

नेपाल सरकार गृह-मन्त्रालय नेपाल ए.पी.एफ. अस्पताल

ICU Ward को लागी मेडिकल मेशिनरी औजार खरिद गर्न प्रस्ताब आब्हानको सूचना

सूचना प्रकाशित मितिः २०७६/१२/२० गते

COVID-19 व्यवस्थापन गर्न स्थापना हुने ICU Ward को लागी अति आवश्यक विभिन्न मेडिकल मेशिनरी औजार विशेष परिस्थितिमा खरिद गर्नपर्ने भएको हुँदा लिखित दररेट सहितको प्रस्ताव आव्हान गरिएको छ । खरिद गरिने समाग्रीको विवरण, गणस्तर तथा स्पेशिफिकेसन, पालना गर्नपर्ने सर्तहरु उल्लेख भएको बोलपत्र कागजात सशस्त्र प्रहरी बल नेपालको बेबसाईट www.apf.gov.np बाट डाउनलोड गर्न वा यस अस्पतालको प्रवन्ध शाखाबाट नि:शुल्क प्राप्त गर्न सिकने छ । ईच्छक फर्म, कम्पनी, आपूर्तिकर्ताहरुले यस अस्पतालको प्रवन्ध शाखाबाट जारी भएको बोलपत्र कागजातमा उल्लेख भए बमोजिमको बिबरण भरी आधिकारीक सहि छाप सहित मिति २०७६/१२/२३ गते दिनको १२:०० बजे भित्र यस अस्पतालको प्रबन्ध शाखामा शिलवन्दी रुपमा दाखिला वा apfhospital.procurement@gmail.com मा प्रेशित गर्नहन अनुरोध गरिएको छ । दर्ता हुन आएका शिलबन्दी तथा ईमेलबाट प्राप्त प्रस्ताबहरु मिति २०७६/१२/२३ गते दिनको १:०० बजे नेपाल ए.पी.एफ. अस्पताल प्रबन्ध शाखामा खोलिनेछ । यस सम्बन्धि थप जानकारीको लागी यस अस्पतालमा सम्पर्क गर्न हन यो सचना प्रकाशन गरिएको छ ।



Government of Nepal Ministry of Home Affairs Nepal APF Hospital Balambu, Kathmandu, Nepal

BIDDING DOCUMENT

For

Procurement of Medical Machinery Equipment For COVID-19 ICU Ward

Issued on:02 April 2020

Bid Document issued to: All Eligible bidders

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हरिश्चन्द्र अधिकारी सशस्त्र प्रहरी नायव महानिरीक्षक अस्पताल प्रमुख

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Section I. Invitation for Special Contingency Procurement (SCP)

Name of Supplier/Bidder

Address of the Supplier:

Nepal APF Hospital, Procurement Section invites Priced Quotations for the supply and delivery of **Medical Machinery Equipment For COVID-19 ICU Ward** as detailed in attached Specifications and the Schedule of Requirements provided herein.

- 1. The Price Quotation submitted by the Bidder shall comprise the following:
 - a. Quotation and Price Schedules
 - b. Schedule of Requirements
 - c. Technical Specifications
- 2. Priced Quotations must be submitted to the office of Nepal APF Hospital, Procurement Section on or before 12: 00 Hour on Sunday, 05-April-2020.
- 3. The Bidders shall indicate on the Price Schedule the unit prices (where applicable) and total price of the goods to be supplied under the contract. All duties, taxes and other levies payable by the Supplier/Bidder under the contract shall be included in the rates, prices and total Bid Price submitted by the Bidder.
- Price quoted by the Bidder shall remain fixed and valid until completion of the Contract Performance and will not be subject to variation in any account.
- Submitted Priced Quotations must remain valid for a period of 15 days after the deadline for submission date.
- 6. The Bidder shall furnish, as part of its bid, documents establishing the Supplier's/ Bidder's eligibility to bid and qualification to perform the contract if the bid is accepted. Documents to establish such eligibility shall be but not limited to the following:
 - a) Up to date Firm/Company Registration Certificate
 - b) VAT and PAN Registration Certificates
 - c) Tax Clearance Certificate of FY 2075-76
 - d) Power of Attorney
 - e) Product Catalogue/supporting document
- The goods supplied under this contract shall confirm to the Schedule of Requirements and the standards mentioned in the Technical Specification.
- 8. If the last date of submission and opening falls on a government holiday then the next working day shall be considered the last day. The price quotation will be opened on Sunday, 05-April-2020.,13:00 Hour at Nepal APF Hospital, Procurement Section.
- The Purchaser reserves the right to accept or reject the Sealed Quotations without assigning any reason, whatsoever.

Yours sincerely,

Procurement Section
Nepal APF Hospital, Procurement Section, Kathmandu.
Tel-01-4315224, 4313600, email-apfhospital.procurement@gmail.com.



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Section II. Conditions of Contract

1.	Definitions
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- 1.1 In this contract, the following terms shall be interpreted as indicated:
 - a. "The Contract" means the 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;
 - b. "The Contract Price" means the price payable to the Supplier under the contract for the full and proper performance of its contractual obligation;
 - "The Goods" means Equipment and related Accessories and spare-parts or any other materials which the Supplier is required to supply to the Purchaser under the contract;
 - d. "Services" means services ancillary to the supply of the goods such as transportation and insurance including the installation, commissioning and the operational and maintenance training of the supplied equipment.
 - e. "The Purchaser" means the procuring entity purchasing the goods;
 - "The Supplier" means the organization supplying the goods and services under this contract.

2. Technical Specification

2.1 The goods supplied under this contract shall confirm to the standards mentioned in the Technical Specification.

3. Patent Right

3.1 The Supplier shall indemnify the Purchaser against all third-party claims of infringement of patent, trademark or industrial design rights arising from use of goods or any part thereof in the Purchaser's country.

4. Inspection and Tests

4.1 The Purchaser or its Representative shall have the right to inspect and/or test the goods to confirm their conformity to the Technical Specification and the quality of performance after the supply and delivery of good to the Purchaser's premises.

5. Packing

- 5.1 The Supplier shall provide such packing of the goods as is required to prevent their damage or deterioration during transmit to their final destination as indicated in the contract.
- 5.2 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.
- 5.3 The packing, marking and documentation within and outside the packages shall comply strictly with such special requirements as shall be expressly provided in accordance with international standard and practice.

6. Delivery of Goods

6.1 Delivery of the goods shall be made by the Supplier in accordance with the terms specified by the Purchaser in its Schedule of Requirements,

7. Warranty

- 7.1 The Supplier warrants that all the goods supplied under the contract shall fully comply with the specification laid down in the contract.
- 7.2 The warranty shall remain valid for one year after the goods have been delivered to the final destination indicated in the contract, and accepted by the Purchaser after installation and commissioning of

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equipment by the Supplier.

- 7.3 The Purchaser shall promptly notify the Supplier in writing of any claims arising under this warranty.
- 7.4 Upon receipt of such notice, the Supplier shall, with all reasonable speed, replace the defective goods without cost to the Purchaser. The Supplier will be entitled to remove, at its own risk and cost, the defective goods.
- 8. Payment
- 8.1 Payment of the goods supplied shall be made in Nepali Rupees after the delivery and installation and commissioning of goods to the satisfaction of the Purchaser.
- 8.2 Payment shall be made within fifteen (15) days of receipt of the goods and upon submission of claim supported by the acceptance certificate issued by the Purchaser. .
- 9. Prices
- 9.1 Prices charged by the Supplier for goods delivered under the contract shall not vary from the prices quoted by the Supplier in its price quotation.
- 10. Insurance

The Purchaser will be responsible for taking out any appropriate insurance coverage.

- 11. Governing Language
- 11.1 The Governing Language shall be: Nepali or English
- 12. Applicable Law
- 12.1 The applicable law shall be Laws of Nepal.
- 13. Notices
- 13.1Purchaser's address for notice purposes:
 Nepal APF Hospital, Procurement Section, Kathmandu.
 Tel-01-4315224, 4313600, email-apfhospital.procurement@gmail.com.
- 13.2 Supplier's address for notice purposes:
- 14. Taxes and Duties
- 14.1 The Supplier shall be entirely responsible for all taxes, duties, licence fees and other such levies imposed by the GoN.
- 15. Operation, Maintenance and Spare-parts Manuals
- 15.1 The successful Supplier shall supply 2 copies of manufacturer's operation, maintenance and spare-part manuals of the goods (Equipment).
- 16.Conduct of Suppliers
- 16.1 The Supplier shall be responsible to fulfil his obligations as per the requirement of the Contract Agreement, Bidding documents, GoN's Procurement Act and Regulations.
- 16.2 The Supplier shall not carry out or cause to carryout the following acts with an intention to influence the implementation of the procurement process or the procurement agreement:
 - a. give or propose improper inducement directly or indirectly,
 - b. distortion or misrepresentation of facts
 - c. engaging or being involved in corrupt or fraudulent practice
 - d. interference in participation of other prospective bidders.
 - coercion or threatening directly or indirectly to impair or harm, any party or the property of the party involved in the procurement proceedings,
 - f. collusive practice among bidders before or after submission of bids for distribution of works among bidders or fixing



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artificial/uncompetitive bid price with an intention to deprive united the Purchaser the benefit of appen competitive bid price... contacting the Purchaser with an intention to influence the

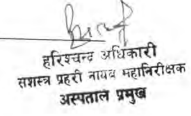
g. contacting the Purchaser with an intention to influence the Purchaser with regards to the bid or interference of any kind in examination and evaluation of the bids during the period after opening of bids up to the notification of award of contract

17.Blacklisting Supplier

- 17.1 The GoN, Public Procurement and Monitoring Office(PPMO) may blacklist a Supplier for his conduct up to three years on the following grounds and seriousness of the act committed by the supplier:
 - a. if it is proved that the supplier committed acts pursuant to the Sub clause 16.2,
 - b. if it is proved later that the supplier had committed substantial defect in implementation of the contract or had not substantially fulfilled his obligations under the contract or the completed work is not of the specified quality as per the contract,
 - if convicted by a court of law in a criminal offence which disqualifies the supplier from participating in the contract.
- 17.2 A Supplier declared blacklisted and ineligible by the GON shall be ineligible to bid for a contract during the period of time determined by PPMO.

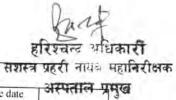
18. Dispute Resolution

18.1 Any dispute arising out of the Contract, which cannot be amicably settled between the parties, shall be referred to adjudication.



Section III. Schedule of Requirements and the state of th

S.N.	Description	Quantity	Place of Delivery	- The series	Supplier
1	. Intensive Care Bed	20 set		As soon as possible from the date of contract sign	
2	. IV Stand	40 set		As soon as possible from the date of contract sign	
3.	Defibrillator	1 set		As soon as possible from the date of contract sign	
4.	Portable x-ray	1 set		As soon as possible from the date of contract sign	
5.	Air mattress	20 pc		As soon as possible from the date of contract sign	
6.	Patient Trolley	2 set		As soon as possible from the date of contract sign	
7.	Wheel chair	4 set		As soon as possible from the date of contract sign	
8.	Bi pap machine	2 set	npu	As soon as possible from the date of contract sign	
9.	Nebulizer	8 set	eathm2	As soon as possible from the date of contract sign	
10	. Suction Machine	8 set	tion, k	As soon as possible from the date of contract sign	
11.	X ray view box	4 set	Sec	As soon as possible from the date	-
12.	ECG machine	1 set	ient	of contract sign As soon as possible from the date	
13.	Bain circuit	4 set	curen	of contract sign As soon as possible from the date of contract sign	
14.	T piece	8 set	al, Pro	As soon as possible from the date of contract sign	
15.	Laryngoscope Set	2 set	Hospit	As soon as possible from the date of contract sign	
16.	Tracheostomy tube Different size	1 set	Nepal APF Hospital, Procurement Section, Kathmandu	As soon as possible from the date of contract sign	
17.	ET tube different size with subglottic suction plug Port	30 set	Nepa	As soon as possible from the date of contract sign	
18.	Pulse oximeter	3 set		As soon as possible from the date of contract sign	
19.	Stethoscope	20 set		As soon as possible from the date of contract sign	
20.	BP apparatus	4 set		As soon as possible from the date of contract sign	
21.	Electric needle destroyer	2 set		As soon as possible from the date of contract sign	
22.	Autoclave 800 Ltr.	1 set		As soon as possible from the date of contract sign	
23.	Electronic weighing machine	1 set		As soon as possible from the date of contract sign	
24.	AMBU bag with reservoir bag	4 set		As soon as possible from the date	



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2	25. hemodialysis unit	1 set	रतान्। अतान्।	As soon as possible from the date of contract sign	
2	26. Resuscitation cart	2 set		As soon as possible from the date of contract sign	
2	7. Closed Suction catheter	30 set		As soon as possible from the date of contract sign	
2	8. Glucometer With Strips	2 set		As soon as possible from the date of contract sign	
2	9. Noninvasive Ventilator Mask	12 set		As soon as possible from the date of contract sign	
30	0. Oropharyngeal airway Different Sizes	30 set	npu	As soon as possible from the date of contract sign	
3	1. nasopharyngeal airway	10 set	Nepal APF Hospital, Procurement Section, Kathmandu	As soon as possible from the date of contract sign	
32	2. Venturi Mask (Different) Fio2	20 set	tion, K	As soon as possible from the date of contract sign	
33	Ventilating Face Mask (Anesthetic Mask) Different Size	10 set	ant Sec	As soon as possible from the date of contract sign	
34	. Refrigerator 185 Ltr	2 set	cureme	As soon as possible from the date of contract sign	
35	. Blood warmer / Fluid Warmer	5 set	d, Proc	As soon as possible from the date of contract sign	
36	. Physiotherapy Chest Vibrator	2 set	lospita	As soon as possible from the date of contract sign	
37.	Intermittent Pneumatic Compression Device	2 set	APF	as soon as possible from the date f contract sign	
38.	Forced Air Warmer Device (patient warmer device)	3 set	Nepal	s soon as possible from the date f contract sign	
39.	Bedside Locker	20 set	A	s soon as possible from the date f contract sign	-
40.	Mayo Table	20 set	A	s soon as possible from the date contract sign	
41.	infrared thermometer (Non Touch)	5 pc	As	s soon as possible from the date contract sign	_
42.	Gum elastic Bongie	2 set	As	s soon as possible from the date contract sign	
43.	Intubatins Stylets	5 set		soon as possible from the date contract sign	



Section IV. Technical Specifications

Bidders must enter their offered specifications against each parameter of this Technical Specifications अस्पताल प्रमुख Form (TSF), comment as necessary, and sign and stamp each page. Failure to complete this statement of compliance may result in the offer being rejected.

The Statement of Compliance must be substantiated with authenticated document /datasheet/manual of the relevant parameters indicated.

The right hand blank side must be completed by the bidder with the technical specifications of the offered product with supplementary documents enclosed.



S.N.		Specifications	Bidder's Proposed Specification	Deviation If Any	Page no in catalog/ Data sheet
	Motorised ICU beds			1	Sheet
	Manufacturer				
	Brand				-
	Type/Model				
	Country of Origin			+	
1	Description of Func	tion			
1.1	Motorised ICU bed is designed for hospitali ICU,CCU or recovery of patient ease. These	a bed specially zed patients in unit who are in need beds have special omfort and well being			
2	Operational Require	ments			-
2.1	It shall have anti-corre treated baked hard epo four sections Monitori	osive and antirust oxy powder coating, sed ICU bed.			
3	System Configuration				
3.1	Monitorised ICU bed, mattress.	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			
4	Technical Specification	ons			
4.1	Dimensions approx.: 2 820H mm (without ma	220Lx995Wx450H-			
4.2	Bed frame shall be ma approx. 50mm x 25mm tubes with proper supp be fitted on the base fra approx. 60X30X1.6mm various supporting link	inly made from 1 x 2mm thick ERW 1 ort. This frame should 2 mainly made of 2 ERW tube on			
1.3	The base frame shall be mm dis non-rusting two Central locking mechan	in wheel castor with			
1.4	Four sections Polyprop fitted on four section to perforated design for ea mattress.	ylene detachable top p bed frame with			
.5	Back rest and leg rest b detachable mattress gua section respectively)	oth shall have ards. (3 nos on each			
.6	Back rest knee rest and approx. (450mm -770m by electromechanical achandset, and additional handset, for operating a function and shall have trendelenburg/reverse trositions.	m) position operated ljustment through nurses' control and locking of above			
7	Simultaneous electrome	chanical adjustment			

		7 70		
S.N.	Purchaser's Specifications	Bidder's Proposed Specification	Deviation If Any	Page no in catalog/ Data sheet
	of back rest and knee rest on both hand set and the additional nurse's hand set.			SHEEK
4.8	One touch key for flattening of the bed at the lowest height for CPR on nurses' hand set.			
4.9	Battery backup with inbuilt battery charger shall be provided.			
4.10	The hand set and nurses' hand set shall have indications for power on and the battery charge.			
4.11	Manual pull lever on both side of bed to quickly bring bed to a flat position.			
4.12	Bed shall have split type swing down railing, two on each side (Head and middle section) made from polymer moulded material.			
4.13	Railings shall avoid any finger and neck entrapments.		1	
4.14	Detachable head/foot board.			
4.15	Bed frame must be sturdy and stable to support weight of at least 250 kg.			
4.16	High quality stainless steel outer covering tube with a knob to mount Syringe pump.			
4.17	There must be suitable buffer mechanism to avoid heating of the bed to the wall.			
4.18	It must have provision of fixing suitable rod for hanging intravenous/irrigation fluid bottle on both sides at head end and foot end.			
4.19	It must have hook on bed frame on both sides for holding urine/ drainage bag (at least 4 nos.)			
4.20	Shall provide with one dual hook 304-grade stainless steel telescopic IV rod.			
4.21	Mattress: Shall provide with one four section mattress of dimensions approx. 85X195 cm with washable cover of good quality. The mattress must be made of high density PU foam of 12cm thickness.			
5	Electrical specification		4	
5.1	Nominal 230V AC			
5.2	Switch mode power supply operating range 90V-300V; 47/63Hz; Max 2A.			
5.3	Should comply with international standards for electric shock protection and liquid ingress protection.			
6	System Configuration Accessories, spares and consumables			
6.1	All standard accessories/consumables/parts required for the proper operation of the above item shall be included in the offer.			9

72.50		Ridder's	mr. 31	
S.N.	Purchaser's Specifications	Bidder's Proposed Specification	Deviation If	Page no in catalog/ Data sheet
	Bidders shall specify, in a separate Excel worksheet, the quantity and details of any items included in this offer, which have not been specified in this Technical Specifications Form.			
6.2	At least 12mm diameter stainless steel SS304 telescopic heavy duty IV rod with two hooks with provision to park when not in use.			
6.3	Urine bag holder - one			
6.4	File holder - one			
7	Operating Environment			
7.1	The system offered shall be designed to store and to operate normally under the conditions of the purchaser's country. The conditions include Climate, Temperature, Humidity, etc.			
8	Standards and Safety Requirements			
8.1	Must submit ISO 9001 or ISO 13485:2003/AC:2007 AND			
8.2	CE or USFDA approved product certificate.			
9	User Training			
9.1	User and maintenance training should be provided to the hospital personnel at the time of installation and any time as per requirement for 5 years.			
10	Warranty			
10.1	Warranty for 2 years after acceptance.	1 C-C		
11	Maintenance Service During Warranty Period			
11.1	Standard warranty conditions are applicable.			
12	Installation and Commissioning			
12.1	Must supply preassembled unit, ready to use.			
13	Documentation			
13.1	Users/Instructions manual shall be provided in English.			



Technical Specification of IV Stand, Four Hooks

S.N.	Purchaser's Specifications	Bidder's Offer
	IV Stand, Four Hooks	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1,1	This IV/saline stand is used for hanging various intravenous	
	items such as blood bag, glucose bottle etc.	
2	Operational Requirements	
2.1	Mobile IV stand on castors with adjustable height.	
3	System Configuration	
3.1	Adjustable IV/saline stand with four hooks and swivels castors.	
4	Technical Specifications	
4.1	Materials:	
4.2	Base: Heavy base on antistatic swivel castors of approx. diameter Ø50mm.	
4.3	Support column: solid mechanism to which the upper pole is fixed; the pole has an adjustable height	
4.4	Hook: Stainless steel 4 hooks welded together on the top of the serum rod.	
4.5	Load capacity:approx.12kg (3kg per hook)	
5	Accessories, spares and consumables	
5.1	Not applicable.	
6	Operating Environment	
6.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Climate, Temperature, Humidity, etc.	
7	Standards and Safety Requirements	
7.1	Must submit ISO 9001 or ISO 13485:2003/AC:2007 AND	
8	User Training	
8.1	Not applicable.	
9	Warranty	
9.1	Warranty for 1 year.	
10	Maintenance Service During Warranty Period	
10.1	Standard warranty conditions are applicable.	
11	Installation and Commissioning	
11.1	Must supply preassembled unit, ready to use.	
12	Documentation	0
12.1	User /instructions manual in English.	/ D



S.N.	o operations	Bidder's Offer/ Statement of Compliance	Page no. of catalogue/ datasheet/ manual
	Automated External Defibrillator (AED)		manual
	Manufacturer		
	Brand		
	Type/Model		
	Country of Origin		
1	Description of Functions		
1.1	Defibrillator to be used to give electrical shocks to the patient's chest assisting theheart to resume its co-ordinated atria-ventricular pump function, in thecontext of advanced cardiac life support.		
2	Operational Requirements		
2.1	It shall operate on internal replaceable batteries.		
3	System Configurations		
3.1	Automated External Defibrillator (AED) with complete accessories, for adult paediatric and infant use.		
4	Technical Specifications		
4.1	It shall be portable Automated External Defibrillator (AED) for immediate operation, self-explanatory and based on intuitivelyunderstood design features.		
4.2	Shock and splash resistant housing allows functioning in demanding environment.		
4.3	Shall perform self-test when device is switched on and shall indicate ready for use. Self-test is performed upon each switched on ready-for-use is indicated		
4.4	It shall have capability of automated assessment and analysis, adequately sensitive and specific forchildren and adults.		
1.5	The device shall have facility of step-by-step guidance from the large pictograms when it is on		
1.6	It shall analyse, shock with self-adhesive external pads, colour coded, each with pictogram.		
.7	It shall have automated direct defibrillation with biphasic waveform, máximum energy approximately 150J.		
.8	It shall have built-in load compensation algorithm adjusts energy delivery according patient's impedance.		b



S.N	and the second s	Bidder's Offer/ Statement of Compliance	Page no. of catalogue/ datasheet/
4.9	8 year or > 25kg) and adults.		manual
4.10	For infants (> 1 year or > 6kg) shall come with attenuation pads, reduction to maximum approximately 50J.		
4.11	It shall have pads with plug and power cord, length approximately 100cm		
4.12	It shall have built-in audible metronome assists Cardiac Pulmonary Resuscitation (CPR)reports, with audio-visual alerts of operational status, malfunctions(electrodes) and low battery status.		
4.13	Facility of internal safety discharge of accumulated energy upon 20sec non-delivery,switch-off or malfunction		
4.14	Battery capacity approximately 100 shocks of 250J.		
5	Accessories, Spare Parts and Consumables		
	 1 x Set of children, adult self-adhesive external pads, colour coded, with pictogram 1 x Set of infant attenuated adhesive external pads, colour coded, with pictogram 1 x CD-ROM with training material 2 x Set of spare batteries 9 V PP3 / 6LR61 (separately packed) or long life lithium manganese oxide battery. 1 x Carry case with storage pocket for leads and other accessories 		
5.2	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified above).		
6	Operating Environment		
.1	The system offered shall be designed to operate normally under the conditions of the purchaser's country. The conditions include Power Supply,		

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S.N	permeations	Bidder's Offer/ Statement of Compliance	Page no. of catalogue/ datasheet/
	Climate, Temperature, Humidity, etc.		manual
6.2	Power supply: It shall operate on internal replaceable batteries, type 9V PP3 / 6LR61 or M5070A type lithium long life battery.		
7	Standards & Safety Requirements		
7.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND		
7.2	CE (93/42 EEC Directives) or USFDA approved product certificate.		
7.3	Comply to AHA & ACLS requirements or shallmeet AAMI DF80 guidelines and AHA recommendations for adult defibrillation (Circulation 1997; 95:1677-1682).		
8	User Training		
8.1	The Supplier shall conduct user training for this equipment to enable operators to use the equipment properly. The training shall include the use of all operational functions of the equipment, as well as routine checks and maintenance expected by users.		
9	Warranty		
9.1	Comprehensive warranty for 2 years after acceptance.		
10	Maintenance Service During Warranty Period		
0.1	During the warranty period supplier must ensure planned preventive maintenance (PPM) along with corrective/breakdown maintenance whenever required.		
11	Installation and Commissioning		
1,1	The bidder must arrange for the equipment to be installed and commissioned by certified or qualified personnel; any prerequisites for installation to be communicated to the purchaser in advance, in detail.		
2	Documentation		
2.1	User (Operating) manual in English		
2.2	Service (Technical / Maintenance) manual in English	n	k



S.N.	Purchaser's Specifications	Bidder's Offer/ Statement of Compliance	Page no. of catalogue/ datasheet/ manual
12.3	List of important spare parts and accessories with their part numbers and costing.		manuai
12.4	Certificate of calibration and inspection from factory.		

N AT	10. Portable X-ray	Bridger Wifer
.N.	Purchaser's Specifications	Bud all the state of the state
	X-Ray Machine Mobile, 6KW or More	
	Manufacturer	
	Brand	
	Type/Model	
	Country Of Origin	
1 ==	Description of Function	
1.1	Mobile X-Ray Unit is required to perform X-Ray studies in Emergency and trauma departments and at bedside in wards and ICU.	
2	Operational requirements	
2.1	Compact, lightweight, easily transportable mobile radiographic unit suitable for bedside X-ray for trauma units (accidental cases), intensive care units, operation theatres and also in the Radiology department for conventional radiography.	
3	System Configuration	
3.1	X-ray Machine Mobile, 6KW or more complete unit and with complete accessories.	
4	Technical Specifications	
4.1	The Generator:	
	 Microprocessor-controlled high frequency generator. Max output: not less than 6kW at 125kv, 100ms Voltage range: 40 - 125kV in more than 25 steps. Max tube current: 250mA mAs range: 0.5 - 200mAs in more than 30 steps Minimum Exposure time: not more than 5ms Soft touch key operations Anatomical Programmable Radiographic mode shall be available. 	
4.2	 X-Ray Tube: Rotating anode type Anode rotation: 2800rpm Anode heat capacity: not less than 100 kHU Dual focal spot: not more than 0.8mm 	
4.3	Collimator:	
4.3	 Manually adjustable multi-leaf collimator rotatable ±90° Collimator light halogen lamp: 180 lux at 1m SID 	
4.4	Tube positioning:	
	 Max tube height: not less than 1800mm, Min tube height: not more than 450 mm Max horizontal extension: not less than 	0 >

S.N.	Purchaser's Specifications	Bidder's Offer
J. 1.	800mm	SURANIE PAR
1.5	The unit shall have counter balanced arm system	3. 0194/A
4.6	Shall have remote control of exposure to protect	
1.0	operator.	
4.7	It shall have cassette compartment of holding	
7.7	about 8 pieces of 35x43cm cassettes.	
4.8	The unit must have an effective braking system	
4.0	for parking, transport and emergency braking.	
4.9	The unit shall come with overload protection	
4.9	device.	
5	Accessories, spares and consumables	
4.1	Accessories:	
4.1	 Lead apron lightweight- 1 nos. Grid(Ratio 6:1) of 12"x15" and 10"x12": 01 each. Remote control kit: 01 no. 	
5.2	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified	
	above).	
6	Operating Environment	
6.1	The system offered shall be designed to operate	
	normally under the conditions of the purchaser's country. The conditions include Power Supply, Climate, Temperature, Humidity, etc.	
6.2	Power supply: 220 - 240 VAC, 50Hz fitted with	
	5m automatic retractable power cable for easy connection to any wall outlet with protective ground conductor.	
7	Standards and Safety Requirements	
7.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND	
7.2	CE (93/42 EEC Directives) or USFDA approved product certificate.	
7.3	Shall meet:	
	 IEC 60601-1-3 - Part 1: General Requirements for safety - Collateral Standard: General Requirements for Radiation Protection in Diagnostic X-Ray Equipment. IEC 60601-2-7 - Part 2-7: Particular Requirements for the Safety of High- Voltage Generators of Diagnostic X-Ray Generators. 	
8	User Training	
8.1	Must provide user training (including how to use and maintain the equipment).	
9	Warranty	
9.1	Comprehensive warranty for 2 years	

S.N.	Purchaser's Specifications	Bidder's Offers
4	afteracceptance.	.(अन् कारमाव
10	Maintenance Service During Warranty Period	
10.1	During the warranty period supplier must ensure preventive maintenance and corrective/breakdown maintenance whenever required.	
11	Installation and Commissioning	
11.1	The bidder must arrange for the equipment to be installed and commissioned by certified or qualified personnel; any prerequisites for installation to be communicated to the purchaser in advance, in detail.	
12	Documentation	
12.1	User (Operating) manual in English.	
12.2	Service (Technical / Maintenance) manual in English.	
12.3	List of important spare parts and accessories with their part number and costing.	
12.4	Certificate of calibration and inspection from factory.	0.



S.N.	Purchaser's Specifications	Bidder's Offers
	Air Mattress with electric air pump	
- 1	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1.1	The <i>medical air mattresses with airpump</i> is useful in preventing or treating serious ailments related to extended bed rest, such as pressure sores and skin shearing.	
2	Operational Requirements	
2.1	Inflatable air mattress with electric air pump	
3	System Configuration	
3.1	Inflatable air mattress with electric pump with complete accessories.	
4	Technical Specifications	
4.1	Should be made of medical grade PVC material, comfortable and skin friendly	
4.2	Should support weight up to 100kg or more.	
4.3	Pump should have low noise	
5	Accessories, spares and consumables	
5.1	High quality electric air pump	
6	Operating Environment	
6.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Climate, Temperature, Humidity, etc.	
7	Standards and Safety Requirements	
7.1	Must submit ISO 9001 or ISO 13485:2003/AC:2007 AND	
8	User Training	
8.1	The Supplier shall conduct user training for this equipment to enable operators to use the equipment properly.	
9	Warranty	
9.1	Warranty for 1 year.	
10	Maintenance Service During Warranty Period	
10.1	Standard warranty conditions are applicable.	
11	Installation and Commissioning	
11.1	Supplier must accomplish proper installation &commissioning of equipment	
12	Documentation	
12.1	User /instructions manual in English.	0-1



S.N.	Purchaser's Specifications	Bidder's Offers
5.11.	Patient Trolley	Didder 5 Offers
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1.1	Patient Trolley is required for Patient transfer to &	
414	froICU/OT/Emergency.	
2	Operational Requirements	
2.1	Patient trolley with pneumatic adjustment for back section & hydraulic adjustment for height.	
3	System Configuration	
3.1	Patient trolley with mattress and with complete accessories.	
4	Technical Specifications	
4.1	Must have three sectional mattress base made of X Ray translucent high pressure laminate with facility to insert X Ray Cassette from either sides & ends of the trolley.	
4.2	Must be able to X Ray the patient from positions along the entire length and width of the trolley.	
4.3	Must have pneumatic step less adjustment for back section, Trendelenburg, reverse Trendelenburg and foot section.	
4.4	Must have hydraulic height adjustment with a foot paddle on either side of the trolley	
4.5	Frame must be made up of epoxy powder coated steel	
4.6	Must have Central braking system with steering facility	
4.7	Must be equipped with 360 deg. swivelling heavy duty castors diameter 150 mm.	
4.8	Must have bumpers at all the four corners of the trolley	
4.9	Must have facility to fix IV rod at all the four corners and middle of mattress base frame.	
4.10	Must have place for fixing 'B' Type Oxygen Cylinder	
4.11	 Dimensions, Approx. ±10%: Max. Length: 2000-2100 mm Max. Width: 730-750 mm Height: 535 – 905 mm Trendelenburg: 14-20° step less Anti Trendelenburg: 7-10° step less X ray viewing area: entire length 	
5	Accessories, spares and consumables	
5.1	Accessories: Anti-static Hygienic, washable Mattress (80mm thick) with pull straps, 01 pc Collapsible Side Rails, 01 pair Stainless steel I.V. Rod 01 pc Cylinder Holder for 'B' Type Oxygen Cylinder.01 pc	
5.2	All standard accessories/consumables/parts required for the proper operation of the above item shall be included in the offer. Bidders shall specify, in a separate Excel worksheet, the quantity and details of any items included in this offer which have not been specified in this	0

Purchaser's Specifications Technical Specifications Form. Operating Environment The system offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The	
The system offered shall be designed to be stored and to operate	
The system offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The	
conditions include Climate, Temperature, Humidity, etc.	
Standards and Safety Requirements	
Must submit ISO13485:2003/AC:2007 for Medical Devices AND	
User Training	
Must provide user training (including how to use and maintain the equipment).	
Warranty	
Comprehensive warranty for 1 year after acceptance.	
Maintenance Service During Warranty Period	
During the warranty period supplier must ensure corrective/breakdown maintenance whenever required.	
Installation and Commissioning	
The supplier must accomplish proper commissioning of the equipment on site.	
Documentation	
User (Operating) manual in English	
Service (Technical / Maintenance) manual in English	
and agating	
NUMERICALITY	Must submit ISO13485:2003/AC:2007 for Medical Devices AND User Training Must provide user training (including how to use and maintain the equipment). Warranty Comprehensive warranty for 1 year after acceptance. Maintenance Service During Warranty Period During the warranty period supplier must ensure corrective/breakdown maintenance whenever required. Installation and Commissioning The supplier must accomplish proper commissioning of the equipment on site. Documentation User (Operating) manual in English Service (Technical / Maintenance) manual in English List of important spare parts and accessories with their part number



S.N.	Purchaser's Specifications	Bidder's Offer	aidders offe
	Wheel Chair (foldable)	- V -	
	Manufacturer	1	
-	Brand		
	Type / Model		
	Country of Origin		
	Description of Function		
1.1	Basic wheelchair for transportation of patients who are unable to stand/walk.		
2	Operational Requirements		
2.1	Basic foldable wheelchair for adult use.		
3	System Configuration		
3.1	Wheel Chair (foldable).		
1	Technical Specifications		
4.1	Heavy carriage mounted on 4 ball-bearing wheels.		
4.2	Front wheels free rolling, 360 degrees swivel.		
4.3	Both rear wheels with brake.		
4.4	Foot lever, integrated in frame, facilitates tilting the wheelchair.		
4.5	Two handles at the rear fit with plastic rims.	F =	
4.6	Swing-away foot and arm supports for easy stepping on/off.		
1.7	Armrests seat and back are upholstered.		
4.8	Materials:		
	 High resistance to corrosion (tropical environment). Frame: Chrome-plated tubular steel. Upholstery: Plastic, flexible highly tear resistant, anti-static, flame retardant, disinfectant- and liquid proof, washable. Tires: Heavy duty solid rubber. 		
4.9	 Dimensions, Approx. ± 10%: Overall: 450 x 500 x 870mm (d x w x h). Back support: 500 x 400mm (w x h). Frame, diameter: 23mm. Wheels, diameter: Front 200mm, Rear 600mm. Carrying capacity: Approximately 150kg. 		
5	Accessories, spares and consumables		
5.1	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication		
	materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified above).		
6	Operating Environment		
6.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include		

S.N.	Purchaser's Specifications ए.पा.ए	Bidder's Offer	Bidders Offers
	Climate, Temperature, Humidity, etc.	Diagers offer	
7	Standards and Safety Requirements		
7.1	Must submit ISO 9001 or ISO 13485:2003/AC:2007 AND		
8	User Training		
8.1	Not applicable.		
9	Warranty		
9.1	Comprehensive warranty for 1 year.		
10	Maintenance Service During Warranty Period		
10.1	Standard warranty conditions are applicable.		
11	Installation and Commissioning		
11.1	Must supply preassembled unit, ready to use.		
12	Documentation		
12.1	User's manual shall be supplied in English.		

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Purchaser's Specifications	Bidder's Offer
BIPAP (Bi-level Positive Airway Pressure)	
in waveform or har scale format the measured and calculated	
DIDAD (Ri-level Positive Airway Pressure), complete unit with all	
B 110 110 110 110 110 110 110 110 110 11	
Mashina shall be besed on the colenoid valve technology and shall	
offer preferably auto track sensitivity and adjustable rise time	
P. A.	
Accessories, spares and consumables	
All standard accessories, consumables and parts required to operate	
the equipment, including all standard tools and cleaning and	
lubrication materials, to be included in the offer. Bidders must	
Operating Environment	
The product offered shall be designed to be stored and to operate	
normally under the conditions of the purchaser's country. The	
etc.	
Power supply: 220 – 240 VAC, 50Hz fitted with appropriate plug.	
The power cable must be at least 3 metre in length.	
Standards and Safety Requirements	
User Training	
equipment).	
Warranty	
Comprehensive warranty for 1 years after acceptance.	
Maintenance Service During Warranty Period	
During warranty period supplier must ensure corrective/breakdown	
maintenance whenever required.	
Installation and Commissioning	
	Manufacturer Brand Type / Model Country of Origin Description of Function

S.N.	Purchaser's Specifications 2.47.94	Bidder's Offer
	onsite.	
12	Documentation	
12.1	User (Operating) manual in English.	
12.2	Service (Technical / Maintenance) manual in English.	
12.3	List of important spare parts and accessories with their part numbers and costing.	



S.N.	Purchaser's Specifications		Bidder's Offer
	Nebuliser		
	Manufacturer		
	Brand		
	Type / Model		
	Country of Origin		
1	Description of Function		
1.1	Nebuliser is a device use	d to administer medication to people in forms of a	
1.1	liquid mist to the airways. It is commonly used in treating cystic fibrosis,		
	asthma, and other respira		
2	Operational Requireme		
2.1	Heavy duty compact Nel	puliser is required.	
3.	System Configuration	ALION TO TO THE MAN TO THE	
3.1	Nebuliser complete unit	with all standard accessories.	
4	Technical Specification		
4.1	Compact, lightweight, lo		
4.2	Durable longlife compre	ssor. Suitable for heavy duty/ institutional (hospital)	
7.2	use must be able to run	ininterruptedly for min one hour.	
4.3	Maximum pressure: 2.0		
4.4	Must produce particle of		
4.5	Aluminium cabinet paint		
4.6	Piston-type electric aspir	ator that offers high performance and great	
4.0	durability.	ator that offere mg. parent	
4.7	Protective thermal cut out relay.		
4.8	Air delivery rate approx.151/min.		
4.9	24 hours continuous wor		
5	Accessories, spares and		
5.1	Accessories:	Consumatives	
3.1		eusable, autoclaveable- 01 no.	
		ace mask reusable, autoclaveable- 02 each.	
	• 1 x 200 cm. tubi	iece, Nosepiece, reusable, autoclaveable- 01 each.	
<i>c</i> 1	Spare filters- 10 nos. All standard accessories, consumables and parts required to operate the		
5.1	equipment, including all		
	materials, to be included		
	every item included in th	eir offer (including items not specified above).	
6	Operating Environmen	t	
6.1	The system offered shall	be designed to store and to operate normally under	
0.1	the conditions of the pur	chaser's country. The conditions include Power	
	Supply, Climate, Tempe	rature, Humidity, etc.	
6.2	Power supply: 220 - 2	40 VAC, 50Hz fitted with appropriate plug. The	
J.2	power cable must be at l	east 3 metre in length.	
7	Standards and Safety I	Requirements	
7.1	Must submit ISO13485:	2003/AC:2007 for Medical Devices AND	
8	User Training		
8.1	Must provide user traini	ng (including how to use and maintain the	
3.1	equipment).		
9	Warranty		0

	The Residence	
S.N.	Purchaser's Specifications	Bidder's Offer
9.1	Comprehensive warranty for 1 year after acceptance.	
10	Maintenance Service during Warranty Period	
10.1	During warranty period supplier must ensure corrective/breakdown maintenance whenever required.	
11	Installation and Commissioning	
11.1	Supplier must accomplish proper commissioning of equipment onsite.	
12	Documentation	
12.1	User (Operating) manual in English.	
12.2	Service (Technical / Maintenance) manual in English.	
12.3	List of important spare parts and accessories with their part number and costing.	



16. Suction Machine

S.N.	Purchaser's Specifications	Bidder's Remarks
	Electric Suction Pump	
	Manufacturer	
	Brand	
	Type / Model	11
	Country of Origin	
1	Description of Function	
1.1	To extract fluid from the body during surgery or emergency treatments.	
2	Operational Requirements	
2.1	An electric double jar suction pump for surgical use.	
3	System Configuration	
3.1	Suction machine with two bottles and accessories.	
4	Technical Specifications	
4.1	It shall be mounted on four robust, fully 360 degree swivelling, antistatic castors, with at least 2 diagonal brakes.	
4.2	Come with suction controller and vacuum gauge / indicator.	
4.3	The pump shall be oil free vacuum pump where the pumped liqu shall be sealed off from the pump.	nid
4.4	Come with overflow control valves. Bidder shall provide technical design and details of the pump withis TSF	th
4.5	Vacuum rate shall be from 0 to not less than 640 mmHg (0.85 bars).	
4.6	Air flow rate shall be at least 25 l/min.	
4.7	The pump shall come fitted with twin unbreakable, transparent, autoclaveable polycarbonate suction bottles minimum 2 litre eac	sh.
4.8	The bottles shall be incorporated with an automatic suction cut-comechanism when they become full.	
4.9	The suction bottles shall come with overflow lid.	
4.10	Noise level: not more than 55 dBA.	
4.11	Air discharge from pump shall be filtered by a 0.3 micron bacter hydrophobic filter.	rial
5	Accessories, spares and consumables	
5.1	Accessories:	sily.
6	Operating Environment	* 1
6.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Power Supply, Climate, Temperature, Humid etc.	
6.2	Must operate on 220-240V AC as well as rechargeable batteries.	
7	Standards and Safety Requirements	
7.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AN	D n

	Durchason's Specifications Did Biddon's Demarks		
S.N.	Purchaser's Specifications		Bidder's Remarks
8	User Training	क्लान कार्या	
8.1	Supplier must provide user training regarding how to equipment.	use the	
9	Warranty		
9.1	Warranty for 1year.		
10	Maintenance Service During Warranty Period		
10.1	During warranty period supplier must ensure corrective maintenance whenever required.	/e/breakdown	
11	Installation, Inspections and Commissioning		
11.1	Must supply preassembled unit, ready to use.		
12	Documentation		
12.1	User (Operating) and Service (Technical/Maintenance be supplied in English.	e) manuals to	
12.3	List of important spare parts and accessories with thei numbers and costing	r part	

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S.N.	Purchaser's Specifications	Bidder's Offer
-	X-ray view box double	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	_
1.1	X-ray view box (double) for viewing MRI, CT and x-ray images	
	in films.	
2	Operational Requirements	
2.1	X-ray view box, complete set	
3	System Configuration	
3.1	X-ray view box, complete set	
4	Technical Specifications	
4.1	Dimension:	
	Frame - approx880*503*29mm	
	Viewing area— approx.736*440mm	
4.2	Should have LED light source	
4.3	Power consumption should be 30watt	
4.4	Adjustable brightness with more than 10000 lux	
4.5	Weight should not be more than 8kg	
4.6	Should have clips for holding on wall	
5	Accessories, spares and consumables	
5.1	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified above).	
6	Operating Environment	
6.1	The system offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Power Supply, Climate, Temperature, Humidity, etc.	
6.2	Power supply: 220 – 240 VAC, 50-60Hz	
7	Standards and Safety Requirements	
7.1	Must submit ISO 9001 or ISO 13485;2003/AC: 2009.	
8	User Training	
8.1	Must provide user training (including how to use and maintain the equipment).	
9	Warranty	
9.1	Comprehensive warranty for lyear.	
10	Installation and Commissioning	
10.1	Supplier must accomplish proper commissioning of the equipment on site.	
11	Documentation	
11.1	User (Operating) / service manual in English	



18. ECG Machine

S.N.	Purchaser's Specifications	Bidder's Offer
	ECG Machine, (3 Channel)	
	Manufacturer	
	Brand	
	Type/Model	
	Country of Origin	
	Manufacturer's Authorization Letter	
1	Description Of Function	
1.1	ECG Machine is primary equipment to record ECG Signal	
	in various configurations.	1
2	Operational Requirements	
2.1	Portable digital ECG machine must be able to acquire	
	Simultaneous 3 channel ECG recording with 12	
	lead simultaneous acquisition with auto summary	
3	System Configuration	
3.1	Portable digital ECG machine with complete accessories	
4	Technical Specifications	
4.1	Simultaneous acquisition of up to 12 leads	
4.2	Should have TFT Color LCD Display	
4.3	Display to Preview signal quality prior to printing thereby	
	saving time and paper	
4.4	Should have patient data entry feature	2
4.5	Should have colour coded keys for ease of operation	
4.6	Should have different mode of printing: Automatic, Manual	
4.7	Should have different sensitivity levels: 2.5,5,10,20,40	
7.1	mm/mV Auto	
4.8	Recording speeds of 5,10, 25 and 50 mm/sec	
4.9	Should have a high frequency Response: 0.05 Hz to 150	
4.2	Hz	
4.10	Should have a sampling frequency of 1000 Hz	
4.11	User selectable filter: AC Filter, EMG filter- 25 or 35,	
75.5	Base Line Filter	
4.12	Printer must compatible with Roll ECG paper	
4.13	Light weight – Less than 1.6 Kg	
4.14	Battery operation – Lithium Ion Battery -minimum 10	
1.4	HRS continuous work.	
4.16	Easy to carry handle	
4.17	Automatic measurement and interpretations of ECGdata.	
4.19	Should have pacemaker detection facility	
4.20	PC interface facility and optional PC interface software	
	(Optional ECG data transferfeature)	
4.21	External storing and retrieving facility through USB	
	storage device	A
5	Accessories, spares and consumables	hat
5.1	Accessories:	हरिश्चन्द्र अधिकारी



S.N.	Purchaser's Specifications	Bidder's Offer
	Power Cable – 1 no;	
	Lead Patient Cable – 1set;	
	Chest Electrodes – 20 nos;	
	Clip-onelectrodes – 8 nos;	
	ECG Gel – 2 bottle;	
	Thermal recording Paper – 5nos	
	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified above).	
6	Operating Environment	
6.1	The system offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Power Supply, Climate, Temperature, Humidity, etc.	
6.2	Power supply: 220–240V AC, 50Hz fitted with appropriate plug type. The power cable must be at least 3 metre in length	
7	Standards and Safety Requirements	
7.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND	
7.2	CE (93/42 EEC Directives) or USFDA approved product certificate	
8	User Training	
8.1	Must provide user training (including how to use and maintain the equipment).	
9	Warranty	
9.1	Comprehensive warranty for 1 years after acceptance	
10	Maintenance Service During Warranty Period	
10.1	During the warranty period supplier must ensure corrective/breakdown maintenance whenever required.	
11	Installation and Commissioning	
11.1	Supplier must accomplish proper installation and commissioning of the equipment on site.	
12	Documentation	
12.1	User (Operating) manual in English	
12.2	Original Brochure Must be submitted	1



S.N.	Purchaser's Specifications	Bidders Offer
	Bain Circuit	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
	Description of Function	
1.1	Bain Circuit is comprises co-axial modification of basic T-piece system which has been developed for facilitating scavenging of waste anesthetic gases. As a tube carrying fresh gas, it travels inside outer reservoir tube to endotracheal tube connector. The process includes patient inspiring fresh gas from the outer reservoir tube and expiring into reservoit tube.	
2	Technical Specifications	
2.1	Should be Compact and inexpensive with low dead-space.	
2.2	Should have Low resistance to breathing	
2.3	Should Facilitates scavenging of waste gases.	
2.4	Should be either Sterile or Non- Sterile or Individual Packed	
3	Standards and Safety Requirements	
3.1	ISO 9001:2003 or CE if applicable	



	ZO. T piece	- 10000000000
S.N.	Purchaser's Specifications	Bidders Offer
	T piece	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Technical Specifications	
1.1	It should be medical grade material.	
1.2	Its tube length should be at least 200cm.	
2	Standards and Safety Requirements	
2.1	ISO 9001:2003 or CE if applicable	

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21. Laryngoscope Set

.N.	Purchaser's Specifications	Bidders Offer
	Laryngoscope Set	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1.1	Laryngoscopy to facilitate tracheal intubation during general anaesthesia or cardiopulmonary resuscitation or for procedures on the larynx or other parts of the upper tracheobronchial tree.	
2	Operational Requirements	
2.1	Battery powered laryngoscope unit (handle to take C-size batteries).	
3	System Configuration	
3.1	Laryngoscope set	
4	Technical Specifications	
4.1	Blades to be made of surgical grade stainless steel.	
4.2	Clip-on quick release mechanism for blades, which also provides electrical contact for blade light. Light to be activated when blade is engaged.	
4.3	Shall operate on C-size batteries.	

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21. Laryngoscope Set

S.N.	Purchaser's Specifications	Bidders Offer
4.4	Handle/battery unit to be made of non-ferrous metal.	
5	Accessories, spares and consumables	
	Accessories:	
	• Spare bulbs: 03 nos.	
	Blades: One each of following sizes:	
5.1	i-Neonate size 00	II.
	ii-Adult small size 3	
	iii-Adult medium size 4	
	iv-Adult large size 5	
	Set of C-sized batteries	
5.2	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified above).	
6	Operating Environment	
5.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Climate, Temperature, Humidity, etc.	
6.2	Battery operated system.	
		A

हरिश्चन्द्र अधिकारी समस्त्र प्रहरी नायत्र महानिरीक्षक अस्पताल प्रमुख



21. Laryngoscope Set

S.N.	Purchaser's Specifications	Bidders Offer
7	Standards and Safety Requirements	
7.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND	
7.2	CE (93/42 EEC Directives) or USFDA approved product certificate.	
8	User Training	
8,1	Not applicable.	
9	Warranty	
9.1	Warranty for 1year after acceptance.	
10	Maintenance Service During Warranty Period	
10.1	Standard warranty conditions are applicable.	
11	Installation and Commissioning	
11.1	Must supply preassembled unit, ready to use.	
12	Documentation	
12.1	User (Operating) manual in English.	
12.2	Service (Technical / Maintenance) manual in English.	
12.3	List of important spare parts and accessories with their part number and costing.	

21. Laryngoscope

S.N.	Purchaser's Specifications	Bidders Offer
	Laryngoscope Set	

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22.Trancheostomy Tube Different Size

5.N	Purchaser's Specifications	Bidder's Offer
	Trancheostomy Tube Different Size	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description Of Function	
1.1	A tracheostomy (trach) tube is a curved tube that is inserted into a tracheostomy stoma (the hole made in the neck and windpipe (Trachea)).	
2	Technical Specifications	
2.1	Should be transparent PVC tube with radiopaque line and low-pressure cuff.	
2.2	Should have adjustable collar and tape	
2.3	This deviceshould be available in different size (size should be clearly mentioned in bidder's offer)	
3	Certification should be provided if applicable	
4	User training/ Technician training if applicable	

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24. ET Tube Different Size with subglottic Suction Plug Port

N	Purchaser's Specifications	Bidder's Offer
	ET Tube Different Size	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description Of Function	
1,1	Endotracheal tube with high quality of valve for relaible control of cuff inflation and pressureand a smooth soft tip reducing the potential of tracheal trauma during intubation	
2	Technical Specifications	
2.1	Should be of single use and sterile	
2.2	Should be latex free	
2.3	Should have depth mark liners to facilitate the placement of tube during intubation	
2.4	Valve should be of good quality for reliable control of cuff inflation and pressure	
2.5	This deviceshould be available in different size (size should be clearly mentioned in bidder's offer)	
3	Certification should be provided if applicable (ISO / CE)	
4	User training/ Technician training if applicable	

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25. Pulse Oximeter

S.N.	Purchaser's Specifications	Bidder's Offer
	Hand Held Pulse Oximeter	
	Manufacturer	
	Brand	
	Type/Model	
	Country of Origin	
1	Description of Function	
1.1	Pulse oximeter for SPO2 and Heart Rate	
2	Operational Requirements	
2.1	Pulse Oximeter with adult and neonate SPO2 Probe	
3	System Configuration	
3.1	Pulse Oximeter with adult and neonate SPO2 Probe	
4	Technical Specifications	
4.1	Should display SPO2 and Pulse Wave form	
4.2	Plethysmograph graph should be displayed on display	
4.3	Should consists of color TFT display with multi directional views with brightness control	
4.4	Should have low battery alarm on display	
4.5	Should have programmable alarms and display	
4.6	Should have pulse sound indication	
4.7	Should have AC charger	
4.8	Should be with Flash memory with 24 data read back.	
4.9	Display direction should be changed.	
4.10	Should have data transmission facilities.	
5.0	System Configuration Accessories, Spares and Consumables.	
5.1	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials to be included in the offer.	
5	Operating Environment	
6.1	The system offered must be designed to operate normally under the condition of the purchaser's country. The conditions include power supply, climate, temperature, and humidity, etc.	
7	Standards and Safety Requirements	
7.1	Must submit ISO 9001 or ISO 13485:2003/AC:2007 AND	
7.2	CE or USFDA approved product certificate	
3	User Training	
3.1	Should provide user training	
)	Warranty	
0.1	Comprehensive warranty for 1 year after acceptance.	
0	Maintenance Service During Warranty Period	
0.1	Standard warranty conditions are applicable.	
1	Installation Inspection and Commissioning	
11.1	Must supply preassembled unit, ready to use.	

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S.N.	Purchaser's Specifications	Bidder's Offer
12	Documentation	
12.1	User/Instructions manual shall be provided in English.	

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26.	Stet	hosco	pe
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N.	Purchaser's Specifications	Bidder's offer
	Stethoscope	
	Manufacturer	
	Brand	-10
	Type / Model	
	Country of Origin	
1	Description of Function	
1,1	The stethoscope is used for listening to the beating heart of a human, or lungs. It is also used for listening to the flow of the blood in the surrounding area of the heart.	the
2	Operational Requirements	
2.1	Dual type stethoscope - Physician's stethoscope.	
3	System Configuration	
3.1	Stethoscope, dual cup/bellTubes	
4	Technical Specifications	
4.1	Dual, cup/bell and diaphragm head	
4.2	Head and ear tube assembly to be made of non-ferrous metal,	
4.3	Tubes to be synthetic material and ear tubes to have shaped plastic cush ends.	ion
5	Accessories, spares and consumables	
5.1	All standard accessories/consumables/parts required for the proper operation of the above item shall be included in the offer. Bidders shall specify, in a separate Excel worksheet, the quantity and details of any ite included in this offer which have not been specified in this Technical Specifications Form.	ems
6	Operating Environment	

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26. Stethoscope

S.N.	Purchaser's Specifications	Bidder's offer
		1
5.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Climate, Temperature, Humidity, etc.	0.0
	Cinnate, Temperature, Trainierey, etc.	
7	Standards and Safety Requirements	
7.1	Must submit ISO 9001 or ISO 13485:2003/AC: 2007.	
8	User Training	
8.1	Not applicable.	
9	Warranty	1/1
9.1	Warranty for 1 year.	
10	Maintenance Service During Warranty Period	
10.1	Standard warranty conditions are applicable.	
11	Installation and Commissioning	
11.1	Must supply preassembled unit, ready to use.	
12	Documentation	
12.1	User's manual in English	

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27. BP apparatus

.N.	Purchaser's Specifications	Bidder's Offer
	Sphygmomanometer (BP apparatus)	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1.1	Sphygmomanometer is a device used to measure blood pressure, composed of an inflatable cuff to restrict blood flow, and a mechanical manometer to measure the pressure.	
2	Operational Requirements	
2.1	Aneroid sphygmomanometer having a dial to show clear numbers and pointer / needle for measurement of pressure.	
3	System Configuration	
	Aneroid sphygmomanometer	
3.1	Cuffs for child size and for adult size (regular)	
	Inflation bulb	
	Carrying pouch	
4	Technical Specifications	
4.1	Packed in easy carrying high quality pouch made of waterproof cloth to accommodate cuff, and inflation bulb.	
4.2	Gauge to be calibrated in 2 mm Hg units.	

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27. BP apparatus

S.N.	Purchaser's Specifications	Bidder's Offer
4.3	Must provide blood pressure cuffs for child size and for adult size (regular).	
5	Accessories, spares and consumables	
5.1	All standard accessories/consumables/parts required for the proper operation of the above item shall be included in the offer. Bidders shall specify, in a separate Excel worksheet, the quantity and details of any items included in this offer which have not been specified in this Technical Specifications Form.	
6	Operating Environment	
6.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Climate, Temperature, Humidity, etc.	
7	Standards and Safety Requirements	
7.1	Must submit ISO 9001 or ISO 13485:2003/AC: 2007.	
8	User Training	
8.1	Not applicable.	
9	Warranty	
9.1	Warranty for 1year.	
10	Maintenance Service During Warranty Period	
10.1	Standard warranty conditions are applicable.	Fe-
11	Installation and Commissioning	
11.1	Must supply preassembled unit, ready to use.	
12	Documentation	
12.1	User's manual in English	

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28. Needle Destroyers

28. Needle Destroyers				
S.N.	Purchaser's Specifications	Bidders Offer		
	Needle Destroyers			
	Manufacturer			
	Brand			
	Type / Model			
	Country of Origin			
1	Description of Function			
1.1	Needle destroyers are used to destroy the needles instantly to prevent reand manage waste management effectively.	euse		
2	Operational Requirements			
2.1	The needle should be completely incinerated without visible sparking a arcing	and		
3	System Configuration			
3.1	Needle Destroyers, complete unit with complete accessories.			
4	Technical Specifications			
3,1	Built in SS sharp blade cutter to cut the nozzle of the syringe			
3.2	Needle destruction rate shall be max. of 2 seconds per needle.			
3.3	Provision of removable and reusable collection receptacle for syringe no and needle debris of approximately 500 syringes.	ozzle		

हरिश्चन्द्र अधिकारी सशस्त्र प्रहरी नायव महानिरीक्षन अस्पताल प्रमुख



28. Needle Destroyers

S.N.	Purchaser's Specifications	Bidders Offer
3.4	Preferably shall have collection receptacle to have a see-through panel to view the waste.	
3.5	During operation and removal of refuses the container shall be designed for safe handling against any injury or spill over / contact with debris.	
3.6	Provision of on/off switch and pilot lamp.	
3.7	Unit shall be made of high grade stainless steel material.	
3.8	Must be able to destroy of all types of needle.	
4.9	Unit shall be shock proof and provided with proper insulation as per international safety standard norms.	
5	Accessories, spares and consumables	
5.1	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified above).	
6	Operating Environment	
6.1	The system offered shall be designed to store and to operate normally under the conditions of the purchaser's country. The conditions include Power Supply, Climate, Temperature, Humidity, etc.	

हरिश्चन्द अधिकारी सरास्त्र प्रहरी नायत्र महानिरीक्षण अस्पताल प्रमुख



28. Needle Destroyers

		1
S.N.	Purchaser's Specifications	Bidders Offer
6.2	Power supply: 220-240 VAC, 50Hz single phase as appropriate fitted with appropriate plug. The power cable must be at least 3 metre in length.	
7	Standards and Safety Requirements	
7.1	Must submit ISO 9001 or ISO13485:2003/AC:2007 AND	
7.2	CE (EEC Directives) or USFDA approved product certificate.	
8	User Training	
8.1	Must provide user training (including how to use and maintain the equipment).	
9	Warranty	
9.1	Comprehensive warranty for 2 years after acceptance.	-
10	Maintenance Service During Warranty Period	
10.1	During the warranty period supplier must ensure corrective/breakdown maintenance whenever required.	
11	Installation and Commissioning	
11,1	The bidder must arrange for the equipment to be installed and commissioned by certified or qualified personnel; any prerequisites for installation to be communicated to the purchaser in advance, in detail.	
12	Documentation	Λ

हरिश्चन्द्र अधिकारी संशस्त्र प्रहरी नायव महानिरीक्षक अस्पताल प्रमुख



28.	Needle	Destroyers
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.N.	Purchaser's Specifications	Bidders Offer
12.1	User (Operating) manual in English.	Bidders Offer
12.2	Service (Technical / Maintenance) manual in English.	
12.3	List of important spare parts and accessories with their part numbers and costing.	

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29. Autoclave, Horizontal, Double Door, 800 litres

S.N.	Purchaser's Specifications	Bidder's Offer Hand
	Autoclave, Horizontal, Double Door, 800 litres	र पी एक
	Manufacturer	रतियु, काठियः
	Brand	
	Type / Model	
	Country of Origin	
	Description of Function	
1.1	CSSD autoclave shall be able to sterilize wrapped instruments, unwrapped instruments, linen, glassware, liquids.	
	Operational Requirements	
2.1	Microprocessor controlled horizontal electrically heated autoclave is required.	
	System Configuration	
3.1	Autoclave, Horizontal, Double Door, 800 litres, with complete accessories.	
- 4	Technical Specifications	
4.1	Shall have fully automatic operation.	
4.2	The sterilizer shall be pneumatically (Compressed Air) operated, fully automatic double door, triple jacketed chamber front loading.	

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4.3	The autoclave shall be designed to operate on various pre select programs such as pre-vacuum cycle, gravity cycle, porous load cycle, liquid cycle and two standard Bowie Dick test and vacuum leak cycle.	महाति प्रमाण के स्टूबर्स के स्टूबर के स्टूब
4.4	Autoclave shall work up to 134-136 °C temperature.	
4.5	It shall come with vertical sliding door, a trolley, a carriage, a steam generator and a dedicated air compressor.	
	Construction:	
	Jacket shall be constructed of heavy duty 304L grade stainless steel.	
4.6	Door shall be constructed of heavy duty 304L grade stainless steel.	
	Chamber shall be constructed of heavy duty 316L Ti grade stainless steel.	
	All the pipes and fittings are made of stainless steel and Brass.	
4.7	Chamber constructed of heavy duty 316L grade stainless steel shall have following features:	
		1

हरिश्चः विकारी संशस्त्र प्रहरी नायन महानिरीक्षक अस्पताल प्रमुख

	Chamber shape: Horizontal rectangular design Chamber shape: Horizontal rectangular design	
	Chamber dimensions: 660 X 950 X 1800mm (W x H x D) approximately.	
	Chamber volume: approx. 800+ litres.	
	Shall come with safety features such as:	
	Door must not open in case chamber is pressurized.	
4.8	Safety valves for chamber/jacket, current overload relays and contactors for vacuum pump.	
	Shall have at least two limit switches at the end of door-close position.	
	The door shall slide down immediately upon sensing an obstruction during closure.	
	Shall have thick glass wool insulation, tight wrapped with thick silver foil around jacket and door to avoid heat exposure.	

हरिश्चन्द्र अधिकारी सशस्त्र प्रहरी नायच महानिरीक्षक अस्पताल प्रमुख

4.9	Chamber is provided with two rails for easy/smooth movement of carriage.	THE STATE OF THE S
4.10	On the front panel of autoclave there are different pressure gauges for depiction of actual pressure in chamber, jacket and pressure on gasket.	
4,11	Trolley shall be made of high quality 316L SS to transfer carriage from one place to another and shall have foot locks and locking mechanism for carriage while resting above the trolley.	
4.12	It shall be high speed microprocessor control for accurate progression of sterilization cycle. Facility to save and create history log files that can be opened with the support of Microsoft based operating system. Facility to view and operate the cycle progression from remote location.	
4.13	Keypad shall be provided which is used for selecting the cycle and to adjust and feed alphanumeric data. Multiple password access levels (specify number) shall be provided to control access/operation of the machine preventing unauthorized access. These access levels shall be user selectable.	
4.14	Approx. 7" touch screen multi-colour LCD display for preselect program information. The information must include cycle stage, chamber temperature, chamber pressure, jacket pressure along with the information about failures and interrupts. It shall have storage capacity of approx. 200 cycles built-in memory.	
4.15	Documentation: The system shall come with real time built-in printer which gives/prints the real time event during the propagation of cycle such as time in hour, minute, second along with date, load no., operator etc. Any failure is indicated via audio-visual alarm and a print out.	

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4.16	Shall come with ring type three phase water pre-vacuum pump for pre-vacuum stage and drying stage. Vacuum shall have an adjustable range between 5 kPa and 75 kPa during preselect of 5 pre-vacuum pulses.	र ज एक अस्ति काठमाण्डे
4,17	Shall come with heat condensation device that cools the condensate emitting from autoclave during the exhaust.	
4.18	Shall have fully automatic steam generator made of 316L chamber to feed steam to autoclave jacket and gasket groove. Water reservoir, water sensing electrodes, pressure switches and safety valve must be part of steam generation unit. It shall come with heating element of 55-65KW made of stainless steel.	
4.19	Exhaust air filtration with condensate sterilization for emission-free sterilization of infectious pathogens, equipped with filter cartridge of 0.2 μm pore size, with easy access for replacement.	
4.20	Air compressor: Shall come with air compressor for all pneumatic operation.	
4.21	Even with a total control failure, all mechanical safety features must be left intact.	
4.22	RS 232 interface for direct connection to a personal computer (PC), and programs for conforming documentation, diagrams, storage, and printout.	
5	Accessories, spares and consumables	
5,1	Accessories:	

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	Spare heating element- 2 set	A OF PULS STEER
	Spare air filters: 5 nos.	काठमाण्डी
	Spare door gaskets: 2 nos.	
5,2	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified above).	
6	Operating Environment	
6.1	The system offered must be designed to store and be operated normally under the condition of the purchaser's Country. The conditions include Climate, temperature and relative humidity.	
6.2	Power supply: 380-440 V (3 Phase), 50Hz fitted with appropriate plug to meet purchaser's country requirements. The power cable must be minimum 3 metres long.	
7	Standards and Safety Requirements	
7.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND	

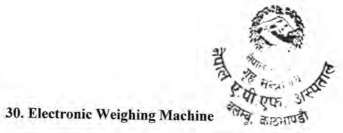
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7.2	CE (93/42 EEC Directives) or USFDA approved product certificate.	and the second s
7.3	Shall meet IEC 61010-2-040 Safety requirements for electrical equipment for measurement, control and laboratory use - Part 2-040: Particular requirements for sterilizers and washerdisinfectors used to treat medical materials.	काउमावडी
8	User Training	
8.1	Must provide user training (including how to use and maintain the equipment).	
9	Warranty	
9.1	Comprehensive warranty for 2 years after acceptance.	
10	Maintenance Service during Warranty Period	
10.1	During warranty period supplier must ensure preventive maintenance and corrective/breakdown maintenance whenever required.	
11	Installation and Commissioning	

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11,1	The bidder must arrange for the equipment to be installed and commissioned by certified or qualified personnel; any prerequisites for installation to be communicated to the purchaser in advance, in detail.	काठमाउँ
12	Documentation	
12.1	User (Operating) manual in English.	
12.2	Service (Technical / Maintenance) manual in English.	
12.3	List of important spare parts and accessories with their part numbers and costing.	
12.4	Certificate of calibration and inspection from factory.	

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S.N.	Purchaser's Specifications Bidder's Offe	
	Electronic Adult Weighing Scale	Didder s Offers
1	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1.1	An electrically operated adult weighing scale	
2	Operational Requirements	
2.1	Electronic Adult Weighing Scale	
3	System Configuration	
3.1	Electronic Adult Weighing Scale on main power as well as	
	battery operated.	
4	Technical Specifications	
4.1	Capacity: 150 kg	
4.2	Accuracy: 100 g	
4.3	Display: LED / LCD	
4.4	TARE facility with zero function.	
4.5	HOLD function to lock the weight.	
4.6	The Scale must have inbuilt rechargeable battery backup	
5	Accessories, spares and consumables	
5.1	All standard accessories, consumables and parts required to	
	operate the equipment, including all standard tools and cleaning	
	and lubrication materials, to be included in the offer. Bidders	
	must specify the quantity of every item included in their offer	
	(including items not specified above).	
5	Operating Environment	
5.1	The system offered shall be designed to be stored and to operate	
	normally under the conditions of the purchaser's country. The	
	conditions include Power Supply, Climate, Temperature,	
	Humidity, etc.	
5.2	Power supply: 220 – 240 VAC, 50Hz Single Phase fitted with	
	appropriate plug. The power cable must be at least 3 metre in	
	length.	
-	Standards and Safety Requirements	
1.1	Must submit ISO 9001 or ISO 13485:2003/AC: 2007.	
0.7	User Training	
.1	Must provide user training (including how to use and maintain	
	the equipment).	
	Warranty	
.1	Comprehensive warranty for 1year.	
0	Maintenance Service During Warranty Period	
0.1	During warranty period supplier must ensure	
	corrective/breakdown maintenance whenever required.	
1	Installation and Commissioning	
1.1	Supplier must accomplish proper commissioning of the	
	equipment on site.	
2	Documentation	
2.1	User (Operating) / service manual in English	



31. AMBU bag with reservoir bag

S.N.	Purchaser's Specifications	Bidder's Remarks
-	Bag Valve Mask (BVM)	
	Manufacturer	
	Brand	
	Type / Model	
4	Country of Origin	
1	Description of Function	
1.1	An Ambu Bag (also known as a Bag Valve Mask or BVM or Ambu bag) is a hand-held device used to provide ventilation to a patient who is not breathing or who is breathing inadequately.	
2	Operational Requirements	
2.1	It shall be self-inflatable and must have pop up valve (non-return valve), attachment for oxygen tube & oxygen reservoir.	
3	System Configuration	
3.1	Ambu bag, complete unit.	
4	Technical Specifications	
4.1	Bag must be made up of medical grade silicon, latex free double layered which retain sensitivity and it must be resistant to rough use.	
4.2	Inlet end of the bag must have separate port for Oxygen supplement.	
4.3	Outer port must be such that re-breathing valve or non-return valve can be attached.	
4.4	Must be supplied with Oxygen reservoir bag of 2000ml and shall deliver tidal volumes of 500-800ml	
4.5	It shall be autoclaveable.	
4.6	It shall be adaptable to all type of facemasks.	
4.7	It shall come with appropriate sized facemasks.	
5	Accessories, spares and consumables	
5.1	Set to be supplied in a heavy duty, re-sealable plastic pouch or clear top plastic box.	
5	Operating Environment	
5.1	The system offered must be designed to store and be operated normally under the condition of the purchaser's Country. The conditions include Climate, temperature and relative humidity.	
7	Standards and Safety Requirements	
7.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND	
1	User Training	
.1	Not applicable.	
	Warranty	
.1	Comprehensive warranty for 1 year after acceptance.	
0	Maintenance Service during Warranty Period	
0.1	Standard warranty conditions are applicable.	
1	Installation, Inspection and Commissioning	
1.1	Must supply preassembled unit ready to use.	
2	Documentation	
2.1	User's manual shall be supplied in English.	

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S.N.	Purchaser's Specifications	Bidder's Offer	Deviation If Any	Remarks
	Hemodialysis Machine			
	Manufacturer			
	Brandw			1
_	Type / Model			+
	Country of Origin			-
1	Description of Function			-
1.1	The haemodialysis unit shall be microprocessor control and capable of providing the following features:			
1.2	Acetate and bicarbonate dialysis with UF accuracy of +/- 1%			
1.3	Volumetric ultrafiltration			
1.4	Sodium and UF profilings			
	Built-in clearance monitoring for real time measurement of effective urea clearance (K) and plasma sodium (Na) for therapy assessment.			
	Built-in blood pressure monitoring for measuring the patient non-invasive blood pressure and pulse rate automatically.			
2	The haemodialysis unit shall have an enlarged and high resolution LCD colour screen for dialysis data display.			
3	The haemodialysis unit shall have a multi- color traffic light located on the top of machine monitor indicating the treatment status.			
4	The keyboard function keys and LCD color display shall provide an immediate overview of the machine status for treatment supervision.			
5	The haemodialysis machine should display informative and context related operator guidance, warning messages and alarm reports.			
6	The haemodialysis machine should include following safety features			
6.1	Closed System Design			
6.2	Volumetric Ultrafiltration			
6.3	Volumetric Concentrate Dilution			
6.4	Startup test		99	
6.5	Self test during treatment			
7	The haemodialysis unit shall have an adjustable arterial blood pump flow rate ranging from 15ml/min to 600ml/min. The unit shall be capable of calculating effective blood flow rate and display in a real-time			

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S.N.	Purchaser's Specifications	Bidder's Offer	Deviation If Any	Remarks
	basis during dialysis automatically.		inj	
8	The haemodialysis unit shall have an adjustable arterial blood pump segment(both for pediatric and adult) for bloodline diameter from 2mm to 10mm.			
9	The haemodialysis machine shall have diagnostic programme for checking individual valves, pumps, and closed loop tightness.			
10.	The haemodialysis machine shall have user-selectable dialysate flow rate of 300,500, 800 ml/min.			
11	The haemodialysis unit shall have adjustable by setting the sodium concentration. The conductivity measurement range should be 12.8 to 15.7 mS/cm			
12	The haemodialysate unit shall have temperature control range from 32.0 to 38.0 degree Centigrade and temperature alarm limits of 33.5 to 38 degree Centigrade.			
13	The haemodialysis machine shall have the following Volumetric Ultrafiltration Control			
13.1	Control Range: 0 to 4L/hr			
13.2	UF Volume: 0 to 9.99L adjustable in 1ml increment.			
13.3	Treatment time: adjustable upto 9 hours 59 min in 1 min increment			
13.4	Isolated ultrafiltration process shall be provided.			
14	The haemodialysis unit shall be capable of online preparation of bicarbonate dialysis Fluid			
15	The haemodialysis unit shall have a hygienic connection for the ultrapure dialysate fluid filter having endotoxin retention capacity not less than 10 ⁶ . The unit shall have to provide a reminder message as the end of filter's service life or maximum number			
	of treatments is about to be reached.			20
16	The measurement of effective urea clearance, dialysis dose (Kt/V) and plasma sodium shall be performed in non-invasive, real time mode without additional disposable required during the treatment.			
17	The haemodialysis unit shall be able to operate and monitor the extracorporeal circuit without interruption for at least 15			

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	कारमावड.	

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S.N.	Purchaser's Specifications	Bidder's Offer	Deviation If Any	Remarks
	min. in case of AC power failure by battery backup.			
18	Concentrate Pump, UF Pump construction shall be stepper motor with diagram.			
19	The haemodialysis unit shall have centrally located function keys for easy use.			
20	The haemodialysis unit shall have the following features with regards to disinfection and cleaning			
20.1	Both chemical and heat disinfection shall be performed.			
20.2	Sodium hypochlorite, diluted formaldehyde or paracetic acid may be used as disinfectant			
20.3	Decalcification shall be possible by using citric acid.			
20.4	Various programmable cleansing cycles can be provided with different phases and timings in accordance with different disinfectants.			
20.5	One-touch fully automatic operation including: pre-rinse, chemical-intake for combined disinfection & decalcification, post-chemical mandatory rinse, and automatic power-off, without extra end-user handling during the whole disinfection process.			
21	The hemodialysis unit shall have the build in non-invasive device for measuring the patient blood pressure automatically with following features:			
21.1	Measuring Range			
21.1.1	Cuff Pressure Range :10-325mmhg or wider choice			
21.1.2	Systolic Range : 30-280mmHg or wider choice			
21.1.3	MAP Range: 20-255mmHg or wider choice			
21.1.4	Diastolic Range: 10-240mmHg or wider choice			
21.1.5	Pulse rate range : 20-245/min or wider choice			
21.2	Alarm Values			
21.2.1	Systolic Range :90& 165 mmHg			
21.2.2	MAP Range: 70& 120mmHg			
21.2.3	Diastolic Range: 50 & 100mmHg			
21.2.4	Pulse range: 40& 150/min			
22	Training Requirement	11		
	The bidder must provide separate class room			

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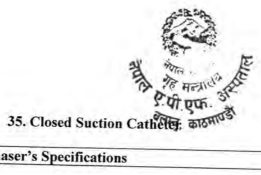
S.N.	Purchaser's Specifications	Bidder's Offer	Deviation If Any	Remarks
	training followed by practical sessions for clinical issues and operation of the machine to the user preferably before the supplied machine are brought into operation.			
	The bidder must provide technician training about the basic technical issues of the machine to the technical person.			
	Training for the User shall be separately included followed by practical session for the clinical issues and operation of the machine.			
23	Spares Availability			
	The bidder shall have ready stock of necessary parts including but not limited to cards, pumps, valves, motors, sensors, and filters for immediate replacement of faulty parts during breakdowns.			
24	The bidder shall confirm the availability of specialized tools (Calibration and other) required for preventive, routine maintenance of offered machine.			
25	Startup Disinfectant			
	The bidder shall provide one canister of recommended disinfectant for each machine supplied as a startup consumable.			
26	Manuals			4-
	The bidder shall include one set each operating manual along with the machine. Technical manual should be provided to each of the participant of the technical training.			
27	Installation			
	The bidder shall install all the machines without any extra cost and shall submit the installation report along with the measurements and set up detail.			
28	Quality Assurance Certificate			
	The bidder shall submit the quality assurance certificate along with the bid. Products with EC/TUV shall be preferred.			
Y	Must submit ISO13485:2003/AC:2007			
29	Warranty			
	2 years of complete maintenance of labour & spares from the date of installation.			

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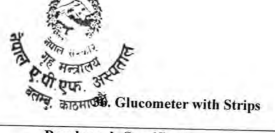
S.N.	-	urchaser's Specifications	Bidder's Offers	
	Resuscitation Cart		Diago o Oneis	
	Manufacturer			
	Brand			
	Type / Model	I P C C		
4	Country of Origin			
1	Description of Functi	on		
1.1	medication/equipment support protocols poten	set of trays/drawers/shelves on wheels used ortation and dispensing of emergency at site of medical/surgical emergency for life intially to save a patient's life.		
2	Operational Requirer	nents		
2.1	Stainless steel trolley of	n stainless steel tubular frame.		
3	System Configuration			
3.1	Resuscitation Cartporta	able with storage units and complete set.		
4	Technical Specification	ns		
4.1	pole assembly	shelf, height adjustable, twin hook/loop, IV		
4.2	moulded plastic).	- at least5 drawers (stainless steel or		
4.3	to have locking/brake n	castors/wheels, with at least one castor/wheel nechanism.		
4.4	Top shelf to have stainl	ess steel guard rail above surface.		
4.5	Fitted with epoxy powd	er coated oxygen cylinder holder.		
4.6	Manual Resuscitator for	r Infant, Children & Adult		
4.7	Should have Manually Infant, Children & Adu	Operated Suction (Foot Suction) Suitable for lt.		
5	Accessories, spares an	d consumables		
5.1	the equipment, including lubrication materials, to	s, consumables and parts required to operate g all standard tools and cleaning and be included in the offer. Bidders must every item included in their offer (including ve).		
5	Operating Environmen			
5.1	The product offered sha normally under the cond conditions include Clim	Il be designed to be stored and to operate litions of the purchaser's country. The ate, Temperature, Humidity, etc.		
	Standards and Safety 1	Requirements		
.1		or ISO 13485:2003/AC:2007 AND		
	User Training			
.1	Not applicable.			
+	Warranty			
.1	Comprehensive warranty	for 1 year.		
0	Maintenance Service D	uring Warranty Period		
0.1	Standard warranty condi			
1	Installation and Comm			
1.1	Must supply preassemble	ed unit, ready to use.		
2	Documentation			
2.1	User's manual shall be s	upplied in English.	1 -	

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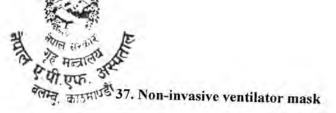
S.N.	Purchaser's Specifications	Bidder's Offers	
	Closed Suction Catheter		
_	Manufacturer		
	Brand		
-	Type / Model		
	Country of Origin		
1	Description of Function		
1.1	Closed suction Catheters used to clear unwanted mucus build-up from the airways of ventilated or tracheostomy patients without the need to disconnect the patient from the ventilator or aerosol source		
2	Operational Requirements		
2.1	Closed Suction Catheter complete set.		
3	System Configuration		
3.1	Closed Suction Catheter complete set.		
4	Technical Specifications		
4.1	Should be made ofnon-toxic, non-irritant medical grade mate	rial	
4.2	Color coded control valve for easy identification of the catheter size.		
1.3	Lockable thumb end cap prevents inadvertent suctioning	ici size.	
4.4	Suction catheter tip should be smooth and soft, rounded to prevent mucosal trauma.		
4.5	Should be supplied in sterile pack		
5	Accessories, spares and consumables		
5.1	Set to be supplied in a heavy duty, re-sealable plastic pouch or clear top plastic box.		
í	Operating Environment		
5.1	The system offered must be designed to store and be operated normally under the condition of the purchaser's Country. The conditions include Climate, temperature and relative humidity.		
	Standards and Safety Requirements		
.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND		
	User Training		
.1	Not applicable.		
	Warranty		
.1	Comprehensive warranty for I year after acceptance.		
0	Maintenance Service during Warranty Period		
0.1	Standard warranty conditions are applicable.		
1	Installation, Inspection and Commissioning		
1.1	Must supply preassembled unit ready to use,		
2	Documentation		
2.1	User's manual shall be supplied in English.		

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S.N.	Purchaser's Specifications	Bidder's Offers
	Glucometer with test strips	
_	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1.1	Glucometer is used to measure blood glucose levels	
2	Operational Requirements	
2.1	Glucometer with test strips, complete set	
3	System Configuration	
3.1	Glucometer with test strips, complete set	
4	Technical Specifications	
4.1	Should be a hand held, Light weight with replaceable battery	
4.2	Should have LCD display with reading range/linearity from approx. 20 to 600 mg/d.	
4.3	Reading time should be of less than 10 seconds	
4.4	Should use a minimum blood sample less than 1.5µl	
4.5	Should have a minimum memory of approx. 50	
4.6	Strips should be available in the local market	
5	Accessories, spares and consumables	
5.1	Test strips –200pieces	
5.2	Covering case – 1no.	
5	Operating Environment	
5.1	The system offered must be designed to store and be operated normally under the condition of the purchaser's Country. The conditions include Climate, temperature and relative humidity.	
7	Standards and Safety Requirements	
7.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND	
	User Training	
3.1	Not applicable.	
	Warranty	
.1	Comprehensive warranty for 1 year after acceptance.	
0	Maintenance Service during Warranty Period	
0.1	Standard warranty conditions are applicable.	
1	Installation, Inspection and Commissioning	
1.1	Must supply preassembled unit ready to use.	
2	Documentation	
2.1	User's manual shall be supplied in English.	

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S.N.	Purchaser's Specifications Non-invasive ventilator mask		Bidder's Offers
			Didder's Offers
	Manufacturer		
	Brand		
	Type / Model		
4	Country of Origin		
1	Description of Funct	ion	
1.1	Non-invasive ventilate	or masks used for non-invasive ventilation therapy	
2	Operational Require	ments	
2.1	Non-invasive ventilate	or mask complete set	
3	System Configuration	1	
3.1	Non-invasive ventilate	r mask complete set	
4	Technical Specification	ons	
4.1	Non-invasive ventilato	r masks should cover the nose and mouth	
4.2	Should provide maximum comfort and effective seal		
4.3	Must be designed to fit with common hospital breathing circuits		
5	Operating Environment		
5.1	The system offered must be designed to store and be operated normally		
	under the condition of	the purchaser's Country The conditions include	
	Climate, temperature a	nd relative humidity.	
6	Standards and Safety Requirements		
5.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND		
7	User Training	1110	
7.1	Not applicable.		
}	Warranty		
3.1	Comprehensive warran	ty for 1 year after acceptance.	
	Maintenance Service	luring Warranty Period	
.1	Standard warranty cond	itions are applicable.	
0	Installation, Inspection	and Commissioning	
0.1	Must supply preassemb	led unit ready to use.	
1	Documentation		
1.1	User's manual shall be s	supplied in English	

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38. Oropharyngeal Airway Different Sizes

S.N.	Purchaser's Specifications	Bidder's Offers	
	Oropharyngeal Airway different sizes		
	Manufacturer		
	Brand		
	Type / Model		
	Country of Origin		
1	Description of Function		
1.1	Oropharyngeal Airway used to maintain or open a patient's airway.		
2	Operational Requirements		
2.1	Oropharyngeal Airway complete set.		
3	System Configuration		
3.1	Oropharyngeal Airway complete set.		
4	Technical Specifications		
4.1	Should be manufactured with medical grade material		
4.2	Should have color coded bite locks for quick identification		
4.3	Sizes from 10-100mm in a set		
4.4	Should be provided in plastic case		
5	Operating Environment		
5.1	The system offered must be designed to store and be opera	ted normally	
	under the condition of the purchaser's Country. The condit	ions include	
	Climate, temperature and relative humidity.		
6	Standards and Safety Requirements		
6.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND		
7	User Training		
7.1	Not applicable.		
8	Warranty		
8.1	Comprehensive warranty for 1 year after acceptance.		
9	Maintenance Service during Warranty Period		
9.1	Standard warranty conditions are applicable.		
10	Installation, Inspection and Commissioning		
10.1	Must supply preassembled unit ready to use.		
11	Documentation		
11.1	User's manual shall be supplied in English.		

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39. Nasopharyngeal Airway

S.N.	Purchaser's Specifications Nasopharyngeal Airway		Bidder's Offers
	Manufacturer		
	Brand		
	Type / Model		
	Country of Origin		
1	Description of Function		
1.1	Nasopharyngeal Airway is a tube that is designed to be inserted into the nasal passageway to secure an open airway		
2	Operational Requirements		
2.1	Nasopharyngeal Airway complete set.		
3	System Configuration		
3.1	Nasopharyngeal Airway complete set		
4	Technical Specifications		
4.1	Should be manufactured with medical grade material		
4.2	Should be used on patients to maintain an airway		
4.3	Should come in sterile package		
4.4	Should have flared end		
5	Operating Environment		
5.1	The system offered must be designed to store and be operated normally under the condition of the purchaser's Country. The conditions include Climate, temperature and relative humidity.		
6	Standards and Safety Requirements		
6.1	Must submit ISO13485:2003/AC:2007 for Medical Devices AND		
7	User Training		
7.1	Not applicable.		
8	Warranty		
8.1	Comprehensive warranty for 1 year after acceptance.		
9	Maintenance Service during Warranty Period		
9.1	Standard warranty conditions are applic	able.	
10	Installation, Inspection and Commiss		
10.1	Must supply preassembled unit ready to use.		
11	Documentation		
11.1	User's manual shall be supplied in Engl	ish.	

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40. Venturi Mask (Different) Fio2

S.N	Purchaser's Specifications	Bidder's Offer
	Venturi Mask (Different) Fio2	- Auden a Chief
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description Of Function	
1.1	Venturi Mask is used to deliver a known oxygen concentration to patient on control oxygen therapy	
2	Technical Specifications	
2.1	Material should be medical PVC	
2.2	Should have adjustable nose clip	
2.3	Oxygen concentration should be 24-50%	
2.4	Should available with anticrush tubing	
2.5	This device should be available in different size (size should be clearly mentioned in bidder's offer)	
3	Certification should be provided if applicable (ISO / CE)	
4	User training/ Technician training if applicable	

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41. Ventilating Face Mask (Anesthetic Mask) Different Size

S.N	Purchaser's Specifications	Bidder's Offer
	Ventilating Face Mask (Anesthetic Mask) Different Size Venturi Mask (Different) Fio2	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description Of Function	
1.1	It is used in connection with medical equipment such as ventilator, oxygen machine, anesthetic apparatus and emergency breathing apparatus.	
2	Technical Specifications	
2.1	Material should be medical grade PVC.	
2.2	Transparent in colour.	
2.3	Should be latex free.	
2.4	Should be hook, ring without check valve.	
2,5	This device should be available in different size (size should be clearly mentioned in bidder's offer).	
3	Certification should be provided if applicable (ISO / CE).	1
4	User training/ Technician training if applicable.	

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42. Refrigerator 185 Ltr

S.N.	Purchaser's Specifications	Bidder's Offer
	Refrigerator	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1.1	Refrigerator 185ltr capacity	
2	Operational Requirements	
2.1	Floor standing model, solid door with lock and handle supplied with two keys.	
3	System Configuration	
3.1	The system consists of:	
	Refrigerator CFC Free	
	PUF insulation	
	Floor standing model	
	Over voltage protection	
4	Technical Specifications	
4.1	Double Compartment:	
	Freezing/ice making compartment.	
	Refrigerator	
4.2	Frost Free	
4.3	Polyurethane (PUF) insulation.	
4.4	Refrigerant Gas: CFC free	
4.5	Over voltage protection: To be supplied complete with mains electric, over-voltage	
	protection unit.	
5	Accessories, spares and consumables	
5.1	Accessories: Adjustable shelves, chiller tray, temperature controller, refrigerator thermometer, auto lamp on/off feature.	
5.2	All standard accessories, consumables and parts required to operate the equipment, including all standard tools and cleaning and lubrication materials, to be included in the offer. Bidders must specify the quantity of every item included in their offer (including items not specified above).	
6	Operating Environment	
5.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Power Supply, Climate, Temperature, Humidity, etc.	
5.2	Power supply: 220 – 240VAC, 50Hz fitted with appropriate plug	
	type D round 3 pins. The power cable must be at least 3 metres in length.	
7	Standards and Safety Requirements	
7.1	Must submit ISO 9001 or ISO 13485:2003/AC: 2007.	
3	User Training	
3.1	Not applicable.	
)	Warranty	
2.1	Comprehensive warranty for 1 year after acceptance.	7

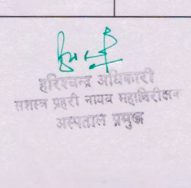
Purchason's Specification of the rate	Bidder's Offer
Maintenance Service During Warranty Period	Didder's Offer
During the warranty period supplier must ensure	
Installation and Commissioning	
Documentation	
User (Operating) manual in English	
Service (Technical / Maintenance) manual in English	
List of important spare parts and accessories with their part numbers and costing.	
	Installation and Commissioning Must supply preassembled unit, ready to use. Documentation User (Operating) manual in English Service (Technical / Maintenance) manual in English List of important spare parts and accessories with their part



S.N.		Purchaser's Specifications	Bidders Offer
	Blood Warmer		Bidders Offer
	Manufacturer		
	Brand		
	Type / Model		
	Country of Origin		
1	Description of Func	tion	
1.1		for warming fluids, crystalloid, colloid or blood g administered to body temperature level.	
2	Technical Specificat	ions	
2.1	Flow Rates should be	from kvo to 150ml/min.	
2.2	Should have temperat	ure range of 36°C to 42°C	
2.3	Should be easily trans	portable	
2.4	Should able to attach	to IV pole and standard electrical sockets	
2.5	Should use dry heat to	echnology	
2.6	Should have audible a	and visual alarms for Temperature	
2.7	Should have automati	c cutoff for set temperature	
2.8	Should be easy to use	and to clean	
2.9		be less than 60 seconds	
2.1	Consumables should l	have built in filter	
5	Accessories, spares a	and consumables	
5.1	Accessories: disposable adult warr		
	disposable pediatric v	varming set-1	



	43. Fluid Warmer		
S.N.	Purchaser's Specifications	Bidders Offer	
5.2	All standard accessories/consumables/parts required for the proper operation of the above item shall be included in the offer. Bidders shall specify, in a separate Excel worksheet, the quantity and details of any items included in this offer which have not been specified in this Technical Specifications Form.		
6	Operating Environment		
6.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Climate, Temperature, Humidity, etc.		
6.2	Power supply: 220-240 VAC, 50Hz single phase as appropriate fitted with appropriate plug. The power cable must be at least 3 metre in length.		
7	Standards and Safety Requirements		
7.1	Should submit ISO 9001 or ISO 13485:2003/AC: 2007 or CE if applicable.		
8	User Training		
8.1	Not applicable.		
9	Warranty		
9.1	Not applicable.		
10	Maintenance Service During Warranty Period		
10.1	Not applicable.		
11	Installation and Commissioning		
11.1	Not applicable.		
12	Documentation		
12.1	User's manual in English		



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44. Physiotherapy chest vibrator

S.N.	Purchaser's Specifications	Bidder's Offers
	Physiotherapy chest vibrator	Didder's Offers
	Manufacturer	-
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Function	
1.1	Physiotherapy chest vibrator used to perform postural drainin COPD patient clear mucus from their airways.	g to help
2	Operational Requirements	
2.1	Physiotherapy chest vibrator complete set.	
3	System Configuration	
3.1	Physiotherapy chest vibrator complete set	
4	Technical Specifications	
4.1	Should be portable and light weight	
4.2	Speed should be approx. 4500rpm	
4.3	Maximum temperature approx. 55°C	
5	Operating Environment	
5.1	The system offered must be designed to store and be operated under the condition of the purchaser's Country. The condition Climate, temperature and relative humidity.	normally s include
5.2	Power supply – 220V, 50Hz	
6	Standards and Safety Requirements	
6.1	Must submit ISO13485:2003/AC:2007 for Medical Devices A	ND
7	User Training	
7.1	Not applicable.	
3	Warranty	
3.1	Comprehensive warranty for 1 year after acceptance.	
)	Maintenance Service during Warranty Period	
0.1	Standard warranty conditions are applicable.	
0	Installation, Inspection and Commissioning	
0.1	Must supply preassembled unit ready to use.	
1	Documentation	
1.1	User's manual shall be supplied in English.	

S.N	Purchaser's Specifications	Bidder's Offer
	Manufacturer	
	Brand	1
	Type / Model	
	Country of Origin	
1	Description of Functions	
1.1	Intermittent pneumatic compression is a therapeutic technique used in medical devices that include an air pump and inflatable auxiliary sleeves, gloves or boots in a system designed to improve venous circulation in the limbs of patients who suffer edema or the risk of deep vein thrombosis or pulmonary embolism	
2	Technical Specifications	
2.1	Should have LCD Screen, touch screen operating, real-time display the parameter of treatment status. treatment part, integrated mode, remaining time, true pressure of every cavity, Inflatable speed etc.	
2.2	Treatment time- 1 min-99 min, adjustable	
2.3	Pressure Range - 0-200mmHg	
2.4	Pressure Holding Time 1s-6s	
2.5	Cycle Interval Time 1s-6s	
2.6	Power Supply AC220V/50HZ	
3	Certification should be provided if applicable (ISO/ CE)	
4	User training/ Technician training if applicable	
		0



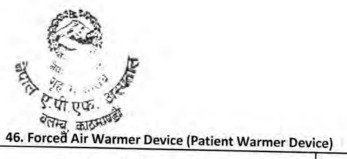
46. Forced Air Warmer Device (Patient Warmer Device)

S.N.	Purchaser's Specifications	Bidders Offer
	Forced air warmer device	
	Manufacturer	
	Brand	
	Type / Model	1
	Country of Origin	
1	Description of Function	
1.1	Forced Air Warming Device is used for the prevention hypothermia.	
2	Operational Requirements	
2.1	For the safe and controlling warming of patients in the Operating Theatre, Emergency Department, ICU and warm setting.	
3	Technical Specifications	
3.1	It must be compact and robust and Unobstructive.	
3.2	The unit must be lightweighted.	
3,3	The unit should have a handle and there should be a mechanism to secure to a drip stand or bed.	
3.4	The control panel must be well sealed preventing entry of fluids to the internal working and circuits of the unit.	
3.5	The Forced air warmer musr have at least 3 selectable temperature settings as follows: i) Low: set value between 30-34°C ii) Medium: set value between 35-39°C iii) High: set value between 39-43°C Accuracy must be within 1.5°C for all setting.	



46. Forced Air Warmer Device (Patient Warmer Device)

S.N.	Purchaser's Specifications	Bidders Offer
3.6	The unit must reach the selected temperture within 60 seconds of the selection of the temperature.	
3.7	The unit must indicate the selected set temperature has been reached and thereafter maintain the selected set temperature.	
3.8	The unit must incorporate visual and audible temperature discrepancy alarms.	, , , , , , , , , , , , , , , , , , ,
3.9	The air supply must incorporate a 0.3 micron (or smaller) HEPA filter. Airflow of at least 19 l/sec must be generated	
3.1	One re usable hose must be supplied with the unit with at least 1.5m and can be easily removal for cleaning or replacement.	
3.11	Forced Air Warmer device Blankets must be disposable, lightweight, soft, radiolucent and latex-free.	
3.12	The Blankets must allow even heat distribution to the patient without the creation of hotspots.	
3.13	There must be a ramge of available blankets.	



S.N.	Purchaser's Specifications	Bidders Offer
2	Accessories, spares and consumables	
4.1	All standard accessories/consumables/parts required for the proper operation of the above item shall be included in the offer. Bidders shall specify, in a separate Excel worksheet, the quantity and details of any items included in this offer which have not been specified in this Technical Specifications Form.	
5	Operating Environment	
5.1	The product offered shall be designed to be stored and to operate normally under the conditions of the purchaser's country. The conditions include Climate, Temperature, Humidity, etc.	
5.2	Power supply: 220-240 VAC, 50Hz single phase as appropriate fitted with appropriate plug. The power cable must be at least 3 metre in length.	
6	Standards and Safety Requirements	
6.1	Must submit ISO 9001 or ISO 13485:2003/AC: 2007.	
6.2	CE or USFDA approved product certificate.	
7	User Training	



46. Forced Air Warmer Device (Patient Warmer Device)

S.N.	Purchaser's Specifications	Bidders Offer
7.1	Must provide user training (including how to use and maintain the equipment).	
8	Warranty	
8.1	Warranty for 2 years.	/
9	Maintenance Service During Warranty Period	
9.1	Standard warranty conditions are applicable.	
10	Installation and Commissioning	
10.1	The bidder must arrange for the equipment to be installed and commissioned by certified or qualified personnel; any prerequisites for installation to be communicated to the purchaser in advance, in detail.	
11	Documentation	
11.1	User (Operating) manual in English.	
11.2	Service (Technical / Maintenance) manual in English.	

47. Bedside Locker

S.N	Purchaser's Specifications	Bidder's Offer
	Bedside Locker	
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Functions	
1.1	Simplify the work of care giver and it enhances comfort and autonomy of the patient in terms of accessability, convenience and storage capacity	
2	Technical Specifications	11
2.1	Over all size should be approx L 40 x W 40 x H82 cms.	
2.2	Should be Machine pressed sheet box with door of heavy guage.	
2.3	Should have CRC tubular legs fitted with PVC stump	
2.4	Should have Stainless steel top.	
2.5	Should be Pre-treated and epoxy powder coated.	
3	Certification should be provided if applicable (ISO/ CE)	
4	User training/ Technician training if applicable	



S.N	Purchaser's Specifications	Bidder's Offer
	Mayo Table	Bluder's Offer
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Functions	
1.1	Mayo Instrument table is an ideal instrument table for most surgical procedures	
2	Technical Specifications	
2.1	Instrument table Mayo with sterilizable stainless steel tray.	
2.2	Height adjustable with telescopic rod with knob from 750 mm to 1400 mm	
2.3	Stainless Steel high polish finish.	
2.4	Mounted on 4 x 50 mm (approx) anti static swivel castors.	
3	Certification should be provided if applicable (ISO/CE)	
4	User training/ Technician training if applicable	



49. Infrared Thermometer (Non-Touch)

S.N.	Purchaser's Specifications	Bidder's Offers
	Infrared Non Touch Hand Thermometer	100000000000000000000000000000000000000
	Manufacturer	
	Brand	
	Type/Model	
	Country of Origin	
	Requirements	
	- Purpose: Non-contact temperature sensing of human body, ideal for individual screening - Measurement Range: At least measure 93.5 F to 109 F	
	- Accuracy: not less than ±0.2°C - Operating Temperature: Human body mode 15°C to 40°C	
	- Display type: LCD with backlight	
	 Response Time: Not more than 3 Seconds, with beep Power Supply: 3 Volt DC (battery operated) Accessories: extra battery 1 pair Warranty: minimum 2 years 	



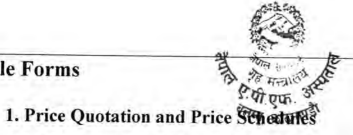
50.Gum Elastic Bongie

S.N	Purchaser's Specifications	Didd-d-off
	Gum Elastic Bongie	Bidder's Offer
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Functions	
1.1	The bougie is device which allows a technique of intubating a patient's airway.	
2	Technical Specifications	
2.1	should be made from low density polyethylene to provides proper stiffness (ease of insertion)	
2.2	Should have Coude tip to facilitates insertion in adults	
2.3	Should be Single use	
2.4	Should be Latex Free	
2.5	Should be calibrated (distance of insertion easily observed for safety)	
3	Certification should be provided if applicable (ISO/CE)	
4	User training/ Technician training if applicable	



s.N	Purchaser's Specifications	Bidder's Offer
	Intubating Stylets	Office
	Manufacturer	
	Brand	
	Type / Model	
	Country of Origin	
1	Description of Functions	
1.1	Intubating Stylets helps to reduce friction for easy insertion into and removal from endotracheal tubes	
2	Technical Specifications	
2.1	Should have Plastic covering extends beyond stylet to minimize the risk of tracheal traum	
2.2	Should be indivivually packed, sterile	
2.3	Should be disposable and single use	
2.4	Should be latex free	
3	Certification should be provided if applicable (ISO/CE)	
4	User training/ Technician training if applicable	

Section V. Sample Forms



Date:			
To: [name and address of	of the Purchaser]		
Gentlemen and/or Ladies	K		
of goods and services in	conformity with the said DP	documents for th	signed, offer to supply and deliver <i>[description</i> he sum of <i>[total amount in words and figures]</i> dule of Prices attached herewith and made parts.
We undertake, if our Pri specified in the Schedule	ce Quotation is accepted, to of Requirements.	deliver the goo	ods in accordance with the delivery schedul
We agree to abide by this Quotation	price Quotation for a Period of	of 15 days from t	the last date fixed for submission of the Price
Until a formal Contract is and your notification of av	prepared and executed, this F ward, shall constitute a binding	Price Quotation, 1 g Contract betwe	together with your written acceptance thereo een us.
We understand that you ar	re not bound to accept the lower	est or any Price (Quotation you may receive.
Dated this	day of	20	
[signature]	[in the capacit	ty of]	
Duly authorized to sign Pri	ice Quotation for and on behal	lf of	

			2. Price	e Schedule	र जिएक अ		
Name of	Supplier			Page of	्रिक्ष एक. अस्य		
Propose	r can quote the price for	single item/	multiple iter	ns.	हरिश्चन अधि		
1	2	3	4	5	खरास्त्र प्रहरी व	ावि प्रकार	
Item No.	Description	Unit	Quantity	Unit price (Site Delivery)	Total price per item (cols. 4 x 5)	ल प्रश्नुश्वर	
İ,	Intensive Care Bed	set	20				
2.	IV Stand	Set	40				
3.	Defibrillator	set	1				
4.	Portable x-ray	set	1				
5.	Air mattress	pc	20				
6.	Patient Trolley	set	2				
7.	Wheel chair	set	4				
8.	Bi pap machine	set	2				
9.	Nebulizer	set	8				
10.	Suction Machine	set	8				
11.	X ray view box	set	4				
12.	ECG machine	set	1				
13.	Bain circuit	set	4				
14.	T piece	set	8				

15. Laryngoscope Set

set

2

हरिश्चन्द्र अधिकारी व्यक्ति नामन महाक्रिके

	200 000			2 %	MA	हरिश्चन्द्र सशस्त्र प्रहरी नाम	न महानिरी
16.	Tracheostomy tube Different size	pack	1	1 7 P	न जार में इ. काटमाई	सशस्त्र प्रहरी नाम अस्पताल	प्रमुख
17.	ET tube different size with subglottic suction plug Port	pack	30	<i>चल</i> ह	बु, क्वारुमा		
18.	Pulse oximeter	set	3				
19.	Stethoscope	set	20				
20.	BP apparatus	set	4				
21.	Electric needle destroyer	set	2				
22.	Autoclave 800 Ltr.	set	1				
23.	Electronic weighing machine	set	1				
24.	AMBU bag with reservoir bag	set	4				
25.	hemodialysis unit	set	1				
26.	Resuscitation cart	set	2				
27.	Closed Suction catheter	set	30				
28.	Glucometer With Strips	set	2				
	Noninvasive Ventilator Mask	set	12				
30.	oropharyngeal airway Different Sizes	set	30				
31.	nasopharyngeal airway	set	10				
	Venturi Mask (Different) Fio2	set	20				

				A	4	F.	प्रास्त्र पहरी क	
33.	Ventilating Face Mask (Anesthetic Mask) Different Size	set	10	T.	ेपी एफ लम्बू, काडम	3/200	अस्पताल अस्पताल	11/10
34.	Refrigerator 185 Ltr	set	2					
35.	Blood warmer / Fluid Warmer	set	5					
36.	Physiotherapy Chest Vibrator	set	2					
37.	Intermittent Pneumatic Compression Device	set	2					
38.	Forced Air Warmer Device (patient warmer device)	set	3					
39.	Bedside Locker	set	20					
40.	Mayo Table	set	20					
41.	infrared thermometer (Non Touch)	pc	5					
42.	Gum elastic Bongie	set	2					
43.	Intubatins Stylets	set	5					
		Total A	Mount					
	Add 13% V	alue Add	led Tax					
	Total	ıg VAT						

Total Price	.(in words)
Signature of Bidder	
Note: In case of discrepancy between unit price and total, the unit price shall prev	ail.

The Bidder has option to bid single or multiple items.



हरिश्चन्द्र अधिकारी सशस्त्र प्रहरी नाम महानिरीक्षक अस्पताल प्रमुख

THIS AGREEMENT made the	day of	20	hetween Iname of Burelon and (harris Commit
"the Purchaser") of the one part and Supplier") of the other part:	[name of Supplier]	of [city	between [name of Purchaser] (hereinafter called and country of Supplier] (hereinafter called "the
WHEREAS the Purchaser invited Pri goods and services] and has accepted the sum of [contract price in words a	a Price Quotation b	v the Sun	Is and ancillary services, viz., <i>[brief description of plier for the supply of those goods and services in the Contract Price?</i>]

NOW THIS AGREEMENT WITNESSETH AS FOLLOWS:

- In this Agreement words and expressions shall have the same meanings as are respectively assigned to them in the Conditions of Contract referred to.
- 2. The following documents shall be deemed to form and be read and construed as part of this Agreement, viz.:
 - a. Price Quotation Form and the Price Schedule submitted by the Supplier;
 - b. The Schedule of Requirements;
 - c. The Technical Specifications;
 - d. The Conditions of Contract; and
 - e. The Purchaser's Notification of Award.
- In consideration of the payments to be made by the Purchaser to the Supplier as hereinafter mentioned, the Supplier hereby covenants with the Purchaser to provide the goods and services and to remedy defects therein in conformity in all respects with the provisions of the Contract.
- 4. The Purchaser hereby covenants to pay the Supplier in consideration of the provision of the goods and services and the remedying of defects therein, the Contract Price or such other sum as may become payable under the provisions of the contract at the times and in the manner prescribed by the Contract.

IN WITNESS whereof the parties hereto have caused this Agreement to be executed in accordance with their respective laws the day and year first above written.

On behalf of the Purchaser	On behalf of the Supplier	
Name:	Name:	
Designation:	Designation:	
Sign:	Sign:	
Seal:	Seal:	