

अनुबंध | Contract



अनुबंध क्रमांक | Contract No: GEMC-511687762353853

अनुबंध तिथि | Generated Date : 07-Jun-2024

| संगठन विवरण Organisation Details | खरीदार विवरण Buyer Details |
|---|---|
| प्ररूप Type : State Government मंत्रालय Ministry : - विभाग Department : Health & Family Welfare Department Gujarat संगठन का नाम Organisation Name : N/A कार्यालय क्षेत्र Office Zone : Sdh Santrampur | पद Designation : Medical Officer संपर्क नंबर Contact No. : 02675-220046- ईमेल आईडी Email ID : mo-sdh-sant@gujarat.gov.in जीएसटीआईएन GSTIN : 24BRDR03571B1DX Lunavada Santrampur Road Santrampur Dist Mahisagar Pin 389260, पता Address : PANCH MAHALS, GUJARAT-389260, 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 : Superintendent वित्तीय अनुमोदन का पदनाम Designation of Financial Approval : Superintendent | Role: BUYER भुगतान का तरीका Payment Mode: Offline पद Designation : Medical Officer ईमेल आईडी Email ID : mo-sdh-sant@gujarat.gov.in जीएसटीआईएन GSTIN : 24BRDR03571B1DX Lunavada Santrampur Road Santrampur Dist Mahisagar Pin 389260, पता Address : PANCHMAHALS, GUJARAT-389260, India |

| विक्रेता विवरण Seller Details |
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| जेम विक्रेता आईडी GeM Seller ID : 8901180000444795 कंपनी का नाम Company Name : AMAR TRADING COMPANY संपर्क नंबर Contact No. : 09898553986 ईमेल आईडी Email ID : abbasvali@yahoo.com पता Address : Shop no.9/496/A/1,Opp. Nuttan High School,Patel wada, Godhra, Gujarat-389001, - एमएसएमई पंजीकरण संख्या MSME Registration number : - जीएसटीआईएन GSTIN: 24AAZFA0698R1ZE (R) |

*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 : Lab Requirement

| # | 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 : KARWA Syphilis Antibody Rapid Test Kit ब्रांड Brand : KARWA ब्रांड प्रकार Brand Type : Registered Brand कैटलॉग की स्थिति Catalogue Status : OEM verified catalogue कैसे बेचा जा रहा है Selling As : OEM verified Reseller श्रेणी का नाम और चतुर्थांश Category Name & Quadrant : Syphilis Rapid Test Kits (Q2) मॉडल Model: KARWA Syphilis Rapid Test Kit एचएसएन कोड HSN Code: HSN not specified by seller | 500 | Test | 15 | NA | 7,500 |
| कुल ऑर्डर मूल्य Total Order Value (in INR) | | | | | | 7,500 |

| क्र.सं. S.No | परोक्षिती 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 |
|----------------|---|--|--------------------|-------------------|---|--|
| 1 | पद Designation : BUYER-CONSIGNEE ईमेल आईडी Email ID : mo-sdh-sant@gujarat.gov.in संपर्क Contact : 02675-220046- जीएसटीआईएन GSTIN : 24BRDR03571B1DX पता Address : Lunavada Santrampur Road Santrampur Dist Mahisagar | KARWA Syphilis Antibody Rapid Test Kit | - | 500 | 07-Jun-2024 | 22-Jun-2024 |

Product Specification for KARWA Syphilis Antibody Rapid Test Kit

| विनिर्देश Specification | उप-विनिर्देश Sub-Spec | मूल्य Value |
|---------------------------|---|--|
| GENERAL FEATURES | Product Description | Syphilis Rapid Test Kit |
| | Clinical Purpose | To provide a quick and accurate diagnosis of syphilis infection in all stages of infection by detecting antibodies to Treponema Pallidum |
| PRODUCT INFORMATION | Type of Kit | Syphilis Antibody Rapid Test Kit |
| | Detects | Total Anti-Treponema Pallidum Antibody (IgG,IgM & IgA) |
| | Detection Type | Qualitative |
| | The assay shall have solid phase coated with synthetic or recombinant type of Treponema Pallidum antigens | Yes |
| | Testing Principle | Lateral Flow Immunochromatographic Assay |
| | Species Reactivity | Human |
| | Type of Sample | Serum,Plasma |
| | Time to Result | ≤ 30 minutes |
| | Ability to evaluate negative or positive test result | Yes |
| | Sensitivity | ≥99% |
| | Specificity | ≥98% |
| | Declared sensitivity and specificity shall be claimed by the manufacturer in the kit literature | Yes |
| | The assay calibrated to WHO reference serum and the same shall be supported by statements in kit insert and certificate from the manufacturer | 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 |
| KIT CONTENTS | Kit Contents | Test Card/Cassette with Desiccant, Sample Dropper, Assay Buffer (if any) |
| | All the components shall be in the quantity as per pack size | Yes |
| | Adequate document detailing principle, components, methodologies, validity criteria, interpretation of results, performance characteristics, bio-safety, limitations of assay, storage condition, mfg & exp date and method of disposal provided with kit | Yes |
| | Positive and negative controls provided with each pack of kit | No |
| | Quantity of positive and negative controls provided | NA |
| | Each test card/strip supplied with sterile auto retractable disposable lancet and disposable alcoholic swabs | Yes |
| PACKAGING | Pack size of kit | 50 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 with Medical Device Rule (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 there under as amended till date | Yes |
| | Valid Drug License Number | MFG/IVD/2020/B/000004 |
| | Manufacturing unit certification | ISO:13485 (Latest) |
| | Additional voluntary certification available | 70943/B/0001/UK/En |
| | 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 in months from the date of manufacture | 24 |

| | | |
|------------------------|--|----------------------------|
| SHELF LIFE | 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 | Yes |
| ADDITIONAL REQUIREMENT | Additional Requirement | NA |

ईपीबीजी विवरण | ePBG Detail

NA

नियम और शर्तें | Terms and Conditions

1. Special terms and conditions- Version:2 effective from 19-05-2023

1.1

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.
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. 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.
4. 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.
5. 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-

- 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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