

Standards for Pharmacy Compounding of Non-sterile Preparations Information Sheet

Dates to Remember

Phase one	October 1, 2020
Phase two	October 1, 2020
Phase three	January 1, 2021
Phase four	April 1, 2021

Important Resources

Where do I start? Guidance Document

NAPRA Guidance Document

Pharmacy Quality Assurance Self-Assessment

Please note: The Pharmacy Quality Assurance Self-Assessment is a tool for gap analysis and should not be submitted to the College for approval or review.

Decision Algorithm for Risk Assessment

The following phases are outlined in the NAPRA_Model Standards for Pharmacy Compounding of Non-sterile Preparations

Phase one

Section 4 - Assessing risk for compounding non-sterile preparations Section 5 - Requirements for all levels of non-sterile compounding activities Section 6.5 - Conduct of personnel Section 8.1 - Level A requirements Phase two Section 6.1 - BUD and dating methods Section 6.2 - Master formulation record Section 6.3 - Ingredients used for compounding Section 6.4 - Compounding record

Phase three

Section 6.6 - Verification

Section 6.7 - Labeling and packaging

Section 6.8 - Storage

Section 6.9 - Transport and delivery

Section 6.10 - Product recalls

Section 6.11 - Incidents and accidents

Section 7 - Quality Assurance

Section 8.2 - Level B requirements

Phase four

Section 9 - Requirements for hazardous preparation

Section 8.3 - Level C requirements