

## अनुबंध | Contract



अनुबंध क्रमांक | Contract No: GEMC-511687725156094

अनुबंध तिथि | Generated Date : 12-Apr-2024

## संगठन विवरण | Organisation Details

प्ररूप | Type : State Government  
मंत्रालय | Ministry : -  
विभाग | Department : Health and Family Welfare Department Delhi  
संगठन का नाम | Organisation Name : N/A  
कार्यालय क्षेत्र | Office Zone: LBS HOSPITAL EAST DELHI

## खरीदार विवरण | Buyer Details

पद | Designation : purchase officer  
संपर्क नंबर | Contact No. : 011-21200729-  
ईमेल आईडी | Email ID : buyer1.hfw.del@gembuyer.in  
जीएसटीआईएन | GSTIN : -  
पता | Address : Lal Bahadur Shastri Hospital, Khichripur, EAST DELHI, DELHI-110091, India

## वित्तीय स्वीकृति विवरण | Financial Approval Detail

आईएफडी सहमति | IFD Concurrence : No  
प्रशासनिक अनुमोदन का पदनाम | Designation of Administrative Approval: Medical Superintendent  
वित्तीय अनुमोदन का पदनाम | Designation of Financial Approval : Medical Superintendent

## भुगतान प्राधिकरण विवरण | Paying Authority Details

Role: PAO  
भुगतान का तरीका | Payment Mode: Offline  
पद | Designation : AAO/DDO  
ईमेल आईडी | Email ID : indra.giri22@gov.in  
जीएसटीआईएन | GSTIN : N  
पता | Address: Lal Bahadur Shastri Hospital, Khichripur, EAST DELHI, DELHI-110091, India

## विक्रेता विवरण | Seller Details

जेम विक्रेता आईडी | GeM Seller ID : XWT6220006449658  
कंपनी का नाम | Company Name : JAIMANTI LIFECARE MEDICAL DEVICES PRIVATE LIMITED  
संपर्क नंबर | Contact No. : 09862782078  
ईमेल आईडी | Email ID : jaimantilifecare@gmail.com  
पता | Address : flat no.-218, Block - F, Narela, North West delhi, DELHI-110040, -  
एमएसएमई पंजीकरण संख्या | MSME Registration number : UDYAM-DL-06-0074970  
एमएसई सामाजिक श्रेणी | MSE Social Category : General  
एमएसई लिंग श्रेणी | MSE Gender : Male  
जीएसटीआईएन | GSTIN: 08AAECJ4541L1ZC (R), (M)

\*जिसके नाम के पक्ष में GST/TAX इनवॉइस पेश किया जाएगा | GST / Tax invoice to be raised in the name of - Buyer

## वितरण निर्देश | Delivery Instructions : NA

## उत्पाद विवरण | Product Details

#	आइटम विवरण   Item Description	आइटम विवरण   Ordered Quantity	इकाई   Unit	इकाई मूल्य (INR)   Unit Price (INR)	कर विभाजन (INR)   Tax Bifurcation (INR)	मूल्य (INR में सभी शुल्क और कर सहित)   Price (Inclusive of all Duties and Taxes in INR)
1	उत्पाद का नाम   Product Name : JML Intravenous Cannulas With injection port and Short bevel cut needle ब्रांड   Brand : JML ब्रांड प्रकार   Brand Type : Registered Brand कैटलॉग की स्थिति   Catalogue Status: OEM verified catalogue कैसे बेचा जा रहा है   Selling As : OEM श्रेणी का नाम और चतुर्थांश   Category Name & Quadrant : Intravenous Cannulas as per IS 10555 - 5 (Q2) मॉडल   Model: JML/PTFE/SBC-20G एचएसएन कोड   HSN Code: HSN not specified by seller	24,000	pieces	5.05	NA	121,200
कुल ऑर्डर मूल्य   Total Order Value (in INR)						121,200

## परेषिती विवरण | Consignee Detail

क्र.सं.   S.No	परेषिती   Consignee	वस्तु   Item	लॉट नंबर   Lot No.	मात्रा   Quantity	दिनांक के बाद डिलीवरी शुरू करना है   Delivery Start After	वितरण पूरा कब तक करना है   Delivery To Be Completed By
	पद   Designation : Store Officer equipment & surgical ईमेल आईडी   Email ID : con33.hfw.delhi@gembuyer.in संपर्क   Contact : -	JML Intravenous Cannulas With injection				

1	जीएसटीआईएन   GSTIN : N पता   Address : Lal Bahadur Shastri Hospital, Khichripur, EAST DELHI, DELHI-110091, India	port and Short bevel cut needle	-	24,000	12-Apr-2024	27-Apr-2024
Product Specification for JML Intravenous Cannulas With injection port and Short bevel cut needle						
विनिर्देश   Specification		उप-विनिर्देश   Sub-Spec			मूल्य   Value	
Performance Parameters	Conforming standards for the I V Canulla			IS / ISO 10555-5		
	Whether IV Canulla is with safety features for preventing needle stick injuries			No		
	Flowrate of inside Catheter in ml/min as per IS/ISO 10555-5			54 ml/min		
	Needle Point Finish			Short bevel cut		
	Needle Hub			Complying with ISO 594-1		
	Needle hub fitting with 6% luer Taper			Yes		
	Injection Port			With		
	Wings			With		
	Luer Lock Plug/Cap			Yes		
	Disposable			Yes		
	Radio opaque line feature			Yes		
	Shelf Life (years)			5		
	Removable Vent Fitting			Yes		
	Sterile			Yes		
	Non-Toxic and Non-Pyrogenic			Yes		
	Provision for Recalling product			yes		
Dimensional and Material Parameters	Material of Needle			Stainless Steel Complying with ISO 9626		
	Material of Catheter			Polytetrafluoroethylene (PTFE)		
	Nominal outside Diameter of catheter tube and color coding as per IS/ISO 10555-5			Pink - 20G		
	Effective Length of Catheter Tube in mm as per IS/ISO 10555-5			32 mm		
PACKING PARAMETERS	Type of packing			blister packing		
	Packing as per specification and provision of Drug & cosmetic act.			yes		
	All supplies shall have a remaining self life at least five by six (5/6) of the stipulated shelf life at the time of delievery			yes		
CERTIFICATE AND REPORTS	Availability of valid drug licence			yes		
	Drug License no & Date of manufacturers and in case of resller, Drug License no & Date (for sale) of Authorized reseller also to be indicated			Drug License no: MFG/MD/2021/000166 & Date: 17 June 2021		
	Manufacturers and the Seller must not be under Conviction in terms of provisions of Drug and Cosmetic act			yes		
	Availability of Latest non conviction certificate issued by concerned Drug authorities			yes		
	Availability of Certificate for Manufacturing such as GMP under revised Schedule-'M' of Drugs & Cosmetics Act 1940 Or WHO-GMP or COPP for imports.			NA		
	Details of above Mentioned Certificate such as Type of certificate, Number, date and validity			NA		
	Availability of any other certification such as CE/FDA/CSA/PQS /ISO etc...			Yes		
	Details of the above mentioned such as Type of certificate, number, date and validity, if Yes otherwise indicate NA			ISO Certificate: ISO13485:2016 Date: 14 APR 2021, Validity: 13 APR 2024		
	Copies of batch in house Test report to be forwarded with each supply			yes		
	Copies of certificates to be provided to buyer on demand after placement of order			yes		
ईपीबीजी विवरण   ePBG Detail						
NA						
नियम और शर्तें   Terms and Conditions						
1. Special terms and conditions- Version:3 effective from 25-11-2022						
1.1	1. All Provisions of Drugs and Cosmetics Act, 1940 and Rules made there under as amended till date will always be applicable. This will include all notifications issued by					

Central Drugs Standard Control Organisation (CDSCO), Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers time to time in this regard.

2. The sellers are registered on GeM based on self-declaration of valid Drug License, product certification, test reports etc. However, buyers must check and validate the details at their end for all applicable licenses and certifications e.g., validity and authenticity/genuineness of drug license, product certification, manufacturer certification/licenses, test reports etc.
3. The price offered by the seller shall not, in any case exceed the DPCO controlled price, if any, fixed by the Central/State Government, the Maximum Retail Price (MRP) and the selling price. The seller must reduce the prices if there is any reduction in DPCO ceiling price, if any.
4. Any other Terms and Conditions which is not included or at variance with the conditions specified in STC/GTC, may be added by the buyer through Additional Terms and Conditions in the bid to ensure items are procured from authentic/validated source with appropriate and applicable quality. The above terms and conditions are in reverse order of precedence i.e., ATC shall supersede specific STC which shall supersede General Terms and Conditions ("GTC"), whenever there are any conflicting provisions.

#### 1.2 Special Terms and Conditions for Medical Devices and Consumables covered under Provisions of Drug and Cosmetic Act

1.2.1 For items wherever Drug Licence requirements are applicable all provisions of Drug and Cosmetic Act 1940 as amended up to date and Rules made there under will be applicable in addition to any other terms and conditions specified in the Portal.

1.2.2 Drug License: For indigenous products offered in the market, Manufacturer should have valid Drug License as per Drugs and Cosmetic Act 1940 issued by concerned State Drug Control authorities. The Seller if different from the manufacturer shall also be required to be holding Drug License for sale. In case of imported products Manufacturer shall be registered under Form no 10 with Central Drug Authorities (CDSCO) and the Seller offering imported products should be also holding valid Sales License issued by the local drug authorities. For imported products, certificate from the OEM that product is being used in the Country of Origin should be available with the Seller. It shall be the responsibility of the Seller to ensure that the Drug License is valid for the product offered and due to any reason the drug control authorities have cancelled or suspended Drug License or convicted the manufacturer or Seller for any offence under the provisions of Drug and Cosmetic Act, Seller should immediately withdraw the product and also intimate the Buyers in case of pending orders for supplies as well as the GeM administration regarding the matter.

1.2.3 Manufacturing & Marketing Experience: Sellers offering the Products in the Portal either as Manufacturers or as Authorized Seller shall ensure that the Products offered are being Manufactured and Marketed in the country (for Indigenous Products) and Marketed (for Imported Products) continuously at least for the last 2 years

1.2.4 Certifications: Manufacturers of offered product (Offered by Manufacturers or by Authorized Seller) should be holding valid Good Manufacturing Practices Certificate (GMP) as per revised Schedule-M of Drug and Cosmetic Act 1940 as amended up to date or WHO-GMP as per norms amended up to date issued by the Licensing Authority or certificate which is at par with WHO-GMP issued by the authorities of exporting countries / COPP certificate.

1.2.5 Non Conviction Certificate: Sellers either Manufacturers or Authorized Sellers are required to ensure that they are not under conviction in terms of the provisions of Drugs & Cosmetic Act and any other law applicable in relation to the same. In case at any point of time, the Manufacturer or Authorized Seller is convicted under provisions of Drug and Cosmetic Act, it shall be their responsibility to withdraw the product immediately from the market.

1.2.6 Banning and Blacklisting: Seller either Manufacturer or Authorized Seller shall ensure that there is no banning or black listing applicable against them for the product offered on the portal due to quality failure and /or fraudulent/illegal practices or for any other reasons

1.2.7 It shall be the responsibility of the Seller either Manufacturer or Authorized Seller to ensure that manufacturer is having own in-house testing lab to carry out all the required tests as per specification and provisions of drug act as amended up to date for the quoted product and shall also forward the copies of the in-house test reports for each batch along with the supplies. For imported products, certificate from OEM regarding availability of all test facilities in house with them should be available with Seller.

1.2.8 Each lot of supplies shall be dispatched under Self Certification scheme duly supported by in house test reports. Consignees shall be at liberty to draw control Samples and send it to approved Laboratories for testing and in case of any failure, entire responsibility shall rest with Seller in addition to any penalties under the provisions of Drug Act including removal of Goods from the Consignee place. Further administrative actions as per terms and conditions Gem Portal shall also be applicable.

1.2.9 Packing shall be as per relevant clause of Standard Specifications applicable as indicated in the Catalogue Parameters indicated in the Portal and as per provisions of Drug & Cosmetic Act as amended up to date.

1.2.10 Expiry Date: All supplies must indicate the Month of Manufacture and Expiry. In addition all supplies shall have a remaining shelf life of at least 5/6th of the stipulated shelf life at the time of delivery.

1.2.11 Recalls: If any batch is to be recalled because of problems with product quality or adverse reaction Seller will be responsible to notify the Buyer full details about the reason leading to the recall and shall take steps to replace the product in question at its own cost with a fresh batch of acceptable quality or give a refund of the value of the goods

#### 2. General Terms and Conditions-

2.1 This contract is governed by the [General Terms and Conditions](#), conditions stipulated to this Product/Service as provided in the Marketplace.

2.2 This Contract between the Seller and the Buyer, is for the supply of the Goods and/ or Services, detailed in the schedule above, in accordance with the General Terms and Conditions (GTC) unless otherwise superseded by Goods / Services specific Special Terms and Conditions (STC) and/ or BID/Reverse Auction Additional Terms and Conditions (ATC), as applicable

नोट: यह सिस्टम जनरेटेड फाइल है। कोई हस्ताक्षर की आवश्यकता नहीं है। इस दस्तावेज़ का प्रिंट आउट भुगतान/लेनदेन उद्देश्य के लिए मान्य नहीं है।

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