

बिड दस्तावेज़ / Bid Document

बिड विवरण/Bid Details	
बिड बंद होने की तारीख/समय /Bid End Date/Time	02-03-2026 11:00:00
बिड खुलने की तारीख/समय /Bid Opening Date/Time	02-03-2026 11:30:00
बिड पेशकश वैधता (बंद होने की तारीख से)/Bid Offer Validity (From End Date)	90 (Days)
मंत्रालय/राज्य का नाम/Ministry/State Name	Ministry Of Labour And Employment
विभाग का नाम/Department Name	Na
संगठन का नाम/Organisation Name	Employees State Insurance Corporation (esic)
कार्यालय का नाम/Office Name	Model Hospital , Rkl
कुल मात्रा/Total Quantity	5750
वस्तु श्रेणी /Item Category	Ayurvedic Classical Medicines - Vati and Gutika (Q1) , Ayurvedic Classical Medicines - Arishta (Q1) , Ayurvedic Classical Medicines - Avaleha and Pak (Q1) , Ayurvedic Classical Medicines - Guggulu (Q1) , Ayurvedic Classical Medicines - Rasa (Q1) , Ayurvedic Classical Medicines - Choorna (Q1) , Ayurvedic Classical Medicines - Taila (Q1)
एमएसएमई के लिए अनुभव के वर्षों और टर्नओवर से छूट प्रदान की गई है/MSE Relaxation for Years of Experience and Turnover	No
स्टार्टअप के लिए अनुभव के वर्षों और टर्नओवर से छूट प्रदान की गई है /Startup Relaxation for Years of Experience and Turnover	No
बिड लगाने की समय सीमा स्वतः नहीं बढ़ाने के लिए आवश्यक बिड की संख्या। / Minimum number of bids required to disable automatic bid extension	1
दिनों की संख्या, जिनके लिए बिड लगाने की समय-सीमा बढ़ाई जाएगी। / Number of days for which Bid would be auto-extended	5
ऑटो एक्सटेंशन अधिकतम कितनी बार किया जाना है। / Number of Auto Extension count	2
बिड से रिवर्स नीलामी सक्रिय किया/Bid to RA enabled	No
बिड का प्रकार/Type of Bid	Single Packet Bid
प्राथमिक उत्पाद श्रेणी/Primary product category	Ayurvedic Classical Medicines - Vati and Gutika

बिड विवरण/Bid Details

तकनीकी मूल्यांकन के दौरान तकनीकी स्पष्टीकरण हेतु अनुमत समय /Time allowed for Technical Clarifications during technical evaluation	5 Days
निरीक्षण आवश्यक (सूचीबद्ध निरीक्षण प्राधिकरण /जेम के साथ पूर्व पंजीकृत एजेंसियों द्वारा)/Inspection Required (By Empanelled Inspection Authority / Agencies pre-registered with GeM)	No
मूल्यांकन पद्धति/Evaluation Method	Total value wise evaluation
मध्यस्थता खंड/Arbitration Clause	No
सुलह खंड/Mediation Clause	No

ईएमडी विवरण/EMD Detail

आवश्यकता/Required	No
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ईपीबीजी विवरण /ePBG Detail

आवश्यकता/Required	No
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बोली विभाजन लागू नहीं किया गया/ Bid splitting not applied.

एमआईआई खरीद वरीयता/MII Purchase Preference

एमआईआई खरीद वरीयता/MII Purchase Preference	No
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एमएसई के लिए आरक्षित / Reserved for MSE

एमएसई के लिए आरक्षित / Reserved for MSE	Yes
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1. Procurement under this bid is reserved for purchase from Micro and Small Enterprises having valid Udyam Registration and whose credentials are validated online through Udyam Registration portal. If the bidder wants to avail themselves of the reservation benefit, the bidder must be the manufacturer / OEM of the offered product on GeM. Traders are excluded from the purview of Public Procurement Policy for Micro and Small Enterprises and hence resellers offering products manufactured by some other OEM are not eligible to participate in this bid. In respect of bid for Services, the bidder must be the Service provider of the offered Service. Relevant documentary evidence in this regard shall be uploaded along with the bid in respect of the offered product or service, and Buyer will decide eligibility based on documentary evidence submitted, while evaluating the bid. Benefits of MSE will be allowed only if seller is validated on-line in GeM profile as well as validated and approved by Buyer after evaluation of documents submitted.

Ayurvedic Classical Medicines - Vati And Gutika (250 pieces)**तकनीकी विशिष्टियाँ /Technical Specifications**

* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Arogyavardhini Gutika
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	10 Grams
	Number of container/bottle in a pack (Secondary Packing)	50
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Vati And Gutika (250 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Chandraprabha Vati
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	10 Grams
	Number of container/bottle in a pack (Secondary Packing)	50
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	3, 5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Arishta (250 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Abhayarishta
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	200 ml
	Number of container/bottle in a pack (Secondary Packing)	12
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Avaleha And Pak (250 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

[* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification](#)

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Haridra Khanda
	Medicine form	Granule
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	100 Grams
	Number of container/bottle in a pack (Secondary Packing)	50
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	3, 5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Guggulu (250 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

[* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification](#)

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Singhnad Guggulu
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	20 Grams
	Number of container/bottle in a pack (Secondary Packing)	25
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.N o.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Rasa (250 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

[* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification](#)

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Ekangavira Rasa
	Medicine form	Tablet
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	5 Grams
	Number of container/bottle in a pack (Secondary Packing)	25
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Choorna (1000 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Ashwagandha Choorna
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	50 Grams
	Number of container/bottle in a pack (Secondary Packing)	50
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	2, 3, 5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	1000	30

Ayurvedic Classical Medicines - Rasa (500 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

[* जेम कैटेगरी विशिष्टि के अनुसार / As per GeM Category Specification](#)

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Lakshmvilasa Rasa (Nardiya)
	Medicine form	Tablet
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	5 Grams
	Number of container/bottle in a pack (Secondary Packing)	50
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	3, 5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	500	30

Ayurvedic Classical Medicines - Choorna (500 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

[* जेम कैटेगरी विशिष्टि के अनुसार / As per GeM Category Specification](#)

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Avipattikar Choorna
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	50 Grams
	Number of container/bottle in a pack (Secondary Packing)	50
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	2, 3, 5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.N o.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	500	30

Ayurvedic Classical Medicines - Taila (250 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Br. Marichyadi Taila
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	50 ml
	Number of container/bottle in a pack (Secondary Packing)	25
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	3, 5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Choorna (250 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

* जेम केटेगरी विशिष्टि के अनुसार / As per GeM Category Specification

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Triphala Choorna
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	50 Grams
	Number of container/bottle in a pack (Secondary Packing)	50
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	2, 3, 5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.N o.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Guggulu (250 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

[* जेम कैटेगरी विशिष्टि के अनुसार / As per GeM Category Specification](#)

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Kaishore Guggulu
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	20 Grams
	Number of container/bottle in a pack (Secondary Packing)	25
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.N o.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	250	30

Ayurvedic Classical Medicines - Taila (1500 pieces)

तकनीकी विशिष्टियाँ /Technical Specifications

[* जेम कैटेगरी विशिष्टि के अनुसार / As per GeM Category Specification](#)

विवरण/Specification	विशिष्टि का नाम /Specification Name	बिड के लिए आवश्यक अनुमत मूल्य /Bid Requirement (Allowed Values)
GENERAL FEATURES	Medicine name	Panchaguna Taila
PACKAGING & LABELLING	Quantity of medicine in one container/bottle (Unit pack size) (A/U) (Primary packing)	50 ml
	Number of container/bottle in a pack (Secondary Packing)	25
	Additional marking requirement on the label and other cartons	GOVT. SUPPLY, NOT FOR SALE
SHELF LIFE	Shelf life from the date of manufacture as per Drugs and Cosmetic Act (Years)	3, 5, 10 Or higher (year)

परेषिती/रिपोर्टिंग अधिकारी तथा मात्रा/Consignees/Reporting Officer and Quantity

क्र.सं./S.N o.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Nawal Kishor Murmu	769012,MEDICAL SUPERINTENDENT ESIC MODEL HOSPITAL ,NEAR GOVT. ITI JAIL ROAD ROURKELA-4	1500	30

Special terms and conditions-Version:3 effective from 14-03-2024 for category Ayurvedic Classical Medicines - Vati and Gutika

1.

- All Provision of Drugs & Cosmetic Act 1940 as amended till date and rules made there under and all rules and regulations issued by Ministry of AYUSH will always be applicable.
- Only CPSEs/State PSUs/State Pharmacies/Co-operatives are allowed to upload their products for sale based on authorization letter of these manufacturers from State Ayush Department/National Ayush Mission at the time of vendor assessment process.
- The sellers are allowed to register on GeM and exempted from the Vendor Assessment process based on the submitted copy of a valid Manufacturing Drug License for the medicine(s) certified by the issuing authority. Buyers must mandatorily ask for submitting the relevant valid drug license and other regulatory documents applicable with the bid. Buyers must also check and validate the details e.g., validity, authenticity/genuineness, name of the drug/medicine under procurement, the license issuing authority etc. at their end.
- The purchase shall be made through Bidding/RA only irrespective of the value.
- Manufacturer shall have a valid own manufacturing license issued by the competent drug licensing authority defined under the Drugs and Cosmetics Act 1940 and Rules made there under as amended till date. (If revalidation of drug license has been applied for, the buyer shall be informed accordingly and the copy of application to State Drug / Licensing authority must be submitted with a certificate that application for renewal was made within time frame as per Drug and Cosmetic Act as amended up to date and that has not been deleted by drug licensing authority.)
- Loan license arrangement shall not be allowed under any circumstances.

7. Medicines must fully comply in all respect with the technical specifications and in accordance with the Pharmacopoeia standards wherever applicable.
8. Each batch of the medicines shall be got tested by the seller from the laboratories approved by State Drug controllers / run by State Government/NABL accredited lab/Government Approved Lab and shall be dispatched along with these test reports.
9. INSPECTION & QUALITY TESTING

a) Medicines shall continue to conform to the description and the quality during the shelf life from the date of delivery of the medicines to the buyer and notwithstanding the fact that the buyer may have inspected and or approved the said medicines.

b) The buyer has the right to inspect, test and where necessary reject the medicines after arrival at the final destination shall in no way be limited or waived by reason of the medicines having previously been inspected, tested and passed by the buyer or his authorized representative prior to the medicines dispatch from the place of manufacture or arrival as the case may be.

c) If any inspected or tested medicines fail to conform to the specifications, the buyer may reject them and the seller will remove the rejected medicines at its own cost.

d) During the shelf-life period, the consignees shall be at liberty to draw samples and send it to laboratories approved by State Drugs controllers / run by State Government/NABL approved laboratories/ Government Approved Lab without any intimation to the seller. If found "Not of Standard Quality", (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality upon testing. The seller shall have to replace the rejected batches (unused quantity) with fresh batches within 3 months or refund the cost of the rejected medicines to the buyer, if so, decided by him. In the event of replacement of rejected medicines by the seller, all the above mentioned provision shall apply to the new medicines replaced from the date of replacement thereof, otherwise the seller shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained and the facts will be notified to the Drugs Controller of India/State Drug Controller for taking necessary action.

e) In case any medicines are found substandard either at the inspection stage or during the shelf life of the medicines, the report of the Government approved/NABL accredited laboratory shall be accepted by the seller. If the same is disputed by the seller giving the reasons, the sample will be sent to the designated appellate Lab (Pharmacopoeia commission for Indian medicine & Homeopathy) for the purpose and the report of the same will only be accepted as final and conclusive report. De-registration / debarment action will be taken against the seller according to the category-A and category-B defects as per guidelines issued by the Ministry of Health & Family Welfare.

f) The cost of post-delivery inspection and testing will be borne by the buyer. However, inspection & testing charges for the failed batches shall be borne by the seller.

g) In the event of the samples of medicines supplied fails in quality tests or found to be not as per specifications, the buyer will send second sample to the 2nd Govt. approved /NABL accredited lab. If the second sample fails, the batch will be rejected but if the second sample passes then third sample will be sent to the designated Appellate lab for the medicines and decision of the Appellate lab will be final.

h) In the event of the samples of medicine supplied finally fail in quality tests or found to be not as per specifications, the seller will have to replace the rejected batch with fresh stock duly inspected within 3 months. If not replaced, the buyer will be at liberty to purchase from other source and recovery to be made from the seller and action to blacklist the company/cancellation of the Drug license will also be initiated.

10. Warranty

Each supply shall be accompanied with a "Warranty Certificate" as specified below, duly signed by the Seller as under.

WARRANTY CERTIFICATE:

I/We, _____ (name of the seller), hereby declare that the medicines sold to the _____ (name of the buyer) under this supply order (No. of the supply order with date) are of the best quality (and workmanship) and strictly in accordance with specification and particulars

mentioned and I/we hereby guarantee that the said medicines would continue to conform to the description and the quality for a period as specified in the Gazette of India No. 605, dated 20/10/2009 & 16-08-2016 from the date of delivery of the said medicines to the buyer and that notwithstanding the fact that the buyer may have inspected and or approved the said medicines, if during the aforesaid period, discovered not conforming to the description and quality aforesaid or have deteriorated (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality. On such rejection, the medicines will be at the seller's risk and all provisions herein contained relating to the rejection of medicines etc., shall apply. In the event of replacement of rejected medicines by the seller, the above-mentioned guarantee period shall as apply to the medicines replaced from the date of replacement thereof, otherwise the contractor shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained. Nothing here in contained shall prejudice and other right of the buyer in that behalf under this supply order or otherwise.

(Signature name & designation and date with rubber stamp)

11. The classification of defects into different categories will as per guidelines issued by the Drugs Controller of India and action will be taken accordingly.
12. If the seller, having been notified, fails to replace rejected medicines with fresh medicines within 3 months, the buyer may proceed to take such remedial action as may be necessary at the seller's risk and expense and without prejudice to other rights which the buyer may have against the seller under the contract.
13. Loss or premature deterioration due to biological and other activities during the life potency of the medicines shall have to be made good by the seller free of cost or shall have to refund the cost of rejected medicine.
14. Recalls- If medicines must be recalled because of problems with medicines, the seller will be obliged to notify the buyer, providing full details about the reason leading to the recall, and shall take steps to replace the medicines in question at seller's own cost at the ultimate destination with a fresh batch of acceptable medicine or withdraw and give a full refund if the medicine has been taken off the market due to safety issues.
15. Maximum lead time will be 60 days from date of receipt of order and delivery will commence there after.
16. It is the responsibility of the seller to intimate Government e-Marketplace (GEM) about any quality complaints of the medicines reported by any buyer/consignee.
17. Order should be placed for the quantities in multiples of the primary packing.
18. The seller shall not be blacklisted / debarred / banned by any State Governments U.T. / Central Government/Corporations/Local Government Bodies in the preceding 3 years.
19. Seller shall not sell the product(s) for which the firm / company has been blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies due to quality failure of the medicines.
20. During the period of contract if the firm / Company is blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to buyer along with relevant authentic document by seller within one month.
21. Each supply of medicines shall be accompanied with batchwise quality analysis report from government approved /NABL accredited laboratory. This report shall contain specific tests for Identity, Purity, Quality and Strength of the ingredients used in the medicine as per Ayurvedic Pharmacopoeia in case of Ayurvedic medicines and Unani Pharmacopoeia in case of Unani medicines.
22. Packing and Marking
 - a) All containers meant for packing is required to be secured with pilfer proof seal to ensure genuineness of the product packed. With each consignment the seller should give an undertaking that material used is of food grade / HDPE material if supplied in plastic bottle.
 - b) For secondary packing, material is required to be corrugated boxes having "A" grade paper i.e. Virgin, and packed in first-hand boxes only, with suitable flute, joint, stitching, flap, tape. The box should be of 5 ply with bursting strength of 9Kg / cm²
 - c) Weight/volume of the medicines to be mentioned on the inner packing. Weight & other technical requirements shall adhere to as per the pharmacopoeia standard applicable i.e. A.F.I. in case of Ayurveda and N.F.U.M. in case of Unani medicines.
23. 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 drugs/medicines 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 Special Terms and Conditions (STC) which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

Special terms and conditions-Version:4 effective from 14-03-2024 for category Ayurvedic Classical Medicines - Arishta

1.

1. All Provision of Drugs & Cosmetic Act 1940 as amended till date and rules made there under and all rules and regulations issued by Ministry of AYUSH will always be applicable.
2. Only CPSEs/State PSUs/State Pharmacies/Co-operatives are allowed to upload their products for sale based on authorization letter of these manufacturers from State Ayush Department/National Ayush Mission at the time of vendor assessment process.
3. The sellers are allowed to register on GeM and exempted from the Vendor Assessment process based on the submitted copy of a valid Manufacturing Drug License for the medicine(s) certified by the issuing authority. Buyers must mandatorily ask for submitting the relevant valid drug license and other regulatory documents applicable with the bid. Buyers must also check and validate the details e.g., validity, authenticity/genuineness, name of the drug/medicine under procurement, the license issuing authority etc. at their end.
4. The purchase shall be made through Bidding/RA only irrespective of the value.
5. Manufacturer shall have a valid own manufacturing license issued by the competent drug licensing authority defined under the Drugs and Cosmetics Act 1940 and Rules made there under as amended till date. (If revalidation of drug license has been applied for, the buyer shall be informed accordingly and the copy of application to State Drug / Licensing authority must be submitted with a certificate that application for renewal was made within time frame as per Drug and Cosmetic Act as amended up to date and that has not been deleted by drug licensing authority.)
6. Loan license arrangement shall not be allowed under any circumstances.
7. Medicines must fully comply in all respect with the technical specifications and in accordance with the Pharmacopoeia standards wherever applicable.
8. Each batch of the medicines shall be got tested by the seller from the laboratories approved by State Drug controllers / run by State Government/NABL accredited lab/Government Approved Lab and shall be dispatched along with these test reports.
9. INSPECTION & QUALITY TESTING

a) Medicines shall continue to conform to the description and the quality during the shelf life from the date of delivery of the medicines to the buyer and notwithstanding the fact that the buyer may have inspected and or approved the said medicines.

b) The buyer has the right to inspect, test and where necessary reject the medicines after arrival at the final destination shall in no way be limited or waived by reason of the medicines having previously been inspected, tested and passed by the buyer or his authorized representative prior to the medicines dispatch from the place of manufacture or arrival as the case may be.

c) If any inspected or tested medicines fail to conform to the specifications, the buyer may reject them and the seller will remove the rejected medicines at its own cost.

d) During the shelf-life period, the consignees shall be at liberty to draw samples and send it to laboratories approved by State Drugs controllers / run by State Government/NABL approved laboratories/ Government Approved Lab without any intimation to the seller. If found "Not of Standard Quality", (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality upon testing. The seller shall have to replace the rejected batches (unused quantity) with fresh batches within 3 months or refund the cost of the rejected medicines to the buyer, if so, decided by him. In the event of replacement of rejected medicines by the seller, all the above mentioned provision shall apply to the new medicines replaced from the date of replacement thereof, otherwise the seller shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained and the facts will be notified to the Drugs Controller of India/State Drug Controller for taking necessary action.

e) In case any medicines are found substandard either at the inspection stage or during the shelf life of the medicines, the report of the Government approved/NABL accredited laboratory shall be accepted by the seller. If the same is disputed by the seller giving the reasons, the sample will be sent to the designated appellate Lab (Pharmacopoeia commission for Indian medicine & Homeopathy) for the purpose and the report of the same will only be accepted as final and

conclusive report. De-registration / debarment action will be taken against the seller according to the category-A and category-B defects as per guidelines issued by the Ministry of Health & Family Welfare.

f) The cost of post-delivery inspection and testing will be borne by the buyer. However, inspection & testing charges for the failed batches shall be borne by the seller.

g) In the event of the samples of medicines supplied fails in quality tests or found to be not as per specifications, the buyer will send second sample to the 2nd Govt. approved /NABL accredited lab. If the second sample fails, the batch will be rejected but if the second sample passes then third sample will be sent to the designated Appellate lab for the medicines and decision of the Appellate lab will be final.

h) In the event of the samples of medicine supplied finally fail in quality tests or found to be not as per specifications, the seller will have to replace the rejected batch with fresh stock duly inspected within 3 months. If not replaced, the buyer will be at liberty to purchase from other source and recovery to be made from the seller and action to blacklist the company/cancellation of the Drug license will also be initiated.

10. Warranty

Each supply shall be accompanied with a "Warranty Certificate" as specified below, duly signed by the Seller as under.

WARRANTY CERTIFICATE:

I/We, _____ (name of the seller), hereby declare that the medicines sold to the _____ (name of the buyer) under this supply order (No. of the supply order with date) are of the best quality (and workmanship) and strictly in accordance with specification and particulars mentioned and I/we hereby guarantee that the said medicines would continue to conform to the description and the quality for a period as specified in the Gazette of India No. 605, dated 20/10/2009 & 16-08-2016 from the date of delivery of the said medicines to the buyer and that notwithstanding the fact that the buyer may have inspected and or approved the said medicines, if during the aforesaid period, discovered not conforming to the description and quality aforesaid or have deteriorated (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality. On such rejection, the medicines will be at the seller's risk and all provisions herein contained relating to the rejection of medicines etc., shall apply. In the event of replacement of rejected medicines by the seller, the above-mentioned guarantee period shall as apply to the medicines replaced from the date of replacement thereof, otherwise the contractor shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained. Nothing here in contained shall prejudice and other right of the buyer in that behalf under this supply order or otherwise.

(Signature name & designation and date with rubber stamp)

11. The classification of defects into different categories will as per guidelines issued by the Drugs Controller of India and action will be taken accordingly.
12. If the seller, having been notified, fails to replace rejected medicines with fresh medicines within 3 months, the buyer may proceed to take such remedial action as may be necessary at the seller's risk and expense and without prejudice to other rights which the buyer may have against the seller under the contract.
13. Loss or premature deterioration due to biological and other activities during the life potency of the medicines shall have to be made good by the seller free of cost or shall have to refund the cost of rejected medicine.
14. Recalls- If medicines must be recalled because of problems with medicines, the seller will be obliged to notify the buyer, providing full details about the reason leading to the recall, and shall take steps to replace the medicines in question at seller's own cost at the ultimate destination with a fresh batch of acceptable medicine or withdraw and give a full refund if the medicine has been taken off the market due to safety issues.
15. Maximum lead time will be 60 days from date of receipt of order and delivery will commence there after.
16. It is the responsibility of the seller to intimate Government e-Marketplace (GEM) about any quality complaints of the medicines reported by any buyer/consignee.

17. Order should be placed for the quantities in multiples of the primary packing.
18. The seller shall not be blacklisted / debarred / banned by any State Governments U.T. / Central Government/Corporations/Local Government Bodies in the preceding 3 years.
19. Seller shall not sell the product(s) for which the firm / company has been blacklisted/debarred/de-registered/banned by any State Government / Central Government / its Drug procurement agencies due to quality failure of the medicines.
20. During the period of contract if the firm / Company is blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to buyer along with relevant authentic document by seller within one month.
21. Each supply of medicines shall be accompanied with batchwise quality analysis report from government approved /NABL accredited laboratory. This report shall contain specific tests for Identity, Purity, Quality and Strength of the ingredients used in the medicine as per Ayurvedic Pharmacopoeia in case of Ayurvedic medicines and Unani Pharmacopoeia in case of Unani medicines.
22. Packing and Marking
 - a) All containers meant for packing is required to be secured with pilfer proof seal to ensure genuineness of the product packed. With each consignment the seller should give an undertaking that material used is of food grade / HDPE material if supplied in plastic bottle.
 - b) For secondary packing, material is required to be corrugated boxes having "A" grade paper i.e. Virgin, and packed in first-hand boxes only, with suitable flute, joint, stitching, flap, tape. The box should be of 5 ply with bursting strength of 9Kg / cm²
 - c) Weight/volume of the medicines to be mentioned on the inner packing. Weight & other technical requirements shall adhere to as per the pharmacopoeia standard applicable i.e. A.F.I. in case of Ayurveda and N.F.U.M. in case of Unani medicines.
23. 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 drugs/medicines 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 Special Terms and Conditions (STC) which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

Special terms and conditions-Version:4 effective from 14-03-2024 for category Ayurvedic Classical Medicines - Avaleha and Pak

1.

1. All Provision of Drugs & Cosmetic Act 1940 as amended till date and rules made there under and all rules and regulations issued by Ministry of AYUSH will always be applicable.
2. Only CPSEs/State PSUs/State Pharmacies/Co-operatives are allowed to upload their products for sale based on authorization letter of these manufacturers from State Ayush Department/National Ayush Mission at the time of vendor assessment process.
3. The sellers are allowed to register on GeM and exempted from the Vendor Assessment process based on the submitted copy of a valid Manufacturing Drug License for the medicine(s) certified by the issuing authority. Buyers must mandatorily ask for submitting the relevant valid drug license and other regulatory documents applicable with the bid. Buyers must also check and validate the details e.g., validity, authenticity/genuineness, name of the drug/medicine under procurement, the license issuing authority etc. at their end.
4. The purchase shall be made through Bidding/RA only irrespective of the value.
5. Manufacturer shall have a valid own manufacturing license issued by the competent drug licensing authority defined under the Drugs and Cosmetics Act 1940 and Rules made there under as amended till date. (If revalidation of drug license has been applied for, the buyer shall be informed accordingly and the copy of application to State Drug / Licensing authority must be submitted with a certificate that application for renewal was made within time frame as per Drug and Cosmetic Act as amended up to date and that has not been deleted by drug licensing authority.)
6. Loan license arrangement shall not be allowed under any circumstances.
7. Medicines must fully comply in all respect with the technical specifications and in accordance with the Pharmacopoeia standards wherever applicable.
8. Each batch of the medicines shall be got tested by the seller from the laboratories approved by State Drug controllers / run by State Government/NABL accredited lab/Government Approved Lab and shall be dispatched along with these test reports.
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the date of delivery of the medicines to the buyer and notwithstanding the fact that the buyer may have inspected and or approved the said medicines.

b) The buyer has the right to inspect, test and where necessary reject the medicines after arrival at the final destination shall in no way be limited or waived by reason of the medicines having previously been inspected, tested and passed by the buyer or his authorized representative prior to the medicines dispatch from the place of manufacture or arrival as the case may be.

c) If any inspected or tested medicines fail to conform to the specifications, the buyer may reject them and the seller will remove the rejected medicines at its own cost.

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conforming to the said description and quality. On such rejection, the medicines will be at the seller's risk and all provisions herein contained relating to the rejection of medicines etc., shall apply. In the event of replacement of rejected medicines by the seller, the above-mentioned guarantee period shall as apply to the medicines replaced from the date of replacement thereof, otherwise the contractor shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained. Nothing here in contained shall prejudice and other right of the buyer in that behalf under this supply order or otherwise.

(Signature name & designation and date with rubber stamp)

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15. Maximum lead time will be 60 days from date of receipt of order and delivery will commence there after.
16. It is the responsibility of the seller to intimate Government e-Marketplace (GEM) about any quality complaints of the medicines reported by any buyer/consignee.
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19. Seller shall not sell the product(s) for which the firm / company has been blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies due to quality failure of the medicines.
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 - a) All containers meant for packing is required to be secured with pilfer proof seal to ensure genuineness of the product packed. With each consignment the seller should give an undertaking that material used is of food grade / HDPE material if supplied in plastic bottle.
 - b) For secondary packing, material is required to be corrugated boxes having "A" grade paper i.e. Virgin, and packed in first-hand boxes only, with suitable flute, joint, stitching, flap, tape. The box should be of 5 ply with bursting strength of 9Kg / cm²
 - c) Weight/volume of the medicines to be mentioned on the inner packing. Weight & other technical requirements shall adhere to as per the pharmacopoeia standard applicable i.e. A.F.I. in case of Ayurveda and N.F.U.M. in case of Unani medicines.
23. 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 drugs/medicines 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 Special Terms and Conditions (STC) which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

Special terms and conditions-Version:3 effective from 14-03-2024 for category Ayurvedic Classical Medicines - Guggulu

1.

1. All Provision of Drugs & Cosmetic Act 1940 as amended till date and rules made there under and all rules and regulations issued by Ministry of AYUSH will always be applicable.
2. Only CPSEs/State PSUs/State Pharmacies/Co-operatives are allowed to upload their products for sale based on authorization letter of these manufacturers from State Ayush Department/National Ayush Mission at the time of vendor assessment process.
3. The sellers are allowed to register on GeM and exempted from the Vendor Assessment process based on the submitted copy of a valid Manufacturing Drug License for the medicine(s) certified by the issuing authority. Buyers must mandatorily ask for submitting the relevant valid drug license and other regulatory documents applicable with the bid. Buyers must also check and validate the details e.g., validity, authenticity/genuineness, name of the drug/medicine under procurement, the license issuing authority etc. at their end.
4. The purchase shall be made through Bidding/RA only irrespective of the value.
5. Manufacturer shall have a valid own manufacturing license issued by the competent drug licensing authority defined under the Drugs and Cosmetics Act 1940 and Rules made there under as amended till date. (If revalidation of drug license has been applied for, the buyer shall be informed accordingly and the copy of application to State Drug / Licensing authority must be submitted with a certificate that application for renewal was made within time frame as per Drug and Cosmetic Act as amended up to date and that has not been deleted by drug licensing authority.)
6. Loan license arrangement shall not be allowed under any circumstances.
7. Medicines must fully comply in all respect with the technical specifications and in accordance with the Pharmacopoeia standards wherever applicable.
8. Each batch of the medicines shall be got tested by the seller from the laboratories approved by State Drug controllers / run by State Government/NABL accredited lab/Government Approved Lab and shall be dispatched along with these test reports.
9. INSPECTION & QUALITY TESTING

a) Medicines shall continue to conform to the description and the quality during the shelf life from the date of delivery of the medicines to the buyer and notwithstanding the fact that the buyer may have inspected and or approved the said medicines.

b) The buyer has the right to inspect, test and where necessary reject the medicines after arrival at the final destination shall in no way be limited or waived by reason of the medicines having previously been inspected, tested and passed by the buyer or his authorized representative prior to the medicines dispatch from the place of manufacture or arrival as the case may be.

c) If any inspected or tested medicines fail to conform to the specifications, the buyer may reject them and the seller will remove the rejected medicines at its own cost.

d) During the shelf-life period, the consignees shall be at liberty to draw samples and send it to laboratories approved by State Drugs controllers / run by State Government/NABL approved laboratories/ Government Approved Lab without any intimation to the seller. If found "Not of Standard Quality", (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality upon testing. The seller shall have to replace the rejected batches (unused quantity) with fresh batches within 3 months or refund the cost of the rejected medicines to the buyer, if so, decided by him. In the event of replacement of rejected medicines by the seller, all the above mentioned provision shall apply to the new medicines replaced from the date of replacement thereof, otherwise the seller shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained and the facts will be notified to the Drugs Controller of India/State Drug Controller for taking necessary action.

e) In case any medicines are found substandard either at the inspection stage or during the shelf life of the medicines, the report of the Government approved/NABL accredited laboratory shall be accepted by the seller. If the same is disputed by the seller giving the reasons, the sample will be sent to the designated appellate Lab (Pharmacopoeia commission for Indian medicine & Homeopathy) for the purpose and the report of the same will only be accepted as final and conclusive report. De-registration / debarment action will be taken against the seller according to the category-A and category-B defects as per guidelines issued by the Ministry of Health & Family Welfare.

f) The cost of post-delivery inspection and testing will be borne by the buyer. However, inspection & testing charges for the failed batches shall be borne by the seller.

g) In the event of the samples of medicines supplied fails in quality tests or found to be not as per

specifications, the buyer will send second sample to the 2nd Govt. approved /NABL accredited lab. If the second sample fails, the batch will be rejected but if the second sample passes then third sample will be sent to the designated Appellate lab for the medicines and decision of the Appellate lab will be final.

h) In the event of the samples of medicine supplied finally fail in quality tests or found to be not as per specifications, the seller will have to replace the rejected batch with fresh stock duly inspected within 3 months. If not replaced, the buyer will be at liberty to purchase from other source and recovery to be made from the seller and action to blacklist the company/cancellation of the Drug license will also be initiated.

10. Warranty

Each supply shall be accompanied with a "Warranty Certificate" as specified below, duly signed by the Seller as under.

WARRANTY CERTIFICATE:

I/We, _____ (name of the seller), hereby declare that the medicines sold to the _____ (name of the buyer) under this supply order (No. of the supply order with date) are of the best quality (and workmanship) and strictly in accordance with specification and particulars mentioned and I/we hereby guarantee that the said medicines would continue to conform to the description and the quality for a period as specified in the Gazette of India No. 605, dated 20/10/2009 & 16-08-2016 from the date of delivery of the said medicines to the buyer and that notwithstanding the fact that the buyer may have inspected and or approved the said medicines, if during the aforesaid period, discovered not conforming to the description and quality aforesaid or have deteriorated (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality. On such rejection, the medicines will be at the seller's risk and all provisions herein contained relating to the rejection of medicines etc., shall apply. In the event of replacement of rejected medicines by the seller, the above-mentioned guarantee period shall as apply to the medicines replaced from the date of replacement thereof, otherwise the contractor shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained. Nothing here in contained shall prejudice and other right of the buyer in that behalf under this supply order or otherwise.

(Signature name & designation and date with rubber stamp)

11. The classification of defects into different categories will as per guidelines issued by the Drugs Controller of India and action will be taken accordingly.
12. If the seller, having been notified, fails to replace rejected medicines with fresh medicines within 3 months, the buyer may proceed to take such remedial action as may be necessary at the seller's risk and expense and without prejudice to other rights which the buyer may have against the seller under the contract.
13. Loss or premature deterioration due to biological and other activities during the life potency of the medicines shall have to be made good by the seller free of cost or shall have to refund the cost of rejected medicine.
14. Recalls- If medicines must be recalled because of problems with medicines, the seller will be obliged to notify the buyer, providing full details about the reason leading to the recall, and shall take steps to replace the medicines in question at seller's own cost at the ultimate destination with a fresh batch of acceptable medicine or withdraw and give a full refund if the medicine has been taken off the market due to safety issues.
15. Maximum lead time will be 60 days from date of receipt of order and delivery will commence there after.
16. It is the responsibility of the seller to intimate Government e-Marketplace (GEM) about any quality complaints of the medicines reported by any buyer/consignee.
17. Order should be placed for the quantities in multiples of the primary packing.
18. The seller shall not be blacklisted / debarred / banned by any State Governments U.T. / Central Government/Corporations/Local Government Bodies in the preceding 3 years.
19. Seller shall not sell the product(s) for which the firm / company has been blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies due to quality failure of the medicines.
20. During the period of contract if the firm / Company is blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies / convicted by any

Court of law in India, it shall be intimated to buyer along with relevant authentic document by seller within one month.

21. Each supply of medicines shall be accompanied with batchwise quality analysis report from government approved /NABL accredited laboratory. This report shall contain specific tests for Identity, Purity, Quality and Strength of the ingredients used in the medicine as per Ayurvedic Pharmacopoeia in case of Ayurvedic medicines and Unani Pharmacopoeia in case of Unani medicines.
22. Packing and Marking
 - a) All containers meant for packing is required to be secured with pilfer proof seal to ensure genuineness of the product packed. With each consignment the seller should give an undertaking that material used is of food grade / HDPE material if supplied in plastic bottle.
 - b) For secondary packing, material is required to be corrugated boxes having "A" grade paper i.e. Virgin, and packed in first-hand boxes only, with suitable flute, joint, stitching, flap, tape. The box should be of 5 ply with bursting strength of 9Kg / cm²
 - c) Weight/volume of the medicines to be mentioned on the inner packing. Weight & other technical requirements shall adhere to as per the pharmacopoeia standard applicable i.e. A.F.I. in case of Ayurveda and N.F.U.M. in case of Unani medicines.
23. 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 drugs/medicines 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 Special Terms and Conditions (STC) which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

Special terms and conditions-Version:4 effective from 14-03-2024 for category Ayurvedic Classical Medicines - Rasa

1.

1. All Provision of Drugs & Cosmetic Act 1940 as amended till date and rules made there under and all rules and regulations issued by Ministry of AYUSH will always be applicable.
2. Only CPSEs/State PSUs/State Pharmacies/Co-operatives are allowed to upload their products for sale based on authorization letter of these manufacturers from State Ayush Department/National Ayush Mission at the time of vendor assessment process.
3. The sellers are allowed to register on GeM and exempted from the Vendor Assessment process based on the submitted copy of a valid Manufacturing Drug License for the medicine(s) certified by the issuing authority. Buyers must mandatorily ask for submitting the relevant valid drug license and other regulatory documents applicable with the bid. Buyers must also check and validate the details e.g., validity, authenticity/genuineness, name of the drug/medicine under procurement, the license issuing authority etc. at their end.
4. The purchase shall be made through Bidding/RA only irrespective of the value.
5. Manufacturer shall have a valid own manufacturing license issued by the competent drug licensing authority defined under the Drugs and Cosmetics Act 1940 and Rules made there under as amended till date. (If revalidation of drug license has been applied for, the buyer shall be informed accordingly and the copy of application to State Drug / Licensing authority must be submitted with a certificate that application for renewal was made within time frame as per Drug and Cosmetic Act as amended up to date and that has not been deleted by drug licensing authority.)
6. Loan license arrangement shall not be allowed under any circumstances.
7. Medicines must fully comply in all respect with the technical specifications and in accordance with the Pharmacopoeia standards wherever applicable.
8. Each batch of the medicines shall be got tested by the seller from the laboratories approved by State Drug controllers / run by State Government/NABL accredited lab/Government Approved Lab and shall be dispatched along with these test reports.
9. INSPECTION & QUALITY TESTING
 - a) Medicines shall continue to conform to the description and the quality during the shelf life from the date of delivery of the medicines to the buyer and notwithstanding the fact that the buyer may have inspected and or approved the said medicines.
 - b) The buyer has the right to inspect, test and where necessary reject the medicines after arrival at the final destination shall in no way be limited or waived by reason of the medicines having previously been inspected, tested and passed by the buyer or his authorized representative prior to the medicines dispatch from the place of manufacture or arrival as the case may be.

c) If any inspected or tested medicines fail to conform to the specifications, the buyer may reject them and the seller will remove the rejected medicines at its own cost.

d) During the shelf-life period, the consignees shall be at liberty to draw samples and send it to laboratories approved by State Drugs controllers / run by State Government/NABL approved laboratories/ Government Approved Lab without any intimation to the seller. If found "Not of Standard Quality", (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality upon testing. The seller shall have to replace the rejected batches (unused quantity) with fresh batches within 3 months or refund the cost of the rejected medicines to the buyer, if so, decided by him. In the event of replacement of rejected medicines by the seller, all the above mentioned provision shall apply to the new medicines replaced from the date of replacement thereof, otherwise the seller shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained and the facts will be notified to the Drugs Controller of India/State Drug Controller for taking necessary action.

e) In case any medicines are found substandard either at the inspection stage or during the shelf life of the medicines, the report of the Government approved/NABL accredited laboratory shall be accepted by the seller. If the same is disputed by the seller giving the reasons, the sample will be sent to the designated appellate Lab (Pharmacopoeia commission for Indian medicine & Homeopathy) for the purpose and the report of the same will only be accepted as final and conclusive report. De-registration / debarment action will be taken against the seller according to the category-A and category-B defects as per guidelines issued by the Ministry of Health & Family Welfare.

f) The cost of post-delivery inspection and testing will be borne by the buyer. However, inspection & testing charges for the failed batches shall be borne by the seller.

g) In the event of the samples of medicines supplied fails in quality tests or found to be not as per specifications, the buyer will send second sample to the 2nd Govt. approved /NABL accredited lab. If the second sample fails, the batch will be rejected but if the second sample passes then third sample will be sent to the designated Appellate lab for the medicines and decision of the Appellate lab will be final.

h) In the event of the samples of medicine supplied finally fail in quality tests or found to be not as per specifications, the seller will have to replace the rejected batch with fresh stock duly inspected within 3 months. If not replaced, the buyer will be at liberty to purchase from other source and recovery to be made from the seller and action to blacklist the company/cancellation of the Drug license will also be initiated.

10. Warranty

Each supply shall be accompanied with a "Warranty Certificate" as specified below, duly signed by the Seller as under.

WARRANTY CERTIFICATE:

I/We, _____ (name of the seller), hereby declare that the medicines sold to the _____ (name of the buyer) under this supply order (No. of the supply order with date) are of the best quality (and workmanship) and strictly in accordance with specification and particulars mentioned and I/we hereby guarantee that the said medicines would continue to conform to the description and the quality for a period as specified in the Gazette of India No. 605, dated 20/10/2009 & 16-08-2016 from the date of delivery of the said medicines to the buyer and that notwithstanding the fact that the buyer may have inspected and or approved the said medicines, if during the aforesaid period, discovered not conforming to the description and quality aforesaid or have deteriorated (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality. On such rejection, the medicines will be at the seller's risk and all provisions herein contained relating to the rejection of medicines etc., shall apply. In the event of replacement of rejected medicines by the seller, the above-mentioned guarantee period shall as apply to the medicines replaced from the date of replacement thereof, otherwise the contractor shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained. Nothing here in contained shall prejudice and other right of the buyer in that behalf under this supply order or otherwise.

(Signature name & designation and date with rubber stamp)

11. The classification of defects into different categories will as per guidelines issued by the Drugs Controller of India and action will be taken accordingly.
12. If the seller, having been notified, fails to replace rejected medicines with fresh medicines within 3 months, the buyer may proceed to take such remedial action as may be necessary at the seller's risk and expense and without prejudice to other rights which the buyer may have against the seller under the contract.
13. Loss or premature deterioration due to biological and other activities during the life potency of the medicines shall have to be made good by the seller free of cost or shall have to refund the cost of rejected medicine.
14. Recalls- If medicines must be recalled because of problems with medicines, the seller will be obliged to notify the buyer, providing full details about the reason leading to the recall, and shall take steps to replace the medicines in question at seller's own cost at the ultimate destination with a fresh batch of acceptable medicine or withdraw and give a full refund if the medicine has been taken off the market due to safety issues.
15. Maximum lead time will be 60 days from date of receipt of order and delivery will commence there after.
16. It is the responsibility of the seller to intimate Government e-Marketplace (GEM) about any quality complaints of the medicines reported by any buyer/consignee.
17. Order should be placed for the quantities in multiples of the primary packing.
18. The seller shall not be blacklisted / debarred / banned by any State Governments U.T. / Central Government/Corporations/Local Government Bodies in the preceding 3 years.
19. Seller shall not sell the product(s) for which the firm / company has been blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies due to quality failure of the medicines.
20. During the period of contract if the firm / Company is blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to buyer along with relevant authentic document by seller within one month.
21. Each supply of medicines shall be accompanied with batchwise quality analysis report from government approved /NABL accredited laboratory. This report shall contain specific tests for Identity, Purity, Quality and Strength of the ingredients used in the medicine as per Ayurvedic Pharmacopoeia in case of Ayurvedic medicines and Unani Pharmacopoeia in case of Unani medicines.
22. Packing and Marking
 - a) All containers meant for packing is required to be secured with pilfer proof seal to ensure genuineness of the product packed. With each consignment the seller should give an undertaking that material used is of food grade / HDPE material if supplied in plastic bottle.
 - b) For secondary packing, material is required to be corrugated boxes having "A" grade paper i.e. Virgin, and packed in first-hand boxes only, with suitable flute, joint, stitching, flap, tape. The box should be of 5 ply with bursting strength of 9Kg / cm²
 - c) Weight/volume of the medicines to be mentioned on the inner packing. Weight & other technical requirements shall adhere to as per the pharmacopoeia standard applicable i.e. A.F.I. in case of Ayurveda and N.F.U.M. in case of Unani medicines.
23. 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 drugs/medicines 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 Special Terms and Conditions (STC) which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

Special terms and conditions-Version:4 effective from 14-03-2024 for category Ayurvedic Classical Medicines - Choorna

1.

1. All Provision of Drugs & Cosmetic Act 1940 as amended till date and rules made there under and all rules and regulations issued by Ministry of AYUSH will always be applicable.
2. Only CPSEs/State PSUs/State Pharmacies/Co-operatives are allowed to upload their products for sale based on authorization letter of these manufacturers from State Ayush Department/National Ayush Mission at the time of vendor assessment process.
3. The sellers are allowed to register on GeM and exempted from the Vendor Assessment process

based on the submitted copy of a valid Manufacturing Drug License for the medicine(s) certified by the issuing authority. Buyers must mandatorily ask for submitting the relevant valid drug license and other regulatory documents applicable with the bid. Buyers must also check and validate the details e.g., validity, authenticity/genuineness, name of the drug/medicine under procurement, the license issuing authority etc. at their end.

4. The purchase shall be made through Bidding/RA only irrespective of the value.
5. Manufacturer shall have a valid own manufacturing license issued by the competent drug licensing authority defined under the Drugs and Cosmetics Act 1940 and Rules made there under as amended till date. (If revalidation of drug license has been applied for, the buyer shall be informed accordingly and the copy of application to State Drug / Licensing authority must be submitted with a certificate that application for renewal was made within time frame as per Drug and Cosmetic Act as amended up to date and that has not been deleted by drug licensing authority.)
6. Loan license arrangement shall not be allowed under any circumstances.
7. Medicines must fully comply in all respect with the technical specifications and in accordance with the Pharmacopoeia standards wherever applicable.
8. Each batch of the medicines shall be got tested by the seller from the laboratories approved by State Drug controllers / run by State Government/NABL accredited lab/Government Approved Lab and shall be dispatched along with these test reports.
9. INSPECTION & QUALITY TESTING

a) Medicines shall continue to conform to the description and the quality during the shelf life from the date of delivery of the medicines to the buyer and notwithstanding the fact that the buyer may have inspected and or approved the said medicines.

b) The buyer has the right to inspect, test and where necessary reject the medicines after arrival at the final destination shall in no way be limited or waived by reason of the medicines having previously been inspected, tested and passed by the buyer or his authorized representative prior to the medicines dispatch from the place of manufacture or arrival as the case may be.

c) If any inspected or tested medicines fail to conform to the specifications, the buyer may reject them and the seller will remove the rejected medicines at its own cost.

d) During the shelf-life period, the consignees shall be at liberty to draw samples and send it to laboratories approved by State Drugs controllers / run by State Government/NABL approved laboratories/ Government Approved Lab without any intimation to the seller. If found "Not of Standard Quality", (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality upon testing. The seller shall have to replace the rejected batches (unused quantity) with fresh batches within 3 months or refund the cost of the rejected medicines to the buyer, if so, decided by him. In the event of replacement of rejected medicines by the seller, all the above mentioned provision shall apply to the new medicines replaced from the date of replacement thereof, otherwise the seller shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained and the facts will be notified to the Drugs Controller of India/State Drug Controller for taking necessary action.

e) In case any medicines are found substandard either at the inspection stage or during the shelf life of the medicines, the report of the Government approved/NABL accredited laboratory shall be accepted by the seller. If the same is disputed by the seller giving the reasons, the sample will be sent to the designated appellate Lab (Pharmacopoeia commission for Indian medicine & Homeopathy) for the purpose and the report of the same will only be accepted as final and conclusive report. De-registration / debarment action will be taken against the seller according to the category-A and category-B defects as per guidelines issued by the Ministry of Health & Family Welfare.

f) The cost of post-delivery inspection and testing will be borne by the buyer. However, inspection & testing charges for the failed batches shall be borne by the seller.

g) In the event of the samples of medicines supplied fails in quality tests or found to be not as per specifications, the buyer will send second sample to the 2nd Govt. approved /NABL accredited lab. If the second sample fails, the batch will be rejected but if the second sample passes then third sample will be sent to the designated Appellate lab for the medicines and decision of the Appellate lab will be final.

h) In the event of the samples of medicine supplied finally fail in quality tests or found to be not as per specifications, the seller will have to replace the rejected batch with fresh stock duly inspected

within 3 months. If not replaced, the buyer will be at liberty to purchase from other source and recovery to be made from the seller and action to blacklist the company/cancellation of the Drug license will also be initiated.

10. Warranty

Each supply shall be accompanied with a "Warranty Certificate" as specified below, duly signed by the Seller as under.

WARRANTY CERTIFICATE:

I/We, _____ (name of the seller), hereby declare that the medicines sold to the _____ (name of the buyer) under this supply order (No. of the supply order with date) are of the best quality (and workmanship) and strictly in accordance with specification and particulars mentioned and I/we hereby guarantee that the said medicines would continue to conform to the description and the quality for a period as specified in the Gazette of India No. 605, dated 20/10/2009 & 16-08-2016 from the date of delivery of the said medicines to the buyer and that notwithstanding the fact that the buyer may have inspected and or approved the said medicines, if during the aforesaid period, discovered not conforming to the description and quality aforesaid or have deteriorated (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality. On such rejection, the medicines will be at the seller's risk and all provisions herein contained relating to the rejection of medicines etc., shall apply. In the event of replacement of rejected medicines by the seller, the above-mentioned guarantee period shall as apply to the medicines replaced from the date of replacement thereof, otherwise the contractor shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained. Nothing here in contained shall prejudice and other right of the buyer in that behalf under this supply order or otherwise.

(Signature name & designation and date with rubber stamp)

11. The classification of defects into different categories will as per guidelines issued by the Drugs Controller of India and action will be taken accordingly.
12. If the seller, having been notified, fails to replace rejected medicines with fresh medicines within 3 months, the buyer may proceed to take such remedial action as may be necessary at the seller's risk and expense and without prejudice to other rights which the buyer may have against the seller under the contract.
13. Loss or premature deterioration due to biological and other activities during the life potency of the medicines shall have to be made good by the seller free of cost or shall have to refund the cost of rejected medicine.
14. Recalls- If medicines must be recalled because of problems with medicines, the seller will be obliged to notify the buyer, providing full details about the reason leading to the recall, and shall take steps to replace the medicines in question at seller's own cost at the ultimate destination with a fresh batch of acceptable medicine or withdraw and give a full refund if the medicine has been taken off the market due to safety issues.
15. Maximum lead time will be 60 days from date of receipt of order and delivery will commence there after.
16. It is the responsibility of the seller to intimate Government e-Marketplace (GEM) about any quality complaints of the medicines reported by any buyer/consignee.
17. Order should be placed for the quantities in multiples of the primary packing.
18. The seller shall not be blacklisted / debarred / banned by any State Governments U.T. / Central Government/Corporations/Local Government Bodies in the preceding 3 years.
19. Seller shall not sell the product(s) for which the firm / company has been blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies due to quality failure of the medicines.
20. During the period of contract if the firm / Company is blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to buyer along with relevant authentic document by seller within one month.
21. Each supply of medicines shall be accompanied with batchwise quality analysis report from government approved /NABL accredited laboratory. This report shall contain specific tests for Identity, Purity, Quality and Strength of the ingredients used in the medicine as per Ayurvedic Pharmacopeia in case of Ayurvedic medicines and Unani Pharmacopeia in case of Unani medicines.
22. Packing and Marking

- a) All containers meant for packing is required to be secured with pilfer proof seal to ensure genuineness of the product packed. With each consignment the seller should give an undertaking that material used is of food grade / HDPE material if supplied in plastic bottle.
- b) For secondary packing, material is required to be corrugated boxes having "A" grade paper i.e. Virgin, and packed in first-hand boxes only, with suitable flute, joint, stitching, flap, tape. The box should be of 5 ply with bursting strength of 9Kg / cm²
- c) Weight/volume of the medicines to be mentioned on the inner packing. Weight & other technical requirements shall adhere to as per the pharmacopoeia standard applicable i.e. A.F.I. in case of Ayurveda and N.F.U.M. in case of Unani medicines.

23. 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 drugs/medicines 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 Special Terms and Conditions (STC) which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

Special terms and conditions-Version:4 effective from 14-03-2024 for category Ayurvedic Classical Medicines - Taila

1.

1. All Provision of Drugs & Cosmetic Act 1940 as amended till date and rules made there under and all rules and regulations issued by Ministry of AYUSH will always be applicable.
2. Only CPSEs/State PSUs/State Pharmacies/Co-operatives are allowed to upload their products for sale based on authorization letter of these manufacturers from State Ayush Department/National Ayush Mission at the time of vendor assessment process.
3. The sellers are allowed to register on GeM and exempted from the Vendor Assessment process based on the submitted copy of a valid Manufacturing Drug License for the medicine(s) certified by the issuing authority. Buyers must mandatorily ask for submitting the relevant valid drug license and other regulatory documents applicable with the bid. Buyers must also check and validate the details e.g., validity, authenticity/genuineness, name of the drug/medicine under procurement, the license issuing authority etc. at their end.
4. The purchase shall be made through Bidding/RA only irrespective of the value.
5. Manufacturer shall have a valid own manufacturing license issued by the competent drug licensing authority defined under the Drugs and Cosmetics Act 1940 and Rules made there under as amended till date. (If revalidation of drug license has been applied for, the buyer shall be informed accordingly and the copy of application to State Drug / Licensing authority must be submitted with a certificate that application for renewal was made within time frame as per Drug and Cosmetic Act as amended up to date and that has not been deleted by drug licensing authority.)
6. Loan license arrangement shall not be allowed under any circumstances.
7. Medicines must fully comply in all respect with the technical specifications and in accordance with the Pharmacopoeia standards wherever applicable.
8. Each batch of the medicines shall be got tested by the seller from the laboratories approved by State Drug controllers / run by State Government/NABL accredited lab/Government Approved Lab and shall be dispatched along with these test reports.
9. INSPECTION & QUALITY TESTING

a) Medicines shall continue to conform to the description and the quality during the shelf life from the date of delivery of the medicines to the buyer and notwithstanding the fact that the buyer may have inspected and or approved the said medicines.

b) The buyer has the right to inspect, test and where necessary reject the medicines after arrival at the final destination shall in no way be limited or waived by reason of the medicines having previously been inspected, tested and passed by the buyer or his authorized representative prior to the medicines dispatch from the place of manufacture or arrival as the case may be.

c) If any inspected or tested medicines fail to conform to the specifications, the buyer may reject them and the seller will remove the rejected medicines at its own cost.

d) During the shelf-life period, the consignees shall be at liberty to draw samples and send it to laboratories approved by State Drugs controllers / run by State Government/NABL approved laboratories/ Government Approved Lab without any intimation to the seller. If found "Not of Standard Quality", (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not

conforming to the said description and quality upon testing. The seller shall have to replace the rejected batches (unused quantity) with fresh batches within 3 months or refund the cost of the rejected medicines to the buyer, if so, decided by him. In the event of replacement of rejected medicines by the seller, all the above mentioned provision shall apply to the new medicines replaced from the date of replacement thereof, otherwise the seller shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained and the facts will be notified to the Drugs Controller of India/State Drug Controller for taking necessary action.

e) In case any medicines are found substandard either at the inspection stage or during the shelf life of the medicines, the report of the Government approved/NABL accredited laboratory shall be accepted by the seller. If the same is disputed by the seller giving the reasons, the sample will be sent to the designated appellate Lab (Pharmacopoeia commission for Indian medicine & Homeopathy) for the purpose and the report of the same will only be accepted as final and conclusive report. De-registration / debarment action will be taken against the seller according to the category-A and category-B defects as per guidelines issued by the Ministry of Health & Family Welfare.

f) The cost of post-delivery inspection and testing will be borne by the buyer. However, inspection & testing charges for the failed batches shall be borne by the seller.

g) In the event of the samples of medicines supplied fails in quality tests or found to be not as per specifications, the buyer will send second sample to the 2nd Govt. approved /NABL accredited lab. If the second sample fails, the batch will be rejected but if the second sample passes then third sample will be sent to the designated Appellate lab for the medicines and decision of the Appellate lab will be final.

h) In the event of the samples of medicine supplied finally fail in quality tests or found to be not as per specifications, the seller will have to replace the rejected batch with fresh stock duly inspected within 3 months. If not replaced, the buyer will be at liberty to purchase from other source and recovery to be made from the seller and action to blacklist the company/cancellation of the Drug license will also be initiated.

10. Warranty

Each supply shall be accompanied with a "Warranty Certificate" as specified below, duly signed by the Seller as under.

WARRANTY CERTIFICATE:

I/We, _____ (name of the seller), hereby declare that the medicines sold to the _____ (name of the buyer) under this supply order (No. of the supply order with date) are of the best quality (and workmanship) and strictly in accordance with specification and particulars mentioned and I/we hereby guarantee that the said medicines would continue to conform to the description and the quality for a period as specified in the Gazette of India No. 605, dated 20/10/2009 & 16-08-2016 from the date of delivery of the said medicines to the buyer and that notwithstanding the fact that the buyer may have inspected and or approved the said medicines, if during the aforesaid period, discovered not conforming to the description and quality aforesaid or have deteriorated (the decision of the buyer in this behalf will be final and conclusive), the buyer will be entitled to reject the said medicines of such portion thereof as may be discovered not conforming to the said description and quality. On such rejection, the medicines will be at the seller's risk and all provisions herein contained relating to the rejection of medicines etc., shall apply. In the event of replacement of rejected medicines by the seller, the above-mentioned guarantee period shall as apply to the medicines replaced from the date of replacement thereof, otherwise the contractor shall pay to the buyer such damages as may arise by the reasons of the breach of the conditions here in contained. Nothing here in contained shall prejudice and other right of the buyer in that behalf under this supply order or otherwise.

(Signature name & designation and date with rubber stamp)

11. The classification of defects into different categories will as per guidelines issued by the Drugs Controller of India and action will be taken accordingly.
12. If the seller, having been notified, fails to replace rejected medicines with fresh medicines within 3 months, the buyer may proceed to take such remedial action as may be necessary at the seller's risk and expense and without prejudice to other rights which the buyer may have against the seller

under the contract.

13. Loss or premature deterioration due to biological and other activities during the life potency of the medicines shall have to be made good by the seller free of cost or shall have to refund the cost of rejected medicine.
14. Recalls- If medicines must be recalled because of problems with medicines, the seller will be obliged to notify the buyer, providing full details about the reason leading to the recall, and shall take steps to replace the medicines in question at seller's own cost at the ultimate destination with a fresh batch of acceptable medicine or withdraw and give a full refund if the medicine has been taken off the market due to safety issues.
15. Maximum lead time will be 60 days from date of receipt of order and delivery will commence there after.
16. It is the responsibility of the seller to intimate Government e-Marketplace (GEM) about any quality complaints of the medicines reported by any buyer/consignee.
17. Order should be placed for the quantities in multiples of the primary packing.
18. The seller shall not be blacklisted / debarred / banned by any State Governments U.T. / Central Government/Corporations/Local Government Bodies in the preceding 3 years.
19. Seller shall not sell the product(s) for which the firm / company has been blacklisted/debarred/de-registered/banned by any State Government / Central Government / its Drug procurement agencies due to quality failure of the medicines.
20. During the period of contract if the firm / Company is blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to buyer along with relevant authentic document by seller within one month.
21. Each supply of medicines shall be accompanied with batchwise quality analysis report from government approved /NABL accredited laboratory. This report shall contain specific tests for Identity, Purity, Quality and Strength of the ingredients used in the medicine as per Ayurvedic Pharmacopeia in case of Ayurvedic medicines and Unani Pharmacopeia in case of Unani medicines.
22. Packing and Marking
 - a) All containers meant for packing is required to be secured with pilfer proof seal to ensure genuineness of the product packed. With each consignment the seller should give an undertaking that material used is of food grade / HDPE material if supplied in plastic bottle.
 - b) For secondary packing, material is required to be corrugated boxes having "A" grade paper i.e. Virgin, and packed in first-hand boxes only, with suitable flute, joint, stitching, flap, tape. The box should be of 5 ply with bursting strength of 9Kg / cm²
 - c) Weight/volume of the medicines to be mentioned on the inner packing. Weight & other technical requirements shall adhere to as per the pharmacopoeia standard applicable i.e. A.F.I. in case of Ayurveda and N.F.U.M. in case of Unani medicines.
23. 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 drugs/medicines 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 Special Terms and Conditions (STC) which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

क्रेता द्वारा जोड़ी गई बिड की विशेष शर्तें/Buyer Added Bid Specific Terms and Conditions

1. 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 up to 25% of the contracted quantity during the currency of the contract at the contracted rates. The delivery period of quantity shall commence from the last date of original delivery order and in cases where option clause is exercised during the extended delivery period the additional time shall commence from the last date of extended delivery period. The additional delivery time shall be $(\text{Increased quantity} \div \text{Original quantity}) \times \text{Original delivery period (in days)}$, subject to minimum of 30 days. If the original delivery period is less than 30 days, the additional time equals the original delivery period. The Purchaser may extend this calculated delivery duration up to the original delivery period while exercising the option clause. Bidders must comply with these terms.

अस्वीकरण/Disclaimer

The additional terms and conditions have been incorporated by the Buyer after approval of the Competent Authority in Buyer Organization, whereby Buyer organization is solely responsible for the impact of these clauses on the bidding process, its outcome, and consequences thereof including any eccentricity / restriction arising in the bidding process due to these ATCs and due to modification of technical specifications and / or terms and conditions governing the bid. If any clause(s) is / are incorporated by the Buyer regarding following, the bid and resultant contracts shall be treated as null and void and such bids may be cancelled by GeM at any stage of bidding process without any notice:-

1. Definition of Class I and Class II suppliers in the bid not in line with the extant Order / Office Memorandum issued by DPIIT in this regard.
2. Seeking EMD submission from bidder(s), including via Additional Terms & Conditions, in contravention to exemption provided to such sellers under GeM GTC.
3. Publishing Custom / BOQ bids for items for which regular GeM categories are available without any Category item bunched with it.
4. Creating BoQ bid for single item.
5. Mentioning specific Brand or Make or Model or Manufacturer or Dealer name.
6. Mandating submission of documents in physical form as a pre-requisite to qualify bidders.
7. Floating / creation of work contracts as Custom Bids in Services.
8. Seeking sample with bid or approval of samples during bid evaluation process. (However, in bids for [attached categories](#), trials are allowed as per approved procurement policy of the buyer nodal Ministries)
9. Mandating foreign / international certifications even in case of existence of Indian Standards without specifying equivalent Indian Certification / standards.
10. Seeking experience from specific organization / department / institute only or from foreign / export experience.
11. Creating bid for items from irrelevant categories.
12. Incorporating any clause against the MSME policy and Preference to Make in India Policy.
13. Reference of conditions published on any external site or reference to external documents/clauses.
14. Asking for any Tender fee / Bid Participation fee / Auction fee in case of Bids / Forward Auction, as the case may be.
15. Buyer added ATC Clauses which are in contravention of clauses defined by buyer in system generated bid template as indicated above in the Bid Details section, EMD Detail, ePBG Detail and MII and MSE Purchase Preference sections of the bid, unless otherwise allowed by GeM GTC.
16. In a category based bid, adding additional items, through buyer added additional scope of work/ additional terms and conditions/or any other document. If buyer needs more items along with the main item, the same must be added through bunching category based items or by bunching custom catalogs or bunching a BoQ with the main category based item, the same must not be done through ATC or Scope of Work.

Further, if any seller has any objection/grievance against these additional clauses or otherwise on any aspect of this bid, they can raise their representation against the same by using the Representation window provided in the bid details field in Seller dashboard after logging in as a seller within 4 days of bid publication on GeM. Buyer is duty bound to reply to all such representations and would not be allowed to open bids if he fails to reply to such representations.

All GeM Sellers/Service Providers shall ensure full compliance with all applicable labour laws, including the provisions, rules, schemes and guidelines under the four Labour Codes i.e. the Code on Wages, 2019; the Industrial Relations Code, 2020; the Occupational Safety, Health and Working Conditions Code, 2020; and the Code on Social Security, 2020 as and when notified and brought into force by the Government of India.

For all provisions of the Labour Codes that are pending operationalisation through rules, schemes or notifications, the corresponding provisions of the pre-existing labour enactments (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. and relevant State Rules) shall continue to remain applicable.

The Seller/ Service Providers shall, therefore, be responsible for ensuring compliance under:

- **All notified and enforceable provisions of the new Labour Codes as mentioned hereinabove; and**
- **All operative provisions of the erstwhile Labour Laws until their complete substitution.**

All obligations relating to wages, social security, safety, working conditions, industrial relations etc. and any other statutory requirements shall be strictly met by the Seller/ Service Provider. Any non-compliance shall constitute a breach of the contract and shall entitle the Buyer to take appropriate action in accordance with the contract and applicable law.

[यह बिड सामान्य शर्तों के अंतर्गत भी शासित है /This Bid is also governed by the General Terms and Conditions](#)

जेम की सामान्य शर्तों के खंड 26 के संदर्भ में भारत के साथ भूमि सीमा साझा करने वाले देश के बिडर से खरीद पर प्रतिबंध के संबंध में भारत के साथ भूमि सीमा साझा करने वाले देश का कोई भी बिडर इस निविदा में बिड देने के लिए तभी पात्र होगा जब वह बिड देने वाला सक्षम प्राधिकारी के पास पंजीकृत हो। बिड में भाग लेते समय बिडर को इसका अनुपालन करना होगा और कोई भी गलत घोषणा किए जाने व इसका अनुपालन न करने पर अनुबंध को तत्काल समाप्त करने और कानून के अनुसार आगे की कानूनी कार्रवाई का आधार होगा।/In terms of GeM GTC clause 26 regarding Restrictions on procurement from a bidder of a country which shares a land border with India, any bidder from a country which shares a land border with India will be eligible to bid in this tender only if the bidder is registered with the Competent Authority. While participating in bid, Bidder has to undertake compliance of this and any false declaration and non-compliance of this would be a ground for immediate termination of the contract and further legal action in accordance with the laws.

---धन्यवाद/Thank You---