2 August 2019

Dear Supplier

INVITATION TO TENDER – SUPPLY OF PHARMACEUTICALS TO DHB HOSPITALS AND/OR TO COMMUNITY PHARMACIES

PHARMAC invites tenders for the supply of certain pharmaceuticals to DHB hospitals and/or to community pharmacies in New Zealand.

This invitation to tender incorporates the following schedules:

- (a) Schedule 1 sets out the definitions used in this invitation;
- (b) Schedule 2 specifies the pharmaceuticals for which you may submit a Tender Bid in relation to community supply and/or hospital supply;
- (c) Schedule 3 describes the process PHARMAC intends to follow in relation to this tender, and provides instructions on how to submit a Tender Bid in relation to community supply and/or hospital supply;
- (d) Schedule 4 sets out terms that will apply if your Tender Bid in relation to community and/or hospital supply is awarded Sole Supply Status and/or Hospital Supply Status;
- (e) Schedule 5 sets out the additional terms that will apply if your Tender Bid in relation to community supply is awarded Sole Supply Status;
- (f) Schedule 6 sets out the additional terms that will apply if your Tender Bid in relation to hospital supply is awarded Hospital Supply Status; and
- (g) Schedule 7 sets out the additional special terms that will apply if your Tender Bid in relation to a particular pharmaceutical is awarded Sole Supply Status and/or Hospital Supply Status.

This invitation to tender also incorporates the information on the Electronic Portal referred to in this invitation.

If you wish to submit a Tender Bid in relation to community supply and/or hospital supply, you must submit it via the Electronic Portal to PHARMAC no later than **4pm** (New Zealand time) on **Friday 13 December 2019**.

If you have any inquiries about this invitation you should contact the **Tender Analysts** at tender@pharmac.govt.nz (Emma Clarke on (04) 830 1987 and Nerissa Ramlall on (04) 830 3824).

We look forward to receiving your tender.

Yours sincerely,

Lisa Williams Director of Operations

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Schedule 1: Definitions and interpretation

1. Definitions

In this Invitation:

Additional Stock Pharmaceutical (or ASP) means a Pharmaceutical, marked with a "@", for which the supplier of the successful Tender Bid would be required:

- (a) to hold additional stock; and
- (b) to report to PHARMAC on the level of that additional stock each Quarter;

Aggregated Tender Bid means a Tender Bid for more than one Tender Item, which PHARMAC is to consider in aggregate, and can include a Tender Bid for more than one Tender Item of the same Chemical Entity but not aggregation within a single Tender Item;

Agreement means:

- (a) Schedule Four; and
- (b) in relation to a Pharmaceutical with Sole Supply Status, Schedule Five; or
- (c) in relation to a Pharmaceutical with Hospital Supply Status, Schedule Six,

and includes, to the extent applicable, the other Schedules (including but not limited to Schedule Seven) and the information on the Electronic Portal comprising the Invitation;

Alternative Pharmaceutical means an alternative brand of a Pharmaceutical that PHARMAC, following consultation with PTAC or its sub-committees, considers to be an acceptable substitute for that Pharmaceutical;

Back-up Supply Agreement means an alternative agreement or arrangement negotiated by PHARMAC, at its sole discretion, with a supplier other than the supplier with Sole Supply Status and/or Hospital Supply Status in respect of a particular Tender Item, to cover the contingency that Sole Supply Status and/or Hospital Supply Status is suspended or withdrawn under the terms of this Agreement in respect of that Tender Item, or that the Tender Item is otherwise out of stock or unavailable for supply;

Chemical Entity means any pharmaceutical that contains, and is described generically according to, the relevant active ingredient specified in Schedule Two and the Electronic Portal. For the avoidance of doubt, the term Chemical Entity does not include any Medical Device;

Combined Community/Hospital Tender Bid means a Community Tender Bid and a Hospital Tender Bid that you submit in combination for the same Tender Item;

Community Tender Bid means a Tender Bid in relation to community supply;

Confidential Information means all information exchanged between us under this Invitation or in relation to your Tender Bid, including during all negotiations relating to your Tender Bid;

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Consents means all consents, permits, licences and authorisations, whether statutory or otherwise, required for the supply of the Tender Item in New Zealand (including Ministry of Health market approval);

Contract Manufacturer means a manufacturer or a supplier that is a party to a contract with the relevant DHB Hospital to compound Pharmaceuticals, on request from that DHB Hospital;

Cost Brand Source Product means a pharmaceutical where there is no price agreed upon by PHARMAC, but the pharmaceutical is subsidised by the Funder at the price obtained by pharmacies;

Crown Direction means any ministerial direction given to PHARMAC under section 103 of the Crown Entities Act 2004;

CTPP means Containered Trade Product Pack SNOMED CT code, which is the unique identifier that describes the packaged, branded product and the container it is dispensed in, as used within the New Zealand Medicines Terminology;

Deadline means 4 pm, Friday 13 December 2019 (New Zealand time);

Designated Delivery Point means at a DHB Hospital's discretion:

- (a) a delivery point agreed between you and the relevant DHB Hospital, to which delivery point you must supply the Pharmaceutical directly at the Price; and/or
- (b) any delivery point designated by the relevant DHB Hospital or PHARMAC, such delivery point being within 30km of your national distribution centre;

DHB Hospital means a DHB, including its hospital or associated provider unit for which that DHB purchases pharmaceuticals;

District Health Board (or DHB) has the same meaning as in the New Zealand Public Health and Disability Act 2000;

DV Limit means, for a particular Pharmaceutical, the National Discretionary Variance (DV) Limit or the Individual DV Limit;

DV Pharmaceutical means a discretionary variance Pharmaceutical, being an Alternative Pharmaceutical that does not have Hospital Supply Status, and includes a pharmaceutical which (unless PHARMAC specifies otherwise in Schedule Two of this Agreement and the Electronic Portal, or we agree otherwise in writing):

- (a) is listed as a DV Pharmaceutical, in association with the relevant Pharmaceutical having Hospital Supply Status, in the then current Section H of the Pharmaceutical Schedule; or
- (b) in the case of a pharmaceutical that is not a Medical Device, is the same Chemical Entity, at the same strength, and in the same or a similar presentation or form, as the relevant Pharmaceutical with Hospital Supply Status, but which is not yet listed as a DV Pharmaceutical.

For the avoidance of doubt, a pharmaceutical which:

(c) in the case of a pharmaceutical that is not a Medical Device, is a different Chemical Entity from the Pharmaceutical with Hospital Supply Status; and

(d) is not listed as a DV Pharmaceutical in the then current Section H of the Pharmaceutical Schedule,

is not a DV Pharmaceutical;

Electronic Portal means the electronic tender system available via the internet address provided to you by PHARMAC through which you are required to submit your Tender Bid(s);

End Date means the last day of the Hospital Supply Status Period, or Sole Supply Period, as applicable;

Evaluation Committee means a committee established by PHARMAC to evaluate Tender Bids;

Final Transition Period means, in respect of a Pharmaceutical with Sole Supply Status or Hospital Supply Status, as applicable, the period of three calendar months beginning on the day after the relevant End Date;

First Transition Period means, in respect of a Pharmaceutical with Sole Supply Status or Hospital Supply Status, the period beginning on the first day of the month following the Market Notification Date and ending on the day prior to five months from the Start Date (or such different or longer period as PHARMAC determines under clause 1.2 of Schedule Three);

Funder means the body or bodies responsible, pursuant to the New Zealand Public Health and Disability Act 2000, for the funding of pharmaceuticals listed on the Pharmaceutical Schedule (which may be, without limitation, one or more District Health Boards and/or the Ministry of Health) and their successors;

GTIN means the Global Trade Item Number for a Pharmaceutical;

Hospital Supply Status means the status of being the brand of the relevant Pharmaceutical listed in Section H of the Pharmaceutical Schedule as having such status, which Pharmaceutical DHB Hospitals must (or in the case of Medical Devices, may) purchase, subject to any DV Limit for that Pharmaceutical, for the Hospital Supply Status Period;

Hospital Supply Status Period means the period beginning on the day after the end of the First Transition Period and ending on 30 June 2023;

Hospital Tender Bid means a Tender Bid in relation to hospital supply;

Individual DV Limit means, for:

- (a) a particular Pharmaceutical; and
- (b) a particular DHB Hospital,

the discretionary variance limit, being a percentage of the Individual Total Market Volume, which equals the percentage of the National DV Limit for that Pharmaceutical, up to which that DHB Hospital may purchase DV Pharmaceuticals of that Pharmaceutical. The Individual DV Limit is set:

- (c) for the number of months during which the Hospital Supply Status Period applies during the period ending on 30 June 2021 and
- (d) the number of months during which the Hospital Supply Status Period applies during the twelve month period ending on 30 June 2022 and

Schedule 1

(e) the number of months during which the Hospital Supply Status Period applies during the twelve month period ending on 30 June 2023.

Individual Total Market Volume means for:

- (a) a particular Pharmaceutical; and
- (b) a particular DHB Hospital,

in any given period, in accordance with data available to PHARMAC, the sum of:

- (c) the total number of Units of the relevant Pharmaceutical with Hospital Supply Status purchased by the relevant DHB Hospital; and
- (d) the total number of Units of all the relevant DV Pharmaceuticals, listed in Section H in association with that Pharmaceutical, purchased by that DHB Hospital;

Invitation means this invitation to tender and includes the cover letter, each of the Schedules and the information on the Electronic Portal referred to in this invitation;

Lead Time means the number of months (being whole months only) indicated on your Tender Bid that, if your Tender Bid is accepted, you would require following the Successful Tenderer Notification Date in order to source sufficient stock of your brand of the Tender Item to meet the entire market demand for the Tender Item as at the Start Date. For the avoidance of doubt, the Lead Time does not affect, and should incorporate the extra time needed to allow for, your obligations in clause 3.1 of Schedule 5 and clause 3.1 of Schedule 6;

Market Notification Date means the date on which PHARMAC notifies the market that your Tender Bid, in respect of a particular Tender Item, has been accepted, being greater than one month prior to the Start Date;

Medical Device means a medical device as that term is defined in the Medicines Act 1981;

National DV Limit means, for a particular Pharmaceutical, the discretionary variance limit, being the specified percentage of the National Total Market Volume up to which all DHB Hospitals may collectively purchase DV Pharmaceuticals of that Pharmaceutical. The National DV Limit is set for DHB Hospitals nationally:

- (a) for the number of months during which the Hospital Supply Status Period applies during the period ending on 30 June 2021; and
- (b) the number of months during which the Hospital Supply Status Period applies during the twelve month period ending on 30 June 2022; and
- (c) the number of months during which the Hospital Supply Status Period applies during the twelve month period ending on 30 June 2023.

National Total Market Volume means, for a particular Pharmaceutical in any given period, in accordance with data available to PHARMAC, the sum of:

- (a) the total number of Units of the relevant Pharmaceutical with Hospital Supply Status purchased by all DHB Hospitals; and
- (b) the total number of Units of all the relevant DV Pharmaceuticals, listed in Section H in association with that Pharmaceutical, purchased by all DHB Hospitals;

Offer Letter means the letter of offer which must be attached to your Tender Submission Form, in the form set out in the Electronic Portal;

OPPs means PHARMAC's then current Operating Policies and Procedures and any relevant supplements, as applicable;

PCT means Pharmaceutical Cancer Treatment. Tender Items that are PCTs are indicated with "PCT" in the list in clause 2 of Schedule Two and the Electronic Portal;

Pharmaceutical means the relevant Tender Item (which may be a Medical Device) for which you have submitted, and PHARMAC has accepted on behalf of the Funder, a Tender Bid;

Pharmacode means the unique six or seven digit identifier assigned to a pharmaceutical and notified to you by the Pharmacy Guild. Suppliers must apply to the Pharmacy Guild of New Zealand to receive a Pharmacode for each presentation of their pharmaceutical before it is listed;

Potential Out-of-Stock Event means:

- (a) in relation to community or hospital supply, your stock of the Pharmaceutical in New Zealand falls below two-thirds of your most recent three months' total Unit sales of the Tender Item, or, where the Pharmaceutical is designated an ASP, your stock of the Pharmaceutical in New Zealand falls below your most recent four months' total Unit sales of the Tender Item; or
- (b) in relation to community or hospital supply, forecast sales demand in respect of the next two-month period is greater than your stock of the Pharmaceutical, or, where the Pharmaceutical is designated an ASP, forecast sales demand in respect of the next four-month period is greater than your stock of the Pharmaceutical; or
- (c) in relation to hospital supply, your stock of the Pharmaceutical in New Zealand falls below the average volume of stock of the Pharmaceutical required to supply the entire New Zealand DHB Hospital market for the Pharmaceutical for any given two-month period, or, where the Pharmaceutical is designated an ASP, your stock of the Pharmaceutical in New Zealand falls below the average volume of stock of the Pharmaceutical required to supply the entire New Zealand DHB Hospital market for the Pharmaceutical for any given four-month period; or
- (d) in relation to community supply, your stock of the Pharmaceutical in New Zealand falls below one-sixth of the Unit Volume, or, where the Pharmaceutical is designated an ASP, your stock of the Pharmaceutical in New Zealand falls below one-third of the Unit Volume;
- (e) in relation to community or hospital supply, your stock of the Pharmaceutical in New Zealand is insufficient to enable you to fully fill all orders as they are received (without restricting quantities that may be ordered); or
- (f) in relation to New Zealand manufactured products if either:
 - (i) forecast sales demand in respect of the next two-month period is greater than your stock of the Pharmaceutical; or
 - (ii) you have insufficient stock to enable you to fully fill all orders as they are received; or

 (iii) your stock of the active pharmaceutical ingredient taking into account manufacturing and stock on hand falls below two months stock for the Pharmaceutical in New Zealand;

For the avoidance of doubt, references to 'your stock' in (a) to (f) above refer to stock physically held by you or on your behalf in New Zealand and do not include stock held in New Zealand by wholesalers or other parties;

Price means the price (in New Zealand dollars and exclusive of GST) at which the Pharmaceutical is to be supplied, or made available for sale and supply, by you to:

- (a) in relation to community supply, wholesalers and other such distributors, and at which the Pharmaceutical is to be subsidised by the Funder, being the price specified in your successful Tender Submission Form, unless there has been a subsequent price change in accordance with the terms of the Invitation, in which case the Price will be the price notified to you by PHARMAC upon acceptance of your Tender Bid; or
- (b) in relation to hospital supply, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding), being the price specified in your successful Tender Submission Form, unless there has been a subsequent price change in accordance with the terms of the Invitation, in which case the Price will be the price notified to you by PHARMAC upon acceptance of your Tender Bid;

PTAC means the Pharmacology and Therapeutics Advisory Committee;

Quarter means the periods:

- (a) 1 January until 31 March;
- (b) 1 April until 30 June;
- (c) 1 July until 30 September; and
- (d) 1 October until 31 December;

Section B means the relevant section or sections of the Pharmaceutical Schedule relating to community pharmaceuticals;

Section H means the relevant section or sections of the Pharmaceutical Schedule identified as such, which relate to pharmaceuticals for use in hospitals;

Sole Supply Period means the period beginning on the day after the expiry of the First Transition Period and ending on 30 June 2023;

Sole Supply Status means, in relation to community supply, the status of being the sole subsidised supplier of the particular Tender Item for the Sole Supply Period;

Start Date means:

- (a) in relation to a Tender Item for which your Tender Bid has been accepted unconditionally, the first day of the month following the date that represents:
 - (i) the Successful Tenderer Notification Date; plus
 - (ii) the Lead Time; or

- (b) in relation to a Tender Item for which your Tender Bid has received conditional acceptance, in terms of clause 7.4 of Schedule Three, the first day of the month following the date that represents:
 - (i) the date that such acceptance ceases to be conditional; plus
 - (ii) the Lead Time; or
- (c) such other date that is negotiated between you and PHARMAC under clause 1.6 of Schedule Three;

Successful Tenderer Notification Date means the date on which PHARMAC notifies you, in relation to a Tender Item for which you have submitted a Tender Bid, that your Tender Bid has been accepted;

Tender Bid means the Offer Letter together with the Tender Submission Form submitted through the Electronic Portal for a particular Tender Item, including the Lead Time, and includes a Community Tender Bid, a Hospital Tender Bid and a Combined Community/Hospital Tender Bid;

Tender Item means:

- (a) in the case of a pharmaceutical that is not a Medical Device, the form and strength of a Chemical Entity (or entities, if applicable) for which you may submit a Tender Bid; or
- (b) in respect of a Medical Device, an item conforming to the individual specifications described for such item in the product list in clause 2 of Schedule Two for which you may submit a Tender Bid;

Tender Submission Form means the form on which you must submit your bid for each Tender Item, as set out in the Electronic Portal;

Transition Periods collectively refers to the First and Final Transition Periods;

Unit means an individual unit of a Tender Item (e.g. tablet, 1 ml of an oral liquid, ampoule, syringe, bag, suture or needle, roll or a dressing);

Unit Price means the relevant Price specified for a pack (or equivalent grouping for any Medical Device) of that Tender Item in Section H of the Pharmaceutical Schedule, divided by the number of Units in the pack specified in the Pharmaceutical Schedule as being the listed pack size for that Tender Item (and where that Tender Item is not listed on the Pharmaceutical Schedule, the price and pack size in the most recent issue of the Pharmaceutical Schedule published prior to that Tender Item being delisted);

Unit Subsidy means the subsidy specified for a pack of that Tender Item in Sections A to G of the Pharmaceutical Schedule, divided by the number of Units in the pack specified in the Pharmaceutical Schedule as being the subsidised pack size for that Tender Item (and where that Tender Item is not listed on the Pharmaceutical Schedule, the subsidy and pack size specified in the most recent issue of the Pharmaceutical Schedule published prior to that Tender Item being delisted); and

Unit Volume means, in relation to community supply, the approximate number of Units of the Tender Item subsidised by PHARMAC, and claimed for by community pharmacies, in one year, as specified in Schedule Two and the Electronic Portal.

2. Interpretation

In the construction of this Invitation, unless the context otherwise requires:

- (a) a reference to a clause or a Schedule is a reference to a clause of, or a Schedule to, this Invitation;
- (b) a reference to a statute, code of conduct or other law includes any regulations and other instruments under it and consolidations, amendments, re-enactments or replacements of any of them (whether before or after the date of this Agreement);
- (c) the singular includes the plural and vice versa;
- (d) the word person includes an individual, a body corporate, an association of persons (whether corporate or not), a trust, a state and an agency of state, in each case, whether or not having a separate legal personality;
- (e) a reference to a person includes a reference to the person's executors, administrators, successors, substitutes, (including, but not limited to, persons taking by novation) and permitted assignees;
- (f) words importing one gender include the other genders;
- (g) headings in this Agreement or in the Electronic Portal are for convenience only and have no legal effect; and
- (h) unless the context requires otherwise, references to the "listing" of a Pharmaceutical:
 - (i) in relation to hospital supply, are to the listing of that Pharmaceutical in Section H of the Pharmaceutical Schedule and are deemed to include any written notification by PHARMAC of that Pharmaceutical being the subject of a national supply contract negotiated by PHARMAC on behalf of DHBs, where such written notification is in advance of the actual listing of that Pharmaceutical in Section H of the Pharmaceutical Schedule (and references to "list", "listed", "delist", "delisted", and "delisting" are to be interpreted accordingly);
 - (ii) in relation to community supply, are to the actual listing of that Pharmaceutical in Sections A to G of the Pharmaceutical Schedule (and references to "list", "listed", "delist", "delisted", and "delisting" are to be interpreted accordingly).

Schedule 2: Products to be tendered

1. Information about Tender Items

1.1 List of Tender Items

This Schedule sets out the Tender Items and information about the Tender Items. While PHARMAC has taken all reasonable care in preparing the information contained in this Schedule, it accepts no liability for any errors or omissions in the information.

1.2 Patents

- (a) Where possible, PHARMAC has identified Tender Items that it understands may be the subject of a patent that it believes is due to expire after the Deadline.
- (b) Where PHARMAC has been advised of the existence of a patent prior to sending out this Invitation, it has shown this in the attached list by the use of a + symbol.
- (c) However, PHARMAC makes no representation as to the patent status of the Tender Items and accepts no liability for any patent infringement that might occur as a result of this tender process or PHARMAC's acceptance of a Tender Bid, including infringement of process patents.

1.3 Unit Volume and market value figures

- (a) Except where indicated otherwise, the Unit Volume figures, in relation to community supply, are based on actual volumes for the year ending 30 June 2019.
- (b) Market value figures, in relation to community supply, are expressed as the Unit Volume in the year ending 30 June 2019, multiplied by the Unit Subsidy as at 1 July 2019.
- (c) The figures referred to in paragraphs (a) and (b):
 - (i) are approximate and indicative only. PHARMAC makes no representation as to the accuracy of these figures or as to the level of sales or likely sales of any Tender Item. In particular, if these figures change at any time during the period from PHARMAC's pre-tender consultation until decisions have been made about the acceptance of Tender Bids for all Tender Items, PHARMAC is not obliged to notify you of any such change; and
 - (ii) unless specified by PHARMAC do not include DHB Hospital volumes. For the avoidance of doubt, PHARMAC makes no representation as to the size of the DHB Hospital market for any Tender Item, in relation to hospital supply.
- (d) You acknowledge and agree that in submitting your Tender Bid you will rely on your own knowledge, skill and independent advice or assessment of the market size for any Tender Item and PHARMAC is to have no liability in that regard.

1.4 **Special terms**

Where there are any special terms relating to a particular Tender Item, those terms are indicated in the column entitled "Comments" in the list and/or Schedule Seven. Special Authority restrictions have been noted for Tender Items where applicable in the list. Further restrictions

Schedule 2

on the supply of Tender Items within the Pharmaceutical Schedule may apply. You acknowledge and agree that in submitting your Tender Bid you will rely on your own knowledge and assessment of any restrictions applicable to a Tender Item within the Pharmaceutical Schedule.

1.5 Subsidies

- (a) The level at which each Tender Item, in relation to community supply, is specified in the attached list as being subsidised per Unit as at 1 July 2019.
- (b) Subsidies of Tender Items, in relation to community supply, may change before a Tender Bid is accepted.
- (c) Where a "*" symbol is indicated next to the Unit Subsidy in the attached list, there is no fully funded product available, in relation to community supply, for that Tender Item as at 1 July 2019.

1.6 **DV Limits**

Where there is a DV Limit relating to a particular Tender Item, in relation to hospital supply, that limit is indicated as a percentage amount in the column entitled "DV Limit" in the attached list and is also shown in the Electronic Portal.

1.7 Tender Items subject to sole supply arrangements

Where a Tender Item is underlined in the list of products below, that item is subject to a sole supply contract as at the date of this Invitation. Accordingly, the subsidy for those items is fixed until 30 June 2020 (unless otherwise indicated) and, for items that are the subject of a sole supply contract, the listing of a new brand, in relation to community supply, could only occur after that date. This information is not available in the Electronic Portal.

1.8 Hospital only products

Where an "H" is indicated, you may submit a Tender Bid for Hospital Supply Status for that Tender Item.

1.9 **Community only Products**

Where a "C" is indicated, you may submit a Tender Bid for Sole Supply Status for that Tender Item.

1.10 **Community and Hospital Products**

Where a "C" and an "H" are indicated, you may submit a Tender Bid for Sole Supply Status and/or a Tender Bid for Hospital Supply Status for that Tender Item. You may also submit a Combined Community/Hospital Tender Bid in accordance with clause 2.5 of Schedule Three.

1.11 PCTs

Where a "PCT" is indicated, you may submit a Tender Bid for Hospital Supply Status for that Tender Item on the basis that, if PHARMAC accepts your Tender Bid, the Tender Item would be listed in Section B and/or Part II of Section H of the Pharmaceutical Schedule subject to clause 11 of Schedule Six. This information is also shown in the Electronic Portal.

Where a Tender Item is indicated as being a "PCT" product, it is the preference of PHARMAC that products have post-compounding stability data greater than 48 hours.

1.12 Capsule and tablet form

Unless otherwise stated, where a Tender Item specifies either:

- (a) a capsule; or
- (b) a tablet,

form of the Chemical Entity, your brand of the relevant Chemical Entity for which you submit a bid may be in either tablet or capsule form, provided that:

- (c) your brand of the relevant Chemical Entity is the same strength as the Tender Item; and
- (d) where the Tender Item specifies both the tablet and capsule form of that Chemical Entity as separate line items, you must submit a bid for the same form and strength for each line item.

1.13 **Pack size preference**

Where a Tender Item is specified as being available for a Tender Bid for Sole Supply Status, it is the preference of PHARMAC that the pack size for such a Tender Item is a 30 or 90 day pack where the Tender Item is in a tablet or capsule form.

Notwithstanding the preference of PHARMAC for Tender Items to be in pack sizes as specified above, pack sizes may be specified in the comments column in the attached list or you may submit, and PHARMAC will consider and may accept, a Tender Bid for any pack size, including larger pack sizes, following its evaluation of Tender Bids under clause 5 of Schedule Three.

1.14 **Pack size for use in DHB Hospitals**

Where a Tender Item is specified as being available for a Tender Bid for Hospital Supply Status, it is the preference of DHB Hospitals that the pack size for such a Tender Item is:

- (a) 500 ml or less, where the Tender Item is in liquid form;
- (b) 200 tablets or capsules, where the Tender Item is in tablet or capsule form; and
- (c) 10 injections, where the Tender Item is in injection form.

Notwithstanding the preference of DHB Hospitals for Tender Items to be in pack sizes as specified in paragraphs (a) to (c) above, pack sizes may be specified in the comments column in the attached list or you may submit, and PHARMAC will consider and may accept, a Tender Bid for any pack size, including larger pack sizes, following its evaluation of Tender Bids under clause 5 of Schedule Three. For the avoidance of doubt, DHB Hospitals do not have a pack (or other equivalent grouping) size preference for Medical Devices and you may submit, and PHARMAC will consider and may accept, a Tender Bid for any pack size (or other equivalent grouping) following its evaluation of Tender Bids under clause 5 of Schedule Three.

1.15 **Pack size for oral contraceptives**

Where an oral contraceptive is included in Schedule Two and on the Electronic Portal, 21 and 28 calendar packs would be considered as different Tender Items (where applicable).

S	CHEDULE	TWO: PRO	DDUCTS TO	D BE TENDERED
Chemical Name			Unit	DV
Line Item	Units	Cost	Subsidy	Limit Comments
Acetazolamide				
<u>Tab 250 mg</u>	674,232	\$114,822	\$0.1703	CH 1%
Acitretin				
<u>Cap 10 mg</u>	483,243	\$143,847	\$0.2977	C H 1% Special Authority restrictions may apply. There may be a preference for a blister pack.
<u>Cap 25 mg</u>	110,664	\$76,284	\$0.6893	C H 1% Special Authority restrictions may apply. There may be a preference for a blister pack.
Ajmaline				
Inj 50 mg per 10 ml				H 1%
Albendazole				
Tab	728	\$5,693	\$7.8200	C H 1% Current restrictions may apply. Note units and unit subsidy corresponds to the currently listed tab 400 mg presentation. All strengths received as tender bids will be reviewed.
Alfacalcidol				
<u>Cap 0.25 mcg</u>	210,847	\$55,495	\$0.2632	C H 1%
<u>Cap 1 mcg</u>	59,093	\$51,990	\$0.8798	C H 1%
<u>Oral drops 2 mcg per ml</u>	14,240	\$43,204	\$3.0340	CH 1%
Alfentanil				
<u>lnj 0.5 mg per ml, 2 ml ampoule</u>				H 1%
Allopurinol				
<u>Tab 100 mg</u>	24,840,484	\$225,552	\$0.0091	C H 1% Preference for a scored tablet.
<u>Tab 300 mg</u>	18,222,228	\$377,200	\$0.0207	C H 1% Preference for a scored tablet.
Alprostadil				
Inj 10 mcg				H 1% Not currently listed in the Pharmaceutical Schedule. Restrictions may apply.
Inj 20 mcg				H 1% Not currently listed in the Pharmaceutical Schedule. Restrictions may apply.
Ambrisentan (current access)				
Tab 10 mg	8,726	\$1,333,624	\$152.8333#	C H 1% A confidential rebate applies. Longer transition periods may apply to this product. Current restrictions may apply. PHARMAC would only award a tender for either current or widened access
Tab 5 mg	2,064	\$315,448	\$152.8333#	C H 1% A confidential rebate applies. Longer transition periods may apply to this product. Current restrictions may apply. PHARMAC would only award a tender for either current or widened access
Ambrisentan (widened access)				
Tab 10 mg				C H 1% PHARMAC would only award a tender for either current or widened access
Tab 5 mg				C H 1% PHARMAC would only award a tender for either current or widened access
Amikacin				
Inj 15 mg per ml				H 1% Current restrictions may apply.
lnj 5 mg per ml				H 1% Current restrictions may apply.
Amiloride				
Tab 5 mg				C H 1% Not currently listed in Section B of the Pharmaceutical Schedule.
Amiloride with Frusemide				
Tab 5 mg with frusemide 40 mg	130,984	\$40,370	\$0.3082	C H 1%
Amiloride with Hydrochlorothiazide Tab 5 mg with hydrochlorothiazide 50 mg	776,892	\$77,689	\$0.1000	СН 1%
Aminophylline	-,	. ,	· · · · ·	
lnj 25 mg per ml, 10 ml	1,522	\$37,858	\$24.8740	СН 1%
sole supply	.,	÷=:,000	, .	#=rebate *=part charge @=ASP +=pate

S	CHEDULE	TWO: PRO	DUCTS 1		NDERE	ED
Chemical Name			Unit		DV	
Line Item	Units	Cost	Subsidy		Limi	t Comments
Amisulpride						
Oral liq	16,187	\$17,679	\$1.0922	С	H 1%	Units and subsidy shown are for oral liq 100 mg per ml presentation.
Amitriptyline						
<u>Tab 10 mg</u>	21,155,458	\$414,647	\$0.0196	С	H 1%	
<u>Tab 25 mg</u>	7,671,899	\$116,613	\$0.0152	С	H 1%	
<u>Tab 50 mg</u>	2,686,424	\$67,429	\$0.0251	С	H 1%	
Amlodipine						
<u>Tab 2.5 mg</u>	9,693,962	\$166,736	\$0.0172	С	H 1%	
<u>Tab 5 mg</u>	19,075,798	\$254,090	\$0.0133	С	H 1%	
<u>Tab 10 mg</u>	9,983,680	\$175,713	\$0.0176	С	H 1%	
Amorolfine						
Nail soln 5%	102,145	\$325,843	\$3.1900	С	H 1%	
Amoxicillin						
Grans for oral liq 125 mg per 5 ml	10,717,993	\$128,616	\$0.0120	С	H 1%	Preference may be given to products which include a measuring device.
Grans for oral liq 250 mg per 5 ml	52,349,459	\$685,778	\$0.0131	С	H 1%	Preference may be given to products which include a measuring device.
<u>Inj 250 mg</u>	83	\$89	\$1.0670	С	H 1%	
<u>Inj 500 mg</u>	1,039	\$1,289	\$1.2410	С	H 1%	
<u>Inj 1 g</u>	6,575	\$11,368	\$1.7290	С	H 1%	
Amoxicillin clavulanate						
Tab amoxycillin 500 mg with potassium clavulanate 125 mg	12,430,916	\$1,168,506	\$0.0940	@ C	H 1%	
Amphotericin B						
Inj 50 mg (non-liposomal)					H 1%	Current restrictions may apply.
Liposomal inj 50 mg					H 1%	Current restrictions may apply.
Anastrozole						
Tab 1 mg	836,608	\$140,550	\$0.1680	С	H 1%	
		+ - ,		-		
Atropine sulphate Eye drops 1%	116,370	\$134,678	\$1.1573	С	H 1%	
	110,010	φ104,010	ψ1.1070	U	11 170	
Aztreonam Inj 1 g					⊔ 10/	Current restrictions may apply
nij i g					H 1%	Current restrictions may apply.
Baclofen			• • • • • • • • •			
lnj 0.05 mg per ml, 1 ml			\$11.5500	С	H 1%	
Bee venom allergy treatment						
Inj with diluent	281	\$85,705	\$305.0000	С		
Maintenance kit	7	\$1,995	\$285.0000	С	H 1%	
Bendroflumethiazide [Bendrofluazide]						
<u>Tab 2.5 mg</u>	18,669,512	\$466,738	\$0.0250	С	H 1%	
<u>Tab 5 mg</u>	2,694,522	\$110,044	\$0.0408	С	H 1%	
Benzatropine Mesylate						
lnj 1 mg per ml, 2 ml	1,517	\$28,823	\$19.0000	С	H 1%	
Benzoyl peroxide						
Soln/Gel/Cream/Lotion 5%				С	H 1%	Note this product may require Medsafe registration prior to an award. Not currently listed in Section B of the Pharmaceutical Schedule.
Benzylpenicillin sodium [Penicillin G] Inj 600 mg	6,996	\$7,241	\$1.0350	С	H 1%	
Betahistine dihydrochloride						
Tab 16 mg	3,382,438	\$116,356	\$0.0344	С	H 1%	
-	-,552,100	÷,		5	1,5	
Betamethasone Inj 4 mg per ml, 1 ml				С	H 1%	Not currently listed in Section B of the Pharmaceutical Schedule.
sole supply					#	erebate *=part charge @=ASP +=pate

SC	HEDULE	TWO: PRO	DUCTS TO	D BE TEND	DERE	Ð
Chemical Name Line Item	Units	Cost	Unit Subsidy		DV Limi	t Comments
Betamethasone						
Tab 500 mcg				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Betamethasone Dipropionate				Н	1%	
Crm 0.05% (pack size 30 g or less)	205 050	¢50.400	¢0 4072	СН		
Crm 0.05% (pack size greater than 30 g)	295,950	\$58,400 \$28,200	\$0.1973	СН	1%	
Crm 0.05% in propylene glycol base	195,450	\$28,209	\$0.1443	Н	1%	
Oint 0.05% (pack size 30 g or less)	200,400	#05.040	¢0.4070	СН	1%	
Oint 0.05% (pack size greater than 30 g)	329,460	\$65,012	\$0.1973		1%	
Oint 0.05% in propylene glycol base	22,995	\$3,319	\$0.1443	СН	1%	
Betamethasone Sodium Phosphate with B Inj 3.9 mg with betamethasone acetate 3	etamethason 1,477	e Acetate \$5,672	\$3.8400	СН	1%	
mg per ml, 1ml						
Betaxolol hydrochloride						
Eye drops 0.25%	17,435	\$41,147	\$2.3600	СН	1%	For products containing BAK, PHARMAC reserves the right to list a BAK or preservative free product for a restricted market. Units and unit subsidy expressed as "per ml".
Eye drops 0.5%	7,990	\$11,985	\$1.5000	СН	1%	For products containing BAK, PHARMAC reserves the right to list a BAK or preservative free product for a restricted market. Units and unit subsidy expressed as "per ml".
Bicalutamide						
<u>Tab 50 mg</u>	588,300	\$79,838	\$0.1357	СН	1%	
Bismuth trioxide Tab	58,995	\$17,120	\$0.2902	СН	1%	Units and subsidy shown based on tab 120 mg presentation. All strengths reviewed as tender bids will be reviewed.
Bisoprolol Fumarate						
Tab 2.5 mg	7,453,023	\$292,308	\$0.0392	СН	1%	
Tab 5 mg	4,265,927	\$244,096	\$0.0572	СН	1%	
Tab 10 mg	1,540,958	\$160,938	\$0.1044	СН	1%	
Budesonide						
Nebuliser soln 250 mcg per ml, 2 ml ampoule				н	1%	
Metered aqueous nasal spray, 50 mcg per dose	3,292,200	\$42,634	\$0.0130	СН	1%	
Metered aqueous nasal spray, 100 mcg per dose	16,498,200	\$236,749	\$0.0144	СН	1%	
Nebuliser soln, 500 mcg per ml, 2 ml				н	1%	
Budesonide (current access)						
Cap 3 mg modified release	347,257	\$642,425	\$1.8500	СН	1%	Current restrictions may apply. PHARMAN would only award a tender for either current or widened access.
Budesonide (widened access)						
Cap 3 mg modified release				СН	1%	Restrictions may apply. PHARMAC would only award a tender for either current or widened access.
Bupivacaine hydrochloride						
Inj 2.5 mg per ml, 20 ml ampoule				н	1%	
Inj 5 mg per ml, 20 ml ampoule				н	1%	
Inj 1.25 mg per ml, 100 ml bag				н	1%	
Inj 1.25 mg per ml, 200 ml bag				Н	1%	
lnj 1.25 mg per ml, 500ml bag				Н	1%	
<u>lnj 2.5 mg per ml, 100 ml bag</u>				Н	1%	
lnj 2.5 mg per ml, 200 ml bag				Н	1%	
<u>lnj 5 mg per ml, 4 ml</u>				Н	1%	
sole supply					#	=rebate *=part charge @=ASP +=pater

SCHEDULE TWO: PRODUCTS TO BE TENDERED

	SCHEDULE 1	rwo: Pro	DUCTS TO	BE TEND	DERE	D
Chemical Name Line Item	Units	Cost	Unit Subsidy		DV Limi	t Comments
Bupivacaine hydrochloride			-			
Inj 0.25%, 20 ml sterile pack				Н	1%	
Inj 0.5%, 10 ml sterile pack				н	1%	
Inj 0.5%, 20 ml sterile pack				Н	1%	
Bupropion hydrochloride Tab modified-release 150 mg	2,330,945	\$854,687	\$0.3667	СН	1%	
Buserelin Acetate	,	**** /***				
lnj 1 mg per ml, 5.5 ml				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Calamine Lotn, BP	2 546 927	\$22,948	\$0.0065	СН	10/	Professore for a 200 ml pack or loss. Note
LUII, BP	3,546,827	¢∠2,946	\$U.UU65	СП	1%	Preference for a 200 ml pack or less. Note this product requires Medsafe registration prior to an award.
Calcipotriol						
Oint 50 mcg per g	1,488,200	\$669,690	\$0.4500	СН	1%	
Calcium carbonate						
Tab	7,608,693	\$228,869	\$0.0301	СН	1%	Preference for tab 1.25 g - 1.5 g.
Calcium folinate						
Tab 15 mg	15,421	\$160,779	\$10.4260	PCT C H	1%	
Calcium Gluconate						
Gel 2.5%				Н	1%	Preference for a pack size of 40 g - 100 g
lnj 10%, 10 ml	769	\$2,461	\$3.2000	СН	1%	
		φ=,	\$0. <u>_</u> 000	•	. ,0	
Capsaicin (current access) Crm 0.025%	1,604,450	\$363,087	\$0.2263#	СН	1%	Current restrictions may apply. Please
0.023 /6	1,004,430	ψ 303,00 7	ψ0.2203#	θH	170	note the widening of access refers to the crm 0.075% presentation only.
Crm 0.075%	347,580	\$96,551	\$0.2778	СН	1%	Current restrictions may apply. PHARMAC would only award a tender for either current or widened access.
Capsaicin (widened access)						
Crm 0.025%	1,604,450	\$363,087	\$0.2263#	СН	1%	Current restrictions may apply. Please note the widening of access refers to the crm 0.075% presentation only. This tender item is listed under widened access for the purpose of aggregate bidding in the e- portal.
Crm 0.075%				СН	1%	PHARMAC would only award a tender for either current or widened access. Widened access would result in the removal of all restrictions.
Captopril Oral liq 5 mg per ml	157,415	\$157,398	\$0.9999	СН	1%	Current restrictions may apply.
Carbimazole						
Tab 5 mg	4,364,939	\$471,413	\$0.1080	СН	1%	
Carboprost						
lnj 250 mcg per ml, 1ml				Н	1%	
Carmellose sodium						
Paste	25,888	\$7,871	\$0.3040 *	СН	1%	Preference for product containing gelatin and pectin. Units subsidy and cost shown is weighted.
Powder	1,288	\$390	\$0.3029	CH	1%	Preference for product containing gelatin and pectin.
Eye drops 0.5%				CH	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Eye drops 1%				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Carvedilol						
Tab 12.5 mg	2,299,938	\$88,157	\$0.0383	СН	1%	
Tab 25 mg	2,736,669	\$134,562	\$0.0492	СН	1%	
sole supply		·				=rebate *=part charge @=ASP +=patent

SCHEDULE TWO: PRODUCTS TO BE TENDERED						
Chemical Name			Unit	DV		
Line Item	Units	Cost	Subsidy	Limit Comments		
Carvedilol						
<u>Tab 6.25 mg</u>	2,410,457	\$89,982	\$0.0373	C H 1%		
Cefazolin sodium						
<u>Inj 500 mg</u>	130	\$88	\$0.6780	CH 1%		
<u>lnj 1 g</u>	25,845	\$17,006	\$0.6580	C H 1%		
Cefotaxime						
Inj 500 mg				H 1% H 1%		
lnj 1 g				H 1%		
Cefoxitin Sodium Inj 1 g				Н 1%		
				11 170		
Ceftaroline fosamil Inj 600 mg vial				H 1% Current restrictions may apply.		
Ceftazidime Inj 1 g				H 1% Current restrictions may apply.		
Cefuroxime sodium Inj 750 mg				Н 1%		
<u>lni 1.5 g</u>				H 1%		
Chloramphenicol						
Inj 1 g				H 1% Current restrictions may apply.		
Chlorhexidine gluconate						
Crm 1% obstetric				H 1% Preference for a small tube for single use		
				under 100 g. Note that this product requires Medsafe registration prior to an award.		
Lotn 1% obstetric				H 1% Note that this product requires Medsafe registration prior to an award.		
Soln 4%	2,358,497	\$18,774	\$0.0080	C H 1% 500mL pump preferred. Note that this product requires Medsafe registration pric to an award.		
Chlorhexidine with ethanol						
Soln 2% with ethanol 70%, non-staining (pink) (pack size less than 50 ml)				H 1%		
Soln 2% with ethanol 70%, non-staining (pink) 100 ml				H 1%		
Soln 2% with ethanol 70%, staining (red) (pack size less than 50 ml)				H 1%		
Soln 2% with ethanol 70%, staining (red) 100 ml				H 1%		
Chloroquine phosphate						
Tab 250 mg				H 1% Current restrictions may apply.		
Chlorpheniramine Maleate						
Inj 10 mg per ml	0 705 100	• • • • -= •	AC 015	H 1%		
Oral liq 2 mg per 5 ml	2,765,123	\$44,574	\$0.0161	CH 1%		
Choline salicylate	00.400	¢40 700	#0.4070			
Oral gel	93,120	\$12,788	\$0.1373	CH 1%		
Cidofovir				H 1% Current restrictions may apply		
lnj 75 mg per ml, 5 ml				H 1% Current restrictions may apply.		
Ciprofloxacin Ear drops 0.3%				C H 1% Not currently listed in the Pharmaceutical Schedule. Any listing may be subject to a Special Authority restriction		
<u>Tab 250 mg</u>	183,732	\$9,515	\$0.0518	C H 1%		
<u>Tab 500 mg</u>	1,047,036	\$9,515 \$74,413	\$0.0518 \$0.0711	СН 1%		
<u>Tab 750 mg</u>	24,706	\$2,779	\$0.1125	CH 1%		

S	CHEDULE	TWO: PRO	DUCTS T	O BE TENDERED
Chemical Name	•		Unit	DV
Line Item	Units	Cost	Subsidy	Limit Comments
Citric acid with magnesium oxide and so	dium picosulfa	ate		
Powder for oral soln 12 g with magnesiun oxide 3.5 g and sodium picosulfate 10 mg per sachet	n I			H 1%
Clarithromycin				
lnj 500 mg				H 1% Current restrictions may apply.
<u>Tab 250 mg</u>	151,953	\$43,199	\$0.2843	C H 1% Current restrictions may apply.
<u>Tab 500 mg</u>	127,685	\$94,852	\$0.7429	C H 1% Current restrictions may apply.
Clobazam				
Liq				C H 1% Not currently listed in the Pharmaceuti Schedule.
Clobetasone butyrate				
Crm 0.05% (pack size 30 g or less)	450,780	\$80,838	\$0.1793	C H 1%
Clonidine				
TDDS 2.5 mg, 100 mcg per day	53,888	\$99,693	\$1.8500	C H 1%
TDDS 5 mg. 200 mcg per day	22,490	\$56,450	\$2.5100	C H 1%
TDDS 7.5 mg, 300 mcg per day	15,264	\$47,089	\$3.0850	C H 1%
Clotrimazole				
<u>Crm 1%</u>	3,523,060	\$123,307	\$0.0350	C H 1%
Soln 1%	13,240	\$2,886	\$0.2180	C H 1%
Codeine phosphate				
Tab 15 mg	10,532,470	\$605,617	\$0.0575	C H 1%
Tab 30 mg	25,125,868	\$1,708,559	\$0.0680	CH 1%
Tab 60 mg	1,665,297	\$224,815	\$0.1350	CH 1%
Colecalciferol				
Cap/Tab 1.25 mg	2,784,346	\$580,063	\$0.2083	CH 1%
Colistin Sulphomethate				
lnj 150 mg	6,133	\$398,645	\$65.0000	C H 1% Current restrictions may apply.
Condoms				
Female, non-latex				C Not currently listed in Section B of the Pharmaceutical Schedule.
Male 55 mm - 58 mm, non-latex				C Not currently listed in Section B of the Pharmaceutical Schedule. Special Authority restrictions may apply.
Cyclopentolate hydrochloride				
Eye drops 1%	81,210	\$47,427	\$0.5840	CH 1%
Cycloserine				
Cap 250 mg	5,185	\$67,120	\$12.9450	C H 1% Current restrictions may apply. Unit subsidy and cost shown is weighted.
Cyproterone acetate with ethinyloestradi		A	A A A A A A	
Tab 2 mg with ethinyloestradiol 35 mcg and 7 inert tablets	10,692,612	\$297,255	\$0.0278	CH 1%
Danazol				
Cap 100 mg	17,703	\$12,096	\$0.6833	C H 1%
Cap 200 mg	9,996	\$9,779	\$0.9783	C H 1%
Darunavir				
<u>Tab 400 mg</u>	93,157	\$520,126	\$5.5833	C H 1% Current restrictions may apply.
<u>Tab 600 mg</u>	33,847	\$268,519	\$7.9333	C H 1% Current restrictions may apply.
Desmopressin				
lnj 15 mcg per ml, 1 ml				H 1%
Inj 4 mcg per ml, 1 ml	4,984	\$33,483	\$6.7180	C H 1% Current restrictions may apply.
Nasal drops 100 mcg per ml	6,093	\$95,116	\$15.6120	C H 1%
Nasal spray 10 mcg per dose	87,816	\$350,532	\$3.9917	C H 1%
Tab 100 mcg	92,684	\$77,236	\$0.8333	C H 1% Current restrictions may apply.
Tab 200 mcg	38,877	\$70,562	\$1.8150	C H 1% Current restrictions may apply.
sole supply				#=rebate *=part charge @=ASP +=pa

5	SCHEDULE .	TWO: PRO	DUCTS TO	D BE TENI	DERE	D
Chemical Name Line Item	Units	Cost	Unit Subsidy		DV Limi	t Comments
Dexmedetomidine hydrochloride						
<u>Inj 100 mcg per ml, 2 ml vial</u>				+ H	1%	Patent may exist for this product.
Dextrochlorpheniramine Maleate						
Oral liq 2 mg per 5 ml	8,780	\$155	\$0.0177	СН	1%	Funding restrictions may apply. Not currently listed in Section H of the Pharmaceutical Schedule.
Tab 2 mg	144,440	\$7,294	\$0.0505 *	СН	1%	Funding restrictions may apply. Not currently listed in Section H of the Pharmaceutical Schedule.
Diazepam						
Tab 2 mg	2,961,780	\$89,150	\$0.0301	СН	1%	There may be a preference for a scored tablet.
<u>Tab 5 mg</u>	4,355,885	\$140,956	\$0.0324	СН	1%	
Diazoxide						
Cap 100 mg	5,620	\$15,736	\$2.8000	СН	1%	Current restrictions may apply.
Cap 25 mg	12,263	\$13,489	\$1.1000	СН	1%	Current restrictions may apply.
Inj 15 mg per ml, 20 ml				н	1%	
Oral liq 50 mg per ml	1,020	\$21,080	\$20.6667	СН	1%	Current restrictions may apply.
Diclofenac sodium						· · · ·
lnj 25 mg per ml, 3 ml	26,367	\$69,609	\$2.6400	СН	1%	
Suppos 12.5 mg	6,197	\$1,264	\$0.2040	СН	1%	
Suppos 25 mg	5,132	\$1,252	\$0.2440	СН	1%	
Suppos 50 mg	75,572	\$31,891	\$0.4220	СН	1%	
Suppos 100 mg	107,910	\$75,537	\$0.7000	СН	1%	
Diflucortolone Valerate	,	, 2,20.		0.1		
Crm 0.1%	87,600	\$15,715	\$0.1794	СН	1%	
Fatty oint 0.1%	101,550	\$18,218	\$0.1794 \$0.1794	СН	1%	
Diltiazem hydrochloride	,	¢:0,2:0	<i>Q</i> OILIOI	• • •	. , 0	
Tab 30 mg	569,392	\$26,192	\$0.0460	СН	1%	
Tab 60 mg	248,826	\$21,150	\$0.0850	СН	1%	
Dinoprostone	,	+=-;-==				
Gel 1 mg				н	1%	
Gel 2 mg				н	1%	
Ű					170	
Disulfiram Tab 200 mg	409,347	\$309,343	\$0.7557	СН	1%	
ů.	+00,047	φ000,0 1 0	ψ 0.1001	OT	170	
Docetaxel Inj 20 mg				РСТ Н	1%	Preference for products where stability data >48 hours post-compounding.
<u>lnj 80 mg</u>				РСТ Н	1%	Preference for products where stability data >48 hours post-compounding.
Docusate Sodium						and a second been been been allowed as
Tab 50 mg	624,983	\$14,437	\$0.0231	СН	1%	
<u>Tab 120 mg</u>	1,457,203	\$45,610	\$0.0231	СН	1%	
-	1,101,200	φ10,010	<i>Q</i> 0.0010	011	170	
Donepezil hydrochloride Tab 5 mg	1,239,193	\$59,754	\$0.0482	СН	1%	
<u>Tab 10 mg</u>	1,127,992	\$39,734 \$83,223	\$0.0482 \$0.0738	СН	1%	
Dosulepin [Dothiepin] Hydrochloride	·,·=·,002	Ψ Ο Ο,ΖΖΟ	40.0100	011	170	
Cap 25 mg	2,478,283	\$159,849	\$0.0645	СН	1%	
Tab 75 mg	894,920	\$109,849 \$100,142	\$0.0045 \$0.1119	СН	1%	
-	507,020	φ100,1 1 2	ψ0.1110	OT	170	
Doxazosin mesylate Tab 2 mg	16,503,612	\$222,799	\$0.0135	СН	1%	
<u>Tab 2 mg</u> Tab 4 mg	12,247,469	\$222,799 \$222,659	\$0.0135 \$0.0182	СН	1%	
-	12,277,403	ΨΖΖΖ,009	ψ0.0102	СH	1 /0	
Doxepin hydrochloride	1 107 500	¢74 600	¢0.0690	0.11	4.07	
Cap 10 mg	1,137,560 1,313,518	\$71,666 \$90,107	\$0.0630 \$0.0686	СН СН	1% 1%	
Cap 25 mg	1,313,518	\$90,107	ФО.0080	Сп		
sole supply					#	=rebate *=part charge @=ASP +=paten

sole supply

S	CHEDULE	TWO: PRO	DUCTS T		NDERI	ED
Chemical Name Line Item	Units	Cost	Unit Subsidy		DV Limi	it Comments
Doxepin hydrochloride	• · · · · ·		Cuboluy			
Cap 50 mg	663,982	\$56,770	\$0.0855	С	H 1%	
Doxycycline	,	. ,				
Inj 100 mg per 20 ml					H 1%	
Tab 20 mg - 50 mg	335,923	\$32,474	\$0.0967	С	H 1%	Note units and unit price expressed is for the 50 mg presentation.
Econazole Nitrate						
Crm 1%	26,780	\$1,339	\$0.0500	С		
Foaming soln 1%, 10 ml sachets	59,486	\$196,106	\$3.2967	С	H 1%	
Emulsifying ointment						
Oint BP (pack size of 200 g or less)	00.070.455		* ~ ~~ ~		H 1%	
<u>Oint BP (pack size greater than 200 g)</u>	38,879,455	\$279,154	\$0.0072	С	H 1%	
Ephedrine					11 40/	
Inj 3 mg per ml, 10 ml prefilled syringe Inj 30 mg per ml, 1ml					H 1% H 1%	
					11 170	
Epoprostenol (current access) Inj 1.5 mg vial	330	\$24,159	\$73.2100	С	H 1%	Current restrictions may apply. PHARMAC
ing its ing via	550	ψ24,109	ψ/ 3.2100	C	11 170	would only award a tender for either current or widened access
Inj 500 mcg vial	30	\$1,098	\$36.6100	С	Η	Current restrictions may apply. PHARMAC would only award a tender for either current or widened access.
Epoprostenol (widened access)						
Inj 1.5 mg vial				С	H 1%	PHARMAC would only award a tender for either current or widened access.
Inj 500 mcg vial				С	H 1%	PHARMAC would only award a tender for either current or widened access.
Ergometrine maleate						
Inj 250 mcg per ml, 1 ml ampoule	200			С		
<u>lnj 500 mcg per ml, 1 ml</u>	1,140	\$23,940	\$21.0000	С	H 1%	
Erlotinib hydrochloride		A	*			2
Tab 100 mg	15,260	\$388,628	\$25.4667	+ C		Current restrictions may apply.
Tab 150 mg	22,313	\$852,357	\$38.2000	+ C	H 1%	Current restrictions may apply.
Erythromycin Ethyl Succinate	0.045.050	A 445 750	\$ 0.0500	0	11 404	B (
Grans for oral liq 200 mg per 5 ml Grans for oral liq 400 mg per 5 ml	2,915,052	\$145,753 \$241,121	\$0.0500 \$0.0677	c c		which include a measuring device.
						which include a measuring device.
Tab 400 mg	3,217,984	\$545,448	\$0.1695	С	H 1%	
Erythromycin Lactobionate	000	#0 500	¢40.0000	6	11 404	
lnj 1 g	223	\$3,568	\$16.0000	С	H 1%	
Escitalopram Tab 10 mg	10,418,137	\$412,975	\$0.0396	С	H 1%	, ,
<u>Tab 20 mg</u>	3,883,752	\$263,551	\$0.0679	С	H 1%	tablet.
Ethambutol hydrochloride						
Tab 100 mg	33,493	\$28,714	\$0.8573	С	H 1%	Current restrictions may apply.
Tab 400 mg	124,390	\$109,596	\$0.8811	С	H 1%	Current restrictions may apply.
Ethinyloestradiol with Desogestrel						
Tab 20 mcg with desogestrel 150 mcg	284,704	\$22,438	\$0.0788	С	H 1%	
Tab 30 mcg with desogestrel 150 mcg	274,652	\$21,645	\$0.0788	С	H 1%	
Ethinyloestradiol with levonorgestrel						
Tab 20 mcg with levonorgestrel 100 mcg	6,957,468	\$180,487	\$0.0259	С	H 1%	Note units and unit subsidy expressed are based on a product inclusive of 7 inert tablets (per month).

SCHEDULE TWO: PRODUCTS TO BE TENDERED								
Chemical Name Line Item	Units	Cost	Unit Subsidy		DV Limi	t Comments		
Ethinyloestradiol with levonorgestrel								
Tab 30 mcg with levonorgestrel 150 mcg	L 27,088,040	\$570,858	\$0.0211	СН	1%	Note units and unit subsidy expressed are based on a product inclusive of 7 inert tablets (per month).		
Tab 50 mcg with levonorgestrel 125 mcg	399,868	\$44,985	\$0.1125	СH	1%	Note units and unit subsidy expressed are based on a product inclusive of 7 inert tablets (per month).		
Etoposide								
lnj 20 mg per ml, 5 ml		\$11,091	\$7.9000	РСТ С Н	1%	PHARMAC reserves the right to award a tender for the whole market for either etoposide or etoposide phosphate.		
Etoposide phosphate								
Inj 100 mg (of etoposide base)				PCT H	1%	Preference for products where stability data >48 hours post-compounding. PHARMAC reserves the right to award a tender for the whole market for either etoposide or etoposide phosphate.		
Etravirine								
Tab 200 mg	38,896	\$499,165	\$12.8333#	СН	1%	Special Authority resctrictions may apply.		
Exemestane								
<u>Tab 25 mg</u>	362,963	\$175,431	\$0.4833	СН	1%			
Ezetimibe (current access)								
<u>Tab 10 mg</u>	7,156,369	\$477,115	\$0.0667	СН	1%	Current restrictions may apply. PHARMAG would only award a tender for either current or widened access		
Ezetimibe (widened access)								
Tab 10 mg				СН	1%	PHARMAC would only award a tender for either current or widened access. Widened access would result in the removal of Special Authority criteria.		
Ezetimibe with simvastatin (current acco		A (A) A =	AA 1 - 1 - 1	.	1.07			
Tab 10 mg with simvastatin 10 mg	59,038	\$10,135	\$0.1717	СН	1%	Current restrictions may apply. PHARMAG would only award a tender for either current or widened access		
Tab 10 mg with simvastatin 20 mg	93,135	\$19,093	\$0.2050	CH	1%	Current restrictions may apply. PHARMAG would only award a tender for either current or widened access		
Tab 10 mg with simvastatin 40 mg	200,811	\$47,859	\$0.2383	СН	1%	Current restrictions may apply. PHARMAG would only award a tender for either current or widened access		
Tab 10 mg with simvastatin 80 mg	167,507	\$45,507	\$0.2717	СH	1%	Current restrictions may apply. PHARMAG would only award a tender for either current or widened access		
Ezetimibe with simvastatin (widened ac	cess)							
Tab 10 mg with simvastatin 10 mg				СН	1%	PHARMAC would only award a tender for either current or widened access. Widened access would result in the removal of all restrictions.		
Tab 10 mg with simvastatin 20 mg				СH	1%	PHARMAC would only award a tender for either current or widened access. Widened access would result in the removal of all restrictions.		
Tab 10 mg with simvastatin 40 mg				СH	1%	PHARMAC would only award a tender for either current or widened access. Widened access would result in the removal of all restrictions.		
Tab 10 mg with simvastatin 80 mg				СH	1%	PHARMAC would only award a tender for either current or widened access. Widened access would result in the removal of all restrictions.		
Fentanyl								
Patches 100 mcg per hour	31,214	\$71,168	\$2.2800	СН	1%			
Patches 12.5 mcg per hour	194,227	\$114,594	\$0.5900	СН	1%			
Patches 25 mcg per hour	151,062	\$110,577	\$0.7320	СН	1%			
sole supply					#	=rebate *=part charge @=ASP +=pater		

SCHEDULE TWO: PRODUCTS TO BE TENDERED							D
Chemical Name			Unit			DV	
Line Item	Units	Cost	Subsidy		I	Limi	t Comments
Fentanyl							
Patches 50 mcg per hour	62,028	\$82,497	\$1.3300		СН	1%	
Patches 75 mcg per hour	17,633	\$32,621	\$1.8500		СН	1%	
Fexofenadine							
Tab 120 mg	145,558	\$68,989	\$0.4740 *		СН	1%	
Tab 180 mg					СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Tab 60 mg	68,166	\$14,792	\$0.2170		СН	1%	
Finasteride							
<u>Tab 5 mg</u>	4,438,091	\$213,472	\$0.0481		СН	1%	Preference for a blister pack. May be subject to Special Authority restrictions.
Flucloxacillin sodium							
<u>Inj 250 mg</u>	120	\$108	\$0.9000		СН	1%	
<u>Inj 500 mg</u>	1,474	\$1,386	\$0.9400		СН	1%	
<u>lnj 1 g</u>	4,696	\$4,903	\$1.0440		СН	1%	
Fluconazole							
Powder for oral suspension 10 mg per ml	22,615	\$63,645	\$2.8143		СН	1%	
<u>Cap 50 mg</u>	101,639	\$7,586	\$0.0746		СН	1%	
<u>Cap 150 mg</u>	68,861	\$22,724	\$0.3300		СН	1%	
<u>Cap 200 mg</u>	62,042	\$11,256	\$0.1814		СН	1%	
Flucytosine							
Cap 500 mg					н	1%	Current restrictions may apply.
						. /0	
Fludrocortisone Acetate	1 242 540	¢400.050	¢0.4422		СН	1%	
Tab 100 mcg	1,342,549	\$192,253	\$0.1432		Сп	170	
Flumetasone pivalate with clioquinol Ear drops 0.02% with clioquinol 1%	40,125	\$23,861	\$0.5947		СН	1%	
Fluorometholone							
Eye drops 0.1%	155,570	\$94,140	\$0.6051		СН	1%	For products containing BAK, PHARMAC reserves its right to list a BAK or preservative free product for a restricted market. Unit subsidy shown is weighted and expressed as "per ml"
Flupenthixol Decanoate							
Inj 100 mg per ml, 1 ml	8,044	\$65,752	\$8.1740		СН	1%	
lnj 20 mg per ml, 1 ml	3,823	\$10,047	\$2.6280		СН	1%	
lnj 20 mg per ml, 2 ml	4,438	\$18,551	\$4.1800		СН	1%	
Foscarnet sodium	,	. ,					
Inj 24 mg per ml, 250 ml					н	1%	Current restrictions may apply.
Fosfomycin							
Powder for oral solution, 3 g sachet					Н	1%	Current restrictions may apply.
Framycetin sulphate							
Ear/Eye drops 0.5%	22,152	\$11,436	\$0.5163		СН	1%	
Fulvestrant	,	, ,					
Inj 250 mg per ml				+	СН	1%	Not currently listed in the Pharmaceutical Schedule.
Fusidic acid							
Tab 250 mg	11,023	\$31,691	\$2.8750		СН	1%	Current restrictions may apply.
Gefitinib	, -					-	2 11 2
Tab 250 mg				+	СН	1%	Not currently listed in the Pharmaceutical Schedule. Current restrictions may apply.
Gemeitabine bydrochloride							
Gemcitabine hydrochloride Inj 1 g				PCT	н	1%	
				101		1 /0	
Gemfibrozil	000 470	\$405,000	¢0,0000		C 11	4.07	
Tab 600 mg	386,478	\$125,992	\$0.3260		СН	1%	

S	CHEDULE	TWO: PRO	DDUCTS T		RED
Chemical Name Line Item	Units	Cost	Unit Subsidy	D Lir	V nit Comments
Gentamicin Sulphate					
Inj 10 mg per ml	1,802	\$9,010	\$5.0000	CH 1	%
lnj 40 mg per ml, 2 ml	13,495	\$8,097	\$0.6000	CH 1	%
Gestrinone					
Cap 2.5 mg				H 1'	%
Gliclazide					
<u>Tab 80 mg</u>	27,065,762	\$557,013	\$0.0206	CH 1	%
Glucagon Hydrochloride					
Inj 1 mg syringe kit	20,608	\$659,456	\$32.0000	CH 1	%
Glucose [Dextrose]					
Gel 40%				H 1	
<u>lnj 50%, 10 ml - 20 ml</u>	11,130	\$65,667	\$5.9000	CH 1	% Inj 50%, 10 ml is currently listed on the Pharmaceutical Schedule. Units and subsidy are shown per 10 ml ampoule.
<u>lnj 50%. 90 ml - 100 ml</u>	377	\$5,467	\$14.5000	CH 1	% Inj 50%, 90 ml is currently listed on the Pharmaceutical Schedule. Units and subsidy are shown per 90 ml ampoule.
Solution 15 g				H 1	% Not currently listed in the Pharmaceutical Schedule.
Tab 1.5 g				H 1	%
Tab 3.1 g				H 1	%
Tab 4 g				H 1'	%
Glucose with sucrose and fructose					
Gel 19.7% with sucrose 35% and fructose 19.7%, 18 g sachet				H 1 ⁴	%
Glycerol					
<u>Liq BP</u>	1,432,643	\$9,398	\$0.0066	CH 1	% There may be a preference for a pack size of 500 mL or less.
Glyceryl trinitrate					
Spray 400 mcg per dose	1,515,650	\$33,723	\$0.0223		%
lnj 1 mg per ml, 5 ml - 10 ml				H 1	%
lnj 5 mg per ml, 10 ml				H 1'	%
Goserelin					
Implant 10.8 mg, syringe	22,220	\$3,944,050	\$177.5000	CH 1	
Implant 3.6 mg, syringe	7,886	\$524,261	\$66.4800	CH 1	%
Granisetron					
<u>Inj 1 mg per ml</u>				H 1'	%
Hexamine hippurate					
Tab 1 g	588,841	\$108,347	\$0.1840	CH 1	%
Hydrocortisone					
Powder	54,663	\$109,217	\$1.9980	CH 1	% Current restrictions may apply. Note this product requires Medsafe registration prior to an award.
Hydrocortisone Acetate					
Rectal foam 10%, CFC-free	196,019	\$246,649	\$1.2583	CH 1	%
Hydrocortisone butyrate					
Cream/lipocream 0.1% (pack size 30 g or less)	1,782,150	\$122,077	\$0.0685	СН 1	
Cream/lipocream 0.1% (pack size greater than 30 g)	21,082,000	\$2,403,348	\$0.1140	CH 19	%
Hydrocortisone with wool fat and mineral					
Lotn 1% with wool fat hydrous 3% and mineral oil	10,698,213	\$452,320	\$0.0423	CH 19	% Preference for a pack size of 300 g or less.

sc	HEDULE	TWO: PRO	DUCTS TO	BE TEND	ERE	D
Chemical Name		•	Unit		DV	•
Line Item	Units	Cost	Subsidy		Limi	t Comments
Hydrogen Peroxide						
Crm 1%	2,416,630	\$1,706,701	\$0.7062#	СН	1%	Preference for a pack size of 50 g or more. Preference for tube packaging. Unit subsidy shown is weighted. This product requires Medsafe registration prior to an award.
Soln 3% (10 vol)	20,902	\$293	\$0.0140	СН	1%	Preference for pack size less than 200 ml. This product requires Medsafe registration prior to an award.
Hydroxyurea						
Cap 500 mg	1,180,903	\$375,055	\$0.3176	PCT C H	1%	
Hyoscine N-butylbromide						
Inj 20 mg, 1 ml	162,145	\$310,346	\$1.9140	СН	1%	
<u>Tab 10 mg</u>	5,812,813	\$508,621	\$0.0875	СН	1%	
Hypromellose						
Eye drops 0.3%	3,207,270	\$491,771	\$0.1533	СН	1%	Will consider bids for products with or without dextran. Note units and unit subsidy are expressed for a product with dextran 0.1%.
Eye drops 0.5%	49,530	\$6,604	\$0.1333	СН	1%	
	,500	40,00 r	÷	011	. ,0	
Ibuprofen Tab 200 mg	55,371,352	\$648,399	\$0.0117	СН	1%	
-	55,571,552	ψ040,099	φ0.011 <i>1</i>	UII	1 70	
Imatinib mesilate	00.000	¢ 450 700	* 0 5000 <i>"</i>	0.11	4.07	
<u>Cap 400 mg</u>	68,920	\$453,723	\$6.5833#	СН		For non-GIST indications only.
<u>Cap 100 mg</u>	194,035	\$316,923	\$1.6333#	СН	1%	For non-GIST indications only.
Imipramine Hydrochloride						
Tab 10 mg	800,985	\$87,788	\$0.1096	СН	1%	
Tab 25 mg	433,685	\$76,329	\$0.1760	СН	1%	
Imiquimod						
<u>Crm 5%</u>	466,774	\$422,430	\$0.9050	СН	1%	Units are per sachet
Indapamide						
Tab 2.5 mg	1,491,369	\$43,086	\$0.0289	СН	1%	
Insulin isophane						
Inj insulin human 100 u per ml, 10 ml	47,080	\$83,237	\$1.7680	СН	1%	
Insulin neutral	,	····				
Inj human 100 u per ml, 10 ml	22,240	\$56,178	\$2.5260	СН	1%	
	22,240	<i>4</i> 50,170	ψ2.0200	011	170	
Insulin pen needles	206 200	¢20.071	\$0.1050	С		
29 g x 12.7 mm	286,390 4,209,146	\$30,071 \$404,575	\$0.1050 \$0.1175			
31 g x 5 mm		\$494,575 \$26,072		C		
31 g x 6 mm	283,922	\$26,973 \$452,200	\$0.0950 \$0.1050	C		
31 g x 8 mm	4,307,520	\$452,290 \$462,568	\$0.1050	C		
32 g x 4 mm	4,405,406	\$462,568	\$0.1050	С		
Insulin syringes, disposable with attached	Ineedle					
Syringe 0.3 ml with 29 g x 4 mm needle				С		Not currently listed in the Pharmaceutical Schedule.
Syringe 0.3 ml with 29 g x 6 mm needle				С		Not currently listed in the Pharmaceutical Schedule.
Syringe 0.5 ml with 29 g x 4 mm needle				С		Not currently listed in the Pharmaceutical Schedule.
Syringe 0.5 ml with 29 g x 6 mm needle				С		Not currently listed in the Pharmaceutical Schedule.
Syringe 1 ml with 29 g x 4 mm needle				С		Not currently listed in the Pharmaceutical Schedule.
Syringe 1 ml with 29 g x 6 mm needle				С		Not currently listed in the Pharmaceutical Schedule.
Syringe 0.3 ml with 29 g x 12.7 mm needle	12,879	\$1,674	\$0.1300 *	С		
Syringe 0.3 ml with 31 g x 8 mm needle	121,925	\$15,852	\$0.1300 *	С		

S	CHEDULE	TWO: PRO	DUCTS T	O BE TENI	DERE	Ð
Chemical Name		a (Unit		DV	
Line Item	Units	Cost	Subsidy		Limi	t Comments
Insulin syringes, disposable with attached Syringe 0.5 ml with 29 g x 12.7 mm needle		\$5,081	\$0.1300	* C		
Syringe 0.5 ml with 31 g x 8 mm needle	126,997	\$16,511	\$0.1300	С		
Syringe 1 ml with 29 g x 12.7 mm needle	25,069	\$3,260	\$0.1300	С		
Syringe 1 ml with 31 g x 8 mm needle	46,277	\$6,016	\$0.1300	С		
lodine Soln aqueous BP				Н	1%	
Iodine supplement						
Tab 150 mcg elemental	10,375,823	\$540,684	\$0.0521	СН	1%	
Ipratropium Bromide						
Aqueous nasal spray, 0.03%	818,595	\$251,579	\$0.3073	СН	1%	
Isosorbide mononitrate						
Tab 20 mg	519,256	\$97,620	\$0.1880	СН	1%	
Tab long-acting 40 mg	613,287	\$153,322	\$0.2500	СН	1%	
Tab long-acting 60 mg	7,038,578	\$648,323	\$0.0921	СН	1%	
Ispaghula (psyllium) husk						
Powder/granules	84,651,000	\$1,024,277	\$0.0121	СН	1%	Units and unit subsidy shown are as per the 500 g presentation currently listed and expressed as 'per g'.
Ivabradine (current access)						
Tab 5 mg				Н	1%	
Ivabradine (widened access)						
Tab 5 mg				Н	1%	PHARMAC would only award a tender for either current or widened access.
Ivermectin						
Tab 3 mg	12,656	\$54,421	\$4.3000	СН	1%	Current restrictions may apply.
Ketoconazole						
Shampoo 2%	6,161,300	\$184,223	\$0.0299	СН	1%	Current restrictions may apply.
Tab 200 mg	1,650			РСТ С Н	1%	Current restrictions may apply.
Lacosamide (current access)						
lnj 10 mg per ml, 20 ml				Н	1%	
Tab 50 mg	202,558	\$362,289	\$1.7886	СН	1%	Special Authority restrictions may apply. PHARMAC would only award a tender for either current or widened access. Refer to Schedule 7 for additional special terms.
Tab 100 mg	130,316	\$465,970	\$3.5757	СН	1%	Special Authority restrictions may apply. PHARMAC would only award a tender for either current or widened access. Refer to Schedule 7 for additional special terms.
Tab 150 mg	35,532	\$190,604	\$5.3643	СН	1%	Special Authority restrictions may apply. PHARMAC would only award a tender for either current or widened access. Refer to Schedule 7 for additional special terms.
Tab 200 mg	90,383	\$646,481	\$7.1527	СН	1%	
Lacosamide (widened access)						
Tab 50 mg				СН	1%	, , , ,
						only award a tender for either current or widened access. Widened access would result in a change from sixth to fifth line treatment. Refer to Schedule 7 for additional special terms.
Tab 100 mg				СН	1%	Restrictions may apply. PHARMAC would only award a tender for either current or widened access. Widened access would result in a change from sixth to fifth line treatment. Refer to Schedule 7 for additional special terms.
sole supply						-rehate *-nart charge @-ASP +-naten

SC	HEDULE	TWO: PRO	DDUCTS TO	D BE TENDER	ED
Chemical Name Line Item	Units	Cost	Unit Subsidy	DV Lim	
Lacosamide (widened access)					
Tab 150 mg				СН 1%	Restrictions may apply. PHARMAC would only award a tender for either current or widened access. Widened access would result in a change from sixth to fifth line treatment. Refer to Schedule 7 for additional special terms.
Tab 200 mg				СН 1%	Restrictions may apply. PHARMAC would only award a tender for either current or widened access. Widened access would result in a change from sixth to fifth line treatment. Refer to Schedule 7 for additional special terms.
Lamivudine					
Tab 300 mg				CH 1%	Not currently listed on the Pharmaceutical Schedule. Special Authority restrictions may apply.
<u>Tab 100 mg</u>	167,182	\$25,077	\$0.1500	CH 1%	Special Authority restrictions may apply.
Tab 150 mg	24,705	\$21,617	\$0.8750	CH 1%	 Special Authority restrictions may apply. Not currently listed in Section H of the Pharmaceutical Schedule.
Leflunomide					
<u>Tab 10 mg</u>	543,182	\$52,509	\$0.0967	CH 1%	
<u>Tab 20 mg</u>	724,984	\$70,084	\$0.0967	CH 1%	5
Levocabastine					
Eye drops 0.5 mg per ml	94,116	\$204,938	\$2.1775	CH 1%	For products containing BAK, PHARMAC reserves its right to list a BAK or preservative free product for a restricted market. Units and unit subsidy shown are per ml
Levodopa with Carbidopa					
Tab 100 mg with carbidopa 25 mg	8,765,447	\$1,575,151	\$0.1797	CH 1%	Longer transition periods may apply to this product.
Tab 250 mg with carbidopa 25 mg	381,297	\$124,570	\$0.3267	CH 1%	product.
Tab long-acting 200 mg with carbidopa 50 mg	1,719,226	\$638,692	\$0.3715	CH 1%	Longer transition periods may apply to this product. Pharmacokinetic data required.
Levonorgestrel					
Subdermal implant	14,402	\$1,539,862	\$106.9200	CH 1%	Longer transition periods may apply to this product. Training should be provided to clinicians if awarded.
Crm 4%	420	\$425	\$1.0125	CH 1%	 Current restrictions may apply. Unit subsidy and cost shown is weighted.
Lidocaine [lignocaine] hydrochloride with	adrenaline a	nd tetracaine I	nydrochloride		
Soln 4% with adrenaline 0.1 % and tetracaine hydrochloride 0.5%, 5 ml syringe	1			H 1%	
Lidocaine [lignocaine] hydrochloride with Nasal spray 5% with phenylephrine hydrochloride 0.5%	phenylephrii	ne hydrochlori	de	H 1%	5
Lithium carbonate					
Cap (immediate-release) 250 mg	1,875,606	\$176,682	\$0.0942	CH 1%	
Tab (immediate-release) 250 mg	1,626,076	\$111,549	\$0.0686	CH 1%	
Tab long-acting 400 mg	1,866,078			CH 1%	
Lopinavir with ritonavir Tab 100 mg with ritonavir 25 mg			\$3.0625	CH 1%	 Special Authority restrictions may apply.
Tab 200 mg with ritonavir 50 mg	49,192	\$189,799	\$3.8583	CH 1%	Special Authority restrictions may apply.
Losartan					
Tab 12.5 mg	2,985,540	\$49,411	\$0.0166	CH 1%	
Tab 25 mg	5,753,456	\$111,617	\$0.0194	CH 1%	
Tab 50 mg	9,430,953	\$224,551	\$0.0238	CH 1%	

SC	HEDULE	TWO: PRO	DUCTS TO	BE TENDE	ERE	Ð
Chemical Name			Unit		DV	
Line Item	Units	Cost	Subsidy	L	.imi	t Comments
Losartan						
<u>Tab 100 mg</u>	4,995,536	\$137,377	\$0.0275	СН	1%	
Macrogol 3350 with potassium chloride, s	odium bicarb	onate and sod	lium chloride			
Powder for oral soln 13.125 g with potassium chloride 46.6 mg.sodium bicarbonate 178.5 mg and sodium chloride 350.7 mg	6,126,064	\$1,384,490	\$0.2260	СН	1%	May be subject to Special Authority restrictions.
Magnesium Cap/Tab (150 mg elemental)				Н	1%	
Magnesium hydroxide						
Paste/oral liq	480,586	\$21,732	\$0.0452	СН	1%	Note units and unit subsidy expressed are for the paste presentation.
Magnesium sulphate						
lnj 0.4 mmol per ml, 250 ml bag					1%	
<u>Inj 2 mmol per ml, 5ml</u>	21,314	\$21,762	\$1.0210	СН	1%	
Malathion						
Lotn 0.5%					1%	
Shampoo 1%				Н	1%	
Maprotiline Hydrochloride						
Tab 25 mg	48,225	\$12,085	\$0.2506		1%	
Tab 75 mg	2,251	\$1,576	\$0.7003	СН	1%	
Mebendazole						
Tab 100 mg	62,189	\$62,682	\$1.0079	СН	1%	
Mebeverine hydrochloride Tab 135 mg	2,510,134	\$502,027	\$0.2000	СН	1%	
Medroxyprogesterone Acetate						
lnj 150 mg per ml, 1 ml	155,810	\$1,129,623	\$7.2500	СН	1%	
Tab 2.5 mg	664,389	\$83,049	\$0.1250	СН	1%	
Tab 5 mg	806,272	\$112,878	\$0.1400	СН	1%	
Tab 10 mg	1,712,373	\$408,110	\$0.2383	СН	1%	
Tab 100 mg	23,739	\$23,976	\$1.0100	СН	1%	Not currently listed in Section H of the Pharmaceutical Schedule.
Melatonin						
Tab 3 mg				Н	1%	Current restrictions may apply.
Tab modified-release 2 mg	2,579,653	\$2,426,602	\$0.9407#	СН	1%	Current restrictions may apply.
Menthol						
Crystals	35,997	\$10,655	\$0.2960	СН	1%	Current restrictions may apply.
Mepivacaine hydrochloride						
Inj 3%, 1.8ml dental cartridge				Н	1%	
Inj 3%, 2.2ml dental cartridge				Н	1%	
Meropenem						
<u>lnj 500 mg</u>				Н	1%	Current restrictions may apply.
<u>lnj 1 g</u>				Н	1%	Current restrictions may apply.
Mesalazine						
Tab long-acting 500 mg	9,331,739	\$5,510,392	\$0.5905	СН	1%	
Metaraminol tartrate						
lnj 0.5 mg per ml, 10 ml				Н	1%	Not currently listed in the Pharmaceutical Schedule.
Inj 0.5 mg per ml, 20 ml prefilled syringe				Н	1%	
lnj 0.5 mg per ml, 6 ml				Н	1%	Not currently listed in the Pharmaceutical Schedule.
lnj 1 mg per ml, 1 ml				Н	1%	
Inj 1 mg per ml, 10 ml prefilled syringe				Н	1%	
lnj 10 mg per ml, 1 ml				Н	1%	

	SCHEDULE 1	TWO: PRO	DUCTS TO	D BE TENDE	RE	D
Chemical Name			Unit	D	v	
Line Item	Units	Cost	Subsidy	_	mit	Comments
Methotrexate						
<u>Inj 100 mg per ml, 50 ml</u>		\$68,711	\$79.9900	PCT C H 1		Preference for products where stability data >48 hours post-compounding.
Methylprednisolone aceponate						
Crm 0.1%	664,920	\$219,424	\$0.3300	CH 1	%	
Oint 0.1%	380,565	\$125,586	\$0.3300	CH 1	%	
Metoclopramide hydrochloride						
Oral liq 5 mg per 5 ml				CH 1	%	Not currently listed in Section B of the Pharmaceutical Schedule.
<u>Tab 10 mg</u>	5,656,088	\$73,529	\$0.0130	CH 1	%	
Metronidazole						
Gel 0.75%				CH 1		Not currently listed in Section B of the Pharmaceutical Schedule. Special Authority restrictions for use on odiferous wounds only may apply.
lnj 5 mg per ml, 100 ml				H 1		Preference for a bag presentation with space to add another antibiotic.
Oral liq 200 mg per 5 ml	240,445	\$60,111	\$0.2500	CH 1	%	
Suppos 500 mg	6,291	\$15,401	\$2.4480	CH 1	%	
Tab 200 mg	406,330	\$42,462	\$0.1045	CH 1	%	
Tab 400 mg	1,978,110	\$359,027	\$0.1815	CH 1	%	
Metyrapone						
Cap 250 mg	27,790	\$289,016	\$10.4000	С		
Miconazole nitrate						
<u>Crm 2% (15 g - 20 g pack)</u>	2,767,860	\$136,539	\$0.0493	CH 1	%	
Lotn/tincture 2%	25,080	\$3,645	\$0.1453	CH 1	%	
Vaginal crm 2% with applicator	436,040	\$42,296	\$0.0970	CH 1	%	Single use applicators preferred.
Mifepristone						
Tab 200 mg				H 1	%	
Misoprostol						
Tab 200 mcg	51,124	\$17,680	\$0.3458	CH 1	%	
Mitomycin C						
lnj 2 mg - 5 mg				PCT H 1	%	Units and subsidy shown are per inj 5 mg presentation.
Inj 10 mg - 20 mg				PCT H 1	%	Not currently listed on the Pharmaceutical Schedule.
Morphine						
<u>lnj 1 mg per ml, 10 ml syringe</u>				H 1		Morphine sulphate inj 1 mg per ml, 10 ml syringe is currently listed in Section H of the Pharmaceutical Schedule.
<u>lnj 1 mg per ml, 100 ml bag</u>				H 1		Morphine sulphate inj 1 mg per ml, 100 ml bag is currently listed in Section H of the Pharmaceutical Schedule.
lnj 1 mg per ml, 2 ml syringe				H 1		Morphine sulphate inj 1 mg per ml, 2 ml syringe is currently listed in Section H of the Pharmaceutical Schedule.
<u>lnj 1 mg per ml. 50 ml syringe</u>				H 1	%	Morphine sulphate inj 1 mg per ml, 50 ml syringe is currently listed in Section H of the Pharmaceutical Schedule.
<u>lnj 10 mg per ml, 1 ml</u>	213,500	\$190,869	\$0.8940	СН 1	%	Units and subsidy shown are for morphine sulphate inj 10 mg per ml, 1 ml.
Inj 10 mg per ml, 100 mg cassette				H 1	%	Morphine sulphate inj 10 mg per ml, 100 ml cassette is currently listed in Section H of the Pharmaceutical Schedule.
lnj 10 mg per ml, 100 ml bag				H 1		Morphine sulphate inj 10 mg per ml, 100 ml bag is currently listed in Section H of the Pharmaceutical Schedule.
<u>lnj 15 mg per ml, 1 ml</u>	5,080	\$4,836	\$0.9520	СН 1		Units and subsidy shown are for morphine sulphate inj 15 mg per ml, 1 ml.

S	CHEDULE	TWO: PRO	DDUCTS TO	BE TEND	ERE	D
Chemical Name Line Item	Units	Cost	Unit Subsidy		DV Limit	t Comments
Morphine						
Inj 2 mg per ml, 30 ml syringe				Н	1%	Morphine sulphate inj 2 mg per ml, 30 ml syringe is currently listed in Section H of the Pharmaceutical Schedule.
Inj 200 mcg in 0.4 ml syringe				Н	1%	Morphine sulphate inj 200 mcg in 0.4 ml syringe is currently listed in Section H of the Pharmaceutical Schedule.
<u>lnj 30 mg per ml. 1 ml ampoule</u>	54,816	\$67,862	\$1.2380	СН	1%	Units and subsidy shown are for morphine sulphate inj 30 mg per ml, 1 ml ampoule.
Inj 300 mcg in 0.3 ml syringe				Н	1%	Morphine sulphate inj 300 mcg in 0.3 ml syringe is currently listed in Section H of the Pharmaceutical Schedule.
lnj 80 mg per ml	15,813	\$135,106	\$8.5440	СН	1%	Units and subsidy shown are for morphine tartrate inj 80 mg per ml, 1.5 ml ampoule.
Tab immediate-release 10 mg	3,300,436	\$924,122	\$0.2800	СН	1%	May be a preference for blister packs. Units and subsidy shown are for morphine sulphate tab immediate-release 10 mg.
Tab immediate-release 20 mg	715,695	\$395,064	\$0.5520	СН	1%	May be a preference for blister packs. Units and subsidy shown are for morphine sulphate tab immediate-release 20 mg.
Mucilaginous Laxatives						
Dry	84,651,000	\$1,024,277	\$0.0121	СН	1%	Not currently listed in Section H of the Pharmaceutical Schedule.
Mupirocin						
Intra-nasal oint 2%				СН	1%	Not currently listed on the Pharmaceutical Schedule. Restrictions may apply.
Topical oint 2% (pack size 5 g or less)	311,055	\$136,864	\$0.4400	СН	1%	Current restrictions may apply.
Naltrexone hydrochloride Tab 50 mg	201,243	\$754,995	\$3.7517	СН	1%	Current restrictions may apply.
Neostigmine metisulfate						
Inj 2.5 mg per ml, 1 ml	11,139	\$21,832	\$1.9600	СН	1%	
Nicardipine hydrochloride (current acces	ss)					
lnj 2.5 mg per ml, 10 ml vial				Н	1%	Current restrictions may apply. PHARMAC would only award a tender for either current or widened access.
Nicardipine hydrochloride (widened acc	ess)					
lnj 2.5 mg per ml, 10 ml vial				Н	1%	PHARMAC would only award a tender for either current or widened access. Widened access would result in the removal of the restriction to paediatric patients only.
Nicotinic acid						
<u>Tab 50 mg</u>	581,959	\$23,977	\$0.0412	СН	1%	
<u>Tab 500 mg</u>	496,432	\$88,812	\$0.1789	СН	1%	
Nifedipine						
Tab long-acting 10 mg	803,651	\$142,383	\$0.1772	СН	1%	
Tab long-acting 20 mg	603,119	\$57,839 \$62,082	\$0.0959 \$0.1017	СН	1%	
Tab long-acting 30 mg Tab long-acting 60 mg	601,723 173,501	\$62,982 \$32,792	\$0.1047 \$0.1890	СН СН	1% 1%	
	175,501	ψ32,792	φ0.1090	OII	1 70	
Nitazoxanide Tab 500 mg				СН	1%	Current restrictions may apply. Not
				011	170	currently listed in Section B of the Pharmaceutical Schedule.
Oral 100 mg per 5 ml				Н	1%	Current restrictions may apply.
Nitrofurantoin						
Tab modified-release				СН	1%	Not currently listed in the Pharmaceutical Schedule.
Noradrenaline						
Inj 0.06 mg per ml, 100 ml bag				Н	1%	
Inj 0.1 mg per ml, 100 ml bag				Н	1%	
sole supply						=rebate *=part charge @=ASP +=paten

SC	HEDULE	TWO: PRO	DUCTS	TO BE TENDERED
Chemical Name			Unit	DV
Line Item	Units	Cost	Subsidy	Limit Comments
Noradrenaline				
Inj 0.1 mg per ml, 50 ml syringe				H 1% Not currently listed on the Pharmaceutical Schedule.
Inj 0.12 mg per ml, 50 ml syringe				H 1%
lnj 1 mg per ml, 100 ml bag				Н 1%
Norfloxacin				
Tab 400 mg	116,182	\$156,846	\$1.3500	СН 1%
Nystatin				
<u>Oral liq 100,000 u per ml</u>	1,792,800	\$145,665	\$0.0813	CH 1%
Vaginal crm 100,000 u per 5 g with	137,100	\$8,134	\$0.0593	C H 1% Units and unit subsidy expressed as 'per
<u>applicator(s)</u>				g'.
Octreotide (somatostatin analogue)	5 4 5 4	\$04 504	\$0,4000	
Inj 50 mcg per ml, 1 ml	5,154	\$31,584 \$62,422	\$6.1280 \$2.7280	СН 1% СН 1%
Inj 100 mcg per ml, 1 ml	16,890 4,151	\$63,133 \$60,184	\$3.7380 \$14.5000	СН 1%
Inj 500 mcg per ml, 1 ml	4,101	φυυ, 104	φ14.0000	
Oestradiol Tab 1 mg	356,944	\$52,521	\$0.1471	* CH 1%
Tab 2 mg	241,640	\$35,555	\$0.1471 \$0.1471	* CH 1%
TDDS 25 mcg per day	276,743	\$211,708	\$0.7650	CH 1%
TDDS 50 mcg per day	350,086	\$308,076	\$0.8800	СН 1%
TDDS 75 mcg per day	61,877	\$61,181	\$0.9888	C H 1%
TDDS 100 mcg per day	115,917	\$114,613	\$0.9888	C H 1%
Oestradiol with norethisterone				
Tab 1 mg with 0.5 mg norethisterone acetate	943,404	\$181,945	\$0.1929	C H 1%
Tab 2 mg with 1 mg norethisterone acetate	373,408	\$72,015	\$0.1929	CH 1%
Tab 2 mg with 1 mg norethisterone acetate (10), and 2 mg (oestradiol tab (12) and 1 mg oestradiol tab (6)	199,108	\$38,400	\$0.1929	C H 1%
Oestriol				
Crm 1 mg per g with applicator(s)	2,988,540	\$1,318,932	\$0.4413	C H 1% Applicator(s) should be mulitple use and a measured dose.
Pessaries 500 mcg	375,994	\$171,953	\$0.4573	C H 1%
<u>Tab 1 mg - 2 mg</u>	200,214	\$46,716	\$0.2333	C H 1% Units and unit subsidy are based per 2 mg tablet. Preference for 2 mg tablets to be scored.
Oestrogens				
Conjugated, equine tab 300 mcg	299,986	\$32,248	\$0.1075	C H 1%
Conjugated, equine tab 625 mcg	360,416	\$53,032	\$0.1471	C H 1%
Olanzapine		.	A	
Orodispersible tab 5 mg	324,772	\$14,498 \$26,442	\$0.0446	CH 1%
Orodispersible tab 10 mg	497,365	\$36,412 \$20,711	\$0.0732 \$0.0220	CH 1%
<u>Tab 2.5 mg</u> <u>Tab 5 mg</u>	1,737,142 2,293,670	\$39,711 \$94,201	\$0.0229 \$0.0411	СН 1% СН 1%
<u>Tab 5 mg</u> Tab 10 mg	2,293,670	\$94,201 \$164,811	\$0.0411 \$0.0589	Сн 1%
Omeprazole	_,. 50,. 0 L	÷	÷0.0000	
Cap 10 mg	6,454,257	\$141,994	\$0.0220	СН 1%
	93,790,879	\$2,042,765	\$0.0218	C H 1%
	27,452,284	\$951,771	\$0.0347	C H 1%
Ondansetron				
Tab disp 4 mg	2,696,769	\$256,193	\$0.0950	CH 1%
<u>Tab disp 8 mg</u>	1,209,514	\$172,961	\$0.1430	C H 1%
Ortho-Tolidine				
Compound diagnostic sticks			\$0.1500	С
Oxazepam				
<u>Tab 10 mg</u>	698,205	\$43,079	\$0.0617	СН 1%
sole supply				#=rebate *=part charge @=ASP +=patent

:	SCHEDULE	TWO: PRO	ODUCTS T	O BE TE	ND	ERE	D
Chemical Name Line Item	Units	Cost	Unit Subsidy			DV Limit	Comments
Oxazepam							
<u>Tab 15 mg</u>	216,524	\$18,469	\$0.0853	C	СН	1%	
Oxybutynin Oral liq 1 mg per ml	425,620	\$54,352	\$0.1277	C	СН	1%	Preference for pack size 500 ml or less
Tab 5 mg	7,771,036	\$137,547	\$0.0177	C	СН	1%	
Paclitaxel							
<u>Inj 30 mg</u>				PCT	Η	1%	Preference for products where stability data >48 hours post-compounding.
<u>lnj 100 mg</u>				PCT	н	1%	Preference for products where stability data >48 hours post-compounding.
Inj 150 mg				PCT	H	1%	
<u>Inj 300 mg</u>				PCT	Н	1%	Preference for products where stability data >48 hours post-compounding.
Paliperidone (one-monthly depot inject	ion)						
Inj 25 mg syringe	924	\$179,487	\$194.2500#	+ C	СН	1%	Special Authority restrictions may apply. Training requirements may apply.
Inj 50 mg syringe	2,652	\$721,211	\$271.9500#	+ C	СН	1%	Special Authority restrictions may apply. Training requirements may apply.
Inj 75 mg syringe	5,088	\$1,818,553	\$357.4200#		СН	1%	Special Authority restrictions may apply. Training requirements may apply.
Inj 100 mg syringe	12,067	\$5,250,593	\$435.1200#		С Н	1%	Special Authority restrictions may apply. Training requirements may apply.
Inj 150 mg syringe	13,558	\$5,899,357	\$435.1200#	+ C	СН	1%	Special Authority restrictions may apply. Training requirements may apply.
Paliperidone (three-monthly depot injection in the second se	ction)						
Inj 175 mg syringe				C	СН	1%	Not currently listed on the Pharmaceutical Schedule. Restrictions may apply. Training requirements may apply.
Inj 250 - 300 mg syringe				C	СН	1%	Not currently listed on the Pharmaceutical Schedule. Restrictions may apply. Training requirements may apply.
Inj 350 mg syringe				C	СН	1%	Not currently listed on the Pharmaceutica Schedule. Restrictions may apply. Training requirements may apply.
Inj 500 mg or greater syringe				C	СН	1%	Not currently listed on the Pharmaceutica Schedule. Restrictions may apply. Training requirements may apply.
Palonosetron							
Inj 250 mcg					Н	1%	Not currently listed in the Pharmaceutical Schedule.
Pamidronate disodium							
Inj 3 mg per ml, 10 ml	334	\$1,997	\$5.9800		СН	1%	
<u>lnj 6 mg per ml, 10 ml</u>	29	\$436	\$15.0200		СН	1%	
<u>lnj 9 mg per ml, 10 ml</u>	83	\$1,415	\$17.0500	C	СН	1%	
Paper wasp venom							
Inj with diluent for extract	17	\$5,185	\$305.0000		СН	1%	Current restrictions may apply.
Treatment kit	8	\$2,440	\$305.0000	С	СН	1%	Special Authority restrictions may apply.
Paracetamol							
<u>Tab 500 mg, blister pack</u>	321,927,226					10%	
Tab 500 mg, bottle pack	38,824,793	\$245,373	\$0.0063	C		10%	
Inj 10 mg per ml, 100 ml		A - · -	A		H	1%	
Oral liq 120 mg per 5 ml	59,671,499	\$319,243	\$0.0054			20%	Preference may be given to products which include a measuring device.
<u>Oral liq 250 mg per 5 ml</u>	257,689,578	\$1,497,176	\$0.0058	C		20%	Preference may be given to products which include a measuring device.
Paracetamol with codeine							
Tab paracetamol 500 mg with codeine phosphate 8 mg	45,440,910	\$827,479	\$0.0182	C	СН	1%	

S	CHEDULE	TWO: PRO	DDUCTS TO	D BE TENDERED
Chemical Name Line Item	Units	Cost	Unit Subsidy	DV Limit Comments
Paraffin liquid with soft white paraffin			-	
Eye oint with soft white paraffin	80,378			C H 1% Not currently listed in Section B of the Pharmaceutical Schedule.
Paraffin liquid with wool fat liquid Eye oint 3% with wool fat liq 3%	231,909	\$240,522	\$1.0371	СН 1%
Paraldehyde	201,000	<i>\</i>	QQ	
Inj 5 ml	5	\$1,500	\$300.0000	СН 1%
Paromomycin	-	+ - 1		
Cap 250 mg	1,672	\$13,167	\$7.8750	C H 1% Special Authority restrictions may appl
Pericyazine				
Tab 10 mg	158,174	\$70,311	\$0.4445	СН 1%
Tab 2.5 mg	267,722	\$33,435	\$0.1249	CH 1%
Perindopril	- 1	+,		
Tab 2 mg - 2.5 mg	838,001	\$104,750	\$0.1250	C H Units and subsidy shown are tab 2 mg
		¢.01,100	ţ <u>_</u>	presentation. Bids for erbumine and/or arginine salts will be considered. Preference for the same salt across all strengths.
Tab 4 mg - 5 mg	1,692,676	\$270,828	\$0.1600	C H Units and subsidy shown are per tab 4 presentation. Bids for erbumine and/or arginine salts will be considered. Preference for the same salt across all strengths.
Tab 8 mg - 10 mg				C H Not currently listed in the Pharmaceuti Schedule. Bids for erbumine and/or arginine salts will be considered. Preference for the same salt across all strengths.
Permethrin				,
<u>Crm 5%</u>	814,350	\$134,368	\$0.1650	CH 1%
Lotion 5%	1,713,960	\$210,817	\$0.1230	C H 1%
Pethidine hydrochloride				
lnj 5 mg per ml, 10 ml				H 1% Preservative free desirable
Inj 10 mg per ml, 100 ml bag				H 1%
lnj 10 mg per ml, 50 ml syringe				H 1%
Inj 5 mg per ml, 100 ml bag				H 1%
<u>lnj 50 mg per ml, 1 ml</u>	4,461	\$4,443	\$0.9960	C H 1%
<u>lnj 50 mg per ml, 2 ml</u>	15,893	\$16,274	\$1.0240	C H 1%
Phenelzine sulphate				
Tab 15 mg	97,065	\$114,536	\$1.1800	C H 1%
Phenothrin				
Shampoo 0.5%	4,537,800	\$257,747	\$0.0568#	C H 1%
Phenoxybenzamine				
Cap 10 mg	20,424	\$44,252	\$2.1667	C H 1%
Phenoxybenzamine hydrochloride				
lnj 50 mg per ml, 2 ml ampoule				H 1%
Phentolamine Mesylate				
Inj 10 mg per ml, 1 ml				H 1%
lnj 5 mg per ml, 1 ml				H 1%
Phenylephrine hydrochloride				
lnj 10 mg per ml, 1 ml				H 1%
Phenytoin Sodium		¢40.55	A-7 7000	0.11 494
lnj 50 mg per ml, 2 ml	591	\$10,476 \$14,517	\$17.7260	CH 1%
lnj 50 mg per ml, 5 ml	542	\$14,517	\$26.7840	C H 1%

SC	HEDULE	TWO: PRO	DUCTS TO	BE TENDER	ED
Chemical Name Line Item	Units	Cost	Unit Subsidy	DV Limi	it Comments
Pimecrolimus			-		
Crm 1%				СН 1%	PHARMAC reserves the right to award tenders for one, some or all presentations of topical tacrolimus or pimecrolimus. This item is not currently listed on the Pharmaceutical Schedule. Special Authority restrictions may apply.
Pine tar with triethanolamine lauryl sulpha			* • • • - -	0.11 444	
Soln 2.3% with triethanolamine lauryl sulphate and fluorescein sodium	30,521,428	\$235,625	\$0.0077	CH 1%	
Piperacillin with tazobactam Inj 4 g with tazobactam 500 mg				H 1%	Current restrictions may apply. PHARMAC will consider other 8:1 (piperacillin:tazobactam) presentations if cost effective.
Pirfenidone					
Cap 267 mg	260,770	\$3,520,395	\$13.5000#	CH 1%	
Pivmecillinam					
Tab 200 mg				CH 1%	Not currently listed in Section B of the Pharmaceutical Schedule. Any listing may be subject to Special Authority restrictions.
Podophyllotoxin					
Soln 0.5%	10,735	\$103,051	\$9.6000	CH 1%	
Poloxamer Oral drops 10%	267,450	\$33,699	\$0.1260	CH 1%	
Polyvinyl alcohol					
Eye drops 1.4%	498,855	\$87,135	\$0.1747	СН 1%	For products containing BAK, PHARMAC reserves its right to list a BAK or preservative free product for a restricted market. Units and unit subsidy shown are per ml
Eye drops 3%	261,525	\$64,160	\$0.2453	CH 1%	For products containing BAK, PHARMAC reserves its right to list a BAK or preservative free product for a restricted market. Units and unit subsidy shown are per ml
Poractant Alfa					
Soln 120 mg per 1.5 ml vial				H 1%	
Soln 240 mg per 3 ml vial				H 1%	
Potassium citrate Tab				CH 1%	Schedule. Any listing may be subject to
					Special Authority restrictions.
Potassium permanganate Crystals/Tab				H 1%	Note this product may require Medsafe registration prior to an award.
Povidone lodine					
Oint 10%	601,920	\$78,731	\$0.1308	CH 1%	
Skin preparation, povidone iodine 10% with 30% alcohol (pack size 100 ml or less	3,300)	\$54	\$0.0163 *	CH 1%	Not currently listed in Section H of the Pharmaceutical Schedule.
Skin preparation, povidone iodine 10% with 30% alcohol (pack size greater than 100 ml)	75,275	\$1,506	\$0.0200	CH 1%	
Skin preparation, povidone iodine 10% with 70% alcohol	220,400	\$3,593	\$0.0163#	CH 1%	
Pravastatin					
Tab 10 mg				CH 1%	Not currently listed in Section B of the Pharmaceutical Schedule
<u>Tab 20 mg</u>	3,394,331	\$160,212	\$0.0472	CH 1%	
<u>Tab 40 mg</u>	1,706,545	\$137,547	\$0.0806	CH 1%	

S	CHEDULE	TWO: PRO	DUCTS T	O BE TEND	DERE	ED
Chemical Name			Unit		DV	
Line Item	Units	Cost	Subsidy		Limi	t Comments
Praziquantel						
Tab 600 mg	822	\$6,983	\$8.5000	СН	1%	
Prazosin Hydrochloride						
Tab 1 mg	536,269	\$29,656	\$0.0553	СН	1%	
Tab 2 mg	299,476	\$20,963	\$0.0700	СН	1%	
Tab 5 mg	104,459	\$12,222	\$0.1170	СН	1%	
Prednisone						
<u>Tab 1 mg</u>	11,419,671	\$243,924	\$0.0214	СН	1%	
<u>Tab 2.5 mg</u>	2,178,215	\$52,669	\$0.0242	СН	1%	
<u>Tab 20 mg</u>	6,366,606	\$369,645	\$0.0581	СН	1%	
<u>Tab 5 mg</u>	12,681,001	\$281,265	\$0.0222	СН	1%	
Pregnancy tests - HCG urine						
Pregnancy test - HCG urine - Dipstick				С		Not currently listed on the Pharmaceutical Schedule. PHARMAC reserves the right to award one, some or all of the tenders for pregnancy tests - HCG urine.
Pregnancy tests - HCG urine - Cassette	428,240	\$128,472	\$0.3000	С		PHARMAC reserves the right to award one, some or all of the tenders for pregnancy tests - HCG urine.
Pregnancy tests - HCG urine - Midstream				С		Not currently listed on the Pharmaceutical Schedule. PHARMAC reserves the right to award one, some or all of the tenders for pregnancy tests - HCG urine.
Primaquine phosphate						
Tab 7.5 - 15 mg	484	\$1,011	\$2.0893	СН	1%	Tab 7.5 mg is currently listed in Section B of the Pharmaceutical Schedule. Units shown are for tab 7.5 mg. Special Authority restrictions may apply. There may be a preference for a scored tablet.
Procaine penicillin						
<u>Inj 1.5 mega u</u>	1,141	\$28,183	\$24.7000	СН	1%	
Prochlorperazine						
<u>Tab 5 mg</u>	3,533,778	\$89,758	\$0.0254	СН	1%	
Progesterone						
Cap 100 mg	40,563	\$22,310	\$0.5500	СН	1%	Must be compatible for intra-vaginal use. Current restrictions may apply. PHARMAC reserves the right to amend access.
Propafenone hydrochloride		A				
Tab 150 mg	70,212	\$57,433	\$0.8180	СН	1%	
Propylthiouracil			•			
Tab 50 mg	243,313	\$85,160	\$0.3500	СН	1%	Current restrictions may apply.
Protionamide						
Tab 250 mg	4,194	\$12,792	\$3.0500	СН	1%	Current restrictions may apply.
Pseudoephedrine Hydrochloride Tab 60 mg				Н	1%	
Pyrazinamide						
Tab 500 mg	110,215	\$65,027	\$0.5900	СН	1%	Current restrictions may apply. There may be a preference for a scored tablet or a tablet that is dispersable in water.
Pyridoxine hydrochloride						
<u>Tab 25 mg</u> <u>Tab 50 mg</u>	800,887 1,713,683	\$24,027 \$46,715	\$0.0300 \$0.0273	СН СН	1% 1%	
Pyrimethamine						
Tab 25 mg	1,594	\$1,178	\$0.7390	СН	1%	Current restrictions may apply. There may be preference for a scored tablet or tablet that is dispersable in water.
Quetiapine						
<u>Tab 100 mg</u>	3,747,041	\$143,624	\$0.0383	СН	1%	
sole supply						-robato *-part charge @-ASP (-patent

	SCHEDULE TWO: PRODUCTS TO BE TENDERED						
Chemical Name			Unit	DV			
Line Item	Units	Cost	Subsidy	Limit Comments			
Quetiapine							
<u>Tab 200 mg</u>	1,196,481	\$76,443	\$0.0639	C H 1%			
<u>Tab 25 mg</u>	21,780,013	\$433,204	\$0.0199	C H 1% There may be a preference for a stablet.	scored		
<u>Tab 300 mg</u>	468,008	\$49,922	\$0.1067	C H 1%			
Quinine dihydrochloride Inj 600 mg				H 1% Current restrictions may apply.			
Ranitidine hydrochloride							
Inj	23,156	\$40,523	\$1.7500	C H 1% Units and subsidy shown are for i per ml, 2 ml presentation.	nj 25 mg		
<u>Oral liq 150 mg per 10 ml</u>	1,929,098	\$33,045	\$0.0171	C H 1%			
<u>Tab 150 mg</u>	9,440,456	\$243,753	\$0.0258	C H 1%			
<u>Tab 300 mg</u>	2,761,040	\$100,557	\$0.0364	C H 1%			
Rasagiline							
Tab 1 mg				C H 1% Not currently listed on the Pharma Schedule. Pricing should be simili selegine.			
Remifentanil hydrochloride							
<u>lnj 1 mg</u>				H 1%			
<u>lnj 2 mg</u>				H 1%			
Rifabutin							
Cap 150 mg	7,187	\$65,881	\$9.1667	C H 1%			
Rifampicin							
<u>Cap 150 mg</u>	87,288	\$48,663	\$0.5575	C H 1% Current restrictions may apply.			
<u>Cap 300 mg</u>	161,524	\$187,772	\$1.1625	C H 1% Current restrictions may apply.			
<u>lnj 600 mg</u>				H 1% Current restrictions may apply.			
<u>Oral liq 100 mg per 5 ml</u>	65,874	\$13,175	\$0.2000	C H 1% Current restrictions may apply.			
Rifaximin							
<u>Tab 200 mg - 550 mg</u>	109,607	\$1,223,293	\$11.1607	C H 1% Current restrictions may apply. Ta mg is currently listed on the Pharmaceutical Schedule. Units s are for tab 550 mg.			
Risperidone							
Inj 25 mg vial	4,162	\$565,949	\$135.9800	C H 1% Current restrictions may apply.			
Inj 37.5 mg vial	5,578	\$996,844	\$178.7100	C H 1% Current restrictions may apply.			
Inj 50 mg vial	7,028	\$1,529,012	\$217.5600	C H 1% Current restrictions may apply.			
Oral liquid 1 mg per ml	359,850	\$91,881	\$0.2553	C H 1%			
<u>Tab 0.5 mg</u>	2,978,434	\$92,331	\$0.0310	C H 1%			
<u>Tab 1 mg</u>	2,129,800	\$73,116	\$0.0343	C H 1%			
<u>Tab 2 mg</u>	1,024,326	\$39,099	\$0.0382	C H 1%			
<u>Tab 3 mg</u>	416,418	\$17,352	\$0.0417	C H 1%			
<u>Tab 4 mg</u>	200,667	\$11,472	\$0.0572	C H 1%			
Rizatriptan Tab orodispersible 10 mg	1,727,121	\$302,816	\$0.1753	СН 1%			
Ropivacaine hydrochloride							
lnj 2 mg per ml, 10 ml				H 1%			
<u>lnj 2 mg per ml, 20 ml</u>				H 1%			
<u>lnj 2 mg per ml, 100 ml</u>				H 1%			
<u>lnj 2 mg per ml, 200 ml</u>				H 1%			
<u>lnj 7.5 mg per ml, 10 ml</u>				H 1%			
<u>lnj 7.5 mg per ml, 20 ml</u>				H 1%			
<u>lnj 10 mg per ml, 10 ml</u>				H 1%			
<u>lnj 10 mg per ml, 20 ml</u>				H 1%			

SCHEDULE TWO: PRODUCTS TO BE TENDERED **Chemical Name** Unit ΠV Line Item Units Cost Subsidy Limit Comments Ropivacaine hydrochloride with fentanyl Inj 2 mg with fentanyl 2 mcg per ml, 100 Sterile bag preferred. н 1% ml bag Inj 2 mg with fentanyl 2 mcg per ml, 200 н 1% Sterile bag preferred. ml bag Rosuvastatin Tab 5 mg СН 1% Not currently listed on Pharmaceutical Schedule. Any listing may be subject to a Special Authority restriction. СН Not currently listed on Pharmaceutical Tab 10 mg 1% Schedule. Any listing may be subject to a Special Authority restriction. Tab 20 mg СН 1% Not currently listed on Pharmaceutical Schedule. Any listing may be subject to a Special Authority restriction. Not currently listed on Pharmaceutical Tab 40 mg СН 1% Schedule. Any listing may be subject to a Special Authority restriction. Salicylic Acid 271,534 \$20,506 \$0.0755 СН Powder 1% Note this product may require Medsafe registration prior to an award. Selexipag Not currently listed in the Pharmaceutical Tab 200 mcg СН 1% Schedule. Tab 400 mcg СН 1% Not currently listed in the Pharmaceutical Schedule. Tab 600 mcg Tab 800 mcg СН 1% Not currently listed in the Pharmaceutical Schedule. Not currently listed in the Pharmaceutical Tab 1000 mcg СН 1% Schedule. Not currently listed in the Pharmaceutical Tab 1200 mcg СН 1% Schedule. Tab 1400 mcg СН 1% Not currently listed in the Pharmaceutical Schedule. Tab 1600 mcg СН 1% Not currently listed in the Pharmaceutical Schedule. Silver Sulphadiazine 506,200 \$109,339 \$0.2160 СН Crm 1% 50 g is currently listed on the Crm 1% (pack size of 100 g or less) 1% Pharmaceutical Schedule. Units shown are for crm 1% 50 g. Simvastatin Tab 10 mg 3,905,690 \$41,244 \$0.0106 СН 1% Tab 20 mg 17,783,874 \$300,370 \$0.0169 СН 1% 15,672,177 \$457,941 \$0.0292 1% <u>Tab 40 mg</u> СН Tab 80 mg 1.054.633 \$70.312 \$0.0667 СН 1% Sodium Acid Phosphate Enema 16% with sodium phosphate 8% 52,926 \$132,314 \$2.5000 СН Not currently listed in Section B of the 1% Pharmaceutical Schedule. Sodium alginate with magnesium alginate Powder for oral soln 974,019 СН Note this product is currently described as 1% 'Alginic acid - sodium alginate 225 mg and magnesium alginate 87.5 mg per sachet' in Section B of the Pharmaceutical Schedule. Sodium alginate with sodium bicarbonate and calcium carbonate Oral liq 18.270.721 \$54,812 \$0.0030 СН 1% Note this product is curently described as sodium alginate oral liq 500 mg with sodium bicarbonate 267 mg and calcium carbonate 160 mg per 10 ml'.

SCHEDULE TWO: PRODUCTS TO BE TENDERED							
Chemical Name			Unit		DV		
Line Item	Units	Cost	Subsidy		Limi	t Comments	
Sodium alginate with sodium bicarbon	ate and calcium c	arbonate					
Tab 500 mg	520,054	\$15,602	\$0.0300	СН	1%	Note this product is currently described as sodium alginate tab 500 mg with sodium bicarbonate 267 mg and calcium carbonate 160 mg in Section B of the Pharmaceutical Schedule.	
Sodium Citro-Tartrate							
Grans effervescent 4 g sachets	2,233,544	\$186,657	\$0.0836	СН	1%		
Sodium cromoglicate							
Nasal spray, 4%				Н	1%		
Sodium fusidate [fusidic acid]							
Eye drops 1%	303,875	\$321,500	\$1.0580	СН	1%	Units and unit subsidy expressed as "per ml".	
Streptomycin sulphate							
lnj 400 mg per ml, 2.5 ml				Н	1%	Current restrictions may apply.	
Sulfadiazine Sodium							
Tab 500 mg	368	\$3,570	\$9.7000	СН	1%	Current restrictions may apply. Not currently listed in Section H of the Pharmaceutical Schedule.	
Suxamethonium chloride							
<u>lnj 50 mg per ml, 2 ml</u>				Н	1%	There may be a preference for a pre-filled syringe.	
Tacrolimus				0.11	4.07		
Oint 0.03%				СН	1%	PHARMAC reserves the right to award tenders for one, some or all presentations of topical tacrolimus or pimecrolimus. This item is not currently listed on the Pharmaceutical Schedule. Special Authority restrictions may apply.	
Oint 0.1%				СН	1%	PHARMAC reserves the right to award tenders for one, some or all presentations of topical tacrolimus or pimecrolimus. This item is not currently listed on the Pharmaceutical Schedule. Special Authority restrictions may apply.	
Tadalafil							
Tab/Cap 2.5 mg				СН	1%	Not currently listed on the Pharmaceutical Schedule. The same restrictions as for sildenafil would apply.	
Tab/Cap 5 mg				СН	1%		
Tab/Cap 10 mg				СН	1%		
Tab/Cap 20 mg				СН	1%		
Talc							
Powder				Н	1%		
Soln (slurry) 100 mg per ml, 50 ml				н	1%		
Tamoxifen citrate							
Tab 10 mg	37,721	\$7,387	\$0.1958	СН	1%		
<u>Tab 20 mg</u>	1,890,447	\$176,435	\$0.0933	СН	1%		
Temazepam							
Tab 10 mg	2,949,628	\$149,841	\$0.0508	СН	1%		
Terazosin hydrochloride							
Tab 1 mg	893,747	\$18,831	\$0.0211	СН	1%		
Tab 2 mg	4,311,188	\$64,668	\$0.0150	СН	1%		
Tab 5 mg	3,140,339	\$68,459	\$0.0218	СН	1%		

SCHEDULE TWO: PRODUCTS TO BE TENDERED							
Chemical Name			Unit			DV	
Line Item	Units	Cost	Subsidy		L	imi	Comments
Terbinafine							
<u>Tab 250 mg</u>	2,085,783	\$198,149	\$0.0950	C	СН	1%	
Terlipressin							
Inj 0.1 mg per ml, 8.5 ml ampoule Inj 1 mg per 8.5 ml ampoule					H H	1% 1%	
Testosterone cypionate							
Inj long-acting 100 mg per ml, 10 ml	1,643	\$125,690	\$76.5000	C	СН	1%	
Tetracaine [amethocaine] hydrochloride							
Gel 4%					Н	1%	
Tetracycline							
Cap 500 mg	27,543	\$42,233	\$1.5333	C	СН	1%	Current restrictions may apply.
Tab 250 mg					Н	1%	
Thiamine hydrochloride							
Inj 100 mg per ml, 1 ml					Н		
Tab 100 mg				C	СН	1%	Currently not listed in Section B of the Pharmaceutical Schedule.
<u>Tab 50 mg</u>	7,133,748	\$348,840	\$0.0489	C	СН	1%	
lnj 100 mg per ml, 2 ml					Н	1%	
Thiotepa							
Inj 100 mg				PCT C		1%	
lnj 15 mg				PCT C	СН	1%	
Thyrotropin alfa Inj 900 mcg vial					Н	1%	
Ticagrelor							
Tab 90 mg	3,199,974	\$5,142,806	\$1.6071	+ C	СН	1%	Current restrictions may apply.
Ticarcillin with clavulanic acid							
Inj 3 g with clavulanic acid 0.1 mg vial					Н	1%	Current restrictions may apply.
Timolol							
Tab 10 mg	53,282	\$5,621	\$0.1055	С	СН	1%	This is listed in Section H of the Pharmaceutical Schedule as timolol maleate.
Timolol maleate							
Eye drops 0.25%	149,600	\$42,786	\$0.2860	C	СН	1%	For products containing BAK, PHARMAC reserves the right to list a BAK or preservative free product for a restricted market.
Eye drops 0.5%	127,025	\$36,329	\$0.2860	C	СН	1%	For products containing BAK, PHARMAC reserves the right to list a BAK or preservative free product for a restricted market.
Tobramycin							
Powder BP					Н	1%	Current restrictions may apply.
Solution for inhalation 60 mg per ml, 5 ml	21,981	\$863,539	\$39.2857#	C	СН	1%	A confidential rebate applies. Current restrictions may apply.
lnj 100 mg per ml, 5 ml					Н	1%	Current restrictions may apply.
Tolterodone tartrate							
Tab 1 mg	172,085	\$44,742	\$0.2600	C	СН	1%	Special Authority restrictions may apply.
Tab 2 mg	186,880	\$48,589	\$0.2600	C	СН	1%	Special Authority restrictions may apply.
Tramadol hydrochloride							
Oral soln 10 mg per ml				C	СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
lnj 10 mg per ml, 100 ml bag					Н	1%	
<u>Cap 50 mg</u>	32,104,932	\$722,361	\$0.0225	C	СН	1%	
<u>lnj 50 mg per ml, 1 ml</u>					Н	1%	
<u>lnj 50 mg per ml, 2 ml</u>						40/	
<u>Tab sustained-release 100 mg</u>	4,932,239	\$382,249	\$0.0775		н Эн	1% 1%	

sole supply

#=rebate *=part charge @=ASP +=patent

SCHEDULE TWO: PRODUCTS TO BE TENDERED						
Chemical Name			Unit		DV	
Line Item	Units	Cost	Subsidy		Limi	t Comments
Tramadol hydrochloride						
Tab sustained-release 150 mg	456,866	\$47,971	\$0.1050	CH	1%	
Tab sustained-release 200 mg	357,486	\$49,154	\$0.1375	CH	1%	
Tranylcypromine Sulphate						
Tab 10 mg	200,560	\$92,017	\$0.4588	CH	1 1%	
Travoprost						
Eye drops 0.004%	49,739	\$223,633	\$4.4961	CH	I 1%	This product is currently subject to a confidential rebate. For products containing BAK, PHARMAC reserves the right to list a BAK or preservative free product for a restricted market. Units and unit subsidy shown are per ml.
Triamcinolone acetonide						
0.1% in Dental Paste USP	133,395	\$142,199	\$1.0660	CH	1%	
<u>lnj 10 mg per ml, 1 ml</u>	4,196	\$17,455	\$4.1600	CH	1 5%	PHARMAC reserves the right to list a preservative free product for a restricted market.
<u>lnj 40 mg per ml, 1 ml</u>	76,398	\$780,788	\$10.2200	CH	l 1%	PHARMAC reserves the right to list a preservative free product for a restricted market.
<u>Crm 0.02%</u>	2,382,200	\$150,079	\$0.0630	CH		
<u>Oint 0.02%</u>	1,383,800	\$87,871	\$0.0635	CH	1%	
Triamcinolone acetonide with gramicidin,	neomycin and					
Ear drops 1 mg with nystatin 100,000 u, neomycin sulphate 2.5 mg and gramicidin 250 mcg per g	582,720	\$400,911	\$0.6880	CH	l 1%	
Triazolam						
Tab 125 mcg	1,747,421	\$89,118	\$0.0510	CH		
Tab 250 mcg	1,493,668	\$61,240	\$0.0410	CH	1 1%	
Triclosan						
Soln 1%	2,117,000	\$24,981	\$0.0118	С		Preference for pack size of 500 ml or less. Subsidy by endorsement only. Note this product may require Medsafe registration prior to an award.
Trimethoprim						
Tab 100 mg				CF	I 1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Ursodeoxycholic acid						
<u>Cap 250 mg - 300 mg</u>	907,883	\$344,542	\$0.3795	CH	ł 1%	Current restrictions may apply. Cap 250 mg is currently listed on the Pharmaceutical Schedule. Units shown are for cap 250 mg.
Vancomycin hydrochloride Inj 500 mg	6,863	\$16,265	\$2.3700	CH	I 1%	Current restrictions may apply.
	0,000	ψ10,200	Ψ2.0700	01	170	
Verapamil hydrochloride Inj 2.5 mg per ml, 2 ml	2,481	\$12,405	\$5.0000	CF	I 1%	
inj 2.5 mg per mi, 2 mi Tab 40 mg	2,481	\$12,405 \$16,246	\$5.0000 \$0.0701	C F C F		
Tab 80 mg	231,750 88,072	\$10,240 \$10,340	\$0.0701 \$0.1174	C F		
Vitamins	00,012	¢10,010	4 011111	•	, o	
Cap/tab (fat soluble vitamins A, D, E, K)	218,614	\$85,259	\$0.3900	CH	1 1%	Current restrictions may apply.
Tab (BPC cap strength)	16,594,283	\$174,240	\$0.0105	CH		
Water-based lubricant						
Single use sachets, 4 ml/g or larger				С		Not currently listed in Section B of the Pharmaceutical Schedule. Restrictions may apply. Preference for a small pack size.
Wool fat with mineral oil						
Lotn hydrous 3% with mineral oil, 1000 ml	1,013,381	\$5,675	\$0.0056	* C		
Lotn hydrous 3% with mineral oil, 250 ml	1,408,000	\$7,885	\$0.0056	* CH		Not currently listed in Section H of the Pharmaceutical Schedule.
sole supply					#	=rebate *=part charge @=ASP +=patent

SCHEDULE TWO: PRODUCTS TO BE TENDERED

S	CHEDULE	rwo: Pro	DUCTS TO	O BE TENDE	ERE	D
Chemical Name Line Item	Units	Cost	Unit Subsidy		DV .imit	Comments
Xylometazoline hydrochloride						
Aqueous nasal spray 0.05%				Н	1%	
Aqueous nasal spray 0.1%				Н	1%	
Nasal drops 0.05%				Н	1%	
Nasal drops 0.1%				Н	1%	
Yellow jacket wasp venom						
Inj with diluent for extract	14	\$4,270	\$305.0000	СН	1%	
Treatment kit	1	\$305	\$305.0000	СН	1%	
Zinc						
Crm (pack size 50 g or less)				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Crm (pack size greater than 50 g)				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Oint (pack size 50 g or less)				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Oint (pack size greater than 50 g)				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Paste (pack size 50 g or less)				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Zinc and castor oil						
Crm (pack size 50 g or less)				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Crm (pack size greater than 50 g)				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Oint 5% (pack size 50 g or less)				СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Oint 5% (pack size greater than 50 g)	19,156,765	\$162,832	\$0.0085	СН	1%	Units and subsidy shown are per the 500 g pack size currently listed.
Zolmitriptan						
Nasal Spray				СН	1%	Not currently listed in the Pharmaceutical Schedule. May be subject to Special Authority restrictions.
Zopiclone						
Tab 3.75 mg (pack size 30 tabs or less)				Н	1%	Not currently listed on the Pharmaceutical Schedule.
Tab 7.5 mg (pack size 30 tabs or less)				Н	1%	Tablets should be scored.
Zuclopenthixol Acetate Inj 50 mg per ml, 1 - 2 ml				Н	1%	
Zuclopenthixol Decanoate						
Inj 200 mg per ml	28,568	\$113,129	\$3.9600	СН	1%	
Inj 500 mg per ml			·	СН	1%	Not currently listed in Section B of the Pharmaceutical Schedule.
Zuclopenthixol hydrochloride						
Tab 10 mg	198,665	\$62,480	\$0.3145	СН	1%	
	,	<i>202,100</i>	÷ 3.0 . 10	0.11	. , 5	

1. General

1.1 Sole Supply Period and Hospital Supply Status Period

- (a) Hospital Tender Bids are to be submitted on the basis that if your Hospital Tender Bid is accepted, you will have Hospital Supply Status for the particular Tender Item for the Hospital Supply Status Period.
- (b) Community Tender Bids are to be submitted on the basis that if your Community Tender Bid is accepted, you will have Sole Supply Status for the particular Tender Item for the Sole Supply Period.
- (c) Combined Community/Hospital Tender Bids are to be submitted on the basis that if your Combined Community/Hospital Tender Bid is accepted, you will have Hospital Supply Status for the particular Tender Item for the Hospital Supply Status Period and Sole Supply Status for the particular Tender Item for the Sole Supply Period.

1.2 **Transition Periods**

- (a) In relation to hospital supply:
 - (i) there will be two Transition Periods (the First Transition Period and the Final Transition Period) during which the successful tenderer's brand is to be available for supply and purchase by DHB Hospitals. Additionally, where the successful tenderer's brand of the Pharmaceutical is not listed immediately prior to the First Transition Period, the successful tenderer's brand must be available for supply and purchase by DHB Hospitals from the applicable dates specified in clause 3 of Schedule 6;
 - (ii) the First Transition Period is intended to allow for an orderly transition to the arrangements that will apply during the Hospital Supply Status Period;
 - (iii) the Final Transition Period is intended to allow for an orderly transition to any new arrangements following the end of the Hospital Supply Status Period;
 - (iv) DHB Hospitals may purchase DV Pharmaceuticals at any time within the First Transition Period and Final Transition Period without any requirement to comply with the DV Limit.
- (b) Subject to paragraph (d) below, in relation to community supply:
 - (i) there will be two Transition Periods (the First Transition Period and the Final Transition Period) during which the successful tenderer's brand is to be available for supply and subsidised, but may not be the sole subsidised brand of that Tender Item. Additionally, where the successful tenderer's brand of the Pharmaceutical is not listed immediately prior to the First Transition Period, the successful tenderer's brand must be available for supply from the applicable dates specified in clause 3 of Schedule 5;
 - (ii) the First Transition Period is intended to allow for an orderly transition to the arrangements that will apply during the Sole Supply Period;

- (iii) the Final Transition Period is intended to allow for an orderly transition to any new arrangements following the end of the Sole Supply Period.
- (c) In relation to community and/or hospital supply, PHARMAC may, in its sole discretion:
 - (i) determine a different commencement date for the First Transition Period, including where it considers that a different commencement date is necessary to ensure appropriate stock management or appropriate supply of the Tender Item; and/or
 - extend the period of the First Transition Period, by determining a different end date, and may do so before or after the commencement date of the First Transition Period. For the avoidance of doubt, in the event that PHARMAC extends the First Transition Period under this clause 1.2(c)(ii):
 - (A) the delisting of all other brands of that form and strength of the Chemical Entity is to be deferred until the actual commencement date of the Sole Supply Period, notwithstanding any date previously notified to suppliers by PHARMAC as being the intended date of delisting;
 - (B) all other brands of that form and strength of the Chemical Entity are to remain listed in accordance with the terms of any existing contract between PHARMAC and the particular pharmaceutical supplier in respect of the relevant brand(s) until such time as that supplier's brand of that form and strength of the Chemical Entity is actually delisted.
- (d) In relation to community supply, if the successful tenderer's brand is the only brand of the Tender Item listed on the Pharmaceutical Schedule as at the Market Notification Date, then the First Transition Period and clause 1.1(a) of Schedule Five will not apply.
- (e) For the avoidance of doubt, any notification by PHARMAC of the delisting of all other brands of that form and strength of the Chemical Entity on the first day of the Sole Supply Period operates solely as advance notice of the intended delisting of those pharmaceuticals and does not constitute a notice of termination of any existing contract for the supply of those other brands.

1.3 **Contract**

If PHARMAC accepts your:

- (a) Community Tender Bid, then a contract on the terms and conditions set out in:
 - (i) your Tender Bid (incorporating any variations as a result of negotiations or discussions under clause 1.6 of this Schedule); and
 - (ii) Schedule Four; and
 - (iii) Schedule Five; and
 - (iv) Schedule Seven (as applicable),

will be deemed to have been entered into between you and PHARMAC for Sole Supply Status for the relevant Pharmaceutical and, where applicable, its listing on the Pharmaceutical Schedule;

(b) Hospital Tender Bid, then a contract on the terms and conditions set out in:

- (i) your Tender Bid (incorporating any variations as a result of negotiations or discussions under clause 1.6 of this Schedule); and
- (ii) Schedule Four; and
- (iii) Schedule Six; and
- (iv) Schedule Seven (as applicable),

will be deemed to have been entered into between you and PHARMAC for Hospital Supply Status for the relevant Pharmaceutical and, where applicable, its listing on the Pharmaceutical Schedule;

- (c) Combined Community/Hospital Tender Bid, then:
 - (i) a contract on the terms and conditions set out in:
 - (A) your Tender Bid, to the extent applicable (incorporating any variations as a result of negotiations or discussions under clause 1.6 of this Schedule); and
 - (B) Schedule Four; and
 - (C) for the Community Tender Bid element of that Combined Community/Hospital Tender Bid, Schedule Five; and
 - (D) Schedule Seven (as applicable),

will be deemed to have been entered into between you and PHARMAC for Sole Supply Status for the relevant Pharmaceutical and, where applicable, its listing on the Pharmaceutical Schedule; and

- (ii) a separate contract on the terms and conditions set out in:
 - (A) your Tender Bid, to the extent applicable (incorporating any variations as a result of negotiations or discussions under clause 1.6 of this Schedule); and
 - (B) Schedule Four; and
 - (C) for the Hospital Tender Bid element of that Combined Community/Hospital Tender Bid, Schedule Six; and
 - (D) Schedule Seven (as applicable),

will be deemed to have been entered into between you and PHARMAC for Hospital Supply Status for the relevant Pharmaceutical and, where applicable, its listing on the Pharmaceutical Schedule.

For the avoidance of doubt, the terms and conditions specified in Schedule Four, Schedule Five, Schedule Six and Schedule Seven, as applicable, apply from the date when PHARMAC notifies you in accordance with clause 7.2 of this Schedule of its acceptance of your Tender Bid, and do not apply solely for the Sole Supply Period or Hospital Supply Status Period, as applicable.

1.4 Extension of Hospital Supply Status to include Sole Supply Status

- (a) You acknowledge and agree that if your Hospital Tender Bid is for a Tender Item that is specified in the product list in clause 2 of Schedule Two and the Electronic Portal as being a Tender Item for which you may submit a Tender Bid for Sole Supply Status, you may agree (such consent not to be unreasonably withheld), if so requested by PHARMAC:
 - (i) if PHARMAC has not yet accepted a Hospital Tender Bid for the particular Tender Item, to extend your Tender Bid to cover community supply; or
 - (ii) if PHARMAC has accepted your Hospital Tender Bid for the particular Tender Item, to supply the Tender Item for use in the community under Sole Supply Status as soon as practicable after such requirement is notified to you, and in any case no later than three months after that notification, under a separate contract for Sole Supply Status.
- (b) The Community Tender Bid referred to in paragraph (a)(i) above and the contract for Sole Supply Status referred to in paragraph (a)(ii) above will be:
 - (i) at a price that is equal to the Price specified for that Pharmaceutical in your Hospital Tender Bid; and
 - (ii) on the other terms and conditions set out in your Hospital Tender Bid (incorporating any variations as a result of negotiations or discussions under clause 1.6 of this Schedule), as applicable; and
 - (iii) for supply in accordance with Schedules Four, Five and Seven; and
 - (iv) for such quantities of the Pharmaceutical as are required for use in the community.
- (c) This clause confers a benefit on, and is enforceable by, the Funder in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

1.5 **Extension of Sole Supply Status to include Hospital Supply Status**

- (a) You acknowledge and agree that if your Community Tender Bid is for a Tender Item that is specified in the product list in clause 2 of Schedule Two as being a Tender Item for which you may submit a Tender Bid for Hospital Supply Status, you may agree (such consent not to be unreasonably withheld), if so required by PHARMAC:
 - (i) if PHARMAC has not yet accepted a Community Tender Bid for the particular Tender Item, to extend your Tender Bid to cover hospital supply; or
 - (ii) if PHARMAC has accepted your Community Tender Bid for the particular Tender Item, to supply the Tender Item for use in DHB Hospitals under Hospital Supply Status as soon as practicable after such requirement is notified to you, and in any case no later than three months after that notification, under a separate contract for Hospital Supply Status.
- (b) The Hospital Tender Bid referred to in paragraph (a)(i) above and the contract for Hospital Supply Status referred to in paragraph (a)(ii) above will be:
 - (i) at a price that is equal to the Price specified for that Pharmaceutical in your Community Tender Bid; and

- (ii) on the other terms and conditions set out in your Community Tender Bid (incorporating any variations as a result of negotiations or discussions under clause 1.6 of this Schedule), as applicable; and
- (iii) for supply in accordance with Schedules Four, Six and Seven; and
- (iv) for such quantities of the Pharmaceutical as are required for use in DHB Hospitals.
- (c) This clause confers a benefit on, and is enforceable by, DHB Hospitals in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

1.6 **PHARMAC may initiate limited negotiations**

- (a) Notwithstanding clause 2.7 of this Schedule, PHARMAC may, in its sole discretion, initiate negotiations or discussions with you in relation to your Tender Bid about:
 - (i) any of the terms and conditions to apply if your Tender Bid is accepted;
 - (ii) the proposed packaging or pack size of the Tender Item;
 - (iii) your ability to ensure continued availability of the Tender Item throughout the Hospital Supply Status Period and/or Sole Supply Period, as applicable;
 - (iv) the price of the Tender Item, but only where PHARMAC determines, in its sole discretion, that an increased price for the Tender Item may be necessary for practicality of supply of the Tender Item (for example, because of particular packaging requirements);
 - (v) DV Limits and/or DV Pharmaceuticals, in relation to hospital supply;
 - (vi) the Lead Time and/or the Start Date; or
 - (vii) any other matter that PHARMAC considers necessary or appropriate.
- (b) If PHARMAC initiates negotiations or discussions with you under paragraph (a), and as a result there is a change to any of the terms and conditions relating to the supply of a Tender Item, PHARMAC is not obliged to inform the other tenderers of that change, nor give those tenderers an opportunity to amend their bid for that Tender Item, unless the change is one which would result in the terms and conditions being materially different in scope from those set out in this Invitation.
- (c) The initiation and pursuit of any negotiations or discussions under this clause shall not constitute a counter-offer and your original Tender Bid will remain open for acceptance in accordance with clause 4.2(b) of this Schedule in the absence of agreement on any variation to that Tender Bid.

1.7 **Termination and amendment of Invitation**

PHARMAC may:

- (a) amend this Invitation at any time up to five business days before the Deadline; and/or
- (b) terminate this Invitation at any time before the acceptance of any Tender Bid by giving five business days' written notice.

2. Information about submitting a Tender Bid

2.1 Choice of forms and strengths

Where a Tender Item includes different forms and strengths of a Chemical Entity or entities, your Tender Bid may, but does not need to, include all of the forms and strengths of the Chemical Entity or entities contained in that Tender Item.

2.2 Consents not yet held

You may submit a Tender Bid for a Tender Item where your brand of the Tender Item is yet to obtain all necessary Consents. In those circumstances, you may be required to demonstrate your ability to obtain those consents within a time frame acceptable to PHARMAC. For example, you may be required to demonstrate that you have the dossier for that brand of the Tender Item ready to submit to Medsafe within one month of such a request being made by PHARMAC. For the avoidance of doubt where your brand of the Tender Item is yet to obtain all necessary Consents, any time period to obtain those Consents shall be exclusive of the Lead Time indicated on your Tender Bid.

2.3 Individual Tender Bids

You may submit more than one bid for a Tender Item (for example, you may submit separate bids for different pack sizes (or other equivalent grouping for a Medical Device) of a Tender Item).

2.4 Aggregated Tender Bids

- (a) You may, in addition to submitting a separate Tender Bid for each Tender Item, submit an Aggregated Tender Bid, provided that:
 - (i) in the case of a pharmaceutical that is not a Medical Device, each brand contained in an Aggregated Tender Bid is only a different form and strength of the same Chemical Entity;
 - (ii) you may not aggregate across different chemical entities when submitting a Tender Bid;
 - (iii) you may not aggregate within a single Tender Item (for example, two different brands or pack sizes);
 - (iv) you must also submit a separate Community Tender Bid and/or Hospital Tender Bid, as applicable, for each particular Tender Item.
- (b) Where a Tender Item includes different forms and strengths of a Chemical Entity or different entities (for example, a two-part injection), and you bid for the whole Tender Item, that is not an Aggregated Tender Bid.

2.5 **Combined Community/Hospital Tender Bids**

You may submit a Combined Community/Hospital Tender Bid, provided that you must also submit a separate Community Tender Bid and a separate Hospital Tender Bid for each Tender Item in respect of which you submit a Combined Community/Hospital Tender Bid.

2.6 Aggregated Combined Community/Hospital Tender Bids

You may submit a Tender Bid that is both an Aggregated Tender Bid and a Combined Community/Hospital Tender Bid, provided that you comply with clauses 2.4 and 2.5 above.

2.7 **No conditions**

You cannot make a conditional Tender Bid nor qualify a Tender Bid in any way.

2.8 Separate offers

PHARMAC will treat each Tender Bid as a separate offer.

2.9 **Tender Bid prices**

You must submit, for each Tender Bid, a single price in New Zealand dollars (exclusive of GST), which will be the Price at which you will supply the Tender Item.

3. What to include in your Offer Letter and Tender Submission Form

3.1 **Compulsory use of Offer Letter and Tender Submission Form**

- (a) You must submit your Tender Bid using the Electronic Portal and attach the Offer Letter and a completed Tender Submission Form for each Tender Item for which you wish to submit a bid.
- (b) An electronic version of the Offer Letter is available on the Electronic Portal.

3.2 Information that must be supplied about you

In the Offer Letter, you must supply the following information about you:

- (a) your company structure;
- (b) your management and technical skills;
- (c) your financial resources;
- (d) your (or your supplier's) existing supply commitments;
- (e) your (or your supplier's) previous supply performance; and
- (f) your quality assurance processes, where applicable.

3.3 Information that must be supplied about the Tender Item

In your Tender Submission Form, you must supply the following information about the Tender Item:

(a) in the case of a pharmaceutical that is not a Medical Device, the chemical, form, strength, brand name, pack size and type of packaging;

- (b) for any Pharmaceutical that does not require Consent from Medsafe;
 - evidence and justification as to why Consent from Medsafe is not required for the Tender Item(s);
 - (ii) confirmation that the Tender Item(s) that you are submitting a Tender Bid in respect of meet the relevant standards and/or regulatory requirements for its intended use and what those standards and/or regulatory requirements are; and
 - (iii) details of the Tender Item(s), including excipients and shelf life;
- (c) for any Medical Device:
 - (i) the brand name, pack size (or other equivalent grouping) and type of packaging;
 - details of the Tender Item(s) and any associated services available in relation to the Tender item(s), including training, education and product support;
 - (iii) confirmation that the Tender Item(s) that you are submitting a Tender Bid in respect of meet the relevant standards and regulatory requirements for its intended use;
 - (iv) information on current usage of and expenditure on the Tender Item(s) by DHBs;
 - (v) confirmation that you have a business continuity plan with a brief summary of the plan;
 - (vi) demonstration of experience and knowledge within the healthcare sector, and specifically DHB Hospitals;
 - (vii) the WAND registration number of the Tender Item(s); and
 - (viii) the name of the sponsor of the Tender Item for the purpose of the Medicines (Database of Medical Devices) Regulations 2003;
- (d) a single price in New Zealand dollars (exclusive of GST) at which you will supply the Tender Item:
 - (i) to wholesalers and other distributors during the Sole Supply Period in respect of a Community Tender Bid; or
 - to, at a DHB Hospital's discretion, Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding), in respect of a Hospital Tender Bid;
- (e) whether it has all necessary Consents (and if not, what the status of registration is);
- (f) the Lead Time for supply of the Tender Item;
- (g) the name and location of:
 - (i) the manufacturer(s) of the finished product (and name and location of the packaging site, if different); and

- (ii) the manufacturer(s) of the active ingredients (not required in respect of Medical Devices); and
- (iii) alternative manufacturers of the finished product and active ingredients (if any) (not required in respect of Medical Devices);
- (h) your proposed distribution and supply arrangements for the Tender Item.

3.4 Information that may be supplied about the Tender Item

In your Tender Submission Form, you may supply the following information about the Tender Item:

- (a) For any Pharmaceutical or Medical Device:
 - (i) other markets you currently provide the Pharmaceutical or Medical Device in.

3.5 **PHARMAC may request further information**

- (a) PHARMAC may request such further information as it considers necessary from or about you for the purposes of clarifying or evaluating your Tender Bid, including (but not limited to):
 - (i) information about your credit status;
 - (ii) information on the price of a Tender Item, but only where PHARMAC requires clarification to confirm the exact price being offered, or where PHARMAC initiates negotiations with you under clause 1.6 of this Schedule;
 - (iii) where a Tender Item is a controlled drug, information about the form in which the Tender Item will be supplied, in which case you must supply that information within 10 business days of PHARMAC requesting the information; and
 - (iv) a sample pack or container of the Tender Item (and if you intend supplying it in a different form from that sample pack or container, information about the form in which it will be supplied), in which case you must supply that sample pack or container or information within 10 business days of PHARMAC requesting it.
- (b) If PHARMAC requests further information from or about you it is not obliged to request the same or any other information from or about any other party.

4. How to submit a Tender Bid

4.1 **Submission of Tender Bids**

All Tender Bids must be submitted via the Electronic Portal. Tender Bids or any copies of Tender Bids should not be delivered in person, by courier, by post, by facsimile or by email to PHARMAC.

4.2 Key dates

Your Tender Bid must:

(a) be submitted via the Electronic Portal by no later than the Deadline; and

- (b) be irrevocable and remain open for acceptance by PHARMAC until, as applicable:
 - (i) Friday, 31 July 2020;
 - (ii) the date specified for a Tender Item in Schedule Two or on the Electronic Portal (if any); or
 - (iii) if PHARMAC so requests at any time, such later date as you agree in writing.

5. Evaluation

5.1 **Process of evaluation**

The Evaluation Committee, taking such regulatory, legal, medical and other advice as it considers appropriate, will evaluate all conforming Tender Bids that have been checked for conformity under clause 6(a) of this Schedule, and any non-conforming Tender Bids that are admitted for consideration under clause 6(b) of this Schedule.

5.2 Matters for evaluation

The Evaluation Committee will evaluate Tender Bids in light of PHARMAC's statutory objective which is "to secure for eligible people in need of pharmaceuticals, the best health outcomes that are reasonably achievable from pharmaceutical treatment and from within the amount of funding provided". In doing so the Evaluation Committee will be guided by the Factors for Consideration (Factors) that form part of PHARMAC's then current Operating Policies and Procedures (OPPs), as published on PHARMAC's website (www.pharmac.govt.nz), to the extent applicable. More information on the Factors can be found at www.pharmac.health.nz/factors-for-consideration.

The requirement for PHARMAC to pursue its statutory objective means that particular emphasis will be given to those aspects of proposals which demonstrate "health outcomes", and those aspects of proposals which demonstrate the impact on the "funding provided" for pharmaceuticals. Those Factors which relate directly to these aspects will be given the greatest weight by the Evaluation Committee but all Factors are important.

The information to be taken into account in applying the Factors by the Evaluation Committee will include, in particular:

- (a) your ability to ensure continued availability of the Tender Item throughout the Sole Supply Period and/or Hospital Supply Status Period and each of the Transition Periods, as applicable, taking into account each of the following separate points:
 - (i) your financial resources;
 - (ii) your management and technical skills;
 - (iii) your, or your supplier's, existing supply commitments;
 - (iv) your, or your supplier's, previous supply performance;
 - (v) your quality assurance processes, where applicable;
 - (vi) the site of manufacture and packaging of the Pharmaceutical, and site of manufacture of the active ingredient;

- (vii) alternative manufacturers of the finished product and active ingredients (if any);
- (viii) other markets in which you currently supply the Pharmaceutical;
- (ix) your proposed distribution and supply arrangements for the Tender Item; and
- (x) the Lead Time for supply of the Tender Item;
- (b) the pack size (or other relevant grouping for a Medical Device) of the Tender Item and the type of packaging;
- (c) the price of the Tender Item;
- (d) the amount and timing of savings, including non-pharmaceutical savings accruing to the Funder or PHARMAC during the Hospital Supply Status Period and the Sole Supply Period, as applicable;
- (e) either:
 - (i) evidence that you have obtained, and still have, market approval for your brand of the Tender Item, and all necessary Consents; or
 - (ii) evidence that will enable the Evaluation Committee to form a view on the likelihood and timing of your brand of the Tender Item gaining all necessary Consents;
- (f) the name and location of the manufacturer of the finished product and active ingredients of the Tender Item; and
- (g) any other benefits to the Funder of selecting you as the supplier of the Tender Item.

6. **Conformity**

- (a) PHARMAC may, in its sole discretion, check your Tender Bid for conformity with this Invitation. If PHARMAC does elect to check your Tender Bid, it is not obliged to check all or any other Tender Bids for conformity, provided that in PHARMAC's judgment this would not be unfair to you in comparison to any other party. A Tender Bid will conform if it:
 - (i) is submitted via the Electronic Portal by the Deadline;
 - (ii) is submitted on the Tender Submission Form and an Offer Letter is attached;
 - (iii) has no conditions or qualifications attached;
 - (iv) includes all information required under clauses 3.2 and 3.3 of this Schedule; and
 - (v) otherwise complies, both as to form and substance, with the requirements of this Invitation.
- (b) PHARMAC may, in its sole discretion, provided that in PHARMAC's judgment this would not be unfair to you in comparison to any other party:
 - (i) exclude any non-conforming Tender Bid from consideration; or
 - (ii) consider, and accept, any non-conforming Tender Bid.

7. Decision

7.1 Decision on acceptance of Tender Bid

- (a) The Evaluation Committee will make a recommendation as to which Tender Bid should be accepted to PHARMAC's Board of Directors (or its delegate under Delegated Authority pursuant to Section 73 of the Crown Entities Act 2004, where applicable).
- (b) PHARMAC's Board of Directors (or its delegate, where applicable) will have the sole discretion to decide whether or not to accept a Tender Bid for any Tender Item.
- (c) PHARMAC's Board of Directors (or its delegate, where applicable):
 - (i) will use the Factors in PHARMAC's then current OPPs as applicable, in deciding whether or not to accept a Tender Bid for any Tender Item; and
 - (ii) is not obliged to act in accordance with any recommendation of the Evaluation Committee.

7.2 Notification of acceptance

- (a) Once PHARMAC's Board of Directors (or its delegate, where applicable) has decided under clause 7.1 above which Tender Bid (if any) to accept for a Tender Item, PHARMAC will, within a reasonable period of time, notify the successful tenderer in writing that it has been successful and in addition:
 - (i) subject to paragraph (b) below, if the successful Tender Bid is unconditionally accepted, PHARMAC will, within a reasonable period of time, notify each unsuccessful tenderer in writing of the identity of the successful tenderer; or
 - (ii) subject to paragraph (b) below, if the successful Tender Bid is conditionally accepted, PHARMAC will, within a reasonable period of time of that tender becoming unconditionally accepted, notify each unsuccessful tenderer in writing of the identity of the successful tenderer.
- (b) If for any reason you do not receive written notification from PHARMAC in accordance with paragraph (a) above, you will be deemed to have received the required notification on the date that each Tender Item you bid for is notified in the Pharmaceutical Schedule.

7.3 **PHARMAC's rights reserved**

- (a) PHARMAC reserves the right to accept or reject any Tender Bid.
- (b) While it is PHARMAC's current intention, unless specified otherwise in Schedule Two or the Electronic Portal, to enter into an agreement to award Hospital Supply Status and/or Sole Supply Status for each Tender Item, PHARMAC will not in any circumstances be bound to accept any or all Tender Bids and, in particular, PHARMAC will not be bound to accept the lowest or any other Tender Bid for a Tender Item.
- (c) Acceptance only occurs if, and when, PHARMAC's Board of Directors (or its delegate, where applicable) resolves to accept a Tender Bid and this acceptance is notified to the successful tenderer.

- (d) PHARMAC may take any action, including making any adjustments to the tender process that it considers appropriate, acting reasonably (provided that it notifies tenderers materially affected by such adjustments).
- (e) PHARMAC may, at any time, suspend or cancel in whole or in part, this tender process in order to fulfil its public law obligations through consultation, or otherwise. In this situation PHARMAC may (without limitation) ask you to adapt and resubmit your Tender Bid in light of consultation, or alternatively we may request that new Tender Bids be submitted (or in the case of a suspension PHARMAC may also resume the tender process without further change following the end of the period of suspension).

7.4 Conditional acceptance

- (a) Where the successful tenderer's brand of a Tender Item is yet to receive all necessary Consents:
 - (i) the contract referred to in clause 1.3 of this Schedule will be conditional upon such Consents being received within a time period specified by PHARMAC; and
 - (ii) PHARMAC may terminate the contract if such Consents have not been obtained, or in PHARMAC's view are unlikely to be obtained, within the period specified by PHARMAC.
- (b) Acceptance of a Tender Bid by PHARMAC's Board of Directors (or its delegate, where applicable), and the contract referred to in clause 1.3 of this Schedule may be conditional upon you satisfying PHARMAC that you will have sufficient stock of the Tender Item available to commence supply as at a date reasonably determined by PHARMAC.

8. Back-up supply

Back-up Supply Agreements

- (a) PHARMAC may at any time negotiate a Back-up Supply Agreement with another supplier for any Tender Item.
- (b) PHARMAC may, at its sole discretion, seek proposals for Back-up Supply Agreements under a separate process to this Invitation to Tender. PHARMAC does not seek submissions for Back-up Supply Agreements in response to this Invitation to Tender and is not obliged to consider proposals or bids for back-up supply submitted as part of the tender process.

9. Dealing with information

9.1 **Confidentiality**

Subject to clause 9.2 below, all Confidential Information is confidential to us and our employees, legal advisers, electronic procurement providers and other consultants (including PTAC and its sub-committees), the Ministry of Health, DHBs and the Funder. You acknowledge that it may be necessary or appropriate for PHARMAC to disclose Confidential Information:

- (a) pursuant to the Official Information Act 1982; or
- (b) in publicly notifying any acceptance of your Tender Bid; or

(c) otherwise pursuant to PHARMAC's public law or any other legal obligations.

PHARMAC may consult with you before deciding whether to disclose Confidential Information for the purposes described in paragraphs (a) and (c) above, in order to ascertain any objections you may have to the disclosure of any of the Confidential Information. You acknowledge, however, that it is for PHARMAC to decide, in its absolute discretion, whether it is necessary or appropriate to disclose information for any of the above purposes, provided that PHARMAC shall act in good faith in disclosing any Confidential Information. Outside the circumstances described in paragraphs (a) and (c) above, Confidential Information must not be disclosed by either of us (or by our employees, legal advisers and other consultants) unless:

- (d) the information is publicly available without any cause attributable to the disclosing party; or
- (e) the other party has been reasonably informed prior to disclosure, and the disclosure is:
 - (i) for the purposes of this Agreement; or
 - (ii) required by law; or
 - (iii) in a form, and of content, agreed to by us.

For the avoidance of doubt, information released by PHARMAC in accordance with paragraphs (a) to (c) above ceases to be Confidential Information and you agree that PHARMAC may release that information again at any time in future without consulting with you or obtaining your prior agreement.

9.2 Use of information

Generalised aggregated information regarding your Tender Bid that does not identify you or that cannot reasonably be expected to identify you or lead to the connection of you with your Tender Bid is not Confidential Information and PHARMAC may use and publish such information as it sees fit.

10. Miscellaneous

10.1 **Process contract**

In submitting a Tender Bid, you agree that you and PHARMAC are contractually bound to follow the process and comply with the obligations expressly contained in this Invitation.

10.2 **Costs**

PHARMAC is not liable in any way whatsoever for any direct or indirect costs incurred, or loss (including loss of profit) or damage sustained, by you in respect, or arising out, of this tendering process or the obtaining or granting of Hospital Supply Status and/or Sole Supply Status, as applicable, for your supply of the Tender Item including, without limitation, costs of obtaining all necessary Consents for any Tender Item.

10.3 No reliance

Your Tender Bid is submitted in reliance on your own knowledge, skill and independent advice, and not in reliance on any representations made by PHARMAC (including for these purposes the sales and market information (if any) provided in Schedule Two or on the Electronic Portal).

10.4 **No further liability**

PHARMAC is not, in any event, liable in contract, tort or any other way whatsoever for any direct or indirect loss (including loss of profit), damage or cost of any kind incurred by you or any other person in relation to this tendering process.

10.5 No lobbying

- (a) You are not to initiate any communication with PHARMAC or its advisors, the Minister of Health (or any Associate Ministers), the Ministry of Health (including its operating unit Medsafe), or a District Health Board or any of their officers or directors, at any time with a view to influencing the outcome of the tendering process.
- (b) Failure to comply with this clause will entitle PHARMAC, in its sole discretion, to disqualify you from this tendering process.

10.6 Enquiries

If you have any enquiries about this Invitation you should contact Emma Clarke and Nerissa Ramlall at PHARMAC. Any additional information that PHARMAC gives to you as a result of your enquiry will also be given by PHARMAC to other potential tenderers, if PHARMAC determines that such information is material.

10.7 Jurisdiction and governing law

We each submit to the exclusive jurisdiction of the New Zealand courts and agree that this Invitation is governed by New Zealand law.

Schedule 4: Contract terms for both Sole Supply Status and Hospital Supply Status

1. General

1.1 **Operating Policies and Procedures**

- (a) You acknowledge that:
 - PHARMAC is required to pursue the objectives, carry out the functions, and otherwise comply with the statutory obligations, prescribed for PHARMAC in the New Zealand Public Health and Disability Act 2000;
 - (ii) PHARMAC is subject to other statutory and public law obligations, which govern PHARMAC's decision-making processes;
 - (iii) PHARMAC has OPPs which provide guidance on the way in which PHARMAC carries out its statutory responsibilities in relation to the management of the Pharmaceutical Schedule;
 - (iv) PHARMAC's OPPs may be amended or updated from time to time, following consultation with relevant groups;
 - (v) the actions which PHARMAC may take under its OPPs include (without limitation):
 - (A) listing new pharmaceuticals;
 - (B) changing guidelines or restrictions on the purchasing, prescribing and dispensing of listed pharmaceuticals;
 - (C) changing the subsidy levels and/or market dynamics for pharmaceuticals as a result of PHARMAC adopting one of the strategies set out in the OPPs or by any other means;
 - (D) amending the basis on which pharmaceuticals are classified into therapeutic groups and sub-groups;
 - (E) delisting pharmaceuticals, or delisting all or part of a therapeutic group or sub-group;
 - (vi) any action taken by PHARMAC pursuant to its OPPs may impact on the listing of the Pharmaceutical.
- (b) PHARMAC agrees not to apply, amend or update its OPPs in order to avoid any of PHARMAC's obligations under this Agreement.

1.2 Amendments to Pharmaceutical Schedule

PHARMAC will consult with you before amending the Pharmaceutical Schedule if a proposed amendment would have a material adverse effect on the listing of the Pharmaceutical.

1.3 **Product identification codes**

You agree to obtain and notify PHARMAC by submitting a notification of product changes form of the Pharmacode, the GTIN and the CTPP for the Pharmaceutical as soon as these are notified to you, and in any event:

- (a) for brand changes, no later than the earlier of:
 - (i) 10 business days following the Market Notification Date; or
 - (ii) the 5th of the month immediately prior to the Start Date.

(b) for price changes, on the 12th of the month prior to the Start Date.

For the avoidance of doubt, this requirement does not apply in relation to any Pharmaceutical that is a Medical Device.

1.4 **Stock Reporting**

You shall provide PHARMAC with reports on stock levels for the Pharmaceuticals upon PHARMAC's request during the Sole Supply Period and/or Hospital Supply Status Period.

1.5 **Supplier Code of Conduct**

You shall comply with the New Zealand Government's Supplier Code of Conduct (see <u>https://www.procurement.govt.nz/assets/procurement-property/documents/supplier-code-of-conduct.pdf</u>).

2. Crown Direction

- (a) You acknowledge that PHARMAC must comply with any Crown Direction.
- (b) PHARMAC may terminate or amend the Agreement, or impose restrictions on the prescribing or dispensing of a Pharmaceutical, at any time during the Sole Supply Period or the Hospital Supply Status Period (as applicable) or the Transition Periods, if the termination, amendment or imposition of restrictions is required to give effect to a Crown Direction.
- (c) In the event that a Crown Direction is issued to PHARMAC that requires an amendment to be made to this Agreement to give effect to that direction:
 - PHARMAC will give you as much notice as practicable of the Crown Direction and of any amendments to this Agreement that are required to give effect to that direction;
 - (ii) the Agreement will be deemed to be amended so as to give effect to the Crown Direction from the date when such direction is due to take effect; and
 - (iii) you may terminate this Agreement on not less than six months' written notice to PHARMAC where the effect of the amendment required under sub-paragraph (ii) above is such that it is no longer viable, financially or otherwise, for you to continue supplying the Pharmaceutical or to perform your obligations under this Agreement.

3. Audit

- (a) PHARMAC may, from time to time, review your records and any other information you hold that relates to this Agreement with regard to stock levels, registration information and supply issues, for the purposes of auditing your compliance with this Agreement. In these circumstances, PHARMAC, in consultation with you, will determine the terms and manner of any such audit, which as a minimum, must include the following:
 - (i) the audit will be conducted by an auditor authorised by PHARMAC;
 - (ii) you agree to co-operate fully with PHARMAC and provide PHARMAC and the auditor with all reasonable assistance to ensure that any audit conducted under this clause is fully and properly completed to PHARMAC's satisfaction, including:
 - (A) allowing the auditor access to your premises, records and other information you hold that relates to this Agreement with regard to stock levels, registration information and supply issues, for the purposes of, and during the course of, conducting the audit;
 - (B) answering promptly any questions from PHARMAC or the auditor concerning any aspect of your compliance with this Agreement.
 - (iii) PHARMAC will give you 10 business days' notice of its intention to conduct an audit under this clause and will ensure that the conduct of any such audit, and access in terms of sub-paragraph (A) above, does not unreasonably disrupt your business operations.
- (b) PHARMAC will notify you in writing if an audit under this clause reveals any noncompliance with this Agreement. You agree to remedy any non-compliance within 10 business days of receiving such notice from PHARMAC or such other period as agreed with PHARMAC.
- (c) In relation to Hospital Supply Status, PHARMAC will use its best endeavours to audit compliance by DHB Hospitals with the DV Limits and related requirements set out under this Agreement.

4. Miscellaneous

4.1 Litigation support

If this Agreement or its terms (including the basis on which the Pharmaceutical is listed):

- (a) give rise to proceedings being issued against PHARMAC; or
- (b) result in PHARMAC being made a party to any proceedings issued by a third party,

you will give PHARMAC all assistance it reasonably requires to gather evidence (including expert medical and clinical evidence) for the purpose of those proceedings.

4.2 **Dispute resolution**

If there is a dispute between us arising out of, or in connection with, this Agreement, neither of us is to commence any proceedings relating to that dispute until the following procedure has been complied with:

- (a) the party claiming a dispute has arisen must give written notice to the other party specifying the nature of the dispute;
- (b) we will endeavour, in good faith, to resolve the dispute referred to in the notice by using informal dispute resolution techniques;
- (c) if we do not agree on a dispute resolution technique within 14 days after the date notice of a dispute was given, the dispute is to be mediated according to the standard mediation agreement of LEADR & IAMA (a body corporate incorporated in Australia, registered as an overseas company in New Zealand in accordance with Part 18 of the Companies Act 1993, trading as the Resolution Institute), and the Chair of LEADR & IAMA (or the Chair's nominee) will select the mediator and determine the mediator's remuneration;
- (d) a party seeking urgent interlocutory relief may, by notice to the other party, elect not to comply with the provisions of this clause, but only to the extent of the relief sought, and only for the period required to dispose of the application for interlocutory relief; and
- (e) pending resolution of the dispute, this Agreement will remain in full effect without prejudicing our respective rights and remedies.

For the avoidance of doubt you acknowledge and agree that where a dispute arises in respect of hospital supply, PHARMAC may elect to involve any relevant DHB in any part, or all, of the above procedure.

4.3 Advertising

You must ensure that any Advertisement aimed at consumers of pharmaceuticals which you procure to be published, or in any way participate or assist in publishing, does not breach any applicable:

- (a) statute or regulation, including the Fair Trading Act 1986, Medicines Act 1981 and Medicines Regulations 1984; or
- (b) industry standard, including the Advertising Standards Authority Codes of Practice and Medicines New Zealand Code of Practice.

For the purposes of this clause:

- (c) "Advertisement" means any words, whether written, printed or spoken, any pictorial representation or design, any sounds or visual images, or combination of sounds and visual images, or any other form of communication used or appearing to be used to promote:
 - (i) the sale of a Pharmaceutical; or
 - (ii) the use of a method of treatment involving a Pharmaceutical; and
- (d) references to a statute, regulation or industry standard include that statute, regulation or industry standard as amended or replaced from time to time.

4.4 **No derogation**

For the avoidance of doubt, the express provision of a remedy for, or consequence of, breach of any term of this Agreement does not derogate from any other legal right or remedy available to PHARMAC under this Agreement or otherwise in respect of such breach.

4.5 No waiver

A failure or delay by either of us to exercise any right arising under this Agreement is not a waiver of that right, and a waiver of a breach of this Agreement is not a waiver of any other breach.

4.5 Agreement prevails

Where any of your terms of supply, whether recorded on your invoices or in credit arrangements entered into or elsewhere, conflict with or detract from any of the terms of this Agreement, the terms of this Agreement will prevail and will apply to the exclusion of any of your terms or documentation.

4.6 Entire agreement

This Agreement:

- (a) is the entire agreement between us regarding the terms on which the Pharmaceutical is, as applicable:
 - (i) listed in Section B of the Pharmaceutical Schedule and subsidised by the Funder; and/or
 - (ii) listed in Section H of the Pharmaceutical Schedule and purchased by DHB Hospitals; and
- (b) supersedes and extinguishes, from the Start Date, all prior agreements and understandings between us, and between you and any District Health Board regarding supply of the Pharmaceutical to DHB Hospitals.

4.7 Amendments

Amendments to this Agreement are only effective if in writing and signed by both of us.

4.8 Assignment

You will not permit any part of this Agreement to be transferred, assigned or sub-contracted (either directly or due to a change of ownership or control) without PHARMAC's prior written consent (such consent not to be unreasonably withheld). Any such consent may be given subject to such reasonable conditions as PHARMAC sees fit but no such consent will relieve you from any liability or obligation under the terms of the Agreement, and you will continue to be responsible for the acts, defaults and neglects of your transferee, assignee or sub-contractor.

4.9 Further assurances

We both agree to execute any further documents and do any further acts within our power as may be reasonably necessary from time to time to give effect to the terms and intentions of this Agreement.

4.10 **Contracts Privity**

(a) For the purposes of the Contract and Commercial Law Act 2017, Part 2, Subpart 1, we both acknowledge that your obligations in this Agreement constitute promises which confer or are intended to confer a benefit on the Funder and related persons and/or DHB Hospitals and related persons (as applicable), and are enforceable at the suit of the Funder, any such DHB Hospitals or any related persons.

- (b) Except as expressly provided in paragraph (a) above, the parties do not intend to create rights in, or grant remedies to, any third party as a beneficiary of this Agreement, and all the provisions of this Agreement shall be for the sole and exclusive benefit of the parties.
- (c) For the avoidance of doubt, you acknowledge that PHARMAC may pursue damages or any other claim (including injunctive or other such relief) under this Agreement on its own account and/or on behalf of the Funder and/or DHB Hospitals (as applicable), in respect of any form of loss or damage incurred by PHARMAC and/or the Funder and/or DHB Hospitals.

4.11 Jurisdiction and governing law

We each submit to the exclusive jurisdiction of the New Zealand courts and agree that this Agreement is governed by New Zealand law.

Schedule 5: Additional contract terms for Sole Supply Status

1. Effect of Sole Supply Status

1.1 **Subsidy arrangements**

- (a) Subject to clause 3.1 of this Schedule, the Pharmaceutical will be subsidised, and you must supply it, during the First Transition Period at the Price. If any other brands of the Chemical Entity are listed on the Pharmaceutical Schedule, those brands will continue to be subsidised for the First Transition Period at the subsidy applicable to those brands immediately before the commencement of the First Transition Period.
- (b) All other brands of that form and strength of the Chemical Entity will be delisted on the first day of the Sole Supply Period, with the result that you will have Sole Supply Status for that form and strength of the Chemical Entity during the Sole Supply Period.
- (c) Subject to PHARMAC's other rights under this Agreement in relation to the Pharmaceutical, the Pharmaceutical will continue to be fully subsidised, and you must continue to supply it, at the Price throughout the Sole Supply Period.
- (d) Subject to PHARMAC's other rights under this Agreement in relation to the Pharmaceutical, the Pharmaceutical will not be delisted during the Final Transition Period.

1.2 Exclusivity for the Sole Supply Period

- (a) Subject to PHARMAC's other rights under this Agreement in relation to the Pharmaceutical, PHARMAC will not subsidise another supplier's brand of the Pharmaceutical on the Pharmaceutical Schedule at any time during the Sole Supply Period.
- (b) This clause does not prohibit PHARMAC from entering into negotiations or arrangements with, or inviting tenders from, other suppliers to be the sole subsidised supplier of any forms and strengths of the Chemical Entity, if such supply commences after the end of the Sole Supply Period.
- (c) For the avoidance of doubt, PHARMAC may lower the subsidy applicable to a Pharmaceutical during the Final Transition Period as it sees fit, including lowering the subsidy of a Pharmaceutical as a result of the implementation of new tender arrangements.

1.3 Withdrawal of Sole Supply Status

- (a) PHARMAC may withdraw Sole Supply Status in relation to your supply of the Pharmaceutical (in which case clauses 1.1 and 1.2 of this Schedule will no longer apply), by written notice to you at any time during the Sole Supply Period or (in anticipation) during the First Transition Period if:
 - (i) you have failed to notify PHARMAC as required under clause 5.1 of this Schedule;
 - (ii) you are unable to supply the Pharmaceutical in accordance with this Agreement for a period of 30 days;
 - (iii) any Consent for the Pharmaceutical is withdrawn; or

- (iv) you otherwise fail to supply the Pharmaceutical in accordance with this Agreement.
- (b) In the event that PHARMAC exercises its rights under clause 1.3(a) above in relation to a Pharmaceutical, it may also withdraw Sole Supply Status in relation to your supply of all forms and strengths of that Pharmaceutical (in which case clauses 1.1 and 1.2 of this Schedule will no longer apply), following a recommendation from its clinical advisors, either by the written notice provided under clause 1.3(a) above or by further written notice to you at any time during the Sole Supply Period or (in anticipation) during the First Transition Period.
- (c) Any withdrawal of Sole Supply Status is without prejudice to PHARMAC's rights under clauses 5.2 and 5.3 of this Schedule.

1.4 Suspension of Sole Supply Status

- (a) If, at any time during the Sole Supply Period or (in anticipation) during the First Transition Period, you are unable to meet demand for the Pharmaceutical, or you notify PHARMAC under clause 5.1 of this Schedule of a Potential Out-of-Stock Event, or you otherwise fail to supply the Pharmaceutical in accordance with this Agreement, PHARMAC may suspend Sole Supply Status in relation to your supply of the Pharmaceutical for the period of such inability.
- (b) In the event that PHARMAC exercises its rights under clause 1.4(a) above in relation to a Pharmaceutical, it may also suspend Sole Supply Status in relation to your supply of all forms and strengths of that Pharmaceutical, following a recommendation from its clinical advisors, either by the written notice provided under clause 1.4(a) above or by further written notice to you at any time during the Sole Supply Period or (in anticipation) during the First Transition Period.
- (c) Any suspension of Sole Supply Status is without prejudice to PHARMAC's rights under clauses 5.2 and 5.3 of this Schedule.
- (d) PHARMAC may, at any time, in its sole discretion, notify you of the date on which the suspension of Sole Supply Status under this clause 1.4 ceases and on which date:
 - (i) Sole Supply Status is to be re-implemented in respect of the Pharmaceutical; or
 - (ii) Sole Supply Status is to be withdrawn in accordance with clause 1.3 of this Schedule.

1.5 **Subsidy arrangements after the End Date**

- (a) Subject to paragraphs (b) and (c) below, the Pharmaceutical is to continue to be the subject of a listing agreement between you and PHARMAC with effect from the End Date, and accordingly:
 - (i) you will cease to have Sole Supply Status for that form and strength of the Chemical Entity; and
 - (ii) the Pharmaceutical will remain listed in Section B of the Pharmaceutical Schedule subject to PHARMAC's standard terms of supply for pharmaceuticals used in the community (as recorded in the then current general listing terms Annex of PHARMAC's standard community contract template).
 - (iii) you may increase the price ex-manufacturer (exclusive of GST) at which you supply the Pharmaceutical to wholesalers and other such distributors on giving PHARMAC

six months' written notice of that price increase. You may provide PHARMAC with this written notice at any time after, but not before, the End Date;

- (iv) if PHARMAC does not increase the subsidy for the Pharmaceutical to the new price notified under paragraph (a)(iii) above, you may withdraw the Pharmaceutical from supply on not less than six months' prior written notice;
- (v) if PHARMAC does increase the subsidy for the Pharmaceutical to the new price notified under paragraph (a)(iii) above, you may withdraw the Pharmaceutical from supply on not less than two years' prior written notice (except where the withdrawal is for reasons that PHARMAC considers to be wholly outside of your control, in which case you must first provide to PHARMAC such information as it may require from you in order to satisfy it, in its sole discretion, that you are required to withdraw supply); and
- (vi) if at the time of providing notice under paragraph (a)(v) above, you advise PHARMAC that you are required to purchase a significant quantity of extra stock of the Pharmaceutical to enable you to continue to supply for the two-year period, and you advise PHARMAC of the total cost of that stock, PHARMAC will either:
 - (A) use reasonable endeavours to enter into an agreement to reimburse you for stock that remains unsold at the end of that two-year period; or
 - (B) release you from your obligations to supply under this paragraph (a).
- (b) PHARMAC may at its sole discretion, with effect from the End Date:
 - (i) require that the Pharmaceutical does not continue to be the subject of a listing agreement, in which case PHARMAC will give you written notice not less than three months prior to the End Date; and/or
 - (ii) apply any of the strategies under PHARMAC's then current OPPs to the Pharmaceutical (including delisting the Pharmaceutical after the Final Transition Period).
- (c) In the event PHARMAC applies any of the strategies described in paragraph (b)(ii) above, you may withdraw the Pharmaceutical from supply on not less than six months' prior written notice. You may provide PHARMAC with this written notice at any time after, but not before, the date that the particular strategy takes effect in the Pharmaceutical Schedule.
- (d) Where a Pharmaceutical is designated an ASP, PHARMAC will provide at least two months' written notice of another supplier's brand of the Pharmaceutical being listed on the Pharmaceutical Schedule and a seven-month initial transition period.

2. Consents

2.1 Warranty and indemnity that Consents are held

You warrant that you have, and will maintain, all necessary Consents. If a Consent is not held by you or is withdrawn or the Pharmaceutical is no longer approved for the treatment of any indication for which it is subsidised, then:

(a) PHARMAC is entitled to terminate this Agreement by 14 days' written notice to you; and

(b) whether or not PHARMAC terminates this Agreement under paragraph (a) above, you are to indemnify the Funder for any additional costs incurred by it (or by PHARMAC on its behalf) as a result of that failure to hold all necessary Consents. This clause confers a benefit on (and is enforceable by) the Funder in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

2.2 Changed medicine notification

If the Ministry of Health issues a changed medicine notification in relation to a Pharmaceutical, and as a result a variant of the Pharmaceutical (the "**CMN Pharmaceutical**") is approved:

- (a) you must immediately notify PHARMAC; and
- (b) PHARMAC may take such action as it considers appropriate in relation to that Pharmaceutical or the CMN Pharmaceutical including (but not limited to):
 - (i) withdrawing Sole Supply Status for the Pharmaceutical;
 - (ii) reviewing the terms of listing of that Pharmaceutical; and
 - (iii) determining whether, and the extent to which, the Funder may subsidise the CMN Pharmaceutical.

3. Price

3.1 **Price change**

- (a) Subject to clause 3.1(b)(ii), clause 3.1(b)(iii) and clause 3.1(b)(iv) of this Schedule your brand of the Pharmaceutical must be available for supply, and you must supply the Pharmaceutical, at the Price from the 12th day of the month prior to the Start Date, and the Pharmaceutical will be subsidised at the Price from the Start Date.
- (b) In the event your brand of the Pharmaceutical is currently listed on the Pharmaceutical Schedule at the beginning of the First Transition Period:
 - (i) you must ensure that wholesalers and other such distributors change the price at which they supply the Pharmaceutical to the Price on the 12th day of the month prior to the Start Date, and you shall provide price support to wholesalers and other such distributors for a maximum 4 weeks stock on hand (or such other level of stock as agreed between you and PHARMAC) of the Pharmaceutical held at wholesalers and other such distributors, provided that such wholesalers and other such distributors can provide you with stock on hand reports upon request; or
 - (ii) your brand of the Pharmaceutical must be available for supply, and you must supply the Pharmaceutical, at the Price from the 1st day of the month prior to the Start Date, and the Pharmaceutical will be subsidised at the Price from the Start Date (which is conditional upon you having at least 2 months Lead Time for the Pharmaceutical); and
 - (iii) notwithstanding clauses 3.1(b)(i) or (b)(ii) above, if the Price would result in a price increase for your brand of the Pharmaceutical you must supply the Pharmaceutical at the Price from the 22nd day of the month prior to the Start Date, and the Pharmaceutical will be subsidised at the Price from the Start Date; and

(iv) notwithstanding clauses 3.1(b)(i), (b)(ii) or (b)(iii) above, PHARMAC may agree a process with you, that results in your brand of the Pharmaceutical, which includes a rebate, being available for supply, and you must supply the Pharmaceutical, at the Price from the 22nd day of the month prior to the Start Date, and you shall provide price support to wholesalers and other such distributors for a maximum 4 weeks stock on hand (or such other level of stock as agreed between you and PHARMAC) of the Pharmaceutical held at wholesalers and other such distributors, provided that such wholesalers and other such distributors can provide you with stock on hand reports upon request.

For the avoidance of doubt if you do not notify PHARMAC in your Tender Bid in the electronic portal which of the options stated in clauses 3.1(b)(i) or (b)(ii) above apply to the Pharmaceutical, clause (b)(i) above shall apply.

(c) You shall upon request by PHARMAC, provide information on how you intend to manage the price changes stated in clauses 3.1(b)(i) to (b)(iv) above. PHARMAC may, at its sole discretion, publish this information at the time the Tender Item is notified in the Pharmaceutical Schedule in accordance with clause 7.2 of Schedule 3.

3.2 Supply Price

During each of the First Transition Period, the Sole Supply Period and the Final Transition Period, the price at which the Pharmaceutical is supplied by you must not exceed the Price.

3.3 Warranty that Pharmaceutical is supplied at not less than cost price

You warrant that the price at which you are required to supply the Pharmaceutical under this Agreement is greater than the cost price of the Pharmaceutical (including, without limitation, the costs of manufacturing the Pharmaceutical and of supplying it to you for supply in New Zealand).

3.4 No reference pricing during Sole Supply Period

The subsidy payable for the Pharmaceutical will not be reduced as a result of a reduction in the reference price for the therapeutic sub-group of which it is a member during the Sole Supply Period. For the avoidance of doubt, PHARMAC will not be prevented from applying its reference pricing mechanisms to the Pharmaceutical to reduce the subsidy payable for it from the End Date.

3.5 **Unsold stock following delisting**

You acknowledge and agree that the price at which you are required to supply any Pharmaceutical under this Agreement incorporates, if applicable, any costs incurred by you associated with unsold stock of the Pharmaceutical held by you or any wholesaler or other distributor, after the Pharmaceutical has been delisted or after notification that it will be delisted.

4. Shelf-life of Pharmaceutical

- (a) You will not supply the Pharmaceutical to wholesalers, or other such distributors, or pharmacies if:
 - (i) the remaining shelf-life of the Pharmaceutical is less than six months; or
 - (ii) where the total shelf-life of the Pharmaceutical is less than six months, the remaining shelf-life is less than 75% of the Pharmaceutical's total shelf-life,

without prior written agreement from PHARMAC.

(b) If you have an agreement with PHARMAC to supply the Pharmaceutical, where the total shelf-life of the Pharmaceutical is less than six months and the remaining shelf-life is less than 75% of the Pharmaceutical's total shelf-life, and a particular wholesaler, or other such distributor, or pharmacy does not distribute or dispense that Pharmaceutical before its expiry or use-by date, you agree to allow that wholesaler, or other such distributor, or pharmaceutical to you and to provide that wholesaler, or other such distributor, or pharmacy with a credit for the Pharmaceutical.

5. **Out-of-stock arrangements**

5.1 Notification of Potential Out-of-Stock Event and supply of Alternative Pharmaceutical

- (a) You must notify PHARMAC in writing as soon as you have reasonable cause to believe at any time that you will fail to supply the Pharmaceutical in accordance with this Agreement and, in any event, you must notify PHARMAC if at any time a Potential Outof-Stock Event occurs, including during the Sole Supply Period or the First Transition Period, in which case PHARMAC may suspend Sole Supply Status in relation to your supply of the Pharmaceutical.
- (b) If a Potential Out-of-Stock Event occurs, or your failure to supply the Pharmaceutical in accordance with this Agreement will result in insufficient stock of the Pharmaceutical being available, then at PHARMAC's option:
 - (i) PHARMAC may implement an arrangement with another supplier to supply an Alternative Pharmaceutical (including an arrangement for back-up supply); or
 - (ii) you must use your best endeavours to procure wholesalers and other such distributors to supply, as soon as practicable, an Alternative Pharmaceutical to pharmacies at the Price, and PHARMAC will subsidise the Alternative Pharmaceutical at the Price.

5.2 General indemnity

You agree to indemnify the Funder if you fail to supply the Pharmaceutical in accordance with this Agreement (other than for reasons that PHARMAC reasonably considers, following discussion with you, to be wholly outside your control) whether as a result of:

- (a) your inability to meet demand for supply of the Pharmaceutical;
- (b) your withdrawal of the Pharmaceutical from supply;
- (c) any failure to have and maintain a Consent as specified in clause 2 of this Schedule;
- (d) any failure to notify PHARMAC in accordance with clause 5.1 above; or
- (e) for any other reason.

This indemnity:

(f) covers all additional costs, including without limitation all costs (if any) incurred in securing and subsidising an Alternative Pharmaceutical, incurred by the Funder (or by PHARMAC

on its behalf) as a result of your failure that are additional to any costs specified in clause 5.3; and

(g) confers a benefit on (and is enforceable by) the Funder in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

5.3 Liquidated damages

- (a) If you fail to supply the Pharmaceutical in accordance with this Agreement (other than for reasons that PHARMAC reasonably considers, following discussion with you, to be wholly outside your control) and:
 - (i) you have not notified PHARMAC under clause 5.1 of this Schedule, then without prejudice to PHARMAC's rights under clause 5.2:
 - (A) subject to paragraph (e) below, you must pay to PHARMAC (for the benefit of PHARMAC and the Funder) liquidated damages for the administrative and/or operational costs incurred by PHARMAC as a result of your failure to supply in the amount of \$50,000 per Pharmaceutical in respect of which you failed to notify PHARMAC; and
 - (B) PHARMAC may withdraw Sole Supply Status in relation to your supply of the Pharmaceutical under clause 1.3 of this Schedule; or
 - (ii) you have notified PHARMAC under clause 5.1 of this Schedule, then without prejudice to PHARMAC's rights under clause 5.2:
 - (A) you are not liable to pay any liquidated damages under this clause 5.3; and
 - (B) if you fail to supply the Pharmaceutical in accordance with this Agreement for more than 30 days, PHARMAC may withdraw Sole Supply Status in relation to your supply of the Pharmaceutical under clause 1.3 of this Schedule.
- (b) If, having notified PHARMAC under clause 5.1 of this Schedule, you remain able to, and you continue to, supply the Pharmaceutical, or an Alternative Pharmaceutical in accordance with clause 5.1(b)(ii) of this Schedule, such that there is no interruption to supply of the Pharmaceutical or of the Alternative Pharmaceutical in accordance with this Agreement, you will not be liable for any costs unless PHARMAC, in its sole discretion, has considered it necessary to enter into an arrangement with an alternative supplier under which PHARMAC has agreed to make a payment to that supplier to ensure continuity of supply, in which case you must indemnify the Funder or PHARMAC for that payment. Such indemnity will be limited to an amount of \$10,000.
- (c) You acknowledge and agree that:
 - the amounts of liquidated damages in this clause represent a reasonable estimate of the administrative and operational costs incurred by PHARMAC (including the use of staff and loss of opportunity as a result of use of staff time, and communication costs), the estimate being based on PHARMAC's previous experience; and
 - the amounts referred to as liquidated damages are not intended to include any penalty element nor any amount for costs relating to the securing of an Alternative Pharmaceutical, or the subsidisation of an Alternative Pharmaceutical,

provided that PHARMAC may, in its sole discretion, require you to pay less than the amount specified as liquidated damages if it is satisfied that the actual costs in the particular circumstances are less than the relevant amount so specified.

- (d) Where a Pharmaceutical in respect of which you are liable to pay liquidated damages pursuant to clause 5.3(a)(i)(A) above also has Hospital Supply Status and where you would otherwise be liable to pay the same amount of liquidated damages in respect of any corresponding failure under the terms of such Hospital Supply Status, you will only be required to pay liquidated damages of \$50,000 in total in respect of both supply failures.
- (e) All amounts referred to in this clause are plus GST.

5.4 **Failure to supply**

References in this clause 5 and elsewhere in this Schedule to your failure or inability to supply the Pharmaceutical in accordance with this Agreement, or your inability to meet demand for the Pharmaceutical, or insufficient stock of the Pharmaceutical being available, include, but are not limited to, circumstances where:

- (a) no stock of the Pharmaceutical is physically held by you or on your behalf in New Zealand;
- (b) the only stock of the Pharmaceutical physically held by you or on your behalf in New Zealand is stock to which clause 4(a)(i) or (ii) of this Schedule applies and no agreement has been reached with PHARMAC in terms of clause 4(a) of this Schedule;
- (c) you fail, directly or indirectly, to ensure that all orders for the Pharmaceutical are filled (without restricting quantities that may be ordered), including in particular where, for reasons attributable (wholly or partly) to you, not all patients for whom the Pharmaceutical is prescribed receive the full amount of the Pharmaceutical they require, or to which they are entitled, under their prescriptions, within the required time frames for dispensing under the then current contract, or notice under section 88 of the New Zealand Public Health and Disability Act 2000, in respect of pharmacy services;
- (d) you fail to supply the Pharmaceutical on and from the Start Date.

5.5 **Default interest and recovery costs**

If payment of any amount required to be paid by you under this clause 5 is not made by you, in full, by the due date for payment of that amount as notified to you in writing by PHARMAC, then:

- (a) interest will accrue in such sum as remains unpaid at a rate per annum equal to the relevant SME overdraft rate (weighted average rate) of the Reserve Bank of New Zealand plus five percentage points, calculated and compounded on a daily basis, from the due date until such time as the sum due (including interest) is paid in full. This obligation to pay default interest is to arise without the need for a notice or demand from PHARMAC for such default interest; and
- (b) PHARMAC may take any action, including legal action, without first needing to implement the dispute resolution procedure contained in clause 4.2 of Schedule Four, to recover that amount and you agree to pay to PHARMAC actual enforcement costs incurred in relation to that action.

6. Termination and restrictions

6.1 **Termination and restrictions for clinical reasons**

PHARMAC reserves the right, but only after consultation with you and a relevant medical adviser (being either the Ministry of Health, PTAC or its sub-committees), to:

- (a) terminate this Agreement at any time during the Sole Supply Period or the First Transition Period if the medical adviser determines for clinical reasons that it is no longer appropriate to have either:
 - (i) a sole subsidised supplier of that form and strength of the Chemical Entity; or
 - (ii) the Pharmaceutical as the sole subsidised brand; and/or
- (b) impose at any time during the Sole Supply Period or the Transition Periods restrictions on the prescribing or dispensing of a Pharmaceutical if those restrictions are necessary for clinical reasons.

6.2 Termination following an audit

PHARMAC may terminate the Agreement, or withdraw Sole Supply Status in relation to a Pharmaceutical, at any time during the Sole Supply Period or the Transition Periods, if you fail to remedy any area of non-compliance in accordance with clause 3(b) of Schedule Four.

7. Guarantee

- (a) PHARMAC may require an entity acceptable to it to provide a guarantee (in a form satisfactory to PHARMAC) of your performance obligations under clauses 5.2 and 5.3 of this Schedule including, without limitation, the payment of any sum payable under the indemnity or as liquidated damages pursuant to those clauses for any failure to supply the Pharmaceutical in accordance with this Agreement during the Sole Supply Period.
- (b) The guarantor's liability under such a guarantee will be limited to a total of \$100,000 per Chemical Entity for all claims made by PHARMAC under the guarantee.

Schedule 6: Additional contract terms for Hospital Supply Status

1. Effect of Hospital Supply Status

1.1 **Pricing arrangements**

- (a) Subject to PHARMAC's other rights under this Agreement and clause 3.1 of this Schedule, on and from the Start Date, during the remainder of the First Transition Period and during the Hospital Supply Status Period, the Pharmaceutical is to be:
 - (i) listed at the Price set out in Section H of the Pharmaceutical Schedule;
 - sold by you to, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding), at the Price.
- (b) Where the Pharmaceutical is included in an order by a DHB Hospital for pharmaceuticals where the total value (excluding GST) of the order is less than \$1,000, you may invoice the DHB Hospital, in accordance with clause 4.1 below, for the cost of freight for that particular order. For the avoidance of doubt, this clause 1.1(b) does not entitle you to invoice a DHB Hospital for any other costs in relation to the particular order.
- (c) Subject to PHARMAC's other rights under this Agreement in relation to the Pharmaceutical (including under clause 1.6 of this Schedule), and provided that there are no Alternative Pharmaceuticals listed in Section H of the Pharmaceutical Schedule at the start of the Final Transition Period, the Pharmaceutical:
 - (i) is to continue to be listed, sold and purchased at the Price referred to in clauses 1.1(a)(i) and (ii) above during the Final Transition Period and beyond; and
 - (ii) is not to be delisted during the Final Transition Period.

1.2 **Supplier for Hospital Supply Status Period**

- (a) Subject to:
 - (i) PHARMAC's other rights under this Agreement in relation to the Pharmaceutical; and
 - (ii) clauses 1.4 and 1.5 of this Schedule relating to the DV Limit for the Pharmaceutical,

your brand of the Pharmaceutical will be the brand listed in Section H of the Pharmaceutical Schedule, and purchased by DHB Hospitals at any time during the Hospital Supply Status Period, as the brand having Hospital Supply Status.

(b) This clause does not prohibit PHARMAC (on behalf of DHB Hospitals) from entering into negotiations or arrangements with, or inviting tenders from, other suppliers to be the supplier of any forms and strengths of the particular Pharmaceutical with Hospital Supply Status, or a relevant Alternative Pharmaceutical having a status equivalent to Hospital Supply Status, if notification of such an arrangement (once finalised) occurs, and such supply commences, after the end of the Hospital Supply Status Period.

1.3 **DV Pharmaceuticals**

- (a) PHARMAC may amend the relevant list of DV Pharmaceuticals specified in Section H of the Pharmaceutical Schedule, from time to time, in accordance with this clause 1.3, whereby:
 - (i) PHARMAC is only to remove a pharmaceutical listed as a DV Pharmaceutical if PHARMAC:
 - (A) has first obtained your agreement; or
 - (B) has a direction from Medsafe or its successor, or a recommendation from PTAC or its sub-committees, based on a significant clinical issue;
 - (ii) PHARMAC may add a pharmaceutical to the relevant list of DV Pharmaceuticals specified in Section H of the Pharmaceutical Schedule if such pharmaceutical is identified as a DV Pharmaceutical during the Hospital Supply Status Period or the First Transition Period by PHARMAC following a recommendation from PTAC or its sub-committees.
- (b) PHARMAC must consult with you prior to the removal of any pharmaceutical from the relevant list of DV Pharmaceuticals specified in Section H of the Pharmaceutical Schedule.

1.4 **DV Limit**

- (a) PHARMAC may, from time to time during the Hospital Supply Status Period or the First Transition Period, amend the DV Limit of the Pharmaceutical following what PHARMAC considers to be appropriate consultation with PTAC or its sub-committees, provided that PHARMAC may only increase the DV Limit without your prior agreement if it has a direction from Medsafe or its successor, or a recommendation from PTAC or its subcommittees, based on a significant clinical issue.
- (b) Subject to clause 1.5 of this Schedule you acknowledge and agree that while you have Hospital Supply Status:
 - DHB Hospitals may purchase DV Pharmaceuticals at any time within the First Transition Period and Final Transition Period without any requirement to comply with the DV Limit;
 - (ii) provided that DHB Hospitals collectively do not exceed the National DV Limit for the relevant Pharmaceutical, a DHB Hospital may purchase DV Pharmaceuticals at any time within the Hospital Supply Status Period;
 - (iii) without derogating from any other rights available to PHARMAC or DHB Hospitals under this Agreement or otherwise, if you fail to supply the Pharmaceutical in accordance with this Agreement (other than for a reason that PHARMAC reasonably considers, following discussion with you, to be wholly outside your control) within the Hospital Supply Status Period, then the relevant DHB Hospital is not required to comply with the DV Limit for the Pharmaceutical during that period of non-supply and the calendar month during which that nonsupply occurred will be excluded in any review of the DV Limit in accordance with clause 1.5 below;
 - (iv) if a DHB Hospital's usage of any DV Pharmaceuticals, in percentage terms, reaches or exceeds the percentage at which the Individual DV Limit is set for the relevant Pharmaceutical, that DHB Hospital may negotiate with you to agree to

vary the application of the Individual DV Limit to the DHB Hospital in respect of particular patients with exceptional needs.

1.5 **DV Limit Compliance**

(a) For the purposes of this clause 1.5:

(i) "Relevant Period" means:

- (A) the initial period starting on the day that the Hospital Supply Status Period begins up to and including 30 June 2021; or
- (B) for a Pharmaceutical listed on or prior to 30 June 2021, the period commencing on 1 July 2021 and ending on 30 June 2022 or, for a Pharmaceutical listed after 30 June 2022, the initial period starting on the date that the Hospital Supply Status Period begins up to and including 30 June 2022; or
- (C) for a Pharmaceutical listed on or prior to 30 June 2022, the period commencing on 1 July 2022 and ending on 30 June 2023, or, for a Pharmaceutical listed after 30 June 2022, the initial period starting on the date that the Hospital Supply Status Period begins up to and including 30 June 2023,

provided that for the purposes of carrying out the calculations in this clause 1.5 any calendar months that fall within those periods when there is any failure to supply the Pharmaceutical in accordance with this Agreement will be excluded.

(ii) **"Actual National DV Limit Indicator"** means, for a particular Pharmaceutical in any Relevant Period, such sum, expressed as a percentage, as is equal to:

(Total DV Pharmaceuticals Volume ÷ (Total DV Pharmaceuticals Volume + Total Pharmaceutical Volume)) x 100;

- (iii) "Total DV Pharmaceuticals Volume" means, for a particular Pharmaceutical in any Relevant Period, the total number of Units of all DV Pharmaceuticals of the relevant Pharmaceutical with Hospital Supply Status purchased by DHB Hospitals, as calculated by PHARMAC, following your request in accordance with clause 1.5(b) below, on the basis of the data extracted by PHARMAC from the electronic records used by it; and
- (iv) "Total Pharmaceutical Volume" means, for a particular Pharmaceutical with Hospital Supply Status in any Relevant Period, the total number of Units of that Pharmaceutical purchased by DHB Hospitals, as calculated by PHARMAC following your request in accordance with clause 1.5(b) below, on the basis of the data extracted by PHARMAC from the electronic records used by it.
- (b) If you reasonably believe that DHB Hospitals' percentage usage of DV Pharmaceuticals collectively exceeds the National DV Limit for a particular Pharmaceutical, you may at any time, but not more often than three-monthly, request that PHARMAC carry out calculations in accordance with the procedure set out in this clause 1.5 for the proportion of the Relevant Period that has passed to the date of your request, and PHARMAC may, in its discretion, agree to carry out the calculations for the Total DV Pharmaceuticals Volume, the Total Pharmaceutical Volume and the Actual National DV Limit Indicator, provided that if PHARMAC refuses to carry out such calculations, it will provide you with the reasons for refusing to do so.

- (c) It is acknowledged, for the avoidance of doubt, that if the Actual National DV Limit Indicator is less than the National DV Limit specified for the relevant Chemical Entity in Schedule Two and the Electronic Portal then, regardless of whether an individual DHB Hospital's percentage usage of DV Pharmaceuticals has exceeded the Individual DV Limit percentage for that Pharmaceutical, PHARMAC may decide, in its sole discretion, not to take any further action.
- (d) If the Actual National DV Limit Indicator is greater than the National DV Limit, PHARMAC will use its best endeavours to identify which individual DHB Hospitals' percentage usage of DV Pharmaceuticals have exceeded the Individual DV Limit percentage for that Pharmaceutical. You acknowledge that if PHARMAC cannot do this on the basis of information held by it, it may be necessary to obtain any further information you can provide. If neither of us can establish or quantify non-compliance by an individual DHB Hospital with the Individual DV Limit, then you acknowledge that PHARMAC may not be able to calculate for you, and you may not be able to obtain, financial compensation under clause 1.5(f)(ii) below. In that event you acknowledge, for the avoidance of doubt, that PHARMAC is not liable to pay any financial compensation on behalf of the relevant DHB.
- (e) If an individual DHB Hospital's percentage usage of DV Pharmaceuticals has exceeded the Individual DV Limit percentage for that Pharmaceutical as a result of DV Pharmaceutical usage that has been agreed to by you in accordance with clause 1.4(b)(iv) above then PHARMAC will not take any further action.
- (f) Subject to paragraph (e) above, PHARMAC will address the issue of non-compliance with any individual DHB Hospital or DHB Hospitals identified in accordance with paragraph (d) above by:
 - using its best endeavours to ensure that the relevant DHB Hospital complies with the DV Limit for that Pharmaceutical in the remainder of that Relevant Period (if applicable) and in any subsequent Relevant Period or Relevant Periods; and/or
 - (ii) following the end of a Relevant Period, and only once in respect of any Relevant Period, determining what financial compensation is payable by that DHB for its contribution towards exceeding the National DV Limit (where PHARMAC is able to quantify this based on the information available to it), being the greater amount of \$1,000 or such sum as is equal to:

DHB Deviation x Adjusted Price

where:

- (A) **"Adjusted Price"** means the Unit Price, for a particular Pharmaceutical in any Relevant Period, divided by two;
- (B) **"DHB Deviation"** is equal to:

(Total Contribution for $DHB_x \div Total$ Contribution for Exceeding DHBs) x Total DV Pharmaceuticals Volume in Excess of DV Limit

where:

"Total Contribution for DHB_x" means, for:

(a) a particular Pharmaceutical; and

(b) a particular DHB Hospital,

in any Relevant Period, the total number of Units of all DV Pharmaceuticals of the relevant Pharmaceutical with Hospital Supply Status purchased by that DHB Hospital minus the number of Units of DV Pharmaceuticals that corresponds to the percentage of the Individual Total Market Volume represented by the Individual DV Limit percentage for that Pharmaceutical, as calculated by PHARMAC for such Relevant Period on the basis of the data extracted by PHARMAC from the electronic records used by it;

"Total Contribution for Exceeding DHBs" means, for a particular Pharmaceutical in any Relevant Period, the sum of the Total Contribution for DHB_x for each DHB Hospital identified by PHARMAC in accordance with paragraph (d) above as exceeding the Individual DV Limit for that Relevant Period, as calculated by PHARMAC for such Relevant Period on the basis of the data extracted by PHARMAC from the electronic records used by it;

"Total DV Pharmaceuticals Volume in Excess of DV Limit" means, for a particular Pharmaceutical in any Relevant Period, the total number of Units of all DV Pharmaceuticals of the relevant Pharmaceutical with Hospital Supply Status purchased by DHB Hospitals in excess of the National DV Limit for that Relevant Period, as calculated by PHARMAC on the basis of the data extracted by PHARMAC from the electronic records used by it;

- (iii) PHARMAC will notify you and the relevant DHB in writing of any DV Limit compensation payable in accordance with clause 1.5(f)(ii) above. You may then invoice the relevant DHB for the amount of DV Limit compensation payable, as calculated and notified to you by PHARMAC. You must provide to PHARMAC a copy of any such invoice, and evidence of any payment received from the DHB in respect of that invoice, within 10 business days of sending such invoice or receiving such payment, respectively.
- (iv) If you have not received the amount of any DV Limit compensation payable in accordance with clause 1.5(f)(ii) above from the DHB within 60 business days of invoicing the DHB for the amount owing, then you may take such further actions (other than ceasing to supply) directly with the DHB as you consider appropriate to recover the amount owing to you. In that event you acknowledge, for the avoidance of doubt, that PHARMAC is not liable to pay any financial compensation on behalf of the relevant DHB.
- (v) For the avoidance of doubt, for the purposes of calculating the Total DV Pharmaceuticals Volume, the Total Contribution for DHB_x and the Total DV Pharmaceuticals Volume in Excess of DV Limit in this clause 1.5, if a pharmaceutical is added to, or removed from, the list of DV Pharmaceuticals during the Relevant Period in accordance with clause 1.3 of this Schedule, then only the number of Units of that pharmaceutical purchased by DHB Hospitals during the portion of the Relevant Period in which that pharmaceutical was a DV Pharmaceutical are to be included in those calculations.

1.6 **Supply arrangements after the End Date**

(a) Subject to paragraphs (b) and (c) below, the Pharmaceutical is to continue to be the subject of a listing agreement between you and PHARMAC with effect from the End Date, and accordingly:

- you will cease to have Hospital Supply Status for that form and strength of the Pharmaceutical (in the case of any Pharmaceutical that is not a Medical Device); or
- (ii) you will cease to have Hospital Supply Status in respect of an item conforming to the individual specifications described for the item in the product list in clause 2 of Schedule Two which the Pharmaceutical was listed as conforming with (in the case of any Pharmaceutical that is a Medical Device); and
- (iii) the Pharmaceutical will remain listed in Section H of the Pharmaceutical Schedule subject to PHARMAC's standard terms of supply for pharmaceuticals used in DHB Hospitals (as recorded in the then current general listing terms Annex of PHARMAC's standard hospital contract template);
- (iv) you may increase the price (exclusive of GST) at which you supply the Pharmaceutical to, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding), on giving PHARMAC six months' written notice of that price increase. You may provide PHARMAC with this written notice at any time after, but not before, the End Date;
- (v) you may withdraw the Pharmaceutical from supply on not less than two years' prior written notice (except where the withdrawal is for reasons that PHARMAC considers to be wholly outside of your control, in which case you must first provide to PHARMAC such information as it may require from you in order to satisfy it, in its sole discretion, that you are required to withdraw supply); and
- (vi) if at the time of providing notice under paragraph (a)(iv) above, you advise PHARMAC that you are required to purchase a significant quantity of extra stock of the Pharmaceutical to enable you to continue to supply for the two-year period, and you advise PHARMAC of the total cost of that stock, PHARMAC will either:
 - (A) use reasonable endeavours to enter into an agreement to reimburse you for stock that remains unsold at the end of that two-year period; or
 - (B) release you from your obligations to supply under this paragraph (a).
- (b) PHARMAC may, at its sole discretion, with effect from the End Date:
 - (i) require that the Pharmaceutical does not continue to be the subject of a listing agreement, in which case PHARMAC will give you written notice not less than three months prior to the End Date; and/or
 - (ii) apply any of the strategies under PHARMAC's then current OPPs to the Pharmaceutical (including delisting the Pharmaceutical after the Final Transition Period).
- (c) In the event PHARMAC applies any of the strategies described in paragraph (b)(ii) above, you may withdraw the Pharmaceutical from supply on not less than six months' prior written notice. You may provide PHARMAC with this written notice at any time after, but not before, the date that the particular strategy takes effect in the Pharmaceutical Schedule.

1.7 Withdrawal of Hospital Supply Status

- (a) PHARMAC may withdraw Hospital Supply Status in relation to your supply of the Pharmaceutical (in which case clauses 1.1, 1.2 and 1.3 of this Schedule will no longer apply), by written notice to you at any time during the Hospital Supply Status Period or (in anticipation) during the First Transition Period if:
 - (i) you have failed to notify PHARMAC as required under clause 7.1 of this Schedule;
 - (ii) you fail, for a period of 30 days, to supply the Pharmaceutical in accordance with this Agreement to any of the DHB Hospitals including to, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding);
 - (iii) any Consent for the Pharmaceutical required under clause 2 of this Schedule is withdrawn;
 - (iv) you have failed to comply with clause 6 of this Schedule on more than one occasion; or
 - (v) you otherwise fail to supply the Pharmaceutical in accordance with this Agreement.
- (b) In the event that PHARMAC exercises its rights under clause 1.7(a) above in relation to a Pharmaceutical, it may also withdraw Hospital Supply Status in relation to your supply of all forms and strengths of that Pharmaceutical (or your supply of all other Medical Devices under this Agreement, where PHARMAC has exercised its rights under clause 1.7(a) above in respect of a Medical Device) (in which case clauses 1.1, 1.2 and 1.3 of this Schedule will no longer apply), following a recommendation from its clinical advisors, either by the written notice provided under clause 1.7(a) above or by further written notice to you at any time during the Hospital Supply Period or (in anticipation) during the First Transition Period.
- (c) Any withdrawal of Hospital Supply Status is without prejudice to PHARMAC's rights under clauses 7.2 and 7.3 of this Schedule.

1.8 Suspension of Hospital Supply Status

- (a) If, at any time during the Hospital Supply Status Period or (in anticipation) during the First Transition Period, you are unable to meet demand for the Pharmaceutical, or you notify PHARMAC under clause 7.1 of this Schedule of a Potential Out-of-Stock Event, or you otherwise fail to supply the Pharmaceutical in accordance with this Agreement, then:
 - (i) PHARMAC may suspend Hospital Supply Status in relation to your supply of the Pharmaceutical for the period of such inability; and
 - (ii) DHB Hospitals may purchase DV Pharmaceuticals during the period when Hospital Supply Status is suspended without the requirement to comply with the DV Limit for the relevant Pharmaceutical.
- (b) In the event that PHARMAC exercises its rights under clause 1.8(a) above in relation to a Pharmaceutical, it may also suspend Hospital Supply Status in relation to your supply of all forms and strengths of that Pharmaceutical (or your supply of all other

Medical Devices under this Agreement, where PHARMAC has exercised its rights under clause 1.8(a) above in respect of a Medical Device), following a recommendation from its clinical advisors, either by the written notice provided under clause 1.8(a) above or by further written notice to you at any time during the Hospital Supply Period or (in anticipation) during the First Transition Period.

- (c) Any suspension of Hospital Supply Status is without prejudice to PHARMAC's rights under clauses 7.2 and 7.3 of this Schedule.
- (d) PHARMAC may, at any time, in its sole discretion, notify you of the date on which the suspension of Hospital Supply Status under this clause 1.8 ceases and on which date:
 - (i) Hospital Supply Status is to be re-implemented in respect of the Pharmaceutical; or
 - (ii) Hospital Supply Status is to be withdrawn in accordance with clause 1.7 of this Schedule.

2. Consents

2.1 Warranty and indemnity that Consents are held

You warrant that you have, and will maintain, all necessary Consents. If a Consent is not held by you or is withdrawn or the Pharmaceutical is no longer approved for the treatment of any indication for which it is listed in Section H of the Pharmaceutical Schedule, then:

- (a) PHARMAC is entitled to terminate this Agreement by 14 days' written notice to you; and
- (b) whether or not PHARMAC terminates this Agreement under paragraph (a) above, you are to indemnify the Funder for any additional costs incurred by it (or by PHARMAC on its behalf) as a result of that failure to hold all necessary Consents. This clause confers a benefit on (and is enforceable by) the Funder in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

2.2 Changed medicine notification

If the Ministry of Health issues a changed medicine notification in relation to a Pharmaceutical, and as a result a variant of the Pharmaceutical (the "**CMN Pharmaceutical**") is approved:

- (a) you must immediately notify PHARMAC; and
- (b) PHARMAC may take such action as it considers appropriate in relation to that Pharmaceutical or the CMN Pharmaceutical including (but not limited to):
 - (i) withdrawing Hospital Supply Status for the Pharmaceutical;
 - (ii) reviewing the terms of listing of that Pharmaceutical; and
 - (iii) determining whether, and the extent to which, DHB Hospitals may purchase the CMN Pharmaceutical.

3. Price

3.1 **Price change**

- (a) Subject to clause 3.1(b)(ii), clause 3.1(b)(iii) and clause 3.1(b)(iv) of this Schedule, you must change the price at which you supply the Pharmaceutical to, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding), to the Price with effect from the 12th day of the month prior to the Start Date. If your brand of the Pharmaceutical is not listed on the Pharmaceutical Schedule at the beginning of the First Transition Period, it must be available for supply or sale, and you must supply or sell it, at the Price on and from the 12th day of the month prior to the Start Date.
- (b) In the event your brand of the Pharmaceutical is currently listed on the Pharmaceutical Schedule at the beginning of the First Transition Period:
 - (i) you must ensure that wholesalers and other such distributors change the price at which they supply the Pharmaceutical to the Price on the 12th day of the month prior to the Start Date, and you shall provide price support to wholesalers and other such distributors for a maximum 4 weeks stock on hand (or such other level of stock as agreed between you and PHARMAC) of the Pharmaceutical held at wholesalers and other such distributors, provided that such wholesalers and other such distributors can provide you with stock on hand reports upon request; or
 - (ii) your brand of the Pharmaceutical must be available for supply, and you must supply the Pharmaceutical, at the Price from the 1st day of the month prior to the Start Date, and the Pharmaceutical will be subsidised at the Price from the Start Date (which is conditional upon you having at least 2 months Lead Time for the Pharmaceutical); and
 - (iii) notwithstanding clauses 3.1(b)(i) or (b)(ii) above, if the Price would result in a price increase for your brand of the Pharmaceutical you must supply the Pharmaceutical at the Price from the 22nd day of the month prior to the Start Date, and the Pharmaceutical will be subsidised at the Price from the Start Date; and
 - (iv) notwithstanding clauses 3.1(b)(i), (b)(ii) or (b)(iii) above, PHARMAC may agree a process with you, that results in your brand of the Pharmaceutical, which includes a rebate, being available for supply, and you must supply the Pharmaceutical, at the Price from the 22nd day of the month prior to the Start Date, and you shall provide price support to wholesalers and other such distributors for a maximum 4 weeks stock on hand (or such other level of stock as agreed between you and PHARMAC) of the Pharmaceutical held at wholesalers and other such distributors, provided that such wholesalers and other such distributors can provide you with stock on hand reports upon request.

For the avoidance of doubt if you do not notify PHARMAC in your Tender Bid in the electronic portal which of the options stated in clauses 3.1(b)(i) or (b)(ii) above apply to the Pharmaceutical, clause (b)(i) above shall apply.

(c) You shall upon request by PHARMAC, provide information on how you intend to manage the price changes stated in clauses 3.1(b)(i) to (b)(iv) above. PHARMAC may, at its sole discretion, publish this information at the time the Tender Item is notified in the Pharmaceutical Schedule in accordance with clause 7.2 of Schedule 3.

3.2 Supply price

Subject to clause 3.1 of this Schedule, during each of the First Transition Period, the Hospital Supply Status Period and the Final Transition Period, if applicable in accordance with clause 1.1(b) of this Schedule, the price at which the Pharmaceutical is supplied by you to, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding), must not exceed the Price.

3.3 Supply at lower price

Notwithstanding clauses 3.1 and 3.2 above but subject to clause 3.4 below, you may supply the Pharmaceutical to, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding) at a price lower than the Price, provided that where you decide to supply the Pharmaceutical in respect of any one or more DHB Hospital(s) at a price lower than the Price, you must supply the Pharmaceutical at the same lower price to all DHB Hospitals in respect of which you supply the Pharmaceutical, in which case that lower price will be deemed to be the Price of that Pharmaceutical for the purposes of this Agreement.

3.4 Warranty that Pharmaceutical is supplied at not less than cost price

You warrant that the Price at which you are required to supply the Pharmaceutical under this Agreement is greater than the cost price of the Pharmaceutical (including, without limitation, the costs of manufacturing the Pharmaceutical and of supplying it to you for supply in New Zealand).

3.5 Unsold stock following delisting

You acknowledge and agree that the price at which you are required to supply any Pharmaceutical under this Agreement incorporates, if applicable, any costs incurred by you associated with unsold stock of the Pharmaceutical held by you or any wholesaler or other distributor, after the Pharmaceutical has been delisted or after notification that it will be delisted.

4. Invoicing and Payment

4.1 Invoice

You are to invoice DHB Hospitals at the end of each month, but no later than the tenth day following the month to which the invoice in respect of the Pharmaceutical relates, specifying for the Pharmaceutical supplied during that month:

- (a) your delivery note reference number;
- (b) the particular DHB's purchase order reference number (if applicable);
- (c) the net amount payable in respect of the Pharmaceutical supplied to that DHB in accordance with this Agreement;
- (d) full details in respect of the Pharmaceutical supplied to that DHB in accordance with this Agreement, including the:
 - (i) DHB's item codes;

- (ii) quantity of the Pharmaceutical supplied;
- (iii) price of the Pharmaceutical;
- (iv) cost of freight for orders that included the Pharmaceutical (only where applicable under clause 1.1(b) above);
- (v) total cost for the total amount of the Pharmaceutical supplied; and
- (e) any other information that DHB Hospital requires you to supply.
- (f) The provisions of Clause 4.1 do not apply to the extent that both parties have agreed to alternative or varied invoicing arrangements in respect of a particular Pharmaceutical.

4.2 **Payment**

- (a) Provided that the Pharmaceutical has been supplied in accordance with this Agreement, and the particular DHB receives an invoice in accordance with clause 4.1 above, payment by the DHB Hospital to you of the amount required to be paid by it is expected to occur:
 - (i) by electronic funds transfer or such other method of payment as is designated by that DHB Hospital;
 - (ii) on the 20th day of the month following the month to which the invoice for the Pharmaceutical relates, or, if the 20th day of the month is not a business day, then on the next business day following the 20th of the month.
- (b) Where you invoice a DHB Hospital later than the tenth day following the month to which the invoice in respect of the Pharmaceutical relates then, provided that the Pharmaceutical has been supplied in accordance with this Agreement, and the invoice otherwise accords with clause 4.1 above, payment by the DHB Hospital to you of the amount required to be paid by it is expected to occur:
 - (i) by electronic funds transfer or such other method of payment as is designated by that DHB Hospital;
 - (ii) on the 20th day of the month following the month in which you invoice the DHB for the Pharmaceutical, or, if the 20th day of the month is not a business day, then on the next business day following the 20th of the month.

4.3 **Future payment**

- (a) A particular DHB Hospital's failure to dispute any invoice prior to payment does not prejudice that DHB Hospital's right subsequently to dispute the correctness of such an invoice, nor its ability to recover any amount of overpayment from you.
- (b) A DHB Hospital may withhold, deduct or set off the amount of any overpayment or any amount recoverable by that DHB Hospital from you under this Agreement from any future amount owing to you.

4.4 **Contracts Privity**

This clause 4 confers a benefit on (and is enforceable by) DHB Hospitals in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

5. Emergency and disaster supply

In the event of an emergency or disaster affecting any DHB Hospital, or an emergency or disaster on a national level, you will use your best endeavours to provide such quantities of the Pharmaceutical as are required by the relevant DHB Hospital(s). Your obligations under this clause include, but are not limited to, using your best endeavours to:

- (a) source the Pharmaceutical from other suppliers and distributors within New Zealand; and
- (b) source the Pharmaceutical or a pharmaceutical that is the same brand as the Pharmaceutical from any overseas manufacturer, supplier or distributor, and airfreighting that stock to New Zealand (for which the relevant DHB Hospital will meet all reasonable costs) for supply, either under Medsafe's explicit consent to import, sell or distribute the Pharmaceutical or under section 29 of the Medicines Act 1981, to DHB Hospitals.

6. Defective and short-dated Pharmaceuticals

6.1 **Pharmaceutical recall**

- (a) In the event that you are required by the Ministry of Health or any other authorities to recall the Pharmaceutical or a particular batch of the Pharmaceutical, you will notify PHARMAC and the relevant DHB Hospitals immediately you become aware of the need to recall the Pharmaceutical or that batch of the Pharmaceutical.
- (b) You will use your best endeavours to provide replacement Pharmaceuticals to DHB Hospitals as soon as possible.
- (c) If you fail to provide replacement Pharmaceuticals or an Alternative Pharmaceutical within what DHBs consider to be a reasonable time frame, then DHB Hospital(s) may purchase an Alternative Pharmaceutical elsewhere. Any reasonable additional costs incurred by DHB Hospital(s) in purchasing such an Alternative Pharmaceutical will be met by you on demand by PHARMAC or the DHB Hospital(s) and will be recoverable from you as a debt due to PHARMAC and to the DHB Hospital(s), as applicable.
- (d) In the event that the Pharmaceutical or a particular batch of the Pharmaceutical is recalled as contemplated by paragraph (a) above, you shall immediately refund to the relevant DHB Hospitals all money paid by them to you for or on account of the Pharmaceutical or that batch of the Pharmaceutical and such money will be recoverable from you as a debt due to the relevant DHB Hospitals, unless you have provided a replacement Pharmaceutical to the relevant DHB Hospitals' satisfaction.
- (e) This clause confers a benefit on (and is enforceable by) DHB Hospitals in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

6.2 **Shelf-life of Pharmaceutical**

(a) You will not supply the Pharmaceutical if:

- (i) the remaining shelf-life of the Pharmaceutical is less than six months; or
- (ii) where the total shelf-life of the Pharmaceutical is less than six months, the remaining shelf-life is less than 75% of the Pharmaceutical's total shelf-life,

without prior agreement from the relevant DHB Hospital.

- (b) If you have an agreement with the relevant DHB Hospital to supply the Pharmaceutical, where the total shelf-life of the Pharmaceutical is less than six months and the remaining shelf-life is less than 75% of the Pharmaceutical's total shelf-life, and that DHB Hospital does not use the Pharmaceutical before its expiry or use-by date, you agree to allow that DHB Hospital to return the Pharmaceutical to you and to provide that DHB Hospital with a credit for the Pharmaceutical.
- (c) This clause confers a benefit on (and is enforceable by) DHB Hospitals in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

7. Out-of-stock arrangements

7.1 Notification of Potential Out-of-Stock Event and supply of Alternative Pharmaceutical

- (a) You must notify PHARMAC in writing as soon as you have reasonable cause to believe at any time that you will fail to supply the Pharmaceutical in accordance with this Agreement and, in any event, you must notify PHARMAC and the relevant DHB Hospitals if at any time a Potential Out-of-Stock Event occurs, including during the Hospital Supply Period or the First Transition Period.
- (b) If a Potential Out-of-Stock Event occurs, or your failure to supply the Pharmaceutical in accordance with this Agreement will result in insufficient stock of the Pharmaceutical being available, then at PHARMAC's option:
 - (i) PHARMAC may implement an arrangement with another supplier to supply an Alternative Pharmaceutical (including an arrangement for back-up supply); and/or
 - (ii) you must use your best endeavours to procure, as soon as practicable, an Alternative Pharmaceutical for supply to, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding), at the Price, and if you are unable to do so you will pay to DHB Hospitals any additional costs incurred by DHB Hospitals as a result of the purchase price for the Alternative Pharmaceutical being higher than the Price.

7.2 General indemnity

You agree to indemnify DHB Hospitals and PHARMAC if you fail to supply the Pharmaceutical in accordance with this Agreement (other than for reasons that PHARMAC reasonably considers, following discussion with you, to be wholly outside your control) whether as a result of:

- (a) your inability to meet demand for supply of the Pharmaceutical;
- (b) your withdrawal of the Pharmaceutical from supply;

- (c) any failure to have and maintain a Consent as specified in clause 2 of this Schedule;
- (d) any failure to notify PHARMAC in accordance with clause 7.1 above; or
- (e) for any other reason.

This indemnity:

- (f) covers all additional costs, including without limitation all costs (if any) incurred in securing and purchasing an Alternative Pharmaceutical, incurred by DHB Hospitals (or by PHARMAC on their behalf) as a result of your failure that are additional to any costs specified in clause 7.3; and
- (g) confers a benefit on (and is enforceable by) DHB Hospitals in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

7.3 Liquidated damages

- (a) If you fail to supply the Pharmaceutical in accordance with this Agreement (other than for reasons that PHARMAC reasonably considers, following discussion with you, to be wholly outside your control) and:
 - (i) you have not notified PHARMAC and the relevant DHB Hospitals under clause 7.1 of this Schedule, then without prejudice to PHARMAC's and the relevant DHB Hospitals' rights under clause 7.2 above, but subject to paragraph (e) below, you must pay to PHARMAC (for the benefit of PHARMAC and DHB Hospitals) liquidated damages for the administrative and/or operational costs incurred by PHARMAC and DHB Hospitals as a result of your failure to supply in the amount of \$50,000 per Pharmaceutical in respect of which you failed to notify PHARMAC; or
 - (ii) you have notified PHARMAC and the relevant DHB Hospitals under clause 7.1 of this Schedule, then without prejudice to PHARMAC's and the relevant DHB Hospitals' rights under clause 7.2 above you are not liable to pay any liquidated damages under this clause 7.3.
- (b) If, having notified PHARMAC and the relevant DHB Hospitals under clause 7.1 of this Schedule, you remain able to, and you continue to, supply the Pharmaceutical, or an Alternative Pharmaceutical in accordance with clause 7.1(b)(ii) of this Schedule, such that there is no interruption to supply of the Pharmaceutical or of the Alternative Pharmaceutical in accordance with this Agreement, you will not be liable for any costs unless PHARMAC, in its sole discretion, has considered it necessary to enter into an arrangement with an alternative supplier under which PHARMAC or the relevant DHB Hospitals have agreed to make a payment to that supplier to ensure continuity of supply, in which case you must indemnify the relevant DHB Hospitals and PHARMAC for that payment. Such indemnity will be limited to an amount of \$10,000 per Pharmaceutical.
- (c) You acknowledge and agree that:
 - the amounts of liquidated damages in this clause represent a reasonable estimate of the administrative and operational costs incurred by PHARMAC and DHB Hospitals (including the use of staff and loss of opportunity as a result of use of staff time, and communication costs), the estimate being based on PHARMAC's and DHB Hospitals' previous experience; and

(ii) the amounts referred to as liquidated damages are not intended to include any penalty element nor any amount for costs relating to the securing of an Alternative Pharmaceutical, or the purchasing of an Alternative Pharmaceutical,

provided that PHARMAC may, in its sole discretion, require you to pay less than the amount specified as liquidated damages if it is satisfied that the actual costs in the particular circumstances are less than the relevant amount so specified.

- (d) Where a Pharmaceutical in respect of which you are liable to pay liquidated damages pursuant to clause 7.3(a)(i) above also has Sole Supply Status and where you would otherwise be liable to pay the same amount of liquidated damages in respect of any corresponding failure under the terms of such Sole Supply Status, you will only be required to pay liquidated damages of \$50,000 in total in respect of both supply failures.
- (e) All amounts referred to in this clause are plus GST.

7.4 Failure to supply

References in this clause 7 and elsewhere in this Schedule to your failure or inability to supply the Pharmaceutical in accordance with this Agreement, or your inability to meet demand for supply of the Pharmaceutical, or insufficient stock of the Pharmaceutical being available, include, but are not limited to, circumstances where:

- (a) no stock of the Pharmaceutical is physically held by you or on your behalf in New Zealand;
- (b) the only stock of the Pharmaceutical physically held by you or on your behalf in New Zealand is stock to which clause 6.2(a)(i) or (ii) of this Schedule applies and no agreement has been reached with the relevant DHB Hospital in terms of clause 6.2(a) of this Schedule;
- (c) you fail, directly or indirectly, to ensure that all orders for the Pharmaceutical are filled (without restricting quantities that may be ordered), including in particular where not all patients for whom the Pharmaceutical is prescribed receive the full amount of the Pharmaceutical they require, or to which they are entitled, under their prescriptions, without delay;
- (d) you fail to supply the Pharmaceutical on and from the Start Date.

7.5 **Default interest and recovery costs**

If payment of any amount required to be paid by you under this clause 7 is not made by you, in full, by the due date for payment of that amount as notified to you in writing by PHARMAC, then:

- (a) interest will accrue in such sum as remains unpaid at a rate per annum equal to the relevant SME overdraft rate (weighted average rate) of the Reserve Bank of New Zealand plus five percentage points, calculated and compounded on a daily basis, from the due date until such time as the sum due (including interest) is paid in full. This obligation to pay default interest is to arise without the need for a notice or demand from PHARMAC for such default interest; and
- (b) PHARMAC may take any action, including legal action, without first needing to implement the dispute resolution procedure contained in clause 4.2 of Schedule Four, to recover that amount and you agree to pay to PHARMAC actual enforcement costs incurred in relation to that action.

8. Termination and restrictions

8.1 **Termination and restrictions for clinical reasons**

PHARMAC reserves the right, but only after consultation with you and a relevant medical adviser (being either the Ministry of Health, PTAC or a sub-committee of PTAC), to:

- (a) terminate this Agreement at any time during the Hospital Supply Status Period or the First Transition Period if the medical adviser determines for clinical reasons that it is no longer appropriate to have:
 - (i) in the case of any Pharmaceutical that is not a Medical Device, any Pharmaceutical, including the Pharmaceutical or any relevant Alternative Pharmaceutical, having Hospital Supply Status of that form and strength of the Pharmaceutical with Hospital Supply Status;
 - (ii) in the case of any Pharmaceutical that is a Medical Device, any Pharmaceutical, including the Pharmaceutical or any relevant Alternative Pharmaceutical, having Hospital Supply Status; or
 - (iii) the Pharmaceutical as the brand having Hospital Supply Status; and/or
- (b) impose at any time during the Hospital Supply Status Period or the Transition Periods restrictions on the prescribing or dispensing of a Pharmaceutical if those restrictions are necessary for clinical reasons.

8.2 **Termination following an audit**

PHARMAC may terminate the Agreement, or withdraw Hospital Supply Status in relation to, or revise DV Limits for, a Pharmaceutical, at any time during the Hospital Supply Status Period or the Transition Periods, if you fail to remedy any area of non-compliance in accordance with clause 3(b) of Schedule Four.

9. Guarantee

- (a) PHARMAC may require an entity acceptable to it to provide a guarantee (in a form satisfactory to PHARMAC) of your performance obligations under clauses 7.2 and 7.3 of this Schedule including, without limitation, the payment of any sum payable under the indemnity or as liquidated damages pursuant to those clauses for any failure to supply the Pharmaceutical in accordance with this Agreement during the Hospital Supply Status Period.
- (b) The guarantor's liability under such a guarantee will be limited to a total of \$100,000 per Pharmaceutical for all claims made by PHARMAC under the guarantee.

10. Access by PHARMAC to price and volume data

- (a) You acknowledge that PHARMAC and its agents will require access to price and volume data held by you and DHB Hospitals in respect of the Pharmaceutical covered by this Agreement to assist PHARMAC to carry out its statutory function in relation to managing the purchasing of hospital pharmaceuticals on behalf of DHBs.
- (b) Notwithstanding any other provisions in this Agreement, including clauses 9.1 and 9.2 of Schedule Three regarding confidential information, you agree that where the

circumstances in this clause apply, a DHB Hospital may provide PHARMAC and its agents with any price and volume data held by that DHB Hospital in respect of a Pharmaceutical covered by this Agreement and PHARMAC and its agents may provide such data on DHBs.

(c) You agree that within 10 business days following any request from PHARMAC, you will provide PHARMAC with volume data in respect of the Pharmaceutical covered by this Agreement for each month of the period specified in that request.

11. **PCTs**

11.1 Listing in Section B of the Pharmaceutical Schedule

- (a) Where the Pharmaceutical is a PCT, you acknowledge and agree that PHARMAC may list the Pharmaceutical in Section B of the Pharmaceutical Schedule:
 - (i) at a price that is equal to (or subject to your agreement, less than) the Price;
 - (ii) subject to the rules and restrictions applying to PCTs in Sections A to G of the Pharmaceutical Schedule.
- (b) If PHARMAC lists the Pharmaceutical in Section B of the Pharmaceutical Schedule pursuant to paragraph (a) above, you acknowledge and agree that:
 - such listing will be for reasons relating to claiming and will not, unless otherwise advised in writing by PHARMAC, enable you to supply the Pharmaceutical for use in the community;
 - (ii) listing of the Pharmaceutical in Section B will, at PHARMAC's option, be additional to or instead of listing in Part II of Section H;
 - (iii) references to the "listing" of the Pharmaceutical will, where applicable, be to the listing of the Pharmaceutical in Section B of the Pharmaceutical Schedule (and references to "list", "listed", "delist", "delisted", and "delisting" are to be interpreted accordingly); and
 - (iv) the standard terms of listing of the Pharmaceutical in Section B of the Pharmaceutical Schedule will, except to the extent otherwise advised in writing by PHARMAC, be the terms set out in Schedule Four and this Schedule, and for that purpose all references in Schedule Four and this Schedule to "Section H" will be deemed to be references to "Section B".
- (c) This clause confers a benefit on (and is enforceable by) DHB Hospitals in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.
- (d) Where the Pharmaceutical is a PCT, clause 7.1 of this Schedule will be deleted and replaced by the following:

7.1 Notification of Potential Out-of-Stock Event and supply of Alternative Pharmaceutical.

(a) You must notify PHARMAC in writing as soon as you have reasonable cause to believe that you will fail to supply a Pharmaceutical in accordance with this Agreement and, in any event, you must notify PHARMAC and the relevant DHB Hospitals if at any time a Potential

Out-of-Stock Event occurs, including during the Hospital Supply Status Period or the First Transition Period.

- (b) If you fail to supply a Pharmaceutical in accordance with this Agreement for more than 1 business day to any DHB Hospital, then:
 - you must use your best endeavours to procure, within what the relevant DHB Hospitals consider to be a reasonable period of time, an Alternative Pharmaceutical for supply to, at a DHB Hospital's discretion, any Designated Delivery Points, and/or Contract Manufacturers (expressly for the purpose of compounding) at the Price; and
 - (ii) if you fail to procure an Alternative Pharmaceutical at the Price in accordance with sub-clause (i) above (other than for reasons that PHARMAC reasonably considers, following discussion with you, to be wholly outside your control) then, at PHARMAC's option:
 - (A) you must pay to PHARMAC (for the benefit of PHARMAC and DHB Hospitals) any additional costs that PHARMAC incurs or that the relevant DHB Hospitals incur as a result of the purchase of the Alternative Pharmaceutical; or
 - (B) PHARMAC may implement an arrangement with another supplier to supply an Alternative Pharmaceutical (including an arrangement for backup supply), and you must pay to PHARMAC (for the benefit of PHARMAC and DHB Hospitals) any additional costs that PHARMAC incurs or that the relevant DHB Hospitals incur as a result of the purchase of the Alternative Pharmaceutical.
 - This clause confers a benefit on (and is enforceable by) DHB Hospitals in accordance with the Contract and Commercial Law Act 2017, Part 2, Subpart 1.

(c)

Schedule 7: Additional Special Terms

1. Lacosamide

In respect of the identified Tender Items in Schedule 2 (lacosamide (current access) tab 50 mg, 100 mg, 150 mg and/or 200 mg or lacosamide (widened access) tab 50 mg, 100 mg, 150 mg and/or 200 mg) ("Lacosamide Tender Items") and notwithstanding any other provision in this Agreement concerning the Sole Supply Status of the Lacosamide Tender Items, and without limiting any other actions that it may be necessary or appropriate for PHARMAC to take in response to patient safety concerns, PHARMAC may grant subsidised access, on a strictly limited basis, to an alternative brand of the relevant Lacosamide Tender Items during the Sole Supply Period to the extent that this is necessary to enable a limited number of patients to continue treatment following adverse effects or efficacy concerns in their treatment with the relevant Lacosamide Tender Items, in accordance with the following condition:

• The maximum number of patients who are treated with the alternative brand of Lacosamide Tender Items shall not exceed 30 patients per calendar year, unless otherwise agreed, during the Sole Supply Period.

For the avoidance of doubt, this clause does not apply to the lacosamide (current access) inj 10 mg per ml, 20 ml Tender Item.