

WRHA PHARMACY PROGRAM - VERIFIER QUALITY ASSURANCE RECORD FOR **BATCH REPACKAGING**

FOR MEDICATION OCCURRENCES NOT REACHING THE PATIENT

Date Occurred _____	Site (circle one):								Service Area _____
	Access Transcona	Conc	DLC	GH	HSC	RHC	SOGH	VGH	
Generic Drug Name		Strength			Form				
Batch Repackaging Format (circle ONE):		Repackaging Type (check ONE):				Discovered on Verification of (circle ONE):			
Oral Topical Injectable		<input type="checkbox"/> Strip pack (Euclid, Automed) (01*) <input type="checkbox"/> Blister card/pack or MediDose (02*) <input type="checkbox"/> IV/PO syringe (03*) <input type="checkbox"/> Vial (tabs, caps, powders) (04*) <input type="checkbox"/> Bottle or cup (liquids) (05*) <input type="checkbox"/> Ointment pot (06*) <input type="checkbox"/> Other (specify) _____(07*)				Batch set-up Final product (contents, quantity, quality, container, seal, barcode, label)			
Batch or Batch Record Error Code Key** (circle all that apply):					Label Error Code Key** for batch or final labels (circle all that apply):				
A. Wrong batch record, repackaging card, or compounding recipe B. Wrong drug, other ingredient, or DIN C. Expired drug or other ingredient D. Defective or contaminated drug or ingredient E. Wrong strength or concentration F. Wrong quantity (volume, weight, number of doses, amount of finished product in batch) G. Wrong dosage form H. Wrong lot number documented on batch record I. Expiry date assigned/documentated incorrectly on batch record J. Wrong barcode selected K. Wrong type of container selected L. Container defect (leak, crack, tear, puncture) M. Unacceptable quality of final product (e.g., prepared incorrectly, not dissolved or suspended or distributed uniformly) N. Multiple batches in one batch bin O. Other (describe in Description of Variance)					P. Illegible (not readable) Q. Wrong patient name R. Wrong patient location S. Wrong barcode T. Wrong drug name U. Wrong strength/volume/weight/concentration V. Wrong instructions (frequency, route) W. Incorrect/missing auxiliary label X. Wrong date of preparation/distribution Y. Wrong expiry date Z. Wrong prescriber AA. Wrong label format or font BB. Wrong lot or batch number CC. Wrong manufacturer or manufacturer abbreviation DD. Not labeled (label missing) EE. Other (describe in Description of Variance)				
Description of the Variance:									
Verifier Signature:									

* For audit purposes

** Codes in BOLD meet the definition of Critical Error (2010 MPhA TVT Framework)