number of micro-organisms is reduced to a safe level or eliminated by applying an agent to inanimate objects that destroys disease-causing pathogens or other harmful micro-organisms but may not kill bacterial and fungal spores.

l. "Hazardous drug" is one of those which studies in animals or humans indicate that exposures to them have a potential for causing cancer, development, or reproductive toxicity or harm to organs.

m. "ISO class" is a description of an atmospheric environment characterized by the number of particles of 0.5 microns or larger, within a cubic foot of air. "ISO class 5" atmospheric environment contains less than 100 particles, 0.5 microns or larger in diameter, per cubic foot of air.

n. "Media fill test" refers to tests used to validate aseptic techniques of compounding personnel and of processes that ensure the personnel and processes used are able to produce sterile products without microbial contamination. Testing uses a microbiological growth medium to substitute for actual drug product to simulate admixture compounding in determining the quality of a person's technique.

o. "NDC number" is the national drug code given to each drug separately and specifically approved by the food and drug administration for identification and reporting.

p. "Preparation" is a drug dosage form, dietary supplement, or a finished device. It contains one or more substances formulated for use on or for the patient or consumer.

q. "Primary engineering control (PEC)" refers to a device or room that provides an ISO class 5 or superior environment during the compounding process, including laminar airflow workbenches (LAFWs), biological safety cabinets (BSCs), compounding aseptic isolators (CAIs), and compounding aseptic containment isolators (CACIs).

r. "Product" is a commercially manufactured drug or nutrient that has been evaluated for safety and efficacy by the food and drug administration, accompanied by full prescribing information.

s. "Repackaging" is the transfer of an ingredient from one container to another.

t. "Risk levels" of CSPs determine the level assigned that represent the probability that it will be contaminated with microbial organisms, spores, endotoxins, foreign chemicals, or other physical matter.

u. "Seventy percent sterile isopropyl" or IPA is an antimicrobial used to clean surfaces used in sterile preparations.
v. "Stability" means the extent to which a preparation retains, with specified limits, and throughout its period of storage and use, the same properties and characteristics it possessed at the time of compounding.

w. "US Pharmacopeia (USP)" is the book of official compendia of standards for the United States.

2. General compounding.
   a. Responsibility of the compounder.
      (1) Personnel engaging in compounding must be proficient, capable, and qualified to perform assigned duties in the compounding area while expanding the individual's knowledge of compounding through seminars or appropriate literature.
      (2) Compounding personnel must be familiar with USP standards and North Dakota regulations, including:
         (a) Certifying all prescriptions orders.
         (b) Approving or rejecting all components, drug product containers, closures, in-process materials, and labeling ensuring preparations and ingredients are of acceptable strength, quality, and purity, with appropriate packaging.
         (c) Preparing and reviewing all compounding records to assure that errors have not occurred in the compounding process and the finished product has expected qualities as well as implementing procedures to prevent cross-contamination.
         (d) Assuring the proper maintenance, cleanliness, sanitation, and use of all equipment used in prescription compounding practice, including the direct and contiguous compounding area allowing for the compounding environment to be suitable for its intended purpose.
         (e) Assuring that the drug product and components of drug products are not on the list of federally recognized drug products that have been withdrawn or removed from the market for public health reasons.
      (3) Policies and procedures must be established concerning washing and donning the appropriate clothing specific to the type of process performed to protect the personnel from chemical exposures and prevent drug contamination.

   b. Training. All compounding supervisors and all personnel involved in compounding must be well trained and must participate in current, relevant training programs. All training activities will be covered by standard operating procedures and must be properly documented. Steps in the training procedure include:
      (1) Be familiar with pharmaceutical compounding and nonsterile compounding (USP 795), pharmaceutical compounding and sterile compounding (USP 797), hazardous drug compounding (USP 800), and pharmaceutical calculations in prescription compounding (USP 1160).
      (2) Be familiar with all procedures relating to compounding specific to the individual's facility, equipment, personnel, compounding process, evaluation, packaging, storage, and dispensing.
      (3) Compounding supervisors must be responsible to follow the instructions below to show that personnel are appropriately trained:
         (a) Demonstrate compounding procedures to compounding personnel.
         (b) Guide personnel through the compounding process with assistance.
         (c) Observe personnel performing a compound without assistance but under supervision.
         (d) Review the compound, correct mistakes, and answer questions concerning compounding and associated processes.
(e) Confirm verbal and functional knowledge of the personnel concerning compounding.

(f) Have personnel perform a compounding procedure without supervision, yet checking off the final preparation.

(g) If properly compounded and when satisfied, sign the documentation records confirming appropriate training.

(h) Continually monitor the work of the personnel, including calculations.

(4) The pharmacist on duty and the pharmacist-in-charge are ultimately responsible for the finished product.

c. Procedures and documentation. Procedures must be developed for the facility, equipment, personnel, preparation, packaging, and storage of the compounded preparation to ensure accountability, accuracy, quality, safety, and uniformity in compounding. This allows for a compounder, whenever necessary, to systematically trace, evaluate, and replicate the steps included throughout the preparation process of a compounded preparation.

d. Nonsterile drug compounding must meet the facility, equipment, packaging, storage, and beyond-use date standards set in USP chapter 795. Policies and procedures should be developed to ensure compliance with those standards.

e. Compounding controls for nonsterile preparations.

1. The compounder must ensure that the written procedures for compounding are available electronically or in hard copy and assure the finished products have the correct identity, strength, quality, and purity.

2. Procedures must be established that give a description of the following:
   a. Components and their amounts.
   b. Order of component additives.
   c. Compounding process.
   d. Drug product.
   e. Required equipment and utensils, including container and closure systems.

3. The compounder will accurately weigh, measure, and subdivide all components as appropriate.
   a. The compounder must check and recheck each procedure at each point of the process to ensure that each weight or measure is correct.
   b. If a component is transferred from the original container to another, the new container must be identified with the component, name, weight or measure, the lot or control number, the expiration or beyond-use date, and the transfer date.

4. The compounder must write procedures that describe the tests or examinations that prove uniformity and integrity of the compounded preparations.

5. Control procedures must be established to monitor the output and validate the performance of compounding personnel that affect variability of final preparations, such as:
   a. Capsule weight variation.
   b. Adequacy of mixing to assure uniformity and homogeneity.
   c. Clarity, completeness, or pH of solutions.

6. The compounder must establish an appropriate beyond-use date for each compounded preparation.

7. Facilities engaging in compounding must have a specifically designated and adequate space for orderly compounding, including the placement and storage of equipment and materials.
f. Labeling of nonsterile preparations.
   (1) The compounder's preparation label must contain all information required by
   North Dakota state law and accepted standards of practice found under chapter
   61-04-06, prescription label requirements, plus the beyond-use date and assigned
   lot number.
   (2) The compounder must label any excess compounded products so as to refer to
   the formula used.
   (3) Preparations compounded in anticipation of a prescription prior to receiving a
   valid prescription should be made in a regularly used amount based on the history
   of prescriptions filled and they should be labeled with:
      (a) Complete list of ingredients or preparation time and reference or established
          chemical name or generic name.
      (b) Dosage form.
      (c) Strength.
      (d) Preparation date and time.
      (e) Inactive ingredients.
      (f) Batch or lot number.
      (g) Assigned beyond-use date.
      (h) Storage conditions.
   (4) The compounder must examine the preparation for correct labeling after
   completion.

g. Records and reports for nonsterile preparations.
   (1) Records must be maintained, including a hard copy of the prescription with
       formulation and compounding records.
   (2) Adequate records of controlled substances used in compounds.
   (3) All records must be kept for five years according to North Dakota state law and be
       available for inspection.
   (4) Formulation record provides a consistent source document for preparing the
       preparation to allow another compounder to reproduce the identical prescription at
       a future date and must list:
       (a) Name, strength, and dosage form of the preparation compounded.
       (b) All ingredients and their quantities.
       (c) Equipment needed to prepare the preparation, when appropriate.
       (d) Mixing instructions including order of mixing, mixing temperatures, and other
          valid instructions, such as duration of mixing.
       (e) Assigned beyond-use date.
       (f) Container used in dispensing.
       (g) Storage requirements.
       (h) Any quality control procedures.
   (5) Compounding record documents the actual ingredients in the preparation and the
       person responsible for the compounding activity and includes:
       (a) Name and strength of the compounded preparation.
       (b) The formulation record reference.
       (c) Sources and lot numbers of the ingredients.
       (d) Total number of dosage units compounded.
       (e) Name of compounding personnel who prepared the preparation.
       (f) The date of preparation.
(g) The assigned internal identification number, lot number, and prescription numbers.

(h) Assigned beyond-use date.

(i) Results of all quality control procedures.

(6) Temperature log records the daily monitoring of temperatures in the storage area specifically for the controlled room temperature, refrigerator, freezer, or incubator.

3. Nonsterile compounding. Compounders are to use the following steps to minimize error and maximize the prescriber's intent, specifics can be found in pharmaceutical compounding - nonsterile compounding (USP 795):
   a. Judge the suitability of the prescription of the preparation in terms of safety and intended use.
   b. Perform necessary calculations to establish the amounts of ingredients needed.
   c. Identify equipment and utensils needed.
   d. Don the proper attire and properly wash hands and arms.
   e. Clean the compounding area and needed equipment.
   f. Only one prescription can be compounded at a time in the specified compounding area.
   g. Assess weight variation, adequacy of mixing, clarity, odor, color consistency, and pH as appropriate of the completed preparation.
   h. Annotate the compounding and formulation records.
   i. Label the prescription containers appropriately.
   j. Sign and date the prescription or compounding record affirming that all procedures were carried out to ensure uniformity, identity, strength, quantity, and purity.
   k. Thoroughly clean all equipment immediately when finished.

4. Compounding process for compounded sterile preparations. Compounders are to follow the USP chapter 797 standards and use the following steps to minimize error and maximize the prescriber's intent:
   a. Judge the suitability of the prescription for the compounded sterile preparation in terms of safety and intended use.
   b. Perform necessary calculations to establish the amounts of ingredients needed.
   c. Identify equipment and utensils needed for the preparation of the compounded sterile preparation.
   d. Sterile compounding areas and critical areas must be structurally isolated from other areas designated to avoid unnecessary traffic and airflow disturbances according to USP chapter 797, separate from nonsterile compounding areas, and restricted to qualified compounding personnel.
   e. Policies and procedures must be established in accordance with USP chapter 797 for personnel cleaning and garbing for protection and avoidance of containment.
   f. Clean and sanitize the compounding area and needed equipment according to USP chapter 797.

5. Facilities for sterile compounding should conform with USP chapter 797.

6. Equipment specific for sterile compounding should conform with USP chapter 797.


8. Suitable current reference sources either in book or electronic data form (available in the pharmacy or online) which might include the United States Pharmacopeia and National Formulary, the United States Pharmacopeia Dispensing Information, Facts & Comparisons, Micro Medex, the ASHP Formulary, Clinical Pharmacology or other suitable references determined by the board which are pertinent to the practice carried on in the licensed pharmacy.
9. Compounding for office use.
   a. It is acceptable to compound human drug products to be used by North Dakota practitioners in their office for administration to patients provided they are prepared by a facility licensed as an outsourcing facility in accordance to North Dakota Century Code section 43-15.3-13 or by a resident North Dakota pharmacy.
   b. It is acceptable for any licensed pharmacy to compound veterinary drug products to be used by veterinarians in their office for administration to client's animals. These compounded office use products may be dispensed to clients for use in a single treatment episode, not to exceed a one hundred twenty-hour supply.
   c. Sales to other pharmacies, veterinarians, clinics, or hospitals are manufacturing and are not allowed. It is the responsibility of the pharmacy and pharmacist involved in the compounding to ensure compliance with this section for the products they compound.

10. Compounding of Hazardous drugs as compounded sterile products (CSPs).
   a. Hazardous drugs, when prepared for administration only, shall be prepared under conditions that protect the health care worker and other personnel in the preparation and storage areas according to USP chapter 800. Appropriate personnel protective equipment shall be worn when compounding hazardous drugs according to USP chapter 800.
   b. Hazardous drugs shall be stored and prepared separately from other nonhazardous drugs in a manner to prevent contamination and personnel exposure according to USP chapter 800.
   c. Hazardous drugs shall be handled by the pharmacy according to USP chapter 800 with caution at all times using appropriate chemotherapy gloves during receiving, distribution, stocking, inventorying, preparation for administration, and disposal.
   d. Hazardous drugs shall be prepared in an ISO class 5 environment with protective engineering controls in place and following aseptic practices specified for the appropriate contamination risk levels specified in this chapter.
   e. All hazardous drugs shall be prepared in a biological safety cabinet (BSC) or a compounding aseptic containment isolator (CACI). The BSC or CACI shall be placed in an ISO class 7 area that is physically separated (i.e., a different area from other preparation areas) and with negative pressure to adjacent positive pressure ISO class 7 or better anteareas. If the CACI is used outside of a buffer area, the compounding area shall maintain a minimum negative pressure of 0.03 inch water column and have a minimum of twelve air changes per hour.
   f. All personnel who compound hazardous drugs shall be fully trained in the storage, handling, and disposal of these drugs according to USP chapter 800. This training shall occur prior to preparing or handling hazardous drugs and this training shall be by testing specific hazardous drug-handling techniques. Such training shall be documented for each person at least annually.

The state board of pharmacy recognizes that the equipment needed will depend on the type of pharmaceutical services offered, and therefore, variations for required equipment may be granted by the state board of pharmacy.

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