

3 November 2025

Dear Supplier

INVITATION TO TENDER – SUPPLY OF PHARMACEUTICALS FOR COMMUNITY SUPPLY AND/OR HOSPITAL SUPPLY

Pharmac invites tenders for the supply of certain pharmaceuticals for community supply and/or hospital supply 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 Principal Supply Status; and
- (e) Schedule 5 sets out the additional special terms that will apply if your Tender Bid in relation to a particular pharmaceutical is awarded Principal 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 **4 pm** (New Zealand time) on **Monday**, **15 December 2025**.

If you have any enquiries about this invitation please contact the **Tender Analysts** at tender@pharmac.govt.nz.

We look forward to receiving your tender.

Yours sincerely

Adrienne Martin

Udrine Mahi

Acting, Director, Pharmaceuticals

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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;

Advertisement means any advertisement as defined in the Medicines Act 1981;

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 Schedule Four and includes, to the extent applicable, the other Schedules (including Schedule Five) and the information on the Electronic Portal comprising the Invitation:

Alternative Brand Allowance means the alternative brand allowance relating to a particular Tender Item, in relation to hospital and/or community supply, as indicated as a percentage amount of the Total Pharmaceutical Volume, in the column entitled "ABA Limit" in the list of products included in Schedule Two;

Alternative Pharmaceutical means an alternative Pharmaceutical that Pharmac has expressly agreed in writing constitutes 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 Principal Supply Status in respect of a particular Tender Item, to cover the contingency that Principal 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;

Brand Allowance Indicator means the actual percentage of Brand Allowance Pharmaceuticals subsidised in the community and/or purchased by Health NZ Hospitals relative to the Total Pharmaceutical Volume in a Relevant Period;

Brand Allowance Pharmaceuticals means an alternative supplier's brand of the Pharmaceutical. For the avoidance of doubt, a Brand Allowance Pharmaceutical shall not be interpreted to be an Alternative Pharmaceutical for the purposes of the Agreement;

Brand Compensation means the compensation payable to you in accordance with clause 3.3 of Schedule 4;

Brand Differential means the difference between the Brand Allowance Indicator and the Alternative Brand Allowance:

Business Day means a day of the week, excluding Saturday, Sunday, and national public holidays in New Zealand. A Business Day starts at 8.30 am and ends at 5.00 pm;

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, but excludes:

- (a) information regarding the Pharmaceutical that does not identify you, or that cannot reasonably be expected to identify you, and you agree that such information is not Confidential Information and that Pharmac may use and publish such information; and
- (b) information released by Pharmac in accordance with clause 9 of Schedule Three of this Invitation, and you agree that such information ceases to be Confidential Information and that Pharmac may release that information again at any time in future without consulting with you or obtaining your prior agreement;

Consent means registrations, consents, permits, licences and authorisations, whether statutory or otherwise;

Crown Direction means any direction given to Pharmac under statutory authority;

Data Sheet means the Pharmaceutical data sheet published by Medsafe on your behalf;

Deadline means 4 pm, Monday 15 December 2025 (New Zealand time);

Default Interest Rate means the base rate of ASB Bank Limited plus 5% per annum;

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);

Eligible Volume means the Volume Multiplier multiplied by the Brand Differential, being a volume of Pharmaceuticals eligible for Brand Compensation in Units of that Tender Item;

End Date means the last day of the Principal Supply Period;

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

Final Transition Period means, in respect of a Pharmaceutical with Principal Supply Status, 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 Principal 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);

Force Majeure Event means any cause preventing the affected party from performing any or all of its obligations under this Agreement which arises from or is attributable to acts, events, omissions or accidents beyond the reasonable control of the affected party, which:

(a) was not reasonably foreseeable;

- (b) could not have been avoided or mitigated through the exercise of good industry practice and due care, skill and diligence; and
- (c) was not caused by the affected party, its affiliates, officers, Personnel or suppliers,

but does not include any lack of finance or financial means or any changes in market conditions;

Health NZ means Health New Zealand I Te Whatu Ora, a Crown agent established under section 11 of the Pae Ora (Healthy Futures) Act 2022;

Health NZ Hospital means Health NZ, including its hospital or associated provider unit for which Health NZ purchases pharmaceuticals;

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

In-Use Shelf-Life means the length of time that the Pharmaceutical has been demonstrated to stay effective and safe to use after the packaging of the original container is opened and stored under defined conditions;

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 2.3 of Schedule Four;

Market Approval means regulatory approval for sale and marketing in New Zealand;

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;

Medsafe means the business unit within the Ministry of Health that has responsibilities in relation to the safety of medicines and medical devices used and supplied in New Zealand;

New Zealand Government's Supplier Code of Conduct means the New Zealand Government's supplier code of conduct (as updated from time to time);

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

OPPs means Pharmac's Operating Policies and Procedures (as updated from time to time);

PCT means a Pharmaceutical for which Health NZ Hospitals are eligible to claim a subsidy through the Pharmaceutical Schedule. Tender Items that are PCTs are indicated with "PCT" in the list in clause 2 of Schedule Two and the Electronic Portal:

Personnel means all individuals engaged by the relevant party, including the parties' employees, contractors, representatives, legal advisors, clinical advisors and other consultants;

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

Pharmaceutical Schedule means the schedule listing all the medicines and medical devices funded for New Zealanders (as updated from time to time);

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

- (a) in relation to community supply, any wholesalers and other such distributors, and at which the Pharmaceutical is to be subsidised, 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, Health NZ Hospitals, any wholesalers and other such distributors, 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;

Principal Supplier means a supplier which has had a Tender Bid accepted for a Tender Item in relation to community and/or hospital supply, being the principal supplier of the relevant Tender Item (subject to the Alternative Brand Allowance provisions);

Principal Supply Period means the period beginning on the day after the expiry of the First Transition Period and ending on 30 June 2029;

Principal Supply Status means the status of being the Principal Supplier for community and/or hospital supply of a Pharmaceutical for the Principal Supply Period;

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;

Relevant Period means the periods:

- (a) beginning on the day after the expiry of the First Transition Period and ending on 30 June 2027;
- (b) 1 July 2027 until 30 June 2028; and
- (c) 1 July 2028 until 30 June 2029.

Shelf-Life means the length of time that the Pharmaceutical has been demonstrated to stay effective and safe to use when packaged in the original container and stored under defined conditions;

Special Authority (or SA) means a designation in relation to a Pharmaceutical which means that the Pharmaceutical is only eligible for subsidy or additional subsidy on approval

of an application for a named person which meets the criteria specified in the Pharmaceutical Schedule:

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
- 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;

Supply Issue means an event which may result, or has resulted, in a failure to supply the Pharmaceutical in accordance with this Agreement, including but not limited to:

- (a) your stock of the Pharmaceutical held by you in New Zealand falls below the minimum stock holding recorded in clause 5.1 of Schedule Four;
- (b) you recall (or have reason to believe you may recall), or are (or have reason to believe you may be) required by Medsafe or any other authorities to recall, the Pharmaceutical:
- (c) any Consent or Market Approval, required in accordance with clause 6.2 of Schedule Four is withdrawn, revoked, suspended or withheld;
- (d) you become aware of any issue that may impact on your ability to fulfil any orders for the Pharmaceutical;
- (e) you plan to withdraw the Pharmaceutical from supply; and/or
- (f) you fail to supply (or have reason to believe you may fail to supply) the Pharmaceutical from the Start Date;

Supply Issues Report means a report provided by you to Pharmac in accordance with clause 4.2 of Schedule Four of this Agreement;

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 electronic form in which you must enter and submit your bid(s) for each Tender Item, as set out in the Electronic Portal;

Total Brand Allowance Pharmaceutical Volume means the total volume of Brand Allowance Pharmaceuticals subsidised in the community and/or purchased by Health NZ Hospitals in a Relevant Period, specified in Units of that Tender Item;

Total Pharmaceutical Volume means the total volume of the Pharmaceutical (inclusive of Brand Allowance Pharmaceuticals) subsidised in the community and/or purchased by Health NZ Hospitals in a Relevant Period, specified in Units of that Tender Item;

Transition Periods collectively refers to the First Transition Period and the Final Transition Period:

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 the Pharmaceutical Schedule for the hospital setting, 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 the Pharmaceutical Schedule for the community setting, 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);

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;

Unique Product Identifiers means for each Pharmaceutical:

- (a) the 'CTPP', which is the 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;
- (b) the 'GTIN' (if available), which is the Global Trade Item Number for a Pharmaceutical;
- (c) the 'Pharmacode', which is the unique identifier assigned to a Pharmaceutical and notified to you by the Pharmacy Guild; and
- (d) the 'Supplier Code', which is the unique product identifier assigned by you to the Pharmaceutical, if applicable; and

Volume Multiplier means the Total Pharmaceutical Volume divided by one hundred (100) (which shall equate to 1% of the Total Pharmaceutical Volume), specified in Units of that Tender Item.

2. Interpretation

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

- (a) references to "Health NZ" encompass Health NZ Hospitals;
- (b) references to "Health NZ Hospitals" may reflect that certain operational matters can in practice occur at a local hospital level notwithstanding that Health NZ Hospitals are part of, and not separate legal entities from, Health NZ;
- (c) references to clauses are to clauses in this Invitation;
- (d) the headings to clauses will be ignored in construing this Invitation;
- (e) the plural includes the singular and vice versa;
- (f) any organisations (including government agencies) referenced in this Invitation include their successors;
- (g) a reference to any statute includes that statute, and regulations made under it, as amended from time to time:
- (h) a reference to any statute includes any statute passed in substitution for that statute;
- (i) an obligation not to do anything includes an obligation not to suffer, permit or cause that thing to be done;
- (j) derivatives of any defined word or term have a corresponding meaning;
- (k) all references to dollars are references to New Zealand dollars unless provided otherwise;
- (I) "including" and similar words do not imply any limitation;
- (m) references to "you" include any third parties acting on your behalf, including subcontractors;
- (n) references to the "listing" of a Pharmaceutical are to the listing of that
 Pharmaceutical on the Pharmaceutical Schedule (and references to "list", "listed",
 "delist", "delisted", and "delisting" are to be interpreted accordingly); and
- (o) none of the terms are to be construed against a party by reason of the fact that that term was first proposed or was drafted by that party.

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 2025.
- (b) Market value figures, in relation to community supply, are expressed as the Unit Volume in the year ending 30 June 2025 multiplied by the Unit Subsidy as at 1 July 2025.
- (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 Health NZ Hospital volumes. For the avoidance of doubt, Pharmac makes no representation as to the size of the Health NZ 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 Five. Special Authority restrictions have been noted for Tender Items where applicable in the list. Further restrictions 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 2025.
- (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 2025.

1.6 Alternative Brand Allowance

The Alternative Brand Allowance relating to a particular Tender Item, in relation to hospital and/or community supply, is indicated as a percentage amount in the column entitled "ABA Limit" in the attached list and is also shown in the Electronic Portal.

1.7 Tender Items subject to principal supply arrangements

Where a Tender Item is underlined in the list of products below, that item is subject to a principal supply contract as at the date of this Invitation. Accordingly, the subsidy for those items is fixed until 30 June 2026 (unless otherwise indicated) and, for items that are the subject of a principal 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 Principal Supply Status for hospital supply for that Tender Item.

1.9 Community only Products

Where a "C" is indicated, you may submit a Tender Bid for Principal Supply Status for community supply 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 Principal Supply Status for community supply and/or hospital supply 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 Principal Supply Status for hospital supply for that Tender Item on the basis that, if Pharmac accepts your Tender Bid, the Tender Item would be listed in the Pharmaceutical Schedule subject to clause 6.6 of Schedule Four. This information is also shown in the Electronic Portal.

Where a Tender Item is indicated as being a "PCT" product, and is in a form intended to be compounded, 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 Vial and ampoule form

Unless otherwise stated, where a Tender Item specifies either:

- (a) an ampoule; or
- (b) a vial,

form of the Chemical Entity, your brand of the relevant Chemical Entity for which you submit a bid may be in either ampoule or vial 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 ampoule and vial form of that Chemical Entity as separate line items, you must submit a bid for the same form and strength for each line item.

1.14 Pack size preference

Where a Tender Item is specified as being available for a Tender Bid for Principal Supply Status for community supply, 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; or
- (b) specified in the comments column in the attached list.

Notwithstanding the preference of Pharmac for Tender Items to be in pack sizes as specified above, 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, in the event of any conflict between the pack size preference in the comments column in the attached list and the pack size preference in this clause 1.14, the pack size preference in the comments column in the attached list will prevail.

1.15 Pack size for use in Health NZ Hospitals

Where a Tender Item is specified as being available for a Tender Bid for Principal Supply Status for hospital supply, it is the preference of Pharmac that the pack size for such a Tender Item is:

- (a) 500 ml or less, where the Tender Item is in liquid form;
- (b) 30 or 90 day pack where the Tender Item is in a tablet or capsule form; and
- (c) 10 or less injections, where the Tender Item is in injection form.

Notwithstanding the preference of Health NZ 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, Health NZ 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.

SCHEDULE TWO: PRODUCTS TO BE TENDERED									
Chemical Name Line Item	Units	Cost	Unit Subsidy	ABA Limit Comments					
ACE Inhibitor									
Oral liquid	73,900	\$63,554	\$0.8600	C H 5% Units and subsidy shown are for the currently listed captopril oral liq.					
Acitretin									
Cap 10 mg	584,090	\$255,055	\$0.4367	C H 5% Preference for a blister pack.					
Cap 25 mg	104,150	\$99,585	\$0.9562	C H 5% Preference for a blister pack.					
Adapalene (current access)									
Crm 0.1%				C H 5% Same restrictions as adapalene gel 0. would apply. Not currently listed in the community on the Pharmacetical Schedule. Pharmac would only award tender for current or widened access.					
Gel 0.1%	1,008,120	\$769,196	\$0.7630#	C H 5% Current restrictions would apply. Pharm would only award a tender for current widened access.					
Adapalene (widened access)									
Crm 0.1%				C H 5% Current maximum per-prescription limi would not apply. Not currently listed in community on the Pharmaceutical Schedule. Pharmac would only award tender for current or widened access.					
Gel 0.1%				C H 5% Current maximum per-prescription limi would not apply. Pharmac would only award a tender for current or widened access.					
Alendronate sodium									
Tab 70 mg	321,966	\$249,524	\$0.7750	C H 5%					
Alfacalcidol									
Cap 0.25 mcg	234,839	\$61,810	\$0.2632	C H 5%					
Cap 1 mcg	54,421	\$47,880	\$0.8798	C H 5%					
Oral drops 2 mcg per ml	14,540	\$44,114	\$3.0340	C H 5%					
Alfentanil Inj 0.5 mg per ml, 2 ml ampoule				@ H 5%					
Allopurinol									
<u>Tab 100 mg</u>	32,711,787	\$588,485	\$0.0180	C H 5% Preference for a scored tablet.					
<u>Tab 300 mg</u>	24,056,173	\$1,082,528	\$0.0450	C H 5% Preference for a scored tablet.					
Ambrisentan (current access)									
Tab 10 mg	22,072	\$147,147	\$6.6667	C H 5% Current restrictions apply. Pharmac wo only award a tender for current or wide access.					
Tab 5 mg	7,377	\$49,180	\$6.6667	C H 5% Current restrictions apply. Pharmac woonly award a tender for current or wide access.					
Ambrisentan (widened access) Tab 10 mg				C H 5% Same restrictions as bosentan would apply. Pharmac would only award a te for current or widened access.					
Tab 5 mg				C H 5% Same restrictions as bosentan would apply. Pharmac would only award a te for current or widened access.					
Amiloride hydrochloride with furosem	ide								
Tab 5 mg with furosemide 40 mg	111,034	\$34,222	\$0.3082	C H 5%					
Amitriptyline									
Tab 10 mg	27,088,413	\$809,944	\$0.0299	C H 5% Preference for a scored and/or dispers tablet.					
Tab 25 mg	7,980,255	\$158,807	\$0.0199	C H 5%					
Tab 50 mg	3,343,982	\$105,001	\$0.0314	C H 5%					
Amlodipine									
<u>Tab 2.5 mg</u>	19,373,721	\$312,111	\$0.0161	C H 5% Preference for a pack size of 30 or 90 tablets in a bottle pack.					
Principal Supply Status				#=rebate *=part charge @=ASP +=part					

SCHEDULE TWO: PRODUCTS TO BE TENDERED										
Chemical Name Line Item	Units	Cost	Unit Subsidy		A	ВА	Limit	Comments		
Amlodipine										
Tab 5 mg	35,114,218	\$471,935	\$0.0134		СН	5%		nce for a pack size of 30 or 90 in a bottle pack.		
<u>Tab 10 mg</u>	18,425,989	\$268,282	\$0.0146		СН	5%	Prefere	nce for a pack size of 30 or 90 in a bottle pack.		
Amorolfine										
Nail soln 5%	292,245	\$1,278,280	\$4.3740		СН	5%				
Amoxicillin										
Grans for oral liq 125 mg per 5 ml	3,826,427	\$84,947	\$0.0222	@	СН	5%	measur	nce for a product which includes a ing device and has a shelf life of a days after reconstitution.		
Grans for oral liq 250 mg per 5 ml	54,485,086	\$1,531,031	\$0.0281	@	СН	5%	measur	nce for a product which includes a ing device and has a shelf life of a days after reconstitution.		
Amoxicillin clavulanate										
Tab 500 mg with potassium clavulanate 125 mg	11,245,085	\$1,787,969	\$0.1590	@	СН	5%				
Anastrozole										
Tab 1 mg	570,341	\$83,458	\$0.1463		СН	5%				
Aripiprazole										
Tab 5 mg	1,212,046	\$424,216	\$0.3500		СН	5%				
Tab 10 mg	895,884	\$313,559	\$0.3500		СН	5%				
Tab 15 mg	325,932	\$114,076	\$0.3500		СН	5%				
Tab 20 mg	214,894	\$75,213	\$0.3500		СН	5%				
Tab 30 mg	99,107	\$34,687	\$0.3500		СН	5%				
Aspirin										
Tab dispersible or soluble 300 mg	623,360	\$35,220	\$0.0565		СН	5%				
Tab EC 100 mg	87,070,120	\$1,112,756	\$0.0128		СН	5%				
Atomoxetine										
Cap 10 mg	366,489	\$563,085	\$1.5364		СН	5%				
Cap 18 mg	128,519	\$209,165	\$1.6275		СН	5%				
Cap 25 mg	328,278	\$519,382	\$1.5821		СН	5%				
Cap 40 mg	490,051	\$808,761	\$1.6504		СН	5%				
Cap 60 mg	116,247	\$213,023	\$1.8325		СН	5%				
Cap 80 mg	92,645	\$215,730	\$2.3286		СН	5%				
<u>Cap 100 mg</u>	47,819	\$112,221	\$2.3468		СН	5%				
Atracurium besylate										
Inj 10 mg per ml, 2.5 ml				@	Н	5%				
Atropine sulphate										
Eye drops 1%	135,930	\$165,563	\$1.2180		СН	5%				
	,	* 1 2 2 , 2 2 2	¥							
Bendroflumethiazide [Bendrofluazide] Tab 2.5 mg	15,649,927	\$1,611,942	\$0.1030		СН	5%				
Tab 5 mg	2,533,207	\$309,051	\$0.1030		СН	5%				
-		# 000,001	-0.1220		٠.,	2,0				
Benzyl benzoate (funded with restriction) 10% - 25%					СН	5%	for seco	ed to scabies hyper-infestation of and line treatment where topical has failed. Not currently listed or		
Control bonzoete (funded without rectaint	ion)						would o	rmaceutical Schedule. Pharmac inly award a tender for benzyl te funded with or without restriction		
Senzyl benzoate (funded without restrict) 10% - 25%	ion)				СН	5%	Not cur	rently listed on the Pharmaceutica		
1076 - 2376					OII	J /0	Schedu tender f	le. Pharmac would only award a for benzyl benzoate funded with o restriction.		
Benzylpenicillin sodium [Penicillin G]										
Inj 600 mg	7,339	\$12,109	\$1.6500	@	СН	5%				
Principal Supply Status						#	=rebate	*=part charge @=ASP +=pate		

S	SCHEDULE TWO: PRODUCTS TO BE TENDERED								
Chemical Name Line Item	Units	Cost	Unit Subsidy		ABA	Limit	Comments		
Betahistine dihydrochloride									
Tab 16 mg	5,136,126	\$190,037	\$0.0370	СН	5%				
Betamethasone dipropionate Crm 0.05% (pack size greater than 30 g) Oint 0.05% (pack size greater than 30 g)	639,530 420,760	\$460,462 \$302,947	\$0.7200 \$0.7200	C H					
Bicalutamide									
Tab 50 mg	729,323	\$108,881	\$0.1493	СН	5%				
Bisoprolol fumarate	20, 400, 204	#200.220	CO 0454	6.1	5 0/	Duefeue			
<u>Tab 2.5 mg</u>	20,406,204	\$308,338	\$0.0151	CH	5%		nce for a scored tablet, and for a ze of 30 or 90 tablets in a bottle		
Tab 5 mg	10,097,674	\$214,273	\$0.0212	СН		tablets	nce for a pack size of 30 or 90 in a bottle pack.		
<u>Tab 10 mg</u>	3,200,080	\$96,354	\$0.0301	СН	5%		nce for a pack size of 30 or 90 in a bottle pack.		
Bupivacaine hydrochloride Inj 2.5 mg per ml, 20 ml ampoule sterile				Н	5%	Sterile	product required.		
pack Inj 5 mg per ml, 4 ml				H		Oterne	oroduct required.		
Bupivacaine hydrochloride with adrenalii	ne								
Inj 0.25% with 1:400,000 of adrenaline, 20 ml)			H					
Inj 0.5% with 1:200,000 of adrenaline, 20 ml				Н	5%				
Cabergoline									
Tab 0.5 mg (2 pack)	100,929	\$226,332	\$2.2425	CH	5%	would a	Special Authority restrictions apply. Units and cost shown are for ble tab 0.5 mg market.		
Tab 0.5 mg (8 pack)	100,929	\$226,332	\$2.2425	СН	5%	would a	Special Authority restrictions apply. Units and cost shown are for ole tab 0.5 mg market.		
Calcium carbonate									
<u>Tab</u>	9,005,285	\$262,234	\$0.0291	CH	5%	Prefere	nce for tab 1.25 g - 1.5 g.		
Calcium folinate									
Inj 350 mg				PCT F	5%	data >4 note thi on the l	nce for products where stability 8 hours post-compounding. Please s line item is also currently listed Pharmaceutical Schedule as inj 10 ml, 35 ml vial.		
Inj 50 mg		\$362,554	\$22.4400	PCT C H	5%	data >4 note thi on the I	nce for products where stability 8 hours post-compounding. Please s line item is also currently listed Pharmaceutical Schedule as inj 10 ml, 5 ml vial.		
Inj 100 mg				PCT H	5%	data >4 note thi on the I	nce for products where stability 8 hours post-compounding. Please s line item is also currently listed Pharmaceutical Schedule as inj 10 ml, 10 ml vial.		
Inj 300 mg				PCT H	5%	data >4 note thi on the I	nce for products where stability 8 hours post-compounding. Please s line item is also currently listed Pharmaceutical Schedule as inj 10 ml, 30 ml vial.		
Inj 1 g				PCT F	5%	data >4 note thi on the I	nce for products where stability 8 hours post-compounding. Please s line item is also currently listed Pharmaceutical Schedule as inj 10 ml, 100 ml vial.		
Carboprost									
Inj 250 mcg per ml, 1ml				Н	5%				

SCHEDULE TWO: PRODUCTS TO BE TENDERED										
Chemical Name Line Item	Units	Cost	Unit Subsidy		ļ	ABA	Limit	Comments		
Carvedilol			,							
Tab 6.25 mg	2,644,975	\$98,737	\$0.0373		СН	5%				
Tab 25 mg	2,394,782	\$117,751	\$0.0492		СН	5%				
Tab 12.5 mg	2,149,122	\$82,376	\$0.0383		СН	5%				
Cefazolin sodium										
Inj 500 mg	276	\$187	\$0.6780		СН	5%				
<u>lnj 1 g</u>	17,918	\$12,865	\$0.7180		СН	5%				
<u>Inj 2 g</u>	1,094	\$1,551	\$1.4180		СН	5%				
Cefotaxime										
Inj 500 mg					Н	5%				
<u>Inj 1 g</u>					Н	5%				
Ceftazidime										
<u>Inj 1 g</u>					Н	5%				
Cefuroxime sodium										
Inj 250 mg					Н	5%		rently listed on the Pharmaceutical le. Restrictions may apply.		
<u>Inj 750 mg</u>					Н	5%				
<u>Inj 1.5 g</u>					Н	5%				
Cetirizine hydrochloride										
Oral liq 1 mg per ml	21,804,879	\$435,007	\$0.0200		СН	5%				
Tab 10 mg	65,308,763	\$1,116,780	\$0.0171		СН	5%	Prefere	nce for a pack size of 90 tablets.		
Chloramphenicol										
Inj 1 g					Н	5%	Current	restrictions may apply.		
Ciclopirox olamine										
Nail lacquer 8%					СН	5%	the Pha	rently listed in the community on rmaceutical Schedule. Special restrictions may apply.		
Ciprofloxacin										
<u>Tab 250 mg</u>	95,770	\$6,669	\$0.0696		СН			nce for a scored, dispersible tablet.		
<u>Tab 500 mg</u>	658,593	\$72,913	\$0.1107		СН	5%		nce for a dispersible tablet.		
<u>Tab 750 mg</u>	21,809	\$3,739	\$0.1714		СН	5%	Prefere	nce for a dispersible tablet.		
Clarithromycin Inj 500 mg					Н	5%				
Clonidine										
TDDS 2.5 mg, 100 mcg per day	86,958	\$254,352	\$2.9250	@	СН	5%				
TDDS 5 mg, 200 mcg per day	31,694	\$101,421	\$3.2000	@	СН	5%				
TDDS 7.5 mg, 300 mcg per day	25,748	\$115,222	\$4.4750	@	СН	5%				
Clotrimazole										
Soln 1%	10,760	\$2,346	\$0.2180		СН	5%	remova product	ng this Tender Item would result in I of the part charge. 20 ml OP currently listed in the community Pharmaceutical Schedule.		
Colecalciferol										
Cap/Tab 1.25 mg	4,696,797	\$1,428,625	\$0.3042		СН	5%				
Oral liq 10 mcg per drop (400 iu per drop) 434,385	\$781,893	\$1.8000		СН	5%	Schedu	ly listed on the Pharmaceutical le as 'colecalciferol oral liq 188 r ml (7,500 iu per ml)'.		
Colestyramine										
Powder for oral liq 4 g	660,833	\$812,825	\$1.2300		СН	5%				
Colistin sulphomethate										
Inj 1,000,000 - 4,500,000 iu	450	\$9,750	\$21.6670		СН	5%				
Cyclopentolate hydrochloride										
Eye drops 1%	31,725	\$53,213	\$1.6773		СН	5%				

SCHEDULE TWO: PRODUCTS TO BE TENDERED										
Chemical Name			Unit							
Line Item	Units	Cost	Subsidy	F	ABA	Limit	Comments			
Cyclosporine										
Eye drops (strength greater than or equal to 0.05%)				СН	5%	Schedu may ap award a tenders	rently listed on the Pharmaceutical le. Special Authority restrictions ply. Pharmac reserves the right to a single tender, two separate or three separate tenders for the yclosporine market.			
Eye emulsion (strength greater than or equal to 0.05%)				СН	5%	Schedu may ap award a tenders	rently listed on the Pharmaceutical le. Special Authority restrictions ply. Pharmac reserves the right to a single tender, two separate or three separate tenders for the yclosporine market.			
Eye oint (strength greater than or equal to 0.05%)				СН	5%	Schedu may ap award a tenders	rently listed on the Pharmaceutical le. Special Authority restrictions ply. Pharmac reserves the right to a single tender, two separate or three separate tenders for the yclosporine market.			
Cyproterone acetate with ethinyloestradio										
Tab 2 mg with ethinyloestradiol 35 mcg and 7 inert tablets	7,985,544	\$241,483	\$0.0302	СН	5%					
Dapsone										
Tab 50 mg				СН	5%	Schedu may ap	rently listed on the Pharmaceutical le. Special Authority restrictions ply. Preference for a pack size of in 100 tablets.			
Tab 25 mg	136,926	\$367,646	\$2.6850	СН	5%		nce for a pack size of less than lets. Current restrictions may apply.			
Tab 100 mg	44,101	\$145,313	\$3.2950	СН	5%		nce for a pack size of less than lets. Current restrictions may apply.			
Darunavir										
<u>Tab 400 mg</u>	90,476	\$226,190	\$2.5000	СН	5%	Prefere	nce for a dispersible tablet.			
Tab 600 mg	25,670	\$96,263	\$3.7500	СН	5%	Prefere	nce for a dispersible tablet.			
Desmopressin acetate										
Nasal spray 10 mcg per dose	767,724	\$447,199	\$0.5825	СН	5%		nce for a product without ated storage requirements.			
Dexamethasone with framycetin and gram	icidin									
Ear/eye drops 500 mcg with framycetin sulphate 5 mg and gramicidin 50 mcg per ml	494,640	\$278,235	\$0.5625	СН	5%		ance of this Tender item would removal of the part charge.			
Dexmedetomidine hydrochloride										
Inj 100 mcg per ml, 2 ml vial				@ H	5%					
Diazepam										
Tab 2 mg	2,829,827	\$537,667	\$0.1900	CH	5%	Prefere	nce for a scored tablet.			
Tab 5 mg	3,989,504	\$917,586	\$0.2300	СН	5%					
Dimethicone										
Crm 10%	4,096,140	\$40,265	\$0.0098	СН	5%	Prefere	nce for a pump bottle.			
Disulfiram Tab 200 mg	420.041	¢1 016 270	¢2.2640	СН	E0/					
Tab 200 mg	429,941	\$1,016,379	\$2.3640	Сп	5%					
Docetaxel Inj 80 mg				PCT H	5%		nce for products where stability 8 hours post-compounding.			
Docusate Sodium										
Tab 50 mg	996,878	\$31,900	\$0.0320	СН	5%					
<u>Tab 120 mg</u>	1,626,043	\$80,977	\$0.0498	СН	5%					
Donepezil hydrochloride										
Tab 5 mg	1,832,211	\$80,709	\$0.0441	СН	5%					
Tab 10 mg	1,472,279	\$96,405	\$0.0655	СН	5%					
-	• •									

SCHEDULE TWO: PRODUCTS TO BE TENDERED									
Chemical Name Line Item	Units	Cost	Unit Subsidy	A	ВА	Limit (Comments		
Duloxetine									
Cap 30 mg				СН	5%		tly listed on the Pharmaceutical Special Authority restrictions		
Cap 60 mg				СН	5%	Not curren	tly listed on the Pharmaceutical Special Authority restrictions		
Econazole nitrate									
Foaming soln 1%, 10 ml sachets	42,837	\$141,219	\$3.2967	СН	5%		ee of this Tender item would emoval of the part charge.		
Efavirenz with emtricitabine and tenofovir Tab 600 mg with emtricitabine 200 mg and tenofovir disoproxil 245 mg (300 mg as a fumarate)		a fumarate) \$73,744	\$3.5627	СН	5%	only award emtricitabi fumarate)	strictions apply. Pharmac would a tender for Efavirenz with ne and tenofovir disoproxil (as a or Efavirenz with emtricitabine ovir disoproxil (as a maleate).		
Efavirenz with emtricitabine and tenofovi	disoproxil (as a	a maleate)							
Tab 600 mg with emtricitabine 200 mg and tenofovir disoproxil 245 mg (300 mg as a maleate)	d 93,165	\$331,916	\$3.5627	СН	5%	only award emtricitabi fumarate)	strictions apply. Pharmac would d a tender for Efavirenz with ne and tenofovir disoproxil (as a or Efavirenz with emtricitabine ovir disoproxil (as a maleate).		
Emulsifying ointment									
Oint (pack size 200 g or less)				Н	5%	Preference SLS-free p	e for a BP product and for an product.		
Oint (pack size greater than 200 g)	42,152,730	\$263,876	\$0.0063	СН	5%	Preference SLS-free p	e for a BP product and for an product.		
Entecavir Tab 0.5 mg	1,668,722	\$669,708	\$0.4013	СН	5%				
Ephedrine									
Inj 3 mg per ml, 10 ml prefilled syringe Inj 30 mg per ml, 1ml				Н Н	5% 5%				
Ertapenem									
Inj 1 g				Н	5%				
Escitalopram									
Tab 10 mg	18,115,986	\$511,052	\$0.0282	СН	5%	Preference	e for a scored tablet.		
Tab 20 mg	6,589,344	\$350,619	\$0.0532	СН	5%				
Esmolol hydrochloride									
Inj 10 mg per ml, 10 ml vial				Н	5%				
Exemestane									
Tab 25 mg	582,690	\$191,513	\$0.3287	СН	5%				
Ezetimibe									
Tab 10 mg	12,419,731	\$728,666	\$0.0587	СН	5%				
Famotidine Inj 10 mg per ml, 2 ml				СН	5%	the Pharm	tly listed in the community on aceutical Schedule. Special		
Inj 10 mg per ml, 4 ml	15,022	\$85,655	\$5.7020	СН	5%	Note this i	estrictions may apply. s currently listed as a Cost irce (CBS).		
Tab 20 mg	8,197,913	\$398,419	\$0.0486	СН	5%	Pharmac ı	reserves the right to award a der for either famotidine tabs or		
Tab 40 mg	1,516,475	\$155,742	\$0.1027	СН	5%	Pharmac ı	reserves the right to award a der for either famotidine tabs or		
Echuyoctot									
Febuxostat Tab 80 mg	807,635	\$136,434	\$0.1689	СН	5%	Current re	strictions may apply.		
Tab 120 mg	192,656	\$81,052	\$0.4207	СН	5%		strictions may apply.		

	SCHEDULE TWO: PRODUCTS TO BE TENDERED									
Chemical Name Line Item	Units	Cost	Unit Subsidy	ļ	ABA	Limit	Comments			
Fenofibrate			•							
Cap/tab 48 mg				СН	5%	Not cur	rently listed on the Pharmaceutical			
·							ule. Special Authority restrictions			
Cap/tab 145 mg				СН	5%		rently listed on the Pharmaceutical ule. Special Authority restrictions ply.			
Fentanyl					5 0/					
Inj 10 mcg per ml, 100 ml premix	ed bag			@ H	5%					
Fibre supplement	404.040.500	# 4 000 400	00.0440	0.11	5 0/					
Powder/granules	104,212,500	\$4,606,193	\$0.0442	СН	5%	ispaghu listed a	nd subsidy shown as per the ula (psyllium) husk 500 g currently nd expressed as 'per g'. Preference oduct containing psyllium husk.			
Finasteride										
Tab 5 mg	7,679,973	\$367,871	\$0.0479	СН	5%	Prefere	ence for a blister pack.			
Flucloxacillin sodium										
<u>Inj 250 mg</u>	90	\$383	\$4.2600	СН	5%					
<u>Inj 500 mg</u>	608	\$2,774	\$4.5630	СН	5%					
<u>lnj 1 g</u>	4,569	\$5,483	\$1.2000	СН	5%					
Fluconazole										
Inj 2 mg per ml, 50 ml vial				Н	5%	across	ated tender bids cannot be made inj and caps.			
<u>Cap 50 mg</u>	265,580	\$38,889	\$0.1464	СН	5%	Aggreg across	ated tender bids cannot be made inj and caps.			
Inj 2 mg per ml, 100 ml vial				Н		across	ated tender bids cannot be made inj and caps.			
<u>Cap 150 mg</u>	161,162	\$72,523	\$0.4500	СН	5%	across	ated tender bids cannot be made inj and caps.			
<u>Cap 200 mg</u>	83,415	\$26,514	\$0.3179	СН	5%	00 0	ated tender bids cannot be made inj and caps.			
Fludarabine phosphate										
Tab 10 mg	1,779	\$36,647	\$20.6000	PCT C H	5%	Prefere	ence for a blister pack.			
Furosemide [Frusemide]										
Tab 20 mg				СН	5%	Not cur Schedu	rently listed on the Pharmaceutical ule.			
Gemcitabine hydrochloride										
<u>lnj 1 g</u>				РСТ Н	5%	data >4	ence for products where stability 18 hours post-compounding. tions may apply.			
Glibenclamide		** * * * = * :	Ac. 27	÷						
Tab 5 mg	194,941	\$14,621	\$0.0750	СН	5%					
Gliclazide	17.0	# 000 = 55	00.0455	2	FC:	D. (
Tab 80 mg	17,077,321	\$686,508	\$0.0402	СН	5%		ence for a scored tablet, and for a ze of less than 500 tablets.			
Glucagon hydrochloride										
Inj 1 mg syringe kit	22,434	\$717,888	\$32.0000	СН	5%	Prefere syringe	ence for a graduated and/or prefilled			
Glucose [Dextrose]										
<u>Inj 50%, 10 ml - 20 ml</u>	12,229	\$84,992	\$6.9500	СН	5%	Pharma subsidy	s, 10 ml is currently listed on the acceutical Schedule. Units and y are shown per 10 ml ampoule. ence for a bag presentation.			
<u>lnj 50%, 90 ml - 100 ml</u>	386	\$6,755	\$17.5000	СН	5%	Pharma subsidy	o, 90 ml is currently listed on the acceutical Schedule. Units and are shown per 90 ml ampoule. ence for a bag presentation.			
Granisetron										
<u>Inj 1 mg per ml</u>				Н	5%					

	CHEDULE	TWO: PRO	DUCTS T	О ВЕ	TEND	ERE	D	
Chemical Name			Unit					
Line Item	Units	Cost	Subsidy		A	BA	Limit	Comments
Haloperidol								
Inj 5 mg per ml, 1 ml	148,558	\$320,142	\$2.1550	@	СН	5%		
Hydrocortisone with paraffin liquid and la	nolin							
Lotn 1% with paraffin liquid and lanolin (pack size 50 - 250 ml)	13,496,869	\$692,659	\$0.0513		СН	5%		
Hydroxyurea								
<u>Cap 500 mg</u>	1,512,041	\$313,295	\$0.2072	PC	CT C H	5%	Prefere	nce for a blister pack.
Hyoscine (Scopolamine)								
Patch 1 mg per 72 hours	51,218	\$453,279	\$8.8500		СН	5%	alternat mg per	s containing hyoscine base or ive salts with a release profile of 1 72 hours will be considered. nce for a pack size of less than 10 s.
Hyoscine N-butylbromide								
<u>lnj 20 mg, 1 ml</u>	181,358	\$69,279	\$0.3820	@	СН	5%		
Ibuprofen								
Tab 200 mg - blister pack (pack size greater than or equal to 100 and less than or equal to 1,000)	56,723,861	\$1,213,891	\$0.0214	@	СН	5%		
Tab 200 mg - blister pack (pack size less than or equal to 20)				@	Н	5%	Schedu	rently listed on the Pharmaceutical le. Preference for an over the (OTC) pack.
Imatinib mesilate	102.010	¢427.042	¢0.7400		СН	5%		
<u>Cap 100 mg</u> <u>Cap 400 mg</u>	183,010 144,362	\$137,043 \$335,689	\$0.7488 \$2.3253		СН	5% 5%		
	144,302	φ333,069	Φ Ζ.3233		CII	5%		
Indapamide	2 150 902	\$383,792	\$0.1778		СН	5%		
<u>Tab 2.5 mg</u>	2,158,803	φ303, <i>19</i> 2	φυ.1776		CII	5%		
Intra-uterine device (Non-hormonal) IUD Long	2,185	\$72,105	\$33.0000		СН	5%	situ life	nce for a product that carries an in- of 10 years. MDR certification d. Special terms apply see le 5.
IUD Medium	4,240	\$113,632	\$26.8000		СН	5%	situ life	nce for a product that carries an in- of 10 years. MDR certification d. Special terms apply see le 5.
IUD Short	4,217	\$125,667	\$29.8000		СН	5%	situ life	nce for a product that carries an in- of 10 years. MDR certification d. Special terms apply see le 5.
lodine supplement								
Tab 150 mcg elemental	13,899,474	\$925,149	\$0.0666		СН	5%		nd subsidy shown are for the y listed potassium iodate.
Irinotecan				5.0		5 0/	5 (
Inj 20 mg per ml, 5 ml				PC	ст н	5%		nce for products where stability 8 hours post-compounding.
Isosorbide mononitrate	604 400	0150 054	\$0.0040		CII	E0/		
Tab 20 mg Tab long-acting 40 mg	681,420 771,739	\$153,251 \$252,104	\$0.2249 \$0.3267		C H C H	5% 5%		
Tab long-acting 40 mg	6,260,056	\$939,008	\$0.3267 \$0.1500		СН	5% 5%	Require	ement for a scored tablet.
	0,200,000	ψ505,000	φο.1000		011	070	rtoquire	anient for a scored tablet.
Ivabradine					ш	E0/		
Tab 5 mg					Н	5%		
Ketamine Inj 100 mg per ml, 2 ml vial					СН	5%		rently listed in the Community on armaceutical Schedule. Restrictions ply.
Ketoconazole (current access)								
Shampoo 2%	9,106,200	\$294,130	\$0.0323		СН	5%		restrictions apply. Pharmac would ard a tender for current or widened
Principal Supply Status						#	=rebate	*=part charge @=ASP +=patent

S	SCHEDULE TWO: PRODUCTS TO BE TENDERED										
Chemical Name Line Item	Units	Cost	Unit Subsidy			DA I	Limit	Comments			
	Ullits	COSI	Subsidy		,	IDA I		Comments			
Ketoconazole (widened access) Shampoo 2%					СН	5%	restriction prescrip	ng access would remove the on of a maximum of 100 ml per otion. Pharmac would only award a for current or widened access.			
Lamivudine											
Tab 100 mg	55,965	\$24,105	\$0.4307		СН	5%					
<u>Tab 150 mg</u>	47,869	\$78,186	\$1.6333		СН	5%					
Latanoprost with timolol Eye drops 0.005% with timolol 0.5%	169,910	\$336,422	\$1.9800		СН	5%					
Lidocaine [lignocaine] hydrochloride witl	•	4000, .22	ψσσσσ		•	0,0					
Inj 1% with adrenaline 1:100,000, 20 ml v					Н	5%	Prefere	nce for a sterile pack.			
Inj 2% with adrenaline 1:200,000, 20 ml v	ial				Н	5%	Prefere	nce for a sterile pack.			
Lithium carbonate											
Immediate-release tab/cap	2,850,256	\$1,019,822	\$0.3578		СН	5%		nd subsidy shown are for the y listed cap 250 mg.			
Losartan	E 001 620	\$140 E19	<u></u> የሰ ሰንንያ		СН	E0/	Droforo	noo for a nook aiza of 20 or 00			
<u>Tab 12.5 mg</u>	5,901,629	\$140,518	\$0.0238		CII	J /0		nce for a pack size of 30 or 90 in a bottle pack.			
Tab 25 mg	13,699,642	\$373,452	\$0.0273		СН	5%		nce for a pack size of 30 or 90 in a bottle pack.			
Tab 50 mg	25,009,223	\$851,564	\$0.0341		СН	5%	tablets i	nce for a pack size of 30 or 90 in a bottle pack.			
<u>Tab 100 mg</u>	13,698,289	\$745,187	\$0.0544		СН	5%		nce for a pack size of 30 or 90 in a bottle pack.			
Macitentan											
Tab 10 mg					СН	5%	Schedu Pharma	rently listed on the Pharmaceutical le. Restrictions may apply. Ic would consider awarding a that is cost neutral to ambrisentan			
Macrogol 3350 with potassium chloride,	sodium bicarb	onate and sodi	um chloride								
Powder for oral soln 13.125 g with potassium chloride 46.6 mg,sodium bicarbonate 178.5 mg and sodium chlorid 350.7 mg	13,673,241 <u>e</u>	\$3,874,039	\$0.2833		СН	5%					
Macrogol 3350 with sodium sulfate, sodi	um chloride, p	otassium chlor	ide, sodium a	ascorbate	and a	scorb	ic acid				
Powder for oral soln					Н	5%					
Magnesium sulphate	00.055	\$00.500	#0.7500	@	0.11	5 0/					
Inj 2 mmol per ml, 5ml	23,855	\$89,528	\$3.7530	@	СН	5%					
Malathion (funded with restriction) 0.5%					СН	5%	for second therapy the Pha	red to scabies hyper-infestation or and line treatment where topical has failed. Not currently listed on armaceutical Schedule. Pharmac anly award a tender for malathion			
								with or without restriction.			
Malathion (funded without restriction)											
0.5%					СН	5%	Schedu	rently listed on the Pharmaceutical le. Pharmac would only award a for malathion funded with or withou on.			
Mebeverine hydrochloride <u>Tab 135 mg</u>	2,974,199	\$280,883	\$0.0944		СН	5%					
Melphalan											
<u>lnj 50 mg vial</u>				PCT	Н	5%		nce for products where stability 8 hours post-compounding.			
Meropenem											
<u>Inj 500 mg</u> <u>Inj 1 g</u>				@	H H	5% 5%					
Principal Supply Status						#:	=rebate	*=part charge @=ASP +=paten			

SC	SCHEDULE TWO: PRODUCTS TO BE TENDERED									
Chemical Name Line Item	Units	Cost	Unit Subsidy		ABA	Limit Comments				
Metaraminol tartrate										
Inj 10 mg per ml, 1 ml				@ H	5%					
Methotrexate										
Inj 100 mg per ml, 50 ml				PCT H	5%	Preference for products where stability data >48 hours post-compounding.				
Inj 100 mg per ml, 10 ml vial			\$25.0000	PCT C H	5%	Preference for products where stability data >48 hours post-compounding.				
Methylprednisolone aceponate										
<u>Crm 0.1%</u>	538,635	\$177,750	\$0.3300	СН	5%	Preference for a smaller pack size < 30 g.				
Oint 0.1%	299,685	\$98,896	\$0.3300	СН	5%	Preference for a smaller pack size < 30 g.				
Metoclopramide hydrochloride										
Tab 10 mg	6,365,549	\$99,939	\$0.0157	СН	5%					
Metoprolol succinate										
Tab long-acting Very Low Dose (20-25 mg) 23,269,108	\$1,085,969	\$0.0467	СН	5%	Units and subsidy shown are for the currently listed tab 23.75 mg.				
Tab long-acting Low Dose (40-50 mg)	32,715,279	\$1,326,932	\$0.0406	СН	5%	Units and subsidy shown are for the currently listed tab 47.5 mg.				
Tab long-acting Medium Dose (90-100 mg) 19,829,443	\$1,154,470	\$0.0582	СН	5%	Units and subsidy shown are for the currently listed tab 95 mg.				
Tab long-acting High Dose (180-200 mg)	3,575,471	\$387,724	\$0.1084	СН	5%	Units and subsidy shown are for the currently listed tab 190 mg.				
Metronidazole										
<u>Tab 200 mg</u>	301,899	\$31,228	\$0.1034	СН	5%	Preference for a scored, dispersible tablet Preference for a pack size of less than 250 tablets.				
<u>Tab 400 mg</u>	2,186,434	\$446,666	\$0.2043	СН	5%	Preference for a pack size of less than 250 tablets.				
Inj 5 mg per ml, 100 ml				@ H	5%	Preference for a bag presentation with space to add another antibiotic.				
Miconazole nitrate										
<u>Crm 2%</u>	3,527,715	\$211,663	\$0.0600	СН	5%	Preference for a 15 g - 20 g pack size.				
Midazolam										
Inj 10 mg, buccal syringe				СН	5%	Pharmac would consider funding midazolam prefilled syringes subject to funding criteria.				
Inj 2.5 mg, buccal syringe				СН	5%	Pharmac would consider funding midazolam prefilled syringes subject to funding criteria.				
Inj 5 mg, buccal syringe				СН	5%	Pharmac would consider funding midazolam prefilled syringes subject to funding criteria.				
Inj 7.5 mg, buccal syringe				СН	5%	Pharmac would consider funding midazolam prefilled syringes subject to funding criteria.				
Minocycline hydrochloride										
Inj 100 mg - 200 mg				Н	5%	Not currently listed on the Pharmaceutical Schedule. Special Authority restrictions may apply.				
Mitomycin C										
Inj 2 mg - 5 mg				PCT C H	5%	Preference for products where stability data >48 hours post-compounding.				
Morphine										
Oral liq 1 mg per ml	8,179,772	\$777,078	\$0.0950	СН	5%	Requirement for a maximum of a 200 ml pack size. Preference for a product with child-resistant packaging. Preference for an excipient free product. Aggregated tender bids cannot be made across inj and oral liq.				

S	SCHEDULE TWO: PRODUCTS TO BE TENDERED									
Chemical Name Line Item	Units	Cost	Unit Subsidy	ABA I	Limit	Comments				
Morphine										
Oral liq 10 mg per ml	43,045	\$8,663	\$0.2013	C H 5%	pack siz Preferer resistan excipien	ment for a maximum of a 200 ml re in line with legislation. nce for a product with child- t packaging. Preference for an at free product. Aggregated tender nnot be made across inj and oral				
Oral liq 2 mg per ml	259,241	\$30,526	\$0.1178	C H 5%	pack siz Preferer resistan excipien	ment for a maximum of a 200 ml te in line with legislation. Ince for a product with child- t packaging. Preference for an at free product. Aggregated tender anot be made across inj and oral				
Oral liq 5 mg per ml	278,942	\$39,331	\$0.1410	C H 5%	pack siz Preferer resistan excipien	ment for a maximum of a 200 ml te in line with legislation. Ince for a product with child- t packaging. Preference for an at free product. Aggregated tender anot be made across inj and oral				
Inj 1 mg per ml, 10 ml syringe				H 5%		ated tender bids cannot be made nj and oral liq.				
Inj 1 mg per ml, 50 ml syringe				H 5%		ated tender bids cannot be made nj and oral liq.				
<u>lnj 1 mg per ml, 100 ml bag</u>				H 5%		ated tender bids cannot be made nj and oral liq.				
Moxifloxacin										
<u>Inj 400 mg</u>				H 5%						
Tab 400 mg	20,575	\$172,826	\$8.4000	C H 5%						
Multivitamins Tab (BPC cap strength)	18,159,014	\$335,942	\$0.0185	C H 5%						
Mupirocin										
Oint 2%	565,095	\$248,642	\$0.4400	C H 5%		nnce of this Tender item would removal of the part charge.				
Naltrexone hydrochloride										
Tab 50 mg	334,931	\$930,328	\$2.7777	C H 5%						
Nifedipine	, , , , , ,	, ,	•							
Tab long-acting 10 mg	496,057	\$172,027	\$0.3468	C H 5%						
Tab long-acting 20 mg	801,995	\$142,114	\$0.1772	C H 5%						
Tab long-acting 30 mg	693,777	\$236,876	\$0.3414	C H 5%						
Tab long-acting 60 mg	128,993	\$68,121	\$0.5281	C H 5%						
Nintedanib	·									
Cap 100 mg	42,840	\$1,823,556	\$42.5667#	C H 5%	Current	restrictions apply.				
Cap 150 mg	104,640	\$6,749,280	\$64.5000#			restrictions apply.				
Nitrofurantoin	,	,				11.7				
Cap modified-release 100 mg	2,120,525	\$1,721,866	\$0.8120	C H 5%						
	_, ,,,	ψ.,,, <u>2.1,000</u>	ψ0.01 2 0	2 370						
Nortriptyline hydrochloride Tab 10 mg	17,874,346	\$439,709	\$0.0246	C H 5%						
Tab 25 mg	10,058,692	\$351,451	\$0.0240	CH 5%						
-	. 0,000,002	ψ001, 1 01	ψ0.00 N	3 370						
Nystatin Oral liq 100,000 u per ml	1,742,520	\$161,183	\$0.0925	C H 5%						
Vaginal crm 100,000 u per 5 g with applicator(s)	166,800	\$12,677	\$0.0760	C H 5%	Units an	nd subsidy expressed as 'per g'.				
Octreotide Inj 100 mcg per ml, 1 ml prefilled syringe				C H 5%	Schedul	rently listed on the Pharmaceutical le. Special Authority restrictions				
					may app	ory.				

	SCHEDULE	TWO: PRO	DUCTS TO	BE TENDER	ΕD	
Chemical Name Line Item	Units	Cost	Unit Subsidy	ABA	Limit	Comments
Octreotide						
Inj 50 mcg per ml, 1 ml prefilled syringe				C H 5%		rently listed on the Pharmaceutical le. Special Authority restrictions ply.
Inj 500 mcg per ml, 1 ml prefilled syringe				C H 5%		rently listed on the Pharmaceutical le. Special Authority restrictions ply.
Inj 50 mcg per ml, 1 ml	1,414	\$7,800	\$5.5160	C H 5%		
Inj 100 mcg per ml, 1 ml	5,014	\$48,636	\$9.7000	C H 5%		
Inj 500 mcg per ml, 1 ml	1,901	\$43,001	\$22.6200	C H 5%		
Olanzapine						
Orodispersible tab 5 mg	543,405	\$46,966	\$0.0864	C H 5%		ated tender bids cannot be made orodispersible tabs and regular
Orodispersible tab 10 mg	710,752	\$73,357	\$0.1032	C H 5%		ated tender bids cannot be made orodispersible tabs and regular
<u>Tab 2.5 mg</u>	2,409,841	\$112,467	\$0.0467	C H 5%	tablets.	nce for a pack size of 30 or 90 Aggregated tender bids cannot be cross orodispersible tabs and tabs.
<u>Tab 5 mg</u>	3,062,490	\$197,010	\$0.0643	C H 5%	tablets.	nce for a pack size of 30 or 90 Aggregated tender bids cannot be cross orodispersible tabs and tabs.
Tab 10 mg	3,343,443	\$215,084	\$0.0643	C H 5%	tablets.	nce for a pack size of 30 or 90 Aggregated tender bids cannot be cross orodispersible tabs and tabs.
Omeprazole						
Inf 40 mg				H 5%	Pharma Aggrega	ly listed in the hospital on the aceutical Schedule as Inj 40 mg. ated tender bids cannot be made inf, inj and caps.
Inj 40 mg	7,992	\$59,748	\$7.4760	C H 5%	Schedu diluent.	ly listed on the Pharmaceutical le as Inj 40 mg ampoule with Aggregated tender bids cannot be cross inf, inj and caps.
<u>Cap 10 mg</u>	8,804,246	\$201,529	\$0.0229	C H 5%	capsule	nce for a pack size of 30 or 90 is in a bottle pack. Aggregated bids cannot be made across inf, injust.
<u>Cap 20 mg</u>	120,126,898	\$2,695,648	\$0.0224	C H 5%	capsule	nce for a pack size of 30 or 90 s in a bottle pack. Aggregated bids cannot be made across inf, injust.
Cap 40 mg	35,545,039	\$1,255,806	\$0.0353	C H 5%	capsule	nce for a pack size of 30 or 90 es in a bottle pack. Aggregated bids cannot be made across inf, injus.
Ondansetron						
Tab disp 4 mg	4,181,034	\$234,138	\$0.0560	C H 5%		
Tab disp 8 mg	1,468,193	\$132,137	\$0.0900	C H 5%		
Oxybutynin						
Tab 5 mg	4,832,908	\$261,944	\$0.0542	C H 5%		
Oxycodone hydrochloride						
Tab immediate-release 10 mg	1,028,862	\$193,117	\$0.1877	C H 5%		nce for a product with child- it packaging.
Tab immediate-release 20 mg	272,753	\$73,016	\$0.2677	C H 5%	Prefere	nce for a product with child- it packaging.
Tab immediate-release 5 mg	2,587,814	\$356,342	\$0.1377	C H 5%		nce for a product with child- it packaging.

SCHEDULE TWO: PRODUCTS TO BE TENDERED								
Chemical Name			Unit					
Line Item	Units	Cost	Subsidy			ABA	Limit Comments	
Paclitaxel								
<u>Inj 100 mg</u>				PC	т н	5%	Preference for products where stability data >48 hours post-compounding.	
Inj 300 mg				PC	т н	5%	Preference for products where stability data >48 hours post-compounding.	
Paracetamol								
Tab 500 mg - blister pack (pack size greater than or equal to 100 and less the or equal to 1,000)	403,539,681 nan_	\$7,969,909	\$0.0198	@	СН	5%		
Tab 500 mg - bottle pack (pack size greater than or equal to 1,000)	57,347,382	\$1,027,665	\$0.0179	@	СН	5%		
Suppos 125 mg	172,015	\$73,794	\$0.4290		СН	5%	Preference for a product that can be stored at room temperature.	
Suppos 250 mg	111,587	\$60,145	\$0.5390		СН	5%	·	
Suppos 500 mg	123,711	\$40,948	\$0.3310		СН	5%	'	
Paraffin		4.	A			_		
White soft (pack size 2000 g or more) White soft (pack size 500 g or less)	1,979,122 128,146	\$15,041 \$1,350	\$0.0076 \$0.0105		C C H	5% 5%	Only funded in combination.	
Pentamidine	-, -	* ,	*****					
Inj 300 mg					Н	5%		
Perindopril with amlodipine								
Tab 10 mg with amlodipine 10 mg					СН	5%	Not currently listed on the Pharmaceutical Schedule. Pharmac would consider funding perindopril with amlodipine if cost saving or cost neutral to current combined cost of funded ACE inhibitor and calcium channel blocker.	
Tab 10 mg with amlodipine 5 mg					CH	5%	Not currently listed on the Pharmaceutical Schedule. Pharmac would consider funding perindopril with amlodipine if cost saving or cost neutral to current combined cost of funded ACE inhibitor and calcium channel blocker.	
Tab 5 mg with amlodipine 10 mg					СН	5%	Not currently listed on the Pharmaceutical Schedule. Preference for a scored tablet. Pharmac would consider funding perindopril with amlodipine if cost saving or cost neutral to current combined cost of funded ACE inhibitor and calcium channel blocker.	
Tab 5 mg with amlodipine 5 mg					СН	5%	Not currently listed on the Pharmaceutical Schedule. Preference for a scored tablet. Pharmac would consider funding perindopril with amlodipine if cost saving or cost neutral to current combined cost of funded ACE inhibitor and calcium channel blocker.	
Permethrin (combined market)								
Crm 5%					СН	5%	Not currently listed on the Pharmaceutical Schedule. Pharmac reserves the right to award a single tender for the whole permethrin market or two separate tenders for cream and lotion.	
Lotn 5%	6,269,370	\$894,451	\$0.1427		СН	5%	Pharmac reserves the right to award a single tender for the whole permethrin market or two separate tenders for cream and lotion.	
Permethrin (split market)								
Crm 5%					СН	5%	Not currently listed on the Pharmaceutical Schedule. Pharmac reserves the right to award a single tender for the entire permethrin market or two separate tenders for cream and lotion.	

S	CHEDULE "	TWO: PRO	DUCTS TO	BE TENDER	RED	
Chemical Name Line Item	Units	Cost	Unit Subsidy	AB	A Limit	Comments
Permethrin (split market)						
Lotn 5%	6,269,370	\$894,451	\$0.1427	C H 59	single te	c reserves the right to award a ender for the entire permethrin or two separate tenders for cream on.
Pimecrolimus (current access) Crm 1%	183,795	\$404,349	\$2.2000	C H 59		restrictions apply. Pharmac would ard a tender for current or widened
Pimecrolimus (widened access) Crm 1%			\$2.2000	C H 59	requiren eyelid. F	ng access would remove the nent for atopic dermatitis of the Pharmac would only award a or current or widened access.
Pine tar with triethanolamine lauryl sulph						
Soln 2.3% with triethanolamine lauryl sulphate and fluorescein sodium	23,290,907	\$252,008	\$0.0108	C H 59	%	
Poloxamer						
Oral drops 10%	268,470	\$37,317	\$0.1390	C H 59	%	
Potassium chloride Inj 75 mg (1 mmol) per ml, 10 ml ampoule	e 11,246	\$14,620	\$1.3000	C H 59	%	
Prasugrel						
Tab 10 mg				C H 59	Schedu	rently listed on the Pharmaceutical le. Special Authority restrictions oly. Preference for a scored tablet.
Tab 5 mg				C H 59	% Not curr	rently listed on the Pharmaceutical le. Special Authority restrictions
Pravastatin						
Tab 20 mg	3,359,763	\$240,559	\$0.0716	C H 59	%	
Tab 40 mg	1,979,080	\$242,437	\$0.1225	C H 59	%	
Prazosin Hydrochloride						
Tab 1 mg	52,861	\$2,923	\$0.0553	C H 59	%	
Tab 2 mg	34,134	\$2,389	\$0.0700	C H 59		
Tab 5 mg	10,298	\$1,205	\$0.1170	C H 59	%	
Prochlorperazine						
Tab 5 mg	3,944,251	\$394,425	\$0.1000	C H 59	%	
Propranolol						
Oral liq 4 mg per ml	276,193			C H 59		s is currently listed as a Cost cource (CBS). Units are expressed ml'
Propylthiouracil						
Tab 50 mg	282,344	\$98,820	\$0.3500	CH 59	%	
Pyridostigmine bromide						
Tab 60 mg	706,519	\$355,238	\$0.5028	C H 59	%	
Pyridoxine hydrochloride						
Tab 25 mg	1,691,202	\$64,452	\$0.0381	C H 59	%	
Quetiapine						
Tab 25 mg	32,200,039	\$844,285	\$0.0262	C H 59		nce for a scored tablet. Preference ck size of 30 or 90 tablets in a ack.
<u>Tab 100 mg</u>	4,340,829	\$308,676	\$0.0711	C H 59		nce for a pack size of 30 or 90 n a blister pack.
<u>Tab 200 mg</u>	1,137,274	\$138,622	\$0.1219	C H 59		nce for a pack size of 30 or 90 n a blister pack.
<u>Tab 300 mg</u>	492,397	\$86,608	\$0.1759	C H 59		nce for a pack size of 30 or 90 n a blister pack.

SC	HEDULE T	WO: PRO	DUCTS T	O BE	TEND	ERE	D	
Chemical Name Line Item	Units	Cost	Unit Subsidy		ļ	ABA I	Limit	Comments
Ranitidine								
Tab 150 mg					СН	5%	the Pha	rently listed in the community on armaceutical Schedule. Pharmac is the right to award a single tender famotidine tabs or ranitidine tabs.
Tab 300 mg					СН	5%	the Pha	rently listed in the community on irmaceutical Schedule. Pharmac is the right to award a single tender ir famotidine tabs or ranitidine tabs.
Remifentanil hydrochloride								
<u>Inj 1 mg</u>				@	Н	5%		
<u>Inj 2 mg</u>				@	Н	5%		
Rifampicin								
<u>Cap 150 mg</u>	70,327	\$41,169	\$0.5854	@	СН	5%		ated tender bids cannot be made caps, inj and oral liq.
<u>Cap 300 mg</u>	227,849	\$278,112	\$1.2206	@	СН	5%		ated tender bids cannot be made caps, inj and oral liq.
<u>Inj 600 mg</u>					Н	5%		ated tender bids cannot be made caps, inj and oral liq.
Oral liq 100 mg per 5 ml	74,225	\$15,587	\$0.2100	@	СН	5%	00 0	ated tender bids cannot be made caps, inj and oral liq.
Rifampicin with isoniazid and ethambutol	and pyrazinami	ide						
Tab/Cap 150 mg with isoniazid 75 mg and ethambutol 275 mg and pyrazinamide 400 mg					СН	5%	Schedu Pharma	rently listed on the Pharmaceutical le. Restrictions may apply. ac would award a tender for any mulation of this combination.
Rifampicin with isoniazid and pyrazinamid	e							
Tab/Cap 150 mg with isoniazid 75 mg and pyrazinamide 400 mg					СН	5%	Schedu Pharma	rently listed on the Pharmaceutical le. Restrictions may apply. ac would award a tender for any mulation of this combination.
Risperidone								
Oral liq 1 mg per ml	520,356	\$178,482	\$0.3430		СН	5%		
<u>Tab 0.5 mg</u>	4,181,113	\$151,231	\$0.0362		СН	5%	Prefere a blister	nce for a pack size of 60 tablets in r pack.
Tab 1 mg	2,188,093	\$88,990	\$0.0407		СН	5%	Prefere a blister	nce for a pack size of 60 tablets in r pack.
Tab 2 mg	1,039,545	\$47,123	\$0.0453		СН	5%	Prefere a blister	nce for a pack size of 60 tablets in r pack.
Tab 3 mg	426,149	\$31,961	\$0.0750		СН	5%	Prefere a blister	nce for a pack size of 60 tablets in r pack.
<u>Tab 4 mg</u>	206,952	\$21,558	\$0.1042		СН	5%	Prefere a blister	nce for a pack size of 60 tablets in r pack.
Rizatriptan Tab orodispersible 10 mg	2,863,421	\$461,956	\$0.1613		СН	5%		
Ropivacaine hydrochloride								
Inj 2 mg per ml, 10 ml ampoule					Н	5%		
Inj 2 mg per ml, 20 ml					н	5%		
Inj 2 mg per ml, 100 ml					 H	5%		
Inj 2 mg per ml, 200 ml					н	5%		
Inj 7.5 mg per ml, 10 ml					'' H	5%		
Inj 7.5 mg per ml, 20 ml					 H	5%		
Inj 10 mg per ml, 10 ml					'' H	5%		
Inj 10 mg per ml, 20 ml					Н	5%		
Rosuvastatin (current access)								
Tab 10 mg	4,289,049	\$241,602	\$0.0563		СН	5%	would o	restriction would apply. Pharmac only award a tender for current or d access.
Tab 20 mg	5,689,288	\$513,913	\$0.0903		СН	5%	Current would o	restriction would apply. Pharmac only award a tender for current or discress.
Principal Supply Status						#:		*=part charge @=ASP +=patent

;	SCHEDULE 1	TWO: PRO	DUCTS T	O BE	ΓEND	ERE	D	
Chemical Name Line Item	Units	Cost	Unit Subsidy		,	ABA	Limit	Comments
Rosuvastatin (current access)								
Tab 40 mg	5,746,741	\$871,608	\$0.1517		СН	5%	would o	restriction would apply. Pharmac nly award a tender for current or daccess.
<u>Tab 5 mg</u>	2,141,021	\$92,064	\$0.0430		СН	5%	would o	restriction would apply. Pharmac nly award a tender for current or d access.
Rosuvastatin (widened access)								
Tab 10 mg					СН	5%	would o	ions would not apply. Pharmac nly award a tender for current or d access.
Tab 20 mg					СН	5%	would o	ions would not apply. Pharmac nly award a tender for current or d access.
Tab 40 mg					СН	5%	would o	ions would not apply. Pharmac nly award a tender for current or d access.
Tab 5 mg					СН	5%	would o	ions would not apply. Pharmac nly award a tender for current or d access.
Rotigotine								
Transdermal patch 2 mg per 24 hours					СН	5%		rently listed on the Pharmaceutical le. Special Authority restrictions ply.
Transdermal patch 4 mg per 24 hours					СН	5%		rently listed on the Pharmaceutical le. Special Authority restrictions ply.
Transdermal patch 6 mg per 24 hours					СН	5%		rently listed on the Pharmaceutical le. Special Authority restrictions ply.
Transdermal patch 8 mg per 24 hours					СН	5%		rently listed on the Pharmaceutical le. Special Authority restrictions ply.
Roxithromycin								
Tab 150 mg	1,388,947	\$366,404	\$0.2638	@	СН	5%		
Tab 300 mg	746,796	\$373,398	\$0.5000	@	СН	5%		
Sapropterin dihydrochloride (current ac	cess)							
Tab soluble 100 mg	14,760	\$714,728	\$48.4233#		СН	5%	would o	restrictions would apply. Pharmac nly award a tender for current or d access.
Sapropterin dihydrochloride (widened a	iccess)							
Tab soluble 100 mg			\$48.4233#		СН	5%	disorde supplen would n	ple with inherited metabolic rs requiring dietary nentation. Current restrictions ot apply. Pharmac would only a tender for current or widened
Selexipag								
Tab 1,000 mcg					СН	5%	Schedu	rently listed on the Pharmaceutical le. Same restrictions as iloprost er soln would apply.
Tab 1,200 mcg					СН	5%	Schedu	rently listed on the Pharmaceutical le. Same restrictions as iloprost er soln would apply.
Tab 1,400 mcg					СН	5%	Schedu	rently listed on the Pharmaceutical le. Same restrictions as iloprost er soln would apply.
Tab 1,600 mcg					СН	5%	Schedu	rently listed on the Pharmaceutical le. Same restrictions as iloprost er soln would apply.
Tab 200 mcg					СН	5%	Schedu	rently listed on the Pharmaceutical le. Same restrictions as iloprost er soln would apply.

SC	CHEDULE	TWO: PRO	DUCTS T	O BE TEN	DERE	ED	
Chemical Name Line Item	Units	Cost	Unit Subsidy		ABA	Limit	Comments
Selexipag							
Tab 400 mcg				CH	l 5%	Schedu	rently listed on the Pharmaceutical ile. Same restrictions as iloprost er soln would apply.
Tab 600 mcg				C F	I 5%	Not cur Schedu	rently listed on the Pharmaceutical ile. Same restrictions as iloprost er soln would apply.
Tab 800 mcg				CH	I 5%	Not cur Schedu	rently listed on the Pharmaceutical ile. Same restrictions as iloprost er soln would apply.
Simvastatin							
Tab 10 mg	2,272,908	\$42,435	\$0.0187	CH	I 5%		nce for a pack size of 30 or 90 in a bottle pack.
Tab 20 mg	9,261,033	\$261,346	\$0.0282	C H	I 5%	Prefere	nce for a pack size of 30 or 90 in a bottle pack.
Tab 40 mg	7,977,931	\$364,352	\$0.0457	C F	I 5%	Prefere	nce for a pack size of 30 or 90 in a bottle pack.
Tab 80 mg	559,410	\$54,761	\$0.0979	CH	l 5%	Prefere	nce for a pack size of 30 or 90 in a bottle pack.
Sodium nitroprusside						iabiets	п а роше раск.
Inj 50 mg vial				F	l 5%		
Sodium phosphate with phosphoric acid							
Enema 10% with phosphoric acid 6.58%	73,571	\$272,213	\$3.7000	CH	l 5%	Commu Schedu	note this is currently listed in the unity on the Pharmaceutical ale as 'Sodium acid phosphate - 16% with sodium phosphate 8%'.
Sulfadiazine silver							
1%	344,300	\$74,369	\$0.2160	C H	I 5%		restrictions apply. Pharmac would a tender for any topical formulation.
Suxamethonium chloride							
<u>lnj 50 mg per ml, 2 ml</u>				@ F	l 5%	Prefere stability	nce for a product with 30 to 60 day
Inj 50 mg per ml, 2 ml prefilled syringe				F	l 5%		d syringe not currently listed on the aceutical Schedule.
Tacrolimus (current access)							
Oint 0.1%	305,520	\$336,072	\$1.1000	C H	I 5%		restrictions apply. Pharmac would yard a tender for current or widened
Tacrolimus (widened access)			¢4 4000	C.1	L 50/	\\/;doni	an access would remove the
Oint 0.1%			\$1.1000	CH	I 5%	require	ng access would remove the ment for atopic dermatitis of the harmac would only award a tender ent or widened access.
Tadalafil							
Tab 10 mg				CF	l 5%	would a Pharma conside	estrictions as sildenafil tablets apply. Not currently listed on the accutical Schedule. Pharmac would be funding tadalafil if cost saving or utral to sildenafil tablets.
Tab 20 mg				CH	I 5%	would a Pharma conside	estrictions as sildenafil tablets apply. Not currently listed on the accutical Schedule. Pharmac would be funding tadalafil if cost saving or utral to sildenafil tablets.
Tafamidis							
Cap 20 mg				CH	I 5%		rently listed on the Pharmaceutical ile. Special Authority restrictions ply.
Cap 61 mg				C F	I 5%	Not cur Schedu	rently listed on the Pharmaceutical lle. Special Authority restrictions
						may ap	μιy.
Tamoxifen citrate	E0 007	¢4.4.700	\$0.2500	0.1	E0/		
Tab 10 mg Principal Supply Status	58,837	\$14,709	\$0.2500	C F		t=rebate	*=part charge @=ASP +=patent
· · · · · · · · · · · · · · · · · · ·							. •

	SCHEDULE T	TWO: PRO	DUCTS TO	D BE TEND	ERE	D	
Chemical Name Line Item	Units	Cost	Unit Subsidy	Į.	ABA I	Limit	Comments
Tamoxifen citrate			•				
Tab 20 mg	1,634,433	\$144,925	\$0.0887	СН	5%		
-	1,001,100	Ψ111,020	φο.σσοι	.	070		
Telmisartan Tab 40 mg				СН	5%	Schedu funding	rently listed on the Pharmaceutical lle. Pharmac would consider telmisartan if cost saving or cost to a current angiotensin II nist.
Tab 80 mg				СН	5%	Schedu funding	rently listed on the Pharmaceutical le. Pharmac would consider telmisartan if cost saving or cost to a current angiotensin II nist.
Telmisartan with amlodipine							
Tab 40 mg with amlodipine 10 mg				СН	5%	Schedu funding saving cost of	rently listed on the Pharmaceutical ale. Pharmac would consider telmisartan with amlodipine if cost or cost neutral to current combined funded angiotensin II antagonist cium channel blocker.
Tab 40 mg with amlodipine 5 mg				СН	5%	Schedu funding saving cost of	rently listed on the Pharmaceutical ale. Pharmac would consider telmisartan with amlodipine if cost or cost neutral to current combined funded angiotensin II antagonist cium channel blocker.
Tab 80 mg with amlodipine 10 mg				СН	5%	Schedu funding saving cost of	rently listed on the Pharmaceutical lle. Pharmac would consider telmisartan with amlodipine if cost or cost neutral to current combined funded angiotensin II antagonist cium channel blocker.
Tab 80 mg with amlodipine 5 mg				СН	5%	Schedu funding saving cost of	rently listed on the Pharmaceutical ule. Pharmac would consider telmisartan with amlodipine if cost or cost neutral to current combined funded angiotensin II antagonist cium channel blocker.
Temazepam							
Tab 10 mg	2,994,418	\$167,687	\$0.0560	C H	5%	Prefere	nce for a scored tablet.
Terbinafine							
<u>Tab 250 mg</u>	2,470,021	\$263,774	\$0.1068	СН	5%		
Thiamine hydrochloride							
Inj 100 mg per ml, 1 ml				н	5%	single t	ac reserves the right to award a ender for the whole thiamine injusted per ml market, or two separate for the 1 ml and 2 ml tations.
Inj 100 mg per ml, 2 ml				Н	5%	single t	ac reserves the right to award a ender for the whole thiamine inj per ml market, or two separate for the 1 ml and 2 ml tations.
Thiotepa Inj 15 mg				PCT H	5%		nce for products where stability 8 hours post-compounding.
<u>Inj 100 mg</u>				PCT H	5%	Prefere	nce for products where stability 18 hours post-compounding.
Timolol maleate							
Eye drops 0.25%	176,405	\$85,380	\$0.4840	СН	5%	For pro Pharma preserv market	ence for a preservative free product. ducts containing preservatives, ac reserves the right to list a rative free product for a restricted which shall not be subject to the lote this line item is for gel-forming

	CHEDULE	TWO: PRO	DUCTS TO	BE TENDER	ĒD	
Chemical Name Line Item	Units	Cost	Unit Subsidy	ABA	Limit	Comments
Timolol maleate						
Eye drops 0.5%	200,755	\$100,378	\$0.5000	C H 5%	For pro- Pharma preserv market	nce for a preservative free product. ducts containing preservatives, ac reserves the right to list a ative free product for a restricted which shall not be subject to the ote this line item is for gel-forming
Tobramycin						
Solution for inhalation 60 mg per ml, 5 ml	4,900	\$34,562	\$7.0536	C H 5%	bisulfite	nce for a product that is sodium free and/or preservative free suitable for use via nebuliser.
Tramadol hydrochloride						
Cap 50 mg	28,495,572	\$948,903	\$0.0333	C H 5%	a contro	note that tramadol is classified as olled drug. Aggregated tender bids be made across caps, inj and tabs.
Inj 50 mg per ml, 1 ml ampoule				H 5%	a contro	note that tramadol is classified as olled drug. Aggregated tender bids be made across caps, inj and tabs.
<u>lnj 50 mg per ml, 2 ml ampoule</u>				H 5%	a contro	note that tramadol is classified as olled drug. Aggregated tender bids be made across caps, inj and tabs.
Tab sustained-release 100 mg	4,563,999	\$444,990	\$0.0975	C H 5%	a contro	note that tramadol is classified as olled drug. Aggregated tender bids be made across caps, inj and tabs.
Tab sustained-release 150 mg	376,708	\$55,564	\$0.1475	C H 5%	a contro	note that tramadol is classified as olled drug. Aggregated tender bids be made across caps, inj and tabs.
Tab sustained-release 200 mg	254,603	\$48,375	\$0.1900	C H 5%	a contro	note that tramadol is classified as olled drug. Aggregated tender bids be made across caps, inj and tabs.
Triamcinolone acetonide						
0.1% in Dental Paste USP	148,530	\$163,086	\$1.0980	C H 5%		ated tender bids cannot be made dental paste, crm, inj and oint.
<u>lnj 10 mg per ml, 1 ml</u>	5,779	\$24,757	\$4.2840	C H 5%	For pro- Pharma preserv market	nce for a preservative free product. ducts containing preservatives, ac reserves the right to list a ative free product for a restricted which shall not be subject to the ggregated tender bids cannot be cross
<u>Inj 40 mg per ml, 1 ml</u>	80,137	\$843,522	\$10.5260	C H 5%	For pro- Pharma preserv market	nce for a preservative free product. ducts containing preservatives, ac reserves the right to list a ative free product for a restricted which shall not be subject to the ggregated tender bids cannot be cross
Crm 0.02%	6,143,400	\$398,707	\$0.0649	C H 5%		ated tender bids cannot be made dental paste, crm, inj and oint.
Oint 0.02%	3,458,500	\$226,186	\$0.0654	C H 5%		ated tender bids cannot be made dental paste, crm, inj and oint.
Tropicamide						
Eye drops 0.5%	780	\$1,067	\$1.3680	C H 5%	For pro- Pharma preserv market	nce for a preservative free product. ducts containing preservatives, as reserves the right to list a ative free product for a restricted which shall not be subject to the nits and unit subsidy expressed as '.
Eye drops 1%	8,535	\$14,123	\$1.6547	C H 5%	For pro- Pharma preserv market	nce for a preservative free product. ducts containing preservatives, ac reserves the right to list a ative free product for a restricted which shall not be subject to the nits and unit subsidy expressed as '.

	SCHEDULE 1		DUICTS T	O RE .	TEND	EDE	ח	
Chemical Name	SCHEDULE	WO. PRO		OBE	IEND	EKE	ט	
			Unit					
Line Item	Units	Cost	Subsidy		,	ABA	Limit	Comments
Urinary alkaliniser								
Effervescent	2,137,671	\$267,209	\$0.1250		СН	5%		nd subsidy shown are for the y listed sodium citro-tartrate grans
Ursodeoxycholic acid								
<u>Cap 250 mg - 300 mg</u>	1,430,728	\$485,732	\$0.3395		СН	5%		nd subsidy shown are for the y listed cap 200 mg.
Vancomycin hydrochloride								
<u>Inj 500 mg</u>	8,218	\$27,777	\$3.3800	@	СН	5%		
Vancomycin hydrochloride (current ac	cess)							
Cap 125 mg					СН	5%	Schedu may ap	rently listed on the Pharmaceutical ile. Special Authority restrictions ply. Pharmac would only award a for current or widened access.
Cap 250 mg					СН	5%	Schedu may ap	rently listed on the Pharmaceutical ile. Special Authority restrictions ply. Pharmac would only award a for current or widened access.
Vancomycin hydrochloride (widened a	ccess)							
Cap 125 mg					СН	5%	with 202 Infectio currentl Schedu Pharma	atment of Clostridium difficile in line 25 Australasian Society of us Diseases guidelines. Not y listed on the Pharmaceutical ile. Restrictions may apply. ac would only award a tender for or widened access.
Cap 250 mg					СН	5%	with 202 Infectio currentl Schedu Pharma	atment of Clostridium difficile in line 25 Australasian Society of us Diseases guidelines. Not y listed on the Pharmaceutical le. Restrictions may apply. ac would only award a tender for or widened access.
Voriconazole								
Powder for oral suspension 40 mg per i	ml 8,400	\$182,786	\$21.7603		СН	5%		
Water for injection								
Purified for inj, 250 ml bag					Н	5%		
Zonisamide								
Cap 100 mg					СН	5%	Schedu	rently listed on the Pharmaceutical ile, restrictions may apply. iions may be similar to lacosamide
Cap 25 mg					СН	5%	Schedu	rently listed on the Pharmaceutical ile, restrictions may apply. tions may be similar to lacosamide
Cap 50 mg					СН	5%	Schedu	rently listed on the Pharmaceutical lle, restrictions may apply. iions may be similar to lacosamide.

Schedule 3: Tender Process

General

1.1 Principal Supply Period

- (a) Hospital Tender Bids are to be submitted on the basis that if your Hospital Tender Bid is accepted, you will have Principal Supply Status for hospital supply for the particular Tender Item for the Principal Supply Period.
- (b) Community Tender Bids are to be submitted on the basis that if your Community Tender Bid is accepted, you will have Principal Supply Status for community supply for the particular Tender Item for the Principal 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 Principal Supply Status for the particular Tender Item for community and hospital supply for the Principal 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 Health NZ 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 Health NZ Hospitals from the applicable dates specified in clause 2.2 of Schedule Four;
 - (ii) the First Transition Period is intended to allow for an orderly transition to the arrangements that will apply during the Principal Supply Period:
 - (iii) the Final Transition Period is intended to allow for an orderly transition to any new arrangements following the end of the Principal Supply Period.
- (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. 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 2.3 of Schedule Four;
 - the First Transition Period is intended to allow for an orderly transition to the arrangements that will apply during the Principal Supply Period;
 - (iii) the Final Transition Period is intended to allow for an orderly transition to any new arrangements following the end of the Principal Supply Period.
- (c) In relation to community and/or hospital supply, Pharmac may, in its sole discretion:
 - 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

- (ii) 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 (subject to Alternative Brand Allowance arrangements) of other brands of that form and strength of the Chemical Entity is to be deferred until the actual commencement date of the Principal Supply Period, notwithstanding any date previously notified to suppliers by Pharmac as being the intended date of delisting;
 - (B) 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 2.1(a) of Schedule Four will not apply.
- (e) For the avoidance of doubt, any notification by Pharmac of the delisting (subject to Alternative Brand Allowance arrangements) of all other brands of that form and strength of the Chemical Entity on the first day of the Principal 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 (as applicable),

will be deemed to have been entered into between you and Pharmac for Principal Supply Status for community supply 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 Five (as applicable),

will be deemed to have been entered into between you and Pharmac for Principal Supply Status for hospital supply 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) Schedule Five (as applicable),

will be deemed to have been entered into between you and Pharmac for Principal Supply Status for community supply 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) Schedule Five (as applicable),

will be deemed to have been entered into between you and Pharmac for Principal Supply Status for hospital supply 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 and Schedule Five, 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 Principal Supply Period.

1.4 Extension of Principal Supply Status for hospital supply

- (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 Principal Supply Status for community supply, 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:
 - (A) Principal Supply Status for community supply; or
 - (B) a listing in the Pharmaceutical Schedule for the community setting, which does not have Principal Supply Status; 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 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:

- (A) Principal Supply Status for community supply; or
- (B) a listing in the Pharmaceutical Schedule for the community setting, which does not have Principal Supply Status.
- (b) The Community Tender Bid referred to in paragraph (a)(i) above and the contract for community supply 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:
 - (A) Schedules Four and Five in the context of Principal Supply Status; or
 - (B) Pharmac's standard terms of supply for pharmaceuticals used in the community (as recorded in the then Standard Terms Annex of Pharmac's Agreement for the listing of Pharmaceutical(s) on the New Zealand Pharmaceutical Schedule) in the context of a listing, which does not have Principal Supply Status; and
 - (iv) for such quantities of the Pharmaceutical as are required for use in the community.

1.5 Extension of Principal Supply Status for community supply

- (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 Principal Supply Status for hospital supply, 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:
 - (A) Principal Supply Status for hospital supply; or
 - (B) a listing in the Pharmaceutical Schedule for the hospital setting, which does not have Principal Supply Status; or
 - (ii) if Pharmac has accepted your Community Tender Bid for the particular Tender Item, to supply the Tender Item for use in Health NZ Hospitals 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:
 - (A) Principal Supply Status for hospital supply: or
 - (B) a listing in the Pharmaceutical Schedule for the hospital setting, which does not have Principal Supply Status.

- (b) The Hospital Tender Bid referred to in paragraph (a)(i) above and the contract for hospital supply 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:
 - (A) Schedules Four and Five in the context of Principal Supply Status; or
 - (B) Pharmac's standard terms of supply for pharmaceuticals used in Health NZ Hospitals (as recorded in the then Standard Terms Annex of Pharmac's Agreement for the listing of Pharmaceutical(s) on the New Zealand Pharmaceutical Schedule) in the context of a listing, which does not have Principal Supply Status; and
 - (iv) for such quantities of the Pharmaceutical as are required for use in Health NZ Hospitals.

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 Principal Supply Period;
 - (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) the Lead Time and/or the Start Date;
 - (vi) any implementation support if acceptance of your Tender bid resulted in a brand change; 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 Types of Tender Bids

- (a) 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.
- (b) Where you submit a Tender Bid for a Tender Item, you must submit separate Tender Bids for that Tender Item for each unique product you are proposing to supply. For example different pack sizes or brands must be submitted as separate Tender Bids.

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 Market Approval and all necessary Consents. In those circumstances, you may be required to demonstrate your ability to obtain Market Approval and 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 Market Approval and all necessary Consents, any time period to obtain Market Approval and those Consents shall be exclusive of the Lead Time indicated on your Tender Bid;
- (b) is supplied under an exemption under the Medicines Act 1981, the Tender Item shall not be classified as holding Market Approval or a Consent for the purposes of this Invitation.

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:

- 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. You must 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;
- (f) your quality assurance processes, where applicable; and
- (g) how your organisation supports wider social, economic, cultural and environmental outcomes (see New Zealand Government Procurement broader outcomes).

3.3 Information that must be supplied about the Tender Item

You must supply all the information requested in the Tender Submission Form and as otherwise requested by Pharmac, including 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 Market Approval and any other Consent:
 - (i) evidence and justification as to why Market Approval and any other Consent 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, Shelf-Life and In-Use Shelf-Life;
- (c) for any Medical Device:
 - (i) the brand name, pack size (or other equivalent grouping) and type of packaging;
 - (ii) 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 Health NZ;
 - (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 Health NZ 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 any wholesalers and other distributors, in respect of a Community Tender Bid; or
 - (ii) to Health NZ Hospitals, any wholesalers and other distributors, in respect of a Hospital Tender Bid;
- (e) whether it has Market Approval and 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
 - 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, for any Pharmaceutical or Medical Device, information about 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;
 - (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; and

- (v) digital artwork associated with the Tender Item.
- (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, 11 September 2026;
 - (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 Principal Supply 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 Shelf-Life and In-Use Shelf-Life of the Tender Item.
- (d) the price of the Tender Item;
- the amount and timing of savings, including non-pharmaceutical savings accruing to Pharmac during the Principal Supply Period;
- (f) either:
 - (i) evidence that you have obtained, and still have, Market Approval 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 Market Approval and all necessary Consents;
- (g) the name and location of the manufacturer of the finished product and active ingredients of the Tender Item; and
- (h) any other benefits 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 via the electronic Tender Submission Form and an Offer Letter is also submitted;
- (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:
 - 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 Principal Supply Status for community and/or hospital supply 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 Market Approval and all necessary Consents:
 - The contract referred to in clause 1.3 of this Schedule will be conditional upon such Market Approval and Consents being received within a time period specified by Pharmac; and
 - (ii) Pharmac may terminate the contract if such Market Approval and 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 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

- (a) Subject to clause 9.2 below, Confidential Information is confidential to you, Pharmac, Health NZ and those parties' respective Personnel (as applicable).
- (b) You acknowledge that Pharmac may be required to disclose Confidential Information in accordance with:
 - (i) section 12 of the Official Information Act 1982; and
 - (ii) any other legal and administrative obligations,

and you consent to such disclosure.

- (c) Where Pharmac reaches a preliminary view that Confidential Information must be disclosed for the purposes stated in clause 9.1(b)(i) above, Pharmac will consult with you, and will act in good faith, before deciding whether to disclose the Confidential Information.
- (d) To the extent permitted by law, Pharmac will inform you if Confidential Information is disclosed for the purposes stated in clause 9.1(b)(ii) above, including any disclosure to a court, inquiry or ombudsman.
- (e) Confidential Information must not be disclosed by you, Pharmac, Health NZ or those parties' respective Personnel unless:
 - (i) the information is publicly available or enters the public domain through no fault of the applicable parties; or
 - (ii) the disclosure is:
 - (A) required or permitted for the purposes of this Invitation;
 - (B) required or permitted by law; or
 - (C) agreed to between the applicable parties.

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 Principal Supply Status for community and/or hospital supply, as applicable, for your supply of the Tender Item including, without limitation, costs of obtaining Market Approval and 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 Health NZ 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 the Tender Analysts (tender@pharmac.govt.nz) 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 Principal Supply Status for both community and hospital supply

1. Pharmac's Role

1.1 Rights and Responsibilities

- (a) You acknowledge that:
 - (i) Pharmac is required to pursue its statutory objectives, carry out its statutory functions and otherwise comply with its statutory obligations;
 - (ii) Pharmac is subject to a range of legal and administrative 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 role and functions;
 - (iv) the actions which Pharmac may take under its OPPs include (without limitation):
 - (A) listing new pharmaceuticals;
 - (B) changing the terms on which a pharmaceutical is listed; and
 - (C) delisting pharmaceuticals or delisting part or all of a therapeutic group or sub-group; and
 - (v) 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.
- (c) Pharmac may terminate or amend this Agreement at its sole discretion in the following circumstances:
 - (i) Pharmac is issued a Crown Direction;
 - (ii) in accordance with any advice from Medsafe or clinical advice, based on patient safety or any other clinical reasons;
 - (iii) a Supply Issue results in a failure to supply the Pharmaceutical;
 - (iv) any Consent or Market Approval is not held by you or is withdrawn for the Pharmaceutical;
 - (v) a Changed Medicine Notification is approved by Medsafe for the Pharmaceutical; or
 - (vi) the Pharmaceutical is delisted for any reason.
 - (d) In the event that:

- this Agreement is terminated (or notice of termination is given) or amended due to any of the circumstances set out in clause 1.1(c)(i) to (v), Pharmac reserves the right to delist, or suspend or amend the listing of, the Pharmaceutical; or
- (ii) the Pharmaceutical is delisted, or has its listing suspended, for any reason then, unless this Agreement is terminated under clause 1.1(c)(vi), this Agreement shall continue in full force and effect until expiry or termination in accordance with its terms and such delisting or suspension shall not constitute or be construed as a repudiation or breach of the terms of this Agreement by Pharmac. You agree that you do not have, and you expressly waive, any rights, at law, including in equity or under statute, and particularly under Part 2, subpart 3 of the Contract and Commercial Law Act 2017 (Contractual remedies), to terminate this Agreement as a result of the delisting, or suspension of the listing, of the Pharmaceutical.

1.2 Amendments to Pharmaceutical Schedule

- (a) Pharmac retains the right, at its sole discretion, to amend any funding restrictions for the Pharmaceutical, for example where the proposed amendment may increase the market size for the Pharmaceutical. For the avoidance of doubt, an amendment to the funding restrictions may apply during the Principal Supply Period for the Pharmaceutical and where a Tender Bid for current access has been accepted for the Pharmaceutical.
- (b) 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 Supplier Code of Conduct

You must comply with the New Zealand Government's Supplier Code of Conduct as amended or substituted from time to time.

2. Price and Payment

2.1 Subsidy arrangements for community supply

- (a) Subject to clause 2.3 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 (except to the extent any brands remain listed under Alternative Brand Allowance arrangements) on the first day of the Principal Supply Period, with the result that you will have Principal Supply Status in the community for that form and strength of the Chemical Entity during the Principal 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 Principal 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.

2.2 Pricing arrangements for hospital supply

- (a) Subject to Pharmac's other rights under this Agreement and clause 2.3 of this Schedule, on and from the Start Date, during the remainder of the First Transition Period and during the Principal Supply Period, the Pharmaceutical is to be:
 - (i) listed at the Price set out in the Pharmaceutical Schedule;
 - (ii) sold by you to Health NZ Hospitals at the Price.
- (b) Where the Pharmaceutical is included in an order by a Health NZ Hospital for pharmaceuticals where the total value (excluding GST) of the order is less than \$1,000, you may invoice the Health NZ Hospital, in accordance with clause 2.8 below, for the cost of freight for that particular order. For the avoidance of doubt, this clause does not entitle you to invoice a Health NZ 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 3.6 of this Schedule Four), and provided that there are no Alternative Pharmaceuticals listed in 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 2.2(a)(i) and (ii) above during the Final Transition Period and beyond; and
 - (ii) is not to be delisted during the Final Transition Period.

2.3 Price change

- (a) Subject to clause 2.3(c)(ii), clause 2.3(c)(iii) and clause 2.3(c)(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) Subject to clause 2.3(c)(ii), clause 2.3(c)(iii) and clause 2.3(c)(iv) of this Schedule, you must change the price at which you supply the Pharmaceutical to Health NZ Hospitals 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.
- (c) 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 any 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 any 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 any 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 or purchased 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 2.3(c)(i) or (c)(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 or purchased at the Price from the Start Date; and
 - (iv) notwithstanding clauses 2.3(c)(i), (c)(ii) or (c)(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 any 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 any 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 2.3(c)(i) or (c)(ii) above apply to the Pharmaceutical, clause (c)(i) above shall apply.

(d) You shall upon request by Pharmac, provide information on how you intend to manage the price changes stated in clauses 2.3(c)(i) to (c)(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 Three.

2.4 Supply Price

During each of the First Transition Period, the Principal Supply Period and the Final Transition Period, the price at which the Pharmaceutical is supplied by you to any wholesalers, other such distributors and/or to a Health NZ Hospital, must not exceed the Price.

2.5 Pharmaceutical Price

You warrant that the Price is inclusive of all costs, for example but not limited to any costs relating to manufacturing, supply chain and price fluctuations, which may occur during the First Transition Period, the Principal Supply Period and the Final Transition Period.

2.6 No reference pricing during Principal 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 Principal 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.

2.7 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.

2.8 Invoices to hospitals

Where a Health NZ Hospital is to be invoiced for the Pharmaceutical, you are to invoice the particular Health NZ Hospital at the end of each month, but no later than the 10th 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 Health NZ Hospital's purchase order reference number (if applicable);
- (c) the net amount payable in respect of the Pharmaceutical supplied to that Health NZ Hospital in accordance with this Agreement;
- (d) full details in respect of the Pharmaceutical supplied to that Health NZ Hospital in accordance with this Agreement, including the:
 - (i) Health NZ Hospital 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 2.2(b) above);

- (v) total cost for the total amount of the Pharmaceutical supplied; and
- (e) any other information that Health NZ Hospital requires you to supply.
- (f) The provisions of clause 2.8 do not apply to the extent that both parties have agreed to alternative or varied invoicing arrangements in respect of a particular Pharmaceutical.

2.9 Payment by hospitals

- (a) Provided that the Pharmaceutical has been supplied in accordance with this Agreement, and the particular Health NZ Hospital receives an invoice in accordance with clause 2.8 above, payment by the Health NZ 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 Health NZ 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 Health NZ Hospital later than the 10th 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 2.8 above, payment by the Health NZ 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 Health NZ Hospital;
 - (ii) on the 20th day of the month following the month in which you invoice the Health NZ Hospital 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.

2.10 Future payment by hospitals

- (a) A particular Health NZ Hospital's failure to dispute any invoice prior to payment does not prejudice that Health NZ 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 Health NZ Hospital may withhold, deduct or set off the amount of any overpayment or any amount recoverable by that Health NZ Hospital from you under this Agreement from any future amount owing to you.

3. Principal Supply Status

3.1 Principal Supplier

- (a) Subject to:
 - Pharmac's other rights under this Agreement in relation to the Pharmaceutical;
 and
 - (ii) this clause 3 of Schedule Four relating to the Alternative Brand Allowance,

Pharmac will not subsidise another supplier's brand of the Pharmaceutical on the Pharmaceutical Schedule and/or Health NZ Hospitals will not purchase another supplier's brand of the Pharmaceutical, at any time during the Principal 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 principal supplier of any forms and strengths of the Chemical Entity, if such supply commences after the end of the Principal 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.
- (d) You shall have Principal Supply Status during the Principal Supply Period, which shall be subject to the Alternative Brand Allowance, where other supplier brands of the Pharmaceutical may be subsidised in the community and/or purchased by Health NZ Hospitals.
- (e) The Alternative Brand Allowance referred to in paragraph (a) above is specified as a percentage of the Total Pharmaceutical Volume for the Pharmaceutical, that percentage being as set out in Schedule Two.
- (f) You acknowledge and agree that any other supplier brands of the Pharmaceutical may be concurrently listed on the Pharmaceutical Schedule at any time during the First Transition Period, the Principal Supply Period and the Final Transition Period and your rights under this Agreement do not extend to an exclusive listing of the Pharmaceutical on the Pharmaceutical Schedule.

3.2 Exceptions to Principal Supply Status

- (a) Pharmac may, from time to time during the Principal Supply Period or the First Transition Period, amend the Alternative Brand Allowance for the Pharmaceutical after consultation with a relevant medical adviser (being either the Ministry of Health, PTAC or its Specialist Advisory Committees), provided that Pharmac may only increase the Alternative Brand Allowance without your prior agreement if it has a direction to that effect from Medsafe or its successor, or a recommendation that it do so from PTAC or its Specialist Advisory Committees, based on a significant clinical issue.
- (b) Subject to clause 3.3 of this Schedule, you acknowledge and agree that while you have Principal Supply Status:

- (i) other supplier brands of the Pharmaceutical may be subsidised in the community and/or purchased by Health NZ Hospitals, subject to the Alternative Brand Allowance; and
- (ii) without derogating from any other rights available to Pharmac or Health NZ 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) at any time during the Principal Supply Period, then the Alternative Brand Allowance shall not apply and other supplier brands of the Pharmaceutical may be subsidised in the community and/or purchased by Health NZ Hospitals without limitation during that period of non-supply and any calculation performed in accordance with clause 3.3 below shall exclude that period of non-supply.

3.3 Principal Supply Status Monitoring

- (a) If you reasonably believe that the percentage usage of other supplier brands of the Pharmaceutical subsidised in the community and/or purchased by Health NZ Hospitals exceeds the Alternative Brand Allowance for a particular Pharmaceutical during the Principal Supply Period, you may at any date after a three (3) month period following the end of any Relevant Period, request that Pharmac carry out calculations for that Relevant Period in accordance with the procedure set out in this clause 3.3, and Pharmac may, acting reasonably, agree to carry out such calculations, provided that if Pharmac refuses to carry out such calculations, it will provide you with the reasons for refusing to do so. For the avoidance of doubt, where you have Principal Supply Status for both community and hospital supply of a Pharmaceutical, Pharmac will carry out any calculations for those markets in combination, with a single, combined figure to be used for each of Total Pharmaceutical Volume and Total Brand Allowance Pharmaceutical Volume when carrying out the calculations below.
- (b) Within 30 Business Days of Pharmac accepting your request to carry out calculations in accordance with paragraph (a) above, Pharmac shall carry out the following calculations for the Relevant Period in question:
 - (i) (Total Brand Allowance Pharmaceutical Volume / Total Pharmaceutical Volume) x 100 = Brand Allowance Indicator;
 - (ii) Brand Allowance Indicator Alternative Brand Allowance = Brand Differential
- (c) In the event the Brand Differential is a number greater than zero i.e. a positive amount, Pharmac shall carry out the following calculations for the Relevant Period in question:
 - (i) Total Pharmaceutical Volume / 100 = Volume Multiplier;
 - (ii) Volume Multiplier x Brand Differential = Eligible Volume;
 - (iii) (Eligible Volume x Unit Price and/or Unit Subsidy) / 2 = Brand Compensation
- (d) Pharmac will notify you in writing of any Brand Compensation payable or not in accordance with paragraphs (b) and (c) above and will provide you with the details of the relevant party or parties to be invoiced for any Brand Compensation payable. Following such notification to you from Pharmac, you may invoice the relevant party or parties for the Brand Compensation.

- (e) You acknowledge and agree that the data extracted from the records used by Pharmac are the best data and those records are the best records, for the purposes of carrying out the calculations.
- (f) You may, within 10 Business Days following notification of the outcome of the calculations in accordance with paragraph (d) above (the "Calculation"), notify Pharmac in writing that you dispute the particular Calculation and that you require an audit of that Calculation to be carried out. If you do give a notice to Pharmac under this clause within that 10-Business Day period, then the following provisions are to apply:
 - (i) The audit is to be carried out by an independent person jointly approved by us or, if there is no agreement on a mutually acceptable person within 5 Business Days of the date of your notice under this clause, then by an independent person nominated for that purpose by the President for the time being of the Institute of Chartered Accountants of New Zealand.
 - (ii) The independent person is to audit the particular Calculation, which is disputed by you, based on all relevant electronic data which is extracted by Pharmac from the records maintained by it. For the avoidance of doubt, the independent person will have no right to inspect, review or have access to the written prescriptions on which the data extracted by Pharmac from those electronic records are based, nor any right to request copies of those written prescriptions.
 - (iii) In carrying out the audit, the independent person is to be considered as acting as an expert and not as an arbitrator.
 - (iv) The independent person will be required to complete the audit, and to provide us with a written determination in that regard, within 5 Business Days of receiving all the information required by the independent person to make a determination and, in any case, no later than 10 Business Days from the date of his or her acceptance of appointment or nomination under this clause, unless we agree otherwise. The independent person's determination of the particular Calculation is to be final and binding on both of us.
 - (v) The costs incurred by the independent person in completing the audit are to be met by you, irrespective of the outcome of the audit.

3.4 Withdrawal of Principal Supply Status

- (a) Pharmac may withdraw Principal Supply Status in relation to the Pharmaceutical (in which case clauses 3.1 and 3.3 of this Schedule Four will no longer apply), by written notice to you at any time during the Principal Supply Period or (in anticipation) during the First Transition Period in the event of a Supply Issue, or in accordance with any advice from Medsafe or clinical advice, based on patient safety or any other clinical reasons.
- (b) Any withdrawal of Principal Supply Status is without prejudice to Pharmac's rights under clauses 5.5 and 5.6 of this Schedule Four.

3.5 Suspension of Principal Supply Status

(a) Pharmac may suspend Principal Supply Status in relation to the Pharmaceutical, by written notice to you at any time during the Principal Supply Period or (in anticipation) during the First Transition Period in the event of a Supply Issue, or in accordance with any advice from Medsafe or clinical advice, based on patient safety or any other clinical reasons.

- (b) Any suspension of Principal Supply Status is without prejudice to Pharmac's rights under clauses 5.5 and 5.6 of this Schedule Four.
- (c) Pharmac may, at any time, in its sole discretion, notify you of the date on which the suspension of Principal Supply Status under this clause 3.5 ceases and on which date:
 - Principal Supply Status is to be re-implemented in respect of the Pharmaceutical; or
 - (ii) Principal Supply Status is to be withdrawn in accordance with clause 3.4 of this Schedule Four.

3.6 Subsidy and 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:
 - (i) you will cease to have Principal Supply Status for that form and strength of the Chemical Entity or you will cease to have Principal Supply Status for hospital supply 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);
 - the Pharmaceutical will remain listed in the Pharmaceutical Schedule subject to the current standard terms Annex of Pharmac's Agreement for the listing of Pharmaceutical(s) on the New Zealand Pharmaceutical Schedule contract template;
 - (iii) you may increase the price ex-manufacturer (exclusive of GST) at which you sell or supply, or make available for supply or sale, by you, to:
 - in relation to community supply, any wholesalers and other such distributors; or
 - (B) in relation to hospital supply, Health NZ Hospitals, any 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, subject to sub-paragraphs (A) to (C) as follows:

- (A) Pharmac reserves the right to consult on any subsidy increases prior to determining whether to increase the subsidy for the Pharmaceutical to the new price notified under this paragraph (a)(iii);
- (B) Where you increase the price at which you supply the Pharmaceutical under this paragraph (a)(iii), you will not subsequently increase the price at which you supply the Pharmaceutical for at least 12 months from the effective date of the price increase;
- (C) Where you did not obtain Market Approval and all necessary Consents for the Pharmaceutical within 12 months following the Deadline, you may not provide six months' written notice of any price increase until a date on or after 12 months following the End Date.

- (iv) if Pharmac does not increase the subsidy or price 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 or price 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 due to a Force Majeure Event); 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.

4. Reporting

4.1 Information

- (a) You agree to provide any information related to the Pharmaceutical and its listing that Pharmac reasonably requests, in such manner and timeframe as Pharmac reasonably requests.
- (b) In particular, and without limiting the generality of clause 4.1(a) above, you:
 - (i) acknowledge that Pharmac requires the provision of Unique Product Identifiers in order to implement the listing of each Pharmaceutical and you agree to obtain and notify Pharmac of the Unique Product Identifiers of each Pharmaceutical as follows:
 - (A) for brand changes, no later than the earlier of:
 - the 12th of the month following the Market Notification Date; or
 - 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;

- (ii) agree to provide Pharmac with digital photos of the Pharmaceutical (e.g. tablet, vial or patch) and its associated packaging, which will be supplied to the New Zealand market, when the Pharmaceutical is available for distribution in New Zealand. If any changes are made to the Pharmaceutical or its associated packaging whilst the Pharmaceutical is listed on the Pharmaceutical Schedule, you shall provide Pharmac with updated digital photos as soon as practicable following those changes being implemented;
- (iii) agree that in the event that you supply an Alternative Pharmaceutical in accordance with this Agreement, or in the event of a Changed Medicine Notification for a Pharmaceutical, you must notify Pharmac of any changed Unique Product Identifiers (or advise if there is no change) as soon as practicable;
- (iv) acknowledge that in the event the listing of the Pharmaceutical includes Special Authority criteria or any other access criteria, you must, for the duration that the Pharmaceutical is listed:
 - (A) notify Pharmac in the event the Data Sheet is amended in a manner which, when considered in the context of any current Special Authority criteria or other current access criteria, could impact on patient safety; and
 - (B) provide Pharmac with a summary of the amendment to the Data Sheet as set out in clause 4.1(b)(iv)(A) above;

Following the notification in clause 4.1(b)(iv)(A) Pharmac reserves the right at its sole discretion to amend the Special Authority criteria or any other access criteria for the Pharmaceutical based on patient safety;

- (v) acknowledge that Pharmac may require stock reports and batch details held by you for the Pharmaceutical and you agree to provide all such stock reports and batch details to Pharmac upon request;
- (vi) acknowledge that Pharmac may require price and volume data held by you relating to sales of the Pharmaceutical and you agree to provide all such price and volume data to Pharmac upon request; and
- (vii) agree that Health NZ may provide Pharmac and its agents with any price and volume data held by Health NZ in respect of the Pharmaceutical, and Pharmac may share any price and volume data held by Pharmac with Health NZ.

4.2 Supply Issues Reporting

- (a) You must send a Supply Issues Report to Pharmac in accordance with clause 5.3(a)(ii) of this Agreement or otherwise at Pharmac's request.
- (b) The Supply Issues Report must be provided to Pharmac in any form notified by Pharmac to you. Unless notified otherwise, the Supply Issues Report must include the following information:
 - (i) average usage of the Pharmaceutical in New Zealand;
 - (ii) quantity of Pharmaceutical stock:
 - (A) held by you (or on your behalf) in New Zealand;
 - (B) held by you (or on your behalf) in other international markets, and available for supply in New Zealand; and
 - (C) held by any wholesalers in New Zealand;
 - (iii) reason for the Supply Issue;
 - (iv) when the Supply Issue occurred;
 - (v) expected delivery dates of the Pharmaceutical to New Zealand;
 - (vi) expected date of authorised release into the New Zealand market (including the date on which the Pharmaceutical is expected to be available for supply) and any applicable supporting evidence, for example export and import licences or other official authorisations and customs formalities necessary for the exportation and importation of the Pharmaceutical;
 - (vii) the estimated duration of the Supply Issue; and
 - (viii) any steps that you have taken or will take to mitigate the risk that you may fail to supply a Pharmaceutical.
- (c) You acknowledge that Pharmac may wish to engage with you in respect of any steps that you advise Pharmac of under clause 4.2(b)(viii) above or any other steps that may be required to mitigate the risk that you may fail to supply the Pharmaceutical, and you agree that you will engage and cooperate with Pharmac in relation to all such actual and proposed mitigation activities.

5. Supply Obligations and Managing Supply Issues

5.1 Stock Holdings

The minimum stock holding of the Pharmaceutical that must be held by you (or on your behalf) in New Zealand and available for supply is set out in each of the rows in the table below. For the avoidance of doubt:

- (a) each minimum stock holding specified in the table in respect of an ASP or non-ASP Pharmaceutical is independent and separate from the other minimum stock holdings specified in the table, such that a failure to meet any of these minimum stock holding requirements for (as applicable) an ASP or non-ASP Pharmaceutical will be a Supply Issue; and
- (b) the minimum stock holdings set out in the table refer to stock physically held by you or on your behalf in New Zealand and do not include stock held in New Zealand by any wholesalers or other parties.

Stock Holding Type	Community and/or Hospital Supply	Minimum Stock Holding
General (non-ASP) stock holding requirement	Community or hospital supply	Two-thirds of your most recent three months' total Unit sales of the Tender Item
ASP	Community or hospital supply	Your most recent four months' total Unit sales of the Tender Item
General (non-ASP) stock holding requirement	Community or hospital supply	Forecast sales demand in respect of the next two-month period is greater than your stock of the Pharmaceutical
ASP	Community or hospital supply	Forecast sales demand in respect of the next four-month period is greater than your stock of the Pharmaceutical
General (non-ASP) stock holding requirement	Hospital	The average volume of stock of the Pharmaceutical required to supply the entire Health NZ Hospital market for the Pharmaceutical for any given twomonth period
ASP	Hospital	The average volume of stock of the Pharmaceutical required to supply the entire Health NZ Hospital market for the Pharmaceutical for any given fourmonth period
General (non-ASP) stock holding requirement	Community	One-sixth of the Unit Volume

ASP	Community	One-third of the Unit Volume
General stock holding requirement (ASP and non-ASP)	Community or hospital supply	Insufficient to enable you to fully fill all orders as they are received (without restricting quantities that may be ordered)
New Zealand manufactured products (ASP and non-ASP)	Community or hospital supply	Forecast sales demand in respect of the next two-month period is greater than your stock of the Pharmaceutical; or you have insufficient stock to enable you to fully fill all orders as they are received; or 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

5.2 Continuity of Supply

- (a) You must supply, and continue to supply, the Pharmaceutical on the terms set out in this Agreement.
- (b) You warrant that you have entered into all contractual and other arrangements to the extent necessary, including licence and supply agreements with third parties, to ensure that you will meet all of your obligations under clause 5.2(a) above and this Agreement generally.

5.3 Notification

- (a) You must:
 - (i) notify Pharmac as soon as you become aware of a Supply Issue; and
 - (ii) send a Supply Issues Report to Pharmac within 2 Business Days of becoming aware of a Supply Issue.
- (b) In the event that you consider (acting reasonably) that any circumstances or events may result in a Supply Issue you must notify Pharmac in writing as soon as practicable, including (but not limited to) any of the following circumstances:
 - you plan any changes to your supply chain, for example but not limited to a change in manufacturing site, in respect of the Pharmaceutical;
 - (ii) you plan any changes to your ordering or delivery systems;
 - (iii) you plan to re-structure your organisation; or
 - (iv) you plan to change the presentation of the Pharmaceutical, including the brand name, pack size, packaging and strength.

(c) After giving Pharmac notice in accordance with clauses 5.3(a)(i) and/or 5.3(b), you must comply with all reasonable requests by Pharmac in respect of steps that may be required to mitigate the risk that you may fail to supply the Pharmaceutical.

5.4 Managing Supply Issues

- (a) In addition to your obligations set out in clause 5.3 you must comply with the obligations set out in this clause 5.4.
- (b) In the event of:
 - a decision or notification by Medsafe or any other authorities to recall the Pharmaceutical; or
 - (ii) the withdrawal of any Consent or Market Approval for the Pharmaceutical,

you must use your best endeavours to engage and co-operate with Medsafe and any other relevant authorities and must, at all times, meet all your regulatory obligations.

- (c) In the event a Supply Issue actually results in a failure to supply, or you have reason to believe may cause you to fail to supply, the Pharmaceutical in accordance with the terms of this Agreement, then:
 - (i) subject to the prior written consent of Pharmac, you must use your best endeavours to procure, within what Pharmac considers to be a reasonable period of time, an Alternative Pharmaceutical for supply to:
 - (A) any wholesalers and other such distributors; and
 - (B) any Health NZ Hospital,

at the Price; and

- (ii) if you fail to procure an Alternative Pharmaceutical at the Price and within the timeframe in accordance with clause 5.4(c)(i) above then Pharmac may implement an arrangement with another supplier to supply an Alternative Pharmaceutical (including an arrangement for back-up supply) and you must pay to Pharmac any additional costs, fees and/or expenses incurred by Health NZ or Pharmac as a result of the purchase of the Alternative Pharmaceutical over and above the costs that would have been incurred by Pharmac had you supplied the Pharmaceutical.
- (d) In the event Pharmac receives information that indicates that you may fail to supply a Pharmaceutical in accordance with this Agreement (whether you notify Pharmac under this Agreement or otherwise), you agree that Pharmac may inform other interested parties who may be impacted, including providing other suppliers with sufficient information to allow those suppliers to adequately prepare for a potential change in demand.

5.5 Indemnity

You agree to indemnify Pharmac and Health NZ (as applicable) for any damages, liability, loss, cost (operational or otherwise) or expense awarded against, incurred or suffered by Pharmac and/or Health NZ as a result of or arising from a Supply Issue (other than a Supply Issue resulting directly from a Force Majeure Event). This indemnity shall be deemed to indemnify Pharmac and Health NZ for all additional costs, including all costs incurred by

Pharmac and/or Health NZ as a result of the purchase of the Alternative Pharmaceutical that are additional to any costs specified in clause 5.6.

5.6 Liquidated Damages

- (a) Subject to clause 5.6(c) and clause 5.6(d), for each and every Supply Issue which actually results in a failure to supply the Pharmaceutical (other than a Supply Issue resulting directly from a Force Majeure Event) you must pay to Pharmac liquidated damages (plus GST (if any)) of \$50,000 to cover Pharmac's administrative and/or operational costs.
- (b) You acknowledge that Pharmac's right to claim the full liquidated damages amount specified in clause 5.6(a) in these circumstances reflects Pharmac's legitimate interests in securing delivery of the Pharmaceutical by the relevant date and in accordance with the terms of this Agreement and is proportionate to those interests during the period, and in the circumstances, in which the liquidated damages are payable under this clause 5.6.
- (c) Liquidated damages are payable where you have not:
 - (i) notified Pharmac under and in accordance with clause 5.3; and/or
 - (ii) complied with all reasonable requests by Pharmac in respect of steps that may be required to mitigate the risk that you may fail to supply the Pharmaceutical.
- (d) Pharmac may, in its sole discretion, require you to pay less than the amount specified as liquidated damages in clause 5.6(a) if Pharmac is satisfied that the actual costs in the circumstances are less than this amount.

5.7 Interest

If payment of any amount required to be paid by you under clauses 5.5 or 5.6 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 on such sum as remains unpaid at a rate per annum equal to the Default Interest Rate, 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 does not limit any other right or remedy of Pharmac; and
- (b) Pharmac may take any action, including legal action, without first needing to implement the dispute resolution procedure contained in clause 6.4, to recover that unpaid amount and you agree to pay to Pharmac actual enforcement costs incurred in relation to that action.

6. **General Obligations**

6.1 Shelf-life of Pharmaceutical

You will not supply the Pharmaceutical:

- (a) if the remaining shelf-life of that Pharmaceutical is less than 6 months; or
- (b) where the total shelf-life of that Pharmaceutical is less than 6 months, the remaining shelf-life is less than 75% of that Pharmaceutical's total shelf-life,

without prior written agreement from Pharmac or the applicable Health NZ Hospital.

6.2 Consents

- (a) Prior to the Start Date you must obtain:
 - (i) Market Approval for the Pharmaceutical;
 - (ii) any Consent required for the supply of the Pharmaceutical; and
 - (iii) any other Consent Pharmac requires you to have or hold.
- (b) You must maintain Market Approval and any other Consent specified in clauses 6.2(a)(ii) and 6.2(a)(iii) for the Pharmaceutical for the duration the Pharmaceutical is listed.

6.3 Health and Safety

Where delivery of the Pharmaceutical (or provision of any related services described in this Agreement) occurs within the facilities of a Health NZ Hospital, you and your Personnel will comply with all relevant health and safety requirements, including:

- (a) the Health and Safety at Work Act 2015 and all regulations made under that Act; and
- (b) any policies and procedures communicated to you by the Health NZ Hospital.

6.4 Dispute Resolution

If there is a dispute between you and Pharmac arising out of, or in connection with, this Agreement, neither of the parties 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) You and Pharmac will endeavour, in good faith, to resolve the dispute referred to in the notice by using informal dispute resolution techniques.
- (c) If you and Pharmac have not resolved the dispute within 14 days after the date notice of a dispute was given, the parties may agree that the dispute is to be:
 - mediated according to the standard mediation agreement of the Resolution Institute (a body corporate incorporated in Australia and registered as an overseas company in New Zealand), and the Chair of the Resolution Institute

- (or the Chair's nominee) will select the mediator and determine the mediator's remuneration, if you and Pharmac are unable to agree on such matters; or
- (ii) submitted to arbitration in accordance with the Arbitration Act 1996, with such arbitration being conducted by a single arbitrator to be agreed on by the parties or, failing agreement, the Chair of the Resolution Institute (or the Chair's nominee) will select the arbitrator.
- (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.
- (e) Pending resolution of the dispute, this Agreement will remain in full effect without prejudicing the parties' respective rights and remedies (including Pharmac's rights under its OPPs).

6.5 Litigation Support

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

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

you will give Pharmac all assistance it reasonably requires for the purpose of the handling of any negotiations and/or litigation related to those proceedings or any claim.

6.6 Listing in the Pharmaceutical Schedule of a PCT

- (a) Where the Pharmaceutical is a PCT and supplied in a Health NZ Hospital, you acknowledge and agree that Pharmac may list the Pharmaceutical in the community setting 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 the Pharmaceutical Schedule.
- (b) If Pharmac lists the Pharmaceutical in the community setting 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 the community setting will, at Pharmac's option, be additional to or instead of listing in the hospital setting; and
 - (iii) references to the "listing" of the Pharmaceutical will, where applicable, be to the listing of the Pharmaceutical in the community setting of the Pharmaceutical Schedule (and references to "list", "listed", "delist", "delisted", and "delisting" are to be interpreted accordingly).

6.7 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 5.5 and 5.6 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 Principal Supply 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.

7. General Terms

7.1 No Derogation

The express provision of a remedy for, or consequence of, breach of any term of this Agreement does not derogate from, or limit, any other legal right or remedy available to Pharmac under this Agreement or otherwise in respect of such breach.

7.2 No Waiver

A failure or delay by either you or Pharmac 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.

7.3 Remedies Cumulative

Except as is expressly stated otherwise in this Agreement:

- the rights, powers and remedies provided in this Agreement are cumulative and are not exclusive of any rights, powers or remedies provided by law or under this Agreement; and
- (b) the exercise of any of the rights, powers and remedies provided in this Agreement will not prejudice the exercise of any other right, power or remedy under this Agreement or existing at law.

7.4 Entire Agreement

This Agreement:

- (a) is the entire agreement between you and Pharmac regarding the terms on which the Pharmaceutical is listed; and
- (b) supersedes and extinguishes all prior agreements and understandings between you and Pharmac, and between you and Health NZ, and any prior agreements and understandings originally entered into between you and district health boards (as applicable), regarding the Pharmaceutical and the subject matter contained herein.

7.5 Advertising

You must ensure that any Advertisement aimed at consumers of the Pharmaceutical does not breach any applicable statute, regulation or industry standard, including the Advertising Standards Authority Codes of Practice and the Medicines New Zealand Code of Practice.

7.6 Contracts Privity

- (a) You and Pharmac acknowledge that your obligations in this Agreement constitute promises and obligations which confer or are intended to confer a benefit on Health NZ and related persons, and are enforceable by Health NZ and any such persons pursuant to Part 2, subpart 1 of the Contract and Commercial Law Act 2017 (Contractual Privity).
- (b) Except as expressly provided in clause 7.6(a) above, the parties do not intend to create rights in, or grant remedies to, any third party as a beneficiary to this

Agreement, and all of the provisions of this Agreement shall be for the sole and exclusive benefit of the parties.

(c) 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 Health NZ.

7.7 No Reliance

You acknowledge that you have entered into this Agreement in reliance on your own knowledge, skill and independent advice, and not in reliance on any representations made, or any information made available to you, by Pharmac.

7.8 Amendments

Amendments to this Agreement must be in writing.

7.9 Assignment

You will not permit this Agreement, or any part of this Agreement, to be transferred or assigned (either directly or due to a change of 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.

7.10 Further Assurances

You and Pharmac agree to execute any further documents and do any further acts as may be reasonably necessary from time to time to give effect to the terms and intentions of this Agreement.

7.11 Specific Performance

You acknowledge that in the event of any breach or threatened breach of this Agreement by you, damages may not be an adequate remedy and Pharmac may seek specific performance of the terms of this Agreement or injunctive relief or any other similar remedy, in addition to any other rights, powers or remedies provided under this Agreement or by law (including equity).

7.12 Agreement Prevails

Where any of your terms of supply, for example on invoices or any purchase orders, conflict or are inconsistent with any of the terms of this Agreement, the terms of this Agreement will prevail.

Schedule 5: Additional Special Terms

1. Intra-uterine device (Non-hormonal)

You shall provide the following information when submitting a Tender Bid for the Pharmaceutical intra-uterine device [Non-hormonal] (IUD Tender Item):

- the size of the IUD Tender Item, including length and width measurements as well as the diameter of the inserter;
- a description of the material the inserter is made of;
- the duration of the therapeutic effect of the IUD Tender Item; and
- the metals which are contained in the IUD Tender Item.

You shall provide the following Resources at no cost for the IUD Tender Item, for the provision of education, training and support to healthcare professionals in respect of the use of the IUD Tender Item.

For the purposes of this clause "Resources" shall include but not be limited to the:

- provision of training materials (DVDs, pamphlets, leaflets, brochures) to healthcare professionals;
- provision of an information sheet explaining the differences between the current brand of intra-uterine device and your IUD Tender Item; and
- provision of presentations and/or demonstrations on the use of your IUD Tender Item to patients and/or healthcare professionals.