Tender Ref. No.: PMBI/SURGICAL/RC-210/2023 Dated: 15/06/2023



PHARMACEUTICALS & MEDICAL DEVICES BUREAU OF INDIA (PMBI)

(Set up under the Department of Pharmaceuticals, Govt. of India)

8th Floor, Videocon Tower, Block E1 Jhandewalan Extension, New Delhi-110055 Telephone: <u>011-49431800/49431829/49431894/49431854/49431874/49431811</u>

Website: janaushadhi.gov.in

e- TENDER FOR SUPPLY OF SURGICAL/ CONSUMABLES AND MEDICAL DEVICES

TO

PHARMACEUTICALS & MEDICAL DEVICES BUREAU OF INDIA (PMBI) FOR TWO YEARS RATE CONTRACT

LAST DATE FOR ONLINE SUBMISSION OF TENDER: 12/07/2023 till 17:00 hours



PHARMACEUTICALS & MEDICAL DEVICES BUREAU OF INDIA (PMBI)

(SET UP UNDER THE DEPARTMENT OF PHARMACEUTICALS, GOVERNMENT OF INDIA)

Regd. Office: 8th Floor, Videocon Tower, Block E1, Jhandewalan Extension, New Delhi-110055

Working Office: 8th Floor, Videocon Tower, Block E1, Jhandewalan Extension, New Delhi-110055

Telephone: <u>011 - 49431800/49431872/49431894/49431829/49431854.</u>

Website: www.janaushadhi.gov.in,

e-TENDER FOR TWO YEARS RATE CONTRACT

FOR SUPPLY OF SURGICAL/ CONSUMABLES AND MEDICAL DEVICES TO PHARMACEUTICALS & MEDICAL DEVICES BUREAU OF INDIA (PMBI)

Tender Reference	PMBI/SURGICAL/RC-210/2023 Dated 15/06/2023
Tender Website	https://eprocure.gov.in
Date of availability of tender documents on website	On 20/06/2023 (Tuesday)
Doubts and queries regarding Tender document should be sent by e-mail to e-mail id "proc10@janaushadhi.gov.in, procure16@janaushadhi.gov.in, procure14@janaushadhi.gov.in, proc6@janaushadhi.gov.in, proc9@janaushadhi.gov.in" by the likely bidders latest by	Till 26/06/2023 up to 17:00 Hours
Time, date and place of pre-bid meeting	On 27/06/2023 (Tuesday) at 15:00 Hours Pharmaceuticals & Medical Devices Bureau of India (PMBI), 9 th Floor, Videocon Tower, Block-E1, Jhandewalan Extension, New Delhi-110055
Last date and time for submission of Online Bid i.e., Bid Submission End Date and time	On 12/07/2023 up to 17:00 Hours
Last Date and time for submission of EMD and Original Required Documents as per ANNEXURE I (Check List) in physical Form and samples in office of Pharmaceuticals & Medical Devices Bureau of India, 8 th Floor, Videocon Tower, Block-E1, Jhandewalan Extension, New Delhi-110055	On 17/07/2023 by 15:00 Hours
Time and date of opening of Technical Bid	On 17/07/2023 at 16:00 Hours (Wednesday)

PMBI/SURGICAL/RC-210/2023

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Place of opening of tender	Pharmaceuticals & Medical Devices Bureau of India (PMBI), 8 th / 9 th Floor, Videocon Tower, Block-E1, Jhandewalan Extension, New Delhi-110055		
Opening of Tender online on	https://eprocure.gov.in		
Address for Communication	Pharmaceuticals & Medical Devices Bureau of India (PMBI), 8th Floor, Videocon Tower, Block-E1, Jhandewalan Extension, New Delhi-110055		
Cost of the Tender Document	Free of cost		
	1. Sh. Gaurav Kaushik Junior Officer (Procurement) Phone: - 011-49431874 Email: - proc10@janaushadhi.gov.in		
	2. Ms. Priya Executive (Procurement) Phone: - 011-49431872 Email: - procure16@janaushadhi.gov.in		
Contact Person for clarification if any	3. Ms. Vakta Parth Belani Senior Executive (Procurement) Phone: - 011-49431894 Email: - procure14@janaushadhi.gov.in		
	4. Sh. P. K. Thakur Assistant Manager (Procurement) Phone: - 011-49431829 Email: - proc6@janaushadhi.gov.in		
	5. Sh. Manik Bera, Manager (Procurement) Phone: - 011-49431854 Email: - proc9@janaushadhi.gov.in		

The tender document can be downloaded free of cost from the CPPP e-Procurement Portal https://eprocure.gov.in and from the website of PMBI: www.janaushadhi.gov.in

Note: The bidders shall be solely responsible for checking these websites at least 3 days prior to closing date of submission of tender for any addendum/amendment issued subsequently to the bid document and take into consideration the same while preparing and submitting the bids. Bids will be opened online.

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PHARMACEUTICALS & MEDICAL DEVICES BUREAU OF INDIA (PMBI)

e-TENDER FOR RATE CONTRACT FOR THE SUPPLY OF SURGICAL/ CONSUMABLES AND MEDICAL DEVICES TO PHARMACEUTICALS & MEDICAL DEVICES BUREAU OF INDIA (PMBI)

PRADHAN MANTRI BHARTIYA JANAUSHADHI PARIYOJANA (PMBJP) is the initiative of Department of Pharmaceuticals, Ministry of Chemical and Fertilizer, Government of India launching with the noble objective of making quality generic medicines, surgical & consumables, and food products available at affordable prices for all, particularly the poor and disadvantaged, through specialized outlets called PRADHAN MANTRI BHARTRIYA JANAUSHADHI KENDRA (PMBJK). PMBI was established in December 2008 under the Department of Pharmaceuticals, Government of India, with the support of all the CPSUs, and identified as the executing agency for PMBJP.

The Bureau has been registered as an independent society under the Societies Registration Act, 1860, in April 2010.

At present, more than 9500 stores are functional. It is proposed to channelize efforts to popularize PMBJP and ensure availability of the complete basket of quality generic medicines, surgical & consumables and food products at affordable prices.

Tender Inviting Authority – CEO, Pharmaceuticals & Medical Devices Bureau of India, 8th Floor, Videocon Tower, Block E1, Jhandewalan Extension, New Delhi-110055 (hereinafter referred as **Tender Inviting Authority** unless the context otherwise requires).

Tender Accepting Authority – CEO, Pharmaceuticals & Medical Devices Bureau of India (hereinafter referred as **PMBI** unless the context otherwise requires).

Tender Inviting Authority Invites Tender for the supply of Surgical/ Consumables and Medical Devices to Pharmaceuticals & Medical Devices Bureau of India (PMBI) for Two Years.

PHARMACEUTICALS & MEDICAL DEVICES BUREAU OF INDIA (PMBI) was formerly known as BUREAU OF PHARMA PSUs OF INDIA (BPPI).

1.TENDERING SYSTEM:

The Bids are to be submitted in two Parts i.e.

- i. Technical Bid (Cover "A")
- ii. Financial Bid / Price Bid (Cover "B")
- i. The **TECHNICAL BID** shall contain the complete technical details of the firm and the documents to provide the eligibility and competency of the bidder and shall be submitted online only in the manner prescribed in Bid document.
 - The documents like Tender Document and Earnest Money Deposit (EMD) shall be submitted before the specified schedule at the office of PMBI super scribed, "Tender Documents & Earnest Money Deposit for Tender Reference No.-PMBI/SURGICAL/RC-210/2023 dated 15/06/2023 for the procurement of Surgical/ Consumables and Medical Devices for the year 2023-2025". However complete hard copy of uploaded tender shall be provided by the bidding firm along-with the mandatory required documents as per clause 3 of Bid document and Earnest Money Deposit (EMD) for evaluation purpose only. This hard copy shall under no case substitute/modify the provisions of e-tender system.
- **ii.** The **Financial Bid/Price Bid** shall be valid for a period of 150 days from the date of opening of Technical Bid. Prior to the expiry of the bid validity, the Tender Inviting Authority may request the Tenderers to extend the bid validity for further period as deemed fit on their original quoted prices and all terms & conditions. However, PMBI reserves the right to place purchase orders at the quoted rate till such period.
 - a) The Tenderer shall fill in the rate per unit size, % age rate of GST in respective column of BOQ for the items quoted.
 - b) In determining the lowest evaluated price, the rate quoted per unit size exclusive of GST as indicated in column No. 7 of the **BOQ** shall be taken into consideration.
 - c) Tender has been called for in the <u>Generic name of items</u> and the Tenderers should quote the rates for the generic products only. The composition, strength and packing of each product should be as per specifications given in **ANNEXURE-XII**, any variation, if found, will result in rejection of the tender.
 - d) Rates (inclusive of customs duty, packing & forwarding charges, transportation, insurance, and any incidental charges, but exclusive GST) should be quoted for each of the required **Surgical/Consumables and Medical Devices** etc., on door delivery basis according to the unit ordered. Tender for the supply of **Surgical/Consumables and Medical Devices** etc. with cross conditions like "AT CURRENT MARKET RATES" shall not be accepted. Handling, clearing, transport charges etc., will not be paid separately. The delivery should be made as stipulated in the purchase order placed with Tenderers.
 - e) The price quoted by the tenderers shall not, in any case exceed the ceiling price as fixed by NPPA (National Pharmaceutical Pricing Authority) as per the provision of "Drugs Price Control Order (DPCO)" if any.

In case any tenderer quotes higher than the ceiling price, competent authority shall be informed for appropriate action.

2. LAST DATE AND TIME FOR SUBMISSION OF ONLINE TENDER:

- i. (a) Online Bids [in two separate Cover {Technical bid (Cover "A") and price bid (Cover "B")}] shall be submitted till 17.00 hours Up to 12/07/2023 (Wednesday) on CPP portal i.e., https://eprocure.gov.in.
 - (b) Hard copy of complete required documents as Per Clause 3. Eligibility Criteria of Bid and EMD shall be submitted on or before the specified schedule at the below mentioned address of PMBI with super scribed, "Tender Document & Earnest Money Deposit (EMD) for Tender Reference No.-PMBI/SURGICAL/RC-210/2023 dated 15/06/2023 for the procurement of Surgical/ Consumables and Medical Devices for the year 2023-2025".

"To,

The Chief Executive Officer (CEO),
Pharmaceuticals & Medical Devices Bureau of India (PMBI)
8th Floor, Videocon Tower, Block-E1,
Jhandewalan Extension, New Delhi-110055"

ii. **Late Tender: -**There is NO PROVISION of uploading late tender beyond stipulated date & time in the e-tendering system.

Note: Sealed envelope containing tender/bid documents as per Annexure-I (Checklist) must be submitted in the name of Tender Inviting Authority 'The CEO, PMBI' as mentioned above. The documents received in the name of any other than tender inviting authority shall be outrightly rejected.

3. ELIGIBILITY CRITERIA (TECHNICAL BID -COVER "A"):

Minimum Eligibility criteria along with list of documents to be submitted in Cover 'A'. Bidders should meet the following criteria to be eligible for bidding and relevant papers/documents must be submitted by them in their technical bid (Cover- 'A') in support of their eligibility for the tender.

A) Earnest Money Deposit (EMD): EMD of Rs.100000/- (Rupees One Lakh only) as specified in Clause 6 of the Tender document in the form of Bank Guarantee or National Electronic Fund Transfer (NEFT) or Bankers Cheque or Demand Draft from Nationalized/Scheduled Bank favoring "Pharmaceuticals & Medical Devices Bureau of India "payable at Delhi which is to be submitted in original to PMBI, New Delhi on or before the date and time stipulated in tender document. Name & full address of the bidder may be written at the back of the Demand Draft/Pay Order. Signed and scanned soft copy of the EMD instrument must be uploaded (ANNEXURE III) to the e-Procurement portal.

EMD in any other form like Cheque/cash/postal order etc. will not be accepted. The Bid (in case not exempted for EMD as mentioned in tender document) without EMD shall be summarily rejected.

Account Details for National Electronic Fund Transfer (NEFT):

Bank Name: Bank of Baroda, Account No. 05860200001696, IFSC Code: BARB0PARLIA

Note: (i) Tenderers falling under MSEs may be exempted from the payment of EMD, if valid registration certificate from NSIC/MSME/ Udyam Registration Certificate is uploaded and submitted self-attested copy with Technical Bid for the product for which bidder has submitted quotation.

(ii) The prior turnover and prior experience for Start-ups (as defined by Department of Industrial Policy and Promotion) shall not be applicable subject to submission of certificate of recognition as start up by Department of Industrial Policy and Promotion for quoted item.

- B) Documentary evidence of the constitution of the company/firm/Proprietorship such as Memorandum and Articles of Association, Partnership Deed etc. should be submitted with details such as Name, Address, Telephone Number, Fax Number, e-mail address of the firm and of the Managing Director / Partners / Proprietor should be submitted to support the fact that the bidding firm is a manufacturer.
- C) Power of Attorney or Resolution of the Board by which the authorized signatory has been authorized by the bidding firm to sign the documents should be submitted.
- D) Tenderer shall be a manufacturer and shall be registered from Director of Industries/District Industries centres, Ministry of Commerce or NSIC for non-drug items and from Licensing Authority for notified items under Class A & B and non-notified under Class C & D. However, the items under mandatory licensing regime as per 'Circular vide F. No. 29/Misc./03/2022-DC (94) dated 12th April, 2023, the manufacturer has to submit application for manufacturing licence or as the case may be duly issued from licensing authority. For items covered under BIS, tenderer should have BIS certificate. In case, the products are covered under Drugs and Cosmetics Act 1940 / Medical Device Act 2017, the tenderer shall have valid manufacturing licence duly licensed by licensing authorities.

E) Bidders must have: -

- a. The bidder must have a valid manufacturing license and duly acknowledged renewal application with old license issued by the state licensing authority/Central licensing approving authority under Medical Devices Rules 2017/Drugs & Cosmetic Act 1940 wherever applicable.
- b. Manufacturing License / permission along with approved product list issued as per the license issued for quoted Drug item / Medical Devices / Consumable / Surgical as per Medical Devices Rules 2017 / Drugs & Cosmetic Act 1940 must be valid till the last date of the submission of tender.
- c. (i) The medical device shall conform to the standards laid down by the Bureau of Indian Standards established under section 3 of the Bureau of Indian Standards Act, 1985 (63 of 1985) or as may be notified by the Ministry of Health and Family Welfare in the Central Government, from time to time
 - (ii) Where no relevant Standard of any medical device has been laid down under sub-rule (1), such device shall conform to the standard laid down by the International Organization for Standardization (ISO) or the International Electro Technical Commission (IEC), or by any other pharmacopeial standards.
 - (iii) In case of the standards which have not been specified under sub-rule (1) and sub-rule (2), the device shall conform to the validated manufacturer's standards.
- d. In case if intending bidders is bidding for electronic items, they must comply the item specification, quality parameters, Safety and Product Standards mentioned in the detailed specification under Annexure-XII. Bidder may also submit additional certification as per international standard.
- e. Bidder must have valid manufacturing permission for non-drug item(s) where neither the Drugs & Cosmetic Act 1940 and Rules there under 1945 nor the Medical Device Rule 2017 is applicable. Bidder must submit an undertaking/Self declaration as per **Annexure-VII** in their letterhead that the item(s) quoted by them is/are non-drug item(s) i.e., neither covered under Drug & Cosmetic Act 1940 nor Under Medical Device Rule 2017.
- F) Bidder must have Market Standing Certificate of minimum three consecutive years (Latest) issued by the concerned Licensing Authority/Drugs Control Department/Concerned Government Department for the quoted product. In case quoted item is not covered under mandatory license regime under Drug & Cosmetic Act 1940 or Medical Device Rule 2017, Market standing Certificate (MSC) must be declared by the C.A./C.S. certifying at least three batch No. of the quoted items that the firm/company has manufactured and marketed the items for last three consecutive years. Self-attested copies are to be submitted.
- G) Declaration on company's letter head duly signed by authorized person stating that the firm & its quoted product is not blacklisted currently (as on the date of submission of the tender) by Central Government/ Central Government agencies/any State Government or any of the State Government agencies/ or any Drug procurement agencies or by PMBI in prescribed format as per Annexure-XV.

- H) Bidder must submit the Quality Management System (QMS) certificate issued by National Accreditation Board for Certification Bodies (NABCB), India. The QMS certificate should remain valid till the last date of submission of tender. Self-attested copies are to be submitted.
 - Note: a) If Manufacturing License/registration/permission for the quoted product(s) is issued under "for export only" category will not be accepted. Distributors/Suppliers/Marketer/Agents/Importer/Loan Licensee/Non-local suppliers are not eligible to participate in the Tenders. Liaison are strictly not permitted to bid on behalf of any manufacturers.
- J) Tenderer must declare their Maximum Production Capacity (item wise) for quoted item(s) highlighting it in Annexure XIV.
- K) Copies of the Audited Balance Sheet, Profit and Loss statement showing details of their annual average turnover for any three of the last four consecutive financial years i.e., 2018-19, 2019-20, 2020-21 and 2021-22 shall not be less than **2 Crores (Two crore).** Details shall be provided as per **Annexure IV**. Self-attested copies are to be submitted along with original copy of Annexure-IV.
- L) Declaration **on Nonjudicial Stamp Paper duly notarized** for eligibility in participating the tender for quoted items in prescribed format as per **Annexure-II.**
- M) Tenderer shall furnish Company's bank details as per Annexure V (Mandate Form).
- N) Tenderers are required to submit **Annexure-VI** indicating details of manufacturing License/permission and market standing certificate.
- O) Tenderer are required to submit declaration duly signed to supply the items as per the design in enclosure in **Annexure VII** as well as other instructions given in this regard.
- P) Duly attested Checklist as per (ANNEXURE- I) shall be submitted.
- Q) Copy of PAN Card of the bidder company should be submitted (self-attested).
- R) Copy of certificate of valid GST registration of the bidder company should be submitted (self-attested).
- S) Copy of Income Tax Return for any three of last four Consecutive Assessment years should be submitted (self-attested).
- T) Duly attested Copy of valid GS-1 registration certificate from GS1 India.
- U) Purchase preference shall be given to bidder(s) based on their declaration of the percentage (%) of minimum local content used in the manufacturing of quoted product as per Public Procurement (Preference to make in India), Order 2017 notification issued by GoI, Ministry of Commerce and Industry, Department of Industrial Policy and Promotion (DIPP) vide order no. P-45021/2/2017-PP (BE-II) dated 16.09.2020 and order no. 31026/65/2020-MD dated 30.12.2020 issued by Ministry of Chemicals & Fertilizers, Department of Pharmaceuticals and accordingly bidder(s) shall be categorized as per below table:

S. No.	Type of Class	Percentage (%) of minimum local content		
1	Class-I Local Supplier	Local content equal to more than 50%		
2	Class-II Local Supplier	Local content more than 20% but less than 50%		
3	Non-Local Supplier	Local content less than or equal to 20%		

The category of supplier based on the % of local content used against each quoted drug shall be mentioned in Annexure X in accordance with Order issued by Ministry of Chemicals & Fertilizers, Department of Pharmaceuticals vide no. order no. 31026/65/2020-MD dated 30.12.2020 and as per table mentioned above.

- **i. Purchase preference:** The 'margin of Purchase preference' means the maximum extent to which the price quoted by the "Class-I local supplier" above the L1 (landed cost).
- **ii.** "Local Content" means the amount of value added in India which shall be the total value of the item procured (excluding net domestic indirect taxes) minus the value of imported content in the item (including all customs duties) as a proportion of the total value, in percent.
- a) (i) If the participating Micro and Small Enterprises (MSE) meets all the other eligibility criteria and their quoting price is within price band of L1+15 (fifteen) per cent shall also be allowed to supply a portion of requirement by bringing down their price to L1 price in a situation where L1 price is from someone other than a MSE and such MSE shall be allowed to supply up to 25 (twenty-five) per cent of total tendered value. The 25 (twenty-five) per cent quantity is to be distributed proportionately among these bidders, in case there are more than one MSEs within such price band.
 - (ii) Within this 25% (Twenty-five Percent) quantity, a sub-target of 4% earmarked for procurement from MSEs owned by Scheduled Caste (SC)/Scheduled Tribe (ST) entrepreneurs (if they participate in the tender process and match the L1 price). Provided that, in event of failure of such SC/ST MSE to participate in tender process or meet tender requirements and L1 price, 4% sub-target shall be met from other MSE. MSEs would be treated as owned by SC/ST entrepreneurs: a) In case of proprietary MSE, proprietor(s) shall be SC /ST b) In case of partnership MSE, the SC/ST partners shall be holding at least 51% (fifty-one percent) shares in the unit c) In case of Private Limited Companies, at least 51% (fifty-one percent) share shall be held by SC/ST promoters.
 - (iii)Within this 25% (Twenty-five Percent) quantity, a sub-target of 3% earmarked for procurement from MSEs owned by Women entrepreneurs (if they participate in the tender process and match the L1 price). Provided that, in event of failure of such Women entrepreneurs MSE to participate in tender process or meet tender requirements and L1 price, 3% sub-target shall be met from other MSE.

Note: -

- (i) The certificates/ reports / annexure submitted with the bid document should be self-attested by the authorized signatory of the firm with official seal, wherever required.
- ii) Technical evaluation of the Bid will be done on the basis of the above-mentioned criteria and documents mentioned in Clause no.3 (Technical Bid- Cover 'A') Mandatory Documents shall be submitted online only at CPP portal: https://eprocure.gov.in Failing which the bid will not be considered for technical evaluation.
- iii) Hard copy of required documents uploaded shall be submitted along with Earnest Money Deposit (EMD) and other required documents on or before the last day of submission of tender for purely evaluation purposes. However, the submission of hard copy of uploaded tender document submitted shall not substitute/modify the provisions of e-tendering system.
- iv) The technical evaluation shall be done only on the basis of documents/papers submitted by the bidder on https://eprocure.gov.in.
- v) Clear copy of valid manufacturing license highlighting the item code should be uploaded. In case scanned copy of license uploaded is not visible or tempered, PMBI shall not considered the license for such items.
- V) If the procurement for a value is more than Rs. 10 crores, the Class-I Local Supplier / Class-II Local Supplier shall be required to provide a certificate from the statutory auditor or cost auditor of the company (in the case of companies) or from a practicing cost accountant or practicing chartered accountant (in respect of suppliers other than companies) giving the percentage of local content as per pt. no. 9.b of DPIIT order no. P-45021/2/2017-PP(BE-II) dated 16.09.2020.

W) Special Terms & Conditions:

- i. Bidder must quote the product as per specification provided in Annexure XII and other physical parameters.
- ii. Catalog must be attached with the bid for technical evaluation along with dimension of quoted items.
- iii. The supplier may be asked to arranging demonstration of their equipment for which rates have been quoted, to the PMBI, if required. The expenditure incurred for demonstrating the items will be borne by the supplier.
- iv. Bidder shall have dedicated Customer Care Support Team for providing technical assistant at consumer level.
- v. Directive 2011/65/EU Restriction of Hazardous Substances Directive (ROHS) to be complied.
- vi. Bidders has to oblige and fulfill the warranty of the product at point of sale i.e., at kendra level for each item wherever specified in Annexure XII and XII (A).

3.1 ELIGIBILITY OF BIDDERS FROM SPECIFIED COUNTRIES:

Compliance under Rule 144 (xi) of the General Financial Rules (GFRs), 2017 and ORDER NO: F. No. 6/18/2019-PPD DATED 23/07/2020 & 24/07/2020 issued by Department of Expenditure (MoF) restricting procurement from bidders from certain countries that share a land border with India shall apply to this procurement.

- 1. GoI vide Order (Public Procurement No.1, 2 & 3) dated 23 Jul 2020 & 24 Jul 2020 has imposed Restrictions under Rule 144 (xi) of the General Financial Rules (GFRs), 2017 on bidders from a country which shares a land border with India.
- 2. "Bidder" for the purpose of this Order (Public Procurement No.1, 2 & 3) (including the term 'tenderer', 'consultant' 'vendor' or 'service provider' in certain contexts) means any person or firm or company, including any member of a consortium or joint venture (that is an association of several persons, or firms or companies), every artificial juridical person not falling in any of the descriptions of bidders stated hereinbefore, including any agency, branch or office controlled by such person, participating in a procurement process.
- 3. "Bidder from a country which shares a land border with India" for the purpose of this Order (Public Procurement No.1, 2 & 3) means;
- a. An entity incorporated, established or registered in such a country; or
- b. A subsidiary of an entity incorporated, established or registered in such a country; or
- c. An entity substantially controlled through entities incorporated, established or registered in such acountry; or
- d. An entity whose beneficial owner is situated in such a country; or
- e. An Indian (or other) agent of such an entity; or
- f. A natural person who is a citizen of such a country; or
- g. A consortium or joint venture where any member of the consortium or joint venture falls under any ofthe above

Note: "Beneficial owner" for the purpose of above paragraph (3) will be as under:

i. In case of a company or Limited Liability Partnership, the beneficial owner is the natural person(s), who, whether acting alone or together, or through one or more juridical person(s), has a controlling ownership interest or who exercises control through other means.

Explanation

- a) "Controlling ownership interest" means ownership of, or entitlement to, more than twenty-fiveper cent of shares or capital or profits of the company;
- b) "Control" shall include the right to appoint the majority of the directors or to control the management or policy decisions, including by virtue of their shareholding or management rights or shareholder's agreements or voting agreements;

- ii. In case of a partnership firm, the beneficial owner is the natural person(s) who, whether acting alone or together, or through one or more juridical person, has ownership of entitlement to more than fifteen percent of capital or profits of the partnership;
- **iii.** In case of an unincorporated association or body of individuals, the beneficial owner is the natural person(s), who, whether acting alone or together, or through one or more juridical person, has ownership of or entitlement to more than fifteen percent of the property or capital or profits of such association or **body of individuals.**
- iv. Where no natural person is identified under (i) or (ii) or (iii) above, the beneficial owner is the relevant natural person who holds the position of senior managing official.
- v. In case of a trust, the identification of beneficial owner(s) shall include identification of the author of the trust, the trustee, the beneficiaries with fifteen percent or more interest in the trust and any other natural person exercising ultimate effective control over the trust through a chain of control or ownership.
- vi. "Agent" for the purpose of this Order (Public Procurement No.1, 2 & 3) dated 23 Jul 2020 & 24 Jul 2020 is a person employed to do any act for another, or to represent another in dealings with third persons.

Rule: Following shall be complied by the Bidders of the said countries while submitting bids.

- A. Any bidder from a country who shares a land border with India will be eligible to bid in this tender only if the bidder is registered with the <u>Competent Authority</u>. The Competent Authority for the purpose of registration under this Order shall be the Registration Committee constituted by the <u>Department for Promotion of Industry and Internal Trade (DPIIT)</u>. However, Order will not apply to bidders from those countries (even if sharing a land border with India) to which the Government of India has extended lines of credit or in which the Government of India is engaged in development projects. Lists of countries to which lines of credit have been extended or in which development projects are undertaken are given in the website of the Ministry of External Affairs.
- B. The Bidder shall have to submit declaration / certificate as per the attached Format towards compliance of Public Order on Restrictions under Rule 144 (xi) of the General Financial Rules (GFRs), 2017. B). Further as per above the format of declaration is added as Annexure XVII. It shall be furnished by the bidder(s) on duly notarized Non-Judicial Paper.

4. GENERAL CONDITIONS:

- A) Tender bid is invited directly from Manufacturers in India.
 - Loan licensee / Distributors / Agents / Contract Manufacturers / Importers / Non-Local supplier are not eligible to participate in the tender.
- B) Manufacturer has Production & financial capacity to manufacture and deliver the items quoted by the firm in the tender as per quantity mentioned in tender during contract period.
- C) Bidders are advised to quote such items only for which they meet the item specification as mentioned in Annexure XII of the tender document. Do not quote if it differs about any parameter. Bidder(s) shall also submit declaration as per Annexure XIV.
- **D)** The quantities specified in the tender is for the tender purpose only and it represents the basis of unit for ease of pricing. The actual quantity may vary from zero to the maximum required quantity during the contract. The quantity will be drawn from successful tenderers as and when required from time to time during the contract period.
- **E)** STP (Standard Testing Procedure) for the awarded items are required to be submitted within 15 days from the date of issue of Letter of Acceptance.

- F) The manufacturer shall declare the material used in manufacturing against all quoted items and declare that it is internationally accepted when ask by PMBI.
- G) The Tenderer should have not been blacklisted/debarred/de-registered/banned due to quality failure for the quoted product /firm by any State Government / Central Government/ PMBI/Central or State Government's Drug procurement agencies at the time of submission of bid. Further, quoted items have not been failed in inhouse testing or testing by any State Government/Central Government / its Drug procurement agencies/PMBI during last two years. If any tenderer blacklisted/debarred/de-registered/banned due quality failure, such to tenderer their Partner/Director/Owner shall not be permitted to participate in the tender.
- **H)** During the validity of the tender if the firm / Company is blacklisted/debarred/de-registered/banned by any State Government/ Central Government/ PMBI/ Central or State Government's Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to PMBI along with relevant authentic document by the tenderer firm/ company within one month.
- I) During tender or Rate Contract period, if L1 bidder is debarred/deregistered /blacklisted/ banned by any Central Government or State Government or its procurement agencies due to quality failure, PMBI may purchase the items from other bidders at L1 rate or may go for fresh tender as per discretion of PMBI.
- **J**) The PMBI reserves the right to purchase any items from PSUs as per discretion of PMBI. In case of emergencies, PMBI may go to PSUs and price will be as per negotiation and at the discretion of PMBI.
- **K**) The Tenderer should confirm that they have read tender document including Amendment(s) to Tender document (if any) along with terms and condition and these terms and condition of tender document including Amendment(s) to Tender document (if any) are acceptable unconditionally to them.
- **L) Validity of Rate Contract:** -The rate contract will be applicable for 2(two) year from the date of acceptance of LOA. The validity of contract may be extended with mutual consent for some specified period to the maximum of 1 (one) year by PMBI, if necessary.
- M) During the contract period at any stage, it is found that the tender has been successfully obtained by the bidder by submitting forged/ fabricated certificates/ documents/ licenses and/or by concealing the fact about blacklisting/ debarring/ de-registration of the firm by Govt. of India/ Suspension/ Cancellation/ non-renewal of the manufacturing license of the bidder firm, the tender bid/ rate contract may be rejected/ terminated and suitable punitive/ legal action may be taken as per the tender terms and condition and in addition to penal action recovery shall be made (if any) against the firm.
- N) If a company/firm has two or more separate manufacturing units at different sites / States, which are not separate entities then the company will be allowed to submit only one tender for all units but should be declare in Annexure II and necessary document regarding separate manufacturing units will be uploaded as a separate set with the same tender. However, one bidder will be allowed to submit only one offer for one product.
- **O**) Only authorized employee of the Company/Tenderer will be allowed to transact the business with the Tender Inviting Authority.

P) Samples:

i) Three (3) properly labeled samples shall be submitted against each quoted item as per the specification mentioned in Annexure XII of the tender document on or before stipulated date. Failure to do so, it shall entail your quotation being disqualified.

- ii) Sealed sample boxes should contain samples the tenderer quoted for along with duly filled Annexure XVIII.
- iii) Sample should be in the form of pack as specified in tender enquiry, otherwise the quotation against that particular item is liable to be rejected.
- iv) Firm may take back their samples if unapproved/ technically disqualified within 10 days from the date of technical disqualification on CPP Portal. Any claim made thereafter shall not be entertained and PMBI reserves all right to dispose the samples as per the discretion.

5. PRICE BID – "COVER-B" (Financial Bid/BOQ)

A) Cover "B" (Financial Bid/BOQ) contains the Price Bid of the Tenderer. The Tenderer shall fill in the rate per unit size, % age rate of GST in respective column of BOQ for the items quoted.

B) Determination of L1 Bidder:

- a) In determining the lowest evaluated price, the rate quoted per unit size for the given specification, exclusive of GST as indicated in column No. 7 of the BOQ shall be taken into consideration. The rates quoted should be in rupees and paisa up to 2 digits. The Tenderer is not permitted to change/alter specification or unit size given in the ANNEXURE-XII.
- b) GST (Goods and Services Tax)-The Tenderers must indicate the rate of GST applicable and payable by them. In case no information is given, it shall be presumed that rate is inclusive of GST and no GST shall be charged by them under any circumstances.
- b) The bidder is required to indicate rate of GST (%) as digit only in column 9 of BOQ without suffixing the % sign and not to indicate amount of GST in Rs. at particular cell of excel sheet of BOQ.
- c) Purchase preference shall be given over acceptable L1 bidder to bidder offering Products manufactured by using higher % age of Local Content computed on the basis of cost of domestic contents in order to promote "Make in India" subject to matching of acceptable L1 rate as per Public Procurement (Preference to make in India), order 2017.
- d) (i) If the participating Micro and Small Enterprises (MSEs) meets all the other eligibility criteria and their quoted price is within price band of L1+15 (fifteen) per cent shall also be allowed to supply a portion of requirement by bringing down their price to L1 price in a situation where L1 price is from someone other than a MSEs and such MSEs shall be allowed to supply up to 25 (twenty-five) per cent of total tendered value. The 25 (twenty-five) per cent quantity is to be distributed proportionately among these bidders, in case there are more than one MSEs within such price band.
- (iv) Within this 25% (Twenty-five Percent) quantity, a purchase preference of four per cent that is, 25 (twenty-five) per cent out of 25 (twenty-five) per cent will be reserved for MSEs owned by Scheduled Caste (SC)/Scheduled Tribe (ST) entrepreneurs (if they participate in the tender process and match the L1 price) provided that in event of failure of such SC/ST MSEs to participate in tender process or meet tender requirements and L1 price, four per cent sub-target shall be met from other MSEs. MSEs would be treated as owned by SC/ST entrepreneurs: a) In case of proprietary MSEs, proprietor(s) shall be SC/ST b) In case of partnership MSEs, the SC/ST partners shall be holding at least 51% (fifty-one percent) shares in the unit c) In case of Private Limited Companies, at least 51% (fifty-one percent) share shall be held by SC/ST promoters.
- (v) Within this 25% (Twenty-five Percent) quantity, a sub-target of 3% earmarked for procurement from MSEs owned by Women entrepreneurs (if they participate in the tender process and match the L1 price). Provided that, in event of failure of such Women entrepreneurs MSE to participate in tender process or meet tender requirements and L1 price, 3% sub-target shall be met from other MSEs.

6. Earnest Money Deposit (EMD)/Bid Security:

A) The Earnest Money Deposit referred to under Clause 3.A, shall be Rs. 100000.00 (One lakhs). The Earnest Money Deposit shall be paid in the form of Bank Guarantee or National Electronic Fund Transfer (NEFT) or Bankers Cheque/ Demand Draft in favour of Pharmaceuticals & Medical Devices Bureau of India, payable at Delhi. EMD in form of Bank Guarantee, Irrevocable Bank Guarantee in favour of Pharmaceuticals & Medical Devices Bureau of India, from any Nationalized/scheduled Bank should be valid for a period of 12 months from the date of tender opening. The format of Bank Guarantee is at ANNEXURE-X. PMBI will not pay interest on any deposit held in the form of Bankers Cheque or Demand Draft or Electronic Fund Transfer.

Account Details for National Electronic Fund Transfer (NEFT):

Bank Name: Bank of Baroda, Account No. 05860200001696, IFSC Code: BARB0PARLIA

- B) Tenderer may be exempted from the payment of EMD, if valid **registration** certificate from NSIC/MSME is uploaded **for the product for which bidder has submitted quotation.**
- C) PSUs are exempted from the payment of EMD.
- D) The tender submitted without sufficient EMD will be summarily rejected.
- E) Non-payment of EMD (except in cases where payment of EMD is specifically exempted) will result in rejection of the bid.
- F) The Earnest Money Deposit will be refunded to the successful bidders after successful completion of first supply.
- G) The Earnest Money Deposit of the Tender will be forfeited without further notice if:
 - a) If the tenderer withdraws his bid any time after opening of price bid.
 - b) On refusal to supply surgical & consumables after the award of contract/Letter of Acceptance (LOA).
 - c) In case of the lowest bidder (L1 bidder), fails to execute the contract or fails to complete the first supply successfully within the stipulated time.
 - d) If the undertaking as Annexure II is not found correct at any stage during the contract period.

7. GUIDELINES FOR THE PREPARATION OF TENDER:

- A) The bidder shall bear all costs associated with the preparation and submission of its bid and Tender Inviting Authority will in no case be responsible or liable for these costs, regardless of the conduct or outcome of the bidding process.
- B) Language of Bid: The Bid prepared by the bidder and all correspondence and documents relating to the bid exchanged by the bidder and the Tender Inviting Authority, shall be in English language, Supporting documents furnished by the bidder may be in other languages provided they are accompanied by an authenticated (by the authority concerned) accurate translation of the relevant passages in the English language in which case, for purposes of interpretation of the Bid, the English translation shall alone govern. Failure to submit authentic translation of documents would result in rejection of bids. No bid can be partly in one language and partly in another language.
- C) Power of Attorney or Resolution of the Board by which the authorized signatory has been authorized by the bidder firm should sign the documents in cases where person other than the Managing Director/Managing Partner or sole Proprietor signs the document.

8. PERIOD OF VALIDITY OF TENDER:

- **a)** The tender must remain valid for minimum 150 days from the date of opening of Technical Bid. (As mentioned in Clause 1.ii)
- **b**) Prior to the expiration of the bid validity the Tender Inviting Authority may extend the bid validity for further period with mutual consent of the bidder.
- c) The bidder who has extended the bid validity is not required or permitted to modify its bid.

d) The bidder cannot withdraw the bid within validity of Tender.

9. AMENDMENT OF TENDER DOCUMENTS:

At any time prior to the last date of submission of online bid, Tender Inviting Authority may, for any reason, whether on own initiative or in response to a clarification requested by a prospective Tenderer, may modify the condition in Tender documents by uploading an amendment on PMBI website: www.janaushadhi.gov.in; and on CPP portal i.e., https://eprocure.gov.in will be binding on them. In order to provide reasonable time to take the amendment into account in preparing their bid, Tender Inviting Authority may at discretion, extend the date and time for submission of online bid.

- A) Bidders are advised to check the *website of PMBI*: <u>www.janaushadhi.gov.in</u>; and CPP Portal i.e., <u>https://eprocure.gov.in</u>; *regularly* at least 3 days prior to closing date of submission of tender for any corrigendum or amendment to the tender document.
- B) PMBI will not issue separate communication for any corrigendum or amendment.

10. METHOD OF SUBMISSION OF TENDER:

- A) The tender document shall be downloaded from the websites <u>www.janaushadhi.gov.in</u>; and CPP portal i.e., https://eprocure.gov.in. Tender Document is free of cost. No tender cost is to be deposited
- B) Bids shall be submitted online only at CPP Portal i.e., https://eprocure.gov.in; Manual bids shall not be accepted except for the original documents/instruments as mentioned in Clause 3 of tender document.
- C) Bidders are advised to follow the 'Special Instructions to the Contractors/Bidders for the esubmission of the bids online' available through the link 'Help for Contractors' at the e-Procurement Portal https://eprocure.gov.in.
- D) If a particular document/Certificate to be uploaded as specified in bid, if not applicable for a bidder, the bidder shall attach a scanned copy of declaration in the letter head stating that the specific document is not applicable/exempted for the bidder in connection to this tender.
- E) Bidder shall not modify the downloaded tender form including downloaded price Bid template in any manner. In case any tender form/Price bid template is found to be tampered with/modified in any manner, such bid will be summarily rejected, Bid Security would be forfeited, and bidder is liable to be banned from doing business with PMBI.
- F) Interested eligible Tenderer may obtain further information in this regard from the office of the Tender Inviting Authority on all working days between 10:00 AM and 5:00 PM.
- G) Once the bid have been uploaded in the CPP Portal https://eprocure.gov.in the bidder has to make sure that he has uploaded the files in the correct format and the bidder has to download the uploaded files from their own end and has to check whether the files uploaded is in proper format or not, no corrupted files have to be uploaded.
- H) Bidder shall not wait till the last time for the submission of bid on CPP portal. In any case, if bidder fails to submit the bid online, PMBI will not be responsible.

Note: In any case if the prospective bidder fails to uploaded technical bid (Cover A)/ Price bid (Cover B) successfully on the CPP Portal online on or before the last date and time of submission of technical bids, the bid shall be summarily rejected without considering any facts.

11. MODIFICATION AND WITHDRAWAL OF BIDS:

- A) The bidder may modify or withdraw its bid after the bid submission before last time and date of submission of online Technical Bid.
- B) No bid will be allowed to be withdrawn after the last date & time of submission of online Technical Bids.

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Note: Any reason (whatsoever it may be except the Force Majeure events condition defined in the tender document or by Government) for withdrawal of bid or modification in bid or any request pertaining to have quoted wrong rates for any unit size instead of the unit size in tender document/BOQ shall not be considered. Bidder who are making such request shall be penalized as per tender clause no. 27 terms and conditions besides blacklisting for a duration of not less than three (3) years.

12. OPENING OF TENDER:

- A) The opening of the Technical Bid and the Price Bid will be done online as specified. The date of technical bid opening is published in advance. The date of opening of price bid will be announced only after the opening and evaluation of technical bid. The bidder who are found eligible and on satisfying the criteria for technical evaluation/based on undertakings & Declaration, will only be informed the time and date of opening of Price Bid Cover "B" of the tender.
- B) Only authorized employee of tenderer is entitled to be present at the time of opening of Technical Bid Cover "A" of the tender submitted by them.
- C) In case, the date for opening of technical bid is declared holiday, the technical bid shall be opened on next working day at 11.30 A.M.
- D) The original/attested hard copies (as mentioned in Clause no. 3, eligibility criteria) must reach the PMBI Head office on or before stipulated date and time, failing which the bid shall be summarily rejected without considering any fact.

13. EVALUATION OF TENDER:

- E) Technical evaluation of the Bid will be done on the basis of criteria and documents mentioned in clause no. 3 (TECHNICAL BID-COVER A) & Annexure I (Check List) which are present in the CPP Portal i.e., https://eprocure.gov.in.
- F) Bids of firms who have furnished all the required documents for each of the product quoted will be considered.
- G) If at any stage, it is found that the contract has been successfully obtained by the bidder by submitting forged/fabricated certificates/documents/licenses and/or by concealing the fact about blacklisting/debarring/de-registration of the firm by Govt. of India/Suspension/Cancellation/non-renewal of the manufacturing license of the bidder firm, the tender bid/rate contract may be rejected/terminated and suitable punitive action may be taken against the firm.
- H) In event of financial bid opening, due to provisions/compulsion of e-tendering system if complete quoted product list of financial bids of a bidder is opened then only those financial bids of quoted product shall be considered of whose technical bid has been found eligible by the Technical Evaluation Committee.
- I) After evaluation of technical bid of tenderer/bidder, PMBI may ask the objection/clarification from tenderer/bidder.

14. INSPECTION OF MANUFACTURING FACILITIES:

- A) PMBI or its authorized representative(s) has/have the right to inspect the manufacturing premises of Tenderers, before accepting the rate quoted by them or before releasing any purchase order(s) or at any point of time during the continuance of tender and also has the right to reject the tender or terminate/cancel the purchase orders issued and/or not to place further order, based on adverse reports brought out during such inspections. Copy of one full set of the documents submitted for the bid should be made available at the time of inspection.
- B) Originals of all the documents uploaded/submitted in the Technical Bids should be produced for verification during Site inspection and Physical Verification.

15. ACCEPTANCE /REJECTION OF BIDS:

- A) PMBI reserves the right to accept or reject the tender for the supply of all or any one or more items of the items tendered for in a tender without assigning any reason.
- B) Evaluation of the tender and determination of the L1 rate (Lowest rate) will be done based on rate per unit size exclusive of GST as mentioned in column 7 of **BOQ.** PMBI shall have the right to call other eligible bidders those are willing to match L1 rates. If such firms are found, then the order quantity may be dispersed in ratio of: -
- a) "Minimum 50% of the tender quantity may be awarded to the qualified bidder(s) falling under Class I local supplier category, if qualified for award of contract and/or subject to the matching of L1 price for quoted drugs at the discretion of PMBI and remaining 50% of quantity may be awarded to the eligible bidder following the guidelines and respective clauses of DPIIT order no. P-45021/2/2017-PP(BE-II) dated 16.09.2020."

The following possible cases may be considered for the award of contract;

- Case-I: If L1 is Class I local supplier, minimum 50% quantity shall be given to L1 bidder, 25% shall be given to MSEs (if comes within the price band (of L1 + 15%) & qualify) and remaining 25% shall be given to other eligible bidders (if comes within the Margin of Price Preference & qualify).
- Case-II: If L1 is Class-II local supplier, as per PPE-MSE order, initially 25% shall be reserved for MSEs (if comes within the price band (of L1 + 15%) & qualify). Thereafter, preference shall be given to Class-I local supplier to award 50% of tender quantity and at last, if quantity remains balance, 25% quantity shall be given to Class-II L1 bidder following the guidelines and respective clauses of DPIIT and MSME.
- C) However, in case the price quoted by the lowest responsive tenderer (L1) is not reasonable and unacceptable, the price may be negotiated with L1 only as per CVC guidelines and, if it reduces the price to the desirable level, rate contract may be concluded with L1. To meet the demand, PMBI shall conclude parallel rate contract by counter offering the L1 rate to higher eligible bidders as per above provision.
- D) Negotiation if required will be done strictly as per Central Vigilance Commission guidelines.
- E) Letter of acceptance of tenders for Rate Contract will be communicated to the Tenderers in writing as per **ANNEXURE XI.**
- F) **Purchase preference:** The margin of Purchase preference shall be 20%.

16. AWARD OF CONTRACT:

A) The contract will be awarded to the lowest evaluated responsive bidder qualifying to the final round after Technical, Price Bid evaluation will be done as per the clause 5. B.

"Minimum 50% of the tender quantity may be awarded to the qualified bidder(s) falling under Class I local supplier category, if qualified for award of contract and/or subject to the matching of L1 price for quoted drugs at the discretion of PMBI. PMBI would follow the guidelines and respective clauses of DPIIT order no. P-45021/2/2017-PP(BE-II) dated 16.09.2020" as mentioned in clause 15 (B).

B) Letter of Acceptance:

The Tender Inviting Authority shall issue Letter of Acceptance (LOA) as per Annexure-XI to the lowest responsive bidder in respect of the items selected. Communication by e-mail / fax / letter will be deemed as valid communication.

- C) The successful bidder, upon receipt of the Letter of Acceptance (LOA), shall communicate the acceptance of the same to the PMBI and shall furnish the documents, asked if any.
- **D**) The bidder shall not, at any time, assign, sub-let or make over the contract or the benefit thereof or any part thereof to any person or persons whatsoever. Such practices will be deemed as fraudulent practices and also as breach of terms of contract and shall invite punitive action.

17. PERFORMANCE SECURITY DEPOSIT:

- **A)** On being informed about the acceptance of the tender for Rate Contract, the Performance Security Deposit @ 3% will be deducted from each running bills against the total value in the purchase order and accumulated security deposit will be refunded without any interest by PMBI to the tenderer within 60 days following the date of completion of tenderers performance obligations under the contract including the shelf-life obligation.
- B) The Security deposit of supplier will be returned by PMBI only after the supplier has given undertaking to replace such items and indemnify PMBI against any losses on account of quality parameters duly notarized on non-judicial paper.

18. METHODOLOGY FOR PLACING ORDERS:

For the above purpose, the following procedures will be adopted:

- A) After the conclusion of Price Bid opening (Cover B), the rates offered by tenderers for each product are evaluated and lowest acceptable rate (L1 Rate) arrived at is declared and that tenderer is informed.
- B) PMBI reserves right to issue purchase order for any item on any one rate contract holder or more than one rate contract holder for same items.
- C) If two or more than two Tenderer's are declared as lowest suppliers for the same item(s), such Tenderers are eligible for Rate Contract and the placement of Purchase Orders for such item(s) for which they are declared as lowest. L1 quantity will be distributed equally among them as per clause no. 16.A.
- D) The supplier shall supply the Items to any or all the Warehouse (Address/Location) as mentioned in clause 19 (A) or any other place decided by PMBI and supply shall confirm to the conditions mentioned in the provision of tender documents, viz, logo, nomenclature, specification etc. within the stipulated period.
- E) Once The supplier shall supply the Items at any of the PMBI Warehouse as mentioned in purchase order (or any other place decided by PMBI) along with copy of Purchase order, copy of test reports and 3 original copies of Invoice. No payment will be processed without test reports.
- F) A purchase order is placed on supplier for supply of definite quantity in terms of Rate Contract during validity period of Rate Contract that purchase order is valid and binding contract.
- G) No Minimum drawl is in the Rate Contract. The actual quantity may vary from nil to maximum required quantity during validity of Rate Contract.
- H)The Items supplied in excess of the ordered quantity shall not be accepted and the supplier shall take back the excess at their cost. PMBI will not be responsible for the loss to the supplier and will not entertain any demand/claim.
- I) The purchaser reserves the right to conclude one or more than one rate contract for the same item.
- J) The purchaser has the option to renegotiate the price with the rate contract holders. In case of emergency, the purchaser may purchase the same item through Ad hoc contract with a new supplier.
- K)Purchase orders, incorporating definite quantity of items/products to be supplied along with all other required conditions following the rate contract terms, shall be issued for obtaining supplies through the rate contract.

- L) The purchaser is entitled to place purchase orders up to the last day of the validity of the rate contract and, though supplies against such purchase orders will be affected beyond the validity period of the rate contract, all such supplies will be guided by the terms & conditions of the rate contract.
- M) The details of the required items, medical devices, etc. are shown in **ANNEXURE -XII**. The tender quantity mentioned herein is not a fixed procurement quantity and it is only a tentative requirement and may be increased or decreased by the PMBI at its discretion depending on its actual need. Though the tentative quantity is indicated in the Rate Contract, the PMBI, will confirm the actual requirement through purchase order/orders from time to time. The tenderers shall supply the items only on the basis of the purchase order issued time to time within validity of Rate contract period by the PMBI. Any supply without a valid purchase order will not be acceptable to PMBI and the PMBI shall not be responsible for any loss on this account.
- N)However, once the purchase order/orders is/are issued by the PMBI, the tenderer shall not renege from the commitment of supplying the quantity mentioned in the acceptance of tender for Rate Contract.
- O) The rates quoted shall not be varied with the Purchase order quantity during the full contract period.
- P) The rates quoted and accepted will be binding on the Tenderer for the full contract period of two years and any increase in the price will not be entertained till the completion of this contract period. Accordingly, this clause will be applicable for all orders placed during the contract period. However, Rate Contract validity period may be extended for period up to further one year at same rate, terms & conditions with the consent of the supplier.
- Q)No Tenderer shall be allowed at any time and on any ground, whatsoever it may be, to claim revision or modification in the rates quoted by them. Representation to make correction in the tender documents on the ground of Clerical error, typographical error, etc., committed by the Tenderers in the Bids shall not be entertained after submission of the tenders. Cross Conditions such as "SUBJECT TO AVAILABILITY", "SUPPLIES WILL BE MADE AS AND WHEN SUPPLIES ARE RECEIVED" etc., will not be entertained under any circumstances and the tenders of those who have mentioned such conditions shall be treated as incomplete and accordingly the Tender will be summarily rejected.
- R) Supplies should be made directly by the tenderer and not through any other Agency / Dealer / Distributors.
- S) The supplier shall take utmost care in supplying the quality Items and ensure that the batch number mentioned in the packages of the Items tally with the batch number mentioned in the Invoice produced to PMBI for payment. Also, the supplier shall ensure the quantity relevant to the Batch Number of the Items is mentioned in the invoice. Items to be supplied of any batch shall not be accepted with different MRP.
- T) "MRP inclusive of all taxes" is to be printed on each unit/label. MRP will be intimated to successful bidders at the time of placing purchase orders.
- U)The Rate Contract (RC) awarded under the present tender enquiry will be in the nature of standing offer. Purchase Order (PO) may be placed from time to time against Rate Contract (RC).

V)FALL CLAUSE:

If at any time during the execution of the contract, the controlled price becomes lower or the supplier reduces the sale price or sells or offers to sell such stores, as are covered under the contract, to any person / organization including the purchaser or any department of Central government/state Govt. or its procurement agencies at a price lower than the price chargeable under the contract, he shall forthwith notify such reduction or sale or offer of sale to the purchaser and the price payable under the contract for the stores supplied after the date of coming into force of such reduction or sale or offer of sale shall stand correspondingly reduced.

W) Subject to the conditions mentioned in the Purchase Order, Tender Document, Price Agreement/letter of Acceptance and here under, the Supplier is entitled for the payment against

supply. In case of any discrepancy in levy of LD, Penalty, Unexecuted Fine, Short Passing of Bills, such discrepancy shall be intimated within 60 days from the date of receipt of payment, thereafter PMBI will not entertain any claim.

NOTE: PMBI do not give any guarantee of minimum purchase under this Rate Contract.

19. SUPPLY CONDITIONS:

- A) Purchase orders will be issued to the Tenderer(s) at the discretion of the PMBI as per actual requirements. All the supplies shall be received at any or all of the following warehouse of PMBI or any other place decided by PMBI:
 - i) Central Warehouse, Gurugram, (Pharmaceuticals & Medical Devices Bureau of India (PMBI) Sugal Logistic Park, Warehouse No.1, Opp. GITM College, Bilaspur-Tauru Road Village Bilaspur and Khasra No. 60//14/2, 17,24,6,15, 16, 25, 7/1, 14/1, 61//9, 10, 11,62//3/2, 4,10//17, 24, 19//3, 8/2, 9/1/1, 12/2/2/2 min 13/1/1 min. Pin Code 122413 Phone No. 011-49431800
 - ii) Regional Warehouse, Guwahati, (Pharmaceuticals & Medical Devices Bureau of India (PMBI)
 DAG No. 884 of K P PATTA No. 04, Mughuapara, Pamohi Village, Dist. Kamrup (M)
 Guwahati, Assam India 781035.
 Phone No. 011-49431800
 - iii) Regional Warehouse, Chennai, Pharmaceuticals & Medical Devices Bureau of India (PMBI) 79, KIZHMUTHALAMPEDU, PANAPAKKAM, City Tiruvallur, State Tamil Nadu Pin Code 601201 Phone No. 011-49431800
 - iv) Regional Warehouse, Surat, Pharmaceuticals & Medical Devices Bureau of India (PMBI).
 Plot no. A-23/2 & A -24/1,
 Ichhapore Bhatpore GIDC, Ichhapore
 Surat, Gujarat 394510
 - v) Newly Proposed Warehouse:
 Regional Warehouse, Bengaluru, Pharmaceuticals & Medical Devices Bureau of India (PMBI).
 Plot No 162 163, KIADB Industrial Area, Hi Tech Defense Aerospace Park, Devanahalli, Bengaluru, Bengaluru Rural, Karnataka -562110

2 days from the receipt of nurchase orders the Tenderer should inform PMPI through mail

- B) Within 3 days from the receipt of purchase orders the Tenderer should inform PMBI through **mail** about the confirmation for the receipt of the purchase order.
- C) The Tenderer should also fill the details of supply/delivery schedule to PMBI through **PMBI vendor portal** within 7 days from the receipt of the purchase order with expected dispatch/supply date.
 - i. The bidder shall have to fill Advance supply notice (ASN) on **PMBI vendor portal** with all other details i.e., invoice copy, Certificate of Analysis (COA), Batch no. Quantity, Date of Manufacturing (DOM) Date of Expiry (DOE), no. of shipper boxes etc.
 - ii. Once the ASN is accepted by the PMBI, the bidder will be provided the date to execute the supplies at PMBI warehouse as mentioned in purchase order.

Note:

- In case, the supply shall not be made by the date as conveyed by the supplier, supply order shall be cancelled at their risk and cost. If no response is received through **PMBI vendor portal** within 7 days from the supplier / tenderer about supply of items as per purchase order, it shall be presumed that the supplier/tenderer is not interested to supply the items ordered as per purchase order and PMBI shall purchase the items from alternative sources.
- In case of newly awarded bidder, bidder must share their permanent email ID and phone number for **PMBI** vendor portal registration to it1@janaushadhi.gov.in and customercare1@janaushadhi.gov.in.

D) The supplier must supply the ordered quantity as follow delivery schedule mentioned below:

Sl. No.	Nature of Product	Delivery Schedule (Days)
NO.		Schedule (Days)
1	Delivery Schedule against first and subsequent P.O. for all tendered	45 days
	items i.e., Surgical/ Consumables and Medical Devices etc.	

- E) If the delivery date happened to be a holiday for PMBI, the supply should be completed by 5.00 PM on the next working day.
- F) In case of non-execution of the order, PMBI reserves the right to place purchase orders (partially/fully) on alternate source at the risk and cost of the default tenderer(s) without any notice/Information.
- G) If a supplier fails to execute supply as per Purchase Order, the 5% of value of unexecuted quantity of Purchase Order shall be recovered from pending bill or Bank Guarantee/Performance security deposit and their bad performance shall be kept in record of PMBI for future dealing as considered appropriate by PMBI.
- H) If the Tenderer fails to execute the supply within the stipulated time, the PMBI is at liberty to make alternative arrangement for purchase of the items for which the Purchase orders have been placed, from any other sources or in the open market or from any other Tenderer who might have quoted higher rates, at the risk and the cost of the defaulted supplier and in such cases the PMBI has every right to recover the cost and impose Liquidated Damages as mentioned in Clause 25.
- I) The liquidated damages as specified in clause 25(B) of the tender conditions will be levied on the quantity supplied after the schedule as mentions in Clause 19.(D) from the date of issue of purchase order. However, no supplies will be accepted after 30 days of the expiry of delivery date i.e., completion of specified liquidated damages period as per clause 25(B), the purchase order shall be cancelled at the risk and cost of the supplier. However, the supplier must take prior approval from PMBI for supply of items beyond stipulated delivery period in Purchase order.
- J) Supplier shall complete the earliest pending purchase order before commencing the supply of subsequent purchase orders. Further, supplies against a purchase order are to be made in minimum numbers of batches as far as possible and same batch should not be supplied in repeated consignment.
- **K**) Bidder must supply the items with minimum 36 months shelf life. Bidders must declare the required shelf-life detail in **Annexure XIII.**
- L) The Tenderer must submit an Analysis report for every batch of supplied product along with invoice. In case of failure on part of the supplier to furnish such report, the batch of items will be returned to the suppliers, and he is bound to replenish the same with Govt. approved lab test report.

M) Tenderer should supply the product as follow:

- (i) Within 2 months excluding month of manufacture of products having shelf life up to 2 years,
- (ii) Within 3 months excluding month of manufacture of products having shelf life more than 2 years & up to 3 years and
- (iii) Within 4 months excluding month of manufacture of products having shelf life more than 3 years.

Products supplied beyond the above-mentioned period from the date of manufacturing shall levied a LD as Per Clause 25.(E) of tender documents. For example, product having manufacturing of November 2022 must be supplied by 31st January 2023 in case shelf life up to 2 Years.

- N) If at any time the Tenderer has, in the opinion of the PMBI delayed the supply of items due to one or more reasons related to **Force Majeure events** such as riots, mutinies, wars, fire, storm, tempest or other exceptional events at the manufacturing premises, the time for supplying the items may be extended by the PMBI at discretion for such period as may be considered reasonable. However, such extension shall be considered only if a specific written request is made by the Tenderer within 10 days from the date of occurrence of such event with necessary documentary evidence. The supplier shall not be liable to pay LD and forfeiture of Security deposit for the delay in executing the contract on account of the extension of supply period on the ground of force majeure events.
- O) The exceptional events do not include the Scarcity of raw material, Increase in the cost of raw material, Electricity failure, Labour disputes/Strikes, Insolvency, and Closure of the Factory/Manufacturing unit on any grounds etc.
- P) Suppliers are required to supply the items within the delivery period mentioned in the purchase order. In this regard it is informed to the bidders that their performance shall be considered unsatisfactory in case of delayed supply (beyond delivery period) or non-supply of products. PMBI may reject their bid in future tenders considering their unsatisfactory performance of supplies.
- Q) Leaked, soiled, broken containers with damaged labels shall not be accepted.
- R) If PMBI observes some physical defects (like empty blisters, improper labelling) of the supplies during sampling, the batch shall be rejected. If supplier wants to take back the batch for rectification, they can take back at their cost, rectify and send back to PMBI within 15 days otherwise same batch shall not be accepted. Due to rectification, if its shelf-life condition as per tender provision does not meet, it shall be discretion of PMBI depending upon requirement to accept the goods with penalty.
- S) Tenderers shall not supply the items declared banned by Government of India, even if Purchase Order is placed.
- T) If the supplier, or any of its approved items gets debarred/banned/blacklisted by any State Government / Central Government / Central or State Government's Drug procurement agencies after entering into agreement with PMBI, it shall be the responsibility of the supplier to inform PMBI without any delay about the same.

20. LOGOGRAM:

Logogram means, wherever the context occurs, the design as specified in **Enclosure of ANNEXURE-VII.** The name of the product shall be mentioned in English and Hindi as per **Pharmacopoeia/any official Compendium/Medical Device Act 2017** and its strength.

- **A**) Tenders should supply for Items etc., as per the specifications such as name, strength, minimum size and packed with appropriate size of the item etc. as per the design enclosed as per **Enclosure to ANNEXURE –VII.**
- B) All form of the supplied product has to be supplied in packing as specified in product list (ANNEXURE XII). Affixing of stickers and rubber stamps shall not be accepted and supplies will be returned back at supplier's cost.
- C) All supplies/packs containing the items tendered for should also carry the printed PMBJP logogram of proportionate size.
- D) Failure to supply Items etc., with the printed logogram of proportionate size will be treated as breach of the terms of Rate Contract / violation of tender conditions. The purchase order shall be cancelled at the risk and cost of the supplier. However, if such failure continuous despite notice,

- will be viewed as a serious lapse and initiate debarring/blacklisting of the supplier and levied a LD as per clause 25 (D) of tender documents.
- E) Items without GS-1 Standard Barcoding on Primary, Secondary and Tertiary Packaging will not be accepted.

21. PACKING:

- A) The items shall be supplied in the package specified in **ANNEXURE -VIII** and **ANNEXURE -XII** and the package shall carry the logograms of proportionate size specified in **Enclosures to ANNEXURE -VII** along with other guideline in this regard (if any) whether it is applicable.
 - a. Medical devices shall be packed in high quality recyclable/ biodegradable material conforming the Standards and Safety parameters if any.
 - b. If any item is required to be packed in disposable plastic packets, the bidder shall adhere to the Plastic Waste Management Amendment Rules, 2021.

Note: Final product must be supplied with PMBJP logogram on primary, secondary, and tertiary packing as per Enclose to ANNEXURE –VII. PMBI may ask the awardee/supplier to develop customize artwork if the product is decided for launching.

Non-affixing of logograms will be treated as violation of tender conditions and fine will be deducted from the amount payable as per condition in Clause 25 (D) of tender documents.

- B) The items to be supplied by the supplier should not be embossed indicating any code no./logo or name of the company. Failure to comply with this shall lead to non-acceptance of the goods besides imposition of penalties as per clause 25. (D)
- C) The packing in each carton shall be strictly as per the specification mentioned in **Annexure-VIII**. The outer carton/secondary packaging should be of pearl white duplex board (**off white/grey is not acceptable**) with a minimum of 350 GSM with **Gloss laminated** packing. **The material to be used for carton should be from virgin chemical pulp.** Failure to comply with this shall lead to non-acceptance of the goods besides imposition of penalties as per clause 25 (D). Storage conditions must be indicated on outer label.
- D) It should be ensured that only virgin packaging material of uniform size is used for packing.
- E) All primary packing containers should be strictly conforming to the specification included in the relevant pharmacopoeia/official Compendium.
- F) **Packing** should be able to prevent damage or deterioration during transit.
- G) The packings/labels of two different products of a same supplier should be clearly distinct from each other
- H) In the event of items / product supplied found to be **not as per specifications in respect of their packing and logogram**, the PMBI is at liberty to make alternative purchase of the items of items for which the Purchase orders have been placed from any other sources or in the open market or from any other Tenderer who might have quoted higher rates, at the risk and the cost of the supplier. In such cases the PMBI has every right to recover the cost and impose penalty as mentioned in Clause 25 & 26.
- I) Designs of packaging with the logograms shall be subject to approval by PMBI within 3 days of receipt of purchase order or within 30 days of release of letter of acceptance. Text matter of all type of label must be checked and responsibility shall be of manufacturer.
 - In case of failure of PMBI to do so, the supplier may go ahead with the design as per the specification in Enclosure to ANNEXURE VII.
 - STP (Standard Testing Procedure) for the awarded items are required to submit within 15 days from the date of Letter of Acceptance.

Note: The bidder shall be solely responsible for the labeling on the packing /product, complying with the official compendium/notification/guideline.

- J) The colour of the strength must be different from the colour of the generic name of the product on primary and secondary packaging and the approval for the same should be taken from the procurement/ quality/regulatory department while taking artwork approval. The printing ink used should be of good quality (clarity, brightness, contrast) which is easily readable.
- K) Therapeutic code, NABL lab tested and other standard confirming the quality of the product shall be indicated on the primary and secondary packaging and shall be incorporated as per the approval from the quality/regulatory department while taking artwork approval.
- L) Barcodes as per GS-1 standards are required to be printed on products at various packaging levels (Primary, Secondary and Tertiary) as per **Annexure-IX**.

22. QUALITY TESTING & QUALITY CONTROL:

- A. All the batches of the items supplied shall be supported by test/ analysis reports furnished by certified Medical Devices Testing Laboratory under MDR Rules 2017. For non-drug items, test / analysis reports must be from independent NABL Accredited Items Testing Laboratory/Central Drug Testing Laboratory/In House Quality Control Laboratory. The Tender Inviting Authority has the right to get the items tested at the laboratories of his choice for further verifications, from PMBI empanelled laboratories.
- B. Random samples of each supplied batch will be chosen at the point of supply or distribution/storage points for testing. The samples will be sent to different PMBI empanelled laboratories including Government Items Testing Laboratory/NIPER/PSU labs for testing. Handling and testing charges will be deducted by PMBI for the above purpose, as specified in Clause 24.
- C. STP (Standard Testing Procedure) for awarded items are required to be submitted within 15 days from the date of Letter of Acceptance by mail to Quality and Regulatory officer of PMBI with artwork approval for design of packaging with the logogram as per Clause 21.K.
- D. The Items shall have the ingredients at the prescribed level as indicated in official compendiums throughout the shelf-life period of the supplied item. The samples will be drawn periodically throughout the shelf-life period and if found "Not of Standard Quality", the cost of entire batch paid will be recovered whether consumed fully/partially. Also, action will be initiated for blacklisting as per clause 26 irrespective of the period of supply. The supplies will be deemed to be completed only upon receipt of the quality certificates from the laboratories. Samples which do not meet quality requirement shall render the relevant batches liable to be rejected. If the sample is declared to be "Not of Standard Quality" or spurious or adulterated or misbranded, such batch/batches will be deemed to be rejected goods.
- E. In the event of the samples of Items supplied fails in quality tests or found to be not as per specifications, the PMBI is at liberty to make alternative purchase of the items of items for which the Purchase orders have been placed from any other sources or in the open market or from any other Tenderer who might have quoted higher rates, at the risk and the cost of the supplier and in such cases the PMBI has every right to recover the cost and impose penalty as mentioned in Clause 26(I).
- F. If the product / sample fails in quality test, every failed batch shall be taken back by the supplier at their own cost and PMBI shall not be responsible for any damage during this period.
- G. The supplier shall furnish evidence of the basis for shelf life and other stability data concerning the commercial final package on request by the PMBI. In case of any complaint in the field, the B.M.R/B.P.R for the particular batch of the product(s) supplied shall be produced when demanded. For New items, complete stability data of 6 months' period shall be acceptable.

- H. The products should conform to the standards of **ISO/CE/ISI/any equivalent guideline as the case may be.** In case the product is not included in the any of the said standard/compendium/guideline, the supplier upon award of the contract must provide the reference and standard testing protocols for product testing.
- I. The case of admixture of items will be treated as a violation of tender conditions and fine will be levied as per clause 26. If such lapses happen more than twice in a tender period such cases will be treated as "Misbranded Items".

23. PAYMENT PROVISION:

- A) No advance payments towards costs of items will be made to the supplier.
- B) Payments towards the supply of items will be made within 60 days from the date of receipt of goods, strictly as per the tender terms and condition. The payment will be made either by means of a/c payee Cheque or through RTGS (Real Time Gross Settlement System)/Core Banking/NEFT. The Tenderer shall furnish the relevant details in original Mandate form (ANNEXURE -V) to make the payment through RTGS/Core Banking/NEFT.
- C) All bills/Invoices should be raised in triplicate and the bills should be drawn as per GST Rules in the name of Pharmaceuticals & Medical Devices Bureau of India. 8th Floor, Videocon Tower, Block-E1, Jhandewalan Extension, New Delhi-110055 or in the name of any other authority as may be designated.
- D)(i) Payments for supply will be considered only after supply of minimum 50% of Items ordered in the individual Purchase Order provided reports of Standard Quality on samples testing are received from Government Analyst or Approved Laboratories of PMBI.
 - (ii) However, in case of cancellation of a particular purchase order due to failure in delivery, payment for part supplies less than 50% of the purchase order quantity on the date of cancellation of the purchase order may be considered for release of payment subject to the following:
 - a) If the Tenderer have supplied at least 50% of the quantity ordered in the subsequent purchase order within delivery period stipulated in purchase order from the issue of such purchase order.
 - b) If further purchase order is not placed with the supplier due to any reason, not attributable to the supplier, the amount eligible will be paid within 60 days from the date of last supply.
 - c) The payment for part supply as mentioned above will subject to the deduction of liquidated damages, penalty towards unexecuted quantity, risk and cost etc., as per the tender conditions.
- E) If at any time during the period of contract, the price of tendered items is reduced or brought down by any law or Act of the Central or State Government or by the Tenderer himself, the Tenderer shall be bound to inform the PMBI immediately about such reduction in the contracted prices. Tender Inviting Authority is empowered to unilaterally effect such reduction as is necessary in rates in case the Tenderer fails to notify or fails to agree for such reduction of rates.
- F) In case of any increase or decrease in the Taxes/GST after the date of submission of tenders and during the tender period, such variation in the taxes/GST will be to the account of the PMBI. For claiming the additional cost on account of the increase in taxes/GST, the Tenderer should produce the proof of having paid additional amount on this account on the goods supplied to PMBI from the concerned authorities and also must claim the same in the invoice separately. However, the basic price structure and the price of the Items approved under the tender shall not be altered. Similarly, if there is any reduction in the taxes/GST and statutory levies as notified by the Govt., after the date of submission of tender, the Tenderer will be paid based on the unit rate worked out on the basis of the reduced taxes/GST/statutory levies without any change in the basic price or the price structure

- of the items approved under the tender. Any increase or decrease in taxes/GST and statutory levies will be considered based on the notification issued by the Government.
- G)However, if the firm supplies after originally stipulated Delivery period, increase in taxes/GST due to statutory variation in taxes/GST shall be borne by the supplier. In case of decrease in taxes/GST due to statutory variation in taxes/GST, the same shall be passed on by the supplier to the PMBI.

24. HANDLING & TESTING CHARGES:

In all supplies, 1.5% of the supply value shall be deducted towards handling & testing charges.

25. LIQIDATED DAMAGES & OTHER PENALTIES:

- A)All supply should be made within the stipulated time as per the clause 19.D of the Delivery Schedule and quantity as mentioned in the Purchase Order.
- B) If the supply reaches the Warehouses beyond the stipulated time as mentioned in PO/Bid document, liquidated damages will be levied at the rates 2% per week or part thereof, subject to maximum of 10% irrespective of the fact that whether the PMBI has suffered any damage/loss or not, on account of delay in effecting supply.
- C) If the supply is received in damaged condition, it shall not be accepted. The supplier shall have to replace the goods with damage and the penalty equal to the penalty for unexecuted supplies will be levied for the damaged goods and payments will be withheld till proper replacement.
- D)All the Tenderers are required to supply the product(s) with printed MRP as per purchase order and logogram of appropriate size on the pack and with prescribed packing specification. If there are any deviation in these Tender conditions, action will be taken to blacklist the product, and/or a separate damage will be levied @ 5% of value of the defaulted quantity irrespective of the Tender Inviting Authority having suffered any damage/loss or not, without prejudice the rights of alternative purchase specified in Clause No.19F,19 H and 21.
- E) If the supplier supplies the product beyond the manufacturing date as mentioned in clause 19. (M) of supply conditions, a liquidation damage will be levied @ 5% per month subject to maximum 30% (Up to 6 months).
- **F**) In all the above conditions, the decision of the Tender Inviting Authority shall be final and binding.

26. DEDUCTION & OTHER PENALTIES ON ACCOUNT OF QUALITY FAILURE:

- A) If the samples do not conform to statutory standards, the Tenderer will be liable for relevant action under the existing laws and the entire stock in such batch has to be taken back by the Tenderer within a period of 30 days of the issue of the letter from the PMBI. Such stock shall be taken back at the expense of the Tenderer. Further, actual handling and testing charges shall be paid to PMBI by the supplier otherwise these charges shall be recovered from their pending bill/Performance Security Deposit.
- B) The PMBI has the right to destroy such "NOT OF STANDARD QUALITY ITEMS" if the Tenderer does not take back the goods within the stipulated time. The PMBI will arrange to destroy the "NOT OF STANDARD QUALITY ITEMS" after the expiry of 30 days mentioned above without further notice and shall also collect handling charges (in case the product is sent back to supplier on freight to pay basis)/ demurrage charges calculated at the rate of 2% per week on the value of the items rejected till such time stipulated. Further, the cost of disposal shall be recovered from the supplier.
- C) If any item or Items supplied by the Tenderer have been partially or wholly used or consumed after supply and are subsequently found to be in bad odour, unsound, inferior in quality or description (Adulterated/Spurious/Misbranded) or otherwise faulty or unfit for consumption, then the contract price or prices of total such batches supplied will be recovered from the Tenderer, if payment had already been made to him. In other words, the Tenderer will not be entitled to any

- payment whatsoever for Item or items found to be of "NOT OF STANDARD QUALITY" whether consumed or not consumed and the Tender Inviting Authority is entitled to deduct the cost of such batch of items from any amount payable to the Tenderer. On the basis of the nature of failure, action will be initiated to blacklist the product/supplier.
- D) For the supply of Adulterated/Spurious/Misbranded as defined in the Drugs and Cosmetics Act, 1940/ Medical Device Rule 2017 other official compendium to PMBI, PMBI reserves the right to blacklist the supplier. No further supplies shall be accepted from the firm/company.
 - If the tenderer is blacklisted, the tenderer shall not be eligible to participate in tenders of PMBI for supply of Items for a period of 5 years from the date of blacklisting.
 - In case of supply of "NOT OF STANDARD QUALITY" items to PMBI, the product shall be debarred/blacklisted by PMBI, and no further supplies shall be accepted for the particular product. The Tenderer shall also not be eligible to participate in tenders of PMBI for supply of such Items for a period of 2 years from the date of blacklisting.
 - In addition, the Director of Drug Control of concerned State will be informed for initiating necessary action on the Tenderer in their state. Performance Security Deposit will also be forfeited without any intimation.
- E) The Tenderer shall furnish the source of procurement of raw material utilized in the formulations, if required by the PMBI. The PMBI reserves the right to cancel the purchase orders, if the source of supply is not furnished.
- F) The decision of the PMBI or any officer authorized by PMBI, as to the quality of the supplied items shall be final and binding. In such cases, the PMBI will be at liberty to terminate, the contract either wholly or in part on 30 days' notice. The Tenderer will not be entitled for any compensation whatsoever in respect of such termination besides forfeiture of Performance Security Deposit.
- G) For contravention of the stipulations of the contract or for other justifiable reasons, the contract may be terminated by the PMBI, and the Tenderer shall be liable to pay for all losses sustained by the PMBI in consequence of the termination which may be recovered from the Tenderer, as per rules besides forfeiture of Performance Security Deposit.
- H) Non-performance of any of the contract conditions and provisions will disqualify a firm from participating in the tender for the next 2 years/Blacklisting the tenderer.
- I) In the event of making Alternative Purchase, as specified in Clause 19.H, penalty will be imposed on the supplier. The excess expenditure over and above contracted prices incurred by the PMBI in making such purchases from any other sources or in the open market or from any other Tenderer who has quoted higher rates and other losses sustained in the process, shall be recovered from the Performance Security Deposit or from any other money due and become due to the supplier and in the event of such amount being insufficient, the balance will be recovered personally from the supplier as per rules.
- **J**) In all the above conditions, the decision of the PMBI shall be final and binding.

27. BLACKLISTING IN THE EVENT OF WITHDRAWAL FROM THE TENDER, AND NON-ADHERENCE TO THE QUALITY STANDARDS AND SUPPLY SCHEDULE:

A) BLACKLISTING OF PRODUCT/TENDERER ON WITHDRAWAL OF TENDER

- a. If the Tenderer fails to perform the obligations under the tender conditions / commits default in the performance of the contract/LOA, such Tenderers will be blacklisted for a period of 2 years by PMBI from the date of intimation besides forfeiture of Performance security deposit.
- b. The Tenderers who have withdrawn after participating in the tender after the last date and time of submission of online bid, either fully or partially, **the entire firm/company** will be blacklisted

for a period of **2 years** from the date of intimation by PMBI apart from forfeiture of the Security Deposit.

B) BLACKLISTING FOR QUALITY FAILURE IN QUALITY TEST BY THE EMPANELLED LABORATORIES OF PMBI.

- a) Each and every batch of items supplied by the supplier shall be subjected to quality test by the Empaneled laboratories as per the procedure adopted by PMBI.
 - PMBI shall also draw the samples of products supplied in the marketplace and get the same tested to make sure the products are conforming to quality requirements till Self life.
- b) If the sample of any batch fails in quality test and report is received stating "Not of standard quality "in any test the report along with the chromatograms etc. such batch of items shall be rejected.
 - (i) If the supplier challenges and request for retesting, the sample shall be tested at government testing laboratory or reputed govt. institute like NIPER. The test report of govt. lab or NIPER will be final and will be binding to the supplier.
 - (ii) The cost of such Re-testing shall be recovered from the supplier.
 - (vi) If **2** batches of item supplied by the same supplier is reported to NOT OF STANDARD QUALITY in specification, then the firm shall be debarred/blacklisted for 2 years after observing procedure laid down in Para 27.B.(d) besides forfeiture of Performance Security Deposit.
 - (vii) If a single batch of any product(s) supplied by the company/firm declared as Adulterated/spurious/ Misbranded by the Government Authorities during the shelf life of the product supplied irrespective of tender period, the company/firm shall be debarred/blacklisted for a period of 2 years from the date of intimation & forfeiture of security deposit.

c) Quality Test by Statutory Authorities:

- (i) If any item is declared "NOT OF STANDARD QUALITY", by any government agencies or Licensing authority, the issue of available stock of the particular item will be stopped. Further, the available stock of the product in hospitals/JAS will be retrieved.
- (ii) If any batch of any product(s) supplied by the company/firm declared, NOT OF STANDARD QUALITY in specification as defined as per the standard mentioned in ISO/CE/ISI/any equivalent guideline by the Government Authorities during the relevant tender period or during quality check within shelf-life period, the company/firm shall be debarred/blacklisted for a period of 2 **years from the date of blacklisting** after observing procedure laid down in Para 27.B (d).

d) Procedure for Blacklisting:

- (i) On receipt of complaint from Distributer/retailers/customers or report from Govt. Analyst/ Drug Testing Laboratory/ Medical Devices Testing Laboratory certified by CDSCO under MDR 2017 indicating that a particular Item is "NOT OF STANDARD QUALITY/ ADULTERATED/ SPURIOUS/MISBRANDED" (As the case may be), a show cause notice shall be issued to the supplier calling for explanation within 7 days from the date of notice. On receipt of explanation from the supplier, the CEO, PMBI may take appropriate action on merits of the case and impose penalty including the blacklisting of the item of the product/company or firm as deemed fit besides forfeiture of Performance Security Deposit.
- (ii) If a particular item has been blacklisted according to the procedure stated above, the supplier is not eligible to participate in any of the tenders for such item floated by the PMBI until the period of blacklisting is over.

(iii) If a supplier company/firm is blacklisted according to the procedure stated above, such supplier is not eligible to participate in any of the tenders floated by the PMBI until the period of blacklisting is over.

G) BLACKLISTING FOR NON-SUPPLY/PART SUPPLY:

- i. Due to non-supply of item against any purchase order, 5 % value of purchase order shall be recovered from the supplier in addition of other penal like risk purchase. In case of repeated circumstances of non-supply of items i.e., 2 times, the supplier may be blacklisted for 2 years in addition of forfeiture of Performance Security Deposit and other penal action at the discretion of PMBI.
- ii. If the supplier fails to execute at least 50% of the quantity mentioned in a purchase order and such part supply is come into existence in three Purchase orders during the currency of contract period, then supplier shall be liable for debarment for the particular product for two years. Two years period will be reckoned from the date of issuance of such debarment order.
- iii. If a supplier does not supply any quantity against two successive purchase orders, then supplier shall be liable for debarment for the particular product for two years. Two-year period will be reckoned from the date of issuance of such debarment order at the discretion of PMBI.

28. SAVING CLAUSE:

No suit, prosecution or any legal proceedings shall lie against the Tender Inviting Authority or any person for anything that is done in good faith or intended to be done in pursuance of the tender.

29. RESOLUTION OF DISPUTES:

The PMBI and the supplier shall make every effort to resolve, amicably by direct informal negotiation any disagreement or dispute arising between them under or in connection with the contract.

A) ARBITRATION AND JURISDICTION.

Normally, there should not be any scope of dispute between the PMBI and the supplier after entering a mutually agreed valid contract/ Rate Contract.

However, due to various unforeseen reasons, problems may arise during the progress of the contract/ Rate Contract leading to disagreement PMBI and the supplier shall first try to resolve the same amicably by mutual Consultation. If the parties fail to resolve the dispute by such mutual consultation within twenty-one days, then, depending on the position of the case, either the PMBI or the supplier shall give notice to other party of its intension to commence Arbitration procedure as per Indian Arbitration and Conciliation Act, 1996. Such disputes/differences shall be referred to Sole Arbitrator to be appointed by the CEO of PMBI. The venue of Arbitration Shall be at New Delhi. The award published by the Arbitrator shall be final and binding on the parties.

B) In case of a complaint received from any local supplier indicating a need for review / verification of Local content of successful vendor / awarded vendor, for accepting a complaint from such complainant (w.r.t the false declaration given by the successful vendor on the local content), a complaint fee of Rs.2 Lakhs or 1% of the locally manufactured items being procured (subject to a maximum Rs.5 Lakhs), whichever was higher, to be paid by demand draft by the complainant. In case, the complaint is found to be incorrect, the complaint fee shall be forfeited. In case, the complaint is upheld and found to be substantially correct, deposited fee of the complainant would be refunded without any interest.

30. CONTACTING THE PMBI BY THE BIDDER:

- A) No bidder shall contact the PMBI on any matter relating to its bid, from the time of bid opening to the time the contract is awarded.
- B) Any effort by a bidder to influence the PMBI in the Purchaser's bid evaluation, bid comparison or contract award decisions may result in rejection of the bidder's bid.

- C) The bidder shall not make any attempt to establish unsolicited and unauthorized contact with the Tender Accepting Authority, Tender Inviting Authority or Tender Scrutiny Committee after opening of the bids and prior to the notification of award and any attempt by any bidder to bring to bear extraneous pressures on the Tender Accepting Authority, Inviting Authority or Tender Scrutiny Committee, shall be sufficient reason to disqualify the bidder.
- D) Notwithstanding anything contained in clause (C) above the Tender Inviting Authority or the Tender Accepting Authority, may seek bonafide clarifications from bidders relating to the bids submitted by them during the evaluation of bids.

31. FRAUDULENT AND CORRUPT PRACTICES:

A) For Bidders:

It is purchaser's policy to ensure that suppliers and their authorized representatives/agents observe the highest standard of ethics during the procurement and execution of such contracts. (In this context, any action taken by a bidder, supplier, contractor, or by their authorized representatives/agent, to influence the procurement process or contract execution for undue advantage is improper) In pursuance of this policy, the purchaser.

a) Defines, for the purposes of this provision, the terms set forth below as follows:

- (i) "Corrupt practice" is the offering, giving, receiving or soliciting, directly or indirectly, of anything of value to influence improperly the actions of another party ("another party" refers to a public official acting in relation to the procurement process or contract execution). In this context, "public official" includes staff and employees of other organizations taking or reviewing procurement decisions.
- (ii) "Fraudulent practice" is any act or omission, including a misrepresentation, that knowingly or recklessly misleads, or attempts to mislead, a party to obtain a financial or other benefit or to avoid an obligation (a "party" refers to a public official; the terms "benefit" and "obligation" relate to the procurement process or contract execution; and the "act or omission" is intended to influence the procurement process or contract execution).
- (iii) "Collusive practice" is an arrangement between two or more parties designed to achieve an improper purpose, including to influence improperly the actions of another party ["parties" refers to participants in the procurement process (including public officials) attempting to establish bid prices at artificial, non-competitive level].
- (iv) "Coercive practice" is impairing or harming, or threatening to impair or harm, directly or indirectly, any party or the property of the party to influence improperly the actions of a party (a "party" refers to a participant in the procurement process or contract execution).
- (v) "obstructive practice" is (a) deliberately destroying, falsifying, altering or concealing of evidence material to the investigation or making false statements to investigators in order to materially impede a investigation into allegations of a corrupt, fraudulent, coercive or collusive practice; and/or threatening, harassing or intimidating any party to prevent it from disclosing its knowledge of matters relevant to the investigation or from pursuing the investigation; or acts intended to materially impede the exercise of the purchaser's inspection and audit rights provided for under sub-clause (e) below.
- b) Will reject a proposal for award if it determines that the bidder considered for award has, directly or through an agent, engaged in corrupt, fraudulent, collusive, coercive, or obstructive practices in competing for the contract in question.
- c) will cancel the contract if the purchaser determines at any time that the bidder, supplier and contractors and their sub-contractors engaged in corrupt, fraudulent, collusive, or coercive practices.

- d) will sanction a firm or individual, including declaring in eligible, either indefinitely or for a stated period of time, to be awarded a contract if it at any time determines that the firm has, directly or through an agent, engaged in corrupt, fraudulent, collusive, coercive or obstructive practices in competing for, or in executing, a contract; and
- e) will have the right to inspect the accounts and records of the bidders, supplier, and contractors and their subcontractors/authorized representatives and to have them audited by auditors appointed by the purchaser.

B) For Suppliers:

If the PMBI determines that a Supplier has engaged in corrupt, fraudulent, collusive, coercive or obstructive practices, in competing for or in executing the Contract, then the PMBI may, after giving 7 days' notice to the Supplier, terminate the Supplier's engagement under the Contract and cancel the contract, and the procurement will be made at the risk and cost of the supplier besides blacklisting the bidder for 5 years with forfeiture of Performance Security Deposit apart from other penal actions.

a) For the purposes of this Sub-Clause:

- (i) "Corrupt practice" is the offering, giving, receiving, or soliciting, directly or indirectly, of anything of value to influence improperly the actions of another party.
- (ii) "Fraudulent practice" is any act or omission, including a misrepresentation, that knowingly or recklessly misleads, or attempts to mislead, a party to obtain a financial or other benefit or to avoid an obligation.
- (iii) "Collusive practice" is an arrangement between two or more parties designed to achieve an improper purpose, including to influence improperly the actions of another party.
- (iv) "Coercive practice" is impairing or harming, or threatening to impair or harm, directly or indirectly, any party or the property of the party to influence improperly the actions of a party.
- (v) "Obstructive practice" is (aa) deliberately destroying, falsifying, altering or concealing of evidence material to the investigation or making false statements to investigators in order to materially impede a purchaser investigation into allegations of a corrupt, fraudulent, coercive or collusive practice; and/or threatening, harassing or intimidating any party to prevent it from disclosing its knowledge of matters relevant to the investigation or from pursuing the investigation; or (bb)acts intended to materially impede the exercise of the purchaser's inspection and audit rights provided for.

32. JURISDICTION:

In the event of any dispute arising out of the tender such dispute would subject to the jurisdiction of the Civil Court within the city of Delhi only.

ANNEXURE – I Ref. Clause 3 (P)

CHECK-LIST (Whether uploaded the documents)

COVER - A

S.N.	Check List	YES / No	Page No.	Remarks
1	Check list – ANNEXURE – I as per clause 3. P.			
2	Bid Security declaration on non-judicial stamp paper as per ANNEXURE-III (Clause 3. A & 6. A).			
3	NSIC or MSME or SSI certificate (If EMD is exempted) as per Clause No. 3.A.			
4	Scanned copy of certificate of recognition as start up by Department of Industrial Policy and Promotion for quoted item for relaxation of prior turnover and prior experience for Start-ups (as defined by Department of Industrial Policy and Promotion) as per clause no. 3.A.(ii)			
5	Copies of documentary evidence for the constitutions of the company / Firm/ Proprietorship such as Memorandum and Article of Association, Partnership deed with complete address as per Clause 3. B.			
6	Power of attorney or Resolution of board by which the authorized signatory has been authorized by the Tenderer to sign the tender documents as per clause 3. C.			
7	Copy of valid Manufacturing License/permission of the product quoted as per Clause 3. E.			
8	Copy of valid Quality Management System (QMS) certificate issued on behalf of manufacturing unit by by National Accreditation Board for Certification Bodies (NABCB), India as per Clause 3. (H).			
9	Valid Market Standing Certificate of minimum three consecutive years (Latest) issued by the concerned Licensing Authority/Drugs Control Department/Concerned Government Department for the quoted product.			
10	Valid Market standing Certificate (MSC) issued by the C.A. certifying batch No. that the firm/company has manufactured quoted items for last three years.			
11	Self- declaration is to be submitted on company's letter head duly signed by authorized signatory certifying that the firm/company has not been convicted in last three years.			
12	Tenderer must declare their maximum Production Capacity (item wise) issued by concerned Licensing Authority / self-declaration highlighting the quoted product as per Clause no. 3. J.			
13	ANNEXURE –II (Declaration On non-judicial Stamp Paper for eligibility in participating the tender) original Annexure II delivered to PMBI as per clause 3. L.			
14	ANNEXURE IV (Certificate from the C.A. (Chartered Accountant) or Company Secretary. Original Annexure IV delivered to PMBI as per clause 3. K.			
15	ANNEXURE-V (Mandate form) to furnish company bank details as per clause 3 (M) & 23(B)			
16	ANNEXURE-VI indicating manufacturing License/Permission/ Registration , validity of license and market standing certificate details as per clause 3. N.			
17	ANNEXURE-VII (Declaration to supply the items as per the			

	design in enclosure in Annexure VII) as per clause 3(O), 20 & 21			
19	ANNEXURE-XV (Declaration on Non-judicial Stamp Paper duly notarized stating that the firm & its quoted product is not blacklisted currently (as on the date of submission of the tender) by Central Government/ Central Government agencies/any State Government or any of the State Government agencies/ or any Drug procurement agencies or by PMBI as per clause 3. D(f).			
20	ANNEXURE – XVI (Declaration % of Local content used in the manufacturing of quoted product) as per clause 3.U.			
21	Copy of valid GS-1 registration certificate for bar coding as per Clause 3. T.			
22	Copy of Audited Annual Balance sheet and Profit and loss statement showing details of their annual average turn over not less than 2 crores for any three of the last four consecutive financial years as per Clause 3. K.			
23	Self-attested copy of PAN Card of the Bidder Company. As per Clause 3. Q.			
24	Self-attested copy of Certificate of valid GST registration of the bidder company. As per Clause 3. R.			
25	Self-attested copy of Income Tax Return for any three of last four consecutive Assessment years. As per Clause 3. S.			
26	Authorization letter nominating an employee of the tenderer to transact the business with the Tender Inviting Authority as per clause 4.0.			
27	Annexure XVII (Clause No. 3.1) Declaration towards Compliance of Order (Public Procurement No.1, 2 & 3) dated 23 Jul 2020 & 24 Jul 2020 under Rule 144 (xi) of the General Financial Rules (GFRs), 2017 on Non-Judicial Paper duly notarized.			
28	Self-attested copies of Test Reports furnished by certified Medical Devices Testing Laboratory under MDR Rules 2017.			
29	Test / analysis reports of non-drug items from independent NABL Accredited Items Testing Laboratory/Central Drug Testing Laboratory/In House Quality Control Laboratory.			
30	Self-attested copies of Catalogues along with dimension and design of quoted items as per clause no. 3.w.			
31	Declaration on Company letterhead to provide the warranty as required in Annexure XII.			
ANNI the d	E: - ANNEXURE II, ANNEXURE III (EMD), ANNEXURE EXURE VI, ANNEXURE XV, ANNEXURE XVI, ANNEXURE ocument as per checklist duly authorized along with samples e stipulate date as mentioned in the tender document "technical"	XVII in should	origir be sul	nal and rest o

Name of authorized signatory:
Signature of authorized signatory:
Company seal:

<u>ANNEXURE -II</u> (On nonjudicial Stamp Paper)

Ref. Clause No. 3. (L)

DECLARATION

I/We M/s re	presented by it	ts Proprietor/Managing	Partner /Managing	Director
having its registered office a	at			and its
factory	pr	emises		at
			do	hereby
declare as under: -				

- (I) that I/we have carefully read all the terms and conditions of tender in ref. no. **PMBI/SURGICAL/RC-210/2023 dated 15/06/2023** including Amendment(s) to Tender document (if any) issued by Pharmaceuticals & Medical Devices Bureau of India, New Delhi,110055 and accept unconditionally all terms and condition of tender document including Amendment(s) to Tender document (if any).
- (II) A. that I/We are holding and have uploaded (a) valid Quality Management System (QMS) of the manufacturing unit issued by National Accreditation Board for Certification Bodies (NABCB), India (b) valid manufacturing permission/license/registration for quoted drug/non-drug item/medical devices along with all relevant certificate including ISO/CE/ISI and mentioned requirements in the specification (c) valid self-declaration of non-conviction certificate on company's letter head as per Annexure XV (d) valid Market Standing Certificate issued by licensing Authority for medical devices, from C.A/C.S for non-drug items confirming that we have manufactured & marketed two batches in last 3 years, (e) declaration of the internationally accepted material (if any) and (f) the copies of the specifications for all quoted items and STP (standard testing procedure) quoted items and also enclosed all undertaking/declaration as per Annexure mentioned in the tender document.

(II) B. that I/we shall supply the items as per specification, design, logo and packing given in ANNEXURE-XII, ANNEXURE-VII, ANNEXURE-VIII. That we agree to develop artwork if required by PMBI.

On the basis of above undertaking/declaration, the price bid shall be opened subsequently after opening of technical bid. However, any document uploaded with technical bid is not complying as per undertaking, the contract/ Rate Contract shall be cancelled with forfeiture of Performance Security Deposit/Bank guarantee against tender no. PMBI/SURGICAL/RC-210/2023 dated **15/06/2023** along with other penal action.

(III) a.) I/We declare that we possess the valid manufacturing license for PMBI's tendered items as per details below:

Sr. No.	Item Code	Description of Item as per PMBI Tender	U	Date of Issue	Address of Manufacturing
			Registration. No.		Unit

b.) I/We declare that we possess the valid Manufacturing license/Permission/Registration Certificate issued by competent authority and complies and continue to comply with the condition lied there under to manufacture the items.

I am / We are aware of the Tender inviting Authority's right to forfeit the Earnest Money Deposit and /or Performance Security Deposit and blacklist me/us for a period of 5 years if, any

information furnished by us proved to be false at time the of inspection and not complying the condition for a period of five years.

(IV) I do hereby declare that I have uploaded valid GS1 registration certificate for bar coding and will supply the item with bar code as per ANNEXURE IX and as per the design as per enclosures to ANNEXURE VII enclosed with tender document as well as other instruction given in this regard.

(V) that in pursuant to the conditions in Clause No. 6. (A) of the tender, the Earnest Money Deposit can be forfeited by the Tender Inviting Authority in case of violation of any of the conditions and non-performance of the obligation under tender document.

(VI) that our company/applied items have not been blacklisted/debarred/de-registered/banned due to quality failure for the quoted product /firm by any State Government / Central Government/ PMBI/ Central or State Government's Drug procurement agencies for the following products quoted in the tender at the time of submission of bid. Further, quoted items have not been failed in house testing or testing by any State Government/Central Government / its Drug procurement agencies/PMBI during last two years. We are eligible to participate in the tender ref. No. PMBI/SURGICAL/RC-210/2023 dated 15/06/2023 for the following quoted products with mentioned shelf life in Annexure XIII: -

S	S. No.	Item Code	Description of Item as per PMBI Tender	Unit Size	Shelf life as per Annexure XIII

(VII) that I/we have quoted the rates in BOQ for above mentioned drug codes (table under para VI) in my/our full consciousness abiding by the terms and condition laid down in the tender document considering unit size, secondary pack, shelf life, packaging type etc. and declare it too be invariable.

	Signed
	Name:
	Designation
	(Company Seal)
Witness: -	
(1) Signed:	
Name:	
Designation:	
(2) Signed:	
Name:	
Designation:	

To be attested by the Notary

ANNEXURE-III

(Ref: -Clause **3(A)**, 6.A)

DETAILS OF EMD SUBMITTED

(UPLOAD THE SCANNED COPY OF DRAFT/ PAY ORDER/BANK GURANTEE/NEFT RECEIPT)

In case bidder willing to submit Bank Guarantee (BG)

MODEL BANK GUARANTEE (BG) FORMAT FOR FURNISHING EMD (if bidder intends to submit BG)

Whereas
for the supply of Items (hereinafter called the "tender") against the purchaser's tender enquiry No. PMBI/SURGICAL/RC-210/2023 know all men by these presents that we
of India (PMBI) of India New Delhi(hereinafter called the "Purchaser) in the sum of Rs. 100000.00 (One lakh) only for which payment will and truly to be made to the said Purchaser, the Bank binds itself, its successors and assigns by these presents. Sealed with the Common Seal of the said Bank this
The conditions of this obligation are:
(1) If the tenderer withdraws or amends, impairs or derogates from the tender in any respect within the period of validity of this tender.
(2) If the tenderer having been notified of the acceptance of his tender by the Purchaser during the period of its validity: -
(a) Fails or refuses to accept/execute the contract.
We undertake to pay the Purchaser up to the above amount upon receipt of its first written demand, without the Purchaser having to substantiate its demand, provided that in its demand the Purchaser will note that the amount claimed by it is due to it owing to the occurrence of one or both the two conditions, specifying the occurred condition or conditions.
This guarantee will remain in force up to 12 months from the due date of tender i.e.,and any demand in respect thereof should reach the Bank not later than the above date.
Signature of the authorized officer of the Bank
Name of the officer
Designation of the officer
Seal, name & address of the Bank and address of the Branch

PMBI/SURGICAL/RC-210/2023

ANNEXURE- IV

Ref. Clause No. 3. (K)

{Format for a certificate from the C.A. (Chartered Accountant) or Company Secretary}

(I) (II)	Ltd./Propri GST regist returned Shri The annual last four c	tetorship /Partnership company/ firm are tration no	is a Private Ltd./ nd they have PAN no			
	Sl. No.	Financial Year	Turnover in Crores (Rs.)			
	1.	2018-19	₹			
	2.	2019-20	₹			
	3.	2020-21	₹			
	4.	2021-22	₹			
	TOTAL		Rs Crores			
	Average	Turnover per annual	Rs Crores			
(III)	It is certified that M/S					
(IV)	Further, It (MSE) an authorities eligible for	is certified that M/Sd registered with Director of Industron quoted items against PMBI tende	tender. This certificate is based on their al and financial statement.			
(V)	They have three years		ommercial batches of each quoted items in last			

Date:	Name:
	Signature:
	Stamp:
	Registration No.:
NOTE (i) Strike which is not applicable in above of	porti fi anto
(ii) MSMEs would be treated as owned by MSME, proprietor(s) shall be SC /ST by shall be holding at least 51% (fifty-one page 1).	by SC/ST entrepreneurs: a) In case of proprietary b) In case of partnership MSME, the SC/ST partners percent) shares in the unit c) In case of Private Limited ent) share shall be held by SC/ST promoters.

ANNEXURE V

Ref. clause 3 (M) & 23 (B)

MANDATE FORM

Sl. No.	Details Required	Information to be filled for correspondence
1.	Company Name:	
	Postal Address of the Company	
2	GST No.	
2.	Telephone No.	
	Fax No.	
	E-mail ID	
3.	Name of the Managing Director / Director / Manager	
3.	Mobile No. / Phone No	
	E-mail ID	
	Name and Designation of the authorized company official	Name:
4.	company official	Designation:
	Mobile No.	
	E-mail ID	
	Name and Designation of the company official Authorised for communication	Name:
5.	in respect of technical documents / artwork (technical person).	Designation:
	Mobile No.	
	E-mail ID	
	Name and Designation of the company official Authorised for communication	Name:
6	in respect of status of Purchase Orders	Designation:
	/ Logistic support. /Artwork	
	Mobile No.	
	E-mail ID	
-	For Vendor Portal Registration	
7.	Permanent E-mail ID	
	Permanent Mobile No.	
	Bank Details	
	a) Name of the Bank	
	b) Branch Name & address c) Branch Code No.	
	d) Branch Manager Mobile No.	
	e) Branch Telephone no	
	f) Branch E-mail ID	
7.	g) 9-digit MICR code number of the	
	bank and branch appearing on the	
	MICR cheque issued by the bank	
	h) IFSC Code of the Branch	
	i) Type of Account (Current / Savings)	
	j) Account Number (as appear in cheque book)	

	r verification of the above part	please <u>attach the original cancelled cheque</u> issued iculars).
transaction is dewould not hold conditions of the	elayed or not effected at all the Pharmaceuticals & Medical D	es given above are correct and complete. If the ne reasons of incomplete or incorrect information, devices Bureau of India responsible. I have read the d and agree to discharge the responsibility expected essful tenderer.
Date:		Signature :
		Name :
		Designation:
Place:	Company Seal	(Name of the person signing & designation)
	HAT THE PARTICULARS F PER OUR RECORDS.	FURNISHED ABOVE BY THE COMPANY ARI
		FURNISHED ABOVE BY THE COMPANY ARI
CORRECT AS	PER OUR RECORDS.	
CORRECT AS	PER OUR RECORDS.	
CORRECT AS	PER OUR RECORDS.	
Bank Seal with	PER OUR RECORDS. address: address:	
Bank Seal with	PER OUR RECORDS. address: address:	Signature of the authorized official of the bank
Bank Seal with a	PER OUR RECORDS. address: address:	Signature of the authorized official of the bank

Annexure VI Ref Clause No. 3 (N)

3.N.	Item Code (Only Quoted items as mentioned		Item Manufacturing License / Permission				Marketing standing Certificate (MSC)			
	in Annexure II)		Manufacturin g License / Permission/ Registration No.	Manufacturing License / Permission/ Registration Issue date	Manufacturing License / permission/ Registration Renewal Date	License Validity Date		Market Standing Certificate Issue Date	Period of Marketing as per Marketing standing Certificate (MSC)	Page no. of Document in uploaded Scan Copy (Do not put page nos. in range)

N	ote	٠

- (i) In case any details as desired above is missing/not submitted against quoted items, the bid for such items is liable to be rejected.
- (ii) It is strictly, do not put page nos. in range, indicate the page nos. one by one for all respective quoted items codes.

Signature:
Name:
Authorized Signatory:
Seal of the Company:

ANNEXURE -VII

Ref. Clause no. 3(O), 20 & 21

DECLARATION

I/We do hereby declare that I/we will supply the item as per the design in Enclosure to Annexure VII as well as other instruction given in this regard. I/We do hereby also declare that I/we will supply the final product with "PMBJP" logogram on it.

I/we will supply the final product with "PMBJP" logogram on it	
Signature of the Tenderer	
Name:	
Designation:	
	(Company Sea
	(Company Sea

PMBI/SURGICAL/RC-210/2023

Enclosure to ANNEXURE - VII

Ref. Clause No. 20

DESIGN FOR: Mono pack

1. Text Matter Printing on mono pack should be in minimum two colour i.e., Black & red.

However, colour and design of PMBJP (Pradhan Mantri Bhartiya Janaushadhi Pariyojana) logogram in standard colour format & PMBI Item code-XXXX as given in PO as per approval at the time of ARTWORK approval before supply should be as given below.

- 2. PMBJP Logogram should be placed along with the address as given below.
- 3. PMBI helpline number 1800 180 8080 should be printed.
- 4. Font type should in CALIBIRI format for any type of title name of the product.
- 5. Title name of generic item should be **bold** in minimum 12 font size & the strength corresponding to it must be **bold** in minimum 14 font sizes and it may increase respectively according to size of label & the rest text matter should be in minimum 8 font size.
- 6. The stereo printing of batch no./manufacturing /expiry date & other details shouldn't overlap the text matter.
- 7. "Pharmaceuticals & Medical Devices Bureau of India(PMBI)" should be running text only and should not be prominent.



1. Pradhan Mantri Bharitya Janaushadhi Priyojana should be printed in Hindi at side of pack

SPECIMEN LABEL FOR MONO CARTON (Secondary Packing)

Generic Name of Product: XXXXXXX



Manufactured for:

Pharmaceuticals & Medical Devices Bureau of India

8th Floor, Videocon Tower, Block-E1, Jhandewalan Extension, New Delhi-110055

PMBI helpline number 1800 180 8080

PMBI ITEM CODE--XXXX

Note: An additional to any statuary requirement.

ANNEXURE-VIII

Ref. Clause No. 21.(A), 21 (A) .(c)

SCHEDULE FOR PACKAGING OF ITEMS

GENERAL SPECIFICATIONS

<u>A.</u>

i) Primary Package:

Each Primary Package shall contain ordered product in standard packing material with a minimum micron thickness that ensures that the pack does not tear in routine handling (subject to approval of sample by concerned Officer/Committee) which will confirm to size of the product and sealed properly.

(ii) Secondary Package:

- a. The outer carton/secondary packaging should be of pearl white duplex board (**off white/grey is not acceptable**) with a minimum of 350 GSM with **Gloss laminated** packing for the product, it shall be with minimum bursting strength of 9-10 Kg/cm2.
- b. The material to be used for carton should be from virgin chemical pulp. Failure to comply with this shall lead to non-acceptance of the goods besides imposition of penalties as per clause 25 (D). Storage conditions must be indicated on outer label.
- c. One Box shall contain primary packages of supplied products as described in Column 5 of Annexure XII.

ANNEXURE IX (BARCODE REQUIREMENTS)

Reference clause 21(L)

GS1 barcode requirements on Items procured by Pharmaceuticals & Medical Devices Bureau of India (PMBI)

These requirements cover items procured by Pharmaceuticals & Medical Devices Bureau of India (PMBI), New Delhi meant for supply and distribution through PMBI regulated distribution channel.

Barcode based on GS1 identification standards are provided below at various levels of product packaging which includes primary, secondary and shipper/carton levels and need to be complied with while supplying items to PMBI.

GS1 India is unique identification & barcoding standards body setup by Ministry of Commerce & Industry, Govt. of India along with APEDA, BIS, Spices board, IIP and apex industry chambers like CII, FICCI, ASSOCHAM to assist India industry and govt. bodies on adoption of global standards.

Suppliers are also required to provide GS1 subscription validity certificate at the time of supply of items issued by GS1 India. For validity certificate suppliers can contact GS1 India at 011-42890-846.

Barcodes based on GS1 global standards are required to be printed on product packaging at primary, secondary and tertiary packaging levels **in addition** to other, existing statutory labelling & marking requirements.

Technical Specification for GS1 Standards

Tertiary Level Pack:

Is defined as a level of packaging that shall contain one or more secondary/primary levels of packaging and is also considered as the final logistics unit like shippers/pallets.

The Tertiary label will carry two barcodes in GS1-128 format

First Barcode

- a) Unique product identification code (GTIN Global Trade Identification Number)
- b) Manufacturing Date
- c) Expiry date
- d) Batch no.
- e) Quantity

Second Barcode

f) Serial Shipping Container Code (SSCC) –

Note-

- 1) While encoding Manufacturing and expiry date in the barcode, if a specific Manufacturing or expiry date is not printed on the finished pack/ then Supplier should select first day of the month as the Manufacturing date and Last day of the month as expiry date.
 - Example- If Shelf life is 24 months, April 2022 manufacturing date should be encoded as 220401 and March 2024 expiry date as 240331.
- 2) SSCC number of the Tertiary pack should never be reused on another Tertiary pack irrespective the Item, Batch or expiry is different.
- 3) For converting, GTIN-13 into GTIN-14, kindly use "0" as a prefix for all levels of packaging.

Attribute	Description	Length	Nature	Data Type
(02)	Application Identifier to indicate GTIN-14 Brackets not encoded in the barcode	2	Fixed	Numeric
0 8901072 00253 3	Unique Product Number-GTIN-14	14	Fixed	Numeric
(11)	Application Identifier to indicate Manufacturing date Brackets not encoded in the barcode	2	Fixed	Numeric
180101	Expiry Date in YYMMDD format	6	Fixed	Date
(17)	Application Identifier to indicate Expiry date Brackets not encoded in the barcode	2	Fixed	Numeric
220131	Expiry Date in YYMMDD format	6	Fixed	Date

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(10)	Application identifier to indicate Lot/batch number Brackets not encoded in the barcode	2	Fixed	Numeric
BATCH123	Batch No / Lot No	20	Variable	Alphanumeric
(37)	Application identifier to indicate Quantity in Outer Carton	2	Fixed	Numeric
500	No of Primary packs in the tertiary.	Upto 8	Variable	Numeric
(00)	Application identifier to indicate the SSCC Brackets not encoded in the barcode	2	Fixed	Numeric
1 8901072 000000000 6	Unique number of the tertiary pack. It should never be reused.	18	Fixed	Numeric
		Ī		

To, BPPI Mnfd By, AAA Pharma Company 125, SEZ Ahmedabad-382213 Gujrat

Drug Name: Dobucin 500 mg Exp Date: 31 Jan 2022 Batch No: BATCH123

Recommended Barcode – GS-128





Secondary Level Pack:

Is defined as a level of packaging that may contain one or more primary packages usually termed as Mono-carton/carton

Secondary level barcode can be generated using 2D- GS1 Datamatrix or 1D- GS1-128 format.

Note-

- 1) Shrink wrap packaging will not be considered as Secondary level packaging.
- 2) For converting, GTIN-13 into GTIN-14, kindly use "0" as a prefix for all levels of packaging.

Data Attributes Captured in GS1 Datamatrix format

1) Unique product identification code (GTIN)

- 2) Batch No.
- 3) Qty-No of packs

Attribute	Description	Length	Nature	Data Type
(02)	Application Identifier to indicate GTIN-14. Brackets not encoded in the barcode	2	Fixed	Numeric
0 8901072 00253 3	GTIN-14- Unique product code with first digit being the packaging indicator	14	Fixed	Numeric
(10)	Application identifier to indicate Lot/batch Brackets not encoded in the barcode	2	Fixed	Numeric
BATCH123	Batch No / Lot No	Upto 20	Variable	Alphanumeric
(37)	Application Identifier to indicate serial number Brackets not encoded in the barcode	2	Fixed	Numeric
5 Quantity/Units in Secondary pack		Upto 8	Variable	Alphanumeric

Recommended Barcode depending upon the space available – GS1 Data matrix Or

GS1-128



(02) 0 8901072 00255 (10) BATCH123 (37) 5



Primary Level Pack:

Is defined as the first level of packaging in direct contact with the product.

Scenario-I Primary pack with a Mono-carton/Carton/Secondary level pack

For primary packaging packed in a Mono-carton/Secondary pack carton

Unique product identification code (GTIN)

Note-

For converting, GTIN-13 into GTIN-14, kindly use "0" as a prefix for all levels of packaging.

Attribute	Description	Length	Nature	Data Type
(01)	Application Identifier to	2	Fixed	Numeric

	indicate GTIN-14 Brackets not encoded in the barcode			
0 8901072 00253 3	GTIN-14 with first digit being the packaging indicator	14	Fixed	Numeric

Recommended Barcode – GS1 Datamatrix,



Scenario-II Primary pack without Mono-carton/Secondary level pack

For Primary packaging going directly into Tertiary pack without a Carton/Mono-carton/Secondary pack

Unique product identification code (GTIN)

Batch No.

Note-

For converting, GTIN-13 into GTIN-14, kindly use "0" as a prefix for all levels of packaging.



(01)08901072002533 (10)BATCH123

Attribute	Description	Length	Nature	Data Type
(01)	Application Identifier to indicate GTIN-14. Brackets not encoded in the barcode	2	Fixed	Numeric
0 8901072 00253 3	GTIN-14- Unique product code with first digit being the packaging indicator	14	Fixed	Numeric
(10)	Application identifier to indicate Lot/batch Brackets not encoded in the barcode	2	Fixed	Numeric
BATCH123	Batch No / Lot No	Upto 20	Variable	Alphanumeric

Mapping of Manufacturer GTIN with PMBI Item code-

- GS1 has facilitated an online application to link Manufacturer GTIN code with PMBI item code. The manufacturer must update the same before sending the physical consignment to PMBI.
- Kindly contact Mr. Ankit Arora or Mr. Amrit Garg for the same at 011-42890846/42890818 or write email at ankit@gs1india.org or amrit@gs1india.org

Barcode Design and Printing-

- For PMBI suppliers, GS1 India has facilitated an online application to generate the barcode designs for each level of packaging.
- Using the same, the supplier will be able to generate Primary, secondary and Tertiary barcodes as per PMBI format.
- Kindly contact Mr. Ankit Arora or Mr. Amrit Garg for the same at 011-42890846/42890818 or write email at ankit@gs1india.org or amrit@gs1india.org

Please contact GS1 India office for any further assistance –

GS1 India

(Under Min. of Commerce, Govt. of India)
330, 2nd Floor, 'C' Wing, August Kranti Bhawan,
Bhikaji Cama Place, New Delhi - 110066 **T** +91-11-42890890, (D) +91-11-42890846 **F** +91-11-26168730

E ankit@gs1india.org

W http://www.gs1india.org

ANNEXURE -X

(On nonjudicial Stamp Paper)

(Refer Clause no. 3.S)

(To be submitted on Non-judicial Stam paper duly notarized)

DECLARATION OF LOCAL CONTENT

at				, D/o, W/o in the cap	acity of Proprietor/	Managing Partner		
reg	gistered o	ffice at			and	factory premises		
	at							
	S. No.	Item code	Item Name	Details of Location(s) at which value addition is made	Percentage (%) of Local content	Category claimed		
	1							
	3							
Fu acce No sup	ocurement rvices on the rther, the cordance of	17-PP(BE-II) at (Preference behalf of M/s calculations of with the guide 1/4/2018- policity the quoted drufts	dated 16.09.2 to Make in Indicated of local content uselines laid downly dated 01.01.2 ags/medicines.	16/2016-MD policy dated (1020) for the implement (1020) for the implement (1020) for the implement (1020) for the implement (1020) and that I found out (1020) and that I found out (1020) for inviting Authority for the Signature	tation of provision ted to procurement to procurement to the drugs/medic of Pharmaceutical firm under Classian firm under Classian the Tender investigation of assistance of the Tender investigation of the purpose of assistance to produce of the Tender investigation of the purpose of assistance of the tender investigation of the purpose of assistance of the tender investigation of the purpose of assistance of the tender investigation of tender investigation of the tender investigation of tender	cines are done in als order vide F. ss local and belief and on relevant records iting Authority/sessing the local		
				Name:				
				Designation				
				(Company Seal/Stamp				
				(To be furnished by	person in capacity	as per para 1)		

Note: The category of supplier against each quoted drug shall be mentioned in accordance with Order issued by Ministry of Chemicals & Fertilizers, Department of Pharmaceuticals vide no. *order no.* P-45021/2/2017-PP(BE-II) dated 16.09.2020 and as per table mentioned under Clause 3.U.

ANNEXURE-XI

Ref: Clause No. 15.E

Letter of acceptance of tender for Rate Contract

Speed post/e-mail

Ref. No. PMBI/SURGICAL/RC-210/2023	Date:
To,	
M/S	
	

Sub: Tender for the Supply of Surgical/ Consumables and Medical Devices to PMBI for two years: Acceptance tender for Rate Contract.

Ref: Your quotation against PMBI e-Tender No. PMBI/SURGICAL/RC-210/2023 dated: 15/06/2023 opened on (Technical Bid) & on (Price bid).

Please refer to your quotation i.e., technical and price bid (BOQ) along with enclosures/Annexure against subject tender read with your subsequent clarification/confirmation for the supply of Items to PMBI, the rate offered/accepted by your firm has been approved for Rate Contract for two years from the date of issue of this letter.

S. N.	Item Code	Item Name	Unit Size	Rates in Rs. Per unit exclusive of GST	

- 2. The contract will be with financial limit and PMBI can place the Purchase Order with unlimited variation in quantities indicated in the tender.
- 3. The estimated value of the contract awarded to you is Rs.....(in word).
- 4. Performance Security Deposit @3% will be deducted from each bill and accumulated security deposit will be refunded by PMBI to the tenderer within 60 days following the date of completion of tenderers performance obligations under the contract including the shelf-life obligation.
- 5. Approval for Artwork should be obtained from our Quality Control department by you within 30 days of release of this letter. (e-mail id: procure13@janaushadhi.gov.in; quality4@janaushadhi.gov.in; & quality5@janaushadhi.gov.in)
- 6. STP (Standard Testing Procedure) for Non- Pharmacopoeia awarded items are required to submit to Quality Control department (e-mail id: procure13@janaushadhi.gov.in; quality4@janaushadhi.gov.in; & quality5@janaushadhi.gov.in) within 15 days from the date of Letter of Acceptance.
- 7. As per clause 4. L of Tender document, the Rate Contract validity period may be extended for period up to further one year at same rate, terms & conditions with the consent of the supplier.
- 8. The terms and conditions of Rate Contract shall be applicable as mentioned in tender document. By issue of this acceptance letter, the Rate Contract is hereby concluded.

Please acknowledge receipt.

Authorized Signatory, For and on behalf of PMBI

Annexure -XII Clause 18 (M)

Pharmaceuticals & Medical Devices Bureau of India (PMBI), New Delhi Tender No. PMBI/SURGICAL/RC-210/2023 dated 15/06/2023)

(1)	(2)	(3)	(4)	(5)	(6)	(7)
S.N.	Item	Generic Name	Detail Specification of Item	Unit Size	Secondary	
	Code	of Item			Pack	Requiren ent in Unit Size
1	5001	Absorbent Cotton Wool IP 75g	Absorbent Cotton Wool IP Net weight of 75 g Made from 100% clean and soft cotton Natural fibres of average staple length not less than 10 mm, soft and fluffy, high absorption	1's in Paper rolled and in Sealed Poly pack	1's X 10 in sealed poly pack	1000000
2	5002	Absorbent Cotton Wool IP 200 g	Absorbent Cotton Wool IP Net weight of 200 g Made from 100% clean and soft cotton Natural fibres of average staple length not less than 10 mm, soft and fluffy, high absorption	1's in Paper rolled and in Sealed Poly pack	1's X 10 in sealed poly pack	500000
3	5003	Absorbent Cotton Wool IP 500 g	Absorbent Cotton Wool IP Net weight of 500 g Made from 100% clean and soft cotton Natural fibres of average staple length not less than 10 mm, soft and fluffy, high absorption	1's in Paper rolled and in Sealed Poly pack	1's X 10 in sealed poly pack	300000
4	5004	Crepe Bandage B.P. 15 cm x 4 M	Crepe Bandage B.P. 15 cm x 4 M Uniform plain weave and of continous length with no loose salvage threads (both sides fastened edges), Wrap threads- Two fold cotton threads, Not less than 170 threads per 10 cm. Weft threads- Combined cotton & Viscose, Not less than 78 threads per 10 cm. Elasticity- Not more than 80% of fully stretched length. washing regains elasticity-fit for re-use. Weight- Not Less Than 140 g / m2 03 Nos. Loop & Hook Closure	1's	Screw Cap PET bottle	500000
5	5005	Crepe Bandage B. P.10 cm x 4 M	Crepe Bandage B.P. 10 cm x 4 M Uniform plain weave and of continuous length with no loose salvage threads (both sides fastened edges), Wrap threads- Two-fold cotton threads, Not less than 170 threads per 10 cm. Weft threads- Combined cotton & Viscose, Not less than 78 threads per 10 cm. Elasticity- Not more than 80% of fully stretched length. washing regains elasticity-fit for re-use. Weight- Not Less Than 140 g / m2 02 Nos. Loop & Hook Closure	1's	Screw Cap PET bottle	500000
6	5006	Cotton Bandage 7.5 cm x 4 M	Cotton Bandages Size 7.5 cm X 4M, Freedom from optical whiteners and Conforming to the standards as per IS 863-1988, Count of yarn:	1's in Paper rolled and in Sealed Poly pack	1's X 12 in sealed poly pack	700000

Warp -NLT 25 tex Weft- NLT 30 tex Threads: Ends (column)- NLT 150 Picks (Row)- NLT 85		
Threads: Ends (column)- NLT 150 Picks (Row)- NLT 85		
Ends (column)- NLT 150 Picks (Row)- NLT 85		
Picks (Row)- NLT 85		
Picks (Row)- NLT 85		
Mass: NLT - 57 ± 5 g / m ²		
pH: 6.5 to 8.5		
	_	
Cotton Bandages Size 10 cm X 4M, Freedom from 1's in Pape		
optical whiteners and rolled and a		
Conforming to the standards as per IS 863-1988, Sealed Poly	У	
Count of yarn: pack		
Cotton Warp -NLT 25 tex	1's X 12	
7 5007 Bandage 10 Weft- NLT 30 tex	in sealed	700000
cm x 4 M Threads:	poly pack	
Ends (column)- NLT 150		
Picks (Row)- NLT 85		
Mass: NLT - 57 ± 5 g / m ²		
pH: 6.5 to 8.6		
Cotton Bandages Size 15 cm X 4M, Freedom from 1's in Pape	ſ.	
Cotton optical whiteners and Conforming to the standards rolled and		
8 Solve Bandages(No as per IS 863-1988, Count of yarn: Warp -NLT 25 Sealed Poly	17	500000
n Sterile) 15 texWeft- NLT 30 texThreads: Ends (column)-	poly pack	
cm x 4 M NLT 150Picks (Row)- NLT 85 Mass: NLT - 57 ±	pory pack	
5 g / m2 pH: 6.5 to 8.7		
Sterile Adhesive Bandages Wash proof 19mm X 1's in		
70 mm containing Benzalkonium Choride Solution individual		
IP 0.5 % w/w and tinted with tartazine yellow. sealed	1's X 100	
9 5009 Bandages The wound pad size NLT 25 mm and should be tearable	in Screw	800000
Washproof water resistant, provide antiseptic protection with paper	cap PET	
nigh quality PVC as backing material. wrapper	jar.	
Skin friendly & Hypoallergic adhesive, no sticky		
residue.		
1's in		
Disposable Syringe 2 ml, with colour coded (as per individual	1's X 100	
Syringe 2 ml RIS) needle 24G Starilised Luar Mount Non Sealed	in packed	
10 5010 with needle toyic as per Drugs and Cosmetics Act 1940 IS No tearable	in mono	5000000
10258- 2002 with BIS/ CE/ USFDA certification.	carton.	
	carton.	
wrapper		
l's in		
Syringe 5 ml Disposable Syringe5 ml, with colour coded (as per individual	1's X 100	
11 5011 with needle BIS) needle 24G Sternised, Luer Mount, Non - Sealed	in packed	6000000
I foxic as per Driigs and Cosmetics Act 1940 IS No Itearable	in mono	0000000
10258- 2002 with BIS/ CE/ USFDA certification.	carton.	
wrapper		
1's in		
	1'e V 50	
Syringe	1's X 50	
12 5012 ml with per BIS) needle 21G Sterilised, Luer Mount, Non - Sealed	in packed	2500000
14 1/V14 1 HI WILL	in mono	
needle 21G toxic as per Drugs and Cosmetics Act 1940, IS No. tearable	carton.	
toxic as per Drugs and Cosmetics Act 1940, IS No. tearable 10258- 2002 with BIS/ CE/ USFDA certification.		
needle 21G toxic as per Drugs and Cosmetics Act 1940, IS No. tearable		
needle 21G toxic as per Drugs and Cosmetics Act 1940, IS No. tearable 10258- 2002 with BIS/ CE/ USFDA certification. poly		
needle 21G toxic as per Drugs and Cosmetics Act 1940, IS No. tearable 10258- 2002 with BIS/ CE/ USFDA certification. poly wrapper Disposable Syringe 10 ml, with colour coded (as individual)	1's X 50	
needle 21G toxic as per Drugs and Cosmetics Act 1940, IS No. tearable 10258- 2002 with BIS/ CE/ USFDA certification. poly wrapper 1's in Disposable Syringe 10 ml, with colour coded (as individual per BIS) needle 23G Sterilised Lucr Mount, Non. Sealed	1's X 50	
toxic as per Drugs and Cosmetics Act 1940, IS No. tearable 10258- 2002 with BIS/ CE/ USFDA certification. poly wrapper Syringe 20 ml with Disposable Syringe10 ml, with colour coded (as per BIS) needle 23G Sterilised, Luer Mount, Non - Sealed toxic as per Drugs and Cosmetics Act 1940, IS No. tearable toxic as per Drugs and Cosmetics Act 1940, IS No. tearable	in packed	500000
needle 21G toxic as per Drugs and Cosmetics Act 1940, IS No. Itearable 10258- 2002 with BIS/ CE/ USFDA certification. Syringe 20 ml with peedle 23G ml with needle 23G Toxic as per Drugs and Cosmetics Act 1940, IS No. Itearable poly wrapper 1's in individual Sealed toxic as per Drugs and Cosmetics Act 1940, IS No. Itearable poly wrapper	in packed in mono	500000
toxic as per Drugs and Cosmetics Act 1940, IS No. tearable 10258- 2002 with BIS/ CE/ USFDA certification. poly wrapper Syringe 20 ml with Disposable Syringe10 ml, with colour coded (as per BIS) needle 23G Sterilised, Luer Mount, Non - Sealed toxic as per Drugs and Cosmetics Act 1940, IS No. tearable toxic as per Drugs and Cosmetics Act 1940, IS No. tearable	in packed	500000

14	5014	Needle 16G	The needles fit the luer nozzle, Sterile, single use, non-toxic, non-pyrogenic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in blister pack	1's X 100 in packed in mono carton.	250000
15	5015	Needle 18G	The needles fit the luer nozzle, Sterile, single use, non-toxic, non-pyrogenic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in blister pack	1's X 100 in packed in mono carton.	250000
16	5016	Needle 23G	The needles fit the luer nozzle, Sterile, single use, non-toxic, non-pyrogenic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in blister pack	1's X 100 in packed in mono carton.	250000
17	5017	Needle 26G half inch	The needles fit the luer nozzle, Sterile, single use, non-toxic, non-pyrogenic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in blister pack	1's X 100 in packed in mono carton.	250000
18	5019	Paper Adhesive Plaster 1 inch X 9m	Size-1 inch X 9 m per roll Material: Non-woven, non-allergic & Strong adhesive but gentle to skin & painless removal, Porous and breathable CE cerified	1's	1's X 12 in monocart on	600000
19	5020	Paper Adhesive Plaster 2 inch X 9m	Size-2 inch X 9 m per roll Material: Non-woven, non-allergic & Strong adhesive but gentle to skin & painless removal, Porous and breathable CE cerified	1's	1's X 12 in monocart on	400000
20	5021	Paper Adhesive Plaster 3 inch X 9m	Size-3 inch X 9 m per rollMaterial: Non-woven, non-allergic & Strong adhesive but gentle to skin & painless removal, Porous and breathableCE cerified	1's	1's X 12 in monocart on	300000
21	5022	Plaster of Paris BP Bandages 15cm X 2.7m	Plaster of Paris Bandages BP 15 cm X 2.7 m per Roll Joint free single roll Conforming to the standards as per IS 6237 : 1971 (Reaffirmed Year : 2018) Count of yarn: Warp -NLT 21 tex Weft- NLT 21 tex Threads per dm: Ends (column)- NLT 150 Picks (Row)- NLT 75 Mass : NLT - 40 g / m2 Fast Setting, Superior casting Strength, Reliable and longer lifespan, complying with standards as per British Pharmacopoeia.	1's in Paper rolled and in Sealed Poly pack	1's X 06	600000
22	5023	Plaster of Paris BP Bandages 10cm X 2.7m / Roll	Plaster of Paris Bandages BP 10 cm X 2.7 m per Roll Joint free single roll Conforming to the standards as per IS 6237: 1971 (Reaffirmed Year: 2018) Count of yarn: Warp -NLT 21 tex Weft- NLT 21 tex Threads per dm: Ends (column)- NLT 150 Picks (Row)- NLT 75	1's in Paper rolled and in Sealed Poly pack	1's X 06 in sealed poly pack	400000

			Mass: NLT - 40 g / m2 Fast Setting, Superior casting Strength, Reliable and longer lifespan, complying with standards as per British Pharmacopoeia.			
23	5024	Scalp Vein Set (Disposable) (18G)	Scalp Vein Set (Disposable) (18G) Disposable Stain less Steel sharp needle, siliconised, Butterfly wings, Latex free PVC for better handling and fixation. soft flexible PUR/PVC tubing of 150-300 mm Length including lyer connector and Cap. ETO Sterilised with CE certification	1's in blister pack	1's X 06 in monocart on	250000
24	5025	Scalp Vein Set (Disposable) (20G)	Scalp Vein Set (Disposable) (20G) Disposable Stain less Steel sharp needle, siliconised, Butterfly wings, Latex free PVC for better handling and fixation. soft flexible PUR/PVC tubing of 150-300 mm Length including lyer connector and Cap. ETO sterilized with CE certification	1's in blister pack	1's X 06 in monocart on	250000
25	5026	Scalp Vein Set (Disposable) (22G)	Scalp Vein Set (Disposable) (22G) Disposable Stain less Steel sharp needle, siliconized, Butterfly wings, Latex free PVC for better handling and fixation. soft flexible PUR/PVC tubing of 150-300 mm Length including lyer connector and Cap. ETO Sterilized with CE certification	1's in blister pack	1's X 06 in monocart on	250000
26	5027	Scalp Vein Set (Disposable) (24G)	Scalp Vein Set (Disposable) (24G)Disposable Stain less Steel sharp needle, siliconized, Butterfly wings, Latex free PVC for better handling and fixation. soft flexiblePUR/PVC tubing of 150-300 mm Lengthincluding lyer connector and Cap. ETO sterilized with CE certification	1's in blister pack	1's X 06 in monocart on	250000
27	5030	Surgical Blade, No. 22, Sterilized	Surgical Blade, Size No. 22 Stainless-steel, well-defined tip and uniform cutting edge. Pre - sterile with Gamma radiation, confirm to the standards as per IS No.: 3319:1995 with CE certification.	1's in peelable Aluminium foil pack	1's X 50 in monocart on	250000
28	5033	Surgical Blade, No. 15, Sterilized	Surgical Blade, Size No. 15 Stainless-steel, well-defined tip and uniform cutting edge. Pre - sterile with Gamma radiation, Confirm to the standards as per IS No.: 3319:1995 with CE certification.	1's in peelable Aluminium foil pack	1's X 50 in monocart on	250000
29	5037	I.V Cannula (Sterile, Disposable), 26 G	Cannula with Integrated 3-Way stop Cock. Size 26G having radio opaque catheter with CE certification. Sterile Disposable (Single Use) Teflon/ PTFE I.V.	1's in blister pack	1's X 50 in monocart on	250000
30	5039	Sterile Disposable Spinal Needle 22G x 3 ½ inch	Sterile Disposable Spinal Needle 22G x 3 ½ inch. Specially designed to administer lumbar/subarachnoid anesthesia. Transparent hub provides rapid detection of Cerebro-spinal fluid (CSF flashback for confirming accurate placement. Sharp bevel design for low puncture force ensures minimal puncture trauma. Fine gauze needle	1's in blister pack	1's X 50 in monocart on	250000

			design greatly reduces the risk of PDPH (Post			
			Dura Puncture Headache). Color coded as per ISO			
			standard enables rapid size identification.)	41 1 11 .		
31	5040	Sterile Disposable Spinal Needle 25G x 3 ½ inch	Sterile Disposable Spinal Needle 25G x 3 ½ inch. Specially designed to administer lumbar/subarachnoid anesthesia. Transparent hub provides rapid detection of Cerebro-spinal fluid (CSF flashback for confirming accurate placement. Sharp bevel design for low puncture force ensures minimal puncture trauma. Fine gauze needle design greatly reduces the risk of PDPH (Post Dura Puncture Headache). Color coded as per ISO standard enables rapid size identification.)	1's in blister pack	1's X 50 in monocart on	250000
32	5041	Urine Collecting Bag, Disposable, 2000ml	Sterile Disposable Urine Collecting Bag 2000 ml with graduated volume marking, non-toxic pyrogen free, double seek, clinical grade PVC, Kink resistant flexible tubing not less than 90 cm in length, should have non-return valve, Top drainage outlet, with non-return input valve, CE certified.	1's pack in sealed poly pack	1's X 25 in monocart on	600000
33	5046	Endotracheal Tube Plain, Size 2.5, Sterile, Single use	ENDOTRACHEAL TUBE PLAIN SIZE 2.5 Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
34	5047	Endotracheal Tube Plain, Size 3, Sterile, Single use	ENDOTRACHEAL TUBE PLAIN SIZE 3 Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
35	5048	Endotracheal Tube Plain, Size 3.5, Sterile, Single use	ENDOTRACHEAL TUBE PLAIN SIZE 3 Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	100000
36	5049	Endotracheal Tube Plain,	ENDOTRACHEAL TUBE PLAIN SIZE 4 Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
37	5050	Endotracheal Tube Plain,	ENDOTRACHEAL TUBE PLAIN SIZE 4.5 Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
38	5051	Endotracheal Tube Plain, Size 5, Sterile, Single use	ENDOTRACHEAL TUBE PLAIN SIZE 5Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
39	5052	Endotracheal Tube Plain, Size 8.5, Sterile, Single use	ENDOTRACHEAL TUBE PLAIN SIZE 8.5 Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000

40	5053	Endotracheal Tube Cuffed, Size 4, Sterile, Single use	ENDOTRACHEAL TUBE CUFFED SIZE 4 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
41	5054	Endotracheal Tube Cuffed, Size 4.5, Sterile, Single use	ENDOTRACHEAL TUBE CUFFED SIZE 4.5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
42	5055	Endotracheal Tube Cuffed, Size 5, Sterile, Single use	ENDOTRACHEAL TUBE CUFFED SIZE 5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
43	5056	Endotracheal Tube Cuffed, Size 6, Sterile, Single use	ENDOTRACHEAL TUBE CUFFED SIZE 6 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
44	5057	Endotracheal Tube Cuffed, Size 6.5, Sterile, Single use	ENDOTRACHEAL TUBE CUFFED SIZE 6.5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
45	5058	Endotracheal Tube Cuffed, Size 7, Sterile, Single use	ENDOTRACHEAL TUBE CUFFED SIZE 7 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
46	5059	Endotracheal Tube Cuffed, Size 7.5, Sterile, Single use	ENDOTRACHEAL TUBE CUFFED SIZE 7.5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
47	5060	Endotracheal Tube Cuffed,	ENDOTRACHEAL TUBE CUFFED SIZE 8 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
48	5061	Endotracheal Tube Cuffed,	ENDOTRACHEAL TUBE CUFFED SIZE 8.5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
49	5062	Endotracheal Tube Cuffed,	ENDOTRACHEAL TUBE CUFFED SIZE 9 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack	1's X 25 in monocart on	50000
50	5064	Tracheostom y Tube (PVC), Sterilised, single use Size-24	Tracheostomy Tube (PVC), sterilized, single use(Tracheostomy Tube, made frommedical grade PVC, Plain, Sterile, Single Use - Size 24 Soft flexible flange at for easy fixation 15 mmconnector at terminal end which canbe rotated in 360-degree directionNon-irritant, Radio-opaque	1's pack in sealed blister pack	1's X 25 in monocart on	50000

			linethroughout the length. Size: 20 X60mm)Confirming to the standards IS 13179: 1991 (Reaffirmed 2022)			
51	5065	Abdominal Drainage Kit, Sterile (2000ml, 24 Graduate)	Abdominal Drainage Kit, Sterile (2000ml, 24 Graduate) ABDOMINAL DRAIN KIT, STERILE, HAVING DRAINAGE CATHETER AND COLLECTION BAG {2000 ml} size 24 Graduated Bag, Soft drainage catheter 50 cm long, with radio opaque line, Catheter with markings at 2 cm interval	1's		150000
52	5066	Abdominal Drainage Kit, Sterile (2000ml, 28 Graduate)	Abdominal Drainage Kit, Sterile (2000ml, 28 Graduate) ABDOMINAL DRAIN KIT, STERILE, HAVING DRAINAGE CATHETER AND COLLECTION BAG {2000 ml} size 28 Graduated Bag, Soft drainage catheter 50 cm long, with radio opaque line, Catheter with markings at 2 cm interval	1's		150000
53	5067	Corrugated Drainage Sheet	Corrugated Drainage Sheet Nontoxic, non-irritant medical grade extra soft PVC, Radio opaque line, Sterile, individually packed in a H.M polybag PVC does not contribute to local inflammation.	1's in High molecular PVC polybag	1's X 25 in monocart on	250000
54	5070	Sterilized Umbilical Cotton Tape 3mm x 75cm	ETO Sterilized Umbilical Cotton Tape 3mm x 75cm	1's in sealed poly pack	1's X 12 in monocart on	250000
55	5071	Bone Wax, Sterilised 2.5	Sterilized Bone wax 2.5 g	1's in peelable Aluminium foil pack	1's X 12 in monocart on	250000
56	5073	Crepe Bandage B.P. 6 cm x 4 M	Crepe Bandage B.P. 6 cm x 4 M Uniform plain weave and of continous length with no loose salvage threads (both sides fastened edges), Wrap threads- Two fold cotton threads, Not less than 170 threads per 10 cm. Weft threads- Combined cotton & Viscose, Not less than 78 threads per 10 cm. Elasticity- Not more than 80% of fully stretched length.washing regains elasticity-fit for re-use. Weight- Not Less Than 140 g / m2 02 Nos. Loop & Hook Closure	1's	Screw Cap PET bottle	700000
57	5077	Rapid Diagnostic Malaria Test Kit	Rapid Diagnostic Malaria Test Kit should contain all the materils required for performing the test i.e. heparinized capillary tubes (Dia- 1 mm) with relevant markings, reaction tube& stand, Test card (cassette); Sterile lancet, Reagents including buffer solution in a dropping bottle and Alcohol Swab. The test kit should be able to rapidly diagnose both P. falciparum (HRP2) and P. vivax (pLDH) and	1's in hermetically sealed with poly pouch	1's X 10 in monocart on	500000

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58	5078	Dengue Test Kit	based on the principle of capture of parasite antigen from blood using monoclonal antibodies specific for antigen targets. PDS for Pf should be at least 95% sensitivity and specificity and for Pv should be at least 75% sensitivity and 90% specificity both at 200 parasites / μL. the false positive rate should be less than 10% and invalid rate should be less than 5%. Storage- 30oC to 45oCThe product should comply with ISO 9001 ISO 13485 (QMS)/CE/USFDA. Each Lot should PASS the test from ICMR designated labs. The ELISA kit should be designed for qualitative detection of dengue NS1 antigen of all 4 dengue serotypes in human serum. The kit should be provided with the following materials and reagents: a) Anti- NS1 Antibody Coated Breakway Microwells (12*8=96 wells). Desiccant should be provided for storing the unused microwells which are to be resealed immediately. b) Horseradish peroxidase conjugated Anti-NS1 monoclonal antibody with preservatives. c) Chromogenic substrate in buffer d) Positive Control in the form of recombinant antigen with preservatives and antibiotics. e) Negative control in the form of confirmed negative human serum with preservatives and antibiotics. f) Calibrators in the form of recombinant antigen with preservatives and antibiotics. f) Calibrators in the form of recombinant antigen with preservatives and antibiotics. f) Calibrators in the form of dengue NS1 antigen should have a sensitivity of >90% and a specificity of >95% taking RT-PCR as the gold standard. The kit should have a shelf-life of at least 6 months when stored at an ambient temperature of 2°C - 8°C. Transportation should be under cold chain. CE certified		1's X 10 in monocart on	50000
59	5079	HCG- Pregnancy Test Kit	One step rapid qualitative test for detecting HCG pregnancy hormone in urine. Each Test Kit contains: 01 Test device with aperture marked as 'S', 'C' for control and 'T' for test line. 01 Dropper, 01 Clean cup/tube. Reading time 5-10 min Hermetically sealed CE certified		1's X 10 in monocart on	1500000
60	5083	Ear buds 100% Cotton	Ear buds 100% natural Absorbent Cotton IP free from optical whiteners/brighteners,	100's in Screw cap	1's X 20 in	1000000

			Stronge paper/wood stems, Cotton swabs should never debond while using/ cleaning. Manufactured on fully automated machines and Untouched with Hands Size: 8 cm ± 0.5 cm	pet jar	monocart on	
61	5094	Sterile Gauze Pad size 2X2	100% woven natural Absorbent Cotton IP, individually wrapped and sterile. Lint-free and have no loose threads or raw edges. Size: 2 in X 2 in Confirming to the standards as per IS No.758/1988	1's in sealed poly pack	1's X 25 in monocart on	300000
62	5095	Sterile Gauze Pad size 3X3	100% woven natural Absorbent Cotton IP, individually wrapped and sterile. Lint-free and have no loose threads or raw edges. Size: 3 in X 3 in Confirming to the standards as per IS No.758/1989	1's in sealed poly pack	1's X 12 in monocart on	300000
63	5096	Sterile Gauze Pad size 4X4	100% woven natural Absorbent Cotton IP, individually wrapped and sterile.Lint-free and have no loose threads or raw edges.Size: 4 in X 4 inConfirming to the standards as per ISNo.758/1990	1's in sealed poly pack	1's X 10 in monocart on	500000
64	5097	Alcohol Swab (Spirit Swab)	Alcohol Swab (Spirit Swab) Four-layer wrapper provides air-tight seal - prevents leakage and drying out. Isopropyl Alcohol 70% V/V USP for optimum anti-bacterial action.	1's inair- tight sealed wrapper	1's X 100 in monocart on	1500000
65	6001	Surgical Rubber Gloves- Disposable, Sterile, 6.5 inch	One pair of Disposable Sterile Surgical Rubber Gloves, Size 6 ½ Inches, Made of natural hypoallergic Latex, 100% electronically tested for holes, sterilized by Gamma Radiation / ETO, pre-powdered, Tensile strength as per EN 455-2. Powder should be non-allergenic should Conform to IS 13422. ISI marked / CE certified / FDA.	1's pair in sealed poly pouch	1's pair X 25 in mono carton	2500000
66	6002	Surgical Rubber Gloves- Disposable, Sterile, 7 inch	One pair of Disposable Sterile Surgical Rubber Gloves, Size 7 Inches, Made of natural hypoallergic Latex, 100% electronically tested for holes, sterilized by Gamma Radiation / ETO, pre-powdered, Tensile strength as per EN 455-2. Powder should be non- allergenic should Conform to IS 13422. ISI marked / CE certified / FDA.	1's pair in sealed poly pouch	1's pair X 25 in mono carton	3000000
67	6003	Surgical Rubber Gloves- Disposable, Sterile, 7.5 inch	One pair of Disposable Sterile Surgical Rubber Gloves, Size 7.5 Inches, Made of natural hypoallergic Latex, 100% electronically tested for holes, sterilized by Gamma Radiation / ETO, pre-powdered, Tensile strength as per EN 455-2. Powder should be non-allergenic should Conform to IS 13422.	1's pair in sealed poly pouch	1's pair X 25 in mono carton	1500000

			ISI marked / CE certified / FDA.			
68	6004	Surgical Rubber Gloves- Disposable, Sterile, 8 inch	One pair of Disposable Sterile Surgical Rubber Gloves, Size 8 Inches, Made of natural hypoallergic Latex, 100% electronically tested for holes, sterilized by Gamma Radiation / ETO, pre-powdered, Tensile strength as per EN 455-2. Powder should be non- allergenic should Conform to IS 13422. ISI marked / CE certified / FDA.	1's pair in sealed poly pouch	1's pair X 25 in mono carton	300000
69	6009	Surgical Cap, Disposable(fo r Surgeons/Nur ses)	Disposable Surgical Cap, Should be manufactured from non-woven fabric Blue / Green / color, Round upon wearing, with elastic Air permeable / breathable Should retain skin and hair particles	1's	1's pair X 100 in PVC pack	250000
70	6010	Suction Catheter FG (5)	Suction Catheter FG (5), 50 cm, sterilized Non-traumatic tip, non-toxic medical grade PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Color code for identification of size.	1's in sealed poly pack	1's X 25 in mono carton	250000
71	6011	Suction Catheter FG (6), 50 cm, Sterilized	Suction Catheter FG (6), 50 cm, sterilized non-traumatic tip, non-toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Color code for identification of size. CE certified	1's in sealed poly pack	1's X 25 in mono carton	250000
72	6012	Suction Catheter FG (8), 50 cm, Sterilized	Suction Catheter FG (8), 50 cm, Sterilized non-traumatic tip, non-toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Colour code for identification of size.CE certified	1's in sealed poly pack	1's X 25 in mono carton	250000
73	6013	Suction Catheter FG (10), 50 cm, Sterilized	Suction Catheter FG (10), 50 cm, sterilized, non-traumatic tip, non-toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Color code for identification of size, CE certified	1's in sealed poly pack	1's X 25 in mono carton	250000
74	6014	Suction Catheter FG (12), 50 cm, Sterilized	Suction Catheter FG (12), 50 cm, sterilized non-traumatic tip, non-toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Color code for identification of size. CE certified	1's in sealed poly pack	1's X 25 in mono carton	250000
75	6015	Suction Catheter FG (14), 50 cm, Sterilized	Suction Catheter FG (14), 50 cm, sterilized non-traumatic tip, non-toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Color code for identification of size. CE certified	1's in sealed poly pack	1's X 25 in mono carton	250000
76	6016	Suction Catheter FG (16), 50 cm, Sterilized	Suction Catheter FG (16), 50 cm, sterilized non-traumatic tip, non-toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Color code for identification of size. CE certified	1's in sealed poly pack	1's X 25 in mono carton	250000
77	6017	Suction Catheter FG (18), 50 cm, Sterilized	Suction Catheter FG (18), 50 cm, sterilized non-traumatic tip, non-toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Color code for identification	1's in sealed poly pack	1's X 25 in mono carton	250000

			of size. CE certified			
78	6018	Suction Catheter FG (20), 50 cm, Sterilized	Suction Catheter FG (20), 50 cm, sterilized non-traumatic tip, non-toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Color code for identification of size. CE certified	1's in sealed poly pack	1's X 25 in mono carton	250000
79	6020	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (8FG)	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (8FG) 2 Way Silicone Coated Latex Foley Catheter, thin, flexible and sterile tube. Made of Silicone elastomer bonded with Latex. Should have hard plastic valve Smooth distal end with smooth eyes for atraumatic intubation. Symmetrical foley balloon Balloon capacity 3- 5 ml. Should conform to IS 11497. Color coding marking to identify size. Length, wall thickness and balloon capacity should be mentioned as per IS 11497. Specification for B, C, D, E, F, G should be mentioned as per IS 11497 for particular size).	1's in sealed poly pack	1's X 10 in mono carton	250000
80	6021	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (10FG)	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (10FG) 2 Way Silicone Coated Latex Foley Catheter, thin, flexible and sterile tube. Made of Silicone elastomer bonded with Latex. Should have hard plastic valve Smooth distal end with smooth eyes for atraumatic intubation. Symmetrical foley balloon Balloon capacity 3- 5 ml. Should conform to IS 11497. Color coding marking to identify size. Length, wall thickness and balloon capacity should be mentioned as per IS 11497. Specification for B, C, D, E, F, G should be mentioned as per IS 11497 for particular size). CE certified	1's in sealed poly pack	1's X 10 in mono carton	250000
81	6022	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (16FG)	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (16FG) 2 Way Silicone Coated Latex Foley Catheter, thin, flexible and sterile tube. Made of Silicone elastomer bonded with Latex. Should have hard plastic valve Smooth distal end with smooth eyes for atraumatic intubation. Symmetrical foley balloon Balloon capacity 3-5 ml. Should conform to IS 11497. Color coding marking to identify size. Length, wall thickness and balloon capacity should be mentioned as per IS 11497. Specification for B, C, D, E, F, G should be mentioned as per IS 11497 for particular size)	1's in sealed poly pack	1's X 10 in mono carton	250000
82	6023	Sterile Catheter Single Use	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (18FG)2 Way Silicone Coated Latex Foley Catheter, thin,	1's in sealed poly pack	1's X 10 in mono carton	250000

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		for Urinary	flexible and sterile tube.Made of Silicone			
		Drainage	elastomer bonded with Latex. Should have hard			
		(Foley	plastic valve Smooth distal end with smooth eyes			
		Balloon	for atraumatic intubation. Symmetrical foley			
		Catheter), 2	balloon Balloon capacity 3- 5 ml.Should conform			
		way (18FG)	to IS 11497. Color coding marking to identify size.			
		• , ,	Length, wall thickness and balloon capacity should			
			be mentioned as per IS 11497. Specification for			
			B,C,D,E,F,G should be mentioned as per IS 11497			
			for particular size)			
			Sterile Catheter Single Use for Urinary Drainage	1's in sealed		
			(Foley Balloon Catheter), 2 way (20FG)	poly pack		
			2 Way Silicone Coated Latex Foley Catheter, thin,	pory pack		
		Sterile	flexible and sterile tube.			
		Catheter	Made of Silicone elastomer bonded with Latex.			
		Single Use	Should have hard plastic valve Smooth distal end			
		for Urinary	<u>-</u>		1's X 10	
83	6024	•	with smooth eyes for atraumatic intubation.			250000
03	0024	U	Symmetrical foley balloon Balloon capacity 3- 5 ml.		in mono carton	250000
		(Foley Balloon	Should conform to IS 11497.		Carton	
		Catheter), 2				
		way (20FG)	Color coding marking to identify size. Length,			
		way (20FG)	wall thickness and balloon capacity should be			
			mentioned as per IS 11497.			
			Specification for B, C, D, E, F, G should be			
			mentioned as per IS 11497 for particular size)	1's in sealed		
			Sterile Catheter Single Use for Urinary Drainage			
			(Foley Balloon Catheter), 2 way (22FG)	poly pack		
		C4amila	2 Way Silicone Coated Latex Foley Catheter, thin,			
		Sterile	flexible and sterile tube. Made of Silicone elastomer bonded with Latex.			
		Catheter				
		Single Use	Should have hard plastic valve Smooth distal end		1!a V 10	
0.4	6025	for Urinary	with smooth eyes for atraumatic intubation.		1's X 10	250000
84	6025	Drainage	Symmetrical foley balloon Balloon capacity 3-5		in mono	250000
		(Foley	ml.		carton	
		Balloon	Should conform to IS 11497.			
		Catheter), 2	Color coding marking to identify size. Length,			
		way (22FG)	wall thickness and balloon capacity should be			
			mentioned as per IS 11497.			
			Specification for B, C, D, E, F, G should be			
			mentioned as per IS 11497 for particular size)	110 20 1 1		1
			Sterile Catheter Single Use for Urinary Drainage	1's in sealed		
			(Foley Balloon Catheter), 2 way (24FG)	poly pack		
		C4!1 -	2 Way Silicone Coated Latex Foley Catheter, thin,			
		Sterile	flexible and sterile tube.			
		Catheter	Made of Silicone elastomer bonded with Latex.			
		Single Use	Should have hard plastic valve Smooth distal end		11 77 10	
0.7	c02 -	for Urinary	with smooth eyes for atraumatic intubation.		1's X 10	250000
85	6026	Drainage	Symmetrical foley balloon Balloon capacity 3- 5		in mono	250000
		(Foley	ml.		carton	
		Balloon	Should conform to IS 11497.			
		Catheter), 2	Color coding marking to identify size. Length,			
		way (24FG)	wall thickness and balloon capacity should be			
			mentioned as per IS 11497.			
1			I Simportugation ton D. (* IV. II. II. (* abould be	•	i	i .
			Specification for B, C, D, E, F, G should be mentioned as per IS 11497 for particular size)			

86	6027	Infant Feeding Tube, Sterile (Size-10 FG, length 50 cm)	Infant Feeding Tube, Sterile (Size-10 FG, length 50 cm) Sterile, DEHP Free, non-toxic PVC, non-irritant, low friction, non-traumatic. CE certified	1's in sealed poly pack	1's X 10 in mono carton	250000
87	6028	Infant Feeding	Infant Feeding Tube, Sterile (Size-8 FG, length 50 cm) Sterile, DEHP Free, non-toxic PVC, non-irritant, low friction, non-traumatic. CE certified	1's in sealed poly pack	1's X 10 in mono carton	250000
88	6029	Infant Feeding Tube, Sterile (Size-5 FG, length 50 cm)	Infant Feeding Tube, Sterile (Size-5 FG, length 50 cm) Sterile, DEHP Free, non-toxic PVC, non-irritant, low friction, non-traumatic. CE certified	1's in sealed poly pack	1's X 10 in mono carton	250000
89	6031	Sterile Disposable Infusion set with Microdrip (I.V.) (Pediatric use)	Sterile Disposable Perfusion Set (Infusion set) with built in Airway molded chamber and Needle (Pediatric Use). Burette type measured volume chamber of 100 ml, Non - Toxic, Non-Pyrogenic, sterilized by ETO. Drop size of approx. 60 drops per ml Injection port, latex free, for intermittent medication. Floating auto shut off valve (latex free) in burette. Soft and kink resistant PVC tubing 2.7 to 3.00 mm tube with fluid filter. Roller controller for flow control Tube length 150 cm 23G needle. Should conform to IS No. 12655 (part-4 of 2003) with CE certification.	1's in sealed poly pack	1's X 25 in mono carton	250000
90	6032	Sterile Disposable Infusion set with Microdrip (I.V.) (Adult use)	Sterile Disposable Perfusion Set (Infusion set) with built in Airway molded chamber and Needle (Adult Use). Burette type measured volume chamber of 100 ml, Non - Toxic, Non-Pyrogenic, sterilized by ETO. Drop size of approx. 60 drops per ml Injection port, latex free, for intermittent medication. Floating auto shut off valve (latex free) in burette. Soft and kink resistant PVC tubing 2.7 to 3.00 mm tube with fluid filter. Roller controller for flow control Tube length 150 cm 23G needle. Should conform to IS No.12655 (part-4 of 2003) with CE certification.	1's in sealed poly pack	1's X 25 in mono carton	500000
91	6033	Insulin Syringe (40 units) with 30G needle	Insulin syringe (40 units) with (fixed) 30 G needle conform to IS 12227 and CE certified. Sterile and non- toxic	1's in sealed blister pack	1's X 100 in mono carton	5500000
92	6035	I.V Cannula (Sterile, Disposable), 18 G	I.V Cannula (Sterile, Disposable), 18 G Sterile Disposable (Single Use) Teflon/ PTFE I.V. Cannula with Integrated 3-Way stop Cock having radio opaque catheter with CE certification. Size 18G, Should conform to IS 10555	1's in sealed blister pack	1's X 50 in mono carton	500000
93	6036	I.V Cannula (Sterile, Disposable), 20 G	I.V Cannula (Sterile, Disposable), 20 G Sterile Disposable (Single Use) Teflon/ PTFE I.V. Cannula with Integrated 3-Way stop Cock having radio opaque catheter with CE certification. Size 20G, Should conform to IS 10555 Standard	1's in sealed blister pack	1's X 50 in mono carton	500000

94	6037	I.V Cannula (Sterile, Disposable), 22 G	I.V Cannula (Sterile, Disposable), 22 G Sterile Disposable (Single Use) Teflon/ PTFE I.V. Cannula with Integrated 3-Way stop Cock having radio opaque catheter with CE certification. Size 22G, Should conform to IS 10555 Standard	1's in sealed blister pack	1's X 50	500000
95	6038	I.V Cannula (Sterile, Disposable), 24 G	I.V Cannula (Sterile, Disposable), 24 G Sterile Disposable (Single Use) Teflon/ PTFE I.V. Cannula with Integrated 3-Way stop Cock having radio opaque catheter with CE certification. Size 24G, Should conform to IS 10555 Standard	1's in sealed blister pack	1's X 50 in mono carton	500000
96	6039	Nasal Oxygen Cannula set, Twin bore (Adult)	Nasal Oxygen Cannula Set, Twin prong (Accessory for Compressed Air Breathing) Adult General specification: Suitable for easy application and efficient administration of oxygen. soft, light weight non- toxic material, non-irritating even in long term use anatomically fit Twin prong/ nasal tips are designed to ensure equal volume of oxygen to both the air passages. CE certified	1's in sealed blister pack	1's X 25 in mono carton	250000
97	6040	Absorbent Gelatin Sponge, Sterilized (80mm x 50mm x 10mm)	Water-insoluble, hemostatic device prepared from purified skin gelatin, and capable of absorbing up to 45 times its weight of whole blood. Gamma Sterile & Ready To Use, Easy To Cut According To Need.	1's in sterile blister pack enclosed in an outer peelable wrapper	1's X 2 in mono carton	250000
98	6042	Blood Transfusion Set with Filter 170 Micron	Blood Transfusion Set 170 Micron(The soft Kink resistance, translucent tubing is prepared from medical grade PVC material, double drip clearly visible chamber facilitate visual access and rapid adjustment of fluid level. Specially design roller controller offers accurate regulation of infusion rate with self-sealing latex bulb to avoid any contamination and Easy flushing. Length 150 cm.) CE certified	1's in sealed blister pack	1's X 25 in mono carton	600000
99	6045	Ryle's Tube/Nasoga stric Tube (Size 10)	Soft kink, resistance PVC tubing for atraumatic intubation. Closed distal end should be coned with radio opaque Material for accurate intubation. Four lateral eyes for greater efficiency. Radio Opaque Line. Marking at 50, 60,70 cm from trip. Color coded funnel. With luer connecter / closure. Length 105 cm. CE certified	1's in sealed blister pack	1's X 25 in mono carton	250000
100	6046	Ryle's Tube/Nasoga stric Tube (Size 12)	Soft kink, resistance PVC tubing for atraumatic intubation. Closed distal end should be coned with radio opaque Material for accurate intubation. Four lateral eyes for greater efficiency. Radio Opaque Line. Marking at 50, 60,70 cm from trip. Color coded funnel. With luer connecter / closure. Length 105 cm. CE certified	1's in sealed blister pack	1's X 25 in mono carton	250000
101	6047	Ryle's Tube/Nasoga	Soft kink, resistance PVC tubing for atraumatic intubation.	1's in sealed blister pack		250000

		stric Tube	Closed distal end should be coned with radio		carton	
		(Size 14)	opaque Material for accurate intubation. Four lateral eyes for greater efficiency. Radio			
			Opaque Line. Marking at 50, 60,70 cm from trip. Color coded funnel. With luer connecter / closure. Length 105 cm. CE certified			
			Soft kink, resistance PVC tubing for atraumatic intubation. Closed distal end should be coned with radio	1's in sealed blister pack		
102	6048	Ryle's Tube/Nasoga stric Tube (Size 16)	opaque Material for accurate intubation. Four lateral eyes for greater efficiency. Radio Opaque Line. Marking at 50, 60,70 cm from trip. Color coded funnel. With luer connecter / closure. Length 105 cm. CE		1's X 25 in mono carton	250000
			certified Soft kink, resistance PVC tubing for atraumatic intubation.	1's in sealed blister pack		
103	6049	Ryle's Tube/Nasoga stric Tube (Size 18)	Closed distal end should be coned with radio opaque Material for accurate intubation. Four lateral eyes for greater efficiency. Radio Opaque Line. Marking at 50, 60,70 cm from trip. Color coded funnel. With luer connecter / closure. Length 105 cm. CE certified		1's X 25 in mono carton	250000
104	6050	Knee Cap (Small)	Made of interlock weave bi-layered fabric (outer layer made of nylon and inner layer made of fine grade dermophillic & hypoallergenic cotton with inbetween air space), Four way stretchable, Heat resistant rubber with high modulus of elasticity. Anatomically designed shape should ensure no Chondromalacia patella on prolonged use. Better uniform compression & grip, simple pull-on application, easy knee movement, snug fitting. Dimensions: Length - Not less than 33 cm (Unstretched) Net Weight- Not less than 100 g for a pair Product should be latex free and should not deteriorate on contact with oil, balm or on washing.	1's pair in sealed poly pack and in mono carton	1's X 10 in mono carton	250000
105	6051	Knee Cap (Medium)	Made of interlock weave bi-layered fabric (outer layer made of nylon and inner layer made of fine grade dermophillic & hypoallergenic cotton with inbetween air space), Four way stretchable, Heat resistant rubber with high modulus of elasticity. Anatomically designed shape should ensure no Chondromalacia patella on prolonged use. Better uniform compression & grip, simple pull-on application, easy knee movement, snug fitting. Dimensions: Length - Not less than 40 cm (Unstretched) Net Weight- Not less than 120 g for a pair. Product should be latex free and should not deteriorate on contact with oil, balm or on	1's pair in sealed poly pack and in mono carton	1's X 10 in mono carton	500000

			washing.			
106	6052	Knee Cap (Large)	Made of interlock weave bi-layered fabric (outer layer made of nylon and inner layer made of fine grade dermophillic & hypoallergenic cotton with inbetween air space), Four way stretchable, Heat resistant rubber with high modulus of elasticity. Anatomically designed shape should ensure no Chondromalacia patella on prolonged use. Better uniform compression & grip, simple pull-on application, easy knee movement, snug fitting.Dimensions: Length - Not less than 45 cm (Un-stretched)Net Weight- Not less than 140 g for a pair.Product should be latex free and should not deteriorate on contact with oil, balm or on washing.	1's pair in sealed poly pack and in mono carton	1's X 10 in mono carton	250000
107	7001	Hot Water Bag (Small)	Dimensions: Not less than- 25 cm x 19 cm Capacity: Not less than 1.5 Litres Wall thickness: Not less than 1.2 mm Made with high-quality, odourless vulcanized rubber of tensile strength not less than 14 MN/m2 (approx 140 kgf/cm2) and elongation at break of not less than 500 percent. The product should have an in-built hanger at one side, tie in cap and uniform rib pattern on both sides at outer side. The product should comply with IS 1867	1's in sealed poly pack and in mono carton	1's X 10	800000
108	8080	Surgical Face Mask, Disposable	Manufactured of 3 ply construction from non-woven poly prop fabric / SMS / Meltblown. Bacterial filtration efficiency (BFE) should not be less than 99%, should be heat sealed to keep 3 layers together, Standard Size- 17.5 cm x 9 cm, Color- Green / Blue Elastic ear loop of length 7 inch or more on one side and properly attached to avoid detachment while wearing. Should conform to the standards of IS 16289	1's	1's X 50 in sealed poly pack	
109	8082	Umbilical Catheter (For New Born)	UMBILICAL CATHETER (FOR NEWBORN) With female flexible mount. Open tip should be soft Non-toxic, medical grade PVC, Smooth round tip, atraumatic insertion, well-finished surface facilitate, smooth passage in the vein. Sterile, CE certified	1's in sealed blister pack	1's X 12 in mono carton	200000
110	8088	Walking Stick Quadripod	Durable, lightweight, rustproof, single telescopic height adjustable shaft made of high-grade aluminum alloy (corrosion resistant) with a molded handgrip and branching into four tips. Confirming standards to IS 5145 Handle: Made of injection-molded polypropylene of grade 2340PC with rubber handgrips (e.g., injection-molded polypropylene of grade 2340PC). Offset shape design to position your weight directly over the shaft, increasing stability and putting less pressure on the wrists. Net length- Not less than 150 mm	1's in sealed poly pack	1's X 1 in mono carton	100000

			Shaft: Made of extruded anodized aluminum (recyclable). The shaft should be in two parts, telescopic in nature. Height adjustable mechanism to be made out of stainless steel for high strength, low deformation and resistance to abrasion. The pin should be made of stainless steel, at least 6mm in diameter and electroplated to prevent corrosion. Clearance between sliding parts not to exceed 0.5mm.Net length- Not less than 600 mm extendable up to 1000 mm. Tips: Non-slip and replaceable, made of PU/PVC. Anti-rattle system to reduce noise made when walking. Net weight: 700-800 g Extra porous, dermophillic three panel Cotton/nylon/elastic webbing fabric.	1's in sealed poly pack		
111	8089	Abdominal Belt Velcro, Cream color (Large)	Broad hook and loop tape panel NLT 15 cm 02 Nylon reeves each not less than 1.5 cm width to prevent rolling over of belt. Length: Not less than 38 inch Width: Not less than 25 cm Net Weight: Not less than 260 g All sewing shall be done with lock stitches by turning in the free ends to depth. The abdominal belt shall be tailored neatly.	and in mono carton	1's X 10 in mono carton	250000
112	8090	Abdominal Belt Velcro, Cream color (Small)	Extra porous, dermophillic three panel Cotton/nylon/elastic webbing fabric.Broad hook and loop tape panel NLT 15 cm02 Nylon reeves each not less than 1.5 cm width to prevent rolling over of belt.Length: Not less than 30 inch Width: Not less than 25 cmNet Weight: Not less than 220 gAll sewing shall be done with lock stitches by turning in the free ends to depth. The abdominal belt shall be tailored neatly.	1's in sealed poly pack and in mono carton	1's X 10 in mono carton	200000
113	8091	Abdominal Belt Velcro, Cream color (Medium)	Extra porous, dermophillic three panel Cotton/ nylon/ elastic webbing fabric. Broad hook and loop tape panel NLT 15 cm	1's in sealed poly pack and in mono carton	1's X 10 in mono carton	250000
114	8092	Abdominal Belt Velcro, Cream color (XL)	Extra porous, dermophillic three panel Cotton/nylon/elastic webbing fabric. Broad hook and loop tape panel NLT 15 cm 02 Nylon reeves each not less than 1.5 cm width to prevent rolling over of belt. Length: Not less than 42 inch Width: Not less than 25 cm Net Weight: Not less than 280 g All sewing shall be done with lock stitches by turning in the free ends to depth. The abdominal belt shall be tailored neatly.	1's in sealed poly pack and in mono carton	1's X 10 in mono carton	200000

115	8093	Abdominal Belt Velcro, Cream color (XXL)	Extra porous, dermophillic three panel Cotton/nylon/ elastic webbing fabric. Broad hook and loop tape panel NLT 15 cm 02 Nylon reeves each not less than 1.5 cm width to prevent rolling over of belt. Length: Not less than 46 inch Width: Not less than 25 cm Net Weight: Not less than 300 g All sewing shall be done with lock stitches by turning in the free ends to depth. The abdominal belt shall be tailored neatly.		1's X 10 in mono carton	200000
116	8099	Cervical Collar Soft (Small)	Made of high-density polyurethane foam not less than 60kg/m3, reinforced with very thick LDPE sheet not less than 1.5mm with rounded edges and shaped to give a uniform mandible support. Nylon sewing thread 210/6. Eyelets: 3 in Number. Made up of virgin polypropylene diameter not less than 20 mm. Stockinette: Dermophillic and hypoallergenic blend of cotton and rayon which should be free from spinning, weaving and processing defects. Fasteners, Hook and Loop Tape: The tape shall have the minimum width of 5 cm and minimum overlap 2.5 cm. Expandity of the collar should be 5 cm. Total Length: not less than 460 mm (excluding fasteners) Length between centre of extreme eyelets: not less than 150 mm Width between crest: Not less than 100 mm Confirming to the standards of IS 11569	1's in sealed poly pack and in mono carton	1's X 1 in mono carton	250000
117	8100	Cervical Collar Soft (Medium)	Made of high-density polyurethane foam not less than 60kg/m3, reinforced with very thick LDPE sheet not less than 1.5mm with rounded edges and shaped to give a uniform mandible support. Nylon sewing thread 210/6.Eyelets: 3 in Number. Made up of virgin polypropylene diameter not less than 20 mm.Stockinette: Dermophillic and hypoallergenic blend of cotton and rayon which should be free from spinning, weaving and processing defects.Fasteners, Hook and Loop Tape: The tape shall have the minimum width of 5 cm and minimum overlap 2.5 cm. Expandity of the collar should be 5 cm.Total Length: not less than 500 mm (excluding fasteners)Length between centre of extreme eyelets: not less than 150 mmWidth between crest: Not less than 110	1's in sealed poly pack and in mono carton	1's X 1 in mono carton	250000
118	8101	Cervical Collar Soft (Large)	mmConfirming to the standards of IS 11569 Made of high-density polyurethane foam not less than 60kg/m3, reinforced with very thick LDPE sheet not less than 1.5mm with rounded edges and shaped to give a uniform mandible support. Nylon sewing thread 210/6. Eyelets: 3 in Number. Made up of virgin polypropylene diameter not less than 20 mm. Stockinette: Dermophillic and hypoallergenic	1's in sealed poly pack and in mono carton	I'S X I In	250000

Section and rayon which should be free from spinning, wearving and processing defects. Fasteners, Hook and Loop Tape: The tape shall have the minimum width of 5 cm and minimum overlap 2.5 cm. Expandity of the collar should be 5 cm. Total Length: not less than 550 mm (excluding fasteners) Length between centre of extreme eyelets: not less than 150 mm Width between crest: Not less than 15150 mm Width between crest: Not less than 150 mm Width between crest: Not less than 120 mn. Stockinette: Dermophillic and hypoallergenic between class than 1.5 mm with rounded edges and shaped to give a uniform mandible support. Nylon sewing thread 210/6. Eyeles: 3 in Number. Made up of virgin polypropylene diameter not less than 20 mm. Stockinette: Dermophillic and hypoallergenic befects. Extracts: Hot and Loop Tape: The tape shall have the minimum width of 5 cm and minimum overlap 2.5 cm. Expandity of the collar should be free from spinning, weaving and processing defects. Extracters, Hook and Loop Tape: The tape shall have the minimum width of 5 cm and minimum overlap 2.5 cm. Expandity of the collar should be 5 cm. Total Length: not less than 600 mm (excluding fasteners). Length not less than 130 mm Confirming to the standards of IS 11569 Made from breathable 03 layered PUP fused fabric/Soft Neoprene. Pre-shaped and removable malleable metal splint for thumb. Distal brace edge situated under the palmar crease. Parce is contoured at the ulnar aspect of the hand. High coverage of dorsal aspect of the hand. Eyeles is the metacarpophalangeal joint (MCP) of the thumb in a way that does not inhibit the movements of the hands. Wrist Cracle-6", Length-7" Made from breathable 03 layered PUP fused fabric/Soft Neoprene. Pre-shaped and removable malleable metal splint for thu							
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Eyelets: 3 in Number. Made up of virgin polypropylene diameter not less than 20 mm. Stockinette: Dermophillic and hypoallergenic blend of cotton and rayon which should be free from spinning, weaving and processing defects. Fasteners, Hook and Loop Tape: The tape shall have the minimum width of 5 cm and minimum overlap 2.5 cm. Expandity of the collar should be 5 cm. Total Length: not less than 600 mm (excluding fasteners) Length between centre of extreme eyelets: not less than 150 mm Width between crest: Not less than 130 mm Confirming to the standards of IS 11569 Made from breathable 03 layered PUF fused fabric/ Soft Neoprene. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Brace is contoured at the ulnar aspect of the hand. Support the carpometacarpal joint (CMC) and immobilize the metacarpophalangeal joint (MCP) of the thumb in a way that does not inhibit the movements of the hands. Wrist Circle-6", Length-7" Made from breathable 03 layered PUF fused fabric/ Soft Neoprene. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Pre-shaped and removable malleable metal spilnt for thumb. Distal brace edge situated under the palmar crease. Pre-shaped and removable metal spilnt for thumb. Distal brace edge situated under the palmar crease. P				shaped to give a uniform mandible support. Nylon	carton		
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			immobilize the metacarpophalangeal joint (MCP) of the thumb in a way that does not inhibit the movements of the hands. Wrist Circle-6"-7.5", Length-7"			
122	8105	Thumb and Wrist brace (Large)	Made from breathable 03 layered PUF fused fabric/ Soft Neoprene. • Pre-shaped and removable malleable metalsplint for thumb. • Distal brace edge situated under the palmar crease. • Brace is contoured at the ulnar aspect of the hand. • High coverage of dorsal aspect of the hand. • Support the carpometacarpal joint (CMC) and immobilize the metacarpophalangeal joint (MCP) of the thumb in a way that does not inhibit the movements of the hands. Wrist Circle-7.5"-8.5", Length-8"	1's in sealed poly pack and in mono carton	1's X 1 in mono carton	250000
123	8106	Thumb and Wrist brace (Extra Large)	Made from breathable 03 layered PUF fused fabric/ Soft Neoprene. • Pre-shaped and removable malleable metal splint for thumb. • Distal brace edge situated under the palmar crease. • Brace is contoured at the ulnar aspect of the hand. • High coverage of dorsal aspect of the hand. • Support the carpometacarpal joint (CMC) and immobilize the metacarpophalangeal joint (MCP) of the thumb in a way that does not inhibit the movements of the hands. Wrist Circle-8.5" and above, Length-8"	1's in sealed poly pack and in mono carton	1's X 1 in mono carton	200000
124	8113	Hernia Belt (Small)	Made up of soft and dermophillic fabric with adjustable pelvic and leg straps. Molded & removable anatomic ethafoam pads to apply gradual pressure around the affected area with focused pressure on the hernia, which pushes the inguinal hernia back. Colour: Grey	1's in sealed poly pack and in mono carton	1's X 1 in mono carton	100000
125	8114	Knee Brace Belt (Medium)	Knee Brace Belt - Excellent Support and Targeted Pain Relief • May Relieve the Pain Associated with Mild Osteoarthritis (OA) of the Knee • Unilateral Hinge and Strapping Configuration Apply Corrective Force to Offload the Affected Side of the Joint • WrapAround, Patient-Friendly Design - Easy to Apply to Swollen Tender Knees • Neoprene Provides Therapeutic Warmth to Soothe the Aching Joint • Support Straps Allow for Adjustable Compression • Available in Medial and Lateral Designs for Left and Right Knee Applications • Interchangeable Condyle Pads for Day-to-Day Adjustment Size- Medium 16-17 fits		1's X 1 in mono carton	250000
126	8115	Knee Brace Belt (Large)	Knee Brace Belt - Excellent Support and Targeted Pain Relief • May Relieve the Pain Associated with Mild Osteoarthritis (OA) of the Knee • Unilateral Hinge and Strapping Configuration Apply Corrective Force to Offload the Affected Side of the Joint •	carton	1's X 1 in mono carton	250000

			WrapAround, Patient-Friendly Design - Easy to Apply to Swollen Tender Knees • Neoprene Provides Therapeutic Warmth to Soothe the Aching Joint, Support Straps Allow for Adjustable Compression • Available in Medial and Lateral Designs for Left and Right Knee Applications • Interchangeable Condyle Pads for Day-to-Day Adjustment Size- Large 18-19 fits			
127	8116	IV Cannula Fixator (Medium 6cm x 5cm)	IV Cannula Fixer Good aesthetic appeal due to woven fast edges, Moisture responsive High Moisture Vapor Transmission Rate Film, Low allergy grid pattern adhesive, porous adhesive to allow skin breathing, Easy and Painless remove, because of thin-non adhesive edges and leaves no residue after remove. Medium: 6cm x 5cm	1's pack in tearable wrapper	1's X 50 in mono carton	500000
128	8117	Cervical Pillow (Memory foam)	Ergonomically Double Contoured Cervical Pillow confirming to the shape of neck to accommodate both long and short necks. Made from viscoelastic high density, open cell, flexible, polyurethane foam (memory foam). Density: 65-70 Kg/m3, Sag factor: >2.5Ball rebound resilience: < 5% Time to return 95% height: 14 sLength: 38 cm, Width: 47 cm, Crest Height1:12 cm, Crest Height2: 9.5 cm,Inner cover to protect the memory foam. Grey color, Washable, hypoallergenic, durable, soft and smooth Zipper cover.	1's in sealed poly pack and in mono carton	1's X 1 in mono carton	200000
129	8120	Lancets, Round Sterile, 28G	Lancets Round Sterile Tip 28G One time use; 28G, in virgin packing Length: 30mm ± 2mm Compatibility: Should be compatible with Jan Aushadhi glucometer.	50 pcs/Box		500000
130	8123	Adhesive Surgical Paper Tape Size 5cmx5m	Breathable, Hypo-allergenic, latex-free, non-woven and micro-porous backing material. High-quality water-soluble adhesive for gentle bonding, holds well on damp skin and clean removal with minimal residue. Size: Size 5 cm x 5 m	1's	1's X 12 in mono carton	200000
131	8124	Adhesive Surgical Paper Tape Size 2.5cmx5m	Breathable, Hypo-allergenic, latex-free, non-woven and micro-porous backing material. High-quality water-soluble adhesive for gentle bonding, holds well on damp skin and clean removal with minimal residue. Size: 2.5 cm x 5 m	1's	1's X 12 in mono carton	600000
132	8125	Adhesive Surgical Paper Tape Size 1.25cmx5m	Breathable, Hypo-allergenic, latex-free, non-woven and micro-porous backing material. High-quality water-soluble adhesive for gentle bonding, holds well on damp skin and clean removal with minimal residue. Size: 1.25 cm x 5 m	1's	1's X 12 in mono carton	400000
133	8127	Nebulizer Mask with Tubing (Adult)	Made of clear, latex free, non-toxic PVC, medium concentration, adjustable nose clip, non kinkable & non-autoclave Tube Length: 2 m 10cc bowl with graduated volume markings 45-degree elbow, Adult elongated mask Medication port to eliminate medication waste ue	1's pack in sealed poly pack	1's X 25 in mono carton	500000

			Nebulization rate: 3cc / 10 mins. CE certified Packing: Single Piece poly packed.			
134	8131	Plastic Urine Pot collector (For both Male & Female)	Wide mouth, Snap-on leak proof tie-on and glow in dark lid, notched handle made from durable polypropylene Plastic Urine Pot collector, Compatible for Male & Female use. Weight: Not less than 150 g, Graduated volume marking up to 1000 ml. Color: White	1's in sealed LDPE pack	1's X 10 in sealed poly pack	250000
135	8132	Plastic Bedpan	Made of high virgin ABS material Wide plastic guard to prevent spills and a tappered end for easier placement. Portable lid and built-in handle with grip. Nonstick ceramic finish inner layer and smooth bottom. Size: 40 cm X 30 cm 9 cm Weight: Not less than 750 g Color: Beige	1's in sealed LDPE pack	1's X 1 in sealed poly pack	200000
136	8135	Breathing Exerciser (3 Chambers)	Breathing Exerciser, Total 3 Chambers, Device is composed of base and central part divided into three chambers containing three small spheres of different size and color, connecting tube with 12mm OD connector and mouth piece, Flow rates 600ml/sec, 900ml/sec, and 1200ml/sec by using different colors of ball for easy identification of the flow rates. Material Specifications: - ABS for chamber, PE for ball and mouthpiece and EVA for tubing.	1's in sealed poly pack and in mono carton	1's X 1 in	500000
137	8150	3-way stopcock with 10 cm extension line	3-way stopcock with 10 cm extension lineSet Length 15 cm Priming Volume 1.06 ml, Bore 3 mm 1 Blue Stopcock Handle(s), 1 Female Luer(s), 1 Male Luer(s), made of Polyvinyl Chloride (PVC)Lipid Resistant, Natural rubber latex is not part of the material formulation, DEHP Free. CE certified	1's in sealed poly pack and in mono carton	1's X 1 in mono carton	100000
138	8151	Adhesive wound dressings, 25cm x 10cm	Adhesive wound dressings, 25cm x 10cm Consisting of breathable non-woven top layer and a low-adherent absorbent pad Size: 25cm x 10cm, Each dressing is individually wrapped and sterile. Sterilisation is by ethylene oxide. CE certified	1's in sealed peelable wrapper	1's X 50 in mono carton	250000
139	8153	Arm Sling Pouch (Large), Grey	Made from breathable 3 layer bonded fabric-Outer 100% polyster knit wrap fabric with raising, Middle PUF layer for cushioning and wrinle free, Inner 100% polyester soft liner wrap knit. Adjustable and non-stretchable shoulder sling of width not less than 4 cm with cushioned ethafoam shoulder pad not less than 10 cm and length 20 cm. Should have Thumb cradle Designed: For Both Elbow, Color: Grey Ergonomic, smart, sleek and open design provides improved ventilation and enhanced comfort. One touch opening of PP slide buckle. Length: Not less than 42 cm	1's in sealed poly pack and in mono carton	1's X 1 in mono carton	150000
140	8154	Arm Sling Pouch (Medium),	Made from breathable 3 layer bonded fabric- Outer 100% polyster knit wrap fabric with raising, Middle PUF layer for cushioning and wrinle free,	1's in sealed poly pack and in mono	mono	150000

		Grey	Inner 100% polyester soft liner wrap knit. Adjustable and non-stretchable shoulder sling of width not less than 4 cm with cushioned ethafoam shoulder pad not less than 10 cm and length 20 cm. Should have Thumb cradle Designed: For Both Elbow, Color: Grey Ergonomic, smart, sleek and open design provides improved ventilation and enhanced comfort. One touch opening of PP slide buckle. Length: Not less than 38 cm	carton		
141 8	8155	Baby Wipes (Pack of 20 wipes)	Made from extra soft and ultra fine spunlace fabric. Alcohol free, paraben free, SLS free, hypoallergic to infant skin. Infused with 2% Chlorhexidine, Aloe vera and Glycerin, Mild fragrance. Size: 240 mm X 290 mm per wipe pH: 4.5 -6, Mass: > 36 g/m2, Moisture content: > 90 % by 98 % demineralized water. Breaking Strength: Machine direction (wet): > 30 N/min	Pack of 20 wipes packed in sealed LDPE pack with resealable plastic flap	01's X 20 in mono carton	500000
142 8	8156	Bed Bath Towel (Pack of 10 wipes)	Cross direction (wet): > 2.5 N/min Made from extra soft and ultra fine spunlace fabric. Alcohol free, paraben free, SLS free, hypoallergic to infant skin. Infused with Aloe vera and Vitamin E (Veg origin), Mild fragrance, Microwaveable. Size: 200 mm X 150 mm per wipe pH: 4.5 -6, Mass: > 36 g/m2, Moisture content: > 90 % by 98 % demineralized water. Breaking Strength: Machine direction (wet): > 30 N/min Cross direction (wet): > 2.5 N/min	Pack of 10 wipes packed in sealed LDPE pack with resealable plastic flap	01's X 10 in mono carton	250000
143 8	8158	Chlorhexidin e Gauze Dressing B.P 10cmx10cm (Sterile)	CHLORHEXIDINE GAUZE DRESSING B.P. 10cm * 10cm (Sterile) Material- Leno Weave Cotton, Coated with Chlorhexidine 0.5% w/w with white Soft Paraffin, Color White.	1's in sealed peelable wrapper	1's X 10 in mono carton	250000
144 8	8160	Disposable Plastic Hand Gloves (Free size)	Disposable Plastic Hand Gloves (Free size)Disposable gloves made with transparent plastic material, polythene gloves.	Pair		500000
145 8	8161	Elastic gauze bandages 10cm	Elastic gauze bandages 10cm High quality elastic fabric, soft edges and porous adhesive mass, Water repellent, Air permeable, Thinner substrate, non-fray edges, Confirming standards of IS 16111	1's in sealed peelable wrapper	1's X 10 in mono carton	200000
146 8	8162	Elastic gauze bandages 7.5cm	Elastic gauze bandages 7.5cm High quality elastic fabric, soft edges and porous adhesive mass, Water repellent, Air permeable, thinner substrate, non-fray edges, Confirming standards of IS 16111	1's in sealed peelable wrapper	1's X 10 in mono carton	250000
147 8	8167	Karman cannula 4	Flexible plastic cannula and with built-in adaptor Specially designed for aseptic medical termination	1's in sealed blister pack		100000

		mm	of pregnancy, coned shaped distal end with two		carton	
			large lateral eyes, Proximal end fitted with MTP syringe or suction apparatus, individually packed			
			in paper pouch & sterile.			
			Confirming standards of IS 8313			
			Lumbar Spinal Brace (Grey)	1's		
			covers all dorsal, lumbar and sacral vertebras. It			
			supports and immobilizes the spine in a neutral			
			position which allows vital body movement. Dimensions: 55 x 42 x 5 cm			
		Lumbar	Features			
148	8168		Easy to use and comfortable Adjustable shoulder			50000
1 10	0100	(Grey)	straps Immobilizes the spine Elastic abdominal			50000
		(313)	panels offer the required compression.			
			It is manufactured with rigid, anatomic			
			customizable, splints which ensure ideal fitting,			
			enhanced immobilization and accurate spinal			
			posture, firm grip and self-tightening of the brace.			
		Magnetic	Made from high quality 3-ply fabric	1's		
		Posture	front-loading plastic buckles, 12 premium			
149	8169	Corrector	magnets, Velcro straps, firm grip, high-quality			50000
		Back Support	breathable, porous neoprene, reinforced cross,			
		Belt - Posture Fit	double stitching.			
		110	Made from non-toxic medical grade PVC,	1's in sterile		
		~	transparent with leakproof lid, Sterile, free from	sealed poly	1's X 50	
150	8177	Specican 30	foreign particle.	pack	in sealed	100000
		ml	Sticker pasted for writing patient name. Weight:		mono	
			Not less than 25 g.		carton	
			Made from spun bond PP non-woven fabric and	1's		
			spun lace fabrics. The caps should be			
			ultrasonically sealed and lighter gravity than		1's X 50	
151	0170	C:1 C	hydrophobic PP fibers for excellent water		in sealed	250000
151	81/8	Surgical Cap	repellency and air permeability resulting in		LDPE	250000
			providing comfort & dryness resulting in anti- bacterial and anti-fungal resistance		zipper pouch	
			Color: Blue, Size: 21 inches with 02 LP Elastic		pouch	
			Thickness: Not less than 15 gsm			
			Made from fabric that hot roll combining two	1's	11 37 50	
			layers of spun bond non-woven on outer side and		1's X 50	
		Surgical shoe	three layer of melt blown non-woven fabric in the		pairs in sealed	
152	8179	cap	middle. 02 Top LP elastic closure.		LDPE	200000
		Сар	Thickness: Not less than 50 GSM		zipper	
			PP + PE laminated (Waterproof)		pouch	
			Height: 20 cm, Length: 410 mm	41 .	F	
			Each peelable sterile wrapper contains one thin	1's in		
			film backing with a non-latex, hypoallergenic adhesive. The Film with border should be notched	peelable Aluminium		
		Transparent	and reinforced with soft cloth tape to provide a	foil pack		
		film	better seal around catheters and other devices.	ion pack	1's X 10	
153	8181	dressings	The dressing should be waterproof, breathable,		in mono	200000
100	5101	(Sterile)	allowing good oxygen and moisture vapor		carton	
		10cm X	exchange.			
		12cm	The dressing should be impermeable to liquids,			
			bacteria, and viruses of size > 30 nm.			

			CE certified			
154	8182	Vein Compression Stocking Knee length (Extra Large)	Each monocarton should contain:01 glider01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking knee length (open toe type). Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >30 cm Calf >40 cm Popliteal >40 cm Each measuring position should retain not less than 85% compression after 30 wash cycles. Tests	1's pair and glider in sealed poly pack and in mono carton	1's X 10 in mono carton	200000
155	8183	Vein Compression Stocking Knee length (Large)	as per conforming to IS 16467. Each monocarton should contain: 01 glider 01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking knee length (open toe type). Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg)	1's pair and glider in sealed poly pack and in mono carton	1's X 10 in mono carton	150000
			Ankle >27 cm Calf >38 cm Popliteal >38 cm Each measuring position should retain not less than 85% compression after 30 wash cycles. Tests as per conforming to IS 16467.			
156	8184	Vein Compression Stocking Knee length (Medium)	Each monocarton should contain: 01 glider 01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking knee length (open toe type). Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17	1's pair and glider in sealed poly pack and in mono carton	1's X 10 in mono carton	200000

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				dtex f 7.			
				Interrupted band and 2-ply gusset to prevent			
				tourniquet effect at the femoral vein.			
				Inlay circumferential knit, helping to provide one			
				way horizontal stretch.			
				A pressure break at the popliteal vein helps to			
				ensure that blood will continue to flow smoothly			
				through this critical area.			
				A reinforced and defined heel pocket to aid correct			
				placement. Cuff not less than 5cm to hold stocking			
				at place.			
				Circumference Compression (mm of Hg)			
				Ankle >23 cm			
				Calf >35 cm			
				Popliteal > 35 cm			
				Each measuring position should retain not less			
				than 85% compression after 30 wash cycles.			
				Tests as per conforming to IS 16467.			
				Each monocarton should contain:01 glider01 pair	1's pair and		
				of latex free, ultra-thin, dermophillic & breathable	glider in		
				vein compression stocking knee length (open toe	sealed poly		
				type). Made up of plaited knitted PA microfilament	pack and in		
				yarns 60 dtex f 60 and elastane yarn in the count	mono carton		
				22/17 dtex f 7.Interrupted band and 2-ply gusset to			
				prevent tourniquet effect at the femoral vein.Inlay			
				circumferential knit, helping to provide one way			
			1 7.	horizontal stretch. A pressure break at the popliteal			
			Vein	vein helps to ensure that blood will continue to		11 37 10	
	1.57	0105	Compression	flow smoothly through this critical area.A		1's X 10	100000
	15/	8185	Stocking	reinforced and defined heel pocket to aid correct		in mono	100000
			Knee length	placement. Cuff not less than 5cm to hold stocking		carton	
			(Small)	at place.			
				Circumference Compression (mm of Hg)			
				Ankle >19 cm			
				Calf >32 cm			
				Popliteal > 32 cm			
				Each measuring position should retain not less			
				than 85% compression after 30 wash cycles. Tests			
				as per conforming to IS 16467.			
				Each monocarton should contain: 01 glider.	1's pair and		
					glider in		
				01 pair of latex free, ultra-thin, dermophillic &	sealed poly		
				breathable vein compression stocking knee length	pack and in		
				(open toe type).	mono carton		
			Vein	Made up of plaited knitted PA microfilament yarns		:-	
			Compression	60 dtex f 60 and elastane yarn in the count 22/17		1's X 10	
	158	8186	Stocking	dtex f 7.		in mono	100000
			Knee length	Interrupted band and 2-ply gusset to prevent		carton	
			(XXL)	tourniquet effect at the femoral vein.			
				Inlay circumferential knit, helping to provide one			
				way horizontal stretch.			
				A pressure break at the popliteal vein helps to			
				ensure that blood will continue to flow smoothly			
				through this critical area.			
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			A reinforced and defined heel pocket to aid correct			
			placement. Cuff not less than 5cm to hold stocking			
			at place.			
			Circumference Compression (mm of Hg)			
			Ankle >33 cm Calf >45 cm			
			Popliteal >45 cm			
			Each measuring position should retain not less			
			than 85% compression after 30 wash cycles.			
			Tests as per conforming to IS 16467.			
			Each monocarton should contain:	1's pair and		
			01 glider	glider in		
			01 pair of latex free, ultra-thin, dermophillic &	sealed poly		
			breathable vein compression stocking Thigh length	pack and in		
			(open toe type).	mono carton		
			Made up of plaited knitted PA microfilament yarns			
			60 dtex f 60 and elastane yarn in the count 22/17			
			dtex f 7.			
			Interrupted band and 2-ply gusset to prevent			
			tourniquet effect at the femoral vein.			
			Inlay circumferential knit, helping to provide one			
		Vein	way horizontal stretch.			
		Compression	A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly		1's X 10	
159	8187	Stocking	through this critical area.		in mono	100000
		Thigh length	A reinforced and defined heel pocket to aid correct		carton	
		(Extra Large)	placement. Silicone dotted cuff not less than 5cm			
			to hold stocking at place.			
			Circumference Compression (mm of Hg)			
			Ankle >33 cm			
			Calf >45 cm			
			Popliteal > 45 cm			
			Lower Thigh >60 cm Upper Thigh >70 cm			
			Upper Thigh >70 cm Each measuring position should retain not less			
			than 85% compression after 30 wash cycles.			
			Tests as per conforming to IS 16467.			
	†		Each monocarton should contain:01 glider01 pair	1's pair and		
			of latex free, ultra-thin, dermophillic & breathable	glider in		
			vein compression stocking Thigh length (open toe	sealed poly		
			type).Made up of plaited knitted PA microfilament	pack and in		
			yarns 60 dtex f 60 and elastane yarn in the count	mono carton		
			22/17 dtex f 7.Interrupted band and 2-ply gusset to			
		Vein	prevent tourniquet effect at the femoral vein.Inlay		11 37 40	
1.00	0100	Compression	circumferential knit, helping to provide one way		1's X 10	100000
100	8188		horizontal stretch. A pressure break at the popliteal		in mono	100000
		Thigh length (Large)	vein helps to ensure that blood will continue to flow smoothly through this critical area.A		carton	
		(Large)	reinforced and defined heel pocket to aid correct			
			placement. Silcone dotted cuff not less than 5cm to			
			hold stocking at place.			
			Circumference Compression (mm of Hg)			
			Ankle >27 cm			

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			Calf >39 cm Popliteal >39 cm			
			Lower Thigh >55 cm			
			Upper Thigh >63 cm			
			Each measuring position should retain not less			
			than 85% compression after 30 wash cycles. Tests			
			as per conforming to IS 16467.			
			Each monocarton should contain:	1's pair and		
			01 glider	glider in		
			01 pair of latex free, ultra-thin, dermophillic &	sealed poly		
			breathable vein compression stocking Thigh length			
			(open toe type).	mono carton		
			Made up of plaited knitted PA microfilament yarns			
			60 dtex f 60 and elastane yarn in the count 22/17			
			dtex f 7.			
			Interrupted band and 2-ply gusset to prevent			
			tourniquet effect at the femoral vein.			
			Inlay circumferential knit, helping to provide one			
		Main	way horizontal stretch.			
		Vein	A pressure break at the popliteal vein helps to		1's X 10	
16	1 8189	Compression Stocking	ensure that blood will continue to flow smoothly		in mono	100000
10	1 0109	Thigh length	through this critical area.		carton	100000
		(Medium)	A reinforced and defined heel pocket to aid correct		Carton	
		(Wicdium)	placement. Silcone dotted cuff not less than 5cm to			
			hold stocking at place.			
			Circumference Compression (mm of Hg)			
			Ankle >23 cm			
			Calf >36 cm			
			Popliteal > 36 cm			
			Lower Thigh >45 cm Upper Thigh >57 cm			
			Each measuring position should retain not less			
			than 85% compression after 30 wash cycles.			
			Tests as per conforming to IS 16467.			
			Each monocarton should contain:01 glider 01 pair	1's pair and		
			of latex free, ultra-thin, dermophillic & breathable	glider in		
			vein compression stocking Thigh length (open toe	sealed poly		
			type).Made up of plaited knitted PA microfilament	pack and in		
			yarns 60 dtex f 60 and elastane yarn in the count	mono carton		
			22/17 dtex f 7.Interrupted band and 2-ply gusset to			
			prevent tourniquet effect at the femoral vein.Inlay			
			circumferential knit, helping to provide one way			
		Vein	horizontal stretch. A pressure break at the popliteal			
		Compression	vein helps to ensure that blood will continue to		1's X 10	
16	2 8190	Stocking	flow smoothly through this critical area.A		in mono	100000
		Thigh length	reinforced and defined heel pocket to aid correct		carton	
		(Small)	placement. Silcone dotted cuff not less than 5cm to			
			hold stocking at place.			
			Circumforum on Communication (CIII)			
			Circumference Compression (mm of Hg)			
			Ankle >19 cm Calf >33 cm			
			Popliteal > 33 cm Lower Thigh >40 cm			
			Upper Thigh >50 cm			
I └─		<u> </u>	Opportingn /Ju cm	<u> </u>		1

			Each measuring position should retain not less			
			than 85% compression after 30 wash cycles. Tests			
			as per conforming to IS 16467.			
			Each monocarton should contain:	1's pair and		
			01 glider	glider in		
			01 pair of latex free, ultra-thin, dermophillic &	sealed poly		
			breathable vein compression stocking Thigh length	pack and in		
			(open toe type).	mono carton		
			Made up of plaited knitted PA microfilament yarns			
			60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7.			
			Interrupted band and 2-ply gusset to prevent			
			tourniquet effect at the femoral vein.			
			Inlay circumferential knit, helping to provide one			
		Vein	way horizontal stretch.			
		Compression	A pressure break at the popliteal vein helps to		1's X 10	
163	8191	Stocking	ensure that blood will continue to flow smoothly		in mono	100000
		Thigh length	through this critical area.		carton	
		(XXL)	A reinforced and defined heel pocket to aid correct placement. Silicone dotted cuff not less than 5cm			
			to hold stocking at place.			
			to note stocking at place.			
			Circumference Compression (mm of Hg)			
			Ankle >33 cm			
			Calf >45 cm			
			Popliteal > 45 cm Lower Thigh >63 cm			
			Lower Thigh >63 cm Upper Thigh >75 cm			
			Each measuring position should retain not less			
			than 85% compression after 30 wash cycles.			
			Tests as per conforming to IS 16467.			
			Net weight 100 g arranged in zig zag pattern,	20's in		
		7:	square shaped.	sealed	20's X 10	
164	8192	Zig zag cotton	Made from 100% clean and soft cotton Natural fibers of average staple length	LDPE pack	in sealed Mono	250000
		Cotton	not less than 10 mm, soft and fluffy, high		pack	
			absorption		puck	
			Made of interlock weave bi-layered fabric (outer	1's pair in		
			layer made of nylon and inner layer made of fine	sealed poly		
			grade dermophillic & hypoallergenic cotton with	pack and in		
			in between air space), Four way stretchable, heat	mono carton		
			resistant rubber with high modulus of elasticity, Anatomically designed.			
			Silicone patellar ring with padding with anti-slip			
		II. C	coating.		11 37 10	
165	0020	Knee Cap	Multiaxial side splint to prevent roll on and easy		1's X 10	250000
103	9039	with patellar ring (Large)	flexion.		in mono carton	250000
		ing (Laige)	Better compression & grip, simple pull-on		Carton	
			application, easy knee movement.			
			Dimensions: Length - Not less than 40 cm (Unstretched)			
			Net Weight- Not less than 190 g for a pair.			
			Product should be latex free and should not			
1			deteriorate on contact with oil, balm or on			1
			deteriorate on contact with on, bann or on			

166	9040	Knee Cap with patellar ring (Medium)	Made of interlock weave bi-layered fabric (outer layer made of nylon and inner layer made of fine grade dermophillic & hypoallergenic cotton with inbetween air space), Four way stretchable, Heat resistant rubber with high modulus of elasticity, Anatomically designed. Silicone patellar ring with padding with anti-slip coating. Multiaxial side splint to prevent roll on and easy flexion. Better compression & grip, simple pull-on application, easy knee movement. Dimensions: Length - Not less than 40 cm (Un-stretched)Net Weight- Not less than 180 g for a pair. Product should be latex free and should not deteriorate on contact with oil, balm or on washing.	1's pair in sealed poly pack and in mono carton	1's X 10	250000
167	9041	Knee Cap with patellar ring (Small)	Made of interlock weave bi-layered fabric (outer layer made of nylon and inner layer made of fine grade dermophillic & hypoallergenic cotton with inbetween air space), Four way stretchable, Heat resistant rubber with high modulus of elasticity, Anatomically designed. Silicone patellar ring with padding with anti-slip coating. Multiaxial side splint to prevent roll on and easy flexion. Better compression & grip, simple pull-on application, easy knee movement. Dimensions: Length - Not less than 35 cm (Unstretched) Net Weight- Not less than 170 g for a pair. Product should be latex free and should not deteriorate on contact with oil, balm or on washing.	1's pair in sealed poly pack and in mono carton	1's X 10	250000
168	9042	Nebulizer Mask with Tubing (Child)	Made of clear, latex free, non-toxic PVC, medium concentration, adjustable nose clip, non kinkable & non-autoclave Tube Length: 2 m 10cc bowl with graduated volume markings 45-degree elbow, Child elongated mask Medication port to eliminate medication waste ue Nebulization rate: 3cc / 10 mins. CE certified Packing: Single Piece poly packed.	1's pack in sealed poly pack	1's X 25 in mono carton	250000
169	9043	Walker (Sit Assist-to- stand support)	Durable, lightweight, rust proof, foldable, height adjustable shaft made of high grade aluminium alloy (corrosion resistant) with 2 pairs of injection-moulded polypropylene of grade 2340 PC with rubber handgrips. Confirming standards to IS 5145 Shaft: Made of extruded anodized aluminum (recyclable). Each vertical shaft should be in two parts, telescopic in nature. 02 'U' shaped Front horizotal shaft and one with pellet button for folding. Height adjustable mechanism to be made out of stainless steel for high strength, low deformation and resistance to abrasion. The pin should be made of stainless steel, at least 6mm in diameter and electroplated to prevent corrosion. Clearance between sliding parts not to exceed 0.5mm.	1's in sealed poly pack	1's X 1 in mono carton	100000

Г							
	170	9044	Elbow crutch (Adjustable)	Foldable Depth: Not more than 100 mm. Frontal Height: 650 mm extendable upto 900 mm Back Height: minimum 450 mm and extendable at par. Frontal width: Not less than 500 mm Lateral width: Not less than 300 mm Tips: Non-slip and replaceable, made of durable PU/PVC. Anti-rattle system to reduce noise made when walking. Net weight: 1900 g- 2000 g Durable, lightweight, rust proof, height adjustable shaft made of high grade aluminium alloy (corrosion resistant) with a molded handgrip. Confirming standards to IS 5145Handle: Made of injection-moulded polypropylene of grade 2340PC with rubber handgrips (e.g. injection-moulded polypropylene of grade 2340PC). Net length- Not less than 150 mmShaft: Made of extruded anodized aluminum (recyclable). The shaft should be double adjustable – from floor to handle & from handle to cuff in two parts. Height adjustable mechanism to be made out of stainless steel for high strength, low deformation and resistance to abrasion. The pin should be made of stainless steel, at least 6mm in diameter and electroplated to prevent corrosion. Clearance between sliding parts not to exceed 0.5mm. Elbow shaft with flipping elbow grip should be anatomically aligned with handle to bear body weight and decrease fatigue. Net length:- Main shaft- Not less than 600 mm and extendable upto	1's in sealed poly pack	1's X 1 in mono carton	100000
	171	9045	Insulting /Medicine Cooling Case	extendable upto 300 mmTips: Non-slip and replaceable, made of durable PU/PVC.Anti-rattle system to reduce noise made when walking.Net weight: 400-500 g Product dimension: 8" x 4.2" x 2.5" Material: 1. Outer Fabric – 1000 Denier Polyester Polyurethane (PU) coated, (0.5mm), Durable, finely woven and water-resistant fabric. 2. Inner Fabric - 4 x 4 Polyester Matty Soft PU coated, light weight, highly flexible and moisture resistant. 3. Polyester Zipper - Size 5 with Metal Slider. 4. PP (Polypropylene) Tape – covering panels and joints inside the Box. 5. Pippin Wire (Tar) – all around outer side, made of LDPE material. 6. White PP Sheet – 0.45mm, Polypropylene Sheet – High temperature & Corrosion Resistant. 7. Aluminum laminated Fabric – Reflect thermal radiation. 8. PVC Clear Film – 0.15mm – Used for moisture	1 Cooling Case, 2 unit Gel Cool Pack, 1 Unit Insulin Pen Folder and 1 Carry Bag	1's X 1 in mono carton	50000

control and to protect Aluminized Fabric.		
9. Woolen Felt Fabric – high graded for inner		
temperature protection. App. 2 mm thick.		
10. Polyester Net/Mesh - for Inside pockets with		
Polyester Elastic.		
11. XLPE one side Alu Foil laminated Foam 6mm		
(Cross-linked polyethylene) – Highly		
Temperature Resistant, best for thermal insulation.		
12. Pen Folder: Size 6.5" x 7", Folded Size 6.5" x		
3.5"		
A – 4 x 4 Matty Soft PU coated,		
B - Aluminum laminated Fabric – Reflect thermal		
radiation		
C - PVC Clear Film – 0.18mm		
D - Polyester Net/Mesh		
E – Embedded with PP Tape		
F – Metal Snap Button (VT5)		
13. Gel - Ice Pouch Size 3" x 6" – Base Material:		
Carbopol, Triethanolamine (TEA) with		
preservative and water in LDPE pouch.		
14. Gel-Ice Pouch Cover – Moisture resistant Poly		
Fabric		
15. Carry Bag with Strings – Water proof		
Laminated Taffeta Fabric		

Note:

i. Bidders shall consider the specification confirming all the quality parameters, Safety and Product Standards.

Annexure - XIII

{Ref:- clause 19(K)}

(1)	(2)	(3)	(4)	(5)	(6)	(7)
S.N.	Item Code	Generic Name of Item	Detail Specification of Item	Unit Size	Minimum Shelf Life (Must not be less than warranty period in specification or 3 years)	HSN Code of item
1	5001	Absorbent Cotton Wool IP 75 g	Absorbent Cotton Wool IP Net weight of 75 g Made from 100% clean and soft cotton Natural fibres of average staple length not less than 10 mm, soft and fluffy, high absorption	1's in Paper rolled and in Sealed Poly pack		
2	5002	Absorbent Cotton Wool IP 200 g	Absorbent Cotton Wool IP Net weight of 200 g Made from 100% clean and soft cotton Natural fibers of average staple length not less than 10 mm, soft and fluffy, high absorption	I's in Paper rolled and in Sealed Poly pack		
3	5003	Absorbent Cotton Wool IP 500 g	Absorbent Cotton Wool IP Net weight of 500 g Made from 100% clean and soft cotton Natural fibers of average staple length not less than 10 mm, soft and fluffy, high absorption	1's in Paper rolled and in Sealed Poly pack		
4	5004	Crepe Bandage B.P. 15 cm x 4 M	Crepe Bandage B.P. 15 cm x 4 M Uniform plain weave and of continuous length with no loose salvage threads (both sides fastened edges), Wrap threads- Two fold cotton threads, Not less than 170 threads per 10 cm. Weft threads- Combined cotton & Viscose, Not less than 78 threads per 10 cm. Elasticity- Not more than 80% of fully stretched length. washing regains elasticity-fit for re-use. Weight- Not Less Than 140 g / m2 03 Nos. Loop & Hook Closure	1's		
5	5005	Crepe Bandage B. P.10 cm x 4 M	Crepe Bandage B.P. 10 cm x 4 M Uniform plain weave and of continuous length with no loose salvage threads (both sides fastened edges), Wrap threads- Two fold cotton threads, Not less than 170 threads per 10 cm. Weft threads- Combined cotton & Viscose, Not less than 78 threads per 10 cm. Elasticity- Not more than 80% of fully stretched length. washing regains elasticity-fit for re-use. Weight- Not Less Than 140 g / m2 02 Nos. Loop & Hook Closure	1's		

	1	1		
6	5006	Cotton Bandage 7.5 cm x 4 M	Cotton Bandages Size 7.5 cm X 4M, Freedom from optical whiteners and Conforming to the standards as per IS 863-1988, Count of yarn: Warp -NLT 25 tex Weft- NLT 30 tex Threads: Ends (column)- NLT 150 Picks (Row)- NLT 85 Mass: NLT - 57 ± 5 g / m2 pH: 6.5 to 8.5	1's in Paper rolled and in Sealed Poly pack
7	5007	Cotton Bandage 10 cm x 4 M	Cotton Bandages Size 10 cm X 4M, Freedom from optical whiteners and Conforming to the standards as per IS 863-1988, Count of yarn: Warp -NLT 25 tex Weft- NLT 30 tex Threads: Ends (column)- NLT 150 Picks (Row)- NLT 85 Mass: NLT - 57 ± 5 g / m2 pH: 6.5 to 8.6	1's in Paper rolled and in Sealed Poly pack
8	5008	Cotton Bandages(Non Sterile) 15 cm x 4 M	Cotton Bandages Size 15 cm X 4M, Freedom from optical whiteners and Conforming to the standards as per IS 863-1988, Count of yarn: Warp -NLT 25 texWeft- NLT 30 texThreads: Ends (column)- NLT 150Picks (Row)- NLT 85 Mass: NLT - 57 ± 5 g / m2 pH: 6.5 to 8.7	1's in Paper rolled and in Sealed Poly pack
9	5009	Adhesive Bandages Wash proof	Sterile Adhesive Bandages Wash proof 19mm X 70 mm containing Benzalkonium Chloride Solution IP 0.5 % w/w and tinted with tartrazine yellow. The wound pad size NLT 25 mm and should be water resistant, provide antiseptic protection with high quality PVC as backing material. Skin friendly & Hypoallergic adhesive, no sticky residue.	I's in individual sealed tearable paper wrapper
10	5010	Syringe 2 ml with needle 24G	Disposable Syringe2 ml, with colour coded (as per BIS), needle 24G Sterilized, Luer Mount, Non - toxic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in individual Sealed tearable poly wrapper
11	5011	Syringe 5 ml with needle 24G	Disposable Syringe5 ml, with colour coded (as per BIS) needle 24G Sterilized, Luer Mount, Non -toxic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in individual Sealed tearable poly wrapper
12	5012	Syringe 10 ml with needle 21G	Disposable Syringe10 ml, with colour coded (as per BIS) needle 21G Sterilized, Luer Mount, Non - toxic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in individual Sealed tearable poly wrapper

13	5013	Syringe 20 ml with needle 23G	Disposable Syringe10 ml, with colour coded (as per BIS) needle 23G Sterilized, Luer Mount, Non - toxic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification. The needles fit the luer nozzle, Sterile, single	1's in individual Sealed tearable poly wrapper 1's in blister
14	5014	16G	use, non-toxic, non-pyrogenic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	pack
15	5015	Needle 18G	The needles fit the luer nozzle, Sterile, single use, non-toxic, non-pyrogenic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in blister pack
16	5016	Needle 23G	The needles fit the luer nozzle, Sterile, single use, non-toxic, non-pyrogenic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in blister pack
17	5017	Needle 26G half inch	The needles fit the luer nozzle, Sterile, single use, non-toxic, non-pyrogenic as per Drugs and Cosmetics Act 1940, IS No. 10258- 2002 with BIS/ CE/ USFDA certification.	1's in blister pack
18	5019	Paper Adhesive Plaster 1 inch X 9m	Size-1 inch X 9 m per roll Material: Non woven, non-allergic & Strong adhesive but gentle to skin & painless removal, Porous and breathable CE certified	1's
19	5020	Paper Adhesive Plaster 2 inch X 9m	Size-2 inch X 9 m per roll Material: Non woven, non-allergic & Strong adhesive but gentle to skin & painless removal, Porous and breathable CE certified	1's
20	5021	Paper Adhesive Plaster 3 inch X 9m	Size-3 inch X 9 m per rollMaterial: Non woven, non-allergic & Strong adhesive but gentle to skin & painless removal, Porous and breathableCE certified	1's
21	5022	Plaster of Paris BP Bandages 15cm X 2.7m	Plaster of Paris Bandages BP 15 cm X 2.7 m per Roll Joint free single roll Conforming to the standards as per IS 6237: 1971 (Reaffirmed Year: 2018) Count of yarn: Warp -NLT 21 tex Weft- NLT 21 tex Threads per dm: Ends (column)- NLT 150 Picks (Row)- NLT 75 Mass: NLT - 40 g / m2 Fast Setting, Superior casting Strength, E Reliable and longer lifespan, complying with standards as per British Pharmacopoeia.	l's in Paper rolled and in Sealed Poly pack
22	5023	Plaster of Paris BP Bandages 10cm X	Plaster of Paris Bandages BP 10 cm X 2.7 m per Roll Joint free single roll	1's in Paper rolled and in Sealed Poly pack

		2.7m / Doll	1071 (Deoffmand Verm. 2019)		
		2.7m / Roll	1971 (Reaffirmed Year : 2018) Count of yarn:		
			Warp -NLT 21 tex		
			Weft- NLT 21 tex		
			Threads per dm:		
			Ends (column)- NLT 150		
			Picks (Row)- NLT 75		
			Mass: NLT - 40 g / m2		
			Fast Setting, Superior casting Strength, E		
			Reliable and longer lifespan, complying with		
			standards as per British Pharmacopoeia.		
23	5024	Scalp Vein	Scalp Vein Set (Disposable) (18G)	1's in blister	
23	5024	Set Set	Disposable Stain less Steel sharp needle,	pack	
		(Disposabl	siliconized, Butterfly wings, Latex free PVC for		
		e) (18G)	better handling and fixation.		
		(100)	soft flexible		
			PUR/PVC tubing of 150-300 mm Length		
			including lyer connector and Cap.		
			ETO Sterilized with CE certification		
24	5025	Scalp Vein	Scalp Vein Set (Disposable) (20G)	1's in blister	
<u> </u>	5025	Set Set	Disposable Stain less Steel sharp needle,	pack	
		(Disposabl	siliconized, Butterfly wings, Latex free PVC for		
		e) (20G)	better handling and fixation.		
		(200)	soft flexible		
			PUR/PVC tubing of 150-300 mm Length		
			including lyer connector and Cap.		
			ETO Sterilized with CE certification		
25	5026	Scalp Vein	Scalp Vein Set (Disposable) (22G)	1's in blister	
23	5020	Set Set	Disposable Stain less Steel sharp needle,	pack	
		(Disposabl	siliconized, Butterfly wings, Latex free PVC for		
		e) (22G)	better handling and fixation.		
		(223)	soft flexible		
			PUR/PVC tubing of 150-300 mm Length		
			including lyer connector and Cap.		
			ETO Sterilized with CE certification		
26	5027	Scalp Vein	Scalp Vein Set (Disposable) (24G)Disposable	1's in blister	
20	002,	Set	Stain less Steel sharp needle, siliconized,	pack	
		(Disposabl	Butterfly wings, Latex free PVC for better		
		e) (24G)	handling and fixation. soft flexiblePUR/PVC		
		(213)	tubing of 150-300 mm Lengthincluding lyer		
			connector and Cap. ETO Sterilized with CE		
			certification		
27	5030	Surgical	Surgical Blade, Size No. 22	1's in peelable	
_,		Blade, No.	Stainless steel, well defined tip and uniform	Aluminium foil	
		22,	cutting edge.	pack	
		Sterilized	Pre - sterile with Gamma radiation, Confirm to		
		Stermized	the standards as per IS No.: 3319:1995 with		
			CE certification.		
28	5033	Surgical	Surgical Blade, Size No. 15	1's in peelable	
20		Blade, No.	Stainless steel, well defined tip and uniform	Aluminium foil	
		15,	cutting edge.	pack	
		Sterilized	Pre - sterile with Gamma radiation, Confirm to		
		Sterrized	the standards as per IS No.: 3319:1995 with		
			CE certification.		
		I		1	

29	5037	I.V Cannula	Cannula with Integrated 3-Way stop Cock. Size 26G having radio opaque catheter with CE	1's in blister pack
		(Sterile, Disposable	certification. Sterile Disposable (Single Use) Teflon/ PTFE	
), 26 G	I.V.	
30	5039	Sterile Disposable Spinal	Sterile Disposable Spinal Needle 22G x 3 ½ inch. Specially designed to administer lumbar/subarachnoid anesthesia. Transparent hub	1's in blister pack
		Needle 22G x 3 ½	provides rapid detection of Cerebro-spinal fluid (CSF flashback for confirming accurate	
		inch	placement. Sharp bevel design for low puncture force ensures minimal puncture trauma. Fine	
			gauze needle design greatly reduces the risk of PDPH (Post Dura Puncture Headache). Color coded as per ISO standard enables rapid size	
21	5040	C4amila	identification.)	1's in blister
31	5040	Sterile Disposable	Sterile Disposable Spinal Needle 25G x 3 ½ inch. Specially designed to administer lumbar/	pack
		Spinal Needle 25G x 3 ½	subarachnoid anesthesia. Transparent hub provides rapid detection of Cerebro-spinal fluid (CSF flashback for confirming accurate	
		inch	placement. Sharp bevel design for low puncture force ensures minimal puncture trauma. Fine	
			gauze needle design greatly reduces the risk of PDPH (Post Dura Puncture Headache). Color	
			coded as per ISO standard enables rapid size identification.)	
32	5041	Urine Collecting	Sterile Disposable Urine Collecting Bag 2000 ml with graduated volume marking, non toxic	1's pack in sealed poly pack
		Bag, Disposable,	pyrogen free, double seek, clinical grade PVC, Kink resistant flexible tubing not less than 90	puek
		2000ml	cm in length, should have non-return valve, Top drainage outlet, with non return input valve, CE certified.	
33	5046	Endotrache al Tube	ENDOTRACHEAL TUBE PLAIN SIZE 2.5 Single use sterile Standard 15 mm connector at	1's pack in sealed blister
		Plain, Size 2.5, Sterile,	proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	pack
		Single use	per is : 0501 1572 (Realitimed 2000)	
34	5047	Endotrache al Tube	ENDOTRACHEAL TUBE PLAIN SIZE 3 Single use sterile Standard 15 mm connector at	1's pack in sealed blister pack
		Plain, Size 3, Sterile, Single use	proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	
35	5048	Endotrache	ENDOTRACHEAL TUBE PLAIN SIZE 3	1's pack in
		al Tube	Single use sterile Standard 15 mm connector at	sealed blister pack
		Plain, Size 3.5, Sterile, Single use	proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	puek
36	5049	Endotrache	ENDOTRACHEAL TUBE PLAIN SIZE 4	1's pack in
		al Tube	Single use sterile Standard 15 mm connector at	sealed blister pack
i		Plain, Size	proximal end. Confirming to the standards as	pack

37	5050	Endotrache al Tube Plain, Size 4.5, Sterile, Single use	ENDOTRACHEAL TUBE PLAIN SIZE 4.5 Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack
38	5051	Endotrache al Tube Plain, Size 5, Sterile, Single use	ENDOTRACHEAL TUBE PLAIN SIZE 5Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack
39	5052	Endotrache al Tube Plain, Size 8.5, Sterile, Single use	ENDOTRACHEAL TUBE PLAIN SIZE 8.5 Single use sterile Standard 15 mm connector at proximal end. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	l's pack in sealed blister pack
40	5053	Endotrache al Tube Cuffed, Size 4	ENDOTRACHEAL TUBE CUFFED SIZE 4 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack
41	5054	Endotrache al Tube Cuffed, Size 4.5	ENDOTRACHEAL TUBE CUFFED SIZE 4.5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	I's pack in sealed blister pack
42	5055	Endotrache al Tube Cuffed, Size 5	ENDOTRACHEAL TUBE CUFFED SIZE 5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack
43	5056	Endotrache al Tube Cuffed, Size 6	ENDOTRACHEAL TUBE CUFFED SIZE 6 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	I's pack in sealed blister pack
44	5057	Endotrache al Tube Cuffed, Size 6.5	ENDOTRACHEAL TUBE CUFFED SIZE 6.5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	l's pack in sealed blister pack
45	5058	Endotrache al Tube Cuffed, Size 7	ENDOTRACHEAL TUBE CUFFED SIZE 7 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack
46	5059	Endotrache al Tube Cuffed, Size 7.5	ENDOTRACHEAL TUBE CUFFED SIZE 7.5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack
47	5060	Endotrache al Tube Cuffed, Size 8	ENDOTRACHEAL TUBE CUFFED SIZE 8 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as per IS: 6581 - 1972 (Reaffirmed 2008)	1's pack in sealed blister pack
48	5061	Endotrache al Tube Cuffed, Size 8.5	ENDOTRACHEAL TUBE CUFFED SIZE 8.5 Soft cuff towards the distal end Kink resistant inflation tube. Confirming to the standards as	1's pack in sealed blister pack

	1	T			
			per IS : 6581 - 1972 (Reaffirmed 2008)		
40	50.60	D 1 . 1	ENDOWN ACHEAL WIDE CHEED OF	1's pack in	
49	5062	Endotrache	ENDOTRACHEAL TUBE CUFFED SIZE 9	sealed blister	
		al Tube	Soft cuff towards the distal end Kink resistant	pack	
		Cuffed, Size 9	inflation tube. Confirming to the standards as		
			per IS : 6581 - 1972 (Reaffirmed 2008)		
50	5064	Tracheosto	Tracheostomy Tube (PVC), Sterilized, single	1's pack in sealed blister	
		my Tube	use(Tracheostomy Tube, made frommedical	pack	
		(PVC),	grade PVC, Plain, Sterile, Single Use - Size 24		
		Sterilized,	Soft flexibleflange at for easy fixation 15		
		single use	mmconnector at terminal end which canbe		
		Size-24	rotated in 360-degree directionNon-irritant,		
			Radio-opaque linethroughout the length. Size: 20 X60mm)Confirming to the standards IS		
			13179 : 1991 (Reaffirmed 2022)		
51	5065	Abdominal	Abdominal Drainage Kit, Sterile (2000ml, 24	1's	
J1	5005	Drainage	Graduate)		
		Kit, Sterile	ABDOMINAL DRAIN KIT, STERILE,		
		(2000ml,	HAVING DRAINAGE CATHETER AND		
		24	COLLECTION		
		Graduate)	BAG {2000 ml} size 24 Graduated Bag, Soft		
		Í	drainage		
			catheter 50 cm long, with radio opaque line,		
			Catheter with		
			markings at 2 cm interval		
52	5066	Abdominal	Abdominal Drainage Kit, Sterile (2000ml, 28	1's	
		Drainage	Graduate)		
		Kit, Sterile	ABDOMINAL DRAIN KIT, STERILE,		
		(2000ml,	HAVING DRAINAGE CATHETER AND		
		28	COLLECTION RAC (2000 ml) size 28 Creducted Box Soft		
		Graduate)	BAG {2000 ml} size 28 Graduated Bag, Soft drainage		
			catheter 50 cm long, with radio opaque line,		
			Catheter with		
			markings at 2 cm interval		
53	5067	Corrugated	Corrugated Drainage Sheet	1's in High	
		Drainage	Non toxic, non irritant medical grade extra soft	molecular PVC	
		Sheet	PVC, Radio opaque line, Sterile, individually	polybag	
			packed in a H.M polybag PVC does not		
			contribute to local inflammation.		
54	5070	Sterilized	ETO Sterilized Umbilical Cotton Tape 3mm x	1's in sealed	
		Umbilical	75cm	poly pack	
		Cotton			
		Tape 3mm			
	5051	x 75cm	G. 71. 1D. 0.5	11-1-1-1-1	
55	5071	Bone Wax,	Sterilized Bone wax 2.5 g	1's in peelable Aluminium foil	
		Sterilised		pack	
		2.5 g			
56	5073	Crepe	Crepe Bandage B.P. 6 cm x 4 M	1's	
		Bandage	Uniform plain weave and of continous length		
		B.P. 6 cm x	with no loose salvage threads (both sides		
		4 M	fastened edges),		

57	5077	Rapid Diagnostic Malaria Test Kit	Wrap threads- Two fold cotton threads, Not less than 170 threads per 10 cm. Weft threads- Combined cotton & Viscose, Not less than 78 threads per 10 cm. Elasticity- Not more than 80% of fully stretched length washing regains elasticity-fit for re-use. Weight- Not Less Than 140 g / m2 02 Nos. Loop & Hook Closure Rapid Diagnostic Malaria Test Kit should contain all the materials required for performing the test i.e. heparinized capillary tubes (Dia-1 mm) with relevant markings, reaction tube& stand, Test card (cassette); Sterile lancet, Reagents including buffer solution in a dropping bottle and Alcohol Swab. The test kit should be able to rapidly diagnose both P. falciparum (HRP2) and P. vivax (pLDH) and based on the principle of capture of parasite antigen from blood using monoclonal antibodies specific for antigen targets. PDS for Pf should be at least 95% sensitivity and specificity and for Pv should be at least 75% sensitivity and 90% specificity both at 200 parasites / M. the false positive rate should be	1's in hermetically sealed with poly pouch	
			parasites / µL. the false positive rate should be less than 10% and invalid rate should be less than 5%. Storage- 30oC to 45oCThe product should comply with ISO 9001 ISO 13485 (QMS)/CE/USFDA. Each Lot should PASS the test from ICMR designated labs.		
58	5078	Dengue Test Kit	The ELISA kit should be designed for qualitative detection of dengue NS1 antigen of all 4 dengue serotypes in human serum. The kit should be provided with the following materials and reagents: a) Anti- NS1 Antibody Coated Breakway Microwells (12*8=96 wells). Desiccant should be provided for storing the unused microwells which are to be resealed immediately. b) Horseradish peroxidase conjugated Anti-NS1 monoclonal antibody with preservatives. c) Chromogenic substrate in buffer d) Positive Control in the form of recombinant antigen with preservatives and antibiotics. e) Negative control in the form of confirmed negative human serum with preservatives and antibiotics. f) Calibrators in the form of recombinant antigen with preservatives and antibiotics g) Sample diluents h) Wash buffer The time required for performing the test for detection of dengue NS1 antigen by ELISA should range between 2-4 hours. The ELISA kit for detection of dengue NS1	I's in hermetically sealed with poly pouch	

59	5079	HCG-	antigen should have a sensitivity of >90% and a specificity of >95% taking RT-PCR as the gold standard. The kit should have a shelf-life of at least 6 months when stored at an ambient temperature of 2°C - 8°C. Transportation should be under cold chain. CE certified One step rapid qualitative test for detecting	1's in hermetically	
		Pregnancy Test Kit	HCG pregnancy hormone in urine. Each Test Kit contains: 01 Test device with aperture marked as 'S', 'C' for control and 'T' for test line. 01 Dropper, 01 Clean cup/tube. Reading time 5-10 min Hermetically sealed CE certified	sealed with poly pouch	
60	5083	Ear buds 100% Cotton	Ear buds 100% natural Absorbent Cotton IP free from optical whiteners/brightners, Stronge paper/wood stems, Cotton swabs should never debond while using/cleaning. Manufactured on fully automated machines and Untouched with Hands Size: 8 cm ± 0.5 cm	100's in Screw cap pet jar	
61	5094	Sterile Gauze Pad size 2X2	100% woven natural Absorbent Cotton IP, Individually wrapped and sterile. Lint-free and have no loose threads or raw edges. Size: 2 in X 2 in Confirming to the standards as per IS No.758/1988	1's in sealed poly pack	
62	5095	Sterile Gauze Pad size 3X3	100% woven natural Absorbent Cotton IP, individually wrapped and sterile. Lint-free and have no loose threads or raw edges. Size: 3 in X 3 in Confirming to the standards as per IS No.758/1989	l's in sealed poly pack	
63	5096	Sterile Gauze Pad size 4X4	100% woven natural Absorbent Cotton IP, individually wrapped and sterile.Lint-free and have no loose threads or raw edges.Size: 4 in X 4 inConfirming to the standards as per ISNo.758/1990	1's in sealed poly pack	
64	5097	Alcohol Swab (Spirit Swab)	Alcohol Swab (Spirit Swab) Four-layer wrapper provides air-tight seal - prevents leakage and drying out. Isopropyl Alcohol 70% V/V USP for optimum anti-bacterial action.	1's inair-tight sealed wrapper	
65	6001	Surgical Rubber Gloves- Disposable, Sterile, 6.5 inch	One pair of Disposable Sterile Surgical Rubber Gloves, Size 6 ½ Inches, Made of natural hypoallergic Latex, 100% electronically tested for holes, sterilised by Gamma Radiation / ETO, pre-powdered, Tensile strength as per EN 455-2. Powder should be non-allergenic	1's pair in sealed poly pouch	

			should Conform to IS 13422. ISI marked / CE certified / FDA.		
66	6002	Surgical Rubber Gloves- Disposable, Sterile, 7 inch	One pair of Disposable Sterile Surgical Rubber Gloves, Size 7 Inches, Made of natural hypoallergic Latex, 100% electronically tested for holes, sterilised by Gamma Radiation / ETO, pre-powdered, Tensile strength as per EN 455-2. Powder should be non-allergenic should Conform to IS 13422. ISI marked / CE certified / FDA.	1's pair in sealed poly pouch	
67	6003	Surgical Rubber Gloves- Disposable, Sterile, 7.5 inch	One pair of Disposable Sterile Surgical Rubber Gloves, Size 7.5 Inches, Made of natural hypoallergic Latex, 100% electronically tested for holes, sterilised by Gamma Radiation / ETO, pre-powdered, Tensile strength as per EN 455-2. Powder should be non-allergenic should Conform to IS 13422. ISI marked / CE certified / FDA.	1's pair in sealed poly pouch	
68	6004	Surgical Rubber Gloves- Disposable, Sterile, 8 inch	One pair of Disposable Sterile Surgical Rubber Gloves, Size 8 Inches, Made of natural hypoallergic Latex, 100% electronically tested for holes, sterilised by Gamma Radiation / ETO, pre-powdered, Tensile strength as per EN 455-2. Powder should be non-allergenic should Conform to IS 13422. ISI marked / CE certified / FDA.	1's pair in sealed poly pouch	
69	6009	Surgical Cap, Disposable (for Surgeons/N urses)	Disposable Surgical Cap, Should be manufactured from non woven fabric Blue / Green / colour Round upon wearing, with elastic Air permeable / breathable Should retain skin and hair particles	1's	
70	6010	Suction Catheter FG (5)	Suction Catheter FG (5), 50 cm, Sterilized Non-traumatic tip, non toxic medical grade PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Colour code for identification of size.	l's in sealed poly pack	
71	6011	Suction Catheter FG (6), 50 cm, Sterilized	Suction Catheter FG (6), 50 cm, Sterilized non-traumatic tip, non toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Colour code for identification of size. CE certified	1's in sealed poly pack	
72	6012	Suction Catheter FG (8), 50 cm, Sterilized	Suction Catheter FG (8), 50 cm, Sterilized non-traumatic tip, non toxic, PVC, integrated ring type (Open End), Four round lateral eyes facilitate efficient suctioning. Colour code for identification of size.CE certified	l's in sealed poly pack	

73	6013	Suction	Suction Catheter FG (10), 50 cm,	1's in sealed
		Catheter	Sterilizednon-traumatic tip, nontoxic, PVC,	poly pack
		FG (10), 50	integrated ring type (Open End), Four round	
		cm,	lateral eyes facilitate efficient suctioning.	
		Sterilized	Colour code for identification of size.CE	
			certified	
74	6014	Suction	Suction Catheter FG (12), 50 cm, Sterilized	1's in sealed
		Catheter	non-traumatic tip, nontoxic, PVC, integrated	poly pack
		FG (12), 50	ring type (Open End), Four round lateral eyes	
		cm,	facilitate efficient suctioning. Colour code for	
		Sterilized	identification of size. CE certified	
75	6015	Suction	Suction Catheter FG (14), 50 cm, Sterilized	1's in sealed
		Catheter	non-traumatic tip, non toxic, PVC, integrated	poly pack
		FG (14), 50	ring type (Open End), Four round lateral eyes	
		cm,	facilitate efficient suctioning. Colour code for	
		Sterilized	identification of size. CE certified	
76	6016	Suction	Suction Catheter FG (16), 50 cm, Sterilized	1's in sealed
		Catheter	non-traumatic tip, non toxic, PVC, integrated	poly pack
		FG (16), 50	ring type (Open End), Four round lateral eyes	
		cm,	facilitate efficient suctioning. Colour code for	
		Sterilized	identification of size. CE certified	
77	6017	Suction	Suction Catheter FG (18), 50 cm, Sterilized	1's in sealed
		Catheter	non-traumatic tip, non toxic, PVC, integrated	poly pack
		FG (18), 50	ring type (Open End), Four round lateral eyes	
		cm,	facilitate efficient suctioning. Colour code for	
		Sterilized	identification of size.CE certified	
78	6018	Suction	Suction Catheter FG (20), 50 cm, Sterilized	1's in sealed
		Catheter	non-traumatic tip, non toxic, PVC, integrated	poly pack
		FG (20), 50	ring type (Open End), Four round lateral eyes	
		cm,	facilitate efficient suctioning. Colour code for	
		Sterilized	identification of size. CE certified	
79	6020	Sterile	Sterile Catheter Single Use for Urinary	1's in sealed
		Catheter	Drainage (Foley Balloon Catheter), 2 way	poly pack
		Single Use	(8FG)	
		for Urinary	2 Way Silicone Coated Latex Foley Catheter,	
		Drainage	thin, flexible and sterile tube.	
		(Foley	Made of Silicone elastomer bonded with Latex.	
		Balloon	Should have hard plastic valve Smooth distal	
		Catheter), 2	end with smooth eyes for atraumatic intubation.	
		way (8FG)	Symmetrical foley balloon Balloon capacity 3-	
			5 ml.	
			Should conform to IS 11497.	
			Color coding marking to identify size. Length,	
			wall thickness and balloon capacity should be	
			mentioned as per IS 11497.	
			Specification for B, C, D, E, F, G should be	
			mentioned as per IS 11497 for particular size).	
80	6021	Sterile	Sterile Catheter Single Use for Urinary	1's in sealed
	0021	Catheter	Drainage (Foley Balloon Catheter), 2 way	poly pack
		Single Use	(10FG)	
		for Urinary	2 Way Silicone Coated Latex Foley Catheter,	
		Drainage	thin, flexible and sterile tube.	
		(Foley	Made of Silicone elastomer bonded with Latex.	
		Balloon	Should have hard plastic valve Smooth distal	
		Catheter), 2	end with smooth eyes for atraumatic intubation.	
		way	Symmetrical foley balloon Balloon capacity 3-	
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	1	I		T T	
		(10FG)	5 ml.		
			Should conform to IS 11497.		
			Color coding marking to identify size. Length,		
			wall thickness and balloon capacity should be		
			mentioned as per IS 11497.		
			Specification for B,C,D,E,F,G should be		
			mentioned as per IS 11497 for particular size).		
			CE certified		
81	6022	Sterile	Sterile Catheter Single Use for Urinary	1's in sealed poly pack	
		Catheter	Drainage (Foley Balloon Catheter), 2 way	poly pack	
		Single Use	(16FG)		
		for Urinary	2 Way Silicone Coated Latex Foley Catheter,		
		Drainage	thin, flexible and sterile tube.		
		(Foley	Made of Silicone elastomer bonded with Latex.		
		Balloon	Should have hard plastic valve Smooth distal		
		Catheter), 2	end with smooth eyes for atraumatic intubation.		
		way	Symmetrical foley balloon Balloon capacity 3-		
		(16FG)	5 ml.		
			Should conform to IS 11497.		
			Color coding marking to identify size. Length,		
			wall thickness and balloon capacity should be		
			mentioned as per IS 11497.		
			Specification for B, C, D, E, F, G should be		
0.2	6000	G . 11	mentioned as per IS 11497 for particular size)	1's in sealed	
82	6023	Sterile	Sterile Catheter Single Use for Urinary	poly pack	
		Catheter	Drainage (Foley Balloon Catheter), 2 way	Fred Fred	
		Single Use	(18FG)2 Way Silicone Coated Latex Foley		
		for Urinary	Catheter, thin, flexible and sterile tube. Made of		
		Drainage	Silicone elastomer bonded with Latex. Should		
		(Foley	have hard plastic valve Smooth distal end with		
		Balloon	smooth eyes for atraumatic intubation.		
		Catheter), 2	Symmetrical foley balloon Balloon capacity 3-		
		way	5 ml.Should conform to IS 11497.Color coding		
		(18FG)	marking to identify size. Length, wall thickness		
			and balloon capacity should be mentioned as		
			per IS 11497. Specification for B, C, D, E, F, G		
			should be mentioned as per IS 11497 for		
02	6024	Ctorilo	particular size) Storila Cotheter Single Use for Uniners	1's in sealed	
83	6024	Sterile	Sterile Catheter Single Use for Urinary Drainage (Follow Balloon Catheter), 2 years	poly pack	
		Catheter	Drainage (Foley Balloon Catheter), 2 way (20FG)		
		Single Use for Urinary	2 Way Silicone Coated Latex Foley Catheter,		
		Drainage	thin, flexible and sterile tube.		
		(Foley	Made of Silicone elastomer bonded with Latex.		
		Balloon	Should have hard plastic valve Smooth distal		
		Catheter), 2	end with smooth eyes for atraumatic intubation.		
		way	Symmetrical foley balloon Balloon capacity 3-		
		(20FG)	5 ml.		
		(2010)	Should conform to IS 11497.		
			Color coding marking to identify size. Length,		
			wall thickness and balloon capacity should be		
			mentioned as per IS 11497.		
			Specification for B, C, D, E, F, G should be		
			mentioned as per IS 11497 for particular size)		
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84	6025	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (22FG)	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (22FG) 2 Way Silicone Coated Latex Foley Catheter, thin, flexible and sterile tube. Made of Silicone elastomer bonded with Latex. Should have hard plastic valve Smooth distal end with smooth eyes for atraumatic intubation. Symmetrical foley balloon Balloon capacity 3- 5 ml. Should conform to IS 11497. Color coding marking to identify size. Length, wall thickness and balloon capacity should be mentioned as per IS 11497. Specification for B, C, D, E, F, G should be mentioned as per IS 11497 for particular size)	1's in sealed poly pack
85	6026	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (24FG)	Sterile Catheter Single Use for Urinary Drainage (Foley Balloon Catheter), 2 way (24FG) 2 Way Silicone Coated Latex Foley Catheter, thin, flexible and sterile tube. Made of Silicone elastomer bonded with Latex. Should have hard plastic valve Smooth distal end with smooth eyes for atraumatic intubation. Symmetrical foley balloon Balloon capacity 3- 5 ml. Should conform to IS 11497. Color coding marking to identify size. Length, wall thickness and balloon capacity should be mentioned as per IS 11497. Specification for B, C, D, E, F, G should be mentioned as per IS 11497 for particular size)	1's in sealed poly pack
86	6027	Infant Feeding Tube, Sterile (Size-10 FG, length 50 cm)	Infant Feeding Tube, Sterile (Size-10 FG, length 50 cm) Sterile, DEHP Free, non-toxic PVC, non-irritant, low friction, non-traumatic, CE certified	1's in sealed poly pack
87	6028	Infant Feeding Tube, Sterile (Size-8 FG, length 50 cm)	Infant Feeding Tube, Sterile (Size-8 FG, length 50 cm) Sterile, DEHP Free, non-toxic PVC, non-irritant, low friction, non-traumatic, CE certified	1's in sealed poly pack
88	6029	Infant Feeding Tube, Sterile (Size-5 FG, length 50 cm)	Infant Feeding Tube, Sterile (Size-5 FG, length 50 cm) Sterile, DEHP Free, non-toxic PVC, non-irritant, low friction, non-traumatic, CE certified	1's in sealed poly pack

89	6031	Sterile Disposable Infusion set with Microdrip (I.V.) (Pediatric use)	Sterile Disposable Perfusion Set (Infusion set) with built in Airway moulded chamber and Needle (Pediatric Use). Burette type measured volume chamber of 100 ml, Non - Toxic, Non Pyrogenic, sterilized by ETO. Drop size of approx. 60 drops per ml Injection port, latex free, for intermittent medication. Floating auto shut off valve (latex free) in burette. Soft and kink resistant PVC tubing 2.7 to 3.00 mm tube with fluid filter. Roller controller for flow control Tube length 150 cm 23G needle. Should conform to IS No.12655 (part-4 of 2003) with CE certification.	1's in sealed poly pack
90	6032	Sterile Disposable Infusion set with Microdrip (I.V.) (Adult use)	Sterile Disposable Perfusion Set (Infusion set) with built in Airway moulded chamber and Needle (Adult Use). Burette type measured volume chamber of 100 ml, Non - Toxic, Non Pyrogenic, sterilised by ETO. Drop size of approx. 60 drops per ml Injection port, latex free, for intermittent medication. Floating auto shut off valve (latex free) in burette. Soft and kink resistant PVC tubing 2.7 to 3.00 mm tube with fluid filter. Roller controller for flow control Tube length 150 cm 23G needle. Should conform to IS No.12655 (part-4 of 2003) with CE certification.	1's in sealed poly pack
91	6033	Insulin Syringe (40 units) with 30G needle	Insulin syringe (40 units) with (fixed) 30 G needle conform to IS 12227 and CE certified. Sterile and non- toxic	1's in sealed blister pack
92	6035	I.V Cannula (Sterile, Disposable), 18 G	I.V Cannula (Sterile, Disposable), 18 G Sterile Disposable (Single Use) Teflon/ PTFE I.V. Cannula with Integrated 3-Way stop Cock having radio opaque catheter with CE certification. Size 18G, Should conform to IS 10555	1's in sealed blister pack
93	6036	I.V Cannula (Sterile, Disposable), 20 G	I.V Cannula (Sterile, Disposable), 20 G Sterile Disposable (Single Use) Teflon/ PTFE I.V. Cannula with Integrated 3-Way stop Cock having radio opaque catheter with CE certification. Size 20G, Should conform to IS 10555 Standard	1's in sealed blister pack
94	6037	I.V Cannula (Sterile, Disposable), 22 G	I.V Cannula (Sterile, Disposable), 22 G Sterile Disposable (Single Use) Teflon/PTFE I.V. Cannula with Integrated 3-Way stop Cock having radio opaque catheter with CE certification. Size 22G, Should conform to IS 10555 Standard	1's in sealed blister pack
95	6038	I.V Cannula (Sterile, Disposable), 24 G	I.V Cannula (Sterile, Disposable), 24 G Sterile Disposable (Single Use) Teflon/ PTFE I.V. Cannula with Integrated 3-Way stop Cock having radio opaque catheter with CE certification. Size 24G, Should conform to IS 10555 Standard	1's in sealed blister pack

96	6039	Nasal Oxygen Cannula set, Twin bore (Adult)	Nasal Oxygen Cannula Set, Twin prong (Accessory for Compressed Air Breathing) Adult General specification: Suitable for easy application and efficient administration of oxygen. soft, light weight non-toxic material, non-irritating even in long term use anatomically fit Twin prong/ nasal tips are designed to ensure equal volume of oxygen to both the air passages. CE certified	1's in sealed blister pack
97	6040	Absorbent Gelatin Sponge, Sterilized (80mm x 50mm x 10mm)	Water-insoluble, hemostatic device prepared from purified skin gelatin, and capable of absorbing up to 45 times its weight of whole blood. Gamma Sterile & Ready to Use, Easy To Cut According To Need.	1's in sterile blister pack enclosed in an outer peelable wrapper
98	6042	Blood Transfusio n Set with Filter 170 Micron	Blood Transfusion Set 170 Micron(The soft Kink resistance, translucent tubing is prepared from medical grade PVC material, double drip clearly visible chamber facilitates visual access and rapid adjustment of fluid level. Specially design roller controller offer accurate regulation of infusion rate with self-sealing latex bulb to avoid any contamination and Easy flushing. Length 150 cm.) CE certified	1's in sealed blister pack
99	6045	Ryle's Tube/Naso gastric Tube (Size 10)	Soft kink, resistance PVC tubing for atraumatic intubation. Closed distal end should be coned with radio opaque Material for accurate intubation. Four lateral eyes for greater efficiency. Radio Opaque Line. Marking at 50, 60,70 cm from trip. Color coded funnel. With luer connecter / closure. Length 105 cm. CE certified	1's in sealed blister pack
	6046	Ryle's Tube/Naso gastric Tube (Size 12)	Soft kink, resistance PVC tubing for atraumatic intubation. Closed distal end should be coned with radio opaque Material for accurate intubation. Four lateral eyes for greater efficiency. Radio Opaque Line. Marking at 50, 60,70 cm from trip. Color coded funnel. With luer connecter / closure. Length 105 cm. CE certified	1's in sealed blister pack
101	6047	Ryle's Tube/Naso gastric Tube (Size 14)	Soft kink, resistance PVC tubing for atraumatic intubation. Closed distal end should be coned with radio opaque Material for accurate intubation. Four lateral eyes for greater efficiency. Radio Opaque Line. Marking at 50, 60,70 cm from trip. Color coded funnel. With luer connecter / closure. Length 105 cm. CE certified	1's in sealed blister pack
102	6048	Ryle's Tube/Naso gastric Tube (Size	Soft kink, resistance PVC tubing for atraumatic intubation. Closed distal end should be coned with radio opaque	1's in sealed blister pack

		16)	Material for accurate intubation.		
			Four lateral eyes for greater efficiency. Radio		
			Opaque Line.		
			Marking at 50, 60,70 cm from trip. Color coded		
			funnel.		
			With luer connecter / closure. Length 105 cm.		
			CE certified		
103	6049	Ryle's	Soft kink, resistance PVC tubing for atraumatic	1's in sealed blister pack	
		Tube/Naso	intubation.	blister pack	
		gastric	Closed distal end should be coned with radio		
		Tube (Size	opaque		
		18)	Material for accurate intubation.		
			Four lateral eyes for greater efficiency. Radio		
			Opaque Line. Marking at 50, 60,70 cm from		
			trip. Color coded funnel. With luer connecter /		
104	6050	W C	closure. Length 105 cm. CE certified	1's pair in	
104	6050	Knee Cap	Made of interlock weave bi-layered fabric	sealed poly	
		(Small)	(outer layer made of nylon and inner layer made	pack and in	
			of fine grade dermophillic & hypoallergenic cotton with in between air space), Four way	mono carton	
			stretchable, Heat resistant rubber with high		
			modulus of elasticity.		
			Anatomically designed shape should ensure no		
			Chondromalacia patella on prolonged use.		
			Better uniform compression & grip, simple		
			pull-on application, easy knee movement, snug		
			fitting.		
			Dimensions: Length - Not less than 33 cm (Un-		
			stretched)		
			Net Weight- Not less than 100 g for a pair		
			Product should be latex free and should not		
			deteriorate on contact with oil, balm or on		
			washing.		
105	6051	Knee Cap	Made of interlock weave bi-layered fabric	1's pair in	
		(Medium)	(outer layer made of nylon and inner layer made	sealed poly pack and in	
			of fine grade dermophillic & hypoallergenic	mono carton	
			cotton with in between air space), Four way		
			stretchable, Heat resistant rubber with high		
			modulus of elasticity.		
			Anatomically designed shape should ensure no		
			Chondromalacia patella on prolonged use.		
			Better uniform compression & grip, simple		
			pull-on application, easy knee movement, snug		
			fitting.		
			Dimensions: Length - Not less than 40 cm (Un-		
			stretched)		
			Net Weight- Not less than 120 g for a pair.		
			Product should be latex free and should not		
			deteriorate on contact with oil, balm or on		
106	6052	Knee Cap	washing. Made of interlock weave bi-layered fabric	1's pair in	
100	0032	(Large)	(outer layer made of nylon and inner layer made	sealed poly	
		(Large)	of fine grade dermophillic & hypoallergenic	pack and in	
			cotton with in between air space), Four way	mono carton	
			stretchable, Heat resistant rubber with high		
			modulus of elasticity. Anatomically designed		
<u> </u>	1	<u> </u>	modulus of classicity. Thintomically designed	<u>l</u>	

107	7001	Hot Water Bag (Small)	shape should ensure no Chondromalacia patella on prolonged use. Better uniform compression & grip, simple pull-on application, easy knee movement, snug fitting.Dimensions: Length - Not less than 45 cm (Un-stretched)Net Weight-Not less than 140 g for a pair.Product should be latex free and should not deteriorate on contact with oil, balm or on washing. Dimensions: Not less than 25 cm x 19 cm Capacity: Not less than 1.5 Litres Wall thickness: Not less than 1.2 mm Made with high-quality, odourless vulcanized rubber of tensile strength not less than 14	1's in sealed poly pack and in mono carton
			MN/m2 (approx 140 kgf/cm2) and elongation at break of not less than 500 percent. The product should have an in-built hanger at one side, tie in cap and uniform rib pattern on both sides at outer side. The product should comply with IS 1867	
108	8080	Surgical Face Mask, Disposable	Manufactured of 3 ply construction from non-woven poly prop fabric / SMS / Meltblown. Bacterial filtration efficiency (BFE) should not be less than 99%, should be heat sealed to keep 3 layers together, Standard Size- 17.5 cm x 9 cm, Color- Green / Blue Elastic ear loop of length 7 inch or more on one side and properly attached to avoid detachment while wearing. Should conform to the standards of IS 16289	1's
109	8082	Umbilical Catheter (For New Born)	UMBILICAL CATHETER (FOR NEWBORN) With female flexible mount. Open tip should be soft Non-toxic, medical grade PVC, Smooth round tip, atraumatic insertion, well-finished surface facilitate, smooth passage in the vein. Sterile, CE certified	1's in sealed blister pack
110	8088	Walking Stick Quadripod	Durable, lightweight, rustproof, single telescopic height adjustable shaft made of high grade aluminium alloy (corrosion resistant) with a molded handgrip and branching into four tips. Confirming standards to IS 5145 Handle: Made of injection-moulded polypropylene of grade 2340PC with rubber handgrips (e.g. injection-moulded polypropylene of grade 2340PC). Offset shape design to position your weight directly over the shaft, increasing stability and putting less pressure on the wrists. Net length- Not less than 150 mm Shaft: Made of extruded anodized aluminum (recyclable). The shaft should be in two parts, telescopic in nature. Height adjustable mechanism to be made out of stainless steel for high strength, low deformation and resistance to abrasion. The pin should be made of stainless	1's in sealed poly pack

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		steel, at least 6mm in diameter and electroplated to prevent corrosion. Clearance between sliding parts not to exceed 0.5mm.Net length- Not less than 600 mm extendable upto 1000 mm. Tips: Non-slip and replaceable, made of PU/PVC. Anti-rattle system to reduce noise made when walking. Net weight: 700-800 g		
111 80	Abdominal Belt Velcro, Cream color (Large)	Extra porous, dermophillic three panel Cotton/ nylon/ elastic webbing fabric. Broad hook and loop tape panel NLT 15 cm 02 Nylon reeves each not less than 1.5 cm width to prevent rolling over of belt. Length: Not less than 38 inch Width: Not less than 25 cm Net Weight: Not less than 260 g All sewing shall be done with lock stitches by turning in the free ends to depth. The abdominal belt shall be tailored neatly.	1's in sealed poly pack and in mono carton	
112 80	Abdominal Belt Velcro, Cream color (Small)	Extra porous, dermophillic three panel Cotton/nylon/elastic webbing fabric.Broad hook and loop tape panel NLT 15 cm02 Nylon reeves each not less than 1.5 cm width to prevent rolling over of belt.Length: Not less than 30 inch Width: Not less than 25 cmNet Weight: Not less than 220 gAll sewing shall be done with lock stitches by turning in the free ends to depth. The abdominal belt shall be tailored neatly.	1's in sealed poly pack and in mono carton	
113 80	Abdominal Belt Velcro, Cream color (Medium)	Extra porous, dermophillic three panel Cotton/nylon/elastic webbing fabric. Broad hook and loop tape panel NLT 15 cm 02 Nylon reeves each not less than 1.5 cm width to prevent rolling over of belt. Length: Not less than 34 inch Width: Not less than 25 cm Net Weight: Not less than 240 g All sewing shall be done with lock stitches by turning in the free ends to depth. The abdominal belt shall be tailored neatly.	l's in sealed poly pack and in mono carton	
114 80	Abdominal Belt Velcro, Cream color (XL)	Extra porous, dermophillic three panel Cotton/nylon/elastic webbing fabric. Broad hook and loop tape panel NLT 15 cm 02 Nylon reeves each not less than 1.5 cm width to prevent rolling over of belt. Length: Not less than 42 inch Width: Not less than 25 cm Net Weight: Not less than 280 g All sewing shall be done with lock stitches by turning in the free ends to depth. The abdominal belt shall be tailored neatly.	1's in sealed poly pack and in mono carton	
115 80	Abdominal Belt Velcro, Cream	Extra porous, dermophillic three panel Cotton/ nylon/ elastic webbing fabric. Broad hook and loop tape panel NLT 15 cm 02 Nylon reeves each not less than 1.5 cm	1's in sealed poly pack and in mono carton	

		color	width to prevent rolling over of belt.			
		(XXL)	Length: Not less than 46 inch			
			Width: Not less than 25 cm			
			Net Weight: Not less than 300 g			
			All sewing shall be done with lock stitches by			
			turning in the free ends to depth. The abdominal			
			belt shall be tailored neatly.			
116	8099	Cervical	Made of high-density polyurethane foam not	1's in sealed		
		Collar Soft	less than 60kg/m3, reinforced with very thick	poly pack and		
		(Small)	LDPE sheet not less than 1.5mm with rounded	in mono carton		
			edges and shaped to give a uniform mandible			
			support. Nylon sewing thread 210/6.			
			Eyelets: 3 in Number. Made up of virgin			
			polypropylene diameter not less than 20 mm.			
			Stockinette : Dermophillic and hypoallergenic			
			blend of cotton and rayon which should be free			
			from spinning, weaving and processing defects.			
			Fasteners, Hook and Loop Tape: The tape shall			
			have the minimum width of 5 cm and minimum			
			overlap 2.5 cm. Expandity of the collar should			
			be 5 cm.			
			Total Length: not less than 460 mm (excluding			
			fasteners)			
			Length between centre of extreme eyelets: not			
			less than 150 mm			
			Width between crest: Not less than 100 mm			
			Confirming to the standards of IS 11569			
117	8100	Cervical	Made of high-density polyurethane foam not	1's in sealed		
		Collar Soft	less than 60kg/m3, reinforced with very thick	poly pack and		
		(Medium)	LDPE sheet not less than 1.5mm with rounded	in mono carton		
			edges and shaped to give a uniform mandible			
			support. Nylon sewing thread 210/6. Eyelets: 3			
			in Number. Made up of virgin polypropylene			
			diameter not less than 20 mm. Stockinette:			
			Dermophillic and hypoallergenic blend of			
			cotton and rayon which should be free from			
			spinning, weaving and processing			
			defects. Fasteners, Hook and Loop Tape: The			
			tape shall have the minimum width of 5 cm and			
			minimum overlap 2.5 cm. Expandity of the			
			collar should be 5 cm. Total Length: not less			
			than 500 mm (excluding fasteners)Length			
			between centre of extreme eyelets: not less than			
			150 mmWidth between crest: Not less than 110			
			mmConfirming to the standards of IS 11569			
118	8101	Cervical	Made of high-density polyurethane foam not	1's in sealed		
		Collar Soft	less than 60kg/m3, reinforced with very thick	poly pack and		
		(Large)	LDPE sheet not less than 1.5mm with rounded	in mono carton		
			edges and shaped to give a uniform mandible			
			support. Nylon sewing thread 210/6.			
			Eyelets: 3 in Number. Made up of virgin			
			polypropylene diameter not less than 20 mm.			
			Stockinette : Dermophillic and hypoallergenic			
			blend of cotton and rayon which should be free			
			from spinning, weaving and processing defects.			
			Fasteners, Hook and Loop Tape: The tape shall			
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			have the minimum width of 5 cm and minimum overlap 2.5 cm. Expandity of the collar should		
			be 5 cm.		
			Total Length: not less than 550 mm (excluding		
			fasteners)		
			Length between center of extreme eyelets: not		
			less than 150 mm		
			Width between crest: Not less than 125 mm		
119	8102	Cervical	Confirming to the standards of IS 11569 Made of high-density polyurethane foam not	1's in sealed	
11)	0102	Collar Soft	less than 60kg/m3, reinforced with very thick	poly pack and	
		(Extra	LDPE sheet not less than 1.5mm with rounded	in mono carton	
		Large)	edges and shaped to give a uniform mandible		
			support. Nylon sewing thread 210/6.		
			Eyelets: 3 in Number. Made up of virgin		
			polypropylene diameter not less than 20 mm.		
			Stockinette: Dermophillic and hypoallergenic		
			blend of cotton and rayon which should be free from spinning, weaving and processing defects.		
			Fasteners, Hook and Loop Tape: The tape shall		
			have the minimum width of 5 cm and minimum		
			overlap 2.5 cm. Expandity of the collar should		
			be 5 cm.		
			Total Length: not less than 600 mm (excluding		
			fasteners)		
			Length between center of extreme eyelets: not less than 150 mm		
			Width between crest: Not less than 130 mm		
			Confirming to the standards of IS 11569		
120	8103	Thumb and	Made from breathable 03 layered PUF fused	1's in sealed	
		Wrist brace	fabric/ Soft Neoprene.	poly pack and in mono carton	
		(Small)	Pre-shaped and removable malleable metal		
			splint for thumb.		
			• Distal brace edge situated under the palmar		
			crease.Brace is contoured at the ulnar aspect of the		
			hand.		
			• High coverage of dorsal aspect of the hand.		
			• Support the carpometacarpal joint (CMC) and		
			immobilize the metacarpophalangeal joint		
			(MCP) of the thumb in a way that does not		
			inhibit the movements of the hands.		
121	8104	Thumb and	Wrist Circle-6", Length-7" Made from breathable 03 layered PUF fused	1's in sealed	
121	0107	Wrist brace	fabric/ Soft Neoprene.	poly pack and	
		(Medium)	Pre-shaped and removable malleable metal	in mono carton	
			splint for thumb.		
			Distal brace edge situated under the palmar		
			crease.		
			• Brace is contoured at the ulnar aspect of the hand.		
			High coverage of dorsal aspect of the hand.		
			• Support the carpometacarpal joint (CMC) and		
			immobilize the metacarpophalangeal joint		
			(MCP) of the thumb in a way that does not		
			inhibit the movements of the hands.		

			Wrist Circle-6"-7.5", Length-7"	
122	8105	Thumb and	Made from breathable 03 layered PUF fused	1's in sealed poly pack and
		Wrist brace	fabric/ Soft Neoprene. • Pre-shaped and	in mono carton
		(Large)	removable malleable metalsplint for thumb.•	
			Distal brace edge situated under the palmar	
			crease. • Brace is contoured at the ulnar aspect	
			of the hand. High coverage of dorsal aspect of	
			the hand. • Support the carpometacarpal joint (CMC) and immobilize the	
			metacarpophalangeal joint (MCP) of the thumb	
			in a way that does not inhibit the movements of	
			the hands. Wrist Circle-7.5"-8.5", Length-8"	
123	8106	Thumb and	Made from breathable 03 layered PUF fused	1's in sealed
		Wrist brace	fabric/ Soft Neoprene.	poly pack and in mono carton
		(Extra	Pre-shaped and removable malleable metal	III IIIOIIO Carton
		Large)	splint for thumb.	
			Distal brace edge situated under the palmar	
			crease.	
			• Brace is contoured at the ulnar aspect of the	
			hand.	
			• High coverage of dorsal aspect of the hand.	
			• Support the carpometacarpal joint (CMC) and immobilize the metacarpophalangeal joint	
			(MCP) of the thumb in a way that does not	
			inhibit the movements of the hands.	
			Wrist Circle-8.5" and above, Length-8"	
124	8113	Hernia Belt	Made up of soft and dermophillic fabric with	1's in sealed
		(Small)	adjustable pelvic and leg straps.	poly pack and in mono carton
			Moulded & removable anatomic ethafoam pads	
			to apply gradual pressure around the affected	
			area with focused pressure on the hernia, which	
			pushes the inguinal hernia back.	
125	8114	Knee Brace	Colour: Grey Knee Brace Belt - Excellent Support and	1's in sealed
143	0114	Belt	Targeted Pain Relief • May Relieve the Pain	poly pack and
		(Medium)	Associated with Mild Osteoarthritis (OA) of the	in mono carton
		(1.10010111)	Knee • Unilateral Hinge and Strapping	
			Configuration Apply Corrective Force to	
			Offload the Affected Side of the Joint • Wrap	
			around, Patient-Friendly Design - Easy to	
			Apply to Swollen Tender Knees • Neoprene	
			Provides	
			Therapeutic Warmth to Soothe the Aching Joint	
			• Support Straps Allow for Adjustable	
			Compression • Available in Medial and Lateral Designs for Left and Right Knee Applications •	
			Interchangeable Condyle Pads for Day-to-Day	
			Adjustment Size- Medium 16-17 fits	
126	8115	Knee Brace	Knee Brace Belt - Excellent Support and	1's in sealed
		Belt	Targeted Pain Relief • May Relieve the Pain	poly pack and in mono carton
		(Large)	Associated with Mild Osteoarthritis (OA) of the	
			Knee • Unilateral Hinge and Strapping	
			Configuration Apply Corrective Force to	

127 8116	IV Cannula Fixator (Medium	Offload the Affected Side of the Joint • Wrap Around, Patient-Friendly Design - Easy to Apply to Swollen Tender Knees • Neoprene Provides Therapeutic Warmth to Soothe the Aching Joint, Support Straps Allow for Adjustable Compression • Available in Medial and Lateral Designs for Left and Right Knee Applications • Interchangeable Condyle Pads for Day-to-Day Adjustment Size- Large 18-19 fits IV Cannula Fixer Good aesthetic appeal due to woven fast edges, Moisture responsive High Moisture Vapor Transmission Rate Film, Low	1's pack in tearable wrapper	
	6cm x 5cm)	allergy grid pattern adhesive, porous adhesive to allow skin breathing, Easy and Painless remove, because of thin-non adhesive edges and leaves no residue after remove. Medium: 6cm x 5cm		
128 8117	Cervical Pillow (Memory foam)	Ergonomically Double Contoured Cervical Pillow confirming the shape of neck to accommodate both long and short necks. Made from visco-elastic high density, open cell, flexible, polyurethane foam (memory foam). Density: 65-70 Kg/m3, Sag factor: >2.5Ball rebound resilience: < 5% Time to retourn 95% height: 14 sLength: 38 cm, Width: 47 cm, Crest Height1:12 cm, Crest Height2: 9.5 cm,Inner cover to protect the memory foam. Grey colour, Washable, hypoallergenic, durable, soft and smooth Zipper cover.	1's in sealed poly pack and in mono carton	
129 8120	Lancets, Round Sterile, 28G	Lancets Round Sterile Tip 28G One time use; 28G, in virgin packing Length: 30mm ± 2mm Compatibility: Should be compatible with Jan Aushadhi glucometer.	50 pcs/Box	
130 8123	Adhesive Surgical Paper Tape Size 5cmx5m	Breathable, Hypo-allergenic, latex-free, non-woven and micro-porous backing material. High-quality water-soluble adhesive for gentle bonding, holds well on damp skin and clean removal with minimal residue. Size: Size 5 cm x 5 m	1's	
131 8124	Adhesive Surgical Paper Tape Size 2.5cmx5m	Breathable, Hypo-allergenic, latex-free, non-woven and micro-porous backing material. High-quality water-soluble adhesive for gentle bonding, holds well on damp skin and clean removal with minimal residue. Size: 2.5 cm x 5 m	1's	
132 8125	Adhesive Surgical Paper Tape Size 1.25cmx5 m	Breathable, Hypo-allergenic, latex-free, non-woven and micro-porous backing material. High-quality water-soluble adhesive for gentle bonding, holds well on damp skin and clean removal with minimal residue. Size: 1.25 cm x 5 m	1's	
133 8127	Nebulizer Mask with Tubing (Adult)	Made of clear, latex free, non-toxic PVC, medium concentration, adjustable nose clip, non kinkable & non-autoclave Tube Length: 2 m	l's pack in sealed poly pack	

			10cc bowl with graduated volume markings			
			45-degree elbow, Adult elongated mask			
			Medication port to eliminate medication waste			
			Nebulization rate: 3cc / 10 mins. CE certified			
			Packing: Single Piece poly packed.			
134	8131	Plastic	Wide mouth, snap-on leak proof tie-on and	1's in sealed		
		Urine Pot	glow in dark lid, notched handle made from	LDPE pack		
		collector	durable polypropylene Plastic Urine Pot			
		(For both	collector, Compatible for Male & Female use.			
		Male &	Weight: Not less than 150 g, Graduated volume			
		Female)	marking upto 1000 ml. Colour: White			
135	8132	Plastic	Made of high virgin ABS material	1's in sealed		
		Bedpan	Wide plastic guard to prevent spills and a	LDPE pack		
			tapered end for easier placement. Portable lid			
			and built-in handle with grip. Nonstick ceramic			
			finish inner layer and smooth bottom.			
			Size: 40 cm X 30 cm 9 cm			
			Weight: Not less than 750 g			
			Colour: Biege			
136	8135	Breathing	Breathing Exerciser, Total 3 Chambers, Device	1's in sealed		
		Exerciser	is composed of base and central part divided	poly pack and in mono carton		
		(3	into	in mono carton		
		Chambers)	three chambers containing three small spheres			
		,	of			
			different size and color, connecting tube with			
			12mm OD connector and mouth piece, Flow			
			rates 600ml/sec, 900ml/sec, and 1200ml/sec by			
			using different colors of ball for easy			
			identification of the flow rates.			
			Material Specifications: - ABS for chamber, PE			
			for ball and mouthpiece and EVA for tubing.			
137	8150	3-way	3-way stopcock with 10 cm extension lineSet	1's in sealed		
		stopcock	Length 15 cm Priming Volume 1.06 ml, Bore 3	poly pack and in mono carton		
		with 10 cm	mm 1 Blue Stopcock Handle(s), 1 Female	in mono carton		
		extension	Luer(s), 1 Male Luer(s), made of Polyvinyl			
		line	Chloride (PVC)Lipid Resistant, Natural rubber			
			latex is not part of the material formulation,			
			DEHP Free. CE certified			
138	8151	Adhesive	Adhesive wound dressings, 25cm x 10cm	1's in sealed		
		wound	Consisting of breathable non-woven top layer	peelable wrapper		
		dressings,	and a low-adherent absorbent pad	wiappei		
		25cm x	Size: 25cm x 10cm, each dressing is			
		10cm	individually wrapped and sterile. Sterilization is			
			by ethylene oxide. CE certified			
139	8153	Arm Sling	Made from breathable 3 layer bonded fabric-	1's in sealed		
		Pouch	Outer 100% polyester knit wrap fabric with	poly pack and		
		(Large),	raising, Middle PUF layer for cushioning and	in mono carton		
		Grey	wrinkle free, Inner 100% polyester soft liner			
			wrap knit.			
			Adjustable and non-stretchable shoulder sling			
			of width not less than 4 cm with cushioned			
			ethafoam shoulder pad not less than 10 cm and			
			length 20 cm.			
			Should have Thumb cradle			
			Designed: For Both Elbow, Color: Grey			
			Ergonomic, smart, sleek and open design			
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			provides improved ventilation and enhanced comfort.		
			One touch opening of PP slide buckle.		
			Length: Not less than 42 cm		
140	8154	Arm Sling Pouch (Medium), Grey	Made from breathable 3 layer bonded fabric-Outer 100% polyester knit wrap fabric with raising, Middle PUF layer for cushioning and wrinkle free, Inner 100% polyester soft liner wrap knit. Adjustable and non-stretchable shoulder sling of width not less than 4 cm with cushioned ethafoam shoulder pad not less than 10 cm and length 20 cm. Should have Thumb cradle Designed: For Both Elbow, Color: Grey Ergonomic, smart, sleek and open design provides improved ventilation and enhanced comfort.	1's in sealed poly pack and in mono carton	
			One touch opening of PP slide buckle.		
			Length: Not less than 38 cm		
141	8155 8156	Baby Wipes (Pack of 20 wipes) Bed Bath Towel (Pack of 10 wipes)	Made from extra soft and ultra fine spunlace fabric. Alcohol free, paraben free, SLS free, hypoallergic to infant skin. Infused with 2% Chlorhexidine, Aloe vera and Glycerin, Mild fragrance. Size: 240 mm X 290 mm per wipe pH: 4.5 -6, Mass: > 36 g/m2, Moisture content: > 90 % by 98 % demineralized water. Breaking Strength: Machine direction (wet): > 30 N/min Cross direction (wet): > 2.5 N/min Made from extra soft and ultra fine spunlace fabric. Alcohol free, paraben free, SLS free, hypoallergic to infant skin. Infused with Aloe vera and Vitamin E (Veg origin), Mild fragrance, Microwaveable.	Pack of 20 wipes packed in sealed LDPE pack with resealable plastic flap Pack of 10 wipes packed in sealed LDPE pack with resealable plastic flap	
1/12	0150	Chlorhavid	Size: 200 mm X 150 mm per wipe pH: 4.5 -6, Mass: > 36 g/m2, Moisture content: > 90 % by 98 % demineralized water. Breaking Strength: Machine direction (wet): > 30 N/min Cross direction (wet): > 2.5 N/min	1's in sealed	
143	8158	Chlorhexid ine Gauze Dressing B.P 10cmx10c m (Sterile)	Chlorhexidine Gauze Dressing B.P. 10cm * 10cm (Sterile) Material- Leno Weave Cotton, Coated with Chlorhexidine 0.5% w/w with white Soft Paraffin, Color White.	peelable wrapper	
144	8160	Disposable Plastic Hand Gloves (Free size)	Disposable Plastic Hand Gloves (Free size)Disposable gloves made with transparent plastic material, polythene gloves.	Pair	

145 8	3161	Elastic gauze bandages 10cm	Elastic gauze bandages 10cm High quality elastic fabric, soft edges and porous adhesive mass, Water repellent, Air permeable, Thinner substrate, non-fray edges, Confirming standards of IS 16111	l's in sealed peelable wrapper
146 8	3162	Elastic gauze bandages 7.5cm	Elastic gauze bandages 7.5cm High quality elastic fabric, soft edges and porous adhesive mass, Water repellent, Air permeable, thinner substrate, non-fray edges, Confirming standards of IS 16111	1's in sealed peelable wrapper
147 8	3167	Karman cannula 4 mm	Flexible plastic cannula and with built-in adaptor Specially designed for aseptic medical termination of pregnancy, coned shaped distal end with two large lateral eyes, Proximal end fitted with MTP syringe or suction apparatus, individually packed in paper pouch & sterile. Confirming standards of IS 8313	1's in sealed blister pack
148 8	8168	Lumbar Spinal Brace (Grey)	Lumbar Spinal Brace (Grey) covers all dorsal, lumbar and sacral vertebras. It supports and immobilizes the spine in a neutral position which allows vital body movement. Dimensions: 55 x 42 x 5 cm Features Easy to use and comfortable Adjustable shoulder straps Immobilizes the spine Elastic abdominal panels offer the required compression. It is manufactured with rigid, anatomic customizable, splints which ensure ideal fitting, enhanced immobilization and accurate spinal posture, firm grip and self-tightening of the brace.	1's
149 8	8169	Magnetic Posture Corrector Back Support Belt - Posture Fit	Made from high quality 3-ply fabric front-loading plastic buckles, 12 premium magnets, Velcro straps, firm grip, high-quality breathable, porous neoprene, reinforced cross, double stitching.	1's
150 8	8177	Specican 30 ml	Made from non-toxic medical grade PVC, transparent with leakproof lid, Sterile, free from foreign particle. Sticker pasted for writing patient name. Weight: Not less than 25 g.	1's in sterile sealed poly pack
151 8	3178	Surgical Cap	Made from spun bond PP non-woven fabric and spun lace fabrics. The caps should be ultrasonically sealed and lighter gravity than hydrophobic PP fibres for excellent water repellency and air permeability resulting in providing comfort & dryness resulting in antibacterial and anti-fungal resistance Colour: Blue, Size: 21 inches with 02 LP Elastic Thickness: Not less than 15 gsm	1's

152 8179	Surgical shoe cap	Made from fabric that hot roll combining two layers of spun bond non-woven on outer side and three layer of melt blown non-woven fabric in the middle. 02 Top LP elastic closure. Thickness: Not less than 50 GSM PP + PE laminated (Waterproof) Height: 20 cm, Length: 410 mm	1's
153 8181	Transparen t film dressings (Sterile) 10cm X 12cm	Each peelable sterile wrapper contains one thin film backing with a non-latex, hypoallergenic adhesive. The Film with border should be notched and reinforced with soft cloth tape to provide a better seal around catheters and other devices. The dressing should be waterproof, breathable, allowing good oxygen and moisture vapor exchange. The dressing should be impermeable to liquids, bacteria, and viruses of size > 30 nm. Size: 10cm X 12cm CE certified	1's in peelable Aluminium foil pack
154 8182	Vein Compressi on Stocking Knee length (Extra Large)	Each monocarton should contain:01 glider01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking knee length (open toe type). Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >30 cm Calf >40 cm Popliteal >40 cm Each measuring position should retain not less than 85% compression after 30 wash cycles. Tests as per conforming to IS 16467.	l's pair and glider in sealed poly pack and in mono carton
155 8183	Vein Compressi on Stocking Knee length (Large)	Each monocarton should contain: 01 glider 01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking knee length (open toe type). Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to	1's pair and glider in sealed poly pack and in mono carton

156 8184 Vein Compressi on Stocking Knee length (Medium)	ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >27 cm Calf >38 cm Popliteal >38 cm Each measuring position should retain not less than 85% compression after 30 wash cycles. Tests as per conforming to IS 16467. Each monocarton should contain: 01 glider 01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking knee length (open toe type). Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm Calf >35 cm Popliteal > 35 cm Each measuring position should retain not less	1's pair and glider in sealed poly pack and in mono carton	
157 8185 Vein Compressi on Stocking Knee length (Small)	than 85% compression after 30 wash cycles. Tests as per conforming to IS 16467. Each monocarton should contain:01 glider01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking knee length (open toe type). Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg)Ankle >19 cm	1's pair and glider in sealed poly pack and in mono carton	

			Calf >32 cm		
			Popliteal > 32 cm		
			Each measuring position should retain not less		
			than 85% compression after 30 wash		
			cycles.Tests as per conforming to IS 16467.		
158	8186	Vein	Each monocarton should contain: 01 glider.	1's pair and glider in sealed	
		Compressi		poly pack and	
		on	01 pair of latex free, ultra-thin, dermophillic &	in mono carton	
		Stocking	breathable vein compression stocking knee		
		Knee	length (open toe type).		
		length	Made up of plaited knitted PA microfilament		
		(XXL)	yarns 60 dtex f 60 and elastane yarn in the		
			count 22/17 dtex f 7.		
			Interrupted band and 2-ply gusset to prevent		
			tourniquet effect at the femoral vein.		
			Inlay circumferential knit, helping to provide		
			one way horizontal stretch.		
			A pressure break at the popliteal vein helps to		
			ensure that blood will continue to flow		
			smoothly through this critical area.		
			A reinforced and defined heel pocket to aid		
			correct placement. Cuff not less than 5cm to		
			hold stocking at place.		
			Circumference Compression (mm of Hg)		
			Ankle >33 cm		
			Calf >45 cm		
			Popliteal > 45 cm		
			Each measuring position should retain not less		
			than 85% compression after 30 wash cycles.		
			Tests as per conforming to IS 16467.		
159	8187	Vein	Each monocarton should contain:	1's pair and	
		Compressi	01 glider	glider in sealed poly pack and	
		on	01 pair of latex free, ultra-thin, dermophillic &	in mono carton	
		Stocking	breathable vein compression stocking Thigh		
		Thigh	length (open toe type).		
		length	Made up of plaited knitted PA microfilament		
		(Extra	yarns 60 dtex f 60 and elastane yarn in the		
		Large)	count 22/17 dtex f 7.		
			Interrupted band and 2-ply gusset to prevent		
			tourniquet effect at the femoral vein.		
			Inlay circumferential knit, helping to provide		
			one way horizontal stretch.		
			A pressure break at the popliteal vein helps to		
			ensure that blood will continue to flow		
			smoothly through this critical area.		
			A reinforced and defined heel pocket to aid		
			correct placement. Silcone dotted cuff not less		
			than 5cm to hold stocking at place.		
			Circumference Compression (mm of Hg)		
			Ankle >33 cm		
			Calf >45 cm		
			Popliteal > 45 cm		
			Lower Thigh >60 cm		
			Upper Thigh >70 cm		
				·	

			Each measuring position should retain not less		
			than 85% compression after 30 wash cycles.		
			Tests as per conforming to IS 16467.		
160	8188	Vein	Each monocarton should contain:01 glider01	1's pair and	
		Compressi	pair of latex free, ultra-thin, dermophillic &	glider in sealed poly pack and	
		on	breathable vein compression stocking Thigh	in mono carton	
		Stocking	length (open toe type). Made up of plaited		
		Thigh	knitted PA microfilament yarns 60 dtex f 60		
		length	and elastane yarn in the count 22/17 dtex f		
		(Large)	7.Interrupted band and 2-ply gusset to prevent		
			tourniquet effect at the femoral vein.Inlay		
			circumferential knit, helping to provide one		
			way horizontal stretch. A pressure break at the		
			popliteal vein helps to ensure that blood will		
			continue to flow smoothly through this critical		
			area. A reinforced and defined heel pocket to aid		
			correct placement. Silcone dotted cuff not less		
			than 5cm to hold stocking at place.		
			Circumference Compression (mm of Hg)		
			Ankle >27 cm		
			Calf >39 cm		
			Popliteal > 39 cm		
			Lower Thigh >55 cm		
			Upper Thigh >63 cm		
			Each measuring position should retain not less		
			than 85% compression after 30 wash		
			cycles. Tests as per conforming to IS 16467.		
161	8189	Vein	Each monocarton should contain:	1's pair and	
		Compressi	01 glider	glider in sealed poly pack and	
		on	01 pair of latex free, ultra-thin, dermophillic &	in mono carton	
		Stocking	breathable vein compression stocking Thigh		
		Thigh	length (open toe type).		
			Made up of plaited knitted PA microfilament		
		length	Made up of planted kineted 111 interofficialient		
		length (Medium)	yarns 60 dtex f 60 and elastane yarn in the		
		_	<u> </u>		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7.		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch.		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area.		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place.		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg)		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm Calf >36 cm		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm Calf >36 cm Popliteal >36 cm		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm Calf >36 cm Popliteal >36 cm Lower Thigh >45 cm		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm Calf >36 cm Popliteal >36 cm Lower Thigh >45 cm Upper Thigh >57 cm		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm Calf >36 cm Popliteal >36 cm Lower Thigh >45 cm Upper Thigh >57 cm Each measuring position should retain not less		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm Calf >36 cm Popliteal >36 cm Lower Thigh >45 cm Upper Thigh >57 cm Each measuring position should retain not less than 85% compression after 30 wash cycles.		
		_	yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >23 cm Calf >36 cm Popliteal >36 cm Lower Thigh >45 cm Upper Thigh >57 cm Each measuring position should retain not less		

163	8190	Vein Compressi on Stocking Thigh length (Small) Vein Compressi on Stocking Thigh length (XXL)	Each monocarton should contain:01 glider01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking Thigh length (open toe type).Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7.Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein.Inlay circumferential knit, helping to provide one way horizontal stretch.A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area.A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >19 cm Calf >33 cm Popliteal >33 cm Lower Thigh >40 cm Upper Thigh >50 cm Each measuring position should retain not less than 85% compression after 30 wash cycles.Tests as per conforming to IS 16467. Each monocarton should contain: 01 glider 01 pair of latex free, ultra-thin, dermophillic & breathable vein compression stocking Thigh length (open toe type). Made up of plaited knitted PA microfilament yarns 60 dtex f 60 and elastane yarn in the count 22/17 dtex f 7. Interrupted band and 2-ply gusset to prevent tourniquet effect at the femoral vein. Inlay circumferential knit, helping to provide one way horizontal stretch. A pressure break at the popliteal vein helps to ensure that blood will continue to flow smoothly through this critical area. A reinforced and defined heel pocket to aid correct placement. Silcone dotted cuff not less than 5cm to hold stocking at place. Circumference Compression (mm of Hg) Ankle >33 cm Calf >45 cm Popliteal >45 cm Lower Thigh >63 cm Upper Thigh >75 cm Each measuring resition schould retain not less Each measuring resition schould retain not less Each measuring resition schould retain not less than 5cm to hold stocking at place.	1's pair and glider in sealed poly pack and in mono carton 1's pair and glider in sealed poly pack and in mono carton	
			Upper Thigh >75 cm Each measuring position should retain not less		
			than 85% compression after 30 wash cycles.		
			Tests as per conforming to IS 16467.		

	8192	Zig zag cotton	Net weight 100 g arranged in zig zag pattern, square shaped. Made from 100% clean and soft cotton Natural fibres of average staple length not less than 10 mm, soft and fluffy, high absorption	20's in sealed LDPE pack	
	9039	Knee Cap with patellar ring (Large)	Made of interlock weave bi-layered fabric (outer layer made of nylon and inner layer made of fine grade dermophillic & hypoallergenic cotton with inbetween air space), Four way stretchable, heat-resistant rubber with high modulus of elasticity, anatomically designed. Silicone patellar ring with padding with antislip coating. Multiaxial side splint to prevent roll on and easy flexion. Better compression & grip, simple pull-on application, easy knee movement. Dimensions: Length - Not less than 40 cm (Unstretched) Net Weight- Not less than 190 g for a pair. Product should be latex free and should not deteriorate on contact with oil, balm or on washing.	1's pair in sealed poly pack and in mono carton	
166	9040	Knee Cap with patellar ring (Medium)	Made of interlock weave bi-layered fabric (outer layer made of nylon and inner layer made of fine grade dermophillic & hypoallergenic cotton with in between air space), Four way stretchable, heat resistant rubber with high modulus of elasticity, Anatomically designed. Silicone patellar ring with padding with antislip coating. Multiaxial side splint to prevent roll on and easy flexion. Better compression & grip, simple pull-on application, easy knee movement. Dimensions: Length - Not less than 40 cm (Unstretched) Net Weight- Not less than 180 g for a pair. Product should be latex free and should not deteriorate on contact with oil, balm or on washing.	I's pair in sealed poly pack and in mono carton	
167	9041	Knee Cap with patellar ring (Small)	Made of interlock weave bi-layered fabric (outer layer made of nylon and inner layer made of fine grade dermophillic & hypoallergenic cotton with inbetween air space), Four way stretchable, heat resistant rubber with high modulus of elasticity, Anatomically designed. Silicone patellar ring with padding with antislip coating. Multiaxial side splint to prevent roll on and easy flexion. Better compression & grip, simple pull-on application, easy knee movement. Dimensions: Length - Not less than 35 cm (Unstretched) Net Weight- Not less than 170 g for a pair.	1's pair in sealed poly pack and in mono carton	

			Product should be latex free and should not deteriorate on contact with oil, balm or on washing.		
168	9042	Nebulizer Mask with Tubing (Child)	Made of clear, latex free, non-toxic PVC, medium concentration, adjustable nose clip, non kinkable & non-autoclave Tube Length: 2 m 10cc bowl with graduated volume markings 45-degree elbow, Child elongated mask Medication port to eliminate medication waste ue Nebulization rate: 3cc / 10 mins. CE certified Packing: Single Piece poly packed.	1's pack in sealed poly pack	
169	9043	Walker (Sit Assist-to- stand support)	Durable, lightweight, rust proof, foldable, height adjustable shaft made of high grade aluminium alloy (corrosion resistant) with 2 pairs of injection-moulded polypropylene of grade 2340 PC with rubber handgrips. Confirming standards to IS 5145 Shaft: Made of extruded anodized aluminum (recyclable). Each vertical shafts should be in two parts, telescopic in nature. 02 'U' shaped Front horizontal shaft and one with pellet button for folding. Height adjustable mechanism to be made out of stainless steel for high strength, low deformation and resistance to abrasion. The pin should be made of stainless steel, at least 6mm in diameter and electroplated to prevent corrosion. Clearance between sliding parts not to exceed 0.5mm. Foldable Depth: Not more than 100 mm. Frontal Height: 650 mm extendable up to 900 mm Back Height: minimum 450 mm and extendable at par. Frontal width: Not less than 500 mm Lateral width: Not less than 300 mm Tips: Non-slip and replaceable, made of durable PU/PVC. Anti-rattle system to reduce noise made when walking.	1's in sealed poly pack	
170	9044	Elbow crutch (Adjustable)	Net weight: 1900 g- 2000 g Durable, lightweight, rust proof, height adjustable shaft made of high-grade aluminium alloy (corrosion resistant) with a molded handgrip. Confirming standards to IS 5145Handle: Made of injection-moulded polypropylene of grade 2340PC with rubber handgrips (e.g. injection-moulded polypropylene of grade 2340PC). Net length-Not less than 150 mmShaft: Made of extruded anodized aluminum (recyclable). The shaft should be double adjustable – from floor to handle & from handle to cuff in two parts. Height adjustable mechanism to be made out of	1's in sealed poly pack	

stainless steel for high strength, low deformation and resistance to abrasion. The pin should he made of stainless steel, at least 6mm in diameter and electroplated to prevent corrosion. Clearance between sliding parts not to exceed 0.5mm. Elbow shaft with flipping elbow grip should be anatomically aligned with handle to bear body weight and decrease fatigue. Net length: Main shaft. Not less than 600 mm and extendable up to 1000 mm. Elbow Shaft. Not less than 600 mm and extendable up to 300 mm. Tips: Non-slip and replaceable, made of durable PU/PVC.Amt-rattle system to reduce noise made when walking. Net weight: 400-500 g. 171 9045 Insulting/M edicine Material: Linux Hamiltonian Material: Linux Hamil					
edicine Cooling Case Naterial: 1. Outer Fabric – 1000 Denier Polyester Polyurethane (PU) coated, (0.5mm), Durable, finely woven and water-resistant fabric. 2. Inner Fabric - 4 x 4 Polyester Matty Soft PU coated, light weight, highly flexible and moisture resistant. 3. Polyester Zipper - Size 5 with Metal Slider. 4. PP (Polypropylene) Tape – covering panels and joints inside the Box. 5. Pippin Wire (Tar) – all around outer side, made of LDPE material. 6. White PP Sheet – 0.45mm, Polypropylene Sheet – High temperature & Corrosion Resistant. 7. Aluminum laminated Fabric – Reflect thermal radiation. 8. PVC Clear Film – 0.15mm – Used for moisture control and to protect Aluminized Fabric. 9. Woolen Felt Fabric – high graded for inner temperature protection. App. 2 mm thick. 10. Polyester Net/Mesh - for Inside pockets with Polyester Elastic. 11. XLDE one side Alu Foil laminated Foam 6mm (Cross-linked polyethylene) – Highly Temperature Resistant, best for thermal insulation. 12. Pen Folder: Size 6.5" x 7", Folded Size 6.5" x 3.5" A - 4 x 4 Matty Soft PU coated, B - Aluminum laminated Fabric – Reflect thermal radiation C - PVC Clear Film – 0.18mm D - Polyester Net/Mesh E - Embedded with PP Tape F - Metal Snap Button (VTS)			deformation and resistance to abrasion. The pin should be made of stainless steel, at least 6mm in diameter and electroplated to prevent corrosion. Clearance between sliding parts not to exceed 0.5mm. Elbow shaft with flipping elbow grip should be anatomically aligned with handle to bear body weight and decrease fatigue.Net length:- Main shaft- Not less than 600 mm and extendable up to 1000 mm.Elbow Shaft- Not less than 200 mm and extendable up to 300 mmTips: Non-slip and replaceable, made of durable PU/PVC.Anti-rattle system to reduce noise made when walking.Net weight: 400-500 g		
	9045	edicine Cooling	Material: 1. Outer Fabric – 1000 Denier Polyester Polyurethane (PU) coated, (0.5mm), Durable, finely woven and water-resistant fabric. 2. Inner Fabric - 4 x 4 Polyester Matty Soft PU coated, light weight, highly flexible and moisture resistant. 3. Polyester Zipper - Size 5 with Metal Slider. 4. PP (Polypropylene) Tape – covering panels and joints inside the Box. 5. Pippin Wire (Tar) – all around outer side, made of LDPE material. 6. White PP Sheet – 0.45mm, Polypropylene Sheet – High temperature & Corrosion Resistant. 7. Aluminum laminated Fabric – Reflect thermal radiation. 8. PVC Clear Film – 0.15mm – Used for moisture control and to protect Aluminized Fabric. 9. Woolen Felt Fabric – high graded for inner temperature protection. App. 2 mm thick. 10. Polyester Net/Mesh - for Inside pockets with Polyester Elastic. 11. XLPE one side Alu Foil laminated Foam 6mm (Cross-linked polyethylene) – Highly Temperature Resistant, best for thermal insulation. 12. Pen Folder: Size 6.5" x 7", Folded Size 6.5" x 3.5" A – 4 x 4 Matty Soft PU coated, B - Aluminum laminated Fabric – Reflect thermal radiation C - PVC Clear Film – 0.18mm D - Polyester Net/Mesh E – Embedded with PP Tape F – Metal Snap Button (VT5)	Case, 2 unit Gel Cool Pack, 1 Unit Insulin Pen Folder and	

Material: Carbopol, Triethanolamine (TEA)	
with	
preservative and water in LDPE pouch.	
preservative and water in LDFE pouch.	
14. Gel-Ice Pouch Cover – Moisture resistant	
Poly Fabric	
15. Carry Bag with Strings – Water proof	
Laminated Taffeta Fabric	

ANNEXURE XIV

[Ref. clause no. 4 (C)]

(To be submitted on firm's letterhead duly authorized)

DECLARATION FOR NON-DRUG ITEM(S)

I/we			in the capacity of Proprietor/Managing
Partner /	Managing	Director in M/s	
having i	ts registere	ed office at	and its factory premises
at			do hereby declare
that the o	quoted iten	n(s) are neither covered under Dru	gs & Cosmetics Act 1940 and Rule their
under noi	Under Me	dical Device Rule 2017.	
That I/w	e are eligil	ble to participate in the tender no.	PMBI/SURGICAL/RC-210/2023 for the
	_	• •	d down in the tender document along with
_			nentioned by various ministry/department
	n the subje		ionizoned by various ministry, apparement
referred f	ii tiie suoje	et tender.	
Sl. No.	Item No.	Specification of the Item	Production Capacity (Per Annum)
		_	ty's right to forfeit the Performance Security period of 5 years if, any information furnished
-	-	se at any time during the contract period	· · · · · · · · · · · · · · · · · · ·
		, , ,	
		Signed	
		Name: .	
		Designa	tion
		(Compa	ny Seal)
		(Above sh	all be furnished by Authorized Signatory
		•	

ANNEXURE-XV

[Ref. clause no. 3(H)]

(To be submitted on non-judicial stamp paper dully notarized)

DECLARATION FOR NON-BLACKLISTING

I/we			, am/are ir	n the	capa	city	of Proprietor/Ma	naging
Partner /	Managing Director	in M/s						
having its	registered office at .				•••••		and its factory pr	emises
at							do hereby decla	re that
our compa	any/applied items hav	ve not been blacklis	ted/debarred/d	de-regi	stered	/banne	d due to quality	failure
for the qu	oted product /firm l	by any State Gover	nment / Cen	tral Go	overni	ment/	PMBI/ Central o	r State
submissio	ent's Drug procurer n of bid. e are eligible to part	J		•	Î	oted in	the tender at the t	ime of
S.N.	ITEM NO.	GENERIO	C NAME OI	F ITE	M		UNIT SIZE	,
		Si	gned					
		N	ame:		••••			
		D	esignation					
		(0	Company Seal)				
		(A	bove shall b	e furi	nishe	d by A	Authorized Sign	atory)

ANNEXURE- XVI

Enclosure-I

(Ref. Clause No. - 3.U)

(To be submitted on Non-judicial Stam paper duly notarized)

DECLARATION OF LOCAL CONTENT

							Resident
/Mai	naging Direct stered office	etor in M/s.					and factory premises
			content for th				
	S. No.	Item code	Item Name	Details Location which addition	value	Percentage (%) of Local content	Category of Bidder
_	1						
-	3						
in guidelines issued by Department of Pharmaceuticals, Ministry of Chemicals & fertilizers, Government of India vide F. No. 31026/36/2016-MD dated 09.11.2020 and DPIIT order no. P-45021/2/2017-PP(BE-II) dated 16.09.2020 for the implementation of provisions of Public Procurement (Preference to Make in India) Order (PPO) 2017 related to procurement of Goods and Services on behalf of M/s							
				Signa	ture		
				Name	: :		
				Desig	gnation		
				(Com	pany Seal	/Stamp)	
				(Abo	ve shall b	e furnished by	Authorized Signatory)

ANNEXURE- XVII [Ref. Clause No. 3.1]

Declaration by Authorized Signatory towards the Compliance of Order (Public Procurement No.1, 2 & 3

(On Non-Judicial Paper duly notarized)

I, the undersigned,	· · · · · · · · · · · · · · · · · · ·
1) The facts contained herein are within my own person	onal knowledge.
2) I have read the Order (Public Procurement No.1, 2 subject of Restrictions under Rule 144 (xi) of the Gerestrictions on procurement from a bidder of a count comply to all the provisions of the Order.	eneral Financial Rules (GFRs), 2017 regarding
3) I certify that M/s	ke out whichever is not applicable), has been by that this SUPPLIER fulfils all requirements in applicable, evidence of valid registration by the and / or if certificate / declaration given by bidder entity) is found to be false, this would action in accordance with law as per Clause 12
Signatu	re
Name:	
Designa	ntion
(Compa	any Seal/Stamp)
(Above sh	all be furnished by Authorized Signatory)

ANNEXURE- XVIII [Ref. Clause No. 4. P (ii)]

SAMPLE RECEIPT

(To be submitted in triplicate on firm's letterhead along with samples)

TENDER NO						Dated			
	Please	receive	e following Sam	ples of Sur	gical/ Consumable	es and Medical D	evices.		
	M/s	• • • • • • • • • • • • • • • • • • • •	•••••	•••••					
	•••	•••••	• • • • • • • • • • • • • • • • • • • •	•••••					
	S.N	Item No.	Product Name	Batch no.	Date of Manufacturing	Date of Expiry	License No	Number of units submitted	
T	otal nui	mber of	Items submitte	d:	•••••				
T	otal nui	mber of	Boxes submitte	d:	••••				
T	otal Nu	mber of	f signed pages of	f Pilot stud	ies / Publications:	•••			
P	lace:								
D	ate:								
ע	ate.				C' 1				
					Name:				
					Designation				
					(Company Seal)				
					(Abov	e shall be furnished	by Authoriz	ed Signatory)	
							Yours	s faithfully,	
								G 11	

Sd/-DGM (Procurement & Quality) For & on behalf of PMBI Ph: 011-49431800(811)