INVITATION TO BID



HP07-2026DAI

SUPPLY AND DELIVERY OF PHARMACEUTICAL PRODUCTS: DROPS, AEROSOLS AND INHALED MEDICINES TO THE DEPARTMENT OF HEALTH FOR THE PERIOD 1 SEPTEMBER 2026 TO 31 AUGUST 2029

BID VALIDITY PERIOD: 180 DAYS

NON-COMPULSORY ONLINE BRIEFING SESSION:
MS TEAMS: 10 OCTOBER 2025 AT 10:00



Private Bag X828, PRETORIA, 0001. DR AB Xuma Building, 1112 Voortrekker Road, Pretoria Townlands 351-JR, PRETORIA 0187 Directorate: Affordable Medicines

Ref: HP07-2026DAI e-mail: tenders@health.gov.za

INVITATION TO BID: HP07-2026DAI

SUPPLY AND DELIVERY OF PHARMACEUTICAL PRODUCTS: DROPS, AEROSOLS AND INHALED MEDICINES TO THE DEPARTMENT OF HEALTH FOR THE PERIOD 1 SEPTEMBER 2026 TO 31 AUGUST 2029

- 1. Kindly furnish the Department of Health with a tender for the supplies shown on the attached forms.
- 2. Included are the General Conditions of Contract (GCC), Special Requirements and Conditions of Contract (SRCC) as well as the Standard Bidding Document (SBD) and Pharmaceutical Bidding Document (PBD) forms listed on the annexure hereto. The Bid Response Document is available as a separate Excel file.
- 3. The Invitation to Bid document, with all pages and forms <u>completed in detail</u>, must be returned with your bid (marked Set 1). Include a USB flash drive with a scanned copy of the completed bid (marked Set 2). Scanned files in Set 2, must be in the exact compilation sequence as per index. All Excel spreadsheets as Set 3, must be on USB flash drive for uploading purposes.
- **4.** All sets to be in a single sealed package with the following information on the outside of the package: Bid number and Closing date of bid, Full name and address of the bidder, Return address and Name of Contact person.
- 5. The bid must be addressed to the Director-General, Department of Health, and be deposited into the pharmaceutical tender box as indicated on the SBD1 form not later than the closing date and time of the bid. The tender box is located at the main entrance of the Department of Health, DR AB Xuma Building, located at 1112 Voortrekker Road, Pretoria Townlands 351-JR, PRETORIA.

Ms K Jamaloodien

& Jana100 Tren

CHIEF-DIRECTOR: SECTOR-WIDE PROCUREMENT

For: Director-General Date: 26 September 2025

Authorised Signatory: Sign or Initial _____ HP07-2026DAI Page 2 of 126

CONTACT PERSONS AT THE NATIONAL DEPARTMENT OF HEALTH

Please direct any queries relating to the bidding process to tenders@health.gov.za

BID DOCUMENTS FOR COMPLETION AND SUBMISSION

To ensure accurate completion of this bid, please adhere to the requirements specified in the Special Requirements and Conditions of Contract section below:

Bid Document Checklist : Paragraph 3

Bid Documents : Paragraph 4.1 (4.1.1 to 4.1.9)

Consortiums Joint Ventures (incorporated

or unincorporated), and Partnerships : Paragraph 4.3

Submission of bids : Paragraph 9

• Completion of documents and

Bid submission : Paragraph 10

INVITATION TO BID SBD 1

YOU ARE HEREBY INVITED TO BID FOR REQUIREMENTS OF THE NATIONAL DEPARTMENT OF HEALTH

BID NUMBER: HP07-2026DAI

CLOSING DATE: 24 NOVEMBER 2025 CLOSING TIME: 11:00

DESCRIPTION: HP07-2026DAI: SUPPLY AND DELIVERY OF PHARMACEUTICAL

PRODUCTS: DROPS, AEROSOLS AND INHALED MEDICINES TO THE DEPARTMENT OF HEALTH FOR THE PERIOD 01 SEPTEMBER 2026 TO

31 AUGUST 2029

Bid documents must be addressed as follows and delivered before the closing date and time:

Addressed to:

The Director-General: Health Dr AB Xuma Building 1112 Voortrekker Road PRETORIA

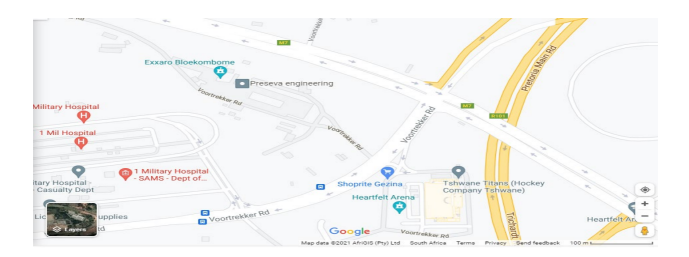
Delivered to:

Directorate: Affordable Medicines Dr AB Xuma Building 1112 Voortrekker Road, Block A Pretoria Townlands 351-JR PRETORIA 0187

Bidders should ensure that bids are delivered on time to the correct address and deposited in the Tender Box. Late bids will not be accepted for consideration

The Pharmaceutical Tender Box is generally accessible during working hours.

See below for map locating Dr AB Xuma Building within Pretoria.



ALL BIDS MUST BE SUBMITTED ON THE OFFICIAL FORMS

This competitive bidding process is subject to the Preferential Procurement Policy Framework Act and the Preferential Procurement Regulations, 2022, the General Conditions of Contract (GCC) and, if applicable, any other Special Requirements and Conditions of Contract.

PART A INVITATION TO BID

YOU ARE HEREBY INVI			E (NAME OF					
	2026DAI	CLOSING DATE:			NOVEMBER 2025		OSING TIME:	11:00
		OF PHARMACEUTICAL PRO NOD 1 SEPTEMBER 2026 TO				NHALED	MEDICINES TO	THE DEPARTMENT
BID RESPONSE DOCUMENTS	MAY BE DEPOSITED	IN THE BID BOX SITUATED A	T 1112 VOORT	REK	KER ROAD, PRETORI	A TOWNL	ANDS 351-JR, PRE	TORIA
PHARMACEUTICAL TEN	PHARMACEUTICAL TENDER BOX							
RECEPTION AREA								
NATIONAL DEPARTMENT OF HEALTH								
DR AB XUMA BUILDING								
BIDDING PROCEDURE ENQUIRIES MAY BE DIRECTED TO TECHNICAL ENQUIRIES MAY BE DIRECTED TO:								
CONTACT PERSON			CONTACT	PER	RSON			
TELEPHONE NUMBER			TELEPHO	NE N	IUMBER			
FACSIMILE NUMBER			FACSIMILI	E NU	IMBER			
E-MAIL ADDRESS SUPPLIER INFORMATION	tenders@health	n.gov.za	E-MAIL AD	DRE	SS		tenders@	health.gov.za
NAME OF BIDDER								
POSTAL ADDRESS								
STREET ADDRESS								
TELEPHONE NUMBER	CODE			NU	MBER			
CELLPHONE NUMBER								
FACSIMILE NUMBER	CODE			NU	MBER			
E-MAIL ADDRESS								
VAT REGISTRATION NUMBER								
SUPPLIER COMPLIANCE STATUS	TAX COMPLIANCE				CENTRAL SUPPLIER			
COMI LIANOL CTATOO	SYSTEM PIN:		OR		DATABASE			
ARE YOU THE					No:	MAAA		
ACCREDITED			ARF YOU	A FC	REIGN BASED			
REPRESENTATIVE IN SOUTH AFRICA FOR	☐Yes	□No	SUPPLIER	FOF	R THE GOODS		Yes	□No
THE GOODS		_	/SERVICE	SOF	·FEKEU?		[IF YES, ANSW	
/SERVICES OFFERED?	[IF YES ENCLO	SE PROOFJ					QUESTIONNAI	RE BELOW]
QUESTIONNAIRE TO BI	DDING FOREIGN	SUPPLIERS						
IS THE ENTITY A RESID	ENT OF THE REP	UBLIC OF SOUTH AFRIC	CA (RSA)?				☐ YE	S NO
DOES THE ENTITY HAV	E A BRANCH IN T	HE RSA?					☐ YE	S 🗌 NO
DOES THE ENTITY HAV	E A PERMANENT	ESTABLISHMENT IN TH	E RSA?				☐ YE	S NO
DOES THE ENTITY HAV	E ANY SOURCE C	OF INCOME IN THE RSA?	•				☐ YE	S NO
DOES THE ENTITY HAVE ANY SOURCE OF INCOME IN THE RSA? IS THE ENTITY LIABLE IN THE RSA FOR ANY FORM OF TAXATION? IF THE ANSWER IS "NO" TO ALL OF THE ABOVE, THEN IT IS NOT A REQUIREMENT TO REGISTER FOR A TAX COMPLIANCE STATUS SYSTEM PIN CODE FROM THE SOUTH AFRICAN REVENUE SERVICE (SARS) AND IF NOT REGISTER AS PER 2.3 BELOW.								

Version 1 of 2023

PART B TERMS AND CONDITIONS FOR BIDDING

1. BID SUBMISSION:

- 1.1. BIDS MUST BE DELIVERED BY THE STIPULATED TIME TO THE CORRECT ADDRESS. LATE BIDS WILL NOT BE ACCEPTED FOR CONSIDERATION.
- 1.2. ALL BIDS MUST BE SUBMITTED ON THE OFFICIAL FORMS PROVIDED (NOT TO BE RE-TYPED) OR IN THE MANNER PRESCRIBED IN THE BID DOCUMENT.
- 1.3. THIS BID IS SUBJECT TO THE PREFERENTIAL PROCUREMENT POLICY FRAMEWORK ACT, 2000 AND THE PREFERENTIAL PROCUREMENT REGULATIONS, THE GENERAL CONDITIONS OF CONTRACT (GCC) AND, IF APPLICABLE, ANY OTHER SPECIAL CONDITIONS OF CONTRACT.
- 1.4. THE SUCCESSFUL BIDDER WILL BE REQUIRED TO FILL IN AND SIGN A WRITTEN CONTRACT FORM (SBD7).

2. TAX COMPLIANCE REQUIREMENTS

- 2.1 BIDDERS MUST ENSURE COMPLIANCE WITH THEIR TAX OBLIGATIONS.
- 2.2 BIDDERS ARE REQUIRED TO SUBMIT THEIR UNIQUE PERSONAL IDENTIFICATION NUMBER (PIN) ISSUED BY SARS TO ENABLE THE ORGAN OF STATE TO VERIFY THE TAXPAYER'S PROFILE AND TAX STATUS.
- 2.3 APPLICATION FOR TAX COMPLIANCE STATUS (TCS) PIN MAY BE MADE VIA E-FILING THROUGH THE SARS WEBSITE WWW.SARS.GOV.ZA.
- 2.4 BIDDERS MAY ALSO SUBMIT A PRINTED TCS CERTIFICATE TOGETHER WITH THE BID.
- 2.5 IN BIDS WHERE CONSORTIA / JOINT VENTURES / SUB-CONTRACTORS ARE INVOLVED; EACH PARTY MUST SUBMIT A SEPARATE TCS CERTIFICATE / PIN / CSD NUMBER.
- 2.6 WHERE NO TCS PIN IS AVAILABLE BUT THE BIDDER IS REGISTERED ON THE CENTRAL SUPPLIER DATABASE (CSD), A CSD NUMBER MUST BE PROVIDED.
- 2.7 NO BIDS WILL BE CONSIDERED FROM PERSONS IN THE SERVICE OF THE STATE, COMPANIES WITH DIRECTORS WHO ARE PERSONS IN THE SERVICE OF THE STATE, OR CLOSE CORPORATIONS WITH MEMBERS PERSONS IN THE SERVICE OF THE STATE."

NB: FAILURE TO PROVIDE / OR COMPLY WITH ANY OF THE ABOVE PA	RTICULARS MAY RENDER THE BID INVALID.
SIGNATURE OF BIDDER:	
CAPACITY UNDER WHICH THIS BID IS SIGNED: (Proof of authority must be submitted e.g. company resolution)	
DATE:	

BID SIGNATURE AUTHORISATION (PBD3)

To confirm the authorised signatory for this bid

1. SINGLE BIDDING ENTERPRISE TYPE

Please indicate by ticking the appropria	te box:							
COMPANY	CLOSE CORPORATION INCORPO			RPORATED J\	1			
2. AUTHORISATION					-			
Single Bidding Enterprise Name:			Single Bidding Enterprise Registration Number:					
I/We, the undersigned, in our capacity as I hereby grant authority to the individual(s) (therefrom. Table 1 - Directors / Members / Owners	Table 2) to sig	n all documents pe	rtaining to th	,	•	•	•	
Name(s) (Print)				Signature)	Date	Date	
(Add rows if needed)								
Table 2 - Authorised Signatory/Signatory	ories							
Name(s) (Print) Authorised Signatory	ID numbe	er	Signature	Initial	Position of Author Signatory in Sing Enterprise		Date	

BID SIGNATURE AUTHORISATION (PBD3.1)

To confirm the authorised signatory for this bid

1. MULTI-ENTITY BIDDING ENTERPRISE TYPE

Please indicate by ticking the appropriate box:

DADTNEDQUID

PARTINERSHIP	JOINT VENTORE DIVINGO	MECKATED		CONSORTION		
2. AUTHORISATION						
Legal Name(s) of Entities in Multi-entity Bidding Enterprise	E	Registration Numbe Entities in Multi-ent Bidding Enterprise	tity			

IOINT VENTURE LININGORDORATED

I/We, the undersigned, in our capacity as Directors / Members / Owners / Partners (Table 1), duly authorised to represent the multi-entity bidding enterprise, hereby grant authority to the individual(s) (Table 2) to sign all documents pertaining to this bid on behalf of the said enterprise and any contract resulting therefrom.

Table 1 - Directors / Members / Owners / Partners of the Bidding Enterprise							
Name(s) (Print)	ID number	Signature	Bidding Entity Represented	Date			

(Add rows if needed)

CONSODTIUM

Table 2 - Authorised Signator	Table 2 - Authorised Signatory/Signatories							
Name(s) (Print) Authorised Signatory	ID number	Signature	Initial	Bidding Entity Represented	Position of Authorised Signatory in the Multi- entity Bidding Enterprise	Date		



PBD4.1

Private Bag X828, PRETORIA, 0001. DR AB Xuma Building, 1112 Voortrekker Road, Pretoria Townlands 351-JR, PRETORIA 0187. Directorate: Access to Affordable Medicines Tel: (012) 395 8130 Fax: (012) 395 8823/4

CONTRACT	T NUMB	ER:		-			人'	
SUPPLIER DETAILS:								
		Departments of pletion of Provi		quire separate reg ocuments.	gistration	of Supplie	ers on their	Databases
If a contract	is awar	ded, full detail fo	r supplier regi	stration or verifica	ation will	be reques	sted.	
		etail provided be ng with detail of s		lease advise the).	National	Departme	ent of Heal	:h
CONTACT	DETAIL			(2)	Ņ			
1. Supplier F Legal entity / with bank	corresp	onding						
2. Contact p	erson re	egarding contra	ct enquiries (to be printed on	contrac	t cover)		
Name & Surname			7.	e-mail		,		
Telephone			Ø√	Fax				
Cell		(ر ار	Other				
3. Contact re	egarding	g orders	•					
Address for p	oosting c	of order		Fax				
				Tel (confirmation)				
	1)		EDI				
Order	Name	& surname:		Tel				
enquiries	V			e-mail				
4. Nadonark	ey Acco	ount Manager (d	or Tender Mar	nager)				
Name				e-mail				
Telephone				Cell				

TENDER NO			LEGAL NAME OF SINGL	LE BIDDING ENTERPRISE				PBD9-1
Directors - Full Names	Surname	Nationality	Identity Number or Passport Number (foreigner)		Executive or Non-	Ownership or Director in related enterprise/s whether or not such enterprise/s are bidding in this tender? (Y/N)	If Yes, specify	
MARK WITH AN X IN THE OPEN C	ELL NEXT TO THE RELEVA	I NT SINGLE BIDDING EN	NTERPRISE					
CLOSE CORPORATION		COMPANY		INCORPORATED JOINT VENTURE		LISTED COMPANY		
OWNERSHIP IN THE SINGLE BIDD	ING ENTERPRISE HELD BY	HDI INDIVIDUALS						
Owners - (Individuals) Full Names	Surname	Nationality	RSA Identity Number or Foreign Passport Number	Race Categorisation	Is this owner HDI as per SRCC HDI definition YES/NO	Gender		% Shareholding in bidding enterprise
OWNERSHIP IN THE SINGLE BIDD	ING ENTERPRISE HELD BY	LEGAL ENTITIES						
Ownership - Legal Entities Name	Company Reg No	% Shareholding						

Authorised Signatory: Sign or Initial: ______ Page 1

Authorised Signatory: Sign or Initial _____

HP07-2026DAI

					ENTITY A (As per SBD1)		CSD REGISTRATION (MAAA NUMBER)		
TENDER NO		LEG		ULTI-ENTITY BIDDING RPRISE	ENTITY B		CSD REGISTRATION (MAAA NUMBER)		PBD9-2
					ENTITY C		CSD REGISTRATION (MAAA NUMBER)		
				Provide details of all dir	ectors for each entity in the b	pidding enterprise			
Directors - Full Names	Surname	Entity A,B or C	Nationality	Identity Number or Passport Number (foreigner)	Are you appointed as a Director? Y/N	Executive or Non-Executive director	Do you hold any directorships or ownership in other enterprises? (Y/N)	If Yes, specify	
ENTERPRISE CATEGORISATION:	SELECT AND MARK WITH A	N X, IN 7	THE OPEN CELI	NEXT TO THE RELEVAN	 	 G ENTERPRISE			
			IT VENTURE						
PARTNERSHIP			CORPORATED		CONSORTIUM		LISTED COMPANY		
OWNERSHIP IN THE MULTI-ENTIT	Y BIDDING ENTERPRISE HE	LD BY H	I INDIVIDUALS	•		•		1	

HP07-2026DAI

				ENTITY A (As per SBD1)		CSD REGISTRATION (MAAA NUMBER)		
TENDER NO			LEGAL NAMES OF MULTI-ENTITY BIDDING ENTERPRISE			CSD REGISTRATION (MAAA NUMBER)		PBD9-2
				ENTITY C		CSD REGISTRATION (MAAA NUMBER)		
			Provide details of all o	owners for each entity in the b	idding enterprise			
Owners - (Individuals) Full Names	Surname	RSA Identity Number or Foreign Passport Number	Race Categorisation	Is this owner HDI as per SRCC HDI definition YES/NO	Gender		% Shareholding in bidding enterprise	Nationality
		Provide	details of all Legal Entities	holding ownership in each e	ntity of the bidding enterprise			
OWNERSHIP IN THE MULTI-ENTIT	TY BIDDING ENTERPRISE HE	ELD BY LEGAL ENTITIES						
Ownership - Legal Entities Name	Company Reg No	% Shareholding	Entity A (As per SBD1)					
	0 0 0	0/ 01 1 1 1						
Ownership - Legal Entities Name	Company Reg No	% Shareholding	Entity B					
Ownership - Legal Entities Name	Company Reg No	% Shareholding	Entity C					

BIDDER'S DISCLOSURE

1. PURPOSE OF THE FORM

Any person (natural or juristic) may make an offer or offers in terms of this invitation to bid. In line with the principles of transparency, accountability, impartiality, and ethics as enshrined in the Constitution of the Republic of South Africa and further expressed in various pieces of legislation, it is required for the bidder to make this declaration in respect of the details required hereunder.

Where a person/s are listed in the Register for Tender Defaulters and / or the List of Restricted Suppliers, that person will automatically be disqualified from the bid process.

2. Bidder's declaration

- 2.1 Is the bidder, or any of its directors / trustees / shareholders / members / partners or any person having a controlling interest1 in the enterprise, employed by the state?

 YES/NO
- 2.1.1 If so, furnish particulars of the names, individual identity numbers, and, if applicable, state employee numbers of sole proprietor/ directors / trustees / shareholders / members/ partners or any person having a controlling interest in the enterprise, in table below.

Full Name	Identity Number	Name of institution	State

2.2 Do you, or any person connected with the bidder, have a relationship

¹ the power, by one person or a group of persons holding the majority of the equity of an enterprise, alternatively, the person/s having the deciding vote or power to influence or to direct the course and decisions of the enterprise.

BD4

	SI with any person who is employed by the procuring institution? YES/NO
2.2.1	If so, furnish particulars:
2.3	Does the bidder or any of its directors / trustees / shareholders / members / partners or any person having a controlling interest in the enterprise have any interest in any other related enterprise whether or not they are bidding for this contract? YES/NO
2.3.1	If so, furnish particulars:
3	DECLARATION
	I, the undersigned, (name)
3.1 3.2 3.3	I have read and I understand the contents of this disclosure; I understand that the accompanying bid will be disqualified if this disclosure is found not to be true and complete in every respect; The bidder has arrived at the accompanying bid independently from, and without consultation, communication, agreement or arrangement with any competitor. However, communication between partners in a joint
3.4	venture or consortium2 will not be construed as collusive bidding. In addition, there have been no consultations, communications, agreements or arrangements with any competitor regarding the quality, quantity, specifications, prices, including methods, factors or formulas used to calculate prices, market allocation, the intention or decision to submit or not to submit the bid, bidding with the intention not to win the bid and conditions or delivery particulars of the products or services to which this bid invitation relates.
3.4	The terms of the accompanying bid have not been, and will not be, disclosed by the bidder, directly or indirectly, to any competitor, prior to the date and time of the official bid opening or of the awarding of the contract.
3.5	There have been no consultations, communications, agreements or arrangements made by the bidder with any official of the procuring
	nt venture or Consortium means an association of persons for

the purpose of combining their expertise, property, capital, efforts, skill and knowledge in an activity for the execution of a contract.

SBD4

institution in relation to this procurement process prior to and during the bidding process except to provide clarification on the bid submitted where so required by the institution; and the bidder was not involved in the drafting of the specifications or terms of reference for this bid.

3.6 I am aware that, in addition and without prejudice to any other remedy provided to combat any restrictive practices related to bids and contracts, bids that are suspicious will be reported to the Competition Commission for investigation and possible imposition of administrative penalties in terms of section 59 of the Competition Act No 89 of 1998 and or may be reported to the National Prosecuting Authority (NPA) for criminal investigation and or may be restricted from conducting business with the public sector for a period not exceeding ten (10) years in terms of the Prevention and Combating of Corrupt Activities Act No 12 of 2004 or any other applicable legislation.

I CERTIFY THAT THE INFORMATION FURNISHED IN PARAGRAPHS 1. 2 and 3 ABOVE IS CORRECT.

I ACCEPT THAT THE STATE MAY REJECT THE BID OR ACT AGAINST ME IN TERMS OF PARAGRAPH 6 OF PFMA SCM INSTRUCTION 03 OF 2021/22 ON PREVENTING AND COMBATING ABUSE IN THE SUPPLY CHAIN MANAGEMENT SYSTEM SHOULD THIS DECLARATION PROVE TO BE FALSE.

Signature	Date
Position	Name of bidder

PBD 8

DECLARATION OF COMPLIANCE WITH THE SPECIAL REQUIREMENTS AND CONDITIONS OF CONTRACT & THE GENERAL CONDITIONS OF CONTRACT

To be signed by the appointed Authorised Signatory in terms of this bid.

l,				
,	(Full name)			
with the following ide	with the following identity number			
being the Authorised	being the Authorised Signatory of			
	(Organisation/Company	Legal Name)		
hereby declares that				
	(Organisation/Company	Legal Name)		
will comply with all th	ne requirements and condi	tions as stipulated in the Special		
Requirements and C	Conditions of Contract (SR	CC) and the General Conditions of		
Contract (GCC)				
Signature Authorised Signatory	(Signed at Location)	(on date)		
 Witness Signature	(Signed at Location)	(on date)		

PREFERENCE POINTS CLAIM FORM IN TERMS OF THE PREFERENTIAL PROCUREMENT REGULATIONS 2022

This preference form must form part of all tenders invited. It contains general information and serves as a claim form for preference points for specific goals.

NB: BEFORE COMPLETING THIS FORM, TENDERERS MUST STUDY THE GENERAL CONDITIONS, DEFINITIONS AND DIRECTIVES APPLICABLE IN RESPECT OF THE TENDER AND PREFERENTIAL PROCUREMENT REGULATIONS, 2022

1. GENERAL CONDITIONS

- 1.1 The following preference point systems are applicable to invitations to tender:
 - the 90/10 system for requirements with a Rand value above R50 000 000 (all applicable taxes included).
- 1.2 The applicable preference point system for this tender is the 90/10 preference point system.
- 1.3 Points for this tender (even in the case of a tender for income-generating contracts) shall be awarded for:
 - (a) Price; and
 - (b) Specific Goals.
- 1.4 The maximum points for this tender are allocated as follows:

	POINTS
PRICE	90
SPECIFIC GOALS	10
Total points for Price and SPECIFIC GOALS	100

- 1.5 Failure on the part of a tenderer to submit proof or documentation required in terms of this tender to claim points for specific goals with the tender, will be interpreted to mean that preference points for specific goals are not claimed.
- 1.6 The organ of state reserves the right to require of a tenderer, either before a tender is adjudicated or at any time subsequently, to substantiate any claim in regard to preferences, in any manner required by the organ of state.
- 1.7 The company must submit ID Copies of Director/Owner/Shareholder/Trustee and Beneficiary with their bid document to substantiate points claimed. The share certificate(s) reflecting the number of shares held by each Director/Owner/Shareholder/Trustee and Beneficiary must be submitted. In the case

of claiming points for disability the company must submit a registered Doctor's note or document as evidence of the disability

2. **DEFINITIONS**

- (a) "tender" means a written offer in the form determined by an organ of state in response to an invitation to provide goods or services through price quotations, competitive tendering process or any other method envisaged in legislation;
- (b) "price" means an amount of money tendered for goods or services, and includes all applicable taxes less all unconditional discounts;
- (c) "rand value" means the total estimated value of a contract in Rand, calculated at the time of bid invitation, and includes all applicable taxes;
- (d) "tender for income-generating contracts" means a written offer in the form determined by an organ of state in response to an invitation for the origination of income-generating contracts through any method envisaged in legislation that will result in a legal agreement between the organ of state and a third party that produces revenue for the organ of state, and includes, but is not limited to, leasing and disposal of assets and concession contracts, excluding direct sales and disposal of assets through public auctions; and
- (e) "the Act" means the Preferential Procurement Policy Framework Act, 2000 (Act No. 5 of 2000).

3. FORMULAE FOR PROCUREMENT OF GOODS AND SERVICES

3.1. POINTS AWARDED FOR PRICE

3.1.1 THE 90/10 PREFERENCE POINT SYSTEMS

A maximum of 90 points is allocated for price on the following basis:

90/10

$$Ps = 90\left(1 - \frac{Pt - Pmin}{Pmin}\right)$$

Where

Ps = Points scored for price of tender under consideration

Pt = Price of tender under consideration

Pmin = Price of lowest acceptable tender

3.2. FORMULAE FOR DISPOSAL OR LEASING OF STATE ASSETS AND INCOME GENERATING PROCUREMENT

3.2.1. POINTS AWARDED FOR PRICE

A maximum of 90 points is allocated for price on the following basis:

90/10

$$Ps = 90\left(1 + \frac{Pt - P max}{Pmax}\right)$$

Where

Ps = Points scored for price of tender under consideration

Pt = Price of tender under consideration

Pmax = Price of highest acceptable tender

4. POINTS AWARDED FOR SPECIFIC GOALS

4.1. In terms of Regulation 4(2); 5(2); 6(2) and 7(2) of the Preferential Procurement Regulations, preference points must be awarded for specific goals stated in the tender. For the purposes of this tender the tenderer will be allocated points based on the goals stated in table 1 below as may be supported by proof/ documentation stated in the conditions of this tender:

Table 1: Specific goals for the tender and points claimed are indicated per the table below.

Note

- The 90/10 preference point system is applicable, corresponding points must also be indicated as such.
- The tenderer must indicate how they claim points for each preference point system.

The specific goals allocated points in terms of this tender	Number of points claimable	Number of points claimed. (To be completed by the tenderer)	Percentage ownership equity claimed (To be completed by the tenderer)
HDI: No Franchise	4		
HDI: Women	2		
HDI: People with Disabilities	2		
RDP: The Promotion of South African owned enterprises	2		

DECLARATION WITH REGARD TO COMPANY/FIRM

4.2.	Name of company/firm
4.3.	Company registration number:
4.4.	TYPE OF COMPANY/ FIRM

4.5.

		nership/Joint Venture / Consortium e-person business/sole propriety
		se corporation
		lic Company
		sonal Liability Company) Limited
	Non	-Profit Company
□ [Tio		e Owned Company LICABLE BOX]
L	J. ()	
		ersigned, who is duly authorised to do so on behalf of the company/firm,
	•	It the points claimed, based on the specific goals as advised in the tender,
qua		he company/ firm for the preference(s) shown and I acknowledge that:
i)		formation furnished is true and correct;
ii)		reference points claimed are in accordance with the General Conditions as ited in paragraph 1 of this form;
iii)	in para	event of a contract being awarded as a result of points claimed as shown agraphs 1.4 and 4.2, the contractor may be required to furnish documentary to the satisfaction of the organ of state that the claims are correct;
iv)	of the	specific goals have been claimed or obtained on a fraudulent basis or any conditions of contract have not been fulfilled, the organ of state may, in on to any other remedy it may have –
	(a)	disqualify the person from the tendering process;
	(b)	recover costs, losses or damages it has incurred or suffered as a result of that person's conduct;
	(c)	cancel the contract and claim any damages which it has suffered as a result of having to make less favourable arrangements due to such cancellation;
	(d)	recommend that the tenderer or contractor, its shareholders and directors, or only the shareholders and directors who acted on a fraudulent basis, be restricted from obtaining business from any organ of state for a period not exceeding 10 years, after the <i>audi alteram partem</i> (hear the other side) rule has been applied; and
	(e)	forward the matter for criminal prosecution, if deemed necessary.
		SIGNATURE(S) OF TENDERER(S)
		Olonaronalo, or renderality
SURNA	ME AN	ND NAME:
DATE:		
ADDRE	SS:	

PBD 5

DECLARATION OF COMPLIANCE WITH GOOD MANUFACTURING PRACTICE (GMP)

To be signed by the appointed Authorised Signatory in terms of this bid.

I,		(Full name)	
with t	he following identity nur	nber	
being	g the Authorised Signato	•	
••••	(0	Organisation/Company Legal Na	 те)
hereb	by declares that to the b	est of my knowledge all rea	asonable steps have been
taken	to ensure that:		
a)		ing or impending GMP or I t on the Company's ability	
b)		the legal requirements as s I Substances Act 101 of 19	
c)		tion, I undertake to inform circumstances that may re	the Department of Health at sult in interrupted supply.
	Signature Authorised Signatory	(Signed at Location)	(on date)
	Witness Signature	(Signed at Location)	(on date)

This document must be signed and submitted together with your bid

THE NATIONAL INDUSTRIAL PARTICIPATION PROGRAMME

INTRODUCTION

The National Industrial Participation (NIP) Programme, which is applicable to all government procurement contracts that have an imported content, became effective on the 1 September 1996. The NIP policy and guidelines were fully endorsed by Cabinet on 30 April 1997. In terms of the Cabinet decision, all state and parastatal purchases / lease contracts (for goods, works and services) entered into after this date, are subject to the NIP requirements. NIP is obligatory and therefore must be complied with. The Industrial Participation Secretariat (IPS) of the Department of Trade and Industry (DTI) is charged with the responsibility of administering the programme.

1. PILLARS OF THE PROGRAMME

- 1.1 The NIP obligation is benchmarked on the imported content of the contract. Any contract having an imported content equal to or exceeding US\$ 10 million or other currency equivalent to US\$ 10 million will have a NIP obligation. This threshold of US\$ 10 million can be reached as follows:
 - a) Any single contract with imported content exceeding US\$10 million.
 - b) Multiple contracts for the same goods, works or services each with imported content exceeding US\$3 million awarded to one seller over a 2 year period which in total exceeds US\$10 million.
 - c) A contract with a renewable option clause, where should the option be exercised the total value of the imported content will exceed US\$10 million.
 - d) Multiple suppliers of the same goods, works or services under the same contract, where the value of the imported content of each allocation is equal to or exceeds US\$ 3 million worth of goods, works or services to the same government institution, which in total over a two (2) year period exceeds US\$10 million.
- 1.2 The NIP obligation applicable to suppliers in respect of sub-paragraphs 1.1 (a) to 1.1 (c) above will amount to 30% of the imported content whilst suppliers in respect of paragraph 1.1 (d) shall incur 30% of the total NIP obligation on a *pro-rata* basis.
- 1.3 To satisfy the NIP obligation, the DTI would negotiate and conclude agreements such as investments, joint ventures, sub-contracting, licensee production, export promotion, sourcing arrangements and research and development (R&D) with partners or suppliers.
- 1.4 A period of seven years has been identified as the time frame within which to discharge the obligation.

2 REQUIREMENTS OF THE DEPARTMENT OF TRADE AND INDUSTRY

- 2.1 In order to ensure effective implementation of the programme, successful bidders (contractors) are required to, immediately after the award of a contract that is in excess of R10 million (ten million Rands), submit details of such a contract to the DTI for reporting purposes.
- 2.2 The purpose for reporting details of contracts in excess of the amount of R10 million (ten million Rands) is to cater for multiple contracts for the same goods, works or services; renewable contracts and multiple suppliers for the same goods, works or services under the same contract as provided for in paragraphs 1.1.(b) to 1.1. (d) above.

3 BID SUBMISSION AND CONTRACT REPORTING REQUIREMENTS OF BIDDERS AND SUCCESSFUL BIDDERS (CONTRACTORS)

- 3.1 Bidders are required to sign and submit this Standard Bidding Document (SBD 5) together with the bid on the closing date and time.
- 3.2 In order to accommodate multiple contracts for the same goods, works or services; renewable contracts and multiple suppliers for the same goods, works or services under the same contract as indicated in sub-paragraphs 1.1 (b) to 1.1 (d) above and to enable the DTI in determining the NIP obligation, successful bidders (contractors) are required, immediately after being officially notified about any successful bid with a value in excess of R10 million (ten million Rands), to contact and furnish the DTI with the following information:
 - Bid/contract number.
 - Description of the goods works or services.
 - Date on which the contract was accepted.
 - Name, address and contact details of the government institution.
 - Value of the contract.
 - Imported content of the contract, if possible.
- 3.3 The information required in paragraph 3.2 above must be sent to the Department of Trade and Industry, Private Bag X84, Pretoria, 0001 for the attention of Mr Elias Malapane within five (5) working days after award of the contract. for further details about the programme, contact Ms R Muthan on telephone (012) 394 1288, Mobile (066) 301 2051 or e-mail at amuthan@thedtic.gov.za.

4 PROCESS TO SATISFY THE NIP OBLIGATION

- 4.1 Once the successful bidder (contractor) has made contact with and furnished the DTI with the information required, the following steps will be followed:
 - a) the contractor and the DTI will determine the NIP obligation;
 - b) the contractor and the DTI will sign the NIP obligation agreement;
 - c) the contractor will submit a performance guarantee to the DTI;
 - d) the contractor will submit a business concept for consideration and approval by the DTI;
 - e) upon approval of the business concept by the DTI, the contractor will submit detailed business plans outlining the business concepts;
 - f) the contractor will implement the business plans; and
 - g) the contractor will submit bi-annual progress reports on approved plans to the DTI.
- 4.2 The NIP obligation agreement is between the DTI and the successful bidder (contractor) and, therefore, does not involve the purchasing institution.

Bid number	Closing date:	
Name of bidder		
Postal Address		
	Postal Code	
Name in print		
Position		
Signature:	Date:	

Authorised Signatory: Sign or In	iitial
----------------------------------	--------

PBD1



AUTHORISATION DECLARATION (PBD1)

	NAME O	F THE BID	DER						
4	Are you	sourcing t	the produ	ıcts from a thir	d party?	Υ	es	No	
	-			to the above qu			details in th	e table below of	
1.		•		where the bidde	-	e products fro	om a third p	arty.	•
	1 ne b		•	es the following:		the DDD1 1	attached t	from a third party in	
	1.1			th the terms and			allacrieu, i	from a third party in	
	1.2						ditions of th	e bid and the third par	ty
								sted in the PBD1.1.	
	1.3	The bide	der has r	eceived the att	ached, uncond	itional writte	n undertak	ing from the third par	ty
		to supply	the prod	ucts listed in the	e PBD1.1 in acc	cordance wit	h the terms	and conditions of the	
							been attacl	ned (PBD1.2) that is to	
			-	rpose of the thir		•			
	1.4					•	nents for t	he products have bee	en
		mutually	agreed u	pon between the	e blader and the	tnira party.			
2.	The b	oidder decl	ares that	the information of	contained herein	is true and	correct.		
3.	The	bidder a	cknowledo	ges that the	Department of	Health res	serves the	right to verify the	
	infor	mation con	tained the	erein and if four	nd to be false or	r incorrect m	ay invoke a	any remedies available	
	to it	in the bid o	documents	S.					
_						T	T	T	
S	igned at				on the		day of		
F	ull Names	6			I	ı			
D	Designation								
S	ignature								

List of the products offered sourced from third party

Bid Item No	Brand Name	Name of the company from where the products will be sourced	Address and contact details of the company from where the products will be sourced

List of the products offered sourced from third party - continued

Bid Item No	Brand Name	Name of the company from where the products will be sourced	Address and contact details of the company from where the products will be sourced

(Should the table provided not be sufficient for all the items offered, please provide additional information as an attachment and it must be properly referenced to this document)

Template for unconditional written undertaking from the third party

Note:		
The authorisation le	tter must be on the official letterhead of the third	party
A separate letter mu	ust be included for each third party	
The authorisation le	tter must be addressed to the Bidding Company	′
Name of Bidding Co	ompany:	
Address of Bidding	Company:	
Dear Sir/Madam		
AUTHORISATION	LETTER: CONTRACT NO	_
We,		Name of Third Party)
hereby authorise yo	ou, <i>(</i>	Name of Company) to include the
products listed belo	w in your bid submission for the abovementione	d contract.
We confirm that we	have firm supply arrangements in place, and	nave familiarised ourselves with the
item descriptions, s	pecifications and bid conditions relating to item/s	s listed below.
Item no.	Description of product	Brand name
(Should the table p	rovided not be sufficient for all the items offere	d, please provide additional
information as an a	ttachment and it must be properly referenced to	this document)
Yours faithfully,		
Signature of the Thi	ird Party:	Date:
orginature of the Thi	iu i aity.	Date.

Definition of fields in Bid Res Conditions of Contract	sponse Document, to be read in conjunction with the Special Requirements and
Field Name	Field Definition
FIELI	DS WHICH ARE PRE-FILLED AND MAY NOT BE ALTERED
Item Number	The relevant item number which will be used throughout the contract period. Each item number is linked to a specification
Item Specification	The specification of the item for which a call for bids has been issued, as linked to the item number.
Unit	The unit of measure for the specification. This determines how the estimates are expressed and how the price should be quoted. This may be one injection or one pack of 100 tablets, etc.
Estimate	The estimated quantities associated with the item number and specification, for the full contract period. Estimates are expressed in unit packs.
FIELDS WHICH ARE T	O BE COMPLETED BY THE BIDDER FOR ALL ITEMS ON WHICH BIDS ARE OFFERED
Registered legal name of bidder	The full, registered, legal name of the bidder, as on VAT registration certificate and Medicine Registration Certificate applicant.
Quantity for full period	The volume of the item (expressed in units) which the bidder can provide during the complete period of the tender
Delivered price in ZAR	Final price offered by a bidder for an item number as per specification, which includes VAT and delivery. Must be the price for a unit as advertised .
Registered Product Name	Brand name. Must correspond with Medicine Registration Certificate (1) GW12/7
Conforms to specification?	Confirm whether or not the product on offer conforms exactly to the Item Specification.
If NO : Detail deviation from specification.	Detail exact deviation from Item Specification, as per registration of product on offer.
Product Registration Number	As per Medicine Registration Certificate Certificate(2) GW12/7
License to Manufacture Medicines: License Number , Expiry date	As per License to Manufacture Medicines – this must correspond with the document submitted
Pack Size Offered: Unit pack	Single unit offered according to specification in numbers e.g. each (1) This must correspond with the delivered price.
Pack Size Offered: Shelf Pack	Number of Unit Packs within the smallest wrap (e.g. 10 ampoules)
Standard units in: Shipper Pack	Number of unit packs in a shipper / bulk box
Lead-Time	Interval between receipt of an order until delivery at facility which placed the order. Must not exceed 14 calendar days.
Initial lead time	Interval between award of the tender and ability to fill an order. This must not exceed 75 calendar days.
Minimum Order Quantity	The lowest acceptable quantity for a given purchase order.
Batch size for the bid item, in number of packs	Batch size, expressed in number of units
Monthly batch capacity	Monthly batch capacity that will be assigned for the bid item for the duration of the contract expressed in number of batches.
Technical amendment required?	Do you require a technical amendment to perform according to the conditions of your bid Y/N
If YES: Provide details	Provide all relevant details (can be provided in a covering letter)

Definition of fields in Bid Response Document, to be read in conjunction with the Special Requirements and Conditions of Contract			
Field Name	Field Definition		
EAN 13 Barcode for Unit Pack	Provide Number		
EAN 13 Barcode for Shelf Pack	Provide Number		
ITF14 Barcode for Shipper Pack	Provide Number		
2D Barcode or Similar	Provide Number		
NAPPI Code	Provide Code		
Manufacturer	As per MCC Certificate (8) GW12/7 – List all sources		
SEP Price	The most recently approved Single Exit Price expressed in corresponding unit to bid		
Are any of the listed manufacturers etc. 3rd parties to the bidder?	Y/N If YES - complete PBD1 and include letter(s) of authorisation as applicable		
API Source Full Site Name (x3)	Full name of API source, including company name and site – List all sources		
API Source Full Address	Full physical address of API source – List all sources		
API Source Country	Country of API source – List all sources		
API Source Contact	Listed contact information		
Note: VAT must be apportio	PRICING COMPONENT BREAKDOWN ned equally across all components. Please see pricing section in Special Conditions		
Percentage of Delivered Price attributable to API	The percentage of the Delivered Price associated with API, (the therapeutically active component of the medicine). Should an item be imported as finished product, the component may be reflected as part of formulation cost.		
Imported (API)	Portion of API component attributable to imported expenditure		
Percentage of Delivered Price attributable to Formulation	The percentage of the delivered price associated with Formulation, (includes all operations in the process of which different chemical substances, including the API, are combined to produce a final medicinal product), includes material, processing, production, quality assurance and related controls.		
Local (Formulation)	Portion of Formulation component attributable to local expenditure		
Imported (Formulation)	Portion of Formulation component attributable to imported expenditure		
Packaging	The percentage of the Delivered Price associated with Packaging, where packaging includes all operations in the process of packaging medicine into primary and/or secondary packaging, packaging material and labels.		
Local (Packaging)	Portion of Packaging component attributable to local expenditure		
Imported (Packaging)	Portion of Packaging component attributable to imported expenditure		
Logistics	Percentage of delivered price associated with logistics, where logistics includes all operations, taking place within the Republic of South Africa, relating to the storage, distribution and transportation of medicine to the healthcare facility or pharmaceutical depot.		
Gross Margin	Percentage of delivered price not associated with API, Formulation, Packaging, or Logistics.		
Currency	Primary currency in which manufacturer trades for imported components		

THE NATIONAL TREASURY

Republic of South Africa



GOVERNMENT PROCUREMENT: GENERAL CONDITIONS OF CONTRACT

July 2010

GOVERNMENT PROCUREMENT

GENERAL CONDITIONS OF CONTRACT July 2010

NOTES

The purpose of this document is to:

- (i) Draw special attention to certain general conditions applicable to government bids, contracts and orders; and
- (ii) To ensure that clients be familiar with regard to the rights and obligations of all parties involved in doing business with government.

In this document words in the singular also mean in the plural and vice versa and words in the masculine also mean in the feminine and neuter.

- The General Conditions of Contract will form part of all bid documents and may not be amended.
- Special Conditions of Contract (SCC) relevant to a specific bid, should be compiled separately for every bid (if (applicable) and will supplement the General Conditions of Contract. Whenever there is a conflict, the provisions in the SCC shall prevail.

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General Conditions of Contract

1. Definitions

- 1. The following terms shall be interpreted as indicated:
- 1.1 "Closing time" means the date and hour specified in the bidding documents for the receipt of bids.
- 1.2 "Contract" means the written agreement entered into between the purchaser and the supplier, as recorded in the contract form signed by the parties, including all attachments and appendices thereto and all documents incorporated by reference therein.
- 1.3 "Contract price" means the price payable to the supplier under the contract for the full and proper performance of his contractual obligations.
- 1.4 "Corrupt practice" means the offering, giving, receiving, or soliciting of any thing of value to influence the action of a public official in the procurement process or in contract execution.
- 1.5 "Countervailing duties" are imposed in cases where an enterprise abroad is subsidized by its government and encouraged to market its products internationally.
- 1.6 "Country of origin" means the place where the goods were mined, grown or produced or from which the services are supplied. Goods are produced when, through manufacturing, processing or substantial and major assembly of components, a commercially recognized new product results that is substantially different in basic characteristics or in purpose or utility from its components.
- 1.7 "Day" means calendar day.
- 1.8 "Delivery" means delivery in compliance of the conditions of the contract or order.
- 1.9 "Delivery ex stock" means immediate delivery directly from stock actually on hand.
- 1.10 "Delivery into consignees store or to his site" means delivered and unloaded in the specified store or depot or on the specified site in compliance with the conditions of the contract or order, the supplier bearing all risks and charges involved until the supplies are so delivered and a valid receipt is obtained.
- 1.11 "Dumping" occurs when a private enterprise abroad market its goods on own initiative in the RSA at lower prices than that of the country of origin and which have the potential to harm the local industries in the

RSA.

- 1.12 "Force majeure" means an event beyond the control of the supplier and not involving the supplier's fault or negligence and not foreseeable. Such events may include, but is not restricted to, acts of the purchaser in its sovereign capacity, wars or revolutions, fires, floods, epidemics, quarantine restrictions and freight embargoes.
- 1.13 "Fraudulent practice" means a misrepresentation of facts in order to influence a procurement process or the execution of a contract to the detriment of any bidder, and includes collusive practice among bidders (prior to or after bid submission) designed to establish bid prices at artificial non-competitive levels and to deprive the bidder of the benefits of free and open competition.
- 1.14 "GCC" means the General Conditions of Contract.
- 1.15 "Goods" means all of the equipment, machinery, and/or other materials that the supplier is required to supply to the purchaser under the contract.
- 1.16 "Imported content" means that portion of the bidding price represented by the cost of components, parts or materials which have been or are still to be imported (whether by the supplier or his subcontractors) and which costs are inclusive of the costs abroad, plus freight and other direct importation costs such as landing costs, dock dues, import duty, sales duty or other similar tax or duty at the South African place of entry as well as transportation and handling charges to the factory in the Republic where the supplies covered by the bid will be manufactured.
- 1.17 "Local content" means that portion of the bidding price which is not included in the imported content provided that local manufacture does take place.
- 1.18 "Manufacture" means the production of products in a factory using labour, materials, components and machinery and includes other related value-adding activities.
- 1.19 "Order" means an official written order issued for the supply of goods or works or the rendering of a service.
- 1.20 "Project site," where applicable, means the place indicated in bidding documents.
- 1.21 "Purchaser" means the organization purchasing the goods.
- 1.22 "Republic" means the Republic of South Africa.
- 1.23 "SCC" means the Special Conditions of Contract.
- 1.24 "Services" means those functional services ancillary to the supply of the goods, such as transportation and any other incidental services, such as installation, commissioning, provision of technical assistance, training, catering, gardening, security, maintenance and other such

obligations of the supplier covered under the contract.

1.25 "Written" or "in writing" means handwritten in ink or any form of electronic or mechanical writing.

2. Application

- 2.1 These general conditions are applicable to all bids, contracts and orders including bids for functional and professional services, sales, hiring, letting and the granting or acquiring of rights, but excluding immovable property, unless otherwise indicated in the bidding documents.
- 2.2 Where applicable, special conditions of contract are also laid down to cover specific supplies, services or works.
- 2.3 Where such special conditions of contract are in conflict with these general conditions, the special conditions shall apply.

3. General

- 3.1 Unless otherwise indicated in the bidding documents, the purchaser shall not be liable for any expense incurred in the preparation and submission of a bid. Where applicable a non-refundable fee for documents may be charged.
- 3.2 With certain exceptions, invitations to bid are only published in the Government Tender Bulletin. The Government Tender Bulletin may be obtained directly from the Government Printer, Private Bag X85, Pretoria 0001, or accessed electronically from www.treasury.gov.za

4. Standards

- 4.1 The goods supplied shall conform to the standards mentioned in the bidding documents and specifications.
- 5. Use of contract documents and information; inspection.
- 5.1 The supplier shall not, without the purchaser's prior written consent, disclose the contract, or any provision thereof, or any specification, plan, drawing, pattern, sample, or information furnished by or on behalf of the purchaser in connection therewith, to any person other than a person employed by the supplier in the performance of the contract. Disclosure to any such employed person shall be made in confidence and shall extend only so far as may be necessary for purposes of such performance.
- 5.2 The supplier shall not, without the purchaser's prior written consent, make use of any document or information mentioned in GCC clause 5.1 except for purposes of performing the contract.
- 5.3 Any document, other than the contract itself mentioned in GCC clause 5.1 shall remain the property of the purchaser and shall be returned (all copies) to the purchaser on completion of the supplier's performance under the contract if so required by the purchaser.
- 5.4 The supplier shall permit the purchaser to inspect the supplier's records relating to the performance of the supplier and to have them audited by auditors appointed by the purchaser, if so required by the purchaser.

6. Patent rights

6.1 The supplier shall indemnify the purchaser against all third-party claims of infringement of patent, trademark, or industrial design rights arising from use of the goods or any part thereof by the purchaser.

7. Performance security

- 7.1 Within thirty (30) days of receipt of the notification of contract award, the successful bidder shall furnish to the purchaser the performance security of the amount specified in SCC.
- 7.2 The proceeds of the performance security shall be payable to the purchaser as compensation for any loss resulting from the supplier's failure to complete his obligations under the contract.
- 7.3 The performance security shall be denominated in the currency of the contract, or in a freely convertible currency acceptable to the purchaser and shall be in one of the following forms:
 - (a) a bank guarantee or an irrevocable letter of credit issued by a reputable bank located in the purchaser's country or abroad, acceptable to the purchaser, in the form provided in the bidding documents or another form acceptable to the purchaser; or
 - (b) a cashier's or certified cheque
- 7.4 The performance security will be discharged by the purchaser and returned to the supplier not later than thirty (30) days following the date of completion of the supplier's performance obligations under the contract, including any warranty obligations, unless otherwise specified in SCC.

8. Inspections, tests and analyses

- 8.1 All pre-bidding testing will be for the account of the bidder.
- 8.2 If it is a bid condition that supplies to be produced or services to be rendered should at any stage during production or execution or on completion be subject to inspection, the premises of the bidder or contractor shall be open, at all reasonable hours, for inspection by a representative of the Department or an organization acting on behalf of the Department.
- 8.3 If there are no inspection requirements indicated in the bidding documents and no mention is made in the contract, but during the contract period it is decided that inspections shall be carried out, the purchaser shall itself make the necessary arrangements, including payment arrangements with the testing authority concerned.
- 8.4 If the inspections, tests and analyses referred to in clauses 8.2 and 8.3 show the supplies to be in accordance with the contract requirements, the cost of the inspections, tests and analyses shall be defrayed by the purchaser.
- 8.5 Where the supplies or services referred to in clauses 8.2 and 8.3 do not comply with the contract requirements, irrespective of whether such supplies or services are accepted or not, the cost in connection with these inspections, tests or analyses shall be defrayed by the supplier.
- 8.6 Supplies and services which are referred to in clauses 8.2 and 8.3 and which do not comply with the contract requirements may be rejected.
- 8.7 Any contract supplies may on or after delivery be inspected, tested or

analyzed and may be rejected if found not to comply with the requirements of the contract. Such rejected supplies shall be held at the cost and risk of the supplier who shall, when called upon, remove them immediately at his own cost and forthwith substitute them with supplies which do comply with the requirements of the contract. Failing such removal the rejected supplies shall be returned at the suppliers cost and risk. Should the supplier fail to provide the substitute supplies forthwith, the purchaser may, without giving the supplier further opportunity to substitute the rejected supplies, purchase such supplies as may be necessary at the expense of the supplier.

8.8 The provisions of clauses 8.4 to 8.7 shall not prejudice the right of the purchaser to cancel the contract on account of a breach of the conditions thereof, or to act in terms of Clause 23 of GCC.

9. Packing

- 9.1 The supplier shall provide such packing of the goods as is required to prevent their damage or deterioration during transit to their final destination, as indicated in the contract. The packing shall be sufficient to withstand, without limitation, rough handling during transit and exposure to extreme temperatures, salt and precipitation during transit, and open storage. Packing, case size and weights shall take into consideration, where appropriate, the remoteness of the goods' final destination and the absence of heavy handling facilities at all points in transit.
- 9.2 The packing, marking, and documentation within and outside the packages shall comply strictly with such special requirements as shall be expressly provided for in the contract, including additional requirements, if any, specified in SCC, and in any subsequent instructions ordered by the purchaser.

10. Delivery and documents

- 10.1 Delivery of the goods shall be made by the supplier in accordance with the terms specified in the contract. The details of shipping and/or other documents to be furnished by the supplier are specified in SCC.
- 10.2 Documents to be submitted by the supplier are specified in SCC.

11. Insurance

11.1 The goods supplied under the contract shall be fully insured in a freely convertible currency against loss or damage incidental to manufacture or acquisition, transportation, storage and delivery in the manner specified in the SCC.

12. Transportation

12.1 Should a price other than an all-inclusive delivered price be required, this shall be specified in the SCC.

13. Incidental services

- 13.1 The supplier may be required to provide any or all of the following services, including additional services, if any, specified in SCC:
 - (a) performance or supervision of on-site assembly and/or commissioning of the supplied goods;
 - (b) furnishing of tools required for assembly and/or maintenance of the supplied goods;
 - (c) furnishing of a detailed operations and maintenance manual for each appropriate unit of the supplied goods;

- (d) performance or supervision or maintenance and/or repair of the supplied goods, for a period of time agreed by the parties, provided that this service shall not relieve the supplier of any warranty obligations under this contract; and
- (e) training of the purchaser's personnel, at the supplier's plant and/or on-site, in assembly, start-up, operation, maintenance, and/or repair of the supplied goods.
- 13.2 Prices charged by the supplier for incidental services, if not included in the contract price for the goods, shall be agreed upon in advance by the parties and shall not exceed the prevailing rates charged to other parties by the supplier for similar services.

14. Spare parts

- 14.1 As specified in SCC, the supplier may be required to provide any or all of the following materials, notifications, and information pertaining to spare parts manufactured or distributed by the supplier:
 - (a) such spare parts as the purchaser may elect to purchase from the supplier, provided that this election shall not relieve the supplier of any warranty obligations under the contract; and
 - (b) in the event of termination of production of the spare parts:
 - (i) Advance notification to the purchaser of the pending termination, in sufficient time to permit the purchaser to procure needed requirements; and
 - (ii)following such termination, furnishing at no cost to the purchaser, the blueprints, drawings, and specifications of the spare parts, if requested.

15. Warranty

- 15.1 The supplier warrants that the goods supplied under the contract are new, unused, of the most recent or current models, and that they incorporate all recent improvements in design and materials unless provided otherwise in the contract. The supplier further warrants that all goods supplied under this contract shall have no defect, arising from design, materials, or workmanship (except when the design and/or material is required by the purchaser's specifications) or from any act or omission of the supplier, that may develop under normal use of the supplied goods in the conditions prevailing in the country of final destination.
- 15.2 This warranty shall remain valid for twelve (12) months after the goods, or any portion thereof as the case may be, have been delivered to and accepted at the final destination indicated in the contract, or for eighteen (18) months after the date of shipment from the port or place of loading in the source country, whichever period concludes earlier, unless specified otherwise in SCC.
- 15.3 The purchaser shall promptly notify the supplier in writing of any claims arising under this warranty.
- 15.4 Upon receipt of such notice, the supplier shall, within the period specified in SCC and with all reasonable speed, repair or replace the defective goods or parts thereof, without costs to the purchaser.
- 15.5 If the supplier, having been notified, fails to remedy the defect(s) within the period specified in SCC, the purchaser may proceed to take

such remedial action as may be necessary, at the supplier's risk and expense and without prejudice to any other rights which the purchaser may have against the supplier under the contract.

16. Payment

- 16.1 The method and conditions of payment to be made to the supplier under this contract shall be specified in SCC.
- 16.2 The supplier shall furnish the purchaser with an invoice accompanied by a copy of the delivery note and upon fulfillment of other obligations stipulated in the contract.
- 16.3 Payments shall be made promptly by the purchaser, but in no case later than thirty (30) days after submission of an invoice or claim by the supplier.
- 16.4 Payment will be made in Rand unless otherwise stipulated in SCC.

17. Prices

17.1 Prices charged by the supplier for goods delivered and services performed under the contract shall not vary from the prices quoted by the supplier in his bid, with the exception of any price adjustments authorized in SCC or in the purchaser's request for bid validity extension, as the case may be.

18. Contract amendments

- 18.1 No variation in or modification of the terms of the contract shall be made except by written amendment signed by the parties concerned.
- 19. Assignment
- 19.1 The supplier shall not assign, in whole or in part, its obligations to perform under the contract, except with the purchaser's prior written consent.

20. Subcontracts

20.1 The supplier shall notify the purchaser in writing of all subcontracts awarded under this contracts if not already specified in the bid. Such notification, in the original bid or later, shall not relieve the supplier from any liability or obligation under the contract.

21. Delays in the supplier's performance

- 21.1 Delivery of the goods and performance of services shall be made by the supplier in accordance with the time schedule prescribed by the purchaser in the contract.
- 21.2 If at any time during performance of the contract, the supplier or its subcontractor(s) should encounter conditions impeding timely delivery of the goods and performance of services, the supplier shall promptly notify the purchaser in writing of the fact of the delay, its likely duration and its cause(s). As soon as practicable after receipt of the supplier's notice, the purchaser shall evaluate the situation and may at his discretion extend the supplier's time for performance, with or without the imposition of penalties, in which case the extension shall be ratified by the parties by amendment of contract.
- 21.3 No provision in a contract shall be deemed to prohibit the obtaining of supplies or services from a national department, provincial department, or a local authority.
- 21.4 The right is reserved to procure outside of the contract small quantities or to have minor essential services executed if an emergency arises, the

- supplier's point of supply is not situated at or near the place where the supplies are required, or the supplier's services are not readily available.
- 21.5 Except as provided under GCC Clause 25, a delay by the supplier in the performance of its delivery obligations shall render the supplier liable to the imposition of penalties, pursuant to GCC Clause 22, unless an extension of time is agreed upon pursuant to GCC Clause 21.2 without the application of penalties.
- 21.6 Upon any delay beyond the delivery period in the case of a supplies contract, the purchaser shall, without canceling the contract, be entitled to purchase supplies of a similar quality and up to the same quantity in substitution of the goods not supplied in conformity with the contract and to return any goods delivered later at the supplier's expense and risk, or to cancel the contract and buy such goods as may be required to complete the contract and without prejudice to his other rights, be entitled to claim damages from the supplier.

22. Penalties

22.1 Subject to GCC Clause 25, if the supplier fails to deliver any or all of the goods or to perform the services within the period(s) specified in the contract, the purchaser shall, without prejudice to its other remedies under the contract, deduct from the contract price, as a penalty, a sum calculated on the delivered price of the delayed goods or unperformed services using the current prime interest rate calculated for each day of the delay until actual delivery or performance. The purchaser may also consider termination of the contract pursuant to GCC Clause 23.

23. Termination for default

- 23.1 The purchaser, without prejudice to any other remedy for breach of contract, by written notice of default sent to the supplier, may terminate this contract in whole or in part:
 - (a) if the supplier fails to deliver any or all of the goods within the period(s) specified in the contract, or within any extension thereof granted by the purchaser pursuant to GCC Clause 21.2;
 - (b) if the Supplier fails to perform any other obligation(s) under the contract; or
 - (c) if the supplier, in the judgment of the purchaser, has engaged in corrupt or fraudulent practices in competing for or in executing the contract.
- 23.2 In the event the purchaser terminates the contract in whole or in part, the purchaser may procure, upon such terms and in such manner as it deems appropriate, goods, works or services similar to those undelivered, and the supplier shall be liable to the purchaser for any excess costs for such similar goods, works or services. However, the supplier shall continue performance of the contract to the extent not terminated.
- 23.3 Where the purchaser terminates the contract in whole or in part, the purchaser may decide to impose a restriction penalty on the supplier by prohibiting such supplier from doing business with the public sector for a period not exceeding 10 years.
- 23.4 If a purchaser intends imposing a restriction on a supplier or any

person associated with the supplier, the supplier will be allowed a time period of not more than fourteen (14) days to provide reasons why the envisaged restriction should not be imposed. Should the supplier fail to respond within the stipulated fourteen (14) days the purchaser may regard the intended penalty as not objected against and may impose it on the supplier.

- 23.5 Any restriction imposed on any person by the Accounting Officer / Authority will, at the discretion of the Accounting Officer / Authority, also be applicable to any other enterprise or any partner, manager, director or other person who wholly or partly exercises or exercised or may exercise control over the enterprise of the first-mentioned person, and with which enterprise or person the first-mentioned person, is or was in the opinion of the Accounting Officer / Authority actively associated.
- 23.6 If a restriction is imposed, the purchaser must, within five (5) working days of such imposition, furnish the National Treasury, with the following information:
 - (i) the name and address of the supplier and / or person restricted by the purchaser;
 - (ii) the date of commencement of the restriction
 - (iii) the period of restriction; and
 - (iv) the reasons for the restriction.

These details will be loaded in the National Treasury's central database of suppliers or persons prohibited from doing business with the public sector.

- 23.7 If a court of law convicts a person of an offence as contemplated in sections 12 or 13 of the Prevention and Combating of Corrupt Activities Act, No. 12 of 2004, the court may also rule that such person's name be endorsed on the Register for Tender Defaulters. When a person's name has been endorsed on the Register, the person will be prohibited from doing business with the public sector for a period not less than five years and not more than 10 years. The National Treasury is empowered to determine the period of restriction and each case will be dealt with on its own merits. According to section 32 of the Act the Register must be open to the public. The Register can be perused on the National Treasury website.
- 24. Anti-dumping and countervailing duties and rights
- 24.1 When, after the date of bid, provisional payments are required, or antidumping or countervailing duties are imposed, or the amount of a
 provisional payment or anti-dumping or countervailing right is
 increased in respect of any dumped or subsidized import, the State is
 not liable for any amount so required or imposed, or for the amount of
 any such increase. When, after the said date, such a provisional
 payment is no longer required or any such anti-dumping or
 countervailing right is abolished, or where the amount of such
 provisional payment or any such right is reduced, any such favourable
 difference shall on demand be paid forthwith by the contractor to the
 State or the State may deduct such amounts from moneys (if any)
 which may otherwise be due to the contractor in regard to supplies or
 services which he delivered or rendered, or is to deliver or render in
 terms of the contract or any other contract or any other amount which

may be due to him

25. Force Majeure

- 25.1 Notwithstanding the provisions of GCC Clauses 22 and 23, the supplier shall not be liable for forfeiture of its performance security, damages, or termination for default if and to the extent that his delay in performance or other failure to perform his obligations under the contract is the result of an event of force majeure.
- 25.2 If a force majeure situation arises, the supplier shall promptly notify the purchaser in writing of such condition and the cause thereof. Unless otherwise directed by the purchaser in writing, the supplier shall continue to perform its obligations under the contract as far as is reasonably practical, and shall seek all reasonable alternative means for performance not prevented by the force majeure event.

26. Termination for insolvency

26.1 The purchaser may at any time terminate the contract by giving written notice to the supplier if the supplier becomes bankrupt or otherwise insolvent. In this event, termination will be without compensation to the supplier, provided that such termination will not prejudice or affect any right of action or remedy which has accrued or will accrue thereafter to the purchaser.

27. Settlement of Disputes

- 27.1 If any dispute or difference of any kind whatsoever arises between the purchaser and the supplier in connection with or arising out of the contract, the parties shall make every effort to resolve amicably such dispute or difference by mutual consultation.
- 27.2 If, after thirty (30) days, the parties have failed to resolve their dispute or difference by such mutual consultation, then either the purchaser or the supplier may give notice to the other party of his intention to commence with mediation. No mediation in respect of this matter may be commenced unless such notice is given to the other party.
- 27.3 Should it not be possible to settle a dispute by means of mediation, it may be settled in a South African court of law.
- 27.4 Mediation proceedings shall be conducted in accordance with the rules of procedure specified in the SCC.
- 27.5 Notwithstanding any reference to mediation and/or court proceedings herein.
 - (a) the parties shall continue to perform their respective obligations under the contract unless they otherwise agree; and
 - (b) the purchaser shall pay the supplier any monies due the supplier.

28. Limitation of liability

- 28.1 Except in cases of criminal negligence or willful misconduct, and in the case of infringement pursuant to Clause 6;
 - (a) the supplier shall not be liable to the purchaser, whether in contract, tort, or otherwise, for any indirect or consequential loss or damage, loss of use, loss of production, or loss of profits or interest costs, provided that this exclusion shall not apply to any obligation of the supplier to pay penalties and/or damages to the purchaser; and

(b) the aggregate liability of the supplier to the purchaser, whether under the contract, in tort or otherwise, shall not exceed the total contract price, provided that this limitation shall not apply to the cost of repairing or replacing defective equipment.

29. Governing language

29.1 The contract shall be written in English. All correspondence and other documents pertaining to the contract that is exchanged by the parties shall also be written in English.

30. Applicable law

30.1 The contract shall be interpreted in accordance with South African laws, unless otherwise specified in SCC.

31. Notices

- 31.1 Every written acceptance of a bid shall be posted to the supplier concerned by registered or certified mail and any other notice to him shall be posted by ordinary mail to the address furnished in his bid or to the address notified later by him in writing and such posting shall be deemed to be proper service of such notice
- 31.2 The time mentioned in the contract documents for performing any act after such aforesaid notice has been given, shall be reckoned from the date of posting of such notice.

32. Taxes and duties

- 32.1 A foreign supplier shall be entirely responsible for all taxes, stamp duties, license fees, and other such levies imposed outside the purchaser's country.
- 32.2 A local supplier shall be entirely responsible for all taxes, duties, license fees, etc., incurred until delivery of the contracted goods to the purchaser.
- 32.3 No contract shall be concluded with any bidder whose tax matters are not in order. Prior to the award of a bid the Department must be in possession of a tax clearance certificate, submitted by the bidder. This certificate must be an original issued by the South African Revenue Services.

33. National 33.1 Industrial Participation (NIP) Programme

The NIP Programme administered by the Department of Trade and Industry shall be applicable to all contracts that are subject to the NIP obligation.

34 Prohibition of Restrictive practices

- In terms of section 4 (1) (b) (iii) of the Competition Act No. 89 of 1998, as amended, an agreement between, or concerted practice by, firms, or a decision by an association of firms, is prohibited if it is between parties in a horizontal relationship and if a bidder (s) is / are or a contractor(s) was / were involved in collusive bidding (or bid rigging).
- 34.2 If a bidder(s) or contractor(s), based on reasonable grounds or evidence obtained by the purchaser, has / have engaged in the restrictive practice referred to above, the purchaser may refer the matter to the Competition Commission for investigation and possible imposition of administrative penalties as contemplated in the Competition Act No. 89 of 1998.
- 34.3 If a bidder(s) or contractor(s), has / have been found guilty by the Competition Commission of the restrictive practice referred to above, the purchaser may, in addition and without prejudice to any other remedy provided for, invalidate the bid(s) for such item(s) offered, and / or

terminate the contract in whole or part, and / or restrict the bidder(s) or contractor(s) from conducting business with the public sector for a period not exceeding ten (10) years and / or claim damages from the bidder(s) or contractor(s) concerned.

Js General Conditions of Contract (revised July 2010)



SPECIAL REQUIREMENTS AND CONDITIONS OF CONTRACT

HP07-2026DAI

SUPPLY AND DELIVERY OF PHARMACEUTICAL PRODUCTS: DROPS, AEROSOLS, AND INHALED MEDICINES TO THE DEPARTMENT OF HEALTH FOR THE PERIOD

1 SEPTEMBER 2026 TO 31 AUGUST 2029

BID VALIDITY PERIOD: 180 DAYS

BID ADVERT DATE: 26 SEPTEMBER 2025

CLOSING DATE AND TIME OF BID: 24 NOVEMBER 2025 AT 11H00

NON-COMPULSORY ONLINE BRIEFING SESSION:

MS TEAMS WEBINAR: 10 OCTOBER 2025 @ 10H00



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1. ABBREVIATIONS

API : Active Pharmaceutical Ingredient

BAC : Bid Adjudication Committee

BAU : Business as Usual

CoO : Commissioner of Oaths

CPA : Contract Price Adjustment

CIPC : Companies and Intellectual Property Commission

CSD : Central Supplier Database

DVP : Digital Variation Portal

EAN : European Article Numbering

EU : European Union

GMP : Good Manufacturing Practice

HDI : Historically, Disadvantaged Individual

ID : Identification Document

IVD : In vitro diagnostic

UJV : Unincorporated Joint Venture

IJV : Incorporated Joint Venture

MCC : Medicines Control Council

MHPL : Master Health Products List

MRC : Medicine Registration Certificate

NDoH : National Department of Health

PBD : Pharmaceutical Bidding Documents

PI : Package Insert

PPPFA : Preferential Procurement Policy Framework Act

RoE : Rate of Exchange

RDP : Reconstruction and Development Programme

SAHPRA : South African Health Products Regulatory Authority

SARS : South African Revenue Service

SBD : Standard Bidding Document



SEP : Single Exit Price

SRCC : Special Requirements and Conditions of Contract

: Value Added Tax VAT



2. **DEFINITIONS**

Unless otherwise specified in this Special Requirements and Condition of Contract (SRCC), any word or expression defined in the applicable Act retains the same meaning within this document, where -

- (1) "Complementary medicine" means any substance or mixture of substances that-
 - (a) originates from plants, fungi, algae, seaweeds, lichens, minerals, animals, or other substance as determined by the South African Health Products Regulatory Authority (SAHPRA).
 - (b) is used or purporting to be suitable for use or manufactured or sold for use
 - (i) in maintaining, complementing, or assisting the physical or mental state; or
 - (ii) to diagnose, treat, mitigate, modify, alleviate, or prevent disease or illness or the symptoms or signs thereof or abnormal physical or mental state of a human being or animal; and
 - (c) is used-
 - (i) as a health supplement; or
 - (ii) in accordance with those disciplines as determined by SAHPRA.
- (2) "Consortium" means a contractual collaboration between two or more separate legal entities who combine resources or expertise for a specific tender or project, without forming a new legal entity.
- (3) "Contract" means the agreement that results from the acceptance of a tender.
- (4) "Disability" means, in respect of a person, a permanent impairment of a physical, intellectual, or sensory function, which results in restricted, or lack of, ability to perform an activity in the manner, or within the range, considered normal for a human being.
- (5) "Health supplement" means any substance, extract or mixture of substances as determined by SAHPRA, sold in dosage forms used or purported for use in restoring, correcting, or modifying any physical or mental state by-
 - (a) complementing health



- (b) supplementing the diet; or
- (c) a nutritional effect, and excludes injectable preparations, medicines or substances listed as Schedule 1 or higher in the Medicines Act.
- (6) "Historically Disadvantaged Individual (HDI)" means a South African citizen
 - (i) who, due to the apartheid policy that had been in place, had no franchise in national elections prior to the introduction of the Constitution of the Republic of South Africa,1983 (Act No 110 of 1983) or the Constitution of the Republic of South Africa,1993 (Act No 200 of 1993) ("the Interim Constitution"); and / or
 - (ii) who is a female; and / or
 - (iii) who has a disability:

Provided that a person who obtained South African citizenship on or after the coming to effect of the Interim Constitution, is deemed not to be an HDI.

- (7) "IVD" (in vitro diagnostic) means a medical device, whether used alone or in combination, intended by the manufacturer for the in vitro examination of specimens derived from the human body solely or principally to provide information for diagnostic, monitoring or compatibility purposes.
- (8) "Joint Venture (Incorporated)" means a distinct legal entity formed through the joint ownership of two or more parties, established for contractual collaboration and registered with the Companies and Intellectual Property Commission (CIPC).
- (9) "Joint venture (Unincorporated)" means a project- or bid-specific contractual collaboration between two or more entities, established without creating a separate legal entity.
- (10) "Label", when used as a verb, means brand, mark or otherwise designate or describe, and when used as a noun, means any brand or mark or any written, pictorial, or other descriptive matter appearing on or attached to or packed with and referring to any article or the package containing any article.

uthorised Signatory: Sign or Initial



- "Locally produced product" refers to a product whose formulation and conversion processes, including the use of materials and components to manufacture medicines, occur within the Republic of South Africa. This includes active pharmaceutical ingredients (APIs) (imported or locally produced) and excipients to produce finished products. Locally produced product includes the fill and finish of sterile products (including vaccines) but excludes the fill, finish, and packaging of products such as solids, liquids, sterile drops and semi-solid dosage forms.
- (12) "Management" in relation to an entity or business, means an activity inclusive of control and performed daily, by any person who is a principal executive officer of the company, by whatever name that person may be designated, and whether or not that person is a director.
- (13) "Manufacture" means all operations including purchasing of material, processing, production, packaging, quality control, release and storage of medicinal products and related control.
- (14) "Medical device" means any instrument, apparatus, implement, machine, appliance, implant, reagent for in vitro use, software, material or other similar or related article, including Group III and IV Hazardous Substances contemplated in the Hazardous Substances Act, 1973 (Act No. 15 of 1973)—
 - (a) intended by the manufacturer to be used, alone or in combination, for humans or animals, for one or more of the following:
 - (i) diagnosis, prevention, monitoring, treatment, or alleviation of disease;
 - (ii) diagnosis, monitoring, treatment, alleviation of or compensation for an injury;
 - (iv) investigation, replacement, modification, or support of the anatomy or of a physiological process;
 - (iv) supporting or sustaining life;
 - (v) control of conception;
 - (vi) disinfection of medical devices; or
 - (vii) providing information for medical or diagnostic purposes by means of in vitro examination of specimens derived from the human body; and



(b) which does not achieve its primary intended action by pharmacological, immunological, or metabolic means, in or on the human or animal body, but which may be assisted in its intended function by such means;

All medical devices are categorized based on the risk associated with the intended use of the medical device or IVD. Medical devices, including in-vitro diagnostic (IVD) medical devices and non-IVD medical devices, are grouped into four classes including Class A devices presenting the lowest potential risk (e.g. a tongue depressor) and Class D devices presenting the greatest potential risk (e.g. pacemakers) to patients, users and public health.

	RISK	NON-IVD EXAMPLES	IVD EXAMPLES	PHASE II REQUIREMENTS
Class A	Low individual risk & minimal or no public health risk	retractors/	Reagents, instruments, specimen receptacle. Microbiological culture medium	
Class B	Low-moderate	Hypodermic needle/ suction equipment	Pregnancy self-test kit, urine self-test strips to detect glucose, biochemistry test for gases, hormones, vitamins	manufacture, or import, distribute or wholesale
Class C	Moderate-high	Lung ventilators	Malaria rapid test, human genetic testing, STD test, Prenatal screening test, Tumour markers, self-monitoring blood glucose	manufacture, or import, distribute or wholesale
Class D	High	Heart valves /Implantable defibrillator	Screening for HIV/Hepatitis B, detection of Rhesus markers; testing red blood cell antigen or antibodies within ABO blood group system	manufacture, or import, distribute or wholesale

- (15) "medical device or IVD establishment" means a facility used by a manufacturer, wholesaler, distributor, retailer, service provider or an importer of medical devices or IVDs for conducting business;
- (16) "medicine" means:

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- (a) any substance or mixture of substances used or purporting to be suitable for use or manufactured or sold for use in
 - (i) the diagnosis, treatment, mitigation, modification, or prevention of disease, abnormal physical or mental state or the symptoms thereof in humans; or
 - (ii) restoring, correcting, or modifying any somatic or psychic or organic function in humans; and
 - (b) includes any veterinary medicine.
- (17) "Medicines Act" means the Medicines and Related Substances Control Act, 1965 (Act No. 101 of 1965).
- (18) "Minimum order quantity (MOQ)" means the fewest number of units a supplier is willing to sell to a single Participating Authority/Authorities in a single consignment.
- (19) "Package" means anything in or by which any medicine, complementary, veterinary medicines or scheduled substance is enclosed, covered, contained, or packed.
- (20) "Partnership" means a profit-driven arrangement between two or more persons, governed by the Partnership Act, 1939, and South African common law, in which the partners share liability and do not constitute a separate legal entity.
- (21) "Person" includes reference to a juristic person.
- (22) "Rand value" means the total estimated value of a contract in Rand denomination which is calculated at the time of tender invitations and includes all applicable taxes and excise duties.
- (23) "Single Exit Price" (SEP) is defined in the Regulations Relating to a Transparent Pricing System for Medicines and Scheduled Substances, under the Medicines and Related Substances Act No 101 of 1965. It is the price set by the manufacturer or importer, including the logistics fee and VAT, and is calculated by multiplying the price of the lowest unit of the medicine or substance by the number of units in the pack.
- (24) "Technology transfer" means a systematic and controlled procedure for transferring a manufacturing process, together with its associated documentation, professional expertise, and quality assurance principles, from one site (or entity) to another at any stage of the

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product life cycle—ranging from development, scale-up, and commercial manufacture to post-approval production.

The process involves the structured handover of documented knowledge and demonstrated operational capability from the transferring unit (TU) to the receiving unit (RU), ensuring that the RU can reproducibly perform the critical elements of the transferred technology to the satisfaction of all parties and in compliance with applicable regulatory requirements.

In this contract, technology transfer occurs within arrangements between a marketing authorization holder (applicant) and a local manufacturer (bidder) as part of initiatives to promote domestic pharmaceutical production. Where, the market authorisation holder remains on the Medicines Registration Certificate (MRC), while the local manufacturer—operating under a technology transfer agreement—executes specified manufacturing processes for the supply of a specific item within South Africa.

- (25) "Tender" means a written offer or bid in a prescribed or stipulated form in response to an invitation by an organ of state for the provision of services or goods.
- (26) "Third party manufacturer" refers to any external company or organisation, other than the holder of the Medicines Registration Certificate (MRC), that is responsible for manufacturing the product as indicated on the MRC for the item being offered in the bid. Where such a manufacturer is involved, the bidder must have formal legal agreement in place with the third party and must submit a signed Authorisation Declaration (PBD1.2) from the third party involved.
- (27) "Working days" for the purpose of this document working days refer to Monday to Friday only excluding public holidays.



SECTION A

3. BID DOCUMENT CHECK LIST

- All bid documents listed below should be compiled, indexed, and submitted in the exact sequence specified. Adhering to the suggested compilation sequence is strongly recommended.
- Each document listed must be supported by the relevant annexure, if applicable.
- All bid documents must be duly signed by a person authorised to legally bind the bidder.
- The table below serves as a guide to the documents that should be included in the bid submission.
- The inclusion of "administrative" documents is strongly recommended. While these documents are not considered during the bid evaluation process, they are required for administrative purposes.
- The absence of mandatory documents will impact the bid's responsiveness.
- Non-compliance with any requirements may render a bid non-responsive, resulting in disqualification from further evaluation in accordance with applicable procurement regulations.
- Submission of bid documents is required unless a specific document is not applicable, in which case the bidder must explicitly indicate "N/A" and provide a justification for its exclusion. If a section is blacked out in the "N/A" field, it is not considered a valid selection option for the bidder.

NO	ADMIN CODE	DOCUMENT NAME	MANDATORY DOCUMENT	N/A	YES	NO	REMARK
1	CL	Covering Letter Note: Status relating to TAX, License to Manufacture, Certificates etc.	Administrative				
2	BFI	Bid/File Index.	Administrative				
3	PBD3	PBD3: Bid Signature Authorisation Original or certified (CoO) copy	Mandatory				



NO	ADMIN CODE	DOCUMENT NAME	MANDATORY DOCUMENT	N/A	YES	NO	REMARK
4	SBD1	SBD 1: Invitation to bid.	Mandatory for bidding entity				
5	PBD4.1	PBD 4.1: Contact Details of Bidder.	Administrative				
6	CSD	CSD Registration report	Mandatory				
7	TCP	SARS Tax Clearance Pin	Mandatory	-			
8	CIPC	CIPC registration certificate	Mandatory				
9	CIPC DC	CIPC notice of change in Directors	Administrative				
10	NC	Proof of company ceding mergers, acquisition, and name changes	Administrative				
11	PBD9.1	PBD9.1: Entity Directors Categorisation and entity ownership profile Signed: Original or Certified Copy (CoO)	Administrative, if single bidding entity				
12	ID	Identification documents of Directors/Owners in PBD9.1 or PBD9.2 Certified Copy (CoO)	Administrative				
13	SBD4	SBD 4: Declaration of interest	Mandatory for bidding entity				
14	SBD6	SBD 6.1: Indicate Preference Points Claimed in table and space provided.	Mandatory				
15	ORGANOGRAM	Company Ownership Organogram	Administrative				
16	SHARES	Share Certificate(s) of HDI member/s (For Close Corporations submit CIPC CK documents) Certified copy required	Mandatory, to substantiate SBD6.1 claim				
17	SHARE REGISTER	Certified Share Register of HDI member/s (For Close Corporations submit CIPC CK documents)	Mandatory, to substantiate SBD6.1 claim				



NO	ADMIN CODE	DOCUMENT NAME	MANDATORY DOCUMENT	N/A	YES	NO	REMARK
		Certified copy required					
18	TRUST DEED	Trust /Scheme Deed listing HDI Trustees Beneficiaries and with stipulated benefit. Certified copy required	Mandatory, if applicable to substantiate SBD6.1 claim				
19	TRUST SHARE CERTIFICATE	Share Certificate for Trust ownership in bidding entity	Mandatory, if applicable to substantiate SBD6.1 claim				
20	HDI ID	ID's of HDI with equity ownership Certified copies required	Mandatory, to substantiate SBD6.1 claim t				
21	ID-DISABILITY	ID of HDI disability claimed in SBD 6.1 Certified copies required	Mandatory, to substantiate SBD6.1 claim				
22	DR-NOTE	Medical Certificate detailing the nature and extent of the disability as claimed in SBD 6.1 Certified copies required	Mandatory, to substantiate SBD6.1 claim				
23	IJV	Incorporated Joint Ventures Administrative evaluation i. Certified copy of relevant agreement between entities ii. PBD3: Bid Signature Authorisation iii. SBD1 iv. CSD report v. Tax Clearance Pin vi. CIPC registration vii. SBD 4 viii. SBD5: The National Industrial Participation Programme. ix. SBD 6.1 x. SBD 6.1 x. SBD 6.1 supporting evidence if claiming preference points xi. PBD8: Declaration of compliance SRCC and GCC	Mandatory, if submitting as an incorporated joint venture (Refer to specific requirement under each section in Annexure A and in the SRCC)				



NO	ADMIN CODE	DOCUMENT NAME	MANDATORY DOCUMENT	N/A	YES	NO	REMARK
		xii. SRCC initialed xiii. GCC initialed					
24	IJV	Incorporated Joint Ventures Administrative evaluation i.Cover letter ii.Bid/File index iii.PBD 4.1: Contact Details of Bidder. iv.CIPC notice of change in Directors, if applicable v.Notice of Ceding vi.PBD9.1: Entity Directors Categorisation vii.Identification documents of Directors / Owners	Administrative, if submitting as an incorporated joint venture (Refer to specific requirement under each section in Annexure A and				
25	CONSORTIUMS, UJV AND PARTNERSHIPS	Consortiums, unincorporated joint ventures or partnerships Administrative evaluation i.Certified copy of relevant agreement between entities ii.PBD3.1: Bid Signature Authorisation (Combined – 1 document) iii.SBD1 (Bidder) iv.CSD report (Each participant in the supplied agreement) v.Tax Clearance Pin (Each participant in the supplied agreement) vi.CIPC registration (Each participant in the supplied agreement) vi.SBD 4 (Bidder) vii.SBD 5: The National Industrial Participation Programme. (Bidder)	in the SRCC) Mandatory, if submitting as an consortiums, unincorporated joint ventures or partnerships				



NO	ADMIN CODE	DOCUMENT NAME	MANDATORY	N/A	YES	NO	REMARK
			DOCUMENT				
		ix.SBD 6.1 (Each					
		participant in the					
		supplied agreement)					
		x.SBD 6.1 supporting					
		evidence if claiming					
		preference points <i>(Each</i>					
		participant in the					
		supplied agreement)					
		xi.PBD8: Declaration of					
		compliance SRCC and	(Refer to specific				
		GCC (Bidder)	requirement				
		xii.SRCC initialed	under each				
		(Authorised signatory)	section in				
		xiii.GCC initialed	Annexure A and				
		(Authorised signatory)	in the SRCC)				
26	CONSORTIUMS,	Consortiums,	Administrative, if				
	UJV AND	unincorporated joint	submitting as an				
	PARTNERSHIPS	ventures or partnerships	consortiums,				
		Administrative evaluation	unincorporated				
		i.Cover letter	joint ventures or				
		ii.Bid/File index	partnerships				
		iii.PBD 4.1: Contact					
		Details of Bidder.					
		iv.CIPC notice of change					
		in Directors, if applicable					
		(Each participant in the					
		supplied agreement)					
		v.Notice of Ceding (Each					
		participant in the					
		supplied agreement)					
		vi.PBD9.2: Entity Directors					
		Categorisation and					
		entity ownership profile					
		(Combined – 1					
		document)	(Refer to specific				
		vii.ldentification documents	requirement				
		of Directors / Owners for	under each				
		(Each participant in the	section in				
		supplied agreement)	Annexure A and				
		viii.Organogram	in the SRCC)				

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NO	ADMIN CODE	DOCUMENT NAME	MANDATORY	N/A	YES	NO	REMARK
27	CONSORTIUMS,	Concortiums incorporated	Mandatory if				
21	IJV, UJV AND	Consortiums, incorporated and unincorporated joint	Mandatory, if submitting as an				
	PARTNERSHIPS	ventures or partnerships	consortiums,				
	FARTNERSHIPS	Technical Evaluation					
		i. PBD5: Good	incorporated / unincorporated				
		Manufacturing Practice	joint ventures or				
		(GMP).	partnerships				
		ii. Valid licence to	partiferships				
		manufacture or import					
		(Each participant in the					
		supplied agreement)					
		iii. Valid licence to					
		manufacture or import,					
		including all annexures					
		for local					
		manufacturing sites					
		(Each participant in the					
		supplied agreement)					
		iv. Valid licence to					
		manufacture/import					
		distribute/wholesale a					
		<u>Complementary</u>					
		Medicines (Each					
		participant in the					
		supplied agreement)					
		v. Valid licence to					
		manufacture/import					
		distribute/wholesale a					
		medical device or an					
		in vitro diagnostic					
		(IVD) (Each participant					
		in the supplied	(Refer to specific				
		agreement)	requirement				
		vi. Valid Medicine	under each				
		Registration	section in				
		Certificates (MRC).	Annexure A and				
		(Applicant)	in the SRCC)				
28	CONSORTIUMS,	Consortiums, incorporated	Administrative, if				
	IJV, UJV AND	and unincorporated joint	submitting as an				
	PARTNERSHIPS	ventures or partnerships	consortiums,				
		Technical Evaluation	incorporated /				
	1		unincorporated	l	I	I	1
		i. Valid SAHPRA approved GMP	unincorporated				



NO	ADMIN CODE	DOCUMENT NAME	MANDATORY	N/A	YES	NO	REMARK
	ADMIN CODE	certificate for local manufacturers ii. MRC Annexures, if applicable iii. Valid Variation Summary (MRC), if applicable iv. PBD1: Third Party Authorisation Declaration v. PBD 1.1: List of products offered sourced from third party vi. PBD 1.2: Unconditional written	joint ventures or partnerships	IV/A		NO	KLIVIARK
		undertaking vii.Package Insert viii. Proof of sample submission.	(Refer to specific requirement under each section in Annexure A and				
		ix. Bidder`s item list	in the SRCC)				
29	PBD5	PBD5: Good Manufacturing Practice (GMP). Declaration of compliance. Authorised signatory (PBD 3) to sign	Mandatory				
30	GMP-LM	Valid SAHPRA approved GMP certificate for local manufacturers as specified in PBD5 Certified copies required	Administrative				
31	SBD5	SBD5: The National Industrial Participation Programme.	Mandatory for bidding entity				
32	LICMI	Valid licence to manufacture or import (in the name of the bidder), including all annexures. Certified copies required.	Mandatory				

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NO	ADMIN CODE	DOCUMENT NAME	MANDATORY DOCUMENT	N/A	YES	NO	REMARK
33	LICM	Valid licence to	Mandatory,				
33	LICIVI	manufacture or import,	preference for				
		including all annexures for	locally produced				
		local manufacturing sites	products				
		as listed on the MRC of the	products				
		bidder (applicant). Certified					
		copies required.					
24	LICOM	<u> </u>	Manalatani				
34	LICCM	Valid licence to	Mandatory				
		manufacture/import distribute/wholesale a					
		· ·					
		Complementary Madicines (in the name of					
		Medicines (in the name of					
		the bidder), including all					
		annexures and DA02 product list:					
		Certified copies required					
35	LICMD	Valid licence to	Mandatory				
33	LICIVID	manufacture/import	iviaridatory				
		distribute/wholesale a					
		medical device or an in					
		vitro diagnostic (IVD) (in					
		the name of the bidder),					
		including all annexures:					
		Certified copies required					
36	MRC	Valid Medicine Registration	Mandatory				
		Certificates (MRC). Note:					
		All MRC's must be marked					
		by the bidder with the					
		relevant item number and					
		be sorted and filed in					
		numerical order. Certified					
37	MRC Annexures	copies required. MRC Annexures must be	Administrative				
37	WIRC Affrexures		Administrative				
		submitted only for newly registered products. Note:					
		The conditions of					
		registration must align with					
		the MRC of the newly					
		registered medicine and					
		must be clearly marked.					
38	VARSUM	A valid Variation Summary	Administrative				
50	VAINOUVI	for any changes on the	/ Millinguative				
		ioi arry criariges off the			L	<u> </u>	<u> </u>



NO	ADMIN CODE	DOCUMENT NAME	MANDATORY DOCUMENT	N/A	YES	NO	REMARK
		MRC where applicable as prescribed by SAHPRA GUIDELINE: BAU VARIATIONS COMMUNICATION, latest version - Certified copies required.					
39	PBD1	PBD1: Third Party Authorisation Declaration	Administrative				
40	PBD1.1	PBD 1.1: List of products offered sourced from third party.	Administrative				
41	PBD1.2	PBD 1.2: Unconditional written undertaking from the third party OR alternatively a formal letter from the third party could be included.	Administrative				
42	PI	The original Package Insert (PI), QR code with professional information approved by the MCC or SAHPRA must be submitted for each product offered.	Administrative				
43	PS	Proof of sample submission.	Administrative				
44	BL	Bidder's item list (list of products offered).	Administrative				
45	PBD8	PBD 8: SRCC and GCC. Declaration of compliance. Authorised signatory (PBD 3) to sign	Mandatory for bidding entity				
46	SRCC	Copy of the Special Requirements and Condition of Contract – initial every page	Mandatory				
47	GCC	Copy of General Conditions of Contract – initial every page	Mandatory				

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NO	ADMIN CODE	DOCUMENT NAME	MANDATORY DOCUMENT	N/A	YES	NO	REMARK
48	PRICE	Signed Excel Bid	Mandatory				
		Response I.e. Pricing					
		Schedule.					
		Note: If the Excel Bid					
		response Pricing					
		Schedule is not signed in					
		the space provided, the					
		bid will not be considered					
		for evaluation.					
49	USB	Set 2 & 3 - Universal Serial	Administrative				
		Bus (USB) Flash Drive /					
		Storage Device with digital					
		copy of the completed bid.					
		Note: Each compilation					
		sequence (document)					
		must be saved as a					
		separate file, with index					
		admin code abbreviations					
		used in each file name.					

All bid documents listed above must be sorted, filed, and submitted in the exact order as indicated above

Submission of bid documents is required unless a specific document is not applicable, in which case the bidder must explicitly indicate "N/A" and provide a justification for its exclusion. If a section is blacked out in the "N/A" field, it is not considered a valid selection option for the bidder.

The bid document check list is available as Annexure A in an excel spreadsheet format and should be completed by all bidders and submitted in hard copy and as part of the electronic copies of "Set 3: Electronic version of bid documents".

NOTE: Annexure A is a guidance document intended to facilitate compliance and support ease of bid submission. Each section must be read in conjunction with the relevant provisions of the Special Requirements and Conditions of Contract, as well as the General Conditions of Contract.

The NDoH reserves the right to request any administrative document or information for clarification if it does not change the substance of the bid. The bidder will have seven (7) working days to submit the requested document or information. Scanned copies must be identical to the hard copy submissions. In the event of any discrepancy, the hard copy shall take precedence.

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3.1. LEGISLATIVE AND REGULATORY FRAMEWORK

This bid and all resulting contracts shall be governed by the applicable provisions of the following legislation:

- The Medicine and Related Substances Act, 1965 (Act 101 of 1965).
- The Pharmacy Act, 1974 (Act 53 of 1974).
- The Patents Act, 1978 (Act 57 of 1978), where applicable to intellectual property rights in procurement.
- The Trademarks Act, 1993 (Act 194 of 1993), where relevant to product identification and branding.
- The General Conditions of Contract (GCC), issued in accordance with Treasury Regulation 16A under the Public Finance Management Act, 1999 (Act 1 of 1999).

The Special Requirements and Conditions of Contract (SRCC) shall supplement the GCC. In the event of any conflict between the SRCC and the GCC, the SRCC shall take precedence, except where such conflict contravenes applicable laws, regulations, or Treasury directives.

3.2. BID INFORMATION SESSION

A non-compulsory online briefing session will be held via an MS Teams Webinar on 10 October 2025 at 10H00. Bidders who wish to participate may join using the following link.

Non-Compulsory Briefing Session: HP07-2026DAI and HP08-2026SSP | Meeting-Join | Microsoft Teams

Prospective bidders must send tender-related enquiries to tenders@health.gov.za in time for responses to be received before the tender closing date.



3.3. EVALUATION CRITERIA

The evaluation process will be conducted in phases as follows:

PHASE I	PHASE II	PHASE III	PHASE IV
Administrative	Product technical	Price and Preference	Recommendation
evaluation	evaluation: Legal	Points evaluation	and Award
	and regulatory		
Bidders will be	Bidders will be	Bidders will be	Recommendation and
assessed for	evaluated for	evaluated w.r.t	award
compliance with the	compliance with the	compliance to HDI and	
mandatory	technical mandatory	RDP Goals (Price and	
administrative	requirements, and the	Preference Points) as	
requirements	product will be	per section 6 of this	
	evaluated for	SRCC	
	compliance to the		
	specification.		

4. PHASE I: ADMINISTRATIVE EVALUATION

Bidders are required to submit all applicable documents relevant to the bidding entity or in the event of a consortium, joint venture (incorporated or unincorporated), or partnership, as specified in section 4.3, by the closing date and time of the bid.

Bidders should use the original templates provided in the bid pack. Either the original or a certified copy of these templates must be submitted.

All documents must be completed, preferably in permanent black ink.

Should a copy of bid document be submitted it must be certified by a Commissioner of Oaths as a true copy of the original. No copies of certified copies will be accepted.

All bid documents that require signatures must be duly signed by the individual authorised in PBD3 or PDB3.1.



4.1. BID DOCUMENTS

Bidders are required to submit responsive bids by completing all bid documents and sort and index the bid in the same sequence as per **Annexure A**. The accurate completion of all bidding documents is required.

4.1.1. PBD3 AND PBD3.1: BID SIGNATURE AUTHORISATION

- PBD3 is required for bids submitted by a single bidding entity.
- PBD3.1 is required for bids submitted by a multi-entity bidding enterprise as described in section 4.3 of this SRCC.
- The PBD3 or PBD3.1 must be the original template from the bid pack.
- If a copy of this mandatory document is submitted, it must be certified by a Commissioner of Oaths as a true copy of the original. Failure to comply will render the bid non-responsive.
- The bidder must clearly indicate the type of bidding enterprise.
- The authorised signatories must sign and initial the PBD3 and PBD3.1 in the designated spaces.
- Electronic signatures (typed name, drawn signature, or image inserted into the PDF)
 and digital signatures (certificate-based ID, e.g. PKI, to encrypt and verify the signer's
 identity and document integrity) will be accepted.
- The authorised signatories must sign the PBD5 and PBD8, preferably in black ink.
- Details of all directors/owners involved in a multi-entity bidding enterprise, as described in Section 4.3 of this SRCC, must be recorded on PBD3.1, and a single combined PBD3.1 must be submitted.
- The absence of PBD3 or PBD3.1, where applicable, will render the bid non-responsive.

4.1.3 PBD9.1: DIRECTOR'S CATEGORIZATION AND ENTITY OWNERSHIP PROFILE

- PBD9.1 is required for bids submitted by a single bidding entity.
- PBD9.2 is required for bids submitted by a multi-entity bidding enterprise as described in section 4.3 of this SRCC.
- The PBD9.1 or PBD9.2 must be the original template from the bid pack.



- Should a copy of this document be submitted it must be certified by a Commissioner
 of Oaths as a true copy of the original.
- Bidders are strongly encouraged to retain the structure of the template provided in the bid pack without alteration.
- All columns must be completed in full, and all pages signed.
- Electronic Signatures (typed name, drawn signature, or image inserted into the PDF)
 and Digital Signature (certificate-based ID (e.g. PKI) to encrypt and verify the signer's
 identity and document integrity) will be accepted.
- Multi-entity bidding enterprise as described in section 4.3 of this SRCC should submit a single PBD9.2.
- PBD9.1 and PBD9.2 must be accompanied by certified copies of the identification documents of all Directors/Owners.

Section 4(1)(b)(iii) of the Competition Act, 1998 expressly prohibits collusive tendering. The Department is therefore required to take proactive measures to identify and mitigate such risks.

Where common director(s) are identified across multiple bidding entities that submit bids for the same item(s), those item(s) will be excluded from consideration for award. This measure is implemented to uphold the principles of fairness, transparency, and competitive integrity in the procurement process.

4.1.4 CIPC REQUIREMENTS FOR SINGLE BIDDING ENTITYS

- The latest certified copy of the CIPC Registration Certificate is required.
- For close corporations, a CIPC CK certificate must be submitted.
- Where applicable, CIPC documentation reflecting any notice of change in directors must be submitted.
- In the case of a consortium, unincorporated joint venture, or partnership, refer to Section 4.3 of this SRCC.



4.1.6 PBD5: DECLARATION OF COMPLIANCE WITH GOOD MANUFACTURING PRACTICE

- The authorised signatory as identified in the PBD3 or PBD3.1 must sign the PBD 5.
- The PBD5 must, preferably be completed and signed in black ink.

4.1.7 PBD8: DECLARATION OF COMPLIANCE WITH THE SRCC

- The authorised signatory as identified in the PBD3 or PBD3.1 must sign the PBD 8.
- The PBD5 must, preferably be completed and signed in black ink

4.1.8 EXCEL BID RESPONSE I.E., PRICING SCHEDULE:

Bidders are required to submit fully completed and responsive bids by accurately completing all fields in the Excel Bid Response Document, including pricing information. All prices must be quoted to two (2) decimal places.

Quoted prices must be all-inclusive (including VAT) and reflect the total cost for supply and delivery to the specified destination. The bid price for each product will be deemed applicable to the pack size and unit of measure as specified in the item description.

Bidders are strongly advised to consult the "Definition of Fields" document included in the Bid Response Document for detailed guidance on completing each field correctly. Incomplete or inaccurate submissions may result in the bid being deemed non-responsive and disqualified.

4.1.9. DELIVERED BID PRICES OFFERED.

Final prices submitted must **not exceed** the most recent Single Exit Price (SEP) as recorded on the National Department of Health (NDoH) SEP database. Where the prices offered at the date and time of bid closure exceed the ex-manufacturer component of the SEP, inclusive of VAT, price negotiations will be required, where applicable, in accordance with the relevant regulations.

If, following negotiations, the bidder offers a price below or equal to the Single Exit Price, the award may be considered. However, the bidder will only qualify for contractual price adjustments



up to the most recent Single Exit Price as recorded in the National Department of Health (NDoH) SEP Database.

4.2. TAX COMPLIANCE STATUS

Bidders must be registered on the Government's Central Supplier Database (CSD) and include their full CSD report with their bid submission. The NDoH will verify the bidder's tax compliance status through the CSD.

The CSD and the Tax Compliance Status (TCS) PIN are the approved methods for verifying a bidder's tax compliance. Bidders must submit a valid TCS PIN with their bid. It is a condition of this bid that the bidder's tax matters are in order, or that satisfactory arrangements have been made with SARS to meet the bidder's tax obligations.

If the bidder is found to be non-compliant with tax obligations during any stage of the evaluation process, the bidder will be notified of their non-compliance status. The bidder will be requested to submit, within seven (7) working days:

- a) Proof of tax compliance
- b) Proof must be provided that arrangements have been made with SARS to address any tax compliance issues, ensuring that the bid adjudication process is not delayed.

By submitting this bid, the bidder confirms that SARS may disclose the bidder's tax compliance status at any time during the contract period. Such confirmation is deemed granted by the bidder upon submission of the bid.

In the case of a consortium, joint venture (unincorporated) or partnership, **each party** must be registered on the CSD, and their tax compliance status will be verified through the CSD, as described in section 4.3.

Bidders are responsible for ensuring that their CSD information is updated in accordance with the bid documents submitted.

Foreign suppliers, who do not have South African tax obligations or a history of doing business in South Africa, must complete the questionnaire on the SBD1 form. If a foreign bidder is $28 \mid P \mid a \mid g \mid e$



recommended for award, the NDoH will submit the completed SBD1 to SARS at the email address: GovernmentInstitute@sars.gov.za. SARS will then issue a confirmation letter to the NDoH, confirming whether the foreign entity has any tax obligations in South Africa.

4.3. CONSORTIUMS, JOINT VENTURE (INCORPORATED OR UNINCORPORATED), AND PARTNERSHIPS

If the bidder is not the applicant as required in Section 5.2, but any of the following conditions apply, the bid submission must include a signed agreement between the bidder and the applicant. This requirement applies in the following cases:

- The bidder is not the applicant on the MRC, but both the bidder and the applicant are subsidiaries of a single legal entity (same parent company).
- The bidder is not the applicant on the MRC, but either the bidder or the applicant is fully or partially owned by the other.
- The bidder is not the applicant on the MRC, but the bidder and the applicant are part of a technology transfer arrangement.

In such cases, the following documentation **must** be included with the bid:

For Consortiums, Joint Ventures (incorporated and unincorporated) or Partnerships, a
certified copy of the signed agreement between the bidder and the applicant, outlining
the terms of their relationship.

Each entity as specified in the supplied agreement and part of the consortium, joint venture (unincorporated), or partnership **must** submit all mandatory documents including the following:

- Certified copy of relevant agreement between entities
- PBD3.1: Bid Signature Authorisation One document should be submitted per bid, identifying the authorized person on behalf of the multi-entity bidding enterprise.
- SBD 1: Invitation to bid. only need to be completed by the bidder
- CSD Registration report Each entity specified in the supplied agreement
- SARS Tax Clearance Pin Each entity specified in the supplied agreement
- CIPC registration certificate Each entity specified in the supplied agreement



- SBD 4: Declaration of interest only need to be completed by the bidder
- (Bidder)
- SBD5: The National Industrial Participation Programme. only need to be completed by the bidder
- SBD 6(1) Indicate Preference Points Claimed in table. -Each entity specified in the supplied agreement
- SBD 6.1 supporting evidence if claiming preference points -Each entity specified in the supplied agreement
- PBD8: Declaration of compliance SRCC and GCC only need to be completed by the bidder
- SRCC initialed (Authorised signatory)
- GCC initialed (Authorised signatory)

Incorporated joint ventures are regarded as single bidding entities for the purpose of administrative evaluation. Therefore, the bidding entity, as specified in the supplied agreement, must submit all mandatory documents, including the following:

- PBD3: Bid Signature Authorisation
- SBD 1: Invitation to bid.
- CSD Registration report
- SARS Tax Clearance Pin
- CIPC registration certificate
- SBD 4: Declaration of interest
- SBD5: The National Industrial Participation Programme.
- SBD 6(1) Indicate Preference Points Claimed in table
- SBD 6.1 supporting evidence if claiming preference points
- PBD8: Declaration of compliance SRCC and GCC
- SRCC initialed
- GCC initialed



Additionally, **all parties involved** in the consortium, joint venture (incorporated or unincorporated) or partnership **must** submit the relevant legislative and mandatory documentation as required for this bid, as specified in the SRCC Section 5.1.

For Phase II compliance, the following documents are mandatory for all parties involved in consortium, joint venture (incorporated or unincorporated), or partnership:

- A valid license to manufacture (bidder and applicant), along with certified copies as per section 5.1.1, must be provided for all parties involved in the bid.
- An MRC (Medicines Registration Certificate) as per section 5.1.2, where one of the
 parties in the consortium, joint venture (incorporated or unincorporated) or partnership
 is identified as the applicant.

The bid must be submitted independently and without collusion or prior consultation with competitors. While communication within a consortium, joint venture (incorporated or unincorporated) or partnership is allowed, sharing bid details with external competitors constitutes collusive bidding, which is prohibited.

If participating in a consortium, joint venture (incorporated or unincorporated), or partnership, no party involved; including any director, as referenced in Section 4.1.3 of the SRCC; shall submit a separate or competing bid for the same item.



5. PHASE II: PRODUCT TECHNICAL EVALUATION: LEGAL AND REGULATORY

5.1. LEGISLATIVE REQUIREMENTS RELATING TO THIS BID

5.1.1 LICENSING REQUIREMENTS

5.1.1.1. THE BIDDER OFFERING A MEDICINE:

- Must be the holder of a valid license to manufacture or import medicines, issued in terms of section 22C(1)(b) of the Medicines Act. The bidder must submit a certified copy of the original license, including all annexures.
- Additionally, if the bidder is offering a product manufactured locally, they must submit
 a certified copy of the original valid license to manufacture medicines, including all
 annexures, for all local manufacturing sites listed on the MRC.

5.1.1.2. THE BIDDER OFFERING A CLASS A, B, C, OR CLASS D MEDICAL DEVICE OR AN IN VITRO DIAGNOSTIC (IVD):

- Must be the holder of a valid license to manufacture, import, distribute, or wholesale
 medical devices or IVDs, issued in terms of section 22C(1)(b) of the Medicines Act,
 including all annexures. The bidder must submit a certified copy of the original license,
 including all annexures relevant to the products offered.
- An information leaflet for the unregistered medical device should be supplied, if required by SAHPRA.

5.1.1.3. THE BIDDER OFFERING CATEGORY D COMPLEMENTARY MEDICINES:

- Must be the holder of a valid license to manufacture, import, or export Complementary
 medicines (Category D), issued in terms of section 22C(1)(b) of the Medicines Act,
 including the DA02 Product List as issued by SAHPRA. The bidder must submit a
 certified copy of the original valid license, including all annexures relevant to the
 products offered.
- An information leaflet for the complementary medicines should be supplied, if required by SAHPRA.



5.1.2. IN THE CASE OF A CONSORTIUM, JOINT VENTURE (INCORPORATED OR UNINCORPORATED), AND/OR PARTNERSHIP.:

 All involved parties must be holders of the license to manufacture or import medicines, issued in terms of section 22C(1)(b) of the Medicines Act. Companies must submit certified copies of the respective licenses, as described in section 4.3.

Where SAHPRA issues an electronic certificate or licence, a hard copy must still be submitted. This printed copy must be certified by a Commissioner of Oaths.

5.2. MEDICINE REGISTRATION CERTIFICATE (MRC) REQUIREMENTS AND VARIATION SUMMARIES

Items offered must be registered in terms of Section 15 of the Medicines Act and must comply with the conditions of registration for the duration of the contract.

- In the case of medicines, a certified copy of the original MRC, issued in terms of Section 15(3)(a) of the Medicines Act, must be included with the bid for each item offered.
- The bidder must be indicated as the applicant on each MRC.
- Where there is a variation in the MRC, the bidder should submit the Variation Summary.
- In the event a product offered is not eligible for registration in terms of Section 15(3)(a) of the Medicines Act, refer to section 5.1.1 relating to Medical Devices, and Complementary medicine requirements.
- Where the bidder is not the applicant, refer to Section 4.3 regarding consortium, joint venture (incorporated or unincorporated), or partnership.

5.3. SUBMISSION OF MRC ANNEXURES (CONDITIONS OF REGISTRATION)

Medicine registration may be subject to conditions as determined by SAHPRA in terms of Section 15(6)(a) of the Medicines Act. These conditions, as outlined in the MRC annexures (conditions of registration), should be submitted in the following instances:

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- All newly registered medicines.
- Medicines for which a bid is being placed for the first time.
- In the event of a medicine review or renewal in terms of Section 15(6)(a) of the Medicines Act.

All bidders should submit, where applicable, a valid variation summary as prescribed by the latest version of the SAHPRA GUIDELINE: BAU VARIATIONS COMMUNICATION, along with a certified copy of the original MRC issued by the MCC/SAHPRA.

5.4. THIRD PARTY AUTHORISATION DECLARATION

Only the holder of a valid MRC issued in terms of the Medicines Act may submit a bid.

If the holder of the Medicines Registration Certificate (MRC) is not the manufacturer of the product offered in this bid, a third-party manufacturer authorisation is required. In such cases, the bidder must establish a formal legal agreement with the third-party manufacturer and submit a signed Authorisation Declaration (PBD1.2) from the relevant manufacturer as approved by SAHPRA which must be listed on the MRC.

The NDoH reserves the right to verify any information supplied by the bidder in the Authorisation Declaration. Should any information be found to be false or incorrect, the NDoH may exercise any remedies available to it as outlined in the bid documents.

Failure to submit a duly completed and signed Authorisation Declaration, along with the required annexures, in accordance with these provisions, may result in the invalidation of the bid for the goods or services offered.

No agreement between the bidder and any third party will be binding on the NDoH.

5.5. SAMPLES TO BE SUBMITTED TO SAMPLE EVALUATION SITES

All bidders are required to submit samples, including those who are currently supplying the NDoH with products, to confirm the following:

• Compliance with the specifications set out in the bid document/item specification.



Compliance of the product with the requirements of the Medicines Act.

Failure to submit samples to both institutions listed below will result in the invalidation of the bid for the items offered. Samples must be submitted to each of the depots at the addresses indicated below prior to the closing date and time of the bid:

GAUTENG MEDICAL SUPPLIES DEPOT	CAPE MEDICAL DEPOT
Ms Pretty Nyokong	Mr Nisaar Mia
Contract Manager	Pharmaceutical Policy Specialist
Tel: 011 628 9001/11	Tel: 021 483 5800
Gauteng: Medical Supplies Depot	Western Cape: Department of Health
Store 3	4th Floor, Cape Medical Depot
35 Plunkett Avenue	16 Chiappini Street
Hurst Hill	Cape Town
2092	8001

- No samples are to be sent to the NDoH.
- Samples should be clearly marked with the bid number, item number, and the bidder's name and address.
- All samples must be a true representation of the product that will be supplied.
- Bidders must submit at least one original pack of each offered item for evaluation.
- A mock sample may be accepted for a registered product with SAHPRA that is not yet available on the market. The mock sample must be a true representation of the product to be supplied, should a contract be awarded, and must include the product (tablet, capsule, liquid, etc.) in a form that may not be in the original container, along with the SAHPRA-approved artwork and package insert.
- It is the bidder's responsibility to ensure that samples have been received at the addresses provided above.
- All samples for awarded items will be retained for the duration of the contract.
- For **Schedule 6** medicines only, the primary packaging/artwork and package insert, or professional information must be submitted (do not include the product itself).
- Proof of sample submission, including a signed copy of the item list as received by the sample evaluation site, should be submitted with the bid documents by the closing date and time of the bid.



- All samples submitted should include an eligible package insert, QR code or professional information leaflet (as indicated in section 5.1.1) approved by SAHPRA.
- Both institutions will evaluate the samples submitted to ensure compliance with the specifications.

5.6. COMPLIANCE WITH SPECIFICATIONS

- Items must comply with the specification as detailed in the bid document.
- The Department reserves the right to award a product with a Specification Deviation.
- Where a product is awarded with a specification deviation, no cost-related conversion will be applied to that item.
- 6. PHASE III: PRICE AND PREFERENCE POINTS EVALUATION
- 6.1. CRITERIA USED FOR THE ALLOCATION OF PREFERENTIAL POINTS CLAIMED IN TERMS OF THE REVISED PREFERENTIAL PROCUREMENT REGULATIONS (PPPFA), 2022

Preference points will be evaluated and allocated in accordance with the revised Preferential Procurement Regulations of 2022, issued under sections 2 and 5 of the Act, which aim to promote:

- a) **Empowerment of Historically Disadvantaged Individuals (HDI)**, which refers to South African citizens who:
 - Were denied the right to vote in national elections prior to the introduction of the Constitution of the Republic of South Africa, 1983 (Act No. 110 of 1983) or the Constitution of the Republic of South Africa, 1993 (Act No. 200 of 1993) (referred to as "the Interim Constitution"). – also referred to as individuals without franchise.
 - Are female.
 - Have a disability.
- b) Promotion of specific Reconstruction and Development Programme (RDP) goals, as outlined in section 2(1)(d) of the Act.
 - Selected Goal: The promotion of South African-owned entities, specifically ownership held by South African citizens in the bidding entity.

6.1.1. HDI AND RDP GOAL POINTS CLAIMABLE FOR THIS TENDER

HDI Promotion and points claimable:

NO	DESCRIPTION	CLAIMABLE POINTS
1	Who had no franchise in national elections before the 1983 and 1993 Constitutions	4
2	Who is a female	2
3	Who has a disability	2

RDP Goal for this tender and points claimable:

NO	DESCRIPTION	CLAIMABLE POINTS
1	The promotion of South African owned entities	2

6.1.2. SUBSTANTIATING DOCUMENTS REQUIRED FOR HDI POINTS (SBD6.1)

Each bid must include a SBD6.1.

In the event of consortiums, joint ventures (unincorporated), or partnerships as specified in section 4.3 of this SRCC, each party should complete SBD6.1.

SBD6.1 must be completed in full and in accordance with the requirements to claim preference points. If the SBD6.1 is not completed in accordance with the requirements no preference points will be allocated.

6.1.2.1. ALLOCATION OF PREFERENCE POINTS FOR HDI

Percentage (%) of HDI ownership held in the bidding entity, should be supported by share certificate and share register i.e. if four (4) points is claimable, then two (2) points will be allocated for 50% ownership.

Equity Ownership claims must be supported by substantiating evidence to be considered for points claimed in SBD6.1.



HDI Equity Ownership (without franchise)

NO	HDI DESCRIPTION	CLAIMABLE POINTS
1	Who had no franchise in national elections before the 1983 and 1993 Constitutions	4

The following supporting documents are mandatory to substantiate claims made for HDI equity ownership:

- Certified copies of identification documents (IDs), and
- Certified copies of Share certificates, and
- Certified Share statement or Share Register reflecting the total number of shares issued by the bidding entity and the shares held by each qualifying HDI.

HDI Equity Ownership through Trusts / Employment Scheme or Similar (without franchise)

Where HDI ownership is held through a Trust Deed or Employment Scheme, the individuals concerned must be listed as **both trustee and beneficiary** in the Trust Deed or Employment Scheme, as proof that they are actively involved in the management of such Trust Deed or Employment Scheme.

The following supporting documents are mandatory to substantiate claims made for HDI ownership within a Trust or Employment Scheme or Similar:

- Certified Trust Deed indicating HDIs listed as Trustees and Beneficiaries, and
- Certified share certificate confirming ownership held by Trust in bidding entity, and
- Certified copies of identification documents (IDs) of qualifying Trustees and Beneficiaries.

HDI: Female Ownership

NO	DESCRIPTION	CLAIMABLE POINTS
2	Who is a female	2



The following supporting documents are mandatory to substantiate claims made for female ownership:

- Certified copies of IDs, and
- Certified copies of Share certificate/s, and
- Certified Share statement or Share Register reflecting the total number of shares issued by the bidding entity and indicating the shares held by South African female/s.

Female Equity Ownership through Trusts / Employment Scheme or Similar

Where female ownership is held through a Trust Deed or Employment Scheme, the individuals concerned must be listed as **both trustee and beneficiary** in the Trust Deed or Employment Scheme, as proof that they are actively involved in the management of such Trust Deed or Employment Scheme.

The following supporting documents are mandatory to substantiate claims made for Female ownership within a Trust or Employment Scheme or Similar:

- Certified Trust Deed indicating HDIs listed as Trustees and Beneficiaries, and
- Certified share certificate confirming ownership held by Trust in bidding entity, and
- Certified copies of identification documents (IDs) of qualifying Trustees and Beneficiaries

Disability Equity Ownership

NO	DESCRIPTION	CLAIMABLE POINTS
3	Who has a disability	2

The following supporting documents are mandatory to substantiate claims made for ownership, by individuals with a disability:

- Certified copies of identification documents (IDs), and
- Medical Certificate detailing the nature and extent of the disability required, and



 Certified Share statement / Share Register reflecting the total number of shares issued by the bidding entity and indicating the shares held by a South African(s) with a disability.

Disability Equity Ownership through Trusts / Employment Scheme or Similar

Where disability ownership is held through a Trust Deed or Employment Scheme, the individuals concerned must be listed as **both trustee and beneficiary** in the Trust Deed or Employment Scheme, as proof that they are actively involved in the management of such Trust Deed or Employment Scheme.

The following supporting documents are mandatory to substantiate claims made for HDI equity ownership held by individuals with disabilities:

- Trust Deed indicating listed HDI owner as trustees and beneficiaries, and
- Certified copies of identification documents (IDs), and
- Medical Certificate detailing the nature and extent of the disability.
- Certified share certificate confirming ownership held by Trust in bidding entity.

6.2. RDP GOAL: PROMOTION OF SOUTH AFRICAN OWNED ENTITYS

6.2.1. ALLOCATION OF RDP GOAL PREFERENTIAL POINTS

Percentage (%) of ownership held by South Africans in the bidding entity, supported by share certificate and share register, will be used to calculate claimable points i.e. one (1) point allocated for 50% ownership.

6.2.2. SUBSTANTIATING DOCUMENTS REQUIRED FOR RDP GOAL (SBD6.1)

NO	DESCRIPTION	CLAIMABLE POINTS
4	The promotion of South African owned enterprises	2

RDP Goal



The following supporting documents are mandatory to substantiate claims made for ownership by South African individuals:

- Certified copies of IDs, and
- Certified copies of Share certificate/s, and
- Share statement/Share Register reflecting the total number of shares issued by the bidding entity and indicating shares held by South Africans

RDP Ownership in Trusts / Employment Scheme or Similar

The following supporting documents are mandatory to substantiate claims made for ownership in Trusts / Employment Scheme or Similar:

- Share certificate(s) reflecting ownership of the Trust / Ownership Scheme in the bidding entity.
- Trust Deed indicating those South Africans who are both Trustees and Beneficiaries and as such and who are actively involved in the management of the Trust; and
- Certified copies of IDs the Trustees and Beneficiaries.

6.3. HDI CLAIMS IN CONSORTIUMS, JOINT VENTURES (UNINCORPORATED), OR PARTNERSHIPS

Multiple entities forming part of a single bidding enterprise, such as a consortium, unincorporated joint venture, or partnership, may claim preferential points for qualifying Historically Disadvantaged Individuals (HDIs).

To validate such claims, a **certified copy of the signed agreement** between the participating entities must be submitted with the bid.

To claim preferential points, the agreement should clearly specify:

- The terms of the relationship;
- The percentage (%) stake of each entity in the bidding enterprise for the purpose of executing the tender.

Note: If the percentage stake of each entity is not identified, no preferential points will be allocated.



The claim for preferential points must be aligned with the equity ownership held by qualifying HDI individuals within each participating entity of the bidding enterprise. For example, where a partnership comprises two entities with a 60% and 40% stake in the execution of a tender:

- The qualifying HDI individuals within the 60% partner will contribute towards 60% of the preferential points that may be earned by the bidding enterprise.
- Similarly, the HDI individuals within the 40% partner will contribute towards the remaining 40% of the preferential points.

Accordingly, each participating entity may only claim HDI points, by completing the **SBD 6.1 form**, in direct proportion to its stake in the consortium, unincorporated joint venture, or partnership, as confirmed in the formal agreement between the entities forming the bidding enterprise.

Preferential points are allocated based on the extent of HDI equity ownership and in alignment with the applicable Reconstruction and Development Programme (RDP) Goals.

6.4. FORMULAE - PREFERENCE POINT SYSTEM TO BE APPLIED IN THIS TENDER 6.4.1. FORMULA FOR PRICE (90)

The 90/10 preference point system will be applied in this tender to allocate points for price. This system is applied for acquisition of goods or services with a Rand value **above R50 000 000 (all applicable taxes included).** The points for price shall be allocated in the following manner:

Responsive bids will be adjudicated by the NDoH on the 90/10-preference point system in terms of which points for price will be awarded to bidders based on:

• The bid price (maximum 90 points)

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The following formula will be used to calculate the points for price:

$$Ps = 90\left(1 - \frac{Pt - Pmin}{Pmin}\right)$$

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Ps = Points scored for price of tender under consideration

Pt = Price of tender under consideration

Pmin = Price of lowest acceptable tender

6.4.2. FORMULA FOR HDI PREFERENCE POINTS (10)

 $NEP = \frac{NOP \times EP}{100}$

Where

NEP = Points awarded for equity ownership by an HDI

NOP = The maximum number of points awarded for equity ownership by an

HDI

EP = The percentage of equity ownership of and HDI within the entity of

business, determined in accordance with the Act and specific provisions contained in the revised Preferential Procurement

Regulations, 2022.



7. PREFERENCE FOR LOCALLY PRODUCED PRODUCTS

The NDoH reserves the right to consider locally produced products offered by bidders. Bidders must indicate the manufacturing location of the products in the Excel Bid Response Document.

To provide preference to locally produced products, the definition of a "locally produced product" is limited to the formulation and conversion processes that use materials and components to manufacture medicines (including raw materials, whether imported or locally produced, for active pharmaceutical ingredients (API) and excipients to produce finished products) within the Republic of South Africa. A locally produced product includes the **fill and finish of sterile products** (vaccines, small and large volume parenterals. However, it **excludes** the **fill, finish, and packaging** of **non-sterile dosage forms** such as solids, liquids, sterile drops, and semi-solid formulations.

Providing that awarding locally produced products does not compromise security of supply or affordability, the quantities allocated for award as locally produced products, may be allocated proportionally, aligning with the percentage of the product volume that will be locally produced.

Preference will be given to bidders of locally produced products if:

- A certified copy of the valid License to Manufacture, as per section 22C(1)(b) of the Medicines Act, for the local manufacturing site (including all applicable annexures) for medicines, complementary medicines, and medical devices/IVDs is submitted.
- The local manufacturing site is listed on the MRC issued by SAHPRA, indicating that the manufacturer is located in the Republic of South Africa.
- The Single Exit Price (SEP) published on the SEP database is not exceeded.
- The local manufacturer has demonstrated the capacity to supply the required volumes based on the data provided in the Excel Bid Response Document.
- The bidder complies with all other clauses contained in this SRCC.

If the necessary documentation or evidence is not included in the bid documents, the bid will not qualify for preference as a locally produced product.

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8. VALUE ADDED TAX

All bid prices must be inclusive of 15% Value-Added Tax. Failure to comply with this condition will invalidate the bid.

9. SUBMISSION OF BIDS

All bid documents should be **compiled**, **indexed**, **and submitted** in the **exact sequence** specified. All required documents as specified in this SRCC and supporting evidence should be submitted.

- Submission of bid documents is required unless a specific document is not applicable, in which case the bidder must explicitly indicate "N/A" and provide a justification for its exclusion. If a section in Annexure A is blacked out in the "N/A" field, it is not considered a valid selection option for the bidder.
- All bid documents must be signed or initialed in the spaces provided within the document, preferably in permanent black ink.
- Where certified copies of original documents are submitted, bidders must ensure that the certification is original, signed, and dated by the Commissioner of Oaths.
- If SAHPRA issues an electronic certificate or license, a hard copy must still be provided. This printed version must be certified by a Commissioner of Oaths.
- All SBD bid documents must be fully signed and witnessed, where required, preferably in permanent black ink.
- All mandatory documents as specified in **Annexure A** must be valid at the time of bid closure.
- The NDoH will not accept updated mandatory bid documents after the bid closure date unless the document was valid at the time of bid closure but is set to expire during the bid validity period. In such cases, an updated document may only be submitted if specifically requested by the Department.
- Bidders who do not comply with any of the mandatory requirements will be deemed non-responsive and may not be considered for evaluation.



10. COMPLETION OF DOCUMENTS AND BID SUBMISSION

Bidders are required to submit three sets of bid documents according to the instructions below. All three sets must be submitted not later than the closing date and time in a sealed package.

The full name and address of the bidder, including the return address, the bid number, and the closing date, must be clearly indicated on the package.

The bid must comprise of:

- **Set 1** The original **Hard copy bid**, (signed legal documents, including all certificates and documents requested); bound with tabs indicating section as per Annexure A Checklist.
- **Set 2 (Electronic Copies),** consisting of a scanned PDF of the Hard Copy bid, and saved together with Set 3 on a USB Flash Drive / Storage Device.
- Set 3 (Excel Spreadsheets) comprising of the electronically completed Excel spreadsheets.

All fields must be completed. Where the requested information / documentation is not applicable, indicate 'N/A' and provide a comment explaining the reason for non-applicability.

10.1. Set 1: Hard copy legally binding bid documents.

Bidders must complete all SBD, PBD and Bid Response forms in permanent black ink, or typed. Where no electronic entry field is provided, bidders must complete the forms in permanent black ink, handwritten. All bid documents must be signed in ink in the spaces provided within the document. All bid documents must be initialed at the bottom of each page.

The following must be applied:

- Where certified copies of original documents are submitted, bidders must ensure that the certification is original and dated by the Commissioner of Oaths.
- Where SAHPRA issues an electronic certificate or license, a hard copy must still be provided. This printed version must be certified by a Commissioner of Oaths. Where applicable, all bid documents must be witnessed preferably in permanent black ink.
- The signed hard copy of the bid document will serve as the legal bid document.



- Bidders must submit their complete bid in hard copy format (paper document).
- All pages in the complete bid document must be signed and initiated with preferably permanent black ink.
- The use of correction fluid is not acceptable.
- Any change/s must be clearly indicated and initiated.

Note Set 2 & 3

Bidders must submit a USB flash drive/storage device with a digital copy of the completed bid. Bidders must follow the same compilation sequence as per the index and use the index admin code abbreviation used in the file name.

10.2. Set 2: PDF of Hard Copy signed legal documents. (i.e., PDF of Set 1)

Bidders must submit a PDF version of the entire signed hard copy bid, including all certificates and documents requested.



10.3. Set 3: Electronic version of bid documents

In addition, bidders must submit the electronic versions, Bid Response Document, and other relevant spreadsheets in Excel (not PDF). All three sets of information must be submitted for the bid to be evaluated. Ensure that the bid price is offered for the product as specified.

Bidders must ensure that the **price quoted** for a product (line item) on the Bid Response Document is for the unit pack as specified. No conversion factors will be applied.

11. LATE BIDS

Bids received after the closing date and time at the address indicated in the bid documents will not be accepted for consideration and, where practical, will be returned unopened to the bidder.

12. COUNTER CONDITIONS

Bidders' attention is drawn to the fact that amendments to any of the bid conditions or setting of counter conditions by bidders may result in the invalidation of such bids.

13. FRONTING

The NDoH supports the spirit of the RDP Goals and HDI empowerment and recognizes that true empowerment can only be achieved through individuals and businesses acting in accordance with the Constitution, and in an honest, fair, equitable, transparent, and legally compliant manner. In this regard, the NDoH condemns any form of fronting.

The NDoH encourages bidders to act with honesty during their bid preparation process. Should any fronting, bid rigging, or collusion practices be suspected, the NDoH reserves the right to conduct investigations to verify the accuracy of the representations made in bid documents. Any form of misrepresentation, corruption, or fraudulent practice identified on the part of the bidder may result in serious consequences as specified in the relevant regulations. These consequences may include prohibiting the offending bidder from conducting business with the public sector for a period not exceeding 10 years.



14. SUPPLIER DUE DILIGENCE

The NDoH reserves the right to conduct supplier due diligence prior to the final award. This may involve such steps as the Department, in its sole and absolute discretion, deems necessary to satisfy itself regarding, inter alia, the legal, compliance, financial, and operational status and condition of the bidder, supplier, and/or its affiliates (as the case may be).

This may include site visits to assess whether:

- The item is manufactured at the site specified in the bid documentation;
- The bidder has the capacity to meet their allocated or agreed demand.

15. COMMUNICATION

The NDoH reserves the right to communicate with bidders post bid closure and during the bid validity period, for the purpose of seeking clarification on documents submitted or extending the validity period of the bid, if necessary. All communication between the bidder and the NDoH must be conducted in writing. Any communication between a bidder and any government official, during the bid validity period, is strongly discouraged.

Information obtained during clarification may be shared with relevant committees involved in the tender process, in accordance with applicable procurement protocols and competition regulations.



16. CONTACT DETAILS

Postal address

Directorate: Affordable Medicines

Private Bag X828

PRETORIA

0001

Physical address

Directorate: Affordable Medicines

Dr AB Xuma Building

1112 Voortrekker Road,

Block A Pretoria

Townlands 351-JR

PRETORIA

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Please use the following e-mail address for any queries relating to the bidding process:

tenders@health.gov.za

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SECTION B

17. CONTRACT PERIOD

The contract shall be for the period of three years starting 1 September 2026 to 31 August 2029.

18. PARTICIPATING AUTHORITIES

Participating Authorities on this contract are: Provincial Departments of Health and other entities as approved by the Accounting Officer:

- Department of Correctional Services;
- South African Military Health Services;

Provincial Departments of Health:

- Eastern Cape
 Western Cape
- Northern Cape
- Free State
- KwaZulu-Natal
- Limpopo
- Mpumalanga
- North-West
- Gauteng

Other entities may request to participate in the contract during the contract period. Such requests will only be considered if the awarded suppliers agree and confirm in writing that the inclusion of additional participants will not compromise the security of supply. Participation by other entities is subject to the approval of the Chief Accounting Officer of the NDoH. Appropriate consultation and communication with the contracted suppliers will take place prior to any approval being granted.



19. REGISTRATION ON DATABASES OF PARTICIPATING AUTHORITIES

The contracted suppliers must register on the supplier databases of Participating Authorities within 30 days after the award of the contract.

Failure to meet this requirement will result in the inability to process payment for goods.

20. AWARD CONDITIONS

NDoH reserves the right to:

- Award the same item as a multiple award to various suppliers (two or more) to address high volume requirements, security of supply and product availability.
- Negotiate prices and minimum order quantities and volumes.
- Award an item with a specification deviation.
- Refrain from applying conversion factors when a 30-day dispensing pack size is advertised, and a 28-day dispensing pack size is offered.
- Combine the quantities and award only one item number, where applicable if an item is advertised as a single item but is included in a therapeutic class and is recommended for award within that class.
- Only award one item in a procurement class. The item could be awarded to multiple suppliers as a split award.

In cases where the tender does not achieve the most economically advantageous price, the NDoH reserves the right not to award that item.

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20.1. SPLIT AND MULTIPLE AWARDS

The NDoH reserves the right to issue split or multiple awards, where necessary, to facilitate security of supply. The following will be taken into consideration when contemplating a split or multiple award:

- Source of API and manufacturing site;
- Capacity to meet expected demand as per published estimates in the Bid Response Document;
- Estimated volume to be supplied;
- Risk to public health if the item is not available;
- Past compliance of the bidder with contractual obligations.
- The Minimum Order Quantity (MOQ) for split or multiple awards will be negotiated and aligned to the smallest acceptable value.

Two-way split awards will be made in accordance with the following schedule based on the points scored:

CATEGORY	DIFFERENCE BETWEEN POINTS SCORED	RECOMMENDED PERCENTAGE SPLIT
Α	Equal points	50/50
В	< 5 points	60/40
С	>5-10 points	70/30
D	>10-20 points	80/20
E	>20 points	90/10

Where a split of **three (3) or more** bidders is contemplated, the total score of each will be applied in the following formula to determine the percentage (%) split for each bidder: For example, the percentage split for the highest scoring bidder will be calculated as follows:

$$% Split = T1/(T1+T2+T3)$$

Where:

T1 = Score of highest Scoring Bidder

T2 = Score of second Highest Scoring Bidder

T3 = Score of third Highest Scoring Bidder



20.2. THERAPEUTIC CLASS AWARDS

The Policy for Classifying Medicines into Therapeutic Classes for Purposes of Therapeutic Interchange (as published: https://www.health.gov.za/wp-content/uploads/2021/08/Therapeutic-Interchange-Policy July2021 final.pdf; July 2021) defines a therapeutic class as a group of medicines that contain active ingredients with comparable therapeutic effects. Medicines within a therapeutic class may not necessarily belong to the same pharmacological class, may differ in chemistry or pharmacokinetic properties, and may have different mechanisms of action, adverse reactions, toxicity, and drug interaction profiles. In most cases, however, these medicines exhibit similar efficacy and safety profiles when administered in equipotent doses for a specific indication.

The ministerially appointed National Essential Medicines List Committee (NEMLC) is responsible for formulating and revising the Standard Treatment Guidelines (STGs) and the Essential Medicines List (EML). Therapeutic classes are specified in the "Medicine Treatment" section of the national STGs, which lists a class of medicines followed by examples, such as HMG-CoA reductase inhibitors (Statins) – e.g., simvastatin. These therapeutic classes are designated when no member of the class offers a significant benefit over another for a specific indication. The NEMLC may designate therapeutic classes for a condition, where applicable.

Such therapeutic classes may be utilised during the contracting process to achieve the most economically advantageous contracts, maximize market volume, and increase competition, thereby offering potential cost efficiencies through robust competition. A single member from the therapeutic class may be awarded on the contract.

HP07-2026DAI – Therapeutic Classes		
Therapeutic Class Number	Therapeutic class description	Members of the therapeutic class
Class 1	Anti-allergic eye drops	Cromoglicic Acid 2%, (Sodium Cromoglycate) ophthalmic drops, 10ml
Class 1		vs Azelastine 0.5mg/ml ophthalmic drops, 10ml



HP07-2026DAI – Therapeuti	HP07-2026DAI – Therapeutic Classes	
Therapeutic Class Number	Therapeutic class description	Members of the therapeutic class
		vs Epinastine 0.5mg/ml ophthalmic drops, 5ml
		vs Ketotifen 0.25mg/ml ophthalmic drops, 5ml
		vs Lodoxamide 1mg/ml ophthalmic drops, 10ml
		vs Olopatadine 1mg/ml ophthalmic drops, 5ml
		Betaxolol 5mg/ml, ophthalmic drops, 5ml
Class 2	Beta-blocker	vs
		Timolol 0.5%, ophthalmic drops, 5ml
	Alpha-agonist/sympathomimetic - High Concentration	Brimonidine 1.5mg/ml, ophthalmic drops, 5ml
Class 3		vs
		Brimonidine 2mg/ml, ophthalmic drops, 5ml
	Prostaglandin analogue	Bimatoprost 0.1mg/ml, ophthalmic drops, 3ml
		vs
		Bimatoprost 0.03% ophthalmic drops 3ml
Class 4		VS
01055 4		Latanoprost 50mcg/ml, ophthalmic drops, 3ml
		vs
		Latanoprostene 0,024% ophthalmic drops, 5ml
		vs



HP07-2026DAI – Therapeutic Classes		
Therapeutic Class Number	Therapeutic class description	Members of the therapeutic class
		Travoprost 40mcg/ml, ophthalmic drops, 3ml
		Oxybuprocaine 0.4%, ophthalmic drops, 3ml
Class 5	Local Anaesthetics	vs
		Tetracaine 1%, ophthalmic drops, 10ml
		Oxybuprocaine 0.4%, ophthalmic drops, single use applicators, 0.5ml, Box of 20
Class 6	Local Anaesthetics	vs
		Tetracaine 1%, ophthalmic drops, single use applicators, 0.5ml, Box of 20
		Latanoprost and Timolol 50mcg/5mg per ml, ophthalmic drops, 2.5ml
		vs
Class 7	Prostaglandin analogue/Timolol	Timolol and Bimatoprost, 5mg and 0.3mg per ml, ophthalmic drops, 3ml
		vs
		Travaprost and Timolol 40mcg and 5mg/ml, ophthalmic drops, 2.5ml
		Beclometasone nasal spray, 100mcg per actuation, 200-meter dose
		vs
Class 8	Corticosteroid, nasal spray	Budesonide nasal spray, 100mcg per actuation, 200 metered doses
		vs
		Ciclesonide nasal spray, 50mcg per actuation, 120 metered doses
		vs



HP07-2026DAI – Therapeutic Classes		
Therapeutic Class Number	Therapeutic class description	Members of the therapeutic class
		Fluticasone nasal spray, 50mcg per actuation, 120 metered doses
		vs
		Mometasone nasal spray, 50mcg per actuation, 140 metered doses
		vs
		Triamcinolone nasal spray, 55mcg per actuation, 120 metered doses
		Beclometasone inhaler, 100mcg per actuation, 200 doses
Class 9	Inhaled corticosteroid	vs
		Budesonide inhaler, 100mcg per actuation, 300 doses
		Budesonide, Formoterol inhaler 200/6 mcg per actuation 120 doses
		vs
Class 10	LABA/Corticosteroid - ICS/formoterol Combination - Anti-inflammatory and Reliever	Budesonide, Formoterol;Inhaler, 160/4.5mcg peractuation ,120 Doses
	Plus Maintenance Therapy	vs
		Mometasone furoate, formoterol fumarate dihydrate inhaler, 100/5 mcg per actuation, 120 doses
		Budesonide and Formoterol inhaler, 320mcg and 9mcg per actuation, 60 doses
		vs
Class 11	LABA/Corticosteroid - Chronic management of COPD (Group E eosinophils >/= 0.1)	Fluticasone and Salmeterol inhaler, 250mcg and 50mcg per actuation, 60 doses
		vs
		Formoterol/Mometasone furoate 200/5 mcg per actuation, 120 doses



20.3. SERIES AWARDS

Items will be considered to be awarded in a series where:

Dose titration is required e.g. a single molecule in a class is awarded across all strengths and pack sizes to allow for incremental dosing. Such an approach is required to ensure seamless dose titration, simplify supply and distribution, support healthcare worker use and acceptance, and improve patient adherence.

Series awards not applicable for this tender.

20.4. PROCUREMENT CLASS

A procurement class is a grouping of medicines with the same active ingredient but different pack sizes, strengths, formulations, or dosage forms. It is used when market competition is limited, or specific product requirements are not clinically essential. Placing these items in a procurement class promotes fair comparison, competition, and economies of scale, while allowing flexibility to adapt to market availability and ensure continued access. Only one item specification is awarded within a procurement class, though the award may be split between suppliers if needed. During price evaluation the cost per tablet will be considered

HP07-2026DAI – Procurement Classes		
Procurement Class Number	Number Members of the procurement classes	
	Budesonide inhaler, 200mcg per actuation, 300 doses	
Class 1	vs	
	Beclometasone inhaler, 200mcg per actuation, 200 doses	
	Formoterol inhaler, 12mcg per actuation, 120 doses	
	vs	
Class 2	Formoterol inhaler, 9-12mcg per actuation, 60 doses	
	vs	
	Salmeterol inhaler, 50mcg per actuation 60 Doses	

20.5. REFERENCE PRICING

20.5.1. PRICE THRESHOLD REFERENCE PRICING

A price threshold reference price when included represent the maximum allowable price to be considered during price evaluation for including a clinically recommended medicine or vaccine on contract. This threshold is based on the current standard of care or an allowable variance from a reference product or alternative. If awarded, the item will be published with a benchmark reference price for ongoing price monitoring in future.

Price threshold reference price is not applicable for this tender.

20.5.2. BENCHMARK REFERENCE PRICING

A benchmark reference price by the Department as a proactive cost-containment measure to ensure affordability and long-term sustainability. This price is typically informed by local procurement data, international pricing benchmarks, and relevant market intelligence. The benchmark reference price serves as the recommended price and informs price negotiations and contract award decisions.

Benchmark referencing pricing is not applicable for this tender.

20.6. NEGOTIATIONS

The NDoH reserves the right to negotiate prices, minimum order quantities, and supply volumes with bidders prior to the award of the contract. The negotiation process will be conducted at the discretion of the NDoH and in a manner it deems appropriate.

Proposed minimum order quantities (MOQs) should facilitate direct delivery to health establishments.

Where applicable, if an item is advertised as a single item but is included in a therapeutic class and is recommended for award within that class, the Department reserves the right to combine



the volumes and award only one item number. In such cases, the Department will negotiate the awarding of combined volumes with the preferred bidder/s.

In addition, the NDoH reserves the right to review prices, minimum order quantities, and supply volumes with successful bidders after the contract award, as part of the contract management process. For more information on price adjustments based on systematic review refer to section 22.

20.7. NON-COMMITMENT

The NDoH reserves the right not to award, in part or in full. The Department also reserves the right to withdraw or amend any of the bid conditions, by providing notice in writing to all bidders prior to the closing of the bid or post-award.

If an incorrect award has been made, the NDoH reserves the right to remedy the matter in any manner it deems fit, including the cancellation of the contract.

21. POST AWARD CONDITIONS

Regulation 16(A)6.6 of the Treasury Regulations, issued under the Public Finance Management Act, 1999 (Act No. 1 of 1999), allows the Accounting Officer of a department, constitutional institution, or public entity to request participation in any contract arranged through a competitive bidding process by any state organ. This participation requires written approval from both the state organ and the relevant contracted suppliers.

The NDoH may change treatment protocols and/or product formulations where required, due to emerging clinical evidence, disease profiles, safety or resistance patterns, and the availability of items registered in terms of the Medicines Act at the date and time of bid closure. In these circumstances, the NDoH reserves the right to cancel the contract for an item or adjust the quantity awarded based on projected changes in demand. The Department will notify the contracted supplier within a reasonable time of the expected change. However, where patient safety is a concern, these changes may be implemented with immediate effect.

22. PRICE REVIEW



The NDoH anticipates three types of price review processes that may be implemented during the duration of this contract:

- A routine adjustment to mitigate foreign exchange fluctuations;
- An exceptional adjustment to mitigate significant short-term foreign exchange fluctuations; and
- A systematic review of prices for comparable products available in the local and international marketplaces.

22.1. ELIGIBILITY RELATING TO RATE OF EXCHANGE ADJUSTMENTS

Eligibility for price adjustments relating to foreign exchange risk depends on the submission of a complete price breakdown per instructions below for all relevant products; and assessment of the rationality of this price breakdown by the NDoH.

22.2. INSTRUCTIONS FOR PRICE BREAKDOWN

- The price breakdown must be completed on the signed bid response document as well as the electronic version. The delivered price must be divided across five components.
 - Active Pharmaceutical Ingredient/s (API);
 - Formulation;
 - Packaging;
 - Logistics (this includes transportation, warehousing, and distribution);
 - Gross margin (remaining portion).
- The sum of these categories must be equal to 100% of the delivered price for the line item.
- The local + imported portions of the first three components must add up to 100% within each component (e.g. Portion of API attributable to local + Portion of API attributable to import = 100% of specific API component).
- VAT must be apportioned equally across all components and not regarded as a separate component.
- Labour must be apportioned appropriately across the relevant components.
- Breakdown must be in percentage format to the closest whole percentage (e.g. 20%).



- The NDoH reserves the right to engage with bidders to verify any of the components of the bid price, which may include audit of invoices and related documentation.
- Items for which price breakdowns were not presented in the prescribed format at the time of bid closure, will render such item(s) ineligible for price adjustments.

22.3. PRICE ADJUSTMENTS RELATING TO FOREIGN EXCHANGE RISK

Only the portion of the bid price facing foreign exchange risk will be adjusted. This portion is determined by the price breakdown on the signed bid submission.

Adjustments are always calculated using the original awarded contracted price as the base.

Price adjustments relating to foreign exchange will be based on the percentage change between the relevant base average rate of exchange (RoE) and an adjustment average RoE. Rates are sourced from the Reserve Bank (www.resbank.co.za).

Eligibility for favourable Contractual Price Adjustments may be withdrawn considering evidence of poor compliance with contractual obligations.

Base average RoE for this tender will be as follows, per currency:

CURRENCY	BASE AVERAGE RATES OF EXCHANGE AVERAGE FOR THE PERIOD 01 MARCH 2025 TO 31 AUGUST 2025
Rand per US Dollar	R18.09
Rand per Br Pound	R24.09
Rand per Euro	R20.54
Rand per Yuan Renminbi	R2.51
Rand per Indian Rupee	R0.21
Rand per Swiss Franc	R21.86
Rand per Australian Dollar	R11.62
Rand per Danish Krone	R2.75

Should the bidder make use of any currency not mentioned above, the bidder must stipulate this clearly and submit the calculated average RoE for the period 1 March 2025 to 31 August 2025 using the South African Reserve Bank published rates for the specific currency.



22.4. APPLICATION FOR CONTRACTUAL PRICE ADJUSTMENTS

Official applications for price adjustment consideration must be submitted to the NDoH at cpapharma@health.gov.za before the submission deadlines specified in the tables below.

The application must contain the following information:

- Contract description;
- Date of application;
- CPA cycle applied for;
- Items to be considered for CPA (Item no, NSN and Description).

The application must be submitted on a company letter head, signed, scanned and submitted to the CPA mailbox (cpapharma@health.gov.za) no later than the submission date as indicated in the table below.

Where no application for an adjustment relating to foreign exchange has been received and such an adjustment would be favourable to the Department, this will be implemented automatically.

Foreign exchange adjustments may never result in a price exceeding the current Single Exit Price. With reference to paragraph 4.1, the supplier will only be eligible for contractual price adjustments up to the most recent Single Exit Price value as recorded in the National Department of Health (NDoH) SEP Database.

22.4.1. EXCEPTIONAL PRICE ADJUSTMENTS BEFORE START OF CONTRACT

The contracted supplier may apply for an exceptional price adjustment before the start of the contract. These will be activated if the absolute change between the base RoE and the six-month retrospective average RoE indicated in the table below fluctuates by more than 10%. This adjustment applies to eligible components subject to CPA price adjustments based on the bid closure price.



REVIE	PERIOD FOR CALCULATING ADJUSTMENT AVERAGE ROE	SUBMISSION OF REQUEST FOR PRICE REVIEW TO REACH THE OFFICE BY	DATE FROM WHICH ADJUSTED PRICES WILL BECOME EFFECTIVE
0.01	01 February 2026 – 31 July 2026	03 August 2026	01 September 2026

22.4.2. **ROUTINE PRICE ADJUSTMENTS**

Schedules for routine price reviews, and periods for calculating adjustment average RoE are detailed in the table below:

REVIEW	PERIOD FOR CALCULATING ADJUSTMENT AVERAGE ROE	SUBMISSION OF REQUEST FOR PRICE REVIEW TO REACH THE OFFICE BY	DATE FROM WHICH ADJUSTED PRICES WILL BECOME EFFECTIVE
1	01 September 2026 – 28 February 2027	03 March 2027	01 April 2027
2	01 March 2027 – 31 August 2027	03 September 2027	01 October 2027
3	01 September 2027 – 28 February 2028	03 March 2028	01 April 2028
4	01 March 2028 – 31 August 2028	03 September 2028	01 October 2028
5	01 September 2028 – 28 February 2029	03 March 2029	01 April 2029

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22.4.3. EXCEPTIONAL PRICE ADJUSTMENTS DURING CONTRACT PERIOD

Contracted suppliers may request exceptional price adjustments during the contracted period according to the schedule in the table below. These will be activated if the absolute change between the base RoE and the three-month retrospective average RoE indicated in the table below fluctuates by more than 10%.

REVIEW	PERIOD FOR CALCULATING ADJUSTMENT AVERAGE ROE	SUBMISSION OF REQUEST FOR PRICE REVIEW TO REACH THE OFFICE BY	DATE FROM WHICH ADJUSTED PRICES WILL BECOME EFFECTIVE
0.1	01 September 2026 – 30 November 2026	03 December 2026	01 January 2027
1.1	01 March 2027- 31 May 2027	03 June 2027	01 July 2027
2.1	01 September 2027 – 30 November 2027	03 December 2027	01 January 2028
3.1	01 March 2028- 31 May 2028	03 June 2028	01 July 2028
4.1	01 September 2028 – 30 November 2028	03 December 2028	01 January 2029
5.1	01 March 2029- 31 May 2029	03 June 2029	01 July 2029

Suppliers who received exceptional adjustments will, thereafter, receive routine adjustments based on the average exchange rate over the preceding three months, rather than the standard six-month historical average. The specific periods used to calculate the average rate of exchange (RoE) for these adjustments are outlined in the table below:

REVIEW	PERIOD FOR CALCULATING ADJUSTMENT AVERAGE ROE POST EXCEPTIONAL ADJUSTMENT	SUBMISSION OF REQUEST FOR PRICE REVIEW TO REACH THE OFFICE BY	DATE FROM WHICH ADJUSTED PRICES WILL BECOME EFFECTIVE
1	01 December 2026 – 28 February 2027	03 March 2027	01 April 2027
2	01 June 2027 – 31 August 2027	03 September 2027	01 October 2027
3	01 December 2027 – 28 February 2028	03 March 2028	01 April 2028



REVIEW	PERIOD FOR CALCULATING ADJUSTMENT AVERAGE ROE POST EXCEPTIONAL ADJUSTMENT	SUBMISSION OF REQUEST FOR PRICE REVIEW TO REACH THE OFFICE BY	DATE FROM WHICH ADJUSTED PRICES WILL BECOME EFFECTIVE
4	01 June 2028 – 31 August 2028	03 September 2028	01 October 2028
5	01 December 2028 – 28 February 2029	03 March 2029	01 April 2029

22.5. PRICE ADJUSTMENTS BASED ON A SYSTEMATIC REVIEW

The NDoH reserves the right to review both local and international market prices to identify the lowest comparable pricing. Should this review reveal prices lower than those stipulated in the contract, the Department may initiate price negotiations with the contracted supplier.

If the outcome of this negotiation is deemed unfavourable, the NDoH reserves the right to terminate the award for the item in question.

23. QUALITY

Products and contracted suppliers must conform to the conditions of registration of the product in terms of the Medicines Act for the full duration of this contract. In the event that the product and or contracted supplier does not conform to the conditions of registration of the product, NDOH reserves the right to cancel the contract.

24. DELIVERY AND QUANTITIES

24.1. DELIVERY BASIS

Firm lead times for delivery must be quoted for the duration of the contract period.

Transit and storage conditions applicable to the relevant products must be always adhered to.

The initial lead time, as proposed in the bid response document, will be calculated from the date of award of the contract and not from the date of placement of the first order. This lead time may



not exceed 75 calendar days from the date of award from when the contract circular signed by the National Department of Health has been published.

Lead time within the contract period is defined as the time from the submission of the order to the supplier to the time of receipt by the Department, as confirmed by the Proof of Delivery document. This lead time may not exceed 14 calendar days.

Failure to comply with the contractual lead time may result in penalties being enforced, as per sections 21 and 22 of the General Conditions of Contract (GCC).

24.2. QUANTITIES

The quantities reflected in the bid are estimated and no guarantee, either explicit or implied, is given regarding the actual quantity that will be procured during the contract period. Fluctuations in monthly demand may occur.

The NDoH reserves the right to negotiate MOQs where necessary. In cases where consensus regarding MOQs cannot be reached, the bid may not be awarded.

Suppliers are required to maintain sufficient buffer stock to meet at least two months' demand for all items, in alignment with the needs of Participating Authority/Authorities.



SECTION C

25. SUPPLIER PERFORMANCE MANAGEMENT

Supplier performance management will be the responsibility of the Participating Authorities, with oversight from the NDoH. If supplier performance disputes cannot be resolved between the contracted supplier and the Participating Authority, the NDoH must be informed for corrective action.

The NDoH, in collaboration with Participating Authorities, will monitor the performance of contracted suppliers throughout the duration of this contract. This will include, but is not limited to, the following areas:

- Ongoing supplier performance monitoring through compliance visits
- Adherence to reporting requirements
- Attendance of quarterly supplier meetings
- Execution of orders and delivery performance
- Management of order cancellations and product substitutions
- Identification and correction of irrational or misaligned orders
- Delivery schedule adherence
- Assurance of continuity of supply
- Compliance with the administrative, legislative and regulatory requirements as specified in the SRCC.

25.1. COMPLIANCE WITH REPORTING REQUIREMENTS:

Suppliers must adhere to the reporting schedule and mechanism established by the NDoH. At a minimum, suppliers must submit the following information in the specified format and mechanism, after receiving training provided by the NDoH:

- All transactional data relating to orders
- A monthly age analysis
- Production pipeline data and forecasts, including:
 - Number of units of the item available (stock on hand)
 - Number of units of the item in Quality Assurance, awaiting release



- Number of units of the item in the current month's production plan
- Status of outstanding orders

25.2. ATTENDANCE OF QUARTERLY MEETINGS

The NDoH will schedule and hold quarterly meetings with contracted suppliers. These meetings will include, but not be limited to, a review of supplier performance and the forecasted demand for the next quarter. Suppliers may be required to present continuous improvement initiatives aimed at improving efficiencies in the supply chain, benefiting both suppliers and Participating Authorities.

25.3 ORDER PLACEMENT AND DELIVERY

- Orders will be placed as needed during the contract period, with delivery points specified by the relevant Participating Authority/Authorities.
- The instructions on the official order form regarding supply, dispatch, and submission of invoices must be strictly adhered to.
- Under no circumstances should the contracted supplier deviate from the orders issued by the Participating Authority/Authorities, unless written instruction is received from the relevant participating authority.
- Changes to any quantities ordered may only be made upon receipt of an amended purchase order.
- A Participating Authority is under no obligation to accept any quantity that exceeds the ordered quantity.
- To facilitate the efficient implementation of the direct delivery strategy, contracted suppliers must pack orders according to the purchase order for the relevant health establishment.
- Only orders made using an official, authorized purchase order format are valid.
- Suppliers must acknowledge receipt of all purchase orders received from Participating Authorities in the manner stipulated by the relevant Participating Authority.

Authorised Signatory: Sign or Initial



25.4 ORDER CANCELLATIONS AND SUBSTITUTION

The Participating Authority/Authorities reserve the right to cancel any order if the lead time exceeds 14 days. In such instances, they may, at their discretion, procure supplies of equivalent quality and quantity as a substitute for the goods not delivered in accordance with the contract, in line with Section 21.6 of the General Conditions of Contract.

Should this occur, the Participating Authority may source the item from an alternative supplier, and any cost difference between the contracted supplier's price and the price of the substitute item will be for the account of the contracted supplier.

25.5 IRRATIONAL OR MISALIGNED ORDERS

In cases where an order is received that appears to be irrational or misaligned with estimates, the contracted supplier must consult the relevant Participating Authority prior to processing the order.

In the event of short supply, incorrect delivery, or misaligned orders, the supplier must issue a credit note within 15 calendar days of receiving both the credit request and the relevant supporting documentation from the participating authority.

25.6 DELIVERY ADHERENCE

- Products and related documentation must be delivered in accordance with the terms,
 conditions, and delivery instructions stipulated in the purchase order.
- The information on invoices and documents relating to delivery must comply with the minimum data requirements as defined by the NDoH. The NDoH reserves the right to update these minimum data requirements as needed (Annexure B).
- Invoices must clearly reflect both the "proprietary name" (brand name/trade name), which is unique to a particular medicine and approved under section 15(4) of the Medicines Act, and the item description as it appears in the contract circular and Master Health Product List (MHPL).



- The supplier must ensure that products are delivered in accordance with the appropriate storage conditions, as per the product's conditions of registration. Delivery is deemed complete upon signature of receipt by the delegated official.
- Any discrepancies between the invoice and the physical stock, or damaged stock, must be reported to the contracted supplier within a reasonable time, or as otherwise arranged with the supplier. This period should allow for verification of the quantities received upon delivery.

Contracted suppliers will be responsible for the collection of goods delivered erroneously or in an incorrect condition, as formally arranged in consultation with the Participating Authorities. The Participating Authorities may recoup any expenses associated with the failure to collect such goods in accordance with the agreement.

25.7 CONTINUITY OF SUPPLY

Contracted suppliers must maintain at least two months' supply of the estimated quantity at the start of the contract and ensure a continuous supply throughout the contract's duration. If order fulfilment for a specific item deviate by 20% from the average monthly estimate for three consecutive months on a rolling basis, suppliers must notify the NDoH/Contract Management Unit (CMU) within two weeks of becoming aware of the discrepancy. In such cases, the supplier should engage with the NDoH and the relevant participating authority to update the demand forecast, align supply volumes accordingly, and prevent supply challenges.

Suppliers are expected to engage regularly with Participating Authorities to review demand and plan proactively to ensure uninterrupted supply.

Contracted suppliers must promptly inform all participating authorities and NDoH of any circumstances that may result in an interrupted supply, including but not limited to:

- Regulatory actions that may impact their GMP status or the status of entities on which they rely;
- Anticipated issues with the availability of active pharmaceutical ingredients (API);
- Industrial actions:



- Challenges with the manufacturing pipeline;
- Any other supply-related challenges.

Official communication regarding continuity of supply should be directed to stockalert@health.gov.za, as well as the Participating Authorities.

Official communication regarding payment challenges should be directed to medacc@health.gov.za, as well as the relevant Participating Authorities.

All official communications must include details of corrective actions taken by the contracted supplier to ensure continuous supply.

If the contracted supplier is unable to supply the awarded item, the supplier is required to source an alternative product that meets the same specifications.

In the case of a split or multiple awards, the alternative product must not be sourced from another contracted supplier for the same product. The alternative product must be supplied at the current price of the contracted item.

Prior to supplying an alternative product including items authorised for procurement utilising Section 21 and Section 36 of the Medicines Act, the contracted supplier must seek approval from the NDoH and provide a sample to the two health establishments as outlined in section 5.5 of this SRCC. The contracted supplier must also provide the following information to the NDoH:

- Name of the product to be supplied;
- Quantities to be supplied;
- The period for which the product will be supplied. This provision applies only to emergency supply situations and cannot be used for routine or continuous supply.

If a contracted supplier is unable to supply the contracted item for a period **not exceeding six months**, the NDoH reserves the right to reallocate volumes proportionally to an alternative contracted supplier for the duration of the supply interruption.

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If a contracted supplier is unable to supply a contracted item for a period **exceeding six months** for any reason, the NDoH reserves the right to cancel the contract, as outlined in Section 23 of the General Conditions of Contract (GCC), Clause 21.2.

Suppliers may be penalized for failing to meet the contractual lead time, as stipulated in Section 22 of the GCC.

26. REPORTING

The NDoH will provide the requirements for reporting and successful bidders will be assisted with complying with these requirements. The National Department of Health may, from time to time and within reason, add to the reporting requirements. Any changes to reporting requirements or the reporting mechanism will be communicated in writing by the Directorate: Affordable Medicines.

27. PACKAGING, LABELLING AND BARCODES

27.1. PACKAGING

- Suppliers must ensure that products delivered are received in good order at the point of delivery. Packaging must be suitable for further dispatch, storage and stacking according to Good Wholesaling Practice and Good Distribution Practice.
- Packaging must be suitable for transportation and should prevent exposure to conditions that could adversely affect the stability and integrity of the product.
- The packaging must be uniform for the duration of the contract period. All products must be packaged in acceptable containers, specifically developed for the product.
- Any change to the packaging must be approved by the NDoH.
- All medicines must be supplied in complete, patient-ready packaging in the specified
 pack size, using containers that are properly sealed and labelled in compliance with
 Medicines Act. Packaging must be in a ready-to-dispense format that does not require
 any manipulation, packing, or repacking by the dispensing healthcare workers.
- The number of units per shipper pack or original carton must be completed in the Bid Response Document.
- Where the supplier recommends a particular stacking and storage configuration,



this should be clearly illustrated on the outer packaging.

- Where the contents of the shipper pack represent a standard supply quantity of an item, the following must be adhered to:
 - Outer packaging flanges must be sealed with suitable tape that will clearly display evidence of tampering.
 - The contents must be packed in neat, uniform rows and columns that will facilitate easy counting when opened.
- Where the contents of a shipper pack represent a non-standard supply quantity,
 the following must be adhered to:
- Outer packaging flanges must be sealed with suitable tape that will clearly display evidence of tampering.
- The shipper pack must contain only one product, mixing multiple products in a single shipper is not allowed.
- The outer packaging must be clearly marked as a "Part Box".

27.2. LABELLING

- All containers, packaging and cartons must be clearly labelled. Bulk packs must be labelled in letters not less than font size 48.
- The following information must be clearly and indelibly printed on both corners (length and breadth) all shipper packs, including any part boxes:
 - ltem name as contained in the contract circular and the Master Health Product List (MHPL),
 - Registered product name;
 - Number of units in pack;
 - Batch number;
 - Expiry date;
 - Storage conditions;
 - Barcode.
- Where the contents of the shipper pack require special attention in terms of storage and/or handling, e.g., thermolabile, high-scheduled or cytotoxic products, such instructions must be clearly and visibly indicated on the outer packaging on

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a brightly colored background.

 Unit packs must be labelled in accordance with Regulation 10 of the General Regulations published in terms of the Medicines Act.

27.3. BARCODES

- All unit and shipper packs should be marked with the appropriate barcode.
- The European Article Numbering Code 13 (EAN 13).

27.4. SHELF LIFE

- Unless SAHPRA has approved a shorter shelf life, products must have a shelf-life of at least 12 months upon delivery.
- Contracted suppliers may apply in writing to Participating Authorities to supply a product with a shorter shelf life provided that:
 - Applications are accompanied by an undertaking that such short-dated products
 will be unconditionally replaced or credited before or after expiry and,
 - Applications are approved by the Participating Authorities before execution of orders; and,
 - Upon notification of the remaining expired stock, such products will be collected and disposed of by the supplier at their own cost and,
 - Failure to collect the products within 30 days after written notification to the supplier will result in the disposal of the product by the Participating Authority for the account of the supplier.
- Unless otherwise agreed to, any Participating Authority may, without prejudice, decline to accept the product with a shelf-life of less than 12 months.

28. DISCONTINUATION OF CONTRACTED PRODUCT SUPPLY

It is the responsibility of the contracted supplier to ensure continuous supply of the contracted product until the end date of the contract, as stipulated in the Letter of Acceptance (SDB 7.1).

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If the contracted supplier foresees a potential long-term interruption in supply, the supplier must submit a written letter to the Director-General of Health at least six months prior to the anticipated interruption. The letter must include the following:

- The reason for the long-term interruption.
- The impact this will have on the contract.
- The proposed solution or suggested way forward.

The supplier may only interrupt supply to a Participating Authority after informing the Director-General of Health and receiving written approval from the NDoH. It is the responsibility of the NDoH to communicate the outcome of this matter to the Participating Authorities.

If the contracted supplier decides to discontinue a contracted product with immediate effect, the Department reserves the right to source the item from an alternative supplier. If the price from the alternative supplier exceeds the contracted price, the supplier discontinuing the product will be liable for the price difference for a period of six months.

29. CEDING, MERGERS, TAKE OVERS AND CHANGES IN SUPPLIER DETAILS

If a contracted supplier plans to merge with or be acquired by another entity, or intends to cede the contract to another supplier, the contracted supplier must inform the NDoH in writing as soon as they become aware of such an event.

Should the contracted supplier plan to cede a contracted item to another supplier, they must submit an official request in writing to the NDoH at least three months prior to the proposed effective date. The NDoH reserves the right to either accept or decline the request to transfer the contractual obligations to the new supplier under the current terms of the contract, or to cancel the contract altogether.

The contracted supplier is also required to inform the NDoH as soon as they become aware of any changes to their address, name, or contact details. These updates must also be reflected on the Central Supplier Database (CSD).



30. CANCELLATION OF CONTRACT

A request for the cancellation of a contract from a contracted supplier will only be considered if:

- A formal cancellation request in writing addressed to the Director-General: National Department of Health; and
- Evidence in support of the request is submitted.

The contracted supplier is obligated to continue supplying the contracted item under the existing terms and conditions of the contract until the NDoH has formally approved the cancellation request. Once approved, the NDoH will notify the Participating Authorities of the contract cancellation.

31. THIRD PARTIES

Participating Authorities will not make payment to or consult with a third party. No third party is entitled to put an account of a Participating Authority/Authorities on hold.

END

Item No	Item Specification	Therapeutic Class / Series	UNIT (Use for Estimate & Price)	Estimate
1	Acetic Acid 2% in Alcohol 50%, ear drops, 10ml		Each	418 100
2	Acetylcholine 20mg/2ml, ophthalmic solution, 2ml		Each	9 052
3	Atropine 1%, ophthalmic drops, 5ml		Each	293 618
4	Azelastine 0.5mg/ml ophthalmic drops, 10ml	Class 1	Each	807 962
5	Beclometasone inhaler, 50mcg per actuation, 200 doses		Each	969 800
6	Beclometasone inhaler, 100mcg per actuation, 200 doses	Class 9	Each	1 352 540
7	Beclometasone inhaler, 100mcg per actuation, 200 doses		Each	271 846
8	Beclometasone inhaler, 200mcg per actuation, 200 doses	Procurement Class 1	Each	2 986 970
9	Beclometasone nasal spray, 50mcg per actuation, 150-200 metered doses		Each	181 609
10	Beclometasone nasal spray, 100mcg per actuation, 200 meter dose	Class 8	Each	3 876 540
11	Betaxolol 5mg/ml, ophthalmic drops, 5ml	Class 2	Each	226 182
12	Bimatoprost 0.03% ophthalmic drops 3ml	Class 4	Each	519 592
13	Bimatoprost 0.1mg/ml, ophthalmic drops, 3ml	Class 4	Each	519 592
14	Brimonidine 1.5mg/ml, ophthalmic drops, 5ml	Class 3	Each	992 277
15	Brimonidine 2mg/ml, ophthalmic drops, 5ml	Class 3	Each	992 277
16	Brimonidine and Timolol, 2mg and 5mg/ml, ophthalmic drops, 5ml		Each	297 550
17	Budesonide and Formoterol inhaler, 320mcg and 9mcg per actuation, 60 doses	Class 11	Each	1 647 940
18	Budesonide inhaler, 100mcg per actuation, 300 doses	Class 9	Each	1 352 540
19	Budesonide inhaler, 200mcg per actuation, 300 doses	Procurement Class 1	Each	1 996 380
20	Budesonide nasal spray, 100mcg per actuation, 200 metered doses	Class 8	Each	3 876 540
21	Budesonide, Formoterol inhaler 200/6 mcg per actuation 120 doses	Class 10	Each	5 933 125
22	Budesonide, Formoterol;Inhaler, 160/4.5mcg peractuation ,120 Doses	Class 10	Each	5 933 125
23	Carboxymethylcellulose sodium 5mg/ml, ophthalmic drops, 15ml-20ml		Each	983 000
24	Ciclesonide nasal spray, 50mcg per actuation, 120 metered doses	Class 8	Each	3 876 540

Item No	Item Specification	Therapeutic Class / Series	UNIT (Use for Estimate & Price)	Estimate
25	Cromoglicic Acid 2%, (Sodium Cromoglycate) ophthalmic drops, 10ml	Class 1	Each	807 962
26	Cyclopentolate 1%, ophthalmic drops, 15ml		Each	1 770
27	Cyclopentolate 1%, ophthalmic drops, single use applicators, 0.5ml, Box of 20		Box of 20	478
28	Cyclopentolate and Phenylephrine, 2mg and 10mg/ml, ophthalmic drops, 5ml		Each	83 364
29	Dexamethasone 0.1%, ophthalmic drops, 5ml		Each	565 289
30	Dorzolamide and Timolol, 20mg and 5mg/ml, ophthalmic drops, 5ml		Each	338 667
31	Epinastine 0.5mg/ml ophthalmic drops, 5ml	Class 1	Each	807 962
32	Fluorescein 1mg, ophthalmic sterile strips, Box of 100		Box of 100 strips	396
33	Fluorescein 2%, ophthalmic drops, single use applicators, 0.5ml, Box of 20		Box of 20	5 492
34	Fluticasone and Salmeterol inhaler, 125mcg and 25mcg per actuation,120 doses		Each	37 638
35	Fluticasone and Salmeterol inhaler, 250mcg and 50mcg per actuation, 60 doses	Class 11	Each	1 647 940
36	Fluticasone and Salmeterol, inhaler, 50mcg and 25mcg per actuation, 120 doses		Each	21 965
37	Fluticasone and Salmeterol, inhaler, 250mcg and 25mcg per actuation, 60 doses		Each	128 435
38	Fluticasone nasal spray, 50mcg per actuation, 120 metered doses		Each	841 470
39	Fluticasone nasal spray, 50mcg per actuation, 120 metered doses	Class 8	Each	3 876 540
40	Formoterol inhaler, 9-12mcg per actuation, 60 doses	Procurement Class 2	Each	85 076
41	Formoterol inhaler, 12mcg per actuation, 120 doses	Procurement Class 2	Each	42 538
42	Formoterol/Mometasone furoate 200/5 mcg per actuation, 120 doses	Class 11	Each	1 647 940
43	Hyaluronic Acid 10mg/ml, intraocular injection, 0.55ml		Each	167 685
44	Hyaluronidase 1500iu, powder for solution, ampoule		Each	51 950
45	Hydroxypropylmethylcellulose 3mg/ml, ophthalmic drops, 20ml		Each	1 188 596
46	Inhaler device (Universal spacer) with mouthpiece, for adults, made of a suitable antistatic material with spacer volume 500ml with multi-inhaler adaptor		Each	14 305
47	Inhaler device (Universal spacer) with mask (fixed or removable), for infants, non-return valve, space volume 150ml to 250ml, made of a suitable anti- static material with multi-inhaler adaptor		Each	23 145

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Item No	Item Specification	Therapeutic Class / Series	UNIT (Use for Estimate & Price)	Estimate
48	Inhaler device (Universal spacer) with mouthpiece, for children , non-return valve, space volume between 250ml and 500ml, made of a suitable anti- static material with multi-inhaler adaptor		Each	42 055
49	Ipratropium Bromide 0.25mg/2ml, respirator solution, single dose vials, Box of 60		Box of 60 vials	102 503
50	Ipratropium Bromide and Salbutamol, 0.5mg and 2.5mg/ 2.5ml, respirator solution, Box of 60		Box of 60	223 421
51	Ipratropium Bromide inhaler, 20mcg per actuation, 200 doses		Each	72 394
52	Ipratropium Bromide, 0.5mg/2ml, respirator solution, single dose vials, Box of 60		Box of 60 vials	117 684
53	Ketorolac 5mg/ml, ophthalmic drops, 5ml		Each	619 994
54	Ketotifen 0.25mg/ml ophthalmic drops, 5ml	Class 1	Each	807 962
55	Lanolin 3%, ophthalmic ointment, 3.5g		Each	1 132 463
56	Latanoprost 50mcg/ml, ophthalmic drops, 3ml	Class 4	Each	519 592
57	Latanoprost and Timolol 50mcg/5mg per ml, ophthalmic drops, 2.5ml	Class 7	Each	947 856
58	Latanoprostene 0,024%, ophthalmic drops 5ml	Class 4	Each	519 592
59	Lodoxamide 1mg/ml ophthalmic drops, 10ml	Class 1	Each	807 962
60	Mometasone furoate, formoterol fumarate dihydrate inhaler, 100/5 mcg per actuation, 120 dose	Class 10	Each	5 933 125
61	Mometasone nasal spray, 50mcg per actuation, 140 metered doses	Class 8	Each	3 876 540
62	Olopatadine 1mg/ml ophthalmic drops, 5ml	Class 1	Each	807 962
63	Oxybuprocaine 0.4%, ophthalmic drops, 3ml	Class 5	Each	34 520
64	Oxybuprocaine 0.4%, ophthalmic drops, single use applicators, 0.5ml, Box of 20	Class 6	Box of 20	11 396
65	Oxymetazoline 0.025%, nasal drops, 10ml		Each	2 026 152
66	Oxymetazoline 0.025%, ophthalmic drops, 15ml		Each	3 592 369
67	Oxymetazoline 0.05%, nasal spray, 15ml		Each	2 203 322
68	Pilocarpine 1%, ophthalmic drops, 15ml		Each	19 690
69	Polyacrylic acid (Carbomer) 2 mg/g ophthalmic gel, 10g		Each	1 080
70	Prednisolone 1%, ophthalmic drops, 5ml		Each	266 981

Item No	Item Specification	Therapeutic Class / Series	UNIT (Use for Estimate & Price)	Estimate
71	Salbutamol 0.5%, respirator solution, 20ml		Each	235 816
72	Salbutamol inhaler, 100mcg per actuation, 200 doses		Each	5 802 201
73	Salmeterol inhaler, 50mcg per actuation 60 Doses	Procurement Class 2	Each	85 076
74	Sodium Chloride 0.9% nasal, 20ml-50ml		Each	227 456
75	Sodium Chloride 0.9%, isotonic sterile solution for irrigation, 30ml		Each	60 442
76	Sodium Chondroitin Sulphate and Sodium Hyaluronate, 40mg and 30mg/ml, visco-elastic opthalmic solution, 0.5ml prefilled syringe.		Each	167 685
77	Sterile intraocular Irrigating Solution, containing per ml: Sodium Chloride 6.4mg, Sodium Acetate 3.9mg, Sodium Citrate 1.7mg, Potassium Chloride 0.75mg, Calcium Chloride 0.48mg and Magnesium Chloride 0.3mg, 15ml		Each	284 935
78	Sterile intraocular Irrigating solution, containing per ml: Sodium Chloride 6.4mg, Sodium Acetate 3.9mg, Sodium Citrate 1.7mg, Potassium Chloride 0.75mg, Calcium Chloride 0.48mg and Magnesium Chloride 0.3mg, 500ml		Each	78 370
79	Tetracaine 1%, ophthalmic drops, 10ml	Class 5	Each	34 520
80	Tetracaine 1%, ophthalmic drops, single use applicators, 0.5ml, Box of 20	Class 6	Box of 20	11 396
81	Timolol 0.25%, ophthalmic drops, 5ml		Each	37 800
82	Timolol 0.5%, ophthalmic drops, 5ml	Class 2	Each	226 182
83	Timolol and Bimatoprost, 5mg and 0.3mg per ml, ophthalmic drops, 3ml	Class 7	Each	947 856
84	Travaprost and Timolol 40mcg and 5mg/ml, ophthalmic drops, 2.5ml	Class 7	Each	947 856
85	Travoprost 40mcg/ml, ophthalmic drops, 3ml	Class 4	Each	519 592
86	Tropicamide 1%, ophthalmic drops, 15ml		Each	57 317
87	Triamcinolone nasal spray, 55mcg per actuation, 120 metered doses	Class 8	Each	3 876 540
88	Tropicamide 1%, ophthalmic drops, single use applicators, 0.5ml, Box of 20		Box of 20	1 218