

**657—20.2(124,126,155A) Definitions.** For purposes of this chapter, the following definitions apply:

“*Anticipatory compounding*” means the compounding of preparations in advance of the pharmacy’s receipt of patient-specific prescriptions.

“*Batch preparation compounding*” means anticipatory compounding, compounding preparations intended for multiple disbursements, or compounding preparations in a multiple-dose container for administration to more than one patient.

“*Beyond-use date*” means the date after which a compounded preparation should not be used, determined from the date that the preparation is compounded.

“*Bulk drug substance*” means any substance that is represented for use in a drug and that, when used in the manufacturing, processing, or packaging of a drug, becomes an active ingredient or a finished dosage form of the drug. The term does not include intermediates used in the synthesis of such substances.

“*Compounding*” means the combining, mixing, diluting, pooling, flavoring, or otherwise altering of a drug or bulk drug substance to create a drug. Compounding includes the preparation of drugs or devices in which all bulk drug substances and components are nonprescription products. Compounding does not include the use of a flavoring agent to flavor a drug pursuant to rule 657—20.13(124,126,155A), nor does it include mixing or reconstituting a drug according to the product’s manufacturer label.

“*FDA*” means the Food and Drug Administration of the U.S. Department of Health and Human Services.

“*Flavoring agent*” means a therapeutically inert, nonallergenic substance consisting of inactive ingredients that is added to a drug to improve the drug’s taste and palatability.

“*Office use*” means that a compounded product has been prepared and distributed to a practitioner for administration to a patient by the practitioner in the course of the practitioner’s professional practice. A compounded product distributed to a practitioner for “office use” shall not require a patient-specific prescription and may not be further distributed to another practitioner or dispensed to a patient for self-administration.

“*Outsourcing facility*” or “*facility*” means any compounding facility that is registered as an outsourcing facility, as defined in 21 U.S.C. Section 353b, that distributes sterile compounded human drug products without a patient-specific prescription to an authorized agent or practitioner in this state.

“*USP*” means United States Pharmacopeia.

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