

BID DOCUMENT NUMBER: ZNB6685/3/2023-H

DESCRIPTION: SUPPLY AND DELIVERY OF RESPIRATORY: ENDOTRACHEAL TUBES AND TRACHEOSTOMY TUBES AND ACCESSORIES FOR VARIOUS INSTITUTIONS: PERIOD OF 3 YEARS

Name of Bidder.....

Central Supplier's Database Registration Number.....

Income Tax Reference Number.....

BIDDER TO NOTE THE FOLLOWING

CLOSING DATE AND TIME:

DATE: 28 SEPTEMBER 2023

TIME: 11: 00AM

BID RESPONSE DOCUMENTS MUST BE DEPOSITED IN THE BID BOX SITUATED AT (STREET ADDRESS) Central Supply Chain Management Directorate Old Boys School, 310 Jabu Ndlovu Street Pietermaritzburg 3201

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SECTION A: INVITATION TO BID (SBD1)

PART A

YOU ARE HEREBY IN	VVITED TO	BID FOR REQUIREM	ENTS OF THE K	NAZULU-N/	ATAL DE	EPARTME	NT OF HE	ALTH	
BID NUMBER: ZN	NB 6685/3/2	2023-H: CLOSING [DATE: 28	SEPTEMB	ER 2023	CLC	DSING TIM	E:	11: H 00 AM
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CENTRAL SUPPLY C									
OLD BOYS SCHOOL,	, 310 JABU	NDLOVU SIREEI							
PIETERMARITZBURG	3								
3201									
SUPPLIER INFORMA	TION								
NAME OF BIDDER									
POSTAL ADDRESS									
STREET ADDRESS			1						
TELEPHONE NUMBE	R	CODE				NUMBER			
CELLPHONE NUMBE	R		1						
FACSIMILE NUMBER		CODE				NUMBER			
E-MAIL ADDRESS									
VATREGISTRATION	NUMBER		1						
		TCS PIN:			OR	CSD No:			
STATUS LEVEL		🗌 Yes					· □	í es	
VERIFICATION CERT						US LEVEL			
_ [TICK APPLICABLE BOX] INO SWORN AFFIDAVIT									
IF YES, WHO WAS TH									
CERTIFICATE ISSUE	D BY?								
AN ACCOUNTING OF	FICER			NG OFFICEI	R AS CO	ONTEMPLA	TED IN TH	E CLOSE	CORPORATION
AS CONTEMPLATED	IN THE		ACT (CCA)						
CLOSE CORPORATIO	ON ACT						ED BY	THE SO	UTH AFRICAN
(CCA) AND NAME TH			ACCREDITATIO		I (SANAS	S)			
APPLICABLE IN THE	TICK		A REGISTEREI	D AUDITOR					
BOX			NAME:						
[A STATUS LEVEL VERIFICATION CERTIFICATE/SWORN AFFIDAVIT (FOR EMEs& QSEs) MUST BE SUBMITTED IN ORDER TO QUALIFY									
FOR PREFERENCE P		R]		-					
ARE YOU THE ACC	REDITED					YOU A F		Yes	No
REPRESENTATIVE IN	N SOUTH	□Yes	No			D SUPPLI			
AFRICA FOR THE	GOODS					GOOL		[IF YES	ANSWER PART
/SERVICES	/WORKS					ICES /		B:3 BELO	OW]
OFFERED?		[IF YES ENCLOSE F	PROOF]				WURNS		-
			-			RED?			
SIGNATURE OF BIDE		••••••			DATE				
CAPACITY UNDER									
THIS BID IS SIGNED									
proof of authority to	•								
bid; e.g. resolu	tion of								
directors, etc.)					-				
TOTAL NUMBER O	F ITEMS					L BID PRI	CE (ALL		
				TEALINIA	INCLU				
BIDDING PROCEDURE).			MATION M			
DEPARTMENT KZN Departmer		partment of Health		DEPARTM			Department		
CONTACT PERSON	Demand	Management		CONTACT	PERSON		Groenewald	<u>i</u>	
TELEPHONE NUMBER		8361/8386		TELEPHON	IE NUMB	BER 033 3	95 4200		
E-MAIL ADDRESS	NL ADDRESS SCM.DemandManagement@kznhealth.gov.za			E-MAIL AD	DRESS	eden	dale.anaestl	netics@kznl	health.gov.za

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 ONLINE
1.3.	
	NAMELY: (BUSINESS REGISTRATION/ DIRECTORSHIP/ MEMBERSHIP/IDENTITY NUMBERS; TAX COMPLIANCE STATUS; AND
	BANKING INFORMATION FOR VERIFICATION PURPOSES). CERTIFICATE OR SWORN AFFIDAVIT FOR MUST BE SUBMITTED
	TO BIDDING INSTITUTION.
1.4.	WHERE A BIDDER IS NOT REGISTERED ON THE CSD, MANDATORY INFORMATION NAMELY: (BUSINESS REGISTRATION/
	DIRECTORSHIP/ MEMBERSHIP/IDENTITY NUMBERS; TAX COMPLIANCE STATUS MAY NOT BE SUBMITTED WITH THE BID
	DOCUMENTATION. CERTIFICATE OR SWORN AFFIDAVIT FOR MUST BE SUBMITTED TO BIDDING INSTITUTION.
1.5.	THIS BID IS SUBJECT TO THE PREFERENTIAL PROCUREMENT POLICY FRAMEWORK ACT 2000 AND THE PREFERENTIAL
	PROCUREMENT REGULATIONS, 2022, THE GENERAL CONDITIONS OF CONTRACT (GCC) AND, IF APPLICABLE, ANY OTHER
	LEGISLATION OR SPECIAL CONDITIONS OF CONTRACT AND ANY AMENDMENTS THERETO. 2. TAX COMPLIANCE REQUIREMENTS
2.1	BIDDERS MUST ENSURE COMPLIANCE WITH THEIR TAX OBLIGATIONS.
	BIDDERS MOST ENSURE COMPLIANCE WITH THEIR TAX OBLIGATIONS. BIDDERS ARE REQUIRED TO SUBMIT THEIR UNIQUE PERSONAL IDENTIFICATION NUMBER (PIN) ISSUED BY SARS TO ENABLE
	THE DEPARTMENT TO VIEW THE TAXPAYER'S PROFILE AND TAX STATUS.
	APPLICATION FOR TAX COMPLIANCE STATUS (TCS) OR PIN MAY ALSO BE MADE VIA E-FILING. IN ORDER TO USE THIS
	PROVISION, TAXPAYERS WILL NEED TO REGISTER WITH SARS AS E-FILERS THROUGH THE WEBSITE WWW.SARS.GOV.ZA.
	BIDDERS MAY ALSO SUBMIT A PRINTED TCS TOGETHER WITH THE BID.
2.5	IN BIDS WHERE CONSORTIA/ JOINT VENTURES/ SUB-CONTRACTORS ARE INVOLVED, EACH PARTY MUST SUBMIT A
	SEPARATE PROOF OF TCS / PIN / CSD NUMBER.
2.6	WHERE NO TCS IS AVAILABLE BUT THE BIDDER IS REGISTERED ON THE CENTRAL SUPPLIER DATABASE (CSD), A CSD
	NUMBER MUST BE PROVIDED.
	3. QUESTIONNAIRE TO BIDDING FOREIGN SUPPLIERS
3.1.	IS THE BIDDER A RESIDENT OF THE REPUBLIC OF SOUTH AFRICA (RSA)?
3.2.	DOES THE BIDDER HAVE A BRANCH IN THE RSA?
3.3.	
	DOES THE BIDDER HAVE ANY SOURCE OF INCOME IN THE RSA?
	HE ANSWER IS "NO" TO ALL OF THE ABOVE, THEN, IT IS NOT A REQUIREMENT TO OBTAIN A TAX COMPLIANCE STATUS/ TAX
	IPLIANCE SYSTEM PIN CODE FROM THE SOUTH AFRICAN REVENUE SERVICE (SARS) AND IF NOT REGISTERED AS PER 2.3
ABC	DVE.

4

NB: FAILURE TO PROVIDE ANY OF THE ABOVE PARTICULARS MAY RENDER THE BID INVALID.

SECTION B: SPECIAL INSTRUCTIONS AND NOTICES TO BIDDERS REGARDING THE COMPLETION OF BIDDING FORMS

PLEASE NOTE THAT THIS BID IS SUBJECT TO TREASURY REGULATIONS 16A ISSUED IN TERMS OF THE PUBLIC FINANCE MANAGEMENT ACT, 1999, THE KWAZULU-NATAL SUPPLY CHAIN MANAGEMENT POLICY FRAMEWORK AND THE GENERAL CONDITIONS OF CONTRACT. REFER TO THE GENERAL CONDITIONS OF CONTRACT AT THE FOLLOWING WEB ADDRESS: http://www.treasury.gov.za/divisions/ocpo/ostb/contracts/default.aspx

- 1. Unless inconsistent with or expressly indicated otherwise by the context, the singular shall include the plural and visa versa and with words importing the masculine gender shall include the feminine and the neuter.
- 2. Under no circumstances whatsoever may the bid forms be retyped or redrafted. Photocopies of the original bid documentation may be used, but an original signature must appear on such photocopies.
- 3. The bidder is advised to check the number of pages and to satisfy himself that none are missing or duplicated.
- 4. Bids submitted must be complete in all respects.
- 5. Bids shall be lodged at the address indicated not later than the closing time specified for their receipt, and in accordance with the directives in the bid documents.
- 6. Each bid shall be addressed in accordance with the directives in the bid documents and shall be lodged in a separate sealed envelope, with the name and address of the bidder, the bid number and closing date indicated on the envelope. The envelope shall not contain documents relating to any bid other than that shown on the envelope. If this provision is not complied with, such bids may be rejected as being invalid.
- 7. All bids received in sealed envelopes with the relevant bid numbers on the envelopes are kept unopened in safe custody until the closing time of the bids. Where, however, a bid is received open, it shall be sealed. If it is received without a bid number on the envelope, it shall be opened, the bid number ascertained, the envelope sealed, and the bid number written on the envelope.
- 8. A specific box is provided for the receipt of bids, and no bid found in any other box or elsewhere subsequent to the closing date and time of bid will be considered.
- 9. No bid sent through the post will be considered if it is received after the closing date and time stipulated in the bid documentation, and proof of posting will not be accepted as proof of delivery.
- 10. No bid submitted by telefax, telegraphic or other electronic means will be considered.
- 11. Bidding documents must not be included in packages containing samples. Such bids may be rejected as being invalid.
- 12. Any alteration made by the bidder must be initialled.
- 13. Use of correcting fluid is prohibited.
- 14. Bids will be opened in public as soon as practicable after the closing time of bid.
- 15. Where practical, prices are made public at the time of opening bids.
- 16. If it is desired to make more than one offer against any individual item, such offers should be given on a photocopy of the page in question. Clear indication thereof must be stated on the schedules attached.
- 17. The bidder must initial each and every page of the bid document.

SECTION C: AUTHORITY TO SIGN A BID

A. COMPANIES

If a Bidder is a company, a certified copy of the resolution by the Board of Directors, personally signed by the Chairperson of the Board, authorising the person who signs this bid to do so, as well as to sign any contract resulting from this bid and any other documents and correspondence in connection with this bid and/or contract on behalf of the company must be submitted with this bid, that is before the closing time and date of the bid

AUTHORITY BY BOARD OF DIRECTORS

By resolution passed by the Board of Directors on	
By resolution passed by the Board of Directors on (whose signature appears below) has been duly authorised to sign all d	
(whose signature appears below) has been duly authorised to sign all d	IOCUMENTS IN CONNECTION WITH THIS DID ON DENAIT OF (Name of Company)
IN HIS/ HER CAPACITY AS:	
SIGNED ON BEHALF OF COMPANY:	(PRINT NAME)
	DATE
SIGNATURE OF SIGNATORY:	
WITNESSES: 1	DATE:
2	DATE:
۷۲	

B. SOLE PROPRIETOR (ONE - PERSON BUSINESS)

I, the undersigned	(Full name)	hereby
confirm that I am the sole owner of the business trading as:	, , , , , , , , , , , , , , , , , , ,	
·	(Name of Business)	
	, , , , , , , , , , , , , , , , , , ,	
SIGNATURE	DATE	

C. PARTNERSHIP

The following particulars in respect of every partner must be furnished and signed by every partner:

FULL NAME OF PARTNER	RESIDENTIAL ADDRESS	SIGNATURE

6

We, the undersigned Partners in the business trading as			
		(name of partnership)	
hereby authorise any contract resulting from the bid a of	and any other documents and correspondenc	(full name) to sign this bid as well as the in connection with this bid and/ or contract on behalf	
SIGNATURE	SIGNATURE	SIGNATURE	
DATE	DATE	DATE	

D. CLOSE CORPORATION

In the case of a Close Corporation submitting a bid, a certified copy of the Founding/ Amended Founding Statement of such corporation shall be included with the bid, together with the resolution by its members authorising a member or other official of the corporation to sign the documents on their behalf.

By resolution of r	nembers at a meeting on	
whose signature	appears below, has been authorised to sign all docume	
·		
		(Name of Close Corporation)
Trading as		(Trading name).
	PACITY AS:	
SIGNED ON BEI	HALF OF THE CLOSE CORPORATION:	(PRINT NAME)
SIGNATURE OF	SIGNATORY:	DATE:
WITNESSES:	1	DATE:
	2	DATE:

E. CO-OPERATIVE

A certified copy of the Constitution of the co-operative must be included with the bid, together with the resolution by its members authoring a member or other official of the co-operative to sign the bid documents on their behalf.

By resolution of members at a meeting on	. 20	
(full name) whose sign	ature	
appears below, has been authorised to sign all documents in connection with this bid on behalf of		
	perative)	
ZNB 6685/3/2023-H: SUPPLY AND DELIVERY OF RESPIRATORY: ENDOTRACHEAL TUBES AND TRACHEOSTOMY TUBES AND ACCESSORIES FOR VARIOUS INSTITUTIONS: PERIOD OF 3 YE	ARS	

SIGNATURE OF AUTHORISED REPRESENTATIVE/SIGNATORY:

IN HIS/ HER CAPACITY AS:				
DATE:				
SIGNED ON BEHALF OF CO-OPERATIVE:				
FULL NAME IN BLOCK LETTERS:				
WITNESSES:	1	DATE:		
	2	DATE:		

F. JOINT VENTURE

If a bidder is a Joint Venture, a certified copy of the resolution/ agreement passed/ reached, signed by the duly authorised representatives of the entities, authorising the representatives who sign this bid to do so, as well as to sign any contract resulting from this bid and any other documents and correspondence in connection with this bid and /or contract on behalf of the Joint Venture must be submitted with this bid, before the closing time and date of the bid.

AUTHORITY TO SIGN ON BEHALF OF THE JOINT VENTURE

By resolution/agreement passed/reached by the Joint Venture passed	artners on20
	(Full name)
	(Full name)
	(Full name)
whose signatures appear below have been duly authorised to signatures appear below have been duly authorised to signature the signature of the	
IN HIS/ HER CAPACITY AS:	
SIGNED ON BEHALF OF (ENTITY NAME):	
SIGNATURE:	DATE:
IN HIS/ HER CAPACITY AS:	
SIGNED ON BEHALF OF (ENTITY NAME):	
SIGNATURE:	DATE:
IN HIS/ HER CAPACITY AS:	

SIGNED ON BEHALF OF (ENTITY N/	АМЕ):
SIGNATURE:	DATE:
IN HIS/ HER CAPACITY AS:	
SIGNED ON BEHALF OF (ENTITY N/	AME):
SIGNATURE:	DATE:
IN HIS/ HER CAPACITY AS:	

G. CONSORTIUM

If a bidder is a Consortium, a certified copy of the resolution/ agreement passed/ reached, signed by the duly authorised representatives of concerned entities, authorising the representatives who sign this bid to do so, as well as to sign any contract resulting from this bid and any other documents and correspondence in connection with this bid and/ or contract on behalf of the Consortium must be submitted with this bid, before the closing time and date of the bid.

AUTHORITY TO SIGN ON BEHALF OF THE CONSORTIUM

By resolution/agreement passed/reached by the Consortium on.	
whose signature appears below have been duly authorised to signify with this bid on behalf of:	
	(Name of Consortium)
IN HIS/ HER CAPACITY AS:	
SIGNATURE:	DATE:

SECTION D: BIDDER'S DISCLOSURE (SBD 4)

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 STATE INSTITUTION

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

.....

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.

3 DECLARATION

I, the undersigned, (name)..... in submitting the accompanying bid, do hereby make the following statements that I certify to be true and complete in every respect:

- 3.1 I have read and I understand the contents of this disclosure;
- 3.2 I understand that the accompanying bid will be disqualified if this disclosure is found not to be true and complete in every respect;
- 3.3 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 venture or consortium will not be construed as collusive bidding.
- 3.4 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 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

SECTION E: THE NATIONAL INDUSTRIAL PARTICIPATION PROGRAMME (SBD 5)

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.
 - or
 - (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 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 tenderers (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 TENDER SUBMISSION AND CONTRACT REPORTING REQUIREMENTS OF TENDERERS AND SUCCESSFUL TENDERERS (CONTRACTORS)

Tenderers are required to sign and submit this Standard Tenderding Document (SBD 5) together with the Tender 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 Tenderers (contractors) are required, immediately after being officially notified about any successful Tender with a value in excess of R10 million (ten million Rands), to contact and furnish the DTI with the following information:
 - Tender / 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 X 84, Pretoria, 0001 for the attention of Mr. Elias Malapane within five (5) working days after award of the contract. Mr. Malapane may be contacted on telephone (012) 394 1401, facsimile (012) 394 2401 or e-mail at Elias@thedti.gov.za for further details about the programme.

4 PROCESS TO SATISFY THE NIP OBLIGATION

- 4.1 Once the successful Tenderer (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 Tenderer (contractor) and, therefore, does not involve the purchasing institution.

Tender number:	ZNB6685/3/2023-H
Name of tenderer:	Closing date: 28 SEPTEMBER 2023
Postal address:	
Signature:	Name (in print):
Date:	

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SECTION F: DECLARATION THAT INFORMATION ON CENTRAL SUPPLIER DATABASE IS CORRECT AND UP TO DATE (To be completed by bidder)

This is to certify that I
(name of bidder/authorized representative)
who represents
(state name of bidder)
am aware of the contents of the Central Supplier Database with respect to the bidder's details and registration information, and that the said information is correct and up to date as on the date of submitting this bid, and I am aware that incorrect or outdated information may be a cause for disqualification of this bid from the bidding process, and/ or possible cancellation of the contract that may be awarded on the basis of this bid.

SIGNATURE OF BIDDER OR AUTHORISED REPRESENTATIVE

DATE:

SECTION G: PREFERENCE POINTS CLAIM FORM IN TERMS OF THE PREFERENTIAL PROCUREMENT REGULATIONS 2022: SBD 6.1

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 80/20 system for requirements with a Rand value of up to R50 000 000 (all applicable taxes included); and
 - the 90/10 system for requirements with a Rand value above R50 000 000 (all applicable taxes included).

1.2 To be completed by the organ of state

- a. The applicable preference point system for this tender is the 80/20 preference point system.
- b. The applicable preference point system for this tender is the 90/10 preference point system.
- c. E 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.

1.3 **To be completed by the organ of state:**

The maximum points for this tender are allocated as follows:

	POINTS	POINTS
PRICE	80	90
SPECIFIC GOALS	20	10
Total points for Price and SPECIFIC GOALS	100	100

- 1.4 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.
- 1.5 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.

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 80/20 OR 90/10 PREFERENCE POINT SYSTEMS

A maximum of 80 or 90 points is allocated for price on the following basis:

80/20 or 90/10 $Ps = 80\left(1 - \frac{Pt - P\min}{P\min}\right)$ or $Ps = 90\left(1 - \frac{Pt - P\min}{P\min}\right)$ 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 80 or 90 points is allocated for price on the following basis:

$$Ps = 80\left(1 + \frac{Pt - Pmax}{Pmax}\right)$$
 or $Ps = 90\left(1 + \frac{Pt - Pmax}{Pmax}\right)$

Where

Ps	=	Points scored for price of tender under consideration
Pt	=	Price of tender under consideration
Pmax	=	Price of highest acceptable tender

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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:
- 4.2. In cases where organs of state intend to use Regulation 3(2) of the Regulations, which states that, if it is unclear whether the 80/20 or 90/10 preference point system applies, an organ of state must, in the tender documents, stipulate in the case of—
 - (a) an invitation for tender for income-generating contracts, that either the 80/20 or 90/10 preference point system will apply and that the highest acceptable tender will be used to determine the applicable preference point system; or

(b) any other invitation for tender, that either the 80/20 or 90/10 preference point system will apply and that the lowest acceptable tender will be used to determine the applicable preference point system, then the organ of state must indicate the points allocated for specific goals for both the 90/10 and 80/20 preference point system.

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

(Note to organs of state: Where either the 90/10 or 80/20 preference point system is applicable, corresponding points must also be indicated as such.

The specific goals points in terms of tender		Number of points allocated (90/10 system) (To be completed by the organ of state)	Number of points allocated (80/20 system) (To be completed by the organ of state)	Number of points claimed (90/10 system) (To be completed by the tenderer)	Number of points claimed (80/20 system) (To be completed by the tenderer)
In terms of Departm Preferential Procure Regulation Policy 20 section 8.1.2.1. for H Disadvantaged Indiv The Department allo 20 or 10 points to co who are at least 51% by Black People	ment 023, Historically /iduals. ocate full ompanies	10 Points	20 Points		
Note: CSD will be us verify ownership	sed to				

Note to tenderers: The tenderer must indicate how they claim points for each preference point system.)

DECLARATION WITH REGARD TO COMPANY/FIRM

- 4.3. Name of company/firm.....
- 4.4. Company registration number:
- 4.5. TYPE OF COMPANY/ FIRM
 - Partnership/Joint Venture / Consortium
 - One-person business/sole propriety
 - Close corporation
 - Public Company
 - Personal Liability Company
 - □ (Pty) Limited
 - Non-Profit Company
 - State Owned Company

[TICK APPLICABLE BOX]

- 4.6. I, the undersigned, who is duly authorised to do so on behalf of the company/firm, certify that the points claimed, based on the specific goals as advised in the tender, qualifies the company/ firm for the preference(s) shown and I acknowledge that:
 - i) The information furnished is true and correct;
 - ii) The preference points claimed are in accordance with the General Conditions as indicated in paragraph 1 of this form;
 - iii) In the event of a contract being awarded as a result of points claimed as shown in paragraphs 1.4 and 4.2, the contractor may be required to furnish documentary proof to the satisfaction of the organ of state that the claims are correct;
 - iv) If the specific goals have been claimed or obtained on a fraudulent basis or any of the conditions of contract have not been fulfilled, the organ of state may, in addition 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 *audi alteram partem* (hear the other side) rule has been applied; and
 - (e) forward the matter for criminal prosecution, if deemed necessary.

	SIGNATURE(S) OF TENDERER(S)
SURNAME AND NAME:	
DATE:	
ADDRESS:	

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SECTION H: RECORD OF AMENDMENTS TO BID DOCUMENTS

I / We confirm that the following communications amending the bid documents that I / we received from KwaZulu-Natal Department of Health or their representative before the closing date for submission of bids have been taken into account in this bid.

ADDENDUM NO.	DATE	TITLE OR DETAILS

SIGNATURE: DATE:

(of person authorized to sign on behalf of the Bidder)

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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34. Prohibition of restrictive practices

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 stockactually 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 goodson 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 andnot 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 bemanufactured.
- 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.
- 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 biddingdocuments.

	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 beobtained directly from the Government Printer, Private Bag X85, Pretoria 0001, or accessed electronically from <u>www.treasury.gov.za</u>
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 (allcopies) 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-partyclaims 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 purchaserand 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 andreturned to the supplier not later than thirty (30) days following thedate of completion of the supplier's performance obligations under the contract, including any warranty obligations, unless otherwise specified in SCC.
^{8.} Inspections ,tests and	8.1	All pre-bidding testing will be for the account of the bidder.
analyses	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 thecost and risk of the supplier who shall, when called upon, remove themimmediately 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 thesuppliers 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 otherdocuments 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 maintenanceof the supplied goods; (c) furnishing of a detailed operations and maintenance manualfor 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 theparties 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 allof 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 supplierof 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 theyincorporate 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 fromdesign, materials, or workmanship (except when the design and/ormaterial 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 finaldestination.
 - 15.2 This warranty shall remain valid for twelve (12) months after thegoods, 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 supplierunder		
	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 obligationsstipulated in the contract.		
	16.3 Payments shall be made promptly by the purchaser, but in no case laterthan 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 bemade 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 thesupplier's performance	21.1 Delivery of the goods and performance of services shall be made bythe 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 likelyduration 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 shallbe 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 foror 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 shallcontinue 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) workingdays 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 databaseof 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.
- 24.1 When, after the date of bid, provisional payments are required, or anti-dumping 24. Anti-dumping and or countervailing duties are imposed, or the amount of a provisional payment or countervailing anti-dumping or countervailing right isincreased in respect of any dumped or duties and rights 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

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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 inperformance 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 notifythe purchaser in writing of such condition and the cause thereof. Unless otherwise directed by the purchaser in writing, the suppliershall continue to perform its obligations under the contract as far as is reasonably practical, and shall seek all reasonable alternative means forperformance not prevented by the force majeure event.
26. Termination for insolvency	26.1	The purchaser may at any time terminate the contract by giving writtennotice 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 ofDisputes	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 inthe 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

ZNB 6685/3/2023-H: SUPPLY AND DELIVERY OF RESPIRATORY: ENDOTRACHEAL TUBES AND TRACHEOSTOMY TUBES AND ACCESSORIES FOR VARIOUS INSTITUTIONS: PERIOD OF 3 YEARS

	(b)	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	doo	e contract shall be written in English. All correspondence and other cuments pertaining to the contract that is exchanged by the parties shall o be written in English.	
30. Applicable law		e contract shall be interpreted in accordance with South Africanlaws, ess 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 orto the address notified later by him in writing and such posting shall bedeemed to be proper service of such notice		
	afo	time mentioned in the contract documents for performing any act after such resaid notice has been given, shall be reckoned from the date of posting of ch notice.	
32. Taxes andduties		foreign supplier shall be entirely responsible for all taxes, stamp duties, ense fees, and other such levies imposed outside the purchaser's country.	
		local supplier shall be entirely responsible for all taxes, duties, license fees, tc., incurred until delivery of the contracted goods tothe purchaser.	
	o ta	lo contract shall be concluded with any bidder whose tax matters arenot in rder. Prior to the award of a bid the Department must be in possession of a ax clearance certificate, submitted by the bidder. This certificate must be an riginal issued by the South AfricanRevenue Services.	
 33. National Industrial Participation (NIP) Programme 34 Prohibition of Restrictive practices 		The NIP Programme administered by the Department of Trade and Industry shall be applicable to all contracts that are subject to the NIP obligation.	
	; (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 norizontal relationship and if a bidder (s) is / areor a contractor(s) was / were nvolved in collusive bidding (or bid rigging).	
	(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 benalties 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.
- I have read, understand and accept the above stated General Conditions of the Contract which are binding upon me.

Signature

Date

Name of Bidder

SECTION J: SPECIAL CONDITIONS OF CONTRACT (SCC)

<u>Note:</u> The special conditions of contract referred as (SCC) are supplementary to that of the General Conditions of Contract (GCC). Where, however, the special conditions of contract are in conflict with the General Conditions of Contract, the special conditions of contract (SCC) shall prevail.

1. ADDITIONAL DEFINITIONS

In addition to the definitions contained in paragraph 1 of the GCC, the following terms shall be interpreted as indicated:

"Accounting Officer": means a person described in Section 36 of the Public Finance Management Act, Act No. 1 of 1999 (As amended by Act 29 of 1999).

"Contract Duration": means the period between the commencement and termination of the contract.

"**Confidential Information**": means but is not limited to contents of the contract, or any provision thereof, or any specification, plan, know-how, drawing, pattern, sample, or information furnished by or on behalf of the Department in connection therewith, to any person other than a person employed by contractor or service provider in the performance of the contract.

"Department": means the KwaZulu-Natal Department of Health.

"Head of Department": means the Head of Department for KwaZulu-Natal Department of Health as defined in Schedule 2 Column 1 and 2 of the Public Service Act 1994 (Proclamation 103 of 3 June 1994, as amended).

"Health Facilities": means Head Office, District Offices, Hospitals, Community Health Centres, Specialized centres and Clinics under the auspices of the Department of Health in the Province.

"ISO Standards": means standards recognized by International Standard Organisation

"Parties": means the KwaZulu-Natal Department of Health and Contractor or Service provider

"Province": means the Province of KwaZulu-Natal.

"ROE": means the Rate of Exchange.

"SABS": means the South African Bureau of Standards.

"SANS": means the South African National Standards.

"Vendor": means Contracted Supplier or Service Provider

2. INTERPRETATIONS

In amplification of the provisions of paragraph 2 of the GCC, unless inconsistent with the context, an expression which denotes:

- 2.1 Any gender includes the other genders.
- 2.2 A natural person includes a juristic person and vice versa.
- 2.3 The singular includes the plural and vice versa.
- 2.4 When any number of days is prescribed in this Contract, the same shall be reckoned exclusively of the first and inclusively of the last day unless the last day falls on a Saturday, Sunday or proclaimed public holiday in the Republic of South Africa, in which event the last day shall be the next succeeding day which is not a Saturday, Sunday or public holiday.
- 2.5 Figures are referred to in numerals and in words, if there is any conflict between the two, the words shall prevail.
- 2.6 Any reference in this contract to "goods" includes works and/or services.
- 2.7 The written and signed contract represents the final agreement between the parties and it super cedes any prior oral agreements or discussions of the Contract.
- 2.8 All annexures and appendices shall form part of the contract.
- 2.9 The headings used throughout the Contract do not have any special significance save to ensure the easy reading of the contract.
- 2.10 Words and phrases defined in this Contract shall bear the meaning assigned to them throughout this Contract.
- 2.11 Words and phrases used in this Contract which are defined or used in any statute or regulation which applies to the subject matter, professional person.
- 2.12 The bid is issued in accordance with Section 217 of the Constitution, The Public Finance Management Act, Treasury Regulations 16A and National Treasury regulations and guidelines.

3. ACCEPTANCE OF A BID

- 3.1 The Department of Health Bid Adjudication Committee is under no obligation to accept any bid.
- 3.2 The financial standing of a bidder and its ability to supply goods or render services may be examined before the bid is considered for acceptance.

4. CERTIFICATE OF COMPLIANCE

- 4.1 If the bidder submits offers for items that make reference to South African National Standards (SANS) or South African Bureau of Standards (SABS) or International Organisation for Standardisation (ISO) specifications, a Certificate of Compliance must be submitted with the bid document at the time of closing of the bid. SABS/SANS can be contacted for testing and conformity services at Tel: 031 203 2900/ Fax: 031 203 2907. SANS, SABS AND CKS specifications will be for the account of the prospective bidder. Failure to submit the certificate, where applicable, will result in the bid being disqualified. The Department reserves its rights to contact SABS/SANS/CKS for testing and conformity services.
- 4.2 The South African National Accreditation System (SANAS) is recognized by the South African Government as the single National Accreditation Body that gives formal recognition that Laboratory, Certification Bodies, Inspection Bodies, Proficiency Testing Scheme Providers and Good Laboratory Practice (GLP) test facilities are competent to carry out specific tasks. This organization can be contacted as follows: Tel: 012 3943760: Fax: 012 3940526.
- 4.3 Prior to an award of the bid being made and/or during the evaluation process, the Department of Health reserves the right to conduct inspections of the premises of the most acceptable bidder. Therefore, premises of the bidder shall be open, at reasonable hours, for inspection by a representative of the Department or organization acting on its behalf. Any specification/s and conformity testing will be for the account of the prospective bidder.
- 4.4 Must comply with ISO 5361:2016, ISO 5366 or equivalent standard.

5. COMPLIANCE WITH SPECIFICATION

- 5.1 Offers must comply strictly with the specification. Offers exceeding specification requirements will be deemed to comply with the specification.
- 5.2 The quality of services/ supply must not be less than what is specified.

6. PERFORMANCE STANDARDS

- 6.1 In amplification of paragraph 4 of the GCC, the preferred bidder shall supply the goods in accordance with performance standards set by the Department below:
- 6.1.2 The items to be supplied must meet all the general clauses and technical clauses as per specification issued for the tender. Failure to comply will lead to the award being cancelled.
- 6.1.3 Items to come with an expiry date.
- 6.1.4 Failure to comply with minimum specification or incorrect response will mean the contract be cancelled with no risk of financial loss to the department.

7 QUALITY CONTROL /TESTING OF PRODUCTS AND GUARANTEE

7.1 The Department and/or Institution reserves the right to have any product tested with an accredited agent in the Republic of South Africa. The quality control testing administrative procedures will be undertaken by the Department's Supply Chain Management Contract Management section.

7.2 If it is discovered that the product supplied is not in accordance with the specification the following will occur:

Testing charges will be for the account of Contractor.;

Possible cancellation of the contract with Contractor.;

Reporting such negligence to the Provincial and National Treasury for listing on the Restricted Suppliers Database.

- 7.3 All goods supplied shall be equal in all respects to samples, patterns or specifications where such are provided. Any changes to quality or brands will have to be approved by the Department, as this is a change to the conditions of the contract.
- 7.4 Should the Department, after the award of the Contract and/or during the manufacture of the goods specified, decide on a variation or alteration to the specification, either at the suggestion of Contractor or otherwise, which will be to the Department's advantage, such variation or alteration shall be performed to the Department's satisfaction. Any variation in the Contract Price arising there from shall be subject to agreement between the Department and Contractor. The variation shall comply with thresholds as prescribed by National Treasury regulations.
- 7.5 Contractor shall not be relieved of its obligations with respect to the sufficiency of the materials and workmanship and the quality of the goods supplied by the reason of no objection having been taken thereto by the Department's Representative at the time the goods were delivered.
- 7.6 Contractor 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. Contractor, 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 Department's specifications) or from any act or omission of Contractor., that may develop under normal use of the supplied goods in the conditions prevailing in the country of the final destination.
- 7.7 This warranty shall remain valid for (24) 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.
- 7.8 The Department shall promptly notify Contractor in writing of any claims arising under this warranty. Contractor shall immediately remedy the said defect free of cost to the Department. Should Contractor delay remedial work in excess of time stipulated by the Department's representative, the Department may have such remedial work executed at Contractor expense. Should the Department decide that the defect is such that it cannot be remedied, the goods may be rejected, such rejected goods shall be held at the risk and expense of Contractor and shall, on request of the Department, be removed by Contractor immediately on receipt of notification of rejection. Contractor shall be responsible for any loss the Department may sustain by reason of such action as the Department may take, in terms of this clause.
- 7.9 The risk in respect of the goods purchased by the Department under the contract shall remain with Contractor, until such goods have been delivered to the Department.

- 7.10 The principle feature of the goods is described in the Specification, but the Specification does not purport to indicate every detail of supply, of Goods necessary to meet the requirements. Omission from the Specification of reference to any part or parts shall not relieve Contractor of their responsibility for carrying out the supply of goods as required under the Contract.
- 7.11 If any dispute arises between the Department and Contractor, in connection with the quality and guarantee of the goods, either party may give the other notice in writing of the existence of such dispute, and the same shall thereupon be referred to arbitration in South Africa by a person mutually agreed upon by both parties. The submission shall be deemed to be a submission to arbitration within the meaning of the terms of the arbitration laws in force in the Republic of South Africa.

8. EQUAL BIDS

- 8.1 During the submission of price quotations, the equal bids and criteria for breaking deadlock in scoring will be as follows:
- 8.1.1 If two or more tenderers score an equal total number of points, the contract must be awarded to the tenderer that scored the highest points for specific goals.
- 8.1.2 If two or more tenderers score equal total points in all respects, the award must be decided by the drawing of lots.

9. LATE BIDS

- 9.1 Bids are permissible to be submitted prior to closing date and time this is to avoid unfortunate or unplanned circumstances that could prevent the bidder from arriving on time during the closing date. If the bidder fail to arrive on time the department will not be held liable.
- 9.2 Bids are late if they are received at the address indicated in the bid documents after the closing date and time.

10. MORE THAN ONE OFFER/ COUNTER OFFERS

- 10.1 Should the bidder make more than one offer, where applicable, against any individual item, such offer/s must be detailed in the Schedule of Additional Offer/s. The Department reserves its rights in and to the consideration of any additional offer/s subject to compliance with specification and the bidding conditions.
- 10.2 Bidders' attention is drawn to the fact that counter offers with regard to any of the abovementioned Special Terms and Conditions will invalidate such bids.
- 10.3 Bidders are at liberty to bid for one, a number of items, or bid for all items. If a bidder is not bidding for all the items, the appropriate price page must reflect: 'nil quote'.

11. ONLY ONE OFFER RECEIVED

11.1 Where only 1 offer is received, the Department of Health will determine whether the price is fair and reasonable. Proof of reasonableness will be determined as follows:

Comparison with prices, after discounts, to the bidder's other normal clients and the relative discount that the State enjoys;

Where this is not possible, profit before tax based on a full statement of relevant costs; and

In all cases, comparison with previous bid prices where these are available.

12. AWARD OF BID (S)

12.1 The State reserves the right to award the same item to more than one (1) bidder to address item availability and compatibility. Due diligence will be applied to ensure that pricing is affordable, market related and aligned to end-user requirements.

The following shall be taken into consideration when contemplating a multiple award:

- 12.1.1 Capacity to meet the expected demand according to the end-user requirements;
- 12.1.2 Mitigation of risk if the item is unavailable; and
- 12.1.3 The maximum number of suppliers per item to be awarded will be at the discretion of the BEC.

12.2. Right of Award

The State reserves its following rights -

- 12.2.1.1To award the bid in part or in full;
- 12.2.2. Not to make any award in this bid or accept any bids submitted;
- 12.2.3. Award the bid to more than one (1) bidder for the same item (multiple-award);
- 12.2.4. Request further technical information from any bidder after the closing date;
- 12.2.5. Verify information and documentation of the bidder(s);
- 12.2.6. Not to accept any of the bids submitted;
- 12.2.7. To withdraw or amend any of the bid conditions by notice in writing to all bidders prior to closing of the

bid and post award; and

12.2.8 In the event that an incorrect award has been made to remedy the matter in any lawful manner it may

deem fit.

12.3 Where specifications are annotated as "to be considered in series" these items will be awarded to a single supplier who is compliant to specification and who has scored the highest points.

13. REGISTRATION ON THE CENTRAL SUPPLIER DATABASE (CSD)

- 13.1 A bidder submitting an offer must be registered on the Central Supplier Database. A bidder who has submitted an offer and is not registered on the Central Supplier Database will not be considered.
- 13.2 Each party to a joint venture/ consortium must be registered on the Central Suppliers Database at the time of submitting the bid.

14. EMPLOYEES TRADING WITH THE ORGANS OF THE STATE

- 14.1 The Public Service Act 103 of 1994 indicates in section 30(1) that "No employee shall perform or engage himself or herself to perform remunerative work outside his or her employment in the relevant department, except with the written permission of the executive authority of the department."
- 14.2 Furthermore, in terms of the Public Service Regulations paragraph 13(c), "An employee shall not conduct business with any organ of state or be a director of a public or private company conducting business with an organ of state, unless such employee is in an official capacity a director of a company listed in schedule 2 and 3 of the Public Finance Management Act"

14.3 If a bidder is found to be employed by the state, through the verification from Central Supplier Database (CSD) Registration Report or Department of Public Service and Administration (DPSA) verification system, the bid will be immediately disqualified. If it is discovered that the winning or contracted bidder is employed by the state through other Computer Assisted Audit Technics (CAATS), the award or contract may be immediately terminated.

15 TRUST, CONSORTIUM OR JOINT VENTURE

- 15.1 To ensure compliance with SCM prescripts, a Trust, Consortium or Joint Venture must submit a consolidated Specific Goals for every separate bid. No award will be made to a Trust/ Joint Venture/ Consortium that is not tax compliant at the finalisation of the award. For verification purposes, each party must submit separate proof of TCS/ PIN / CSD number.
- 15.2 A separate B-BBEE Certificate or Sworn Affidavit will be required from each company participating in the awarded Trust, Consortium or Joint Venture during the formal contract stage.
- 15.3 The Joint Venture agreement must clearly specify the percentage of the contract to be undertaken by each company participating therein.
- 15.4 The Joint Venture/Consortium must submit a formal agreement that outlines the roles and responsibilities of each member of the Joint Venture/ Consortium, nomination of an authorised person to represent the Joint Venture or Consortium in all matters relating to this bid and the details of the bank account for payments to be affected.

16. VALIDITY PERIOD OF BID AND EXTENSION THEREOF

16.1 The validity (binding) period for the bid will be <u>180 days</u> from close of bid. However, circumstances may arise whereby the department may request bidders to extend the validity (binding) period. Should this occur, the department will request bidders to extend the validity (binding) period under the same terms and conditions as originally offered for by bidders? This request will be done before the expiry of the original validity (binding) period.

17. CHANGE OF ADDRESS

17.1 Bidders must advise the Department of Health's Central Supply Chain Management Unit, Contract Section, should their ownership and/or address (domicilium citandi et executandi) details change from the time of bidding to the expiry of the contract.

18. DELIVERY, MARKING AND PACKAGING

- 18.1 Basis of delivery of products must be made in accordance with the instruction appearing on the official Order form. The instructions appearing on the official order form regarding the supply, dispatch and submission of invoices must be strictly adhered to.
- 18.2 All deliveries or dispatches must be accompanied by a delivery note stating the official order number against the delivery that has been affected.
- 18.3 In respect of goods and services awarded, the Contractors must adhere strictly to the delivery periods stipulated in the bid document or as agreed with the Department. In case of delays in the supplier's performance, the supplier must inform the department or institution of such delays and comply with conditions as stipulated on the GCC. Should the Contractor fail to supply the goods within the time stated in its bid, or within the extended time allowed to them, the department reserves the right, to cancel the contract and purchase the goods elsewhere and the Contractor shall refund to the department any extra cost incurred over and above the contract price.
- 18.4 All deliveries must take place from Monday to Friday between 08h00 and 14h00. In emergency cases, the department reserves the right to request the successful bidder/s to urgently effect deliveries at any given time including Saturdays, Sundays and public holidays.

- 18.5 Order details must be presented upon delivery on delivery notes. Deliveries not complying with the order form, specifications or samples submitted, will be returned to the Contractor at the Contractor's expense. Goods delivered shall in all cases be accompanied by delivery notes in duplicate, one which will be retained by the Department. The Contractor shall be responsible for the safe delivery as to the quality, quantity and condition of the goods.
- 18.6 All goods shall be crated, packed or battened securely in such a manner as to prevent damage during loading, transport and off-loading. Unless otherwise specified, packing cases and packing materials are included in the Contract Price, and shall be and remain the property of the Department. It is the Contractor's responsibility to off load the delivery vehicle. Delivery packages should be of a durable quality that will allow stacking and for further transportation without breakage.
- 18.7 The following information must appear on the outer packaging of the carton/box:
 - (a) Name of the manufacturer/supplier
 - (b) Description of item
 - (c) Date of manufacture
- 18.8 Where applicable each item in a carton must be individually labelled and the following information must appear on the outer packaging of the carton:

Name of the manufacturer/supplier;

Description of item;

Item number code/catalogue number;

Date of manufacture;

Product expiry date;

Batch No.;

Lot No.

- 18.9 Random inspection and sampling of items will be conducted upon delivery to verify quantity and compare the item against the contract sample and any other quality accreditation or health standards that is prescribed.
- 18.10 No locally manufactured product may be substituted during the contract period with an imported product, and vice versa, without prior approval of Contract Management at Central Supply Chain Management, Department of Health.

19 PERIOD OF CONTRACT

19.1 The period of this contract is Three (03) years.

20 INVOICES AND PAYMENTS

- 20.1 All invoices must be submitted in the original format.
- 20.2 All invoices submitted by the Contractor must contain the word "INVOICE" for non-VAT vendors or "TAX INVOICE" for VAT vendors only. VAT number must be reflected for VAT vendors.
- 20.3 A tax invoice shall be in the currency of the republic of South Africa and shall contain the following particulars:

- (a) The name, address and registration number of the supplier;
- (b) The name and address of the recipient;
- (c) An individual serialized number and the date upon which the tax invoice is issued;
- (d) A description of the goods or services supplied;
- (e) The quantity or volume of the goods or services supplied
- (f) The value of the supply, the amount of tax charged and the consideration for the supply; or
- (g) Where the amount of tax charged is calculated by applying the tax fraction to the consideration, the consideration for the supply and either the amount of the tax charged, or a statement that it includes a charge in respect of the tax and the rate at which the tax was charged.
- 20.4 A Contractor shall be paid by the institution concerned, in accordance with supplies delivered and services rendered. The goods must be accepted and signed off by the relevant delegated official.
- 20.5 Should a Contractor indicate a special discount on his/her account provided payment is made within a certain time, every effort shall be made to take advantage of such discount. Where discounts or rebates received by the Department, the Contractor to provide credit note.
- 20.6 Any query concerning the non-payment of accounts must be directed to the institution concerned. The following protocol will apply if accounts are queried:
 - (i) Contact must be made with the officer-in-charge of Logistics and Accounts Payable;
 - (ii) If there is no response from Logistics and Accounts Payable, the Finance Manager and the

Chief Executive Officer of the institution must be contacted.

(iii) Failing all of the above, the Contractor must contact the Chief Director: Accounting Services supplying the following details:

Name/s of person/s contacted at the Institution and dates; and Details of outstanding account. the Chief Director: Accounting Services will then take the appropriate action.

20.7 The Institutions shall not be responsible for payment of any statutory increases in tariffs or imports or any fluctuations in foreign exchange rate for any item required Contractor, to realise its obligations in terms of this Contract. The rate of exchange, as agreed upon in this Contract is subject to review if stipulated within this contract and as agreed consented by both Parties.

21. STATEMENT OF SUPPLIES AND SERVICES

- 21.1 The Contractor shall, monthly, furnish particulars of supplies delivered or services executed. Such information must be submitted to the Department of Health Supply Chain Management, Contract Management as follows:
- (i) Name of institution.
- (ii) Orders received per each institution, order number, catalogue number, quantity delivered and invoice amount all inclusive.
- 21.2 Historical value and volume reports may be requested by the Department of Health, Supply Chain Management, during the term of the contract for the following:

SUPPLIER MEASURES

Delivery period adherence

Quality adherence

Note: This information will be submitted at the expense of the Contractor.

22. FIRM PRICES AND ESCALATIONS

- 22.1 This bid requires that all bid prices offered are firm for the period of the contract. If a non-firm price is offered, the bidder shall be disqualified for not complying with the conditions of the bid.
- 22.2 It is the responsibility of the bidder to take necessary precautions or to cater or include cover for unfavourable rate of exchange. Therefore, a price adjustment in respect of a rate of exchange claim will not be considered.

23. VALUE ADDED TAX (VAT)

- 23.1 All bid prices must be inclusive of all applicable taxes.
- 23.2 Bidders who make taxable supplies in excess of R1 million in any 12-month consecutive period are

liable for compulsory VAT registration, but an entity may also choose to register voluntarily provided

that the minimum threshold of R50 000 (as of 1 March 2010) has been exceeded in the past 12 month period. Bidders who meet the above requirement must register as VAT vendors, if successful, within one month of award of bid.

- 23.3 VAT will not be included after an award of the bid or during contract management period.
- 23.4 It is compulsory for bidders to be tax complaint, in the event that they are awarded the bid, Bidders must me tax compliant at time of award, upon placing of orders and during the contract period. Failure to be tax compliant or tax affairs not being in order will result in the disgualification of the bidder or cancellation of the contract or order

.24. ENTERING OF HOSPITAL/CLINIC STORES

24.1 No representative from a company shall be permitted to enter the hospital/clinic premises, buildings or containers where stores are kept unless he/she is accompanied by the responsible official in charge of stores. Before entering the hospital/clinic premises, buildings or containers where stores are kept, the company representative must in writing, motivate why entry is necessary and written authority must be obtained to enter from the Head of the Institution or delegated official.

25. DEPARTMENTAL PROPERTY IN POSSESSION OF A CONTRACTOR

- 25.1 The Department's property supplied to a Contractor for the execution of a contract remains the property of the Department and shall at all times be available for inspection by the Department or its representatives. Any such property in the possession of the Contractor on the completion of the contract shall, at the Contractor's expense, be returned to the Department forthwith.
- 25.2 The Contractor shall be responsible at all times for any loss or damages to the Department's property in his possession and, if required, he shall furnish such security for the payment of any such loss or damages as the Department may require.

26 IRREGULARITIES

26.1 Companies are encouraged to advise the Department of Health timeously of any possible irregularities

which might come to their notice in connection with this or other contracts.

27 UNSATISFACTORY PERFORMANCE

- **27.1** In amplification of, unsatisfactory performance occurs when performance is not in accordance with the contract conditions.
- 27.2. The institution shall warn the Contractor by registered/certified mail or email that action will be taken in accordance with the contract conditions unless the Contractor complies with the contract conditions and delivers satisfactory supplies or services within a specified reasonable time (7 days minimum). If the Contractor does not perform satisfactorily despite the warning the institution will:

Take necessary and appropriate action such as termination of contract in terms of its delegated powers.

- 27.3 When correspondence is addressed to the Contractor, reference will be made to the contract number/item number/s and an explanation of the complaint.
- 27.3.1 Treasury Regulation 16A9.2 specifies that "The accounting officer or accounting authority -

(a) may disregard the bid of any bidder if that bidder, or any of its directors -

- (i) have abused the institution's supply chain management system
- (ii) have committed fraud or any other improper conduct in relation to such system; or

(iii) have failed to perform on any previous contract

27.4. In the event that the awarded bidder fails to perform as per the contract conditions, the bidder shall be registered on the Departmental defaulters register and may be deemed failing to perform as per contract and therefore future bids disregarded.

28 RESTRICTION OF BIDDING

The Accounting Officer or his/her delegate must:

- a) Notify the supplier and any other person of the intention to restrict it doing business with Department by registered mail or email. The letter of restriction must provide for:
 - 1. The grounds for restriction;
 - II. The period of restriction which must not exceed 10 years;
 - A period of 14 calendar days for the supplier to provide reasons why the restriction should not be imposed.
- b) The Accounting Officer his/her delegate:
 - May regard the intended penalty as not objected to and may impose such penalty on the supplier, should the supplier fail to respond within the 14 days; and
 - Must assess the reasons provided by the supplier and take the final decision.
- c) If the penalty is imposed, the Accounting Officer must inform National Treasury of the restriction within 7 calendar days and must furnish the following information:
 - The name and address of the entity/ person to be restricted;
 - IL The identity number of individuals and the registration number of the entity; and
 - III. The period of restriction.
- d) National Treasury will load the details on the Database of Prohibited Vendors.
- e) The restriction period applicable will be based on the value of award/s made to the supplier over a financial year. The table below illustrates the restriction period that will be applicable per the award threshold:

29 CONTRACTOR'S LIABILITY

- 29.1 In the event of the contract being cancelled by the Department in the exercise of its rights in terms of these conditions, the Contractor shall be liable to pay to the Department any losses sustained and/or additional costs or expenditure incurred as a result of such cancellation, and the Department shall have the right to recover such losses, damages or additional costs by means of set-off from moneys due or which may become due in terms of the contract or any other contract or from guarantee provided for the due fulfilment of the contract and, until such time as the amount of such losses, damages or additional costs have been determined, to retain such moneys or guarantee or any deposit as security for any loss which the Department may suffer or may have suffered.
- 29.2 The Contractor may be held responsible for any consequential damages and loss sustained which may be caused by any defect, latent or otherwise, in supply or service rendered or if the goods or service as a result of such defect, latent or otherwise, does not conform to any condition or requirement of the contract.

30 RIGHTS TO PROCURE OUTSIDE THE CONTRACT

- 30.1 The Department reserves the right to procure goods outside the contract in cases of urgency or emergency or if the quantities are too small to justify delivery costs, or if the goods are obtainable from another organ of State or if the Contractor's point of supply is not situated at or near the place where the goods are required or if the Contractor's goods are not readily available.
- 30.2 No provision in a contract shall be deemed to prohibit the obtaining of goods or services from a Department or local authority.
- 30.3 If contracted item/s become available from National Treasury transversal contract, the Department reserve a right to cancel the contract with a winning bidder by giving thirty (30) days' notice. If it in the advantage and interest of the department to participate.

31. PATENTS

31.1 The Contractor shall pay all royalties and expenses and be liable for all claims in respect of the use of patent rights, trademarks or other protected rights, and hereby indemnifies the Department against any claims arising there from.

32 WAIVER

- 32.1 The granting by any party of any indulgence or postponement shall not be a waiver of its rights arising from this contract to demand full and specific performance of the contract.
- 32.2 No favour, delay or relaxation or indulgence on the part of any party in exercising any power or right conferred on each party in terms of this contract shall operate as a waiver of such power or right nor preclude any other or further exercises thereof or the exercise of any other power or right under this contract.

33 SUSPENSION

- 33.1 The Department may temporarily suspend whole or part of the supplied goods by providing no less than 5 days written notice to the Contractor, who shall on receipt of such written notice immediately cease the supply the goods. The Department will indicate the date on which the contract will be resumed in the aforementioned notice. No suspension shall exceed a total of 90 days unless otherwise agreed to by the parties in writing.
- 33.2 When the supply of the goods is suspended, the Contractor shall be entitled to pro-rata payment for the goods already delivered and reimbursement of all costs incidental to the prompt and orderly suspension of the contract.
- 33.3 Suspension of the contract shall not prejudice or affect the accrued rights and liabilities of the parties as at the date of suspension.

34 BREACH

- 34.1 Any termination notice referred to in GCC paragraph 23.1 shall be preceded by written notice requiring the defaulting party to remedy a breach of this contract within 14 days of the date of receipt of the notice.
- 34.2 If the defaulting party fails to remedy the breach within the 14 days, the aggrieved party shall be entitled without notice, in addition to any other remedy available to them at law or under this contract:
- 34.3 To claim specific performance of any obligation whether or not the due date for performance has arrived; or
- 34.4 To terminate this contract in accordance with paragraph 23.1 of the GCC, against the defaulting party, in either event without prejudice to the aggrieved party's rights to claim damages.
- 34.5 The Contractor shall immediately advise the Department of the same, upon which the Department shall, in its sole and absolute discretion, decide whether to proceed with this contract or to terminate forthwith. Failure by the Contractor to advise the Department of a conflict of interest shall amount to a material breach of this contract.
- 34.6 A Party shall be deemed to be in breach of this Contract should the Party fail to comply with any material provisions of this Contract.
- 34.7 The aggrieved Party shall be obliged to first attempt to settle the matter by way of consultation with the defaulting Party. If the consultation fails, then the aggrieved Party shall promptly give the defaulting Party fourteen (14) days written notice to remedy the breach. If the defaulting Party fails to comply with such notice, the aggrieved Party may, without prejudice to any other's right at law:
- 34.7.1 Cancel this Contract in the event the defaulting Party committed a material breach.
- 34.7.2 Claim specific performance by the defaulting Party if such is a competent remedy in the circumstance.
- 34.7.3 Claim damages suffered, as limited under this Contract.

35. PREFERENCES

- 35.1 Should the Contractor apply for preferences in the submission of his bid, and it is found at a later stage that these applications were incorrect or made under false pretences, the Department may, at its own right:
 - i. Recover from the Contractor all costs, losses or damages incurred or sustained by the Department as a result of the award of the Contract; and/or
 - ii. Cancel the contract and claim any damages which the Department may suffer by having to make less favourable arrangements after such cancellation.
 - iii. The Department may impose penalties, however, only if provision therefore is made in the Special Conditions of Contract and Bid.

36. SEVERABILITY

36.1 The finding of any invalidity to any provision of the contract shall not render the whole contract a nullity. A court of law or arbitrator may sever the invalid provision and the remainder of the contract shall remain enforceable.

37. EXPORT LICENSES

- 37.1 When orders are placed for goods in respect of which an export licence from the country of origin of supplies is required, Contractor shall:
- 37.1.1 Not incur any direct or indirect costs in connection with the supply or dispatch of such supplies before they have obtained such license;
- 37.1.2 If the government of the country from which the supplies are to be exported refuses, or fails to grant such license within three months of the placing of the order, the order shall be considered to be cancelled and no liability will be accepted for any loss or expenses irrespective of the nature thereof, including loss or expenditure suffered or incurred by Contractor or any other person in respect of the production, supply, transportation or delivery of such supplies.

38 INSURANCE

- 38.1 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.
- 38.2 Any insurance policies taken out by Contractor to cover goods delivered for a contract must be taken out with a company registered in South Africa in terms of relevant insurance and companies acts.
- 38.3 The Department and the Contractor must ensure that the insurance remains in force throughout the contract period.
- 38.4 In the event that the Department requests for such Certificate of Insurance, the Contractor shall submit such Certificate within 5 days, if this was not a mandatory requirement.

39. ESTIMATED QUANTITIES

39.1 The Department is under no obligation to purchase any stock, which is in excess of the indicated quantities of each item. Should there be quantities reflected in the bid forms these will be estimated figures and no guarantee is given or implied as to the actual quantity which will be ordered.

40. EXTENTION OF CONTRACT

- 40.1 This contract may be extended on a month-to-month basis for a period not exceeding six (6) months.
- 40.2 Further extension of the contract, authority will be granted by Head of Department: Health, subject to the provisions of National Treasury Regulations and Instruction Notes.

41. CESSION OF CONTRACT

- 41.1 Cession refers to the transfer of only the rights a service provider has in terms of a contract from it to a third party. In commercial contracts, the main right involved is the right to be paid for services rendered. While the appointed bidder remains the service provider that continues to render the services, the service provider may cede (transfer) its right to be paid for the services it rendered in terms of the contract to a third party. This means that the service provider renders the services to an organ of state, while the organ of state pays for the services rendered to a third party instead, most commonly, a financial institution.
- 41.2 Cession will only be permissible on approval by the Accounting Officer.

42. CONTRACT AMENDMENTS / VARIATIONS

- 42.1 In amplification of paragraph 18 of the GCC, any amendments/variations, of the Contract shall come into effect in terms of the conditions contained in on "**Contract Amendments/Variations Register**". This register must be signed by the duly authorised signatories of winning bidder and the Head of Department: Health or his/her delegated official.
- 42.2 Contracted winning bidder shall not, in performing its obligation, vary from the terms and conditions stated in this Contract whether by way of addition thereto or by way of omission therefrom, without the prior written consent from the Department (Accounting Officer/delegated official), and no claim on the part of winning bidder for any extra payments on the grounds of any alterations or extra work will be entertained.
- 42.3 If, after the commencement of the contract, the cost or duration of the services is altered as a result of changes in, or in additions to, any statute, regulation or by-law, or the requirements of any authority having jurisdiction over any matter in respect of the contract, then the contract price and time for completion shall be adjusted in order to reflect the impact of those changes, provided that, within 14 days of first having become aware of the change, winning bidder shall furnish the Department with a detailed justification for the adjustment to the contract price.

43. SAMPLES

- 43.1. Samples will not be accepted with the closing of the bid document.
- 14.2 A sample meeting will be arranged with selected companies whereby the companies will be invited to forward their samples on a specified date and time.
- 14.3 Samples must be made available for the sample meeting, failure to provide a sample will reject their bid offer.
- 14.4 Samples shall be supplied by the bidder at his/her own cost/risk. Samples must be packaged as per the specification, in its original packagaing. Failure to do so will render the bid invalid.
- 14.5 Representative samples will not be accepted.
- 14.6 The Department reserves the right not to return such samples and to dispose of them at its discretion.
- 14.7 Samples must be clearly marked: Item number:
 - i. Brand Name
 - ii. Name of the Company
 - iii. Bid number
 - iv. Name of the manufacturer/supplier
 - v. Description of item
 - vi. Date of manufacture

14.8 The award of this bid will be based on the sample submitted from a manufacturer based on a letter of undertaking, which is compliant to specification. If, during the contract, the awarded supplier wishes to change the item being supplied, the service provider shall apply to the Department in writing, giving reasons why they want to change the product being supplied, which the Department shall consider. This process will be subject to the sample being submitted to the technical committee for evaluation and if in order, to the adjudication committee for approval. This will be done via the contract management unit of the Department. If there is a change in the product being supplied, and no prior approval has been granted, the Department reserves its right to cancel the contract.

N.B Failure to clearly mark the samples submitted shall result in the samples not being evaluated and eliminated from further consideration

44. INTELLECTUAL PROPERTY

44.1 In amplification of paragraph 6 of the GCC, the intellectual property discovered or created as the direct or indirect result of this contract shall remain the property of the Department.

45. INSOLVENCY

- 45.1 In the event to winning bidder institutes insolvency proceedings or has insolvency proceedings involuntarily instituted against it, the Department may terminate this Contract immediately.
- 45.2 In the event of assets and monies issued to winning bidder in terms of this Contract, such assets and monies shall be excluded from the estate of winning bidder and shall be returned immediately upon clause 40.1 coming into effect.

46. DISPUTE RESOLUTION

46.1 If any dispute arises between the Department and Contractor, in connection with the

Specification and deliverables, either party may give the other notice in writing of the existence of such dispute, and the same shall thereupon be referred to arbitration in South Africa by a person mutually agreed upon by both parties. The submission shall be deemed to be a submission to arbitration within the meaning of the terms of the arbitration laws in force in the Republic of South Africa.

47. DOMICILLIA CITANDI ET EXECUTANDI

For the purpose of this contract, the parties choose their respective domicillia citandi et executandi as follows :

The Department Physical and Postal Address:

Department Name	The KwaZulu- Natal Department of Health
Physical Address	Natalia Building, 330 Langalibalele Street, Pietermaritzburg, 3201
Postal Address:	Private Bag X9051, Pietermaritzburg, 3200
Telephone numbers	033 – 395 2111
Telefax:	Nil

The Contractor or Bidder Physical and Postal Address:

Bidder/ Contractor Name	
Physical Address	
Postal Address:	
Telephone numbers	
Telefax:	
Email Address	

- 47.1 The parties hereby choose <u>domicilium citandi et executandi</u> for all notices and processes to be given and served in pursuance hereof at their respective addresses given on the first page of this Contract. Any notice of any change in such address shall be given in writing by the parties concerned and delivered by hand or sent by registered mail to the other party, upon notification of which address so notified shall serve as the new <u>citandi et executandi</u>.
- 47.2 A party may at any time change that party's domicilium by notice in writing, provided that the new domicilium is in the Republic of South Africa and consists of, or includes, a physical address at which the process can be served.
- 47.3 Any notice to a party:
- 47.3.1 Sent by prepaid registered post in a correctly addressed envelope, to it, shall be deemed to have been received on the 7th (seventh) day after posting unless the contrary is proved);
- 47.3.2 Delivered by hand to a responsible person during ordinary business hours at the physical address chosen as its domicilium, shall be deemed to have been received on the day of delivery; or
- 47.3.3 Sent by telefax or email to its chosen telefax or email number, shall be deemed to have been received on the date of despatch (unless the contrary is proved).

LIST OF ITEMS: ZNB 6685/3/2023: RESPIRATORY AIDS: ENDOTRACHEAL TUBES AND TRACHEOSTOMY TUBES

AND ACCESSORIES

ENDOTRACHEAL TUBES:

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
1	CUFFED IVORY PVC ENDOTRACHEAL TUBES	999952U4778472	IC 1	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 5.0mm
		999952U4778484	IC 2	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 5.5mm
		999952U4778496	IC 3	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 6.0mm
		999952U4778508	IC 4	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 6.5mm
		999952U4778510	IC 5	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 7.0mm
3	UNCUFFED EXTRA-SOFT PLASTICIZED PVC ORAL/NASAL TRACHEAL	999952U4778534	33 125 50	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 2.0mm
	TUBE FOR LONG TERM INTUBATION	999952U4779740	33 125 51	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 2.5mm
		999952U4779765	33 125 52	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 3.0mm
		999952U4779777	33 125 53	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 3.5mm
		999952U4779789	33 125 54	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 4.0mm
		999952U4779791	33 125 55	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 4.5mm
		999952U4779803	33 125 56	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 5.0mm
		999952U4779815	33 125 57	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 5.5mm
		999952U4779827	33 125 58	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 6.0mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
4	SHORT-CUFFED ENDOTRACHEAL TUBE WITH SUBGLOTTIC PRESSURE-FREE ZONE,	999952U4779839	SC 1	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 3.0mm
	MAGILL TIP and NO MURPHY EYE	999952U4781783	SC 2	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 3.5mm
		999952U4781795	SC 3	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 4mm
		999952U4781807	SC 4	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 4.5mm
		999952U4781819	SC 5	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 5mm
		999952U4781860	SC 6	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 5.5mm
5	SHORT-CUFFED ORAL SOUTH-FACING PREFORMED (RAE) ENDOTRACHEAL TUBE WITH A SUBGLOTTIC	999952U4781922	SCOR 1	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 3.0
	PRESSURE-FREE ZONE, MAGILL TIP WITHOUT A MURPHY EYE	999952U4781950	SCOR 2	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 3.5mm
		999952U4781922	SCOR 3	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 4mm
		999952U4781950	SCOR 4	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 4.5mm
		999952U4781922	SCOR 5	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 5mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
		999952U4781950	SCOR 6	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 5.5mm
6	CUFFED ORAL/NASAL TRACHEAL TUBE WITH/ WITHOUT A MURPHY EYE -	999952U4781974	PCN1	Cuffed Oral/Nasal tracheal tube with/without a Murphy eye Size: 2.0 Age: New-born (< 2kg)
	NEW-BORNS	999952U4781986	PCN2	Cuffed Oral/Nasal tracheal tube with/without a Murphy eye Size: 2.5 Age: New-born (< 2.5kg)
7	CUFFED ORAL/NASAL TRACHEAL TUBE WITH A MURPHY EYE - PAEDIATRIC	999952U4781998	PCP 1	Cuffed Oral/Nasal tracheal tube with a Murphy eye ID: 3.0mm Age: Neonates (≥3 kg) to <1yr
		999952U4782001	PCP 2	Cuffed Oral/Nasal tracheal tube with a Murphy eye ID: 3.5mm Age: 1 to < 2yr
8	CUFFED ORAL/NASAL ENDOTRACHEAL TUBE WITH REDUCED TRAUMA TIP	999952U4782013	PCRT 1	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 3.0mm
		999952U4782025	PCRT 2	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 3.5mm
		999952U4782037	PCRT 3	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 4.0mm
		999952U4783039	PCRT 4	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 4.5mm
		999952U4783054	PCRT 5	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 5.0mm
		999952U4783078	PCRT 6	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 5.5mm
		999952U4783092	PCRT 7	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 6.0mm
		999952U4783116	PCRT 8	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 6.5mm
		999952U4783130	PCRT 9	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 7.0mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
		999952U4783142	PCRT 10	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 7.5mm
		999952U4783155	PCRT 11	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 8.0mm
		999952U4783179	PCRT 12	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 8.5mm
9	CUFFED REINFORCED ORAL/NASAL ENDOTRACHEAL TUBE - PAEDIATRIC	999952U4783181	RCP1	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 3.0mm OD: 4.8mm Length ± 184mm
		999952U4783193	RCP2	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 3.5mm OD: 5.5mm Length ± 185mm
		999952U4783205	RCP3	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 4.0mm OD: 6.0mm Length ± 195mm
		999952U4783217	RCP4	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 4.5mm OD: 6.5mm Length ± 198mm
10	UNCUFFED NASAL NORTH- FACING PREFORMED (RAE) ENDOTRACHEAL TUBE -	999952U4783243	NRUN1	Uncuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.0mm
	NEW-BORNS	999952U4783256	NRUN2	Uncuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.5mm
11	CUFFED NASAL NORTH RAE WITH/ WITHOUT MURPHY EYE - NEW-BORNS	999952U4783270	NRCN1	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.0mm
		999952U4783294	NRCN2	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.5mm
12	CUFFED NASAL NORTH RAE WITH MURPHY EYE - PAEDIATRIC	999952U4783306	NRC6	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 3.0mm
		999952U4783318	NRC7	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 3.5mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
		999952U4783460	NRC8	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 4.0mm
		999952U4783484	NRC9	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 4.5mm
		999952U4783496	NRC10	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 5.0mm
		999952U4783508	NRC11	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 5.5mm
13	TRACHEOSTOMY TUBES - UNCUFFED NEONATE	999952U4783546	TUNeo1	Tracheostomy Tube – Uncuffed Neonate ID: 2.5mm OD: ± 4.2mm Length: ± 28mm
		999952U4783559	TUNeo2	Tracheostomy Tube – Uncuffed Neonate ID: 3.0mm OD: ± 4.8mm Length: ± 30mm
		999952U4783561	TUNeo3	Tracheostomy Tube – Uncuffed Neonate ID: 3.5mm OD: ± 5.4mm Length: ± 32mm
		999952U4783573	TUNeo4	Tracheostomy Tube – Uncuffed Neonate ID: 4.0mm OD: ± 6.0mm Length: ± 34mm
		999952U4783585	TUNeo5	Tracheostomy Tube – Uncuffed Neonate ID: 4.5mm OD: ± 6.7mm Length: ± 36mm
14	TRACHEOSTOMY TUBES UNCUFFED PAEDIATRIC	999952U4783597	TUPaed1	Tracheostomy Tube – Uncuffed Paediatrics ID: 2.5mm OD: ± 4.2mm Length: ± 38mm
		999952U4783609	TUPaed2	Tracheostomy Tube – Uncuffed Paediatrics ID: 3.0mm
		999952U4783635	TUPaed3	Tracheostomy Tube – Uncuffed Paediatrics ID: 3.5mm
		999952U4784070	TUPaed4	Tracheostomy Tube – Uncuffed Paediatrics ID: 4.0mm
		999952U4784082	TUPaed5	Tracheostomy Tube – Uncuffed Paediatrics ID: 4.5mm
		999952U4784094	TUPaed6	Tracheostomy Tube – Uncuffed Paediatrics ID: 5.0mm
		999952U4784118	TUPaed7	Tracheostomy Tube – Uncuffed Paediatrics ID: 5.5mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
15	TRACHEOSTOMY TUBES UNCUFFED PAEDIATRIC - LONG	999952U4784132	TULPaed1	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 5.0mm OD: ± 7.1mm Length: ± 50mm
		999952U4784169	TULPaed2	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 5.5mm OD: ± 7.7mm Length: ± 52mm
		999952U4784171	TULPaed3	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 6.0mm OD: ± 8.3mm Length: ± 54mm
		999952U4784183	TULPaed4	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 6.5mm OD: ± 9.0mm Length: ± 56mm
16	TRACHEOSTOMY TUBES - ADULT UNCUFFED FENESTRATED	999952U4784195	TAUF1	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 4.0mm OD: ± 7.2mm Length: ± 59mm
		999952U4784207	TAUF2	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 5.0mm OD: ± 8.6mm Length: ± 66mm
		999952U4784219	TAUF3	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 6.0mm OD: ± 9.2mm Length: ± 72mm
		999952U4784308	TAUF4	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 7.0mm OD: ± 10.4mm Length: ± 74mm
		999952U4784310	TAUF5	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 8.0mm OD: ± 11.4mm Length:± 76mm
		999952U4784322	TAUF6	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 9.0mm OD: ± 12.5mm Length: ± 78mm
		999952U4784334	TAUF7	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 10.0mm OD: ± 13.8mm Length: ± 80mm
17	TRACHEOSTOMY TUBES - ADULT CUFFED FENESTRATED	999952U4784346	TACF1	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 4.0mm OD: ± 7.2mm Length: ± 59mm
		999952U4784359	TACF2	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 5.0mm OD: ± 8.6mm Length: ± 66mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
		999952U4784361	TACF3	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 6.0mm OD: ± 9.2mm Length: ± 72mm
		999952U4784373	TACF4	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 7.0mm OD: ± 10.4mm Length: ± 74mm
		999952U4784385	TACF5	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 8.0mm OD: ± 11.4mm Length:± 76mm
		999952U4784397	TACF6	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 9.0mm OD: ± 12.5mm Length: ± 78mm
		999952U4784409	TACF7	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 10.0mm OD: ± 13.8mm Length: ± 80mm
18	TRACHEOSTOMY TUBES - TAPER CUFFED	999952U4784411	TATC1	Tracheostomy Tube – Taper Cuffed ID: 6.5mm OD: 9.4mm Total Length: 62mm
		999952U4784423	TATC2	Tracheostomy Tube – Taper Cuffed ID: 7.0mm OD: 10.1mm Total Length: 68mm
		999952U4784435	TATC3	Tracheostomy Tube – Taper Cuffed ID: 7.5mm OD: 10.8mm Total Length: 74mm
		999952U4784447	TATC4	Tracheostomy Tube – Taper Cuffed ID: 8.0mm OD: 11.4mm Total Length: 77mm
		999952U4784450	TATC5	Tracheostomy Tube – Taper Cuffed ID: 8.5mm OD: 12.2mm Total Length: 79mm
		999952U4784462	TATC6	Tracheostomy Tube – Taper Cuffed ID: 9.0mm OD: 12.7mm Total Length: 79mm
19	TRACHEOSTOMY TUBES - TAPER CUFFED WITH SUBGLOTTIC SUCTION LINE	999952U4784474	TATCSL1	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 6.5mm OD: 9.4mm Total Length: 62mm
		999952U4784486	TATCSL2	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 7.0mm OD: 10.1mm Total Length: 68mm
		999952U4784500	TATCSL3	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 7.5mm OD: 10.8mm Total Length: 74mm
		999952U4784512	TATCSL4	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 8.0mm OD: 11.4mm Total Length: 77mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
		999952U4784524	TATCSL5	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 8.5mm OD: 12.2mm Total Length: 79mm
		999952U4784536	TATCSL6	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 9.0mm OD: 12.7mm Total Length: 79mm
20	TRACHEOSTOMY TUBES - CUFFED WITH SUBGLOTTIC SUCTION LINE	999952U4784587	TACSL1	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 4.0mm OD: 7.2mm Total Length: 59mm
		999952U4784601	TACSL2	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 5.0mm OD: 8.6mm Total Length: 66mm
		999952U4784613	TACSL3	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 6.0mm OD: 9.2mm Total Length: 72mm
		999952U4784676	TACSL4	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 7.0mm OD: 10.4mm Total Length: 74mm
		999952U4784688	TACSL5	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 8.0mm OD: 11.4mm Total Length: 76mm
		999952U4784690	TACSL6	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 9.0mm OD: 12.5mm Total Length: 78mm
		999952U4784702	TACSL7	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 10.0mm OD: 13.8mm Total Length: 98mm
21	TRACHEOSTOMY TUBE - CUFFED EXTRA-LONG AND SPIRAL REINFORCED	999952U4784714	TASR1	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 7.0mm OD: ± 9.7mm Length: ±104mm Cuff Size ± 22mm
		999952U4784726	TASR2	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 8.0mm OD: ± 11.2mm Length: ± 123mm Cuff Size ± 28mm
		999952U4784738	TASR3	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 9.0mm OD: ± 12.3mm Length: ± 126mm Cuff Size ± 30mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
		999952U4784740	TASR4	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 10.0mm OD: ± 13.7mm Length: ±134mm Cuff Size ± 32mm
22	TRACHEOSTOMY TUBE CUFFED EXTENDED LENGTH PROXIMAL	999952U4784753	TTEP1	Tracheostomy tube - Cuffed , Extended length - Proximal ID: 5.0mm ; OD: 9.6mm Total Length: 90mm Proximal length: 20mm Distal Length: 33mm Cuff diameter: 29mm
		999952U4784765	TTEP2	Tracheostomy tube - Cuffed, Extended length - Proximal ID: 6.0mm ; OD: 11mm Total Length: 95mm Proximal length: 23mm Distal Length: 34mm Cuff diameter: 31mm
		999952U4784777	TTEP3	Tracheostomy tube - Cuffed, Extended length - Proximal ID: 7.0mm OD: 12.3mm Total Length: 100mm Proximal length: 27mm Distal Length 34mm Cuff diameter: 35mm
		999952U4784789	TTEP4	Tracheostomy tube - Cuffed , Extended length - Proximal ID: 8.0mm OD: 13.3mm Total Length: 105mm Proximal length 30mm Distal Length 35mm Cuff diameter: 35mm
23	TRACHEOSTOMY TUBE CUFFED EXTENDED LENGTH DISTAL	999952U4784791	TTED1	Tracheostomy tube - Cuffed, Extended length - Distal ID: 5.0mm OD: 9.6mm Total Length: 90mm Proximal length: 5mm Distal Length: 48mm Cuff diameter: 29mm
		999952U4784803	TTED2	Tracheostomy tube - Cuffed, Extended length - Distal ID: 6.0mm OD: 11mm Total Length: 95mm Proximal length: 8mm Distal Length: 49mm Cuff diameter: 31mm
		999952U4784815	TTED3	Tracheostomy tube - Cuffed, Extended length - Distal ID: 7.0mm OD: 12.3mm Total Length: 100mm Proximal length: 12mm Distal Length: 49mm Cuff diameter: 35mm
		999952U4784827	TTED4	Tracheostomy tube - Cuffed, Extended length - Distal ID: 8.0mm OD: 13.3mm Total Length: 105mm Proximal length: 15mm Distal Length: 50mm Cuff diameter: 35mm
24	PERCUTANEOUS TRACHEOSTOMY KIT WITH TRACHEOSTOMY TUBE -	999952U4792596	PCACF1	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID: 7.0mm OD: ± 10.4mm Length: ± 74mm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
	ADULT CUFFED FENESTRATED	999952U4792608	PCACF2	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID: 8.0mm OD: ± 11.4mm Length:± 76mm
		999952U4792610	PCACF3	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID: 9.0mm OD: ± 12.5mm Length: ± 78mm
25	PERCUTANEOUS TRACHEOSTOMY DILATION KIT	999952U4792622	PCTD	Percutaneous Tracheostomy Dilation Kit
26	TRACHEOSTOMY BIB	999952U4792634	TB1	Tracheostomy Bib 3 Ply
		999952U4792646	TB2	Tracheostomy Bib 6 Ply
		999952U4792659	TB3	Tracheostomy Bib 12 Ply
27	TRACHEOSTOMY DRESSINGS	999952U4792661	TD1	Tracheostomy Dressings for Neonates and Children Size: ± 6.5 x 6.3cm Colour: White
		999952U4792685	TD2	Tracheostomy Dressings for Adults Size: ± 9.0 x 9.8cm Colour: White
28	TRACHEOSTOMY FOAM DRESSINGS	999952U4792697	TF1	Tracheostomy Foam dressings for Neonates and Children Size: ± 6.5 x 6.4cm Colour: Flesh
		999952U4792709	TF2	Tracheostomy Foam dressings for Adults Size: ± 9.0 x 9.8cm Colour: Flesh
29	TRACHEOSTOMY BRUSH - ANGLED	999952U4792854	TBA1	Tracheostomy Brush - Angled for Tube ID: 6mm Bristle diameter: 7mm Total Length: ± 21cm
		999952U4792711	TBA2	Tracheostomy Brush - Angled for Tube ID: 7/8mm Bristle diameter: 9mm Total Length: ± 21cm
		999952U4792723	ТВАЗ	Tracheostomy Brush - Angled for Tube ID: 9/10mm Bristle diameter: 11mm Total Length: ± 21cm
		999952U4792735	TBS1	Tracheostomy Brush - Straight for Tube ID: 5/6mm Bristle diameter: 7mm Total Length: ± 32cm
		999952U4792747	TBS2	Tracheostomy Brush - Straight for Tube ID: 7/8mm Bristle diameter: 9mm Total Length: ± 34cm

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
		999952U4792750	TBS3	Tracheostomy Brush - Straight for Tube ID: 9/10mm Bristle diameter: 11mm Total Length: ± 36cm
30	TRACHEOSTOMY CLEANING SWABS	999952U4792762	TCS1	Tracheostomy Cleaning Swab
31	STOMA BUTTONS	999952U4792774	SBS1	Stoma Button - Short Size: 6 ID: 6mm OD: 9mm Length: 22mm
		999952U4792786	SBS2	Stoma Button - Short Size: 7 ID: 7mm OD: 10mm Length: 22mm
		999952U4792800	SBS3	Stoma Button - Short Size: 8 ID: 8mm OD: 11mm Length: 22mm
		999952U4792812	SBS4	Stoma Button - Short Size: 9 ID: 9mm OD: 12mm Length: 22mm
		999952U4792836	SBS5	Stoma Button - Short Size: 10 ID: 10mm OD: 13mm Length: 22mm
		999952U4792848	SBS6	Stoma Button - Short Size: 11 ID: 11mm OD: 14mm Length: 22mm
		999952U4792863	SBL1	Stoma Button - Long Size: 6 ID: 6mm OD: 9mm Length: 30mm
		999952U4792875	SBL2	Stoma Button - Long Size: 7 ID: 7mm OD: 10mm Length: 30mm
		999952U4792887	SBL3	Stoma Button - Long Size: 8 ID: 8mm OD: 11mm Length: 30mm
		999952U4792899	SBL4	Stoma Button - Long Size: 9 ID: 9mm OD: 12mm Length: 30mm
		999952U4792901	SBL5	Stoma Button - Long Size: 10 ID: 10mm OD: 13mm Length: 30mm
		999952U4792913	SBL6	Stoma Button - Long Size: 11 ID: 11mm OD: 14mm Length: 30mm
32	TRACHEOSTOMY SPEAKING VALVES	999952U4792925	TSV1	Hands-Free Tracheostomy Speaking Valve
33	TRACHEOSTOMY MASK	999952U4792989	TM1	Tracheostomy masks - Paediatric
		999952U4792991	TM2	Tracheostomy masks - Adult

No.	CATEGORY	ICN NO:	ITEM:	DESCRIPTION:
34	TRACHEOSTOMY TUBE GUARD (T – PROTECTOR)	999952U4784839	TTG	Tracheostomy Tube Guard
35	TRACHEOSTOMY TAPE	999952U4784841	TT	Tracheostomy Tape Width: 12.5mm Roll: 1m

ENDOTRACHEAL TUBES:

CUFFED IVORY PVC ENDOTRACHEAL TUBES

Purpose: Cuffed Ivory PVC tube for short term oral or nasal intubation in paediatric patients

Consists of firm and kink resistant Ivory PVC tube with a full Magill curvature.

Must softens at body temperature to conform to anatomy

Must have a radio-opaque line and must be marked with numbered depth markings in 1cm intervals with tube size, length, ID, OD, trade name.

The markings on the tube must be clear and must not rub off when wiped with alcohol.

Must also have marks/reference lines to aid with depth of intubation and guide proper placement.

Must have a secure 15mm OD circuit connector made from medical grade plastic at the proximal end

Must have a bevelled atraumatic tip.

Must have a high-volume-low-pressure polyurethane cuff with pilot cuff

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation.

The cuff must be resistant to Nitrous Oxide effects and the pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

All components must be latex free, sterile and individually packed in peel pouch that is easy to open - For single use only

To comply with ISO 5361:2016 or equivalent

The following must be noted on the packaging:

Trade name; Size and specification; Method of sterilization; Manufacturing site; CE number; Lot number; Expiry date

ICN NO:	ITEM:	DESCRIPTION:
999952U4778472	IC 1	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 5.0mm OD: 7.3mm
999952U4778484	IC 2	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 5.5mm OD: 8.0mm
999952U4778496	IC 3	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 6.0mm OD: 8.8mm
999952U4778508	IC 4	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 6.5mm OD: 9.5mm
999952U4778510	IC 5	Cuffed Ivory PVC Oral/Nasal tracheal tube. ID: 7.0mm OD: 10.2mm

UNCUFFED EXTRA-SOFT PLASTICIZED PVC ORAL/NASAL TRACHEAL TUBE FOR LONG TERM INTUBATION

Purpose: Long term oral or nasal intubation in paediatric patients

Consists of a firm and kink resistant **extra-soft plasticized PVC** with a full Magill curvature that **softens at body temperature** to conform to anatomy

Must have a radiopaque line and must be marked with numbered depth markings in **50mm** or **1cm** intervals and the side of the tube with **tube size**, **length**, **ID**, **OD**, **trade name**.

The markings on the tube must be clear and must not rub off when wiped with alcohol.

Must have a precision band mark to aid with depth of intubation and guide proper placement.

Solid or circumferential precision band markings accepted and must be no more than 20mm from the tip

Proximal end - secure 15mm OD circuit connector made from medical grade plastic

Distal end - Atraumatic left-facing bevelled tip with **NO Murphy eye** Must be bevelled at **37°** with an **atraumatic tip**.All components must be latex and DEHP free, sterile and individually packed in peel pouch that is easy to open For SINGLE use only

To comply with the latest ISO 5361 or equivalent

Considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4778534	33 125 50	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 2.0mm OD: 3.4mm Length: 165mm Depth markings in 0.5cm intervals, starting at 3cm.
999952U4779740	33 125 51	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 2.5mm OD: 4.1mm Length: 165mm Depth markings in 0.5cm intervals, starting at 3cm.
999952U4779765	33 125 52	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 3.0mm OD: 4.6mm Length: 165mm Depth markings in 0.5cm intervals, starting at 3cm.
999952U4779777	33 125 53	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 3.5mm OD: 5.2mm Length: 165mm Depth markings in 0.5cm intervals starting at 3cm.
999952U4779789	33 125 54	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 4.0mm OD: 5.7mm Length: 230mm Depth markings in 1cm intervals from 9 - 21cm.
999952U4779791	33 125 55	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 4.5mm OD: 6.2mm Length: 230mm Depth markings in 1cm intervals from 9 - 21cm
999952U4779803	33 125 56	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 5.0mm OD: 7.0mm Length: 230mm Depth markings in 1cm intervals from 13 - 21cm
999952U4779815	33 125 57	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 5.5mm OD: 8.0mm Length: 270mm Depth markings in 1cm intervals from 15 - 25cm
999952U4779827	33 125 58	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 6.0mm OD: 8.5mm Length: 270mm Depth markings in 1cm intervals from 15 - 25cm

SHORT-CUFFED ENDOTRACHEAL TUBE with SUBGLOTTIC PRESSURE-FREE ZONE, MAGILL TIP and NO MURPHY EYE

Purpose: Short term oral or nasal intubation without a murphy's eye

Consists of a clear, kink-resistant, thermo-sensitive PVC tube that is atraumatic to airways and has a **full Magill curvature** with **no Murphy eye.**

Must have a radio-opaque line and The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name.

The markings on the tube must be clear and must not rub off when wiped with alcohol.

Must have a precision band mark just above the cuff to aid with depth of intubation.

Preference will be given to tubes with further markings to guide proper placement.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic.

Distal end - short atraumatic tip distal to a shortened cuff with no Murphy eye

The high-volume-low- pressure polyvinyl chloride cuff with pilot cuff must be short and cylindrical and made of micro-thin (10µ) polyurethane

Must be placed close to the tip of the tube and allow for a subglottic pressure-free zone.

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation. The deflated cuff must not add to the OD: circumference of the tube

The cuff must be resistant to the effects of Nitrous Oxide

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

All components must be **latex-free**, sterile and individually packed in peel pouch that is easy to open - For SINGLE use only

To comply with ISO 5361:2016 or equivalent

Considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4779839	SC 1	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 3.0mm OD: 4.3 - 5.0mm Length: 160 - 190mm Age: New-borns (≥3 kg) to <1yr Cuff length: ±10mm
999952U4781783	SC 2	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 3.5mm OD: 4.3 - 5.3mm Length: 190 - 220mm Age:1 to <2yr Cuff length: ±12mm
999952U4781795	SC 3	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 4mm OD: 5.6 - 6.0mm Length: 220 - 240mm Age:2 to <4yr Cuff length: ±12mm

ICN NO:	ITEM:	DESCRIPTION:
999952U4781807	SC 4	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 4.5mm OD: 6.2 - 6.3mm Length: 230 - 260mm Age: 4 to <6yr Cuff length:±14mm
999952U4781819	SC 5	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 5mm OD: 6.7 - 6.9mm Age: 6 to < 8yr Cuff length: ±14mm
999952U4781860	SC 6	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 5.5mm OD: 7.3 - 7.5mm Age: 8 to <10yr Cuff length: ±16mm
999952U4779839	SC 1	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 3.0mm OD: 4.3 - 5.0mm Length: 160 - 190mm Age: New-borns (≥3 kg) to <1yr Cuff length: ±10mm

SHORT-CUFFED ORAL SOUTH-FACING PREFORMED (RAE) ENDOTRACHEAL TUBE with a SUBGLOTTIC PRESSURE-FREE ZONE, MAGILL TIP and NO MURPHY EYE

Purpose: Short term oral or nasal intubation without a murphy's eye

Consists of a clear, kink-resistant, thermo-sensitive PVC tube with a **south-facing preformed curve** witho**ut Murphy eye**. Must be **radiopaque** or have a **radiopaque line**.

Must be atraumatic to the airways and able to temporarily straighten to allow the passage of a suction catheter.

The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings must be clear and must not rub off when wiped with alcohol.

Must have a precision band mark just above the cuff to aid with depth of intubation and guide proper placement.

Preference will be given to tubes with further markings to guide proper placement.

Must also have a **distinct marker** at the tube bend.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic

Distal end - short atraumatic tip distal to a shortened cuff without Murphy eye

The high-volume-low-pressure-cuff with pilot cuff must be short and cylindrical and made of micro-thin (10µ) Polyurethane

Must be placed close to the tip of the tube and allow for a **subglottic pressure-free zone**.

The cuff must inflate and deflate easily, and when deflated must conform closely to the tube to minimise the risk of trauma during intubation, and extubation and not increase the OD of the tube.

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

All components must be **latex-free**, **sterile and individually packed** in peel pouch that is easy to open - For SINGLE use only

To comply with the latest ISO 5361

Considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4781922	SCOR 1	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic
		Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 3.0 Age: New-borns (≥3 kg)
		to <1yr
999952U4781950	SCOR 2	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic
		Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 3.5mm Age:1 to <2yr
999952U4781922	SCOR 3	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic
		Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 4mm Age: 2 to <4yr
999952U4781950	SCOR 4	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic
		Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 4.5mm Age: 4 to <6yr
999952U4781922	SCOR 5	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic
		Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 5mm Age: 6 to < 8yr
999952U4781950	SCOR 6	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic
		Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 5.5mm Age: 8 to <10yr

CUFFED ORAL/NASAL TRACHEAL TUBE WITH/ WITHOUT A MURPHY EYE - NEW-BORNS

Purpose: Short term oral or nasal intubation

Consists of a **clear thermo-sensitive Polyvinyl Chloride (PVC)** tube that is kink resistant, atraumatic to airways and has a **full Magill curvature**. Must be radiopaque or have a radiopaque line

The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings must be clear and must not rub off when wiped with alcohol.

Must have a precision band mark just above the cuff to aid with depth of intubation and guide proper placement.

Solid or circumferential precision band markings accepted. **1 or 2** circumferential bands - tubes with a second circumferential band must have bands not more than **1cm** from each other to guide proper placement.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic.

Distal end - Atraumatic left-facing bevelled tip with NO or ONE Murphy eye on the opposite side

Must have a high-volume-low-pressure barrel-shaped polyvinyl chloride cuff with a pilot cuff. Cuff thickness ± 50 microns

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

The cuff must inflate and deflate easily, and when deflated must conform closely to the tube to minimise the risk of trauma during intubation, and extubation and not increase the OD of the tube.

Must be appropriately sized in relation to the tube size and appropriately placed with regards to the distance from the distal tip.

The cuff must be resistant to Nitrous Oxide effects.

All components must be latex-free sterile and individually packed in peel pouch that is easy to open; For SINGLE use only

To comply with the latest ISO 5361

Size 2.0mm and 2.5mm to be awarded INDIVIDUALLY

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4781974	PCN1	Cuffed Oral/Nasal tracheal tube with/without a Murphy eye Size: 2.0 Age: New-born (< 2kg)
999952U4781986	PCN2	Cuffed Oral/Nasal tracheal tube with/without a Murphy eye Size: 2.5 Age: New-born (< 2.5kg)

CUFFED ORAL/NASAL TRACHEAL TUBE WITH A MURPHY EYE - PAEDIATRIC

Purpose: Short term oral or nasal intubation

Consists of a clear thermo-sensitive **Polyvinyl Chloride (PVC)** tube that is kink resistant, atraumatic to airways and has a full Magill curvature. Must be radiopaque or have a radiopaque line

The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings must be clear and must not rub off when wiped with alcohol.

Must have a precision band mark just above the cuff to aid with depth of intubation and guide proper placement.

Solid or circumferential precision band markings accepted. 1 or 2 circumferential bands - tubes with a second circumferential band must have bands not more than 1cm from each other to guide proper placement.

Proximal end - secure OD:15mm circuit connector made from medical grade plastic

Distal end - Atraumatic left-facing bevelled tip with 1 or 2 Murphy eyes on the opposite side

Must have a high-volume-low-pressure barrel-shaped PVC cuff with a pilot cuff. Cuff thickness ± 50 microns

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

The cuff must inflate and deflate easily, and when deflated must conform closely to the tube to minimise the risk of trauma during intubation, and extubation and not increase the OD of the tube.

Must be appropriately sized in relation to the tube size and appropriately placed with regards to the distance from the distal tip.

The cuff must be resistant to Nitrous Oxide effects

All components must be latex-free sterile and individually packed in peel pouch that is easy to open; For SINGLE use only

To comply with the latest ISO 5361

Sizes 3mm - 6mm considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4781998	PCP 1	Cuffed Oral/Nasal tracheal tube with a Murphy eye ID: 3.0mm Age: Neonates (≥3 kg) to <1yr
999952U4782001	PCP 2	Cuffed Oral/Nasal tracheal tube with a Murphy eye ID: 3.5mm Age: 1 to < 2yr

CUFFED ORAL/NASAL ENDOTRACHEAL TUBE WITH REDUCED TRAUMA TIP - PAEDIATRIC

Purpose: Short term oral or nasal intubation with tip designed to reduce airway trauma

Consists of a clear **thermo-sensitive Polyvinyl Chloride (PVC) tube** that is kink resistant, atraumatic to airways and has a full Magill curvature and **two Murphy eyes**

Must be radiopaque or have a radiopaque line

The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings must be clear and must not rub off when wiped with alcohol.

Must have a **precision band mark** just above the cuff to aid with depth of intubation and guide proper placement. **Solid or circumferential** precision band markings accepted.

1 or 2 circumferential bands - tubes with a second circumferential band must have bands not more than **1cm** from each other to guide proper placement.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic

Distal end - Tapered, soft flexible pointed tip with an **upturned leading edge**, configured with the tip point in the midline so as to pass easily between the vocal cords without touching them.

Must have **2 Murphy Eyes.** The tube must be able to demonstrate a reduction in intubation induced airway trauma.

Must have a high-volume-low-pressure PVC barrel shaped cuff with pilot cuff. Cuff thickness ± 50 microns a shore hardness of no more than 86 microns

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation and not increase the OD of the tube.

Cuff must be barrel shaped to facilitate optimal pressure dispersion against the tracheal wall.

The cuff must be resistant to Nitrous Oxide effects

All components must be latex-free sterile and individually packed in peel pouch that is easy to open - For SINGLE use only

To comply with the latest ISO 5361

Size 3.0mm - 6.0mm considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4782013	PCRT 1	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 3.0mm
999952U4782025	PCRT 2	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 3.5mm
999952U4782037	PCRT 3	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 4.0mm
999952U4783039	PCRT 4	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 4.5mm
999952U4783054	PCRT 5	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 5.0mm
999952U4783078	PCRT 6	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 5.5mm
999952U4783092	PCRT 7	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 6.0mm

CUFFED ORAL/NASAL ENDOTRACHEAL TUBE WITH REDUCED TRAUMA TIP - ADULT

Purpose: Short term oral or nasal intubation with tip designed to reduce airway trauma

Consists of a clear **thermo-sensitive PVC tube** that is kink resistant, atraumatic to airways and has a full Magill curvature and two Murphy eyes

Must have a radio-opaque line and The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings on the tube must be clear and must not rub off.

Must have a precision band mark just above the cuff to aid with depth of intubation.

Preference will be given to tubes with a second circumferential band (not more than **2cm** from each other) to guide proper placement.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic

Distal end - Tapered, soft flexible pointed tip with an **upturned leading edge**, configured with the tip point in the midline so as to pass easily between the vocal cords without touching them.

The tube must be able to demonstrate a reduction in intubation induced airway trauma.

Must have a high-volume-low- pressure polyurethane barrel shaped cuff with pilot cuff. Cuff thickness ± 50 microns a shore hardness of no more than 86 microns

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation.

Cuff must be barrel shaped to facilitate optimal pressure dispersion against the tracheal wall

The cuff must be resistant to Nitrous Oxide effects

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

All components must be latex-free sterile and individually packed in peel pouch that is easy to open - For SINGLE use only

To comply with ISO 5361:2016 or equivalent

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4783116	PCRT 8	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 6.5mm
999952U4783130	PCRT 9	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 7.0mm
999952U4783142	PCRT 10	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 7.5mm
999952U4783155	PCRT 11	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 8.0mm
999952U4783179	PCRT 12	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 8.5mm

CUFFED REINFORCED ORAL/NASAL ENDOTRACHEAL TUBE - PAEDIATRIC

Purpose: Oral or nasal intubation where tube patency and flexibility is essential

Consists of a **soft flexible Polyvinyl Chloride (PVC)** tube with **spiral-wound medical grade reinforcing wire** that is sealed tightly against the **bonded 15mm connector**.

Must be kink resistant and atraumatic to airways. Must be radiopaque or have a radiopaque line

The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings must be clear and must not rub off when wiped with alcohol.

Must have a precision band mark to aid with depth of intubation and guide proper placement. **Solid or circumferential** precision band markings accepted

1 or 2 circumferential bands - tubes with a second circumferential band must have bands not more than **1cm** from each other to guide proper placement.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic

Distal end - Atraumatic left-facing bevelled tip with Murphy eye on the opposite side

Must have a high-volume-low-pressure PVC barrel shaped cuff with pilot cuff. Cuff thickness ± 50 microns.

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation and not increase the OD of the tube.

The cuff must be resistant to Nitrous Oxide effects

All components must be latex-free, sterile and individually packed in peel pouch that is easy to open - For SINGLE use only

To comply with the latest ISO 5361

Size 3mm - 6mm considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4783181	RCP1	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 3.0mm OD: 4.8mm Length ± 184mm
999952U4783193	RCP2	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 3.5mm OD: 5.5mm Length ± 185mm
999952U4783205	RCP3	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 4.0mm OD: 6.0mm Length ± 195mm
999952U4783217	RCP4	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 4.5mm OD: 6.5mm Length ± 198mm

UNCUFFED NASAL NORTH-FACING PREFORMED (RAE) ENDOTRACHEAL TUBE - NEW-BORNS

Purpose: Short term oral intubation where tube is North facing and removed from surgical field

Consists of a clear **thermo-sensitive Polyvinyl Chloride (PVC) tube** that is kink resistant, with a **North polar preformed curve** with a **soft flexible pointed tip** and **two Murphy eyes**

The tube must be able to straighten temporarily, to allow the passage of a suction catheter.

Must have a radiopaque line and The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings on the tube must be clear and must not rub off.

Must also have marks/reference lines proximal to cuff to aid with depth of intubation and guide proper placement. **Solid or circumferential** precision band markings accepted

1 or 2 circumferential bands - tubes with a second circumferential band must have bands not more than **1cm** from each other to guide proper placement.

Must also have a **distinct mark** at preformed curve to aid positioning.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic

Distal end - Atraumatic left-facing bevelled tip NO or 1 Murphy eyes on the opposite side

All components must be latex-free sterile and individually packed in peel pouch that is easy to open - For SINGLE use only

To comply with the latest ISO 5361

Size 2.0mm and 2.5mm considered INDIVIDUALLY

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4783243	NRUN1	Uncuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.0mm OD: 4.0mm Length ± 115mm
999952U4783256	NRUN2	Uncuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.5mm OD 4.0mm Length ± 115mm

CUFFED NASAL NORTH RAE WITH/ WITHOUT MURPHY EYE - NEW-BORNS

Purpose: Short term nasal intubation where tube is North facing and removed from surgical field

Consists of a clear **thermo-sensitive PVC tube** that is kink resistant, with a **North polar preformed curve** with/without **Murphy eyes**

The tube must be able to straighten temporarily, to allow the passage of a suction catheter.

Must have a radio-opaque line and the tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings on the tube must be clear and must not rub off.

Must have a precision band mark just above the cuff to aid with depth of intubation.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic

Distal end - Atraumatic left-facing bevelled tip with NO/1 Murphy eye on the opposite side.

Must have a high-volume-low- pressure polyvinyl chloride cuff with pilot cuff. Cuff thickness ± 50 microns.

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation.

The cuff must be resistant to Nitrous Oxide effects

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

All components must be latex-free sterile and individually packed in peel pouch that is easy to open - For SINGLE use only

To comply with ISO 5361:2016 or equivalent

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4783270	NRCN1	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.0mm
999952U4783294	NRCN2	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.5mm

CUFFED NASAL NORTH RAE WITH MURPHY EYE - PAEDIATRIC

Purpose: Short term nasal intubation where tube is North facing and removed from surgical field

Consists of a clear **thermo-sensitive PVC tube** that is kink resistant, with a **North polar preformed curve** and **two Murphy eyes**

The tube must be able to straighten temporarily, to allow the passage of a suction catheter.

Must have a radio-opaque line and The tube must be marked with numbered depth markings in 1cm intervals, tube size, length, ID, OD, trade name. The markings on the tube must be clear and must not rub off.

Must have a precision band mark just above the cuff to aid with depth of intubation.

Proximal end - secure 15mm OD circuit connector made from medical grade plastic

Distal end - Atraumatic left-facing bevelled tip with Murphy eyes on the opposite side.

Must have a high-volume-low- pressure polyvinyl chloride cuff with pilot cuff. Cuff thickness ± 50 microns.

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation.

The cuff must be resistant to Nitrous Oxide effects

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

All components must be latex-free sterile and individually packed in peel pouch that is easy to open - For SINGLE use only

To comply with ISO 5361:2016 or equivalent

Size 3mm - 5.5mm considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4783306	NRC6	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 3.0mm OD 4.0mm Length ± 115mm
999952U4783318	NRC7	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 3.5mm OD 5.3mm Length ± 155mm
999952U4783460	NRC8	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 4.0mm OD 5.3mm Length ± 155mm
999952U4783484	NRC9	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 4.5mm OD 6.0mm Length ± 180mm
999952U4783496	NRC10	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 5.0mm OD 6.7mm Length ± 190mm
999952U4783508	NRC11	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 5.5mm OD 7.3mm Length ± 195mm

TRACHEOSTOMY TUBES - UNCUFFED NEONATE

Purpose: To provide tracheal access in airway management.

Consists of an **uncuffed** radiopaque tube with a built in standard 15mm connector on the proximal end of the tube for direct connection to standard ventilation equipment. **Uncuffed** with **NO** inner tube.

Must come with an obturator/ stylet introducer and neck tape and must have a soft flange for comfort.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

Proof of use in SA tertiary hospital setting or sample to be assessed by end-user.

To comply with ISO 5366 or equivalent.

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4783546	TUNeo1	Tracheostomy Tube – Uncuffed Neonate ID: 2.5mm OD: ± 4.2mm Length: ± 28mm
999952U4783559	TUNeo2	Tracheostomy Tube – Uncuffed Neonate ID: 3.0mm OD: ± 4.8mm Length: ± 30mm
999952U4783561	TUNeo3	Tracheostomy Tube – Uncuffed Neonate ID: 3.5mm OD: ± 5.4mm Length: ± 32mm
999952U4783573	TUNeo4	Tracheostomy Tube – Uncuffed Neonate ID: 4.0mm OD: ± 6.0mm Length: ± 34mm
999952U4783585	TUNeo5	Tracheostomy Tube – Uncuffed Neonate ID: 4.5mm OD: ± 6.7mm Length: ± 36mm

TRACHEOSTOMY TUBES UNCUFFED PAEDIATRIC

Purpose: To provide tracheal access in airway management.

Consists of an uncuffed radiopaque tube with a built in standard **15mm** connector on the proximal end of the tube for direct connection to standard ventilation equipment. **Uncuffed** with **NO inner tube.**

Must come with an obturator/ stylet introducer and neck tape and must have a soft flange for comfort.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

Proof of use in SA tertiary hospital setting or sample to be assessed by end-user.

To comply with ISO 5366 or equivalent.

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4783597	TUPaed1	Tracheostomy Tube – Uncuffed Paediatrics ID: 2.5mm OD: ± 4.2mm Length: ± 38mm
999952U4783609	TUPaed2	Tracheostomy Tube – Uncuffed Paediatrics ID: 3.0mm OD: ± 4.8mm Length: ± 39mm
999952U4783635	TUPaed3	Tracheostomy Tube – Uncuffed Paediatrics ID: 3.5mm OD: ± 5.4mm Length: ± 40mm
999952U4784070	TUPaed4	Tracheostomy Tube – Uncuffed Paediatrics ID: 4.0mm OD: ± 6.0mm Length: ± 41mm
999952U4784082	TUPaed5	Tracheostomy Tube – Uncuffed Paediatrics ID: 4.5mm OD: ± 6.7mm Length: ± 42mm
999952U4784094	TUPaed6	Tracheostomy Tube – Uncuffed Paediatrics ID: 5.0mm OD: ± 7.3mm Length: ± 44mm
999952U4784118	TUPaed7	Tracheostomy Tube – Uncuffed Paediatrics ID: 5.5mm OD: ± 7.9mm Length: ± 46mm

TRACHEOSTOMY TUBES UNCUFFED PAEDIATRIC - LONG

Purpose: To provide tracheal access in airway management.

Consists of an uncuffed radiopaque tube with a built in standard 15mm connector on the proximal end of the tube for direct connection to standard ventilation equipment. **Uncuffed** with **NO inner tube.**

Must come with an obturator/ stylet introducer and neck tape and must have a soft flange for comfort.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

Proof of use in SA tertiary hospital setting or sample to be assessed by end-user.

To comply with ISO 5366 or equivalent.

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784132	TULPaed1	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 5.0mm OD: ± 7.1mm Length: ± 50mm
999952U4784169	TULPaed2	Tracheostomy Tube - Uncuffed Paediatrics Long ID: 5.5mm OD: ± 7.7mm Length: ± 52mm
999952U4784171	TULPaed3	Tracheostomy Tube - Uncuffed Paediatrics Long ID: 6.0mm OD: ± 8.3mm Length: ± 54mm
999952U4784183	TULPaed4	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 6.5mm OD: ± 9.0mm Length: ± 56mm

TRACHEOSTOMY TUBES - ADULT UNCUFFED FENESTRATED

Purpose: For use in providing tracheal access for long-term airway management and to allow easier vocalization.

Consists of an **uncuffed** radio-opaque tube with **multiple fenestrations** on the outer cannula. Must be thin walled to ensure maximum air flow.

Must have **1 x non-fenestrated inner cannula**, **1 x fenestrated inner cannula** with 15mm connectors and with a safe locking system; an occlusion cap.

Must come with a perforated obturator/ stylet introducer and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784195	TAUF1	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 4.0mm OD: ± 7.2mm Length: ± 59mm
999952U4784207	TAUF2	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 5.0mm OD: ± 8.6mm Length: ± 66mm
999952U4784219	TAUF3	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 6.0mm OD: ± 9.2mm Length: ± 72mm
999952U4784308	TAUF4	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 7.0mm OD: ± 10.4mm Length: ± 74mm
999952U4784310	TAUF5	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 8.0mm OD: ± 11.4mm Length:± 76mm
999952U4784322	TAUF6	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 8.0mm OD: ± 12.5mm Length: ± 78mm
999952U4784334	TAUF7	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 10.0mm OD: ± 13.8mm Length: ± 80mm

TRACHEOSTOMY TUBES - ADULT CUFFED FENESTRATED

Purpose: For use in providing tracheal access for long-term airway management and to allow easier vocalization.

Consists of a **cuffed** radio-opaque tube with **multiple fenestrations** on the outer cannula. Must be thin walled to ensure maximum air flow.

Must have **1 x non-fenestrated inner cannula**, **1 x fenestrated inner cannula** with 15mm connectors and with a **safe locking system**; an occlusion cap.

Must come with a **perforated obturator**/ **stylet** introducer and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Must have a low profile low-pressures-high-volume cuff to aid with ease of insertion. When deflated, the cuff must conform closely to the tube to minimize the risk of trauma during insertion and removal.

The pilot cuff must indicate the degree of inflation accurately and have a one-way valve.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784346	TACF1	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 4.0mm OD: ± 7.2mm Length: ± 59mm
999952U4784359	TACF2	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 5.0mm OD: ± 8.6mm Length: ± 66mm
999952U4784361	TACF3	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 6.0mm OD: ± 9.2mm Length: ± 72mm
999952U4784373	TACF4	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 7.0mm OD: ± 10.4mm Length: ± 74mm
999952U4784385	TACF5	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 8.0mm OD: ± 11.4mm Length:± 76mm
999952U4784397	TACF6	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 9.0mm OD: ± 12.5mm Length: ± 78mm
999952U4784409	TACF7	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 10.0mm OD: ± 13.8mm Length: ± 80mm

TRACHEOSTOMY TUBES - TAPER CUFFED

Purpose: For use in providing tracheal access for airway management and to aid with ventilation

Consists of a **cuffed** tracheostomy tube with a radio-opaque outer cannula. Must be thin walled to ensure maximum air flow.

Must have 1 x inner cannula with 15mm connectors and with a safe locking system; an occlusion cap.

Must come with a **perforated obturator**/ **stylet** introducer and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Must have a high-volume-low-pressure PVC taper-shaped cuff with pilot cuff. Cuff thickness ± 50 microns

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation.

Cuff must be **barrel shaped** to facilitate optimal pressure dispersion against the tracheal wall

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784411	TATC1	Tracheostomy Tube – Taper Cuffed ID: 6.5mm OD: 9.4mm Total Length: 62mm
999952U4784423	TATC2	Tracheostomy Tube – Taper Cuffed ID: 7.0mm OD: 10.1mm Total Length: 68mm
999952U4784435	TATC3	Tracheostomy Tube – Taper Cuffed ID: 7.5mm OD: 10.8mm Total Length: 74mm
999952U4784447	TATC4	Tracheostomy Tube – Taper Cuffed ID: 8.0mm OD: 11.4mm Total Length: 77mm
999952U4784450	TATC5	Tracheostomy Tube – Taper Cuffed ID: 8.5mm OD: 12.2mm Total Length: 79mm
999952U4784462	TATC6	Tracheostomy Tube – Taper Cuffed ID: 9.0mm OD: 12.7mm Total Length: 79mm

TRACHEOSTOMY TUBES - TAPER CUFFED WITH SUBGLOTTIC SUCTION LINE

Purpose: For use in providing tracheal access for airway management and to aid with ventilation

Consists of a **cuffed** tracheostomy tube with a radio-opaque outer cannula. Must be thin walled to ensure maximum air flow.

Must have **1 x inner cannula** with 15mm connectors and with a safe locking system; an occlusion cap.

Must come with a **perforated obturator/ stylet** introducer and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Must have a high-volume-low-pressure PVC taper-shaped cuff with pilot cuff. Cuff thickness ± 50 microns

The cuff must inflate and deflate easily and when deflated must conform closely to the tube to minimise the risk of trauma during intubation and extubation.

Cuff must be **barrel shaped** to facilitate optimal pressure dispersion against the tracheal wall

The pilot cuff must have a one-way valve and must indicate the degree of inflation accurately.

Subglottic Suction Line for patients with an expected ventilation period of more than 72 hour

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784474	TATCSL1	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 6.5mm OD: 9.4mm Total Length: 62mm
999952U4784486	TATCSL2	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 7.0mm OD: 10.1mm Total Length: 68mm
999952U4784500	TATCSL3	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 7.5mm OD: 10.8mm Total Length: 74mm
999952U4784512	TATCSL4	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 8.0mm OD: 11.4mm Total Length: 77mm
999952U4784524	TATCSL5	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 8.5mm OD: 12.2mm Total Length: 79mm
999952U4784536	TATCSL6	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 9.0mm OD: 12.7mm Total Length: 79mm

TRACHEOSTOMY TUBES - CUFFED WITH SUBGLOTTIC SUCTION LINE

Purpose: For use in providing tracheal access for airway management and to aid with ventilation and suctioning of secretions to prevent VAP

Consists of a **cuffed tracheostomy tube** with a radio-opaque outer cannula. Must be thin walled to ensure maximum air flow.

Must have 1 x inner cannula with 15mm connectors and with a safe locking system; an occlusion cap.

Must come with a **perforated obturator**/ **stylet introducer** and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Must have a low profile low-pressures-high-volume cuff to aid with ease of insertion. When deflated, the cuff must conform closely to the tube to minimize the risk of trauma during insertion and removal.

The pilot cuff must indicate the degree of inflation accurately and have a one-way valve.

Subglottic Suction Line for patients with an expected ventilation period of more than 72 hours

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784587	TACSL1	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 4.0mm OD: 7.2mm Total Length:
		59mm
999952U4784601	TACSL2	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 5.0mm OD: 8.6mm Total Length:
		66mm
999952U4784613	TACSL3	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 6.0mm OD: 9.2mm Total Length:
		72mm
999952U4784676	TACSL4	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 7.0mm OD: 10.4mm Total
		Length: 74mm
999952U4784688	TACSL5	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 8.0mm OD: 11.4mm Total
		Length: 76mm
999952U4784690	TACSL6	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 9.0mm OD: 12.5mm Total
		Length: 78mm
999952U4784702	TACSL7	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 10.0mm OD: 13.8mm Total
		Length: 98mm

TRACHEOSTOMY TUBE - CUFFED EXTRA-LONG AND SPIRAL REINFORCED

Purpose: For use in providing tracheal access for airway management and to aid with ventilation

Consists of an **extra-long**, **spiral-reinforced**, **cuffed**, **siliconised Tracheostomy Tube**. Must be thin walled to ensure maximum air flow

Must come with a **perforated obturator**/ **stylet introducer** and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Must have a low profile low-pressures-high-volume cuff to aid with ease of insertion. When deflated, the cuff must conform closely to the tube to minimize the risk of trauma during insertion and removal.

The pilot cuff must indicate the degree of inflation accurately and have a one-way valve.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784714	TASR1	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: \pm 7.0mm OD: \pm 9.7mm Length: \pm 104mm Cuff Size \pm 22mm
999952U4784726	TASR2	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 8.0mm OD: ± 11.2mm Length: ± 123mm Cuff Size ± 28mm
999952U4784738	TASR3	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: \pm 9.0mm OD: \pm 12.3mm Length: \pm 126mm Cuff Size \pm 30mm
999952U4784740	TASR4	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: \pm 10.0mm OD: \pm 13.7mm Length: \pm 134mm Cuff Size \pm 32mm

TRACHEOSTOMY TUBE CUFFED EXTENDED LENGTH PROXIMAL

Purpose: For use in providing tracheal access for airway management and to aid with ventilation in larger patients with full or thick necks

Consists of a cuffed tracheostomy tube with a radio-opaque outer cannula – that can adapt to **unusual anatomies**. Must be thin walled to ensure maximum air flow.

Must have a **flexible inner cannula** that conforms to the outer cannula's shape, with 15mm connector with a safe locking system; an occlusion cap. Must come with a perforated obturator/ stylet introducer and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Must have a low profile low-pressures-high-volume cuff to aid with ease of insertion. When deflated, the cuff must conform closely to the tube to minimize the risk of trauma during insertion and removal.

The pilot cuff must indicate the degree of inflation accurately and have a one-way valve.

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784753	TTEP1	Tracheostomy tube - Cuffed , Extended length - Proximal ID: 5.0mm ; OD: 9.6mm Total Length: 90mm Proximal length: 20mm Distal Length: 33mm Cuff diameter: 29mm
999952U4784765	TTEP2	Tracheostomy tube - Cuffed, Extended length - Proximal ID: 6.0mm ; OD: 11mm Total Length: 95mm Proximal length: 23mm Distal Length: 34mm Cuff diameter: 31mm
999952U4784777	TTEP3	Tracheostomy tube - Cuffed , Extended length - Proximal ID: 7.0mm OD: 12.3mm Total Length: 100mm Proximal length: 27mm Distal Length 34mm Cuff diameter: 35mm
999952U4784789	TTEP4	Tracheostomy tube - Cuffed , Extended length - Proximal ID: 8.0mm OD: 13.3mm Total Length: 105mm Proximal length 30mm Distal Length 35mm Cuff diameter: 35mm

TRACHEOSTOMY TUBE CUFFED EXTENDED LENGTH DISTAL

Purpose: For use in providing tracheal access for airway management and to aid with ventilation in patients with long tracheal anatomies, malacia or stenosis

Consists of a cuffed tracheostomy tube with a radio-opaque outer cannula – that can adapt to unusual anatomies. Must be thin walled to ensure maximum air flow.

Must have a flexible inner cannula that conforms to the outer cannula's shape, with 15mm connector with a safe locking system; an occlusion cap. Must come with a perforated obturator/ stylet introducer and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Must have a low profile low-pressures-high-volume cuff to aid with ease of insertion. When deflated, the cuff must conform closely to the tube to minimize the risk of trauma during insertion and removal.

The pilot cuff must indicate the degree of inflation accurately and have a one-way valve.

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4784791	TTED1	Tracheostomy tube - Cuffed, Extended length - Distal ID: 5.0mm OD: 9.6mm Total Length: 90mm Proximal length: 5mm Distal Length: 48mm Cuff diameter: 29mm
999952U4784803	TTED2	Tracheostomy tube - Cuffed, Extended length - Distal ID: 6.0mm OD: 11mm Total Length: 95mm Proximal length: 8mm Distal Length: 49mm Cuff diameter: 31mm
999952U4784815	TTED3	Tracheostomy tube - Cuffed, Extended length - Distal ID: 7.0mm OD: 12.3mm Total Length: 100mm Proximal length: 12mm Distal Length: 49mm Cuff diameter: 35mm
999952U4784827	TTED4	Tracheostomy tube - Cuffed, Extended length - Distal ID: 8.0mm OD: 13.3mm Total Length: 105mm Proximal length: 15mm Distal Length: 50mm Cuff diameter: 35mm

PERCUTANEOUS TRACHEOSTOMY KIT WITH TRACHEOSTOMY TUBE - ADULT CUFFED FENESTRATED

Purpose: Tracheostomy insertion at the bedside

The percutaneous dilation set must contain a scalpel, syringe, introducing needle, Straight dilator, Nitinol guidewire, a guiding catheter with a safety stop and a curved dilator.

The Nitinol Guidewire must be kink resistant and with an introducer and must be marked to assist with the alignment with the guiding catheter.

The Guiding Catheter must have marks that line up with the guidewire and a safety ridge to guide dilator distance and passage

The Curved Dilator must line up to the guiding catheter's safety ridge. Must have an ergonomic handle, must slide easily through the skin and tissue. Must have a skin level dilation diameter guide to prevent dilator from being inserted too deeply

The Tracheostomy tube must sit snuggly on a loading dilator and/or the curved dilator from the dilator set

Consists of a cuffed radio-opaque tube with multiple fenestrations on the outer cannula. Must be thin walled to ensure maximum air flow.

Must have 1 x non-fenestrated inner cannula, 1 x fenestrated inner cannula with 15mm connectors and with a safe locking system; an occlusion cap.

Must come with a perforated obturator/ stylet introducer and must have a soft anatomically shaped neck flange that moves on two axes: vertically and horizontally.

Must have a low profile low-pressures-high-volume cuff to aid with ease of insertion. When deflated, the cuff must conform closely to the tube to minimize the risk of trauma during insertion and removal.

The pilot cuff must indicate the degree of inflation accurately and have a one-way valve.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP PVC/ silicone that softens with body temperature.

Must have a robust design and the materials must be suitable for long term home tracheostomy care when needed.

Must have a wide neck strap manufactured of hypoallergenic material

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

For safety and standardisation purposes, considered in SERIES

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4792596	PCACF1	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID:
		7.0mm OD: ± 10.4mm Length: ± 74mm
999952U4792608	PCACF2	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID:
		8.0mm OD: ± 11.4mm Length:± 76mm
999952U4792610	PCACF3	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID:
		9.0mm OD: ± 12.5mm Length: ± 78mm

PERCUTANEOUS TRACHEOSTOMY DILATION KIT

Purpose: Tracheostomy insertion at the bedside

The percutaneous dilation set must contain a scalpel, syringe, introducing needle, Straight dilator, Nitinol guidewire, a guiding catheter with a safety stop and a curved dilator.

The Nitinol Guidewire must be kink resistant and with an introducer and must be marked to assist with the alignment with the guiding catheter.

The Guiding Catheter must have marks that line up with the guidewire and a safety ridge to guide dilator distance and passage

The Curved Dilator must line up to the guiding catheter's safety ridge. Must have an ergonomic handle, must slide easily through the skin and tissue. Must have a skin level dilation diameter guide to prevent dilator from being inserted too deeply

The appropriately branded Tracheostomy tube must sit snuggly on a loading dilator and/or the curved dilator from the dilator set

Sterile and individually packed in peel pouch that is easy to open, for SINGLE use only

To comply with ISO 5366 or equivalent

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4792622	PCTD	Percutaneous Tracheostomy Dilation Kit

TRACHEOSTOMY BIB

Purpose: To filter and warm the respiratory air and protect the patient's clothing from secretions.

Polyester bib with an adjustable velcro strap

ICN NO:	ITEM:	DESCRIPTION:
999952U4792634	TB1	Tracheostomy Bib 3 Ply
999952U4792646	TB2	Tracheostomy Bib 6 Ply
999952U4792659	ТВЗ	Tracheostomy Bib 12 Ply

TRACHEOSTOMY DRESSINGS

Purpose: Tracheal compresses used to absorb secretions from the tracheostoma and prevent skin irritations.

Compressive dressing with zigzag slit to ensure secure placement and ease of replacement.

For padding between tracheostoma and tracheostomy tube.

Sterile, single use, Latex-free open-pored polyurethane foam.

The following must be noted on the packaging:

ICN NO:	ITEM:	DESCRIPTION:
999952U4792661	TD1	Tracheostomy Dressings for Neonates and Children Size: ± 6.5 x 6.3cm Colour: White
999952U4792685	TD2	Tracheostomy Dressings for Adults Size: ± 9.0 x 9.8cm Colour: White

TRACHEOSTOMY FOAM DRESSINGS

Purpose: Tracheal compresses used to absorb secretions and reduces bacterial contamination

Compressive non-sterile dressing with zigzag slit to ensure secure placement and ease of replacement.

For padding between tracheostoma and tracheostomy tube.

Sterile, single use, Latex-free open-pored polyurethane foam.

The following must be noted on the packaging:

Trade name; Size and specification; Manufacturing site; CE number; Lot number; Expiry date

ICN NO:	ITEM:	DESCRIPTION:
999952U4792697	TF1	Tracheostomy Foam dressings for Neonates and Children Size: ± 6.5 x 6.4cm Colour: Flesh
999952U4792709	TF2	Tracheostomy Foam dressings for Adults Size: ± 9.0 x 9.8cm Colour: Flesh

TRACHEOSTOMY BRUSH - ANGLED

Purpose: To clean encrustations from tracheostomy tubes without damaging the tubes

Soft nylon bristles arrayed around a flexible stainless steel twisted wire base.

Must have a wool tip to remove incrustation, but no sharp edges that scratch the tracheostomy tube.

Must have a plastic handle with a hook to allow the brush to be hung up.

ICN NO:	ITEM:	DESCRIPTION:
999952U47928 54	TBA1	Tracheostomy Brush - Angled for Tube ID: 6mm Bristle diameter: 7mm Total Length: ± 21cm
999952U47927 11	TBA2	Tracheostomy Brush - Angled for Tube ID: 7/8mm Bristle diameter: 9mm Total Length: ± 21cm
999952U47927 23	TBA3	Tracheostomy Brush - Angled for Tube ID: 9/10mm Bristle diameter: 11mm Total Length: ± 21cm

TRACHEOSTOMY BRUSH - STRAIGHT

Purpose: To clean encrustations from tracheostomy tubes without damaging the tubes

Soft nylon bristles arrayed around a flexible stainless steel twisted wire base.

Must have no sharp edges that scratch the tracheostomy tube.

Must have a plastic handle with a hook to allow the brush to be hung up.

ICN NO:	ITEM:	DESCRIPTION:
999952U47927 35	TBS1	Tracheostomy Brush - Straight for Tube ID: 5/6mm Bristle diameter: 7mm Total Length: ± 32cm
999952U47927 47	TBS2	Tracheostomy Brush - Straight for Tube ID: 7/8mm Bristle diameter: 9mm Total Length: ± 34cm
999952U47927 50	TBS3	Tracheostomy Brush - Straight for Tube ID: 9/10mm Bristle diameter: 11mm Total Length: ± 36cm

TRACHEOSTOMY CLEANING SWABS

Purpose: To clean tracheostomy tubes and inner cannula

Consists of a bendable plastic swabs with polyurethane sponge tip for cleaning tracheostomy tubes

Must have no sharp edges that scratch the tracheostomy tube. Single use

ICN NO:	ITEM:	DESCRIPTION:
999952U47927 62	TCS1	Tracheostomy Cleaning Swab

STOMA BUTTON

Purpose: Designed to keep the stoma open and prevent narrowing of the stoma

Radio-opaque medical grade Silicone button with smooth edges. Must allow for normal neck movements and be comfortable.

Manufactured from biocompatible, hypoallergenic, latex-free, non-DEHP Silicone

Must be suitable for use with sterilizing fluids

ICN NO:	ITEM:	DESCRIPTION:
999952U47927 74	SBS1	Stoma Button - Short Size: 6 ID: 6mm OD: 9mm Length: 22mm
999952U47927 86	SBS2	Stoma Button - Short Size: 7 ID: 7mm OD: 10mm Length: 22mm
999952U47928 00	SBS3	Stoma Button - Short Size: 8 ID: 8mm OD: 11mm Length: 22mm
999952U47928 12	SBS4	Stoma Button - Short Size: 9 ID: 9mm OD: 12mm Length: 22mm
999952U47928 36	SBS5	Stoma Button - Short Size: 10 ID: 10mm OD: 13mm Length: 22mm
999952U47928 48	SBS6	Stoma Button - Short Size: 11 ID: 11mm OD: 14mm Length: 22mm
999952U47928 63	SBL1	Stoma Button - Long Size: 6 ID: 6mm OD: 9mm Length: 30mm
999952U47928 75	SBL2	Stoma Button - Long Size: 7 ID: 7mm OD: 10mm Length: 30mm
999952U47928 87	SBL3	Stoma Button - Long Size: 8 ID: 8mm OD: 11mm Length: 30mm
999952U47928 99	SBL4	Stoma Button - Long Size: 9 ID: 9mm OD: 12mm Length: 30mm
999952U47929 01	SBL5	Stoma Button - Long Size: 10 ID: 10mm OD: 13mm Length: 30mm
999952U47929 13	SBL6	Stoma Button - Long Size: 11 ID: 11mm OD: 14mm Length: 30mm

TRACHEOSTOMY SPEAKING VALVES

Purpose: Allows a patient with a tracheostomy to vocalize with heat moisture exchange

Plastic cap to fit securely to the 15mm connector of a fenestrated tracheostomy tube or a stoma button

Must contain a one way valve that allows the patient to inhale air but closes upon exhalation, causing a redirection of exhaled gas into the upper airway, thus allowing speech.

Individually packed; Single use; Latex free

Manufactured from medical grade plastic

The following must be noted on the packaging:

Size and specification; Manufacturing site; Expiry date

ICN NO:	ITEM:	DESCRIPTION:
999952U47929 25	TSV1	Hands-Free Tracheostomy Speaking Valve

TRACHEOSTOMY MASK

Purpose: Mask designed to fit over tracheostomy tube as protector or to deliver oxygen or humidification

Transparent mask shaped to fit over tracheostomy and fit comfortably in neck. Clear for easy visibility. Has an elastic band to secure the mask over the tube

Tubing connector must swivel 360 degrees and accept corrugated tubing with ID: 22 mm

Manufactured from medical grade plastic; Single use, individually packed; Clinically clean

The following must be noted on the packaging:

Trade name; Size and specification; Manufacturing site; CE number; Lot number

ICN NO:	ITEM:	DESCRIPTION:
999952U47929 89	TM1	Tracheostomy Mask - Paediatric
999952U47929 91	TM2	Tracheostomy Mask - Adult

TRACHEOSTOMY TUBE GUARD (T – PROTECTOR)

Purpose: Prevents accidental occlusion of and /or insertion of foreign body into the tracheostomy tube in paediatric patients not on the ventilator.

Medical grade plastic cap to fit securely onto a standard 15 mm connector of the tracheostomy tube. medical grade. Height: < 2cm

Must have multiple (minimum of 6) small openings around and on top of the guard to allow for easy exchange of air and secretion removal.

Single use, Latex free, Individually packed

The following must be noted on the packaging:

Trade name, Description; Manufacturing site; Expiry date

ICN NO:	ITEM:	DESCRIPTION:
999952U47848 39	TTG	Tracheostomy Tube Guard

TRACHEOSTOMY TAPE	
Purpose: To secure the tracheostomy in place.	
White non- fraying, flat cotton tape. Packed in 1m rolls	
Roll packed individually.	
Per meter	

ICN NO:	ITEM:	DESCRIPTION:
999952U47848 41	TT	Tracheostomy Tape Width: 12.5mm Roll: 1m

SECTION L: EVALUATION CRITERIA

The Department will evaluate applications received before the closing date and time using Three (3) evaluation phases these are peremptory requirements, should the applicant fail to comply, the application will be regarded as non-responsive and be disqualified. The criteria are as follows:

- Phase 1: Minimum Compulsory Requirements
- Phase 2: Technical Evaluation
- Phase 3: Price and Preference Points

Phase 1: Minimum Compulsory Requirements

The Bidder shall complete and submit the following returnable schedules and documents:

		COMPULSORY (YES / NO)	COMPULSORY	FOR OFFICIAL USE ONLY		
NO.	SECTION/ SCHEDULE	NON- SUBMISSION WILL RENDER BIDDERS NON- RESPONSIVE	(YES / NO) FOR BID EVALUATION PURPOSES	YES	NO	N/A
Prospec	tive Bidders MUST ensure that the following Section	ons of the bid docur	ment MUST be cor	npleted	in AL	L
respects	to qualify for the next stage of evaluation:					
1.1	Section A: Invitation To Bid	Yes	Yes			
1.2	Section B: Special Instructions and Notices to Bidders	Yes	Yes			
1.3	Section C: Authority To Sign A Bid	Yes	Yes			
1.4	Section D: Bidder's Disclosure (SBD 4)	Yes	Yes			
1.5	Section E: The National Industrial Participation Programme (SBD 5)	Yes	Yes			
1.6	Section F: Central Supplier Database (CSD)	Yes	Yes			
1.7	Section G: Preference Points Claim	Yes	Yes			
1.8	Section H: General Conditions Of Contract(GCC)	Yes	Yes			
1.9	Section I : Special Conditions Of Contract (SCC)	Yes	Yes			
1.10	Section J: Specification	Yes	Yes			
1.11	Section K: Pricing Schedule	Yes	Yes			
Prospec	tive Bidders MUST provide the following as per the	e Mandatory Requir	ements:		•	•
1	Copy of the Consortium/ Joint Venture/ Partnership agreement, if applicable	Yes If Applicable				
2	Relevant compliance certificates/Equivalent applicable to each item as per specification	Yes	Yes			

		COMPULSORY (YES / NO)	COMPULSORY		r offic Se onl	
NO.	SECTION/ SCHEDULE	NON- SUBMISSION WILL RENDER BIDDERS NON- RESPONSIVE	(YES / NO) FOR BID EVALUATION PURPOSES	YES	NO	N/A
3	Letter of undertaking if not the manufacturer of the Equipment, for each item as per specification	Yes	Yes			
4	SAHPRA certification	Yes	Yes			
3. Prosp stage	ective bidder must provide the following as addition	nal Requirement fro	om Main Contract	tor duri	ng cor	ntract
3	 B-BBEE certificate indicating the B-BBEE status level of contributor. The B-BBEE certificate must be issued by a SANAS accredited verification agency; Or A duly completed Sworn Affidavit signed by the deponent and commissioned by the authorized commissioner of oaths. The sworn affidavit must indicate the day, month and year on which the annual total revenue is based on and the level of black ownership that is claimed; Or A sworn affidavit on an accredited template issued by the DTI/CIPC for both EME or QSE, Note: i. Bidders must ensure that the correct sworn affidavit for the Financial Sector are submitted, ii. A trust, consortium, or joint venture (including unincorporated consortia and joint ventures) must submit a consolidated B-BBEE status level certificate. iii. The B-BBEE certificate or sworn affidavit will be required from service provider, during signing of contract 	Yes	Will only be Required from awarded service provider during Contract Management phase			

Phase 2: Technical Evaluation of Bid

The prospective bidder will be required to provide a sample for evaluation purposes as required in terms of the special terms and conditions of the bid. Samples per item must be accompanied with relevant Certification as per specification. Offers must comply strictly with the specification. Offers exceeding specification requirements will be deemed to comply with the specification. The quality of services/ supply must not be less than what is specified.

Phase 3: Price and Preference Points

The value of this bid is estimated to exceed or not to exceed R 50 000 000 (all applicable taxes included) and therefore the 80/20 or 90/10 preference point system shall be applicable.

Points for this bid shall be awarded for:

Price, and Specific Goals

The maximum points for this bid are allocated as follows:

CATEGORY	POINTS
PRICE	80 OR 90
SPECIFIC GOALS	20 OR 10
Total points for Price and must not exceed	100

The Department has identified the following specific goal:

- full points(20 points) to companies who are at least 51% Owned by Black People

Ownership verification will be conducted through Central Suppliers (CSD) Database by National Treasury, through the B-BBEE scorecard attributes or Companies and Intellectual Property Commission (CIPC). Bidders must submit CSD report and CIPC

Failure on the part of a bidder to submit proof of specific goals together with the bid will be interpreted to mean that preference points for specific goals are not claimed.

SECTION M: PRICING SCHEDULE

Name of bidder.....

Bid number: **ZNB 6685/3/2023-H**

Closing Time 11:00

Closing Date: 28 September 2023

ICN NO:	ITEM:	DESCRIPTION:	Unit Price Year 1 (incl. VAT)	Unit Price Year 2 (incl. VAT)	Unit Price Year 3 (incl. VAT)	Total Price Year 1 + year 2 + Year 3 (incl. VAT)	BRAND NAME
999952U4778472		Cuffed Ivory PVC Oral/Nasal tracheal					
	IC 1	tube. ID: 5.0mm					
999952U4778484		Cuffed Ivory PVC Oral/Nasal tracheal					
	IC 2	tube. ID: 5.5mm					
999952U4778496		Cuffed Ivory PVC Oral/Nasal tracheal					
	IC 3	tube. ID: 6.0mm					
999952U4778508		Cuffed Ivory PVC Oral/Nasal tracheal					
	IC 4	tube. ID: 6.5mm					
999952U4778510		Cuffed Ivory PVC Oral/Nasal tracheal					
	IC 5	tube. ID: 7.0mm					
999952U4778534	33 125 50	Uncuffed Extra-soft Plasticized PVC					
		Oral/Nasal tracheal tube for long-term					
99995204778534	33 125 50						

OFFER TO BE VALID FOR 180 DAYS FROM THE CLOSING DATE OF BID

ICN NO:	ITEM:	DESCRIPTION:	Unit Price Year 1 (incl. VAT)	Unit Price Year 2 (incl. VAT)	Unit Price Year 3 (incl. VAT)	Total Price Year 1 + year 2 + Year 3 (incl. VAT)	BRAND NAME
999952U4779740	33 125 51	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 2.5mm					
999952U4779765	33 125 52	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 3.0mm					
999952U4779777	33 125 53	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 3.5mm					
999952U4779789	33 125 54	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 4.0mm					
999952U4779791	33 125 55	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 4.5mm					
999952U4779803	33 125 56	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 5.0mm					
999952U4779815	33 125 57	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 5.5mm					
999952U4779827	33 125 58	Uncuffed Extra-soft Plasticized PVC Oral/Nasal tracheal tube for long-term intubation. ID: 6.0mm					

ICN NO:	ITEM:	DESCRIPTION:	Unit Price Year 1 (incl. VAT)	Unit Price Year 2 (incl. VAT)	Unit Price Year 3 (incl. VAT)	Total Price Year 1 + year 2 + Year 3 (incl. VAT)	BRAND NAME
999952U4779839	SC 1	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 3.0mm					
999952U4781783	SC 2	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 3.5mm					
999952U4781795	SC 3	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 4mm					
999952U4781807	SC 4	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 4.5mm					
999952U4781819	SC 5	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 5mm					
999952U4781860	SC 6	Short-cuffed endotracheal tube with Subglottic Pressure-Free Zone, Magill Tip and NO Murphy Eye ID: 5.5mm					
999952U4781922	SCOR 1	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 3.0					

ICN NO:	ITEM:	DESCRIPTION:	Unit Price Year 1 (incl. VAT)	Unit Price Year 2 (incl. VAT)	Unit Price Year 3 (incl. VAT)	Total Price Year 1 + year 2 + Year 3 (incl. VAT)	BRAND NAME
999952U4781950	SCOR 2	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 3.5mm					
999952U4781922	SCOR 3	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 4mm					
999952U4781950	SCOR 4	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 4.5mm					
999952U4781922	SCOR 5	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 5mm					
999952U4781950	SCOR 6	Short-Cuffed Oral South-Facing Preformed (RAE) Endotracheal Tube with Subglottic Pressure-Free Zone, Magill Tip without a Murphy Eye ID: 5.5mm					

ICN NO:	ITEM:	DESCRIPTION:	Unit Price Year 1 (incl. VAT)	Unit Price Year 2 (incl. VAT)	Unit Price Year 3 (incl. VAT)	Total Price Year 1 + year 2 + Year 3 (incl. VAT)	BRAND NAME
999952U4781974	PCN1	Cuffed Oral/Nasal tracheal tube with/without a Murphy eye Size: 2.0 Age: New-born (< 2kg)					
999952U4781986	PCN2	Cuffed Oral/Nasal tracheal tube with/without a Murphy eye Size: 2.5 Age: New-born (< 2.5kg)					
999952U4781998	PCP 1	Cuffed Oral/Nasal tracheal tube with a Murphy eye ID: 3.0mm Age: Neonates (≥3 kg) to <1yr					
999952U4782001	PCP 2	Cuffed Oral/Nasal tracheal tube with a Murphy eye ID: 3.5mm Age: 1 to < 2yr					
999952U4782013	PCRT 1	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 3.0mm					
999952U4782025	PCRT 2	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 3.5mm					
999952U4782037	PCRT 3	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 4.0mm					
999952U4783039	PCRT 4	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 4.5mm					
999952U4783054	PCRT 5	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 5.0mm					
999952U4783078	PCRT 6	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 5.5mm					

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999952U4783092	PCRT 7	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 6.0mm					
999952U4783116	PCRT 8	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 6.5mm					
999952U4783130	PCRT 9	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 7.0mm					
999952U4783142	PCRT 10	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 7.5mm					
999952U4783155	PCRT 11	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 8.0mm					
999952U4783179	PCRT 12	Cuffed Oral/Nasal endotracheal tube with Reduced Trauma Tip: ID: 8.5mm					
999952U4783181	RCP1	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 3.0mm OD: 4.8mm Length ± 184mm					
999952U4783193	RCP2	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 3.5mm OD: 5.5mm Length ± 185mm					
999952U4783205	RCP3	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 4.0mm OD: 6.0mm Length ± 195mm					

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999952U4783217	RCP4	Cuffed Reinforced Oral/Nasal endotracheal tube - Paediatric ID: 4.5mm OD: 6.5mm Length ± 198mm					
999952U4783243	NRUN1	Uncuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.0mm					
999952U4783256	NRUN2	Uncuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.5mm					
999952U4783270	NRCN1	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.0mm					
999952U4783294	NRCN2	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - New-borns ID: 2.5mm					
999952U4783306	NRC6	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 3.0mm					
999952U4783318	NRC7	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 3.5mm					

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999952U4783460	NRC8	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 4.0mm					
999952U4783484	NRC9	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 4.5mm					
999952U4783496	NRC10	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 5.0mm					
999952U4783508	NRC11	Cuffed Nasal North-Facing Preformed (RAE) Endotracheal Tube - Paediatric ID: 5.5mm					
999952U4783546	TUNeo1	Tracheostomy Tube – Uncuffed Neonate ID: 2.5mm OD: ± 4.2mm Length: ± 28mm					
999952U4783559	TUNeo2	Tracheostomy Tube – Uncuffed Neonate ID: 3.0mm OD: ± 4.8mm Length: ± 30mm					
999952U4783561	TUNeo3	Tracheostomy Tube – Uncuffed Neonate ID: 3.5mm OD: ± 5.4mm Length: ± 32mm					

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999952U4783573	TUNeo4	Tracheostomy Tube – Uncuffed Neonate ID: 4.0mm OD: ± 6.0mm Length: ± 34mm					
999952U4783585	TUNeo5	Tracheostomy Tube – Uncuffed Neonate ID: 4.5mm OD: ± 6.7mm Length: ± 36mm					
999952U4783597	TUPaed1	Tracheostomy Tube – Uncuffed Paediatrics ID: 2.5mm OD: ± 4.2mm Length: ± 38mm					
999952U4783609	TUPaed2	Tracheostomy Tube – Uncuffed Paediatrics ID: 3.0mm					
999952U4783635	TUPaed3	Tracheostomy Tube – Uncuffed Paediatrics ID: 3.5mm					
999952U4784070	TUPaed4	Tracheostomy Tube – Uncuffed Paediatrics ID: 4.0mm					
999952U4784082	TUPaed5	Tracheostomy Tube – Uncuffed Paediatrics ID: 4.5mm					
999952U4784094	TUPaed6	Tracheostomy Tube – Uncuffed Paediatrics ID: 5.0mm					
999952U4784118	TUPaed7	Tracheostomy Tube – Uncuffed Paediatrics ID: 5.5mm					

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999952U4784132	TULPaed1	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 5.0mm OD: ± 7.1mm Length: ± 50mm					
999952U4784169	TULPaed2	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 5.5mm OD: ± 7.7mm Length: ± 52mm					
999952U4784171	TULPaed3	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 6.0mm OD: ± 8.3mm Length: ± 54mm					
999952U4784183	TULPaed4	Tracheostomy Tube – Uncuffed Paediatrics Long ID: 6.5mm OD: ± 9.0mm Length: ± 56mm					
999952U4784195	TAUF1	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 4.0mm OD: ± 7.2mm Length: ± 59mm					
999952U4784207	TAUF2	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 5.0mm OD: ± 8.6mm Length: ± 66mm					
999952U4784219	TAUF3	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 6.0mm OD: ± 9.2mm Length: ± 72mm					

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999952U4784308	TAUF4	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 7.0mm OD: ± 10.4mm Length: ± 74mm					
999952U4784310	TAUF5	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 8.0mm OD: ± 11.4mm Length:± 76mm					
999952U4784322	TAUF6	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 9.0mm OD: ± 12.5mm Length: ± 78mm					
999952U4784334	TAUF7	Tracheostomy Tube – Adult Uncuffed Fenestrated ID: 10.0mm OD: ± 13.8mm Length: ± 80mm					
999952U4784346	TACF1	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 4.0mm OD: ± 7.2mm Length: ± 59mm					
999952U4784359	TACF2	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 5.0mm OD: ± 8.6mm Length: ± 66mm					
999952U4784361	TACF3	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 6.0mm OD: ± 9.2mm Length: ± 72mm					

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999952U4784373	TACF4	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 7.0mm OD: ± 10.4mm Length: ± 74mm					
999952U4784385	TACF5	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 8.0mm OD: ± 11.4mm Length:± 76mm					
999952U4784397	TACF6	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 9.0mm OD: ± 12.5mm Length: ± 78mm					
999952U4784409	TACF7	Tracheostomy Tube – Adult Cuffed Fenestrated ID: 10.0mm OD: ± 13.8mm Length: ± 80mm					
999952U4784411	TATC1	Tracheostomy Tube – Taper Cuffed ID: 6.5mm OD: 9.4mm Total Length: 62mm					
999952U4784423	TATC2	Tracheostomy Tube – Taper Cuffed ID: 7.0mm OD: 10.1mm Total Length: 68mm					
999952U4784435	TATC3	Tracheostomy Tube – Taper Cuffed ID: 7.5mm OD: 10.8mm Total Length: 74mm					
999952U4784447	TATC4	Tracheostomy Tube – Taper Cuffed ID: 8.0mm OD: 11.4mm Total Length: 77mm					

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999952U4784450	TATC5	Tracheostomy Tube – Taper Cuffed ID: 8.5mm OD: 12.2mm Total Length: 79mm					
999952U4784462	TATC6	Tracheostomy Tube – Taper Cuffed ID: 9.0mm OD: 12.7mm Total Length: 79mm					
999952U4784474	TATCSL1	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 6.5mm OD: 9.4mm Total Length: 62mm					
999952U4784486	TATCSL2	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 7.0mm OD: 10.1mm Total Length: 68mm					
999952U4784500	TATCSL3	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 7.5mm OD: 10.8mm Total Length: 74mm					
999952U4784512	TATCSL4	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 8.0mm OD: 11.4mm Total Length: 77mm					

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999952U4784524	TATCSL5	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 8.5mm OD: 12.2mm Total Length: 79mm					
999952U4784536	TATCSL6	Tracheostomy Tube – Taper Cuffed with Subglottic Suction Line ID: 9.0mm OD: 12.7mm Total Length: 79mm					
999952U4784587	TACSL1	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 4.0mm OD: 7.2mm Total Length: 59mm					
999952U4784601	TACSL2	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 5.0mm OD: 8.6mm Total Length: 66mm					
999952U4784613	TACSL3	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 6.0mm OD: 9.2mm Total Length: 72mm					
999952U4784676	TACSL4	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 7.0mm OD: 10.4mm Total Length: 74mm					
999952U4784688	TACSL5	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 8.0mm OD: 11.4mm Total Length: 76mm					

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999952U4784690	TACSL6	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 9.0mm OD: 12.5mm Total Length: 78mm					
999952U4784702	TACSL7	Tracheostomy Tube – Cuffed with Subglottic Suction Line ID: 10.0mm OD: 13.8mm Total Length: 98mm					
999952U4784714	TASR1	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 7.0mm OD: ± 9.7mm Length: ±104mm Cuff Size ± 22mm					
999952U4784726	TASR2	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 8.0mm OD: ± 11.2mm Length: ± 123mm Cuff Size ± 28mm					
999952U4784738	TASR3	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 9.0mm OD: ± 12.3mm Length: ± 126mm Cuff Size ± 30mm					
999952U4784740	TASR4	Tracheostomy Tube – Cuffed Extra-long and Spiral-Reinforced ID: ± 10.0mm OD: ± 13.7mm Length: ± 134mm Cuff Size ± 32mm					
999952U4784753	TTEP1	Tracheostomy tube - Cuffed , Extended length - Proximal ID: 5.0mm ; OD: 9.6mm Total Length: 90mm Proximal					

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		length: 20mm Distal Length: 33mm Cuff diameter: 29mm					
999952U4784765	TTEP2	Tracheostomy tube - Cuffed, Extended length - Proximal ID: 6.0mm ; OD: 11mm Total Length: 95mm Proximal length: 23mm Distal Length: 34mm Cuff diameter: 31mm					
999952U4784777	TTEP3	Tracheostomy tube - Cuffed , Extended length - Proximal ID: 7.0mm OD: 12.3mm Total Length: 100mm Proximal length: 27mm Distal Length 34mm Cuff diameter: 35mm					
999952U4784789	TTEP4	Tracheostomy tube - Cuffed , Extended length - Proximal ID: 8.0mm OD: 13.3mm Total Length: 105mm Proximal length 30mm Distal Length 35mm Cuff diameter: 35mm					
999952U4784791	TTED1	Tracheostomy tube - Cuffed, Extended length - Distal ID: 5.0mm OD: 9.6mm Total Length: 90mm Proximal length: 5mm Distal Length: 48mm Cuff diameter: 29mm					

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999952U4784803	TTED2	Tracheostomy tube - Cuffed, Extended length - Distal ID: 6.0mm OD: 11mm Total Length: 95mm Proximal length: 8mm Distal Length: 49mm Cuff diameter: 31mm					
999952U4784815	TTED3	Tracheostomy tube - Cuffed, Extended length - Distal ID: 7.0mm OD: 12.3mm Total Length: 100mm Proximal length: 12mm Distal Length: 49mm Cuff diameter: 35mm					
999952U4784827	TTED4	Tracheostomy tube - Cuffed, Extended length - Distal ID: 8.0mm OD: 13.3mm Total Length: 105mm Proximal length: 15mm Distal Length: 50mm Cuff diameter: 35mm					
999952U4792596	PCACF1	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID: 7.0mm OD: ± 10.4mm Length: ± 74mm					
999952U4792608	PCACF2	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID: 8.0mm OD: ± 11.4mm Length:± 76mm					

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999952U4792610	PCACF3	Percutaneous Tracheostomy Kit With Tracheostomy Tube - Adult Cuffed Fenestrated ID: 9.0mm OD: ± 12.5mm Length: ± 78mm					
999952U4792622	PCTD	Percutaneous Tracheostomy Dilation Kit					
999952U4792634	TB1	Tracheostomy Bib 3 Ply					
999952U4792646	TB2	Tracheostomy Bib 6 Ply					
999952U4792659	TB3	Tracheostomy Bib 12 Ply					
999952U4792661	TD1	Tracheostomy Dressings for Neonates and Children Size: ± 6.5 x 6.3cm Colour: White					
999952U4792685	TD2	Tracheostomy Dressings for Adults Size: ± 9.0 x 9.8cm Colour: White					
999952U4792697	TF1	Tracheostomy Foam dressings for Neonates and Children Size: ± 6.5 x 6.4cm Colour: Flesh					
999952U4792709	TF2	Tracheostomy Foam dressings for Adults Size: ± 9.0 x 9.8cm Colour: Flesh					

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999952U4792854	TBA1	Tracheostomy Brush - Angled for Tube ID: 6mm Bristle diameter: 7mm Total Length: ± 21cm					
999952U4792711	TBA2	Tracheostomy Brush - Angled for Tube ID: 7/8mm Bristle diameter: 9mm Total Length: ± 21cm					
999952U4792723	TBA3	Tracheostomy Brush - Angled for Tube ID: 9/10mm Bristle diameter: 11mm Total Length: ± 21cm					
999952U4792735	TBS1	Tracheostomy Brush - Straight for Tube ID: 5/6mm Bristle diameter: 7mm Total Length: ± 32cm					
999952U4792747	TBS2	Tracheostomy Brush - Straight for Tube ID: 7/8mm Bristle diameter: 9mm Total Length: ± 34cm					
999952U4792750	TBS3	Tracheostomy Brush - Straight for Tube ID: 9/10mm Bristle diameter: 11mm Total Length: ± 36cm					
999952U4792762	TCS1	Tracheostomy Cleaning Swab					
999952U4792774	SBS1	Stoma Button - Short Size: 6 ID: 6mm OD: 9mm Length: 22mm					

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999952U4792786	SBS2	Stoma Button - Short Size: 7 ID: 7mm OD: 10mm Length: 22mm					
999952U4792800	SBS3	Stoma Button - Short Size: 8 ID: 8mm OD: 11mm Length: 22mm					
999952U4792812	SBS4	Stoma Button - Short Size: 9 ID: 9mm OD: 12mm Length: 22mm					
999952U4792836	SBS5	Stoma Button - Short Size: 10 ID: 10mm OD: 13mm Length: 22mm					
999952U4792848	SBS6	Stoma Button - Short Size: 11 ID: 11mm OD: 14mm Length: 22mm					
999952U4792863	SBL1	Stoma Button - Long Size: 6 ID: 6mm OD: 9mm Length: 30mm					
999952U4792875	SBL2	Stoma Button - Long Size: 7 ID: 7mm OD: 10mm Length: 30mm					
999952U4792887	SBL3	Stoma Button - Long Size: 8 ID: 8mm OD: 11mm Length: 30mm					
999952U4792899	SBL4	Stoma Button - Long Size: 9 ID: 9mm OD: 12mm Length: 30mm					
999952U4792901	SBL5	Stoma Button - Long Size: 10 ID: 10mm OD: 13mm Length: 30mm					

ICN NO:	ITEM:	DESCRIPTION:	Unit Price	Unit Price	Unit Price	Total Price	BRAND NAME				
			Year 1	Year 2	Year 3	Year 1 + year 2 + Year 3					
			(incl. VAT)	(incl. VAT)	(incl. VAT)	(incl. VAT)					
999952U4792913	SBL6	Stoma Button - Long Size: 11 ID:									
		11mm OD: 14mm Length: 30mm									
999952U4792925	TSV1	Hands-Free Tracheostomy Speaking Valve									
999952U4792989	TM1	Tracheostomy masks - Paediatric									
999952U4792991	TM2	Tracheostomy masks - Adult									
999952U4784839	TTG	Tracheostomy Tube Guard									
999952U4784841	TT	Tracheostomy Tape Width: 12.5mm Roll: 1m									
The ar	nnual unit pr	the price that will be used to evaluate the ice will be the applicable (contractual) pri be in accordance with packaging as per	ice per year per	item.							
Required by:			PARTMENT OF H	IEALTH							
At:	VARIOUS INSTITUTIONS										
Country of origi	n										
Brand											
Delivery period (on order) Failure to comply with the above shall invalidate the offer received. Note: All delivery costs must be included in the bid price, for delivery at prescribed destination.											
(Signature of Bi			e of Witness)	Date							