

अनुबंध | Contract



अनुबंध क्रमांक | Contract No: GEMC-511687704758784

अनुबंध तिथि | Generated Date : 26-Mar-2025

बोली/आरए/पीबीपी संख्या | Bid/RA/PBP No.: [GEM/2024/B/5544947](#)

अनुसूची नाम | Schedule Name: Schedule 6

संगठन विवरण Organisation Details	खरीदार विवरण Buyer Details
प्रारूप Type : Central PSU मंत्रालय Ministry : Ministry of Coal विभाग Department : COAL INDIA LIMITED संगठन का नाम Organisation Name : Mahanadi Coalfields Limited कार्यालय क्षेत्र Office Zone : MCL Sambalpur Odisha	पद Designation : Shivendra Vyas Asst Manager MM संपर्क नंबर Contact No. : -9549295368- ईमेल आईडी Email ID : mmpur1d.mcl@coalindia.in जीएसटीआईएन GSTIN : - पता Address : MAHANADI COALFIELDS LIMITED, PO - JAGRUTI VIHAR, BURLA, SAMBALPUR, ODISHA-768020, India

वित्तीय स्वीकृति विवरण Financial Approval Detail	भुगतान प्राधिकरण विवरण Paying Authority Details
आईएफडी सहमति IFD Concurrence : No प्रशासनिक अनुमोदन का पदनाम Designation of Administrative Approval : GM (MM)/HoD वित्तीय अनुमोदन का पदनाम Designation of Financial Approval : AF(MM)	Role: PAO भुगतान का तरीका Payment Mode: Offline पद Designation : B Narasimha AM F ईमेल आईडी Email ID : nb.basanaboina@coalindia.in जीएसटीआईएन GSTIN : 21AABCM5188P1Z3 पता Address : MAHANADI COALFIELDS LIMITED, AT /PO - JAGRUTI VIHAR, BURLA, SAMBALPUR, ODISHA-768020, India

विक्रेता विवरण Seller Details
जेम विक्रेता आईडी GeM Seller ID : OL9F210003877965 कंपनी का नाम Company Name : SUNWEST BIO MEDICAL PRIVATE LIMITED संपर्क नंबर Contact No. : 09873914444 ईमेल आईडी Email ID : sunwestbiomedicalpvtltd@gmail.com पता Address : Plot No - 114,,Sector-7, Village/Town:- IMT Manesar, City:- Gurgaon,Sector-7, Village/Town:- IMT Manesar, City:- Gurgaon, HARYANA-122050, India एमएसएमई पंजीकरण संख्या MSME Registration number : UDYAM-HR-05-0025016 जीएसटीआईएन GSTIN: 06ABFCS9428R1ZG (R) , (M)
खरीदार द्वारा सत्यापित एमएसएमई स्थिति MSME Status as verified by buyer : Verified
एमएसई सामाजिक श्रेणी MSE Social Category : General एमएसई लिंग श्रेणी MSE Gender : Male

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

वितरण निर्देश | Delivery Instructions : NA

#	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 : Sunsure H.Pylori Antibody (IgM/IgG/IgA) Rapid Test Kit ब्रांड Brand : Sunsure 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 : Point of Care Rapid Test Kits (V2) (Q2) मॉडल Model: Sunsure H-Pylori Ab RAPID TEST एचएसएन कोड HSN Code: HSN not specified by seller	200	Test	28	NA	5,600
कुल ऑर्डर मूल्य Total Order Value (in INR)						5,600

परोक्षी विवरण Consignee Detail						
क्र.सं. S.No	परोक्षी Consignee	वस्तु Item	लॉट नंबर Lot No.	मात्रा Quantity	दिनांक के बाद डिलीवरी शुरू करना है	वितरण पूरा कब तक करना है Delivery To Be

					Delivery Start After	Completed By
1	<p>पद Designation :- ईमेल आईडी Email ID : kunachandra.polai@nic.in संपर्क Contact : 0663-2542560- जीएसटीआईएन GSTIN : 21AABCM5188P1Z3 पता Address : Anand Vihar Hospital, At/Po: Jagriti Vihar, Burla, Sambalpur-768020, Sambalpur, ODISHA-768020, India</p>	Sunsure H.Pylori Antibody (IgM/IgG/IgA) Rapid Test Kit	-	200	26-Mar-2025	25-Apr-2025
Product Specification for Sunsure H.Pylori Antibody (IgM/IgG/IgA) Rapid Test Kit						
विनिर्देश Specification	उप-विनिर्देश Sub-Spec	मूल्य Value				
GENERAL FEATURES	Product description	Point of Care Rapid Test Kit				
	Purpose	To provide in-vitro diagnosis of different types of infections and diseases				
PRODUCT INFORMATION	Type of Kit	H.Pylori Antibody (IgM/IgG/IgA) Rapid Test Kit				
	Detection Type	Qualitative				
	Differential detection of antibodies and/or antigen	Yes				
	Testing Principle	Lateral Flow Immunochromatographic Assay				
	Species Reactivity	Human				
	Compatible Specimen	Whole Blood, Serum, Plasma				
	Total procedure time	≤ 30 minutes				
	Ability to evaluate negative or positive test result	Yes				
	Assay Sensitivity (%)	≥ 99				
	Assay Specificity (%)	≥ 99				
	Declared sensitivity and specificity shall be claimed by the manufacturer in the kit literature	Yes				
	Contains an internal control dot/band for the confirmation that the test has been performed correctly	Yes				
	The control dot/band able to detect the presence of human immunoglobulin and not be just a procedural control or meant merely for checking the flow of reagents or integrity of antigens except in lateral flow technology	Yes				
	Maintenance of cold chain by the supplier during storage and transportation of Kits at 2°C to 8°C and placement of cumulative time temperature indicator technology on every pack of kits	No				
KIT CONTENTS	Kit contents	Test card/cassette with dessicant, Sample dropper, Assay Buffer (if any)				
	All the components shall be in the quantity as per pack size	Yes				
	Adequate document in English detailing principle, components, methodologies, validity criteria, interpretation of result, performance characteristic, bio-safety, limitation of assay, storage condition, mfg and Exp date & method of disposal provided	Yes				
	Positive and negative controls provided with each pack of kit	No				
	Quantity of positive and negative control	Sufficient for conducting 20% of the tests (10 % negative and 10% positive controls)				
PACKAGING	Pack size of kit	30 Tests				
	The test kit packed in such a way that there is provision to conduct single test at a time	Yes				
	Each test card/cassette with desiccant individually packed in a hermetically sealed and non-permeable pouch	Yes				
CERTIFICATIONS & REPORTS	Compliance to Medical Device Rules (MDR) 2017 as amended till date	Yes				
	Availability of valid drug license for the product issued from the competent authority defined under Drugs and Cosmetic Act 1940 and Rules made there under as amended till date	Yes				
	Valid Drug License Number	MFG/IVD/2023/000034				
	Manufacturing unit certification	ISO:13485 (Latest)				
	Additional voluntary certification available	CE				
	Availability of Test Report for each supplied batch/product as per Medical Device Rules (MDR) 2017 as amended till date	Yes				
	Submission of all necessary certifications, licenses and test reports to the buyer at the time of bid submission or along with supplies as per buyer requirement	Yes				

SHELF LIFE	Shelf life from the date of manufacture	24 month
	Minimum shelf life of the product at the time of delivery to the consignee	3/4 th of total shelf life
ADVANCE SAMPLE	Agree to provide advance sample of the product for buyer's approval before commencement of supply in case of bidding	No
ADDITIONAL REQUIREMENT	Additional Requirement	NA

ईपीबीजी विवरण | ePBG Detail

सलाहकार बैंक Advisory Bank :	NA
ईपीबीजी प्रतिशत (%) ePBG Percentage(%):	NA

नियम और शर्तें | Terms and Conditions

1. Special terms and conditions- Version:1 effective from 12-12-2023

- 1.1
- All Provisions of Drugs and Cosmetics Act, 1940 and Rules (including Medical Device Rule 2017) made there under as amended till date will always be applicable. This will include all notifications issued by Central Drugs Standard Control Organization (CDSCO), Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers time to time in this regard.
 - The sellers are registered on GeM based on the submission of valid Drug License and self declaration of 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.
 - In case of authorized resellers/distributors, it will be the legal & regulatory liability of the manufacturer to ensure that their resellers/distributors are operating in compliance with all relevant laws and regulations and are properly licensed to sell the manufacturer's products, including verifying the validity and authenticity of drug license held by them.
 - The price offered by the seller/bidder shall not, in any case exceed the DPCO/NPPA controlled price or price fixed by State Government, if any. The seller must reduce the prices if there is any reduction in DPCO/NPPA ceiling price or price fixed by State Government, if any.
 - 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 (ATC) 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.

2. General Terms and Conditions-

- This contract is governed by the [General Terms and Conditions](#), conditions stipulated to this Product/Service as provided in the Marketplace.
- 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
- All GeM Sellers / Service Providers are mandated to ensure compliance with all the applicable laws / acts / rules including but not limited to all Labour Laws such as The Minimum Wages Act, 1948, The Payment of Wages Act, 1936, The Payment of Bonus Act, 1965, The Equal Remuneration Act, 1976, The Payment of Gratuity Act, 1972 etc. Any non-compliance will be treated as breach of contract and Buyer may take suitable actions as per GeM Contract.

3. Buyer Added Bid Specific Terms and Conditions-

3.1 Scope of Supply:

Scope of supply (Bid price to include all cost components) : Only supply of Goods

3.2 Generic

OPTION CLAUSE: The Purchaser reserves the right to increase or decrease the quantity to be ordered up to 25 percent of bid quantity at the time of placement of contract. The purchaser also reserves the right to increase the ordered quantity by up to 25% of the contracted quantity during the currency of the contract at the contracted rates. Bidders are bound to accept the orders accordingly.

3.3 Certificates:

Bidder's offer is liable to be rejected if they don't upload any of the certificates / documents sought in the Bid document, ATC and Corrigendum if any.

3.4 Generic

Shelf Life: The Product/Spare parts to be supplied as part of the services must have minimum

3/4

Shelf Life. On the date of supply, minimum

3/4

usable shelf life should be available / balance.

3.5 Certificates:

To be eligible for award of contract, Bidder / OEM must possess following Certificates / Test Reports on the date of bid opening (to be uploaded with bid):

The bidder should provide any one of these BIS/CE/FDA and ISO certification of the manufacturer.

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।

Note: This is system generated file. No signature is required. Print out of this document is not valid for payment/ transaction purpose.

