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1. Introduction

This document:

a. Has been compiled for Pharmacies when claiming reimbursement for services;

b. Should be read in conjunction with:

i. The Pharmaceutical Schedule

ii. The Pharmaceutical Transactions Data Specification (in relation to matters concerning file formats and data to be provided for processing purposes), and

iii. The Pharmacy Services Agreement for the provision of pharmacy services that came into effect on 1 March 2010.

c. For the purpose of reimbursement, the order of priority for the above documents is as follows:

i. The Pharmaceutical Schedule;

ii. The Pharmaceutical Transactions Data Specification (in relation to matters concerning file formats and data to be provided for processing purposes);

iii. The Pharmacy Services Agreement or section 88 notice for the provision of pharmacy services;

iv. This Procedures Manual.

2. Specific Point to Note

2.1 Pharmacy Closure or Change of Ownership

Pharmacy closure or change of ownership, guidance on appropriate procedures is available from your DHB or Sector Services Agreement Administration phone 0800 281 222 Option 4, then 1

2.2 Online Claiming

Pharmacies which are currently claiming manually and are interested in claiming electronically should liaise with Sector Services Contact Centre. For further information, phone 0800 353 2425.
3. Submission of Batches

3.1 Verification of Claims

All claim items submitted by a Pharmacy must:

a. Be supported by an original prescription form filed in date of dispensing, then unique transaction number (the batch). Any form received by the pharmacy at a later date must be inserted in the original batch containing the corresponding date of dispensing and be filed in date of dispensing, then unique transaction number.

b. Meet all legal and contractual requirements.

Variances between the original prescription and the computer record or supply must be clearly annotated on the prescription form for clarification.

Each batch must be accompanied by a coversheet (in the form approved by Sector Services from time to time) which is to be completed in full and signed on behalf of the pharmacy.

3.2 Date for Submission of Batches

3.2.1 Diskette and On Line Claimants

Pharmacies may retain batches for five months. After the five-month period, batches must be submitted to Sector Services once or twice each month. Batch submission must reflect the claim period of the claim file previously submitted.

3.3 Batch Delivery Instructions

The delivery address for batches Manual Pharmacy Claims, Diskettes and Invoices (unless otherwise advised by Sector Services) is:

Sector Services
179 St Hill Street
WHANGANUI 4500

The delivery address for batches to be archived (unless otherwise advised from Sector Services) is:

Archive Pharmacy Claims
137 London Street
WHANGANUI 4500

Note: If sending diskettes with batches please put diskette at top of batch and clearly annotate “Disk enclosed”.

3.4 Audit

On occasion, our agent may require batches to be submitted early for audit purposes. Pharmacies will be notified when this is necessary and the notified pharmacy shall comply with the time frames and delivery requirements specified by our agent.
4. Legal and Subsidy Issues

Procedure:

For prescriptions other than Controlled Drugs the following is a checklist of the legal requirements that must be on the prescription when it is presented to you for dispensing. Prescriptions submitted for payment must meet certain legal requirements. (For Controlled Drugs – refer to Section 4.3).

Checklist:

- Prescriber’s signature
- Prescribing date
- Prescriber’s physical address
- Title, surname and initials of the patient
- Physical address of the patient
- Date of birth (if the prescription is for a child under 13 years)
- Name of the medicine
- Strength of the medicine (where appropriate)
- Dose and frequency of the dose, for an internal medicine
- Method and frequency of use for an external medicine
- Total quantity to be supplied as a single supply or on each dispensing e.g. 90 tablets or 30 tablets

If the medicine is to be supplied more than once the pharmacist must ensure the prescription is annotated with the following information:

- The number of times the medicine can be supplied; or
- The interval between each supply; or
- The total period of treatment, e.g. 30 x 3 or 30 per month or 3 month’s supply.

The following are the legal requirements, which must be added by the pharmacy:

- Pharmacy stamp – per form
- Dispensing date – per item
- Prescription number (Unique Identifying Number) – per item

4.1 Prescription Requirements

4.1.1 Prescriber Information

In addition to the legal requirements of a prescription, the following information is required for subsidy purposes:

a. Signature

A facsimile signature is not acceptable. Subject to the conditions below, if a prescription is faxed, the original prescription must be obtained or the prescriber must indelibly sign the faxed copy before a claim can be made for the prescription item/s.
However, if the original prescription (or the faxed copy signed by the prescriber as above) has not been received by the pharmacy within four weeks of the date of the original dispensing, reimbursement can be claimed. The signed prescription must be obtained and submitted in due course (see clause 3.2 and 3.3 of the Procedures Manual) with the batch for audit purposes.

If no signed prescription is obtained by the batch submission date then the pharmacy must refund any money previously claimed in respect of this claimed item by crediting the amount against its next claim(s).

If the pharmacy considers that special circumstances apply to a specific claim item it may apply in writing to the DHB. The DHB may, at its discretion allow the pharmacy to retain payment. If no such express written permission is provided then the refund must be made by the pharmacy.

b. Legibility

The prescription must be legibly and indelibly printed and cannot be written in pencil. A reprint of the label for the item attached to a prescription form is not acceptable for claiming payment.

c. Prescriber’s address

The prescriber’s address must be specified and clear. This must be the practice address, not a PO Box or Rural Delivery number. A rural grid number is acceptable.

4.1.2 Patient Information

In addition to the legal requirements of a prescription as set out under clause 4 - Procedure, the following Patient information is required for subsidy purposes:

a. Patient address

The address of the Patient must be specified. This must be the residential address, not a PO Box or Rural Delivery number. A rural grid number is acceptable.

b. Patient category

Patient eligibility must be clearly identified in accordance with the Pharmaceutical Schedule. If the prescriber has included this information on the prescription, the pharmacy may accept these details as correct unless the Patient provides documented evidence to the contrary.

Information on eligibility can be found on the MoH web site at: http://www.moh.govt.nz/eligibility

An “H” code is used for a Patient who is usually a resident in the Hokianga Ward of the Far North District. The prescription must be written by a registered medical practitioner (or other legally authorised prescriber) employed by, and on a form supplied by, the Hokianga Health EnterpriseTrust.
### 4.1.3 Procedure

The patient categories and pharmaceutical co-payments as at 1 September 2008 are:

**Youth (ages 0 to 5 years)**

<table>
<thead>
<tr>
<th>CSC or PHO Status</th>
<th>HUHC Holder / Care Plus Patient</th>
<th>Patient Category</th>
<th>Maximum Pharmaceutical Co-payment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>No PSC</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>With a PSC</td>
</tr>
<tr>
<td>Low-cost PHO Enrolee; Or Eligible person and eligible provider/prescriber</td>
<td>Yes</td>
<td>Y4Z</td>
<td>$0</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>Y4</td>
<td>$0</td>
</tr>
<tr>
<td></td>
<td></td>
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<tr>
<td></td>
<td>No</td>
<td>Y3</td>
<td>$0</td>
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</table>

**Junior (ages 6 to 17 years)**

<table>
<thead>
<tr>
<th>CSC or PHO Status</th>
<th>HUHC Holder / Care Plus Patient</th>
<th>Patient Category</th>
<th>Maximum Pharmaceutical Co-payment</th>
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</thead>
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<tr>
<td></td>
<td></td>
<td></td>
<td>With a PSC</td>
</tr>
<tr>
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</table>
## Adult (ages 18 and above)

<table>
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<tr>
<th>CSC or PHO Status</th>
<th>HUHC Holder / Care Plus Patient</th>
<th>Patient Category</th>
<th>Maximum Pharmaceutical Co-payment</th>
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</tr>
<tr>
<td>Or Eligible person and eligible provider/prescriber</td>
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<td>$3</td>
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<tr>
<td></td>
<td>No</td>
<td>A4</td>
<td>$3</td>
</tr>
<tr>
<td>CSC Holder</td>
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<td>$3</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>A1</td>
<td>$3</td>
</tr>
<tr>
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</tr>
<tr>
<td></td>
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<td>A3</td>
<td>$15</td>
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</tbody>
</table>

Persons usually resident in the Hokianga Ward of the Far North District with a prescription issued by a registered medical practitioner or other legally authorised prescriber employed by, and on a form supplied by, the Hokianga Health Enterprise Trust

<table>
<thead>
<tr>
<th>CSC or PHO Status</th>
<th>HUHC Holder / Care Plus Patient</th>
<th>Patient Category</th>
<th>Maximum Pharmaceutical Co-payment</th>
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<tr>
<td></td>
<td></td>
<td></td>
<td>No PSC</td>
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<tr>
<td>Low-cost PHO Enrolee;</td>
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</tr>
<tr>
<td>CSC Holder</td>
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<td>$0</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>H1</td>
<td>$0</td>
</tr>
<tr>
<td>None of the Above</td>
<td>Yes</td>
<td>H3Z</td>
<td>$0</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>H3</td>
<td>$0</td>
</tr>
</tbody>
</table>
Glossary – for the purposes of clause 4.1

CSC means a Community Services Card

Eligible Person means a person eligible for publicly funded health services under the Eligibility Direction (refer to www.moh.govt.nz/eligibility) who is issued with a prescription.

An Eligible Prescriber is a prescriber who issues a prescription in the following circumstances:

i. when the prescriber is employed by a DHB (e.g. Public Hospital or community-based service);
ii. when the prescriber is subcontracted to a Primary Health Organisation;
iii. when the prescriber is employed by an After Hours provider with a service agreement with a DHB or a PHO;
iv. when the prescriber is providing a fully publicly funded service under a Section 88 notice alone.

Exclusions to the eligible provider or prescriber policy are:

a. Providers providing completely privately funded services;
b. Providers providing services under a Section 88 notice alone that are not solely publicly funded.

HUHC means a High Use Health Card

Low-cost PHO Enrolee means an eligible person who is enrolled in any practice in a Low Cost Primary Health Organisation (PHO) where the prescription has been issued by a prescriber working for the eligible person’s enrolling Low Cost PHO (unless local arrangements are in place).

Patient(s) For the purposes of this document, the term Patient(s) refers to the term Service User as defined in the Pharmacy Services Agreement.

PHO Status means whether or not the eligible person is enrolled in a Primary Health Organisation (PHO). Note: Care Plus patients will be coded either Y4Z, J4Z, A4Z or H4Z.

PSC means a Prescription Subsidy Card

Updating
The Crown may update the co-payments listed here from time to time. Pharmacies will be notified of these changes via the Pharmaceutical Schedule and/or directly by the Ministry of Health or DHBs.

The patient category codes may also be updated from time to time. Any changes to the patient category codes will be linked to an amendment of the Pharmaceutical Transactions Data Specification.

4.1.4 Pharmacy Stamp
As per the Medicine Regulations 1984, the pharmacy stamp is required to be on all prescriptions, supply orders and other types submitted for claiming purposes.

4.1.5 Date of Dispensing
The Date of Dispensing, until DHBs, pharmacy and software vendors have worked on an alternative solution, is the date on which a Prescription Item is processed within the pharmacy’s practice management system [in line with the side letter DHBs issued in December 2009].

The Date of Dispensing must be recorded on all prescriptions for which a subsidy is claimed. This record must be stamped, hand-written or recorded on the third-part label. The Date of Dispensing on the prescription including that on the third part label must be the same as the...
date in the computer record. If the third-part label is used as the only indication of Date of Dispensing the pharmacy has a responsibility to ensure that the quality of the label means that it is fixed to the prescription in a manner that will withstand multiple handling.

The Date of Dispensing must not precede the prescribing date. If the prescribing date returned on a signed telephone/faxed prescription is after the date of dispensing, for the purposes of payment, the signed prescription and the faxed forms must be stapled together or the date may be annotated by the pharmacist to explain the discrepancy between the prescribing date and the date of dispensing.

4.1.6 Prescription Number (Unique Identifying Number)
This numbering system applies to all prescriptions, Supply Orders, and other types.

The appropriate suffix is determined by the prescription. If the prescription is for a single supply (including those items dispensed stat), the suffix used is ‘0’. For the initial dispensing of a prescription where eligible repeats are prescribed, the suffix is ‘1’. Each subsequent dispensing of a repeat on a prescription has the next consecutive number as its suffix.

Prescription numbers should follow the following format:

123456789/1

The prescription number should be adjacent, where possible, to the relevant item on the original prescription form. Sometimes third part labels have a reference to the item, but an effort should be made to place the label next to the item. If working from a faxed or telephone copy, place the third part label on the copy then staple copy to the original when received.

Hand written, legible numbers are for emergency or exceptional circumstances only.

4.2 Specialist Recommendation

Definitions for Specialist, Retail Pharmacy–Specialist and Retail Pharmacy-Specialist Prescription, in relation to a Prescription, are detailed in the Pharmaceutical Schedule. A look-up facility is available on the Medical Council of New Zealand’s website (www.mcnz.org.nz) to attain a prescriber’s specialist status.

A Specialist Recommendation (within their scope of practice) should follow the format below:

“Recommended by [name of the specialist and year of authorisation]”

The recommendation must be written on the prescription form and signed or initialled by the prescriber. The authorisation is valid for two years.

Where the authorisation remains current during this two year period and this information is known to the pharmacist (and can be demonstrated on audit), the pharmacist can annotate the prescription with specialists name and year of authorisation. In these circumstances, the prescription does not need to be returned to the prescriber for annotation.

Prescriptions originating from hospitals for “specialist” prescription items (i.e. Retail Pharmacy- Specialist, and Hospital Pharmacy-Specialist) are deemed to have been prescribed by an appropriate specialist, irrespective of the status of the medical practitioner writing the prescription. For the purposes of the definition, it makes no difference whether or not the Specialist is employed by a hospital.
This however does not apply to prescriptions which are subject to the restrictions “Retail Pharmacy–Specialist Prescription” and “Hospital Pharmacy-Specialist Prescription”, where the Specialist must sign the prescription.

4.3 Controlled Drug Prescriptions

The following is a checklist of the legal requirements that must be on the prescription when it is presented for dispensing:

For CLASS B Controlled Drugs, the prescription must:

- Be written on a form provided by the Director General of Health.
- Be legibly and indelibly written in the Prescriber’s own handwriting.
- Be indelibly signed by the Prescriber personally with his/her usual signature.
- Include the date on which the Controlled Drug prescription is written.
- Include the address of the Prescriber (this can be stamped however the stamped address must be on all three copies of the Controlled Drug prescription).
- Include the surname, initials, and street address of the patient.
- Include the age in years and months (in words) if the patient is under 12 years.
- Include the name and total amount of the Controlled Drug to be dispensed and the number of occasions on which it may be dispensed.
- Set out the name of the Controlled Drug in full or be abbreviated only by the use of BP, BPC or other recognised titles.
- Include the dose and frequency of the dose for an internal use medicine, and for external use have the directions of use.
- Where the prescription has an unusual dose, or what may be regarded as a dangerous dose, the dose should be underlined and initialed by the Prescriber.
- Any alterations must be signed by the Prescriber.

In addition to the above, if the Class B Controlled Drug is Methadone and if the prescriber is authorised by the MOH or its delegate, or works in a place for the time being specified by the Minister under the Misuse of Drugs Act, the prescription Must be legibly and indelibly written, or in a form approved from time to time by the Director-General of Health.

Controlled Drug Prescriptions Written by a Midwife

- The Controlled Drug prescription must include the words “For midwifery use only.”
- Midwives can only prescribe pethidine.
- A prescription written by a midwife for pethidine must be first dispensed not more than 4 days after the date on which the prescription was written
- The medicine can not be supplied on more than 2 occasions.
- The total quantity prescribed cannot exceed 1 month.

Controlled Drug Prescriptions Written by a Dentist

- If the prescriber is a dentist the prescription must include the words “For dental treatment only”.
- A dentist can not write a prescription for a supply of a Controlled Drug in any quantity greater than 7 days.

In accordance with the Pharmaceutical Schedule, only a quantity sufficient to provide 5 days treatment will be reimbursed. The remaining two days are not subsidised.

If for special reasons relating to the protection of the patient or for limiting the quantity of any Controlled Drug in the possession of the patient, the prescriber directs daily dispensing or other dispensing intervals a Controlled Drug may be supplied on that number of occasions.
and not more frequently than the intervals indicated. The total quantity covered by such a prescription can not exceed one month..

**Controlled Drug Prescriptions Written by a Nurse Prescriber**

- A nurse prescriber can not write a prescription for a supply of a Controlled Drug in any quantity greater than 3 days.
- A prescription written by a nurse prescriber must be first dispensed not more than 4 days after the date on which the prescription was written.

### 4.3.1 Annotation of Controlled Drug Prescriptions

All three copies of the prescription form must be annotated with:

a. The prescription number(s); and  
b. Each Date of Dispensing; and  
c. The quantity dispensed; and  
d. The strength dispensed; and  
e. The initials of the dispensing pharmacist; and  
f. Pharmacy Stamp.

The first dispensing (for the supply of Class B Controlled Drugs only) can be claimed as two dispensings if stock is unavailable to dispense the full amount required. This includes situations where both dispensings are supplied on the same day. Subsequent repeats where insufficient stock is available must be claimed as one repeat and an “owe”.

**Procedure:**

- Claim for the first supply as an initial dispensing  
- The second dispensing should be claimed as a repeat dispensing.  
- A note should be made on the Controlled Drug Prescription of the quantities and dates of the dispensing of both supplies

### 4.3.2 Submission of Controlled Drug Prescriptions

On the completion of all dispensings:

a. the top copy (white) is to be retained in the pharmacy;  
b. the 2nd copy (yellow) and 3rd copy (red) are to be filed in the batch on the date of initial dispensing.

### 4.4 Supply Orders

Bulk Supply Orders (BSOs), Practitioner Supply Orders (PSOs) and Wholesale Supply Orders (WSOs) must be supplied in accordance with ‘Miscellaneous Provisions’ of the Pharmaceutical Schedule.

Except antipsychotic injections for mental health day clinics, PSOs will not be reimbursed where the pharmaceuticals are supplied to hospitals or clinics.

BSOs, PSOs and WSOs will not be reimbursed where the pharmaceuticals are supplied to the Armed Services or the Department of Corrections (including prisons).

Supply Order forms are available from Wickliffe Ltd 0800 259 138. The reorder numbers are:
4.5 Specific Prescription Types

All prescriptions, however generated, must have placed on the face of the prescription, the pharmacy stamp (as per the Medicines Regulations 1984).

4.5.1 Prescriptions for Multiple Patients

Prescriptions, such as antifungal or scabies treatments, for multiple Patients on one form should be treated as separate prescriptions. All names are required on the prescription, normal co-payment rules will apply.

4.5.2 Pharmacy Generated Prescription Forms

No claim for payment shall be made in respect of a pharmacy generated prescription form until the pharmacy receives the signed form from the prescriber and meets all legal and contractual requirements and the dispensing process is complete.

4.5.3 Bulk / Merged Prescription Forms

Where a prescription is generated for multiple rest home Patients, the pharmacy must ensure that:

a. The prescriber has initialled beside each patient on the page; and
b. There is this statement acknowledging that each patient is under the prescribers care, ‘I have read and authorised these prescription orders for the above name patients’; and
c. Each page has a full prescriber’s signature and date at the bottom of the page.

4.5.4 Telephone Prescriptions

The circumstances in which an orally communicated prescription may be made are set out in the Medicines Regulations 1984 and include:

“Within 7 days after a communication made by an authorised prescriber to a pharmacist, the prescriber must forward a written prescription confirming the oral communication”

4.5.5 Certified Repeat Copies

A Certified Repeat Copy is a computer generated copy of the record of a repeat Prescription Item. It can be used for dispensing repeat supplies as an alternative to dispensing from the original prescription.

If not dispensing from the original prescription, a Certified Repeat Copy (CRC) must be generated when repeats are dispensed in the following two situations:

- when a repeat is being claimed as uncollected or
- if the conditions of repeat supply are different to those at the first dispensing (e.g. variable quantities; supply of the prescription item for two months for access exemption purposes where the stamp needs to be placed on the CRC).

The change must be annotated on either the original prescription or on the CRC.
The certified repeat copy must be filed in the batch at the date of the repeat dispensing.

For further information regarding repeat supplies refer to clause 4.6.

4.5.6 Certified True Copies

A Certified True Copy must be used only:

d. When the original prescription form has to be made available to the NZ Police, Medsafe, Medical Officer of Health, or the Coroner; or
e. Where an item has to be dispensed by another pharmacy as per conditions set out in the Pharmacy Services Agreement.

Procedure:

- Those items which can be dispensed should be supplied and the pharmacy stamp, date of dispensing, prescription number(s), and items dispensed should be clearly indicated on the original form.
- A Certified True Copy of the complete prescription form should be made by the pharmacy, be retained and submitted in the normal manner as part of the batch.
- The original prescription which has been annotated and signed by a pharmacist is given to the Patient to take to a pharmacy for supply of the undispensed items.
- A Certified True Copy must be annotated with the words: “Certified True Copy” and be signed and dated by the dispensing pharmacist.
- A photocopy is the preferred option for a Certified True Copy. In special circumstances the Certified True Copy can be handwritten or computer generated and the reason annotated.

4.5.7 Certified True Photocopies

A Certified True Photocopy must be made when all items on a multi item prescription are not processed on the same day. A Certified True Photocopy is different from a Certified Repeat Copy and is used for a different purpose.

The copy of the form MUST BE a photocopy

Procedure:

- On the original prescription after the dispensing annotate the items not being claimed on this day
- Take a photocopy of the original prescription. The original prescription form should be inserted into the batch as at the date of dispensing.
- The photocopy is referred to as a Certified True Photocopy and should be retained on file for dispensing of the remaining original items
- A Certified True Photocopy must be annotated with the words: “Certified True Photocopy” and be signed and dated by a pharmacist.
- The Certified True Photocopy MUST include all the items on the original prescription with the items previously dispensed crossed through. Prescription numbers for the items previously dispensed must be included on the photocopy.
- Once a Certified True Photocopy has been created, there must be no changes to the original prescription The Certified True Photocopy must be an exact copy of the original when the original is submitted to Sector Services for claiming.
- If another certified true photocopy is required for a subsequent dispensed date, repeat the process but use the 1st Certified True Photocopy as your starting point. The last Certified True Photocopy created must be a complete picture and show details of all items dispensed from that prescription form.
4.6 Repeat Supplies

To be eligible for subsidy, repeats must be dispensed in accordance with the terms and conditions of the Pharmaceutical Schedule as summarised in the table below:

<table>
<thead>
<tr>
<th>Indicator in the Pharmaceutical Schedule</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prescription details</td>
</tr>
<tr>
<td>No Endorsement (i.e. default)</td>
</tr>
<tr>
<td>Prescriber endorsed: “Certified exemption”</td>
</tr>
<tr>
<td>Patient Certified Access Exemption</td>
</tr>
<tr>
<td>Prescriber endorsed: “Close Control”</td>
</tr>
<tr>
<td>Prescriber or pharmacist endorsed: “Unstable medicine”</td>
</tr>
<tr>
<td>Dentist’s prescription (other than sodium fluoride)</td>
</tr>
</tbody>
</table>

Authorised repeats can be dispensed:

a. When a specific and express request for that repeat is made by the Patient or his/her caregiver, and

b. The pharmacist can reasonably assume that the last preceding supply has been exhausted or substantially exhausted including any previous prescriptions and repeats dispensed by that pharmacy, or for a reason otherwise known to the pharmacist (e.g: the patient is travelling and the patient signs the Access Exemption Declaration).

As a general rule, for a pharmaceutical benefit to apply ‘substantially exhausted’ means that either 2/3rds of the supply period has elapsed since the previous dispensing or 2/3rds of the supply has been used. In special circumstances where the Patient has lost or damaged the previous supply, or has an increased need for the medication due to a change in dose or frequency, the pharmacist can supply the medication earlier. If an earlier supply is made in these circumstances, the reason for the early supply must be annotated on the prescription or CRC for the Patient to be eligible for a subsidy.

4.7 Prescription Item Owing

The pharmacy must comply with the obligations of the Pharmacy Council’s Code of Ethics Obligation 1.7:

“The pharmacist must consult with the patient to achieve a mutually acceptable arrangement when it is not possible to dispense a medicine as prescribed.”

The commentary for this Obligation states:

“This provision will include situations such as part supply on dispensed medicines. In such cases the Charge Pharmacist should ensure that there is a documented procedure detailing the handling of medicines owed on prescriptions”
### Procedure:

- It is preferable to provide the full dispensing. The pharmacist should issue a part supply of a prescription item in cases where the Patient is required to begin the treatment immediately.
- If the full quantity of prescription item is not available, there must be a reference in the computer record, or in an "owes" file and on the prescription form specifying the quantity dispensed and the quantity owing.
- The patient must be provided with written information on the quantity owed and the timeframe for collection for the owed prescription item where known e.g. could be out of stock. The Owed prescription items must be collected or delivered within the period of supply on the prescription.
- The Pharmacist must not re-dispense any item for which a claim has been submitted.
- Payment will only be made for any owed prescription items when supplied to the Patient or his/her caregiver. Dispensing fees will not be paid for these balances.

### 4.8 Uncollected Prescription Items

Except where expressly permitted under the Agreement only prescription item(s) collected by or delivered to the Service User, their caregiver or prescriber can be claimed. Uncollected prescription items may not be claimed unless the provisions of clause H1.2 of the Agreement and the following requirements are met:

"take and document reasonable and auditable steps" as described in clause H1.2(c)(i) of the pharmacy agreement includes but is not limited to the following:

- Notify the Service User on at least two occasions that the Items are available for collection at least 4 weeks prior to the date on which the medicine would otherwise become “Uncollected”
- Notification can be by telephone calls, letters, texts, emails or other methods
- All such notifications must be recorded (preferably in the electronic notes for the patient )
- If an item remains “Uncollected”, the steps taken to notify the patient (including the dates) must be recorded on the Prescription Form
- After 90 days expiry, if it uncollected, claim as uncollected. If the patient comes after 90 days the patient should pay and the item shouldn’t be claimed as uncollected. Pharmacies will be credited back if an uncollected claim has been made and it must be reversed.

"relevant timeframe" as described in clause H1.2(c)(ii)(A) of the Pharmacy Service Agreement shall be the subsidy expiry time frames as set out in the Pharmaceutical Schedule. Only after these time frames have expired may any claim be considered for an Uncollected or Uncollected Repeat prescription item.

After the subsidy expiry time frame, if a prescription item is uncollected, and all of the necessary contractual requirements are met, it may be claimed as uncollected.

If the patient comes to collect the item after the subsidy expiry time frame (but before the item has been claimed as uncollected) the patient should pay the full cost and the item shouldn’t be claimed as uncollected; or

If the patient comes to collect the item after the subsidy expiry time frame (but after the item has been claimed as uncollected) the Pharmacy must request full payment from the patient and then deduct the amount off its next claim.
“capable of being verified in any audit” in relation to clause H1.2(c)(ii)(B) of the Agreement shall mean a contemporaneous written request by the Service User, the Service User’s caregiver or the prescriber relating individually to that request for you to generate that repeat Prescription Form. In relation to clause H1.2(d) of the Pharmacy Service Agreement it means a specific and express request by the Service User, the Service User’s caregiver or the prescriber. No blanket request signed by the Service User, Service User’s caregiver or prescriber will be accepted as a request for any original or repeat claimed as Uncollected.

“reasonably able to be Dispensed instead to another Service User” as described in clause H1.2(c)(ii)(C) and H1.2(c)(iv)(D) of the Agreement means that the Prescription Item can be returned to stock and provided to a different Service User, provided that this clause applies only to the Uncollected and Uncollected Repeat provisions of the Agreement, where the Pharmaceutical Items have remained at all times within the pharmacy premises and under the control of the pharmacy. It is expected that many items can be returned to stock (unless specifically detailed below as unable to be re-issued for dispensing) as the integrity and expiry of the medicine have not been compromised. For example, tablets counted from a larger supply and placed in a new container can be retained in that new container, and then dispensed to a subsequent Service User with the application of a new dispensing label.

Items unable to be re-issued for dispensing to another Service User include:
- Reconstituted antibiotic mixtures and extemporaneously compounded preparations if they are uncollected by the date that the prepared medicine expires or the subsidy expiry timeframes whichever comes sooner.
- Any pharmaceutical item where the integrity and expiry of the item has been compromised, for example:
  - Medicines which are blister packed and are hygroscopic or medicines that may be “contaminated” by touching other medicines in the blister pack.
  - Strip packed medicines that have been de-blistered (removed from their original foil packaging)
  - Tablets that have been halved (or otherwise divided)

For clarity, the following items cannot be Uncollected or Uncollected Repeats and cannot be re-dispensed under any circumstances:
- Any pharmaceutical item that has already previously been dispensed and for which a claim for payment has been made (for example blister pack dispensing to rest homes that is returned is not able to be re-dispensed to another Service User).
- Any Pharmaceutical Item that has been returned by a Service User.
Annotation
If a claim is to be made for an Uncollected Prescription Item or an Uncollected Repeat Prescription Item the original prescription form or CRC is to be annotated by you with the words “Uncollected” or “Uncollected Repeat” respectively. By annotating the script “Uncollected”, you are representing that you have met all of the requirements of clause H1.2 of the Agreement and of this Procedures Manual prior to submitting the claim.

For an Uncollected Prescription Item;

* Annotate on the front of the original prescription form “Uncollected”, then on the back of the original prescription form the item(s) claimed as Uncollected.
* List only the prescription numbers that relate to the “Uncollected” items
* the dates, times and initials of person responsible for “reasonable and auditable steps” performed must be recorded on the back of the original prescription form and a copy of the letter sent to the service user shall be stapled to the original prescription.

For an Uncollected Repeat Prescription Item:

* A CRC is to be printed for each Service User containing all repeat items claimed as “Uncollected Repeat” items.
* Annotate the front of the CRC to indicate these items relate to “Uncollected Repeat” items
* the dates, times and initials of person responsible for “reasonable and auditable steps” performed will be recorded on the CRC.

General matters

A record shall be made in the Service User’s medication record within the software system that the prescription item(s) has been prepared and placed on the uncollected shelf and this item shall be continuously deferred for payment until such time as it is collected or claimed as Uncollected. No claim should be made for an Uncollected item if collected by the service user.

Only one original dispensing or one repeat relating to a unique identifying number should be prepared and placed on the uncollected shelf at a time. No further remaining repeats are to be prepared until the previous original dispensing or repeat has been collected. If a repeat is claimed as uncollected then no further remaining repeats should be prepared or claimed as Uncollected.

Example:
12345/5 was prepared but uncollected. The remaining repeats of /6 to /12 for this prescription item are not to be prepared and placed on the uncollected shelf and later claimed as uncollected.

No original or repeat prescription item should be prepared and later claimed as Uncollected if the pharmacy has some knowledge that the patient would not be able to collect the prescription item within the subsidy timeframes, examples include a service user is moving away, is in hospital, going overseas, or has died.

If requested by the DHB or its agent each pharmacy must prepare and produce a report, either in hardcopy or by electronic, of all Prescription Items claimed as Uncollected and/or Uncollected Repeats. This report is to include the following information:

Patient Name and address, unique identifying number of the Prescription Item, deemed date of dispensing, medicine name (including strength and form), quantity, and claim date.
4.9 Annotations

An annotation is text written by a pharmacist. Any annotation should clearly differentiate the information added by the pharmacist from that written by the prescriber, preferably using green ink unless the prescriber has written in green. If possible all annotations should be adjacent to the prescription item.

Prescriptions should be annotated:
- Where it is required by regulations, or
- Where it is necessary for clarification, or
- Where it is required for subsidy, including those outlined in the Pharmaceutical Schedule e.g. Cost Brand Source, Multiple-Patients.
- Where there is no Patient category code or status code on the prescription or when it is known to be erroneous.

Changes made to the category codes by the pharmacist must be initialled and reflected in the electronic claim file.

Pharmacists may annotate prescriptions with clarifications to:
- Dosage;
- Strength;
- Quantity;
- Brand (the pharmacist may only annotate a change of brand subject to the substitution rules contained in the Medicines Regulations)

Points to note
- The reason for any variance between the original prescription and the electronic record must be annotated on the prescription. Note: “C.B” can be used as an abbreviation for Change Brand.
- Subject to the rules of the Pharmaceutical Schedule, where there is a financial implication for the DHB the pharmacist cannot increase the quantity of a prescription, without the prescriber signing the alteration
- Non-subsidised items should be identified.
- Manual Claimants should annotate the brand dispensed on any generic prescription where more than one brand is subsidised.

4.10 Endorsements

An endorsement is text written by a prescriber on a prescription. The Pharmaceutical Schedule defines the requirements, which may vary from time to time. Where an endorsement is required on a prescription it must either be:

- Hand-written, or computer generated on the prescription by the prescriber; or
- Where it is not hand-written, or computer generated by the prescriber, and where it is specified in the Pharmaceutical Schedule, be initialled by the prescriber; and
- Where it has been altered or added to by the pharmacy, be initialled by the prescriber.

4.11 Alteration to Quantity Dispensed

An alteration made by a pharmacist to the unit quantity dispensed is one that does not affect the end amount of medicine prescribed to the Patient. The Patient will get the same dosage of medicine in the following example: the prescription reads “500 mg, one tablet per day, 30” and the pharmacist dispenses “250 mg tablets, two tablets per day, 60.” In this case, the pharmacist has altered the unit quantity, and subsequent dosage instructions, without changing the total daily dose or frequency ordered by the prescriber.
Alternatively, a change from tablets to mixtures is deemed appropriate as long as both the individual dose and total daily dose is not altered. In the above example, if there is no additional cost to the District Health Board the pharmacist can annotate and sign the changes.

**Procedure:**

For any alteration made by the pharmacist to the quantity dispensed, if there is a financial implication or increased cost to the DHB:

- The pharmacist must annotate and sign the reason for the change
- The change must be authorised and signed by the prescriber.
- In cases where PHARMAC has approved and notified in writing such a change in dispensing of a named pharmaceutical due to an out of stock event or short supply, the pharmacist must annotate and initial the alteration.

### 4.12 Cost, Brand, Source (CBS)

Where CBS is indicated against a medicine in the Pharmaceutical Schedule or if the item is an Exceptional Circumstances medicine not in the Pharmaceutical Schedule (as described in the Pharmacy Agreement), the medicine is eligible for subsidy on the basis of the Pharmacists annotation of purchase price, brand and source of supply. The purchase price should be GST exclusive. The Pharmaceutical Schedule requires that the purchase price, brand and source of supply be annotated. Alternatively a copy of the invoice for the purchase of the Pharmaceutical may be attached to the prescription, in the place of an annotation, in order to be eligible for Subsidy.

The details of the purchase may be subject to audit, and all receipts of purchase must be kept and available for audit.

### 4.13 Original Pack Dispensing

**Procedure:**

- If an item has the letters “OP” in the pack size column of the Pharmaceutical Schedule, then payment is made to the nearest original unit size
- The pack size dispensed should be the closest size to meet the dosage instructions, and will be reimbursed for the total subsidy per OP dispensed.

Example: Collapsible Tube (if defined as ‘OP’ in the Pharmaceutical Schedule): Locoid Lipo cream Apply bd 15g. Even though the prescription only calls for 15g, the pharmacist can claim 1OP or 30g. If the Locoid prescription had called for 50g, the pharmacist can claim 2OP or 60g.

### 4.14 Oral Antibiotic Liquids

Where a prescriber has written a prescription for a reconstitutable oral liquid antibiotic indicated in the Pharmaceutical Schedule as an original pack, and the dispensing of which would require the pharmacist to break into another pack, the pharmacist should reduce the amount dispensed to the quantity contained in a whole pack provided that the reduction in the amount dispensed is less than 10% of an original pack and in the reasonable opinion of the pharmacist will not effect the efficacy of the course of treatment.
Example:
5ml tds for 7 days = 105ml
Dispense 100ml

Example:
10ml stat, 5ml tds for 7 days = 110ml
Dispense 110ml.
Remainder can be claimed if unused

4.14.1 Broken Packs

Where a pharmacist dispenses a part pack of a proprietary product, subsidy is based on the appropriate portion of the pack size listed in the Pharmaceutical Schedule, unless the item lists “OP” in the pack size column of the Schedule.

At the time of dispensing the pharmacist must keep a record of the quantity discarded. To ensure wastage is reduced, the pharmacist should reduce the amount dispensed to make it equal to the quantity contained in a whole pack where:

a) the difference of the amount dispensed and the amount prescribed by the Practitioner is less than 10% (e.g.; if a prescription is for 105mls then a 100ml pack would be dispensed); and

b) in the reasonable opinion of the pharmacist the difference would not affect the efficacy of the course of treatment prescribed by the Practitioner.

4.15 Close Control

The Close Control rule is detailed in the Pharmaceutical Schedule. Below is a copy of the rule and a flow chart explaining how the rule works.

“Close Control” means the dispensing of a Community Pharmaceutical, in accordance with a Prescription, in quantities less than one 90 Day Lot (or for oral contraceptives, less than one 180 Day Lot) for a Community Pharmaceutical referred to in Section F Part I, or in quantities less than a Monthly Lot for any other Community Pharmaceutical, where any of a), b) or c) apply.

a) All of the following conditions are met:
   i) the Community Pharmaceutical has been prescribed for a patient who:
      1) is not a resident in a Penal Institution, Rest Home or Residential Disability Care Institution; and
      2) either of the following:
         i) in the opinion of the prescribing Practitioner is:
            a) frail; or
            b) infirm; or
            c) unable to manage their medication without additional support; or
            d) intellectually impaired; or
            e) requires close monitoring due to recent initiation onto, or dose change for, the Community Pharmaceutical (applicable to the patient’s first changed Prescription only); and
            f) requires that Community Pharmaceutical to be dispensed in a smaller quantity than that for which it is currently funded, or
ii) the Community Pharmaceutical is any of the following:
   a) a tri-cyclic antidepressant; or
   b) an antipsychotic; or
   c) a benzodiazepine; or
   d) a Class B Controlled Drug; and

ii) the prescribing Practitioner has:
   A) endorsed each Community Pharmaceutical on the Prescription clearly with the words “Close Control” or “CC”; and
   B) initialled the endorsement in their own handwriting; and
   C) specified the maximum quantity or period of supply to be dispensed at any one time.

b) All of the following conditions are met:
   i) The Community Pharmaceutical is prescribed for a patient who is a resident in a Rest Home or Residential Disability Care Institution; and
      A) the quantity or period of supply to be dispensed at any one time is not less than 28 days’ supply; and
      B) the prescriber or pharmacist has written the name of the Rest Home or Residential Disability Care Institution on the prescription; and
      C) the prescriber or pharmacist has:
         1) written on the Prescription the words “Close Control” or “CC” (this applies to all medicines prescribed on the prescription), and
         2) initialled the endorsement/annotation in their own handwriting; and
         3) specified the maximum quantity or period of supply to be dispensed at any one time.

c) All of the following conditions are met:
   i) where PHARMAC has approved and notified pharmacists to annotate prescriptions for a specified Community Pharmaceutical(s) “Close Control” without prescriber endorsement for a specified time; and
   ii) the dispensing pharmacist has:
      A) clearly annotated each of the approved Community Pharmaceuticals that appear on the prescription with the words “Close Control” or “CC”; and
      B) initialed the annotation in their own handwriting; and
      C) specified the maximum quantity or period of supply to be dispensed at any one time, as specified by PHARMAC at the time of notification.
Changes to the Close Control Rules from 1 June 2008

**Subsidised medicine**

- **Rest home or residential care**
  - **Patient resident in a rest home or residential disability care institution**
    - YES
    - **Specific Medications**
      - The pharmaceutical is either:
        - a tricyclic antidepressant; or
        - an antipsychotic; or
        - a benzodiazepine; or
        - a class B Controlled Drug
      - YES
      - **Stock Issue**
        - PHARMAC has notified pharmacists to implement Close Control for a specified pharmaceutical for a specified time
        - YES
      - **Penal Institution**
        - Patient resident in a penal institution
        - YES
      - **Requires monitoring**
        - The prescriber determines the patient:
          - Frail, or infirm, or unable to manage their medication without additional support, or intellectually impaired, or requiring close monitoring due to recent initiation or dose adjustment of a pharmaceutical
          - Needs the pharmaceutical to be dispensed in smaller quantities than that for which it is currently funded
        - YES
      - **Close Control dispensing funded if the prescriber**:
        - Endorses each item to be dispensed
        - Close Control with “close control” or “CC”, and
        - Initials each endorsement in their own handwriting; and
        - Specifies maximum quantity/period of supply to be dispensed at any one time
      - **Close Control dispensing funded not funded**

- **NO**
4.16 Multiple Dispensings in Out of Stock Situations

There will be situations where there are limited supplies of subsidised items due to the manufacturer/distributor having insufficient stock to meet demand. PHARMAC will assess these situations and may amend the defined period of supply. (Refer to Section 4.15) In these instances pharmacists will be eligible to endorse the prescription as “close control” and be reimbursed for the extra dispensings.

The procedure to be followed is:

- PHARMAC will notify pharmacies by fax of the effective date of the change. Where appropriate the advice will be reiterated in the following Pharmaceutical Schedule Update.
- Pharmacists may reduce the period of supply in accordance with the instructions issued by PHARMAC.
- The change to the period of supply must be annotated on the prescription as Close Control and the annotation initialled and dated by the pharmacist.
- The change must be consistent with the period authorised by PHARMAC e.g. monthly rather than three monthly.
- Once there is sufficient stock PHARMAC will advise pharmacies of the date when pharmacists may not endorse the pharmaceutical “close control.” This advice will be sent by fax, and reiterated in the following Pharmaceutical Schedule Update.
- Any prescription first dispensed under these arrangements as “close control” may be completed as “close control” unless otherwise directed by PHARMAC or dispensed in accordance with the prescriber’s original instructions.

5. Interpretation of Reimbursement Restrictions

There are several specific rulings that provide an interpretation for pharmacists on the quantity of pharmaceuticals that can be reimbursed under the Pharmaceutical Schedule General Rules, the Pharmacy Services Agreement or the provision of pharmacy services. These rulings are provided to Sector Services, having been considered by the DHBs.

**Procedure:**

- The pharmacy is required to comply with the Pharmaceutical Schedule General Rules and subsidy restrictions, the Pharmacy Services Agreement for the provision of pharmacy services
- Where necessary for clarification, the pharmacist should annotate the prescription.

5.1 Eye Drops

If a prescription is written for a three-month supply of eye drops, at least one original pack will be subsidised per month even if the directions are such that one pack would suffice for the complete three-month course. This follows the requirement to discard eye drops 30 days after opening. The pharmacist must annotate the prescription when they are claiming for quantities in excess of dose and frequency prescribed.
The following guidelines should be used for calculating quantities of eye drops:

a. 12 drops = 1ml
b. 60 drops = 5ml

5.2 Insulin Vials and Cartridges

If a prescription is written for a three-month supply of insulin, at least one vial or one cartridge will be subsidised per month even if the directions are such that one pack would suffice for the complete three-month course. This follows the need to discard vials 30 days after opening. The pharmacist must annotate the prescription when claiming for quantities in excess of dose and frequency prescribed.

5.3 Mucilaginous Laxatives

These products are reimbursed as an original pack. The following guidelines should be used to calculate quantities.

a. One teaspoonful = 7 grams
b. One dessertspoonful = 14 grams
c. One tablespoon = 28 grams

5.4 Bronchodilator Asthma Inhalers

Where a prescription for a bronchodilator inhaler has a “when required” component in the dosing schedule, up to 1200 doses will be reimbursed per 3 months.

Example:
Salbutamol inhaler: 2 puffs bd and prn Or 2 puffs prn

In these circumstances, up to 1200 doses (or 6 x 200 dose inhalers) will be reimbursed. These inhalers should be dispensed in quantities depending upon the Patient’s needs.

Example:
Six inhalers can be dispensed as 2+2+2 or 3+2+1 or 4+1+1

If the dosing schedule does not have a “when required” component, then the quantity supplied must relate to the total number of doses ordered.

Example:
2 puffs qid for 3/12
720 doses (or 4 x 200 dose inhalers)

In instances where a quantity larger than 1200 doses is required and the reason for the extra quantity is annotated on the prescription by the Prescriber, the quantity prescribed will be reimbursed.

5.5 Steroid Asthma Inhalers

For steroid inhalers without a definitive dosing and frequency instruction, only one inhaler can be claimed in each monthly dispensing.
Example:
Beclomethasone inhaler 100mcg/dose 2-4 puffs prn
Prescriptions, which specify both a dose and frequency of dose, will be reimbursed up to the maximum number of inhalers as provided for by the prescriber's instructions.

Example:
Beclomethasone inhaler 100mcg/dose 2-4 puffs bd prn
Up to four 200-dose inhalers would be reimbursed on these instructions

Example:
Beclomethasone inhaler 100mcg/dose 4 puffs bd increasing to 8 puffs bd prn
A maximum of eight 200-dose inhalers would be reimbursed on this prescription

6. **Health Entitlement Cards**

6.1 **Community Services Cards**

Community Services Cards are available to provide targeted subsidies to selected Patients to access Health and Disability Services, in particular pharmaceuticals and general practice services.

If a Patient qualifies for a Community Services Card, he/she will receive an individual card. If the Patient is married (i.e. legally married or living with someone in a relationship which is similar to marriage – reference Health Card Regulations 1993 Reg. 3) both Patients will have their own card. Either card can be used to cover dependent children.

Patients who qualify for NZ Super or a Veteran's Card will have CSC entitlement noted on their SuperGold Card.

For further information, contact Work and Income National Community Services Card Centre on 0508 555 999 between 9.00am - 5.00pm Monday to Friday, with the exception of Wednesday when the hours are 9.30am – 5.00pm.

Community Services Card status should be indicated by a 1 on the prescription form: 1 is used for those Patients covered by a Community Services Card.

6.2 **High Use Health Cards**

High Use Health Cards are for those people who visit their doctor on 12 or more occasions within a year for an ongoing medical condition/s. There are specific requirements necessary for eligibility.

A prescriber applies for, a High Use Health Card on behalf of a Patient. Applications are made to Sector Services’s Wanganui office.

A High Use Health Card is issued to an individual and not a family.

Application forms and information brochures for Patients are available from Wickliffe Ltd 0800 259 138.
6.3 Prescription Subsidy Cards

Each time an initial dispensing of a prescription with repeats or a “stat” dispensing (single supply) is made and all or the initial part of a co-payment ($3, $10 or $15) is paid by a family unit, this should be recorded on either the Patient’s Prescription Record Card or against the individual’s medication history. Please note that where the Patient does not pay a patient co-payment the item does not count towards the family’s PSC count.

Patient Co-payments are not required for the following items/prescriptions:

i. Class B Controlled Drugs, other than methylphenidate hydrochloride or dexamphetamine sulphate

ii. Children aged under 6 years

iii. Patients enrolled with the Hokianga Health Enterprise Trust

iv. Antituberculotic (TB) prescription items

v. Antileprotic prescription items

vi. Repeat supplies if the full co-payment has been made on previous dispensings.

vii. Prescription items that are not subsidised

viii. Medicines for approved Templeton patients

Under co-payment prescriptions must be added to the Prescription Record Card as a Patient co-payment has been paid.
**Glossary**

**Definition of a Family Unit** From The Health Entitlement Cards Regulations:

Part 3 – Prescription Subsidy Cards, Section 22(1)

Family Unit means –

a. A married (or partnered) couple with one or more dependent children:

b. A married (or partnered) couple with no dependent children:

c. One person with one or more dependent children:

d. One person who is not a member of a family unit described in paragraphs (a) to (c) of this definition:

Part 3 – Prescription Subsidy Cards, Section 22(2)

For the purpose of this Part, the Director-General may regard as married any man and woman who, although not legally married or in a civil union, have entered into a relationship in the nature of marriage, and may determine a date on which that relationship is to be taken as having commenced.

Section 2(1)

A dependent child has the meaning given to it by section 3(1) of the Social Security Act 1964; but does not include a child for whom an orphan’s benefit or an unsupported child’s benefit is paid under that Act:

From the Social Security Act 1964

Part 1 – Monetary benefits (Section 3(1)

Dependent child, in relation to any person, means a child –

Whose care primarily the responsibility of that person; and

Who is being maintained as a member of that person’s family; and

Who is financially dependent on that person; and

Who is not a child in respect of whom payments are being made under section 363 of the Children, Young Persons, and Their Families Act 1989

- A member of a family unit may, at any time, request to see a copy of the family prescription record.
- The Pharmacy computer system must maintain accurate links to the prescriptions of the family unit members. These links will be audited.
- Once a family unit has received 20 initial dispensing of single supplies of subsidised pharmaceuticals in the year commencing 1 February to 31 January, the family must be issued with a Prescription Subsidy Card by the pharmacist.
- The 20 prescriptions recorded may have been dispensed by any number of pharmacies. However, prescriptions from another pharmacy must be able to be verified by a printout or receipt from the dispensing pharmacy.
- The PSC must contain the names of the family members eligible to use the card, be signed by the issuing pharmacist, and stamped on the back with the pharmacy stamp as indicated on the card.
- The pharmacist certifying exemption and issuing the Prescription Subsidy Card must sign the printout of the 20 items or the Prescription Record Card. The claimant number must be included.
- On the issue of a Prescription Subsidy Card, the pharmacist must record the number of the Prescription Subsidy Card on the Prescription Record Card and retain the printout or completed Prescription Record Card for 10 years following the date of issue.
- If the pharmacy uses computerised records to register the dispensed items to the family unit, a print out or record of the items from all involved pharmacies should be attached and retained when issuing a Prescription Subsidy Card.
A duplicate card should not be issued under ordinary circumstances. A photocopy of the prescription subsidy card can be used to inform another pharmacy of the family member’s entitlement such as in the case of a child at a boarding school.

The Prescription Subsidy card period is from February 1st in any year until January 31st of the next. All patients noted on the Prescription Subsidy Card are entitled to reduced Co-payment charges until that card expires.

The pharmacist will be provided with a supply of blank PSCs for each period by Wickliffe Ltd on behalf of Sector Services (Reorder Number 74077). Additional supplies are available from Wickliffe Ltd 0800 259 138.

7. **Special Authority**

The Pharmaceutical Schedule specifies “Special Authority” pharmaceuticals and their access criteria. “Special Authority” means that the Community Pharmaceutical is not eligible for Subsidy unless it has been prescribed and dispensed to a Patient in accordance with all the restrictions and instructions specified for that Pharmaceutical in Sections B, C or D of the Schedule. Clinicians submit applications for Special Authorities on behalf of their Patients to Sector Services.

Please refer to your Pharmaceutical Schedule for more information.

A Special Authority number entitles Patients who comply with the relevant criteria to one of the following:

- Subsidy on pharmaceuticals or special foods;
- Manufacturer’s price in cases where a premium would otherwise be payable. The entitlement to full subsidy continues following an increase in price;
- A higher subsidy than would be available without a Special Authority, but possibly still less than the manufacturer’s price. This is known as an alternate subsidy and is sometimes linked to the price of cheaper alternate products. Although the entitlement may at times be equal to the manufacturer’s price, this would not continue following an increase in price;
- Waive a restriction that would otherwise apply, such as a maximum quantity per prescription.

An example of a special authority approval number is CHEM0000078/Jan10. The month and year refer to the expiry date of the Special Authority (the approval will expire on the last day of the stated month).

Special Authority numbers can be verified by:

1. Phoning Sector Services on 0800 243 666. Access to this number is available 8 am – 5 pm Monday to Friday with the exception of Wednesday when the hours are 9:30am – 5pm.

A pharmacist must quote the pharmacy’s claimant number before any information can be obtained. If Sector Services staff cannot identify the pharmacy, they will ask for the query to be forwarded by fax to 0800 100 131. Sector Services will then contact the Pharmacy to discuss the query.
2. Using the on-line special authority look-up tool. Access to the look-up tool is available 24 hours per day, 7 days per week unless notified by email by Sector Services.

If necessary to clarify the expiry date of a Special Authority Approval the pharmacist should quote the approval number or the Patient's name and NHI number (if known) to obtain any Patient information.

Special Authority approvals are not retrospective. Subsidy is applicable from the date of receipt of a valid application at Sector Services. If prescriptions are presented prior to authorisation of the Special Authority approval number, and medication is required urgently, payment arrangements should be made with the patient.

The pharmacist needs to check expiry dates on Special Authority numbers to ensure that it is current. The expiry date forms part of the Special Authority number.

If a three-month prescription is first dispensed before the Special Authority expiry date, the repeats will be reimbursed if they are collected after the Special Authority expiry date unless the medicine has been delisted from the Pharmaceutical Schedule.

**Procedure:**

**If the Special Authority has been annotated on the prescription by the prescriber**

- Check the special Authority is valid for the patient, pharmaceutical and the expiry date given is still current.
- Check the number follows the appropriate format (up to 10 numeric characters)
- Dispense the medication either with a valid Special Authority number, or at a charge to the patient, or as an ethical supply.
- Submit a claim.

The claim system will validate the Special Authority number prior to payment by reference to the Special Authority database. If the number can be found on the database, is still current and is being used for the designated pharmaceutical the item will be reimbursed.

If the claim item is unable to be validated, it will be rejected. In these circumstances refer to the Error Code Booklet. The first action should be ascertain how the error has occurred.

a. Pharmacist Error

If on data entry the Special Authority number is not entered correctly the claim will be rejected. Refer to the original prescription. The item will need to be edited to include the correct number and resent.

b. Prescriber Error

Please review the original prescription and contact the prescriber, or Sector Services on 0800 243 666

The problem here can be ascertained by reference to the prescriber, Sector Services (0800 243 666) or the patient.

The problem here

i. If the patient has a valid Special Authority and the information has not been written on the prescription correctly, eg it is a problem with transposition of numbers, the claim for the item can
be edited to include the right number and resent.

ii. If an application has been submitted, and the medicine has been dispensed prior to the approval being processed, the correct number should be added to the claim item and resubmitted for reimbursement. Please note that the Special Authority is valid from the date of a correct application being received by Sector Services. It is permitted to split the script to make an initial small unsubsidised dispensing pending issue of the Special Authority, which when received will permit the balance of the script to be processed as a subsidised prescription (not to exceed in total the original prescribed amount).

iii. If the problem can not be corrected (eg an application has not been submitted or the information as supplied cannot be validated) contact Sector Services on 0800 243 666, and advise that the correct procedures listed above have been followed, but a correct number has not been available. Sector Services will provide a Risk number after a review of the circumstances.

The Risk number can be added to the claim and the item resent.

**If a prescription is presented without a Special Authority number and the item requires one:**

Contact the prescriber, the patient or Sector Services on 0800 243 666. If a Special Authority application has been made successfully, and the appropriate information (eg number and expiry dates) is available this can be added to the claim for reimbursement.

If there is to be a delay in obtaining the information, (e.g. an application is pending, no application has been received, or there is no ability to contact anyone to ascertain the correct information) the following options should be considered:

iv. Delay supply of the medication until the appropriate documentation can be sourced if the pharmacist is comfortable that the patient can wait; or

v. Supply the medicine at the patient’s cost: or

vi. Supply the medication as an “ethical supply” because in the professional judgement of the pharmacist the patient will be at serious risk without the pharmaceutical. In this situation, if possible, the steps already outlined to obtain the correct Special Authority number before sending in the claim should be followed.

vii. If points iv - vi can not be done, contact Sector Services (0800 243 666) to apply for a RISK number. If a Risk number is approved you can then submit for payment.

**Risk Number Procedure**

1. A RISK Special Authority number to cover “ethical supply” will be issued only in the following circumstances.
   - The prescriber could not be contacted.
   - The prescription was presented during hours where the Sector Services 0800 number was not operational, or you do not have access to the online Special Authority lookup tool..
   - The patient was at serious risk without the medication, such as a life threatening condition or imminent hospitalisation. Examples of where this would apply are an hyperglycaemic event when appropriate insulin is not available, or a risk of kidney graft rejection without immediate availability of immunosuppressants.
2. “Ethical supply does not cover medicines where it is unlikely there would be a serious deterioration in the patient’s condition due to a delay in receiving the medicine, such as a prescription for a statin. It is expected that “ethical supply” under Special Authority will not be a frequent occurrence. It is designed as a last resort to protect patients at risk. These should only be for a supply that is enough to get the Patient through while the clinician is contacted for an application.

3. RISK numbers issued to cover prescriber error or the ethical supply situations are valid for the life of the prescription

It is important with the administration of Special Authority numbers that any additions or changes made by the pharmacist are initialled on the prescription and dated so that in an audit situation it is obvious which information the pharmacist is responsible for.

The following table may be of assistance as a ‘quick reference’ table for Pharmacists.

<table>
<thead>
<tr>
<th>Prefix</th>
<th>SA Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHEM</td>
<td>Special Authority</td>
<td>Allow patients to receive Special Authority medicines through a Community Pharmacy</td>
</tr>
<tr>
<td>EXCP</td>
<td>Community Exceptional Circumstances</td>
<td>Allows a patient to access a subsidy sufficient to fully fund the pharmaceutical. Criteria and application details are described in the Pharmaceutical Schedule.</td>
</tr>
</tbody>
</table>
| HOSP   | Special Authority           | Special Foods  
This prefix indicates that either:  
a. the doctor has requested a complete diet for their Patient, or  
b. the medicine can only be dispensed by a hospital pharmacy. |
| RISK   | Risk Number                 | Available where a Pharmacy has made a dispensing in good faith or if the patient has a life threatening condition. |
| SPEC   | Special Authority           | This prefix is indicates that either:  
a. the Prescriber has requested a supplement diet for their Patient, restricted to 500ml per day or as defined in the Pharmaceutical Schedule, or  
b. to waiver a restriction |
| TEMP   | Templeton                   | Enables subsidy for patients who were residents at the Templeton Centre at the time of closure. The approval numbers cover all medicines required by the patient. |

Note: The prefix (HOSP or SPEC) is only required on a prescription if Special Foods have been prescribed.

7.1 Special Foods

Prefixes are used to identify whether a Special Authority has been allocated for complete diet or for supplementary purposes.

a. A HOSP number for the purposes of a Special Food is an indication that the doctor has requested a complete diet for their Patient.

b. A SPEC number for the purposes of a Special Food is an indication that the doctor requested a supplement diet for their Patient, restricted to 500ml per day or as defined in the Pharmaceutical Schedule.
Refer above for Approval information and to the Error Code book for managing rejected claims.

### 7.2 Exceptional Circumstances

The purpose of the Exceptional Circumstances scheme is to provide funding from the Exceptional Circumstances budget for medication to be used in the community, in circumstances where provision of a funded Community Pharmaceutical is appropriate, but funding from the Pharmaceutical Budget cannot be provided through the Pharmaceutical Schedule.

Further information can be obtained from the Pharmaceutical Schedule.

### 8. Receipts and Late Claims

#### 8.1 Receipts

The following information is required on each receipt issued:

- a. Name of the Patient;
  - And for each prescription item on the receipt:
- b. Name of the prescription item
- c. Total Cost
- d. DHB subsidy
- e. Patient contribution

#### 8.2 Late Claims

**Time limit for receiving Claim Items**

Pursuant to clause H9.1 (and subject to clauses H9.2 and H9.3) of the Pharmacy Services Agreement, all Claim Items must be received within six months after the date when the Pharmaceutical is Dispensed, except for oral contraceptive Pharmaceuticals where the Claim Items must be received within nine months after the date when that Pharmaceutical is Dispensed (**Final Due Dates**).

**Submission out of time**

Where a Claim Item (provided it is for more than $20.00) has not been submitted or resubmitted by the applicable Final Due Date, it may be submitted out of time together with a written explanation of the reason for the delay. This explanation must be submitted to the DHB and copied to Sector Services. Where, in the reasonable opinion of the DHB, reasonable grounds have been established for late submission, the DHB will consider that Claim Item for payment.

**No submission after 12 months**

In no circumstances will any Claim Item submitted or resubmitted more than 12 months after the date of the Dispensing qualify for payment.
9. **Nicotine Replacement Therapy (NRT)**

This is a guideline for the reimbursement of Nicotine Replacement Therapy electronic claims made in accordance with the Pharmacy Services Agreement for the provision of pharmacy services. Claiming should include codes for Quit Cards 88888 or 99999 for a Quit Card issued by provider other than Quit Line.

Nicotine Replacement Therapy may be claimed by either a prescription or via a Quit Line Card.

If prescription is received for Nicotine Replacement Therapy please claim as per legal requirements for a prescription.

If a Quit line card is received please claim as per legal requirements for a prescription. Please use either the following numbers in the Provider registration number:

MC88888 for Quit cards from Quitline

MC99999 for Quit Cards from Quit Card Providers

9.1 **Electronic Claiming**

The procedure for NRT claiming is the same as for normal prescriptions.