



# 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