Schedule to License to Operate Pharmacy Conditions Pharmacy Accuracy Checking Technician Activities

In accordance with the Licence to Operate Pharmacy, every pharmacy must comply with the requirements of the Health and Disability Service Standards Pharmacy Services Standard NZS 8134.7:2010.3. As the role of the PACT is not defined within the Standards, Medsafe issues a Licence to Operate Pharmacy with an additional condition to enable a certified PACT to independently conduct accuracy checks as follows:

- final accuracy check of a dispensed prescription
- accuracy check of a repackaged medicine
- accuracy check of a de-blistered medicine
- accuracy check of each ingredient for a non-aseptically compounded medicine.

When conducting these activities, the following requirements of Standard 5, Health and Disability Services Pharmacy Services Standard NZS 8134.7:2010, may be completed by a PACT, subject to the following limitations:

STANDARD 5.2: GOOD DISPENSING PRACTICE A disciplined dispensing procedure shall ensure that the appropriate product is selected and dispensed accurately and efficiently.	
Criteria 5.2.3 (f)	All prescription forms clearly record who dispensed the prescription and the pharmacist responsible for the final check for completeness and accuracy
Limitation 5.2.3(f)	 The prescription form must record the identity of: the pharmacist conducting the clinical check, and the PACT responsible for conducting the final accuracy check.
STANDARD 5.15: COMPOUNDING PROCEDURES Compounding procedures shall ensure that the medicines prepared are of the required quality	
Criteria 5.15.5	The identity, and weight (or volume) of each ingredient shall be checked by a pharmacist
Limitation 5.15.5	The identity of the PACT accuracy checking the identity and weight (or volume) of each ingredient must be recorded on the compounding document. Note. Reconstitution of antibiotics is not considered compounding.
STANDARD 5.29: RECORDS FOR For regularly repacked pr make the record.	REPACKING OF MEDICINES roducts a copy of the master batch document shall be used to
Criteria 5.29.1(j)	The signature or initials of the pharmacist checking/releasing the finished product shall be recorded
Limitation 5.29.1 (j)	The identity of the PACT accuracy checking/releasing the finished repackaged product must be recorded on the batch document.

specifications and releas	step where the finished (repackaged)product is compared with its sed or rejected
Criteria 5.34.1	The requirements in 5.25 shall be met: 5.25.1 – the decision to release the final product shall be made by the pharmacist taking responsibility for quality. 5.25.2 - the assessment and decision shall be recorded
Limitation 5.34.1	A PACT may make the decision to release a <u>repackaged</u> product. Compliance with criterion 5.25.2 is required.
STANDARD 5.39: DE-BLISTERIN Documented and systen from their original strip po	natic procedures shall be followed when medicines are removed
Criteria 5.39.2(e)	The batch record shall include the identity of the pharmacist checking the end product
Limitation 5.39.2(e)	The identity of the PACT accuracy checking de-blistered medicines must be recorded on the batch record sheet. Note. The new expiry date must be assigned by a pharmacist, refer 5.39.2(f).
Criteria 5.39.6	Only one medicine shall be de-blistered at a time, with the process finished and checked off by the pharmacist before another medicine de-blistering process begins.
Limitation 5.39.6	Only one medicine shall be de-blistered at a time, with the process finished and checked off by the PACT, before another medicine de- blistering process begins.
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