

## बिड दस्तावेज़ / Bid Document

बिड विवरण/Bid Details	
बिड बंद होने की तारीख/समय /Bid End Date/Time	28-05-2026 15:00:00
बिड खुलने की तारीख/समय /Bid Opening Date/Time	28-05-2026 15:30:00
बिड पेशकश वैधता (बंद होने की तारीख से)/Bid Offer Validity (From End Date)	150 (Days)
मंत्रालय/राज्य का नाम/Ministry/State Name	Ministry Of Health And Family Welfare
विभाग का नाम/Department Name	Department Of Health And Family Welfare
संगठन का नाम/Organisation Name	Central Medical Services Society (cmss)
कार्यालय का नाम/Office Name	Central Medical Services Society
कुल मात्रा/Total Quantity	94578000
वस्तु श्रेणी /Item Category	Condoms (Free Supply) for Family Planning Programme (Q1) , Emergency Contraceptives Pills (ECP) for Family Planning Programme (Q1) , Injectable Contraceptive (Antara) for Family Planning Programme (Q1) , IUCD 380A for Family Planning Programme (Q1) , IUCD 375 for Family Planning Programme (Q1) , Tubal Rings for Family Planning Programme (Q1)
बिडर का न्यूनतम औसत वार्षिक टर्नओवर (3 वर्षों का) /Minimum Average Annual Turnover of the bidder (For 3 Years)	25 Lakh (s)
मूल उपकरण निर्माता का औसत टर्नओवर (गत 3 वर्षों का)/OEM Average Turnover (Last 3 Years)	50 Lakh (s)
उन्हीं/समान सेवा के लिए अपेक्षित विगत अनुभव के वर्ष/Years of Past Experience Required for same/similar service	2 Year (s)
टर्नओवर के लिए एमएसई को छूट प्राप्त है / MSE Relaxation for Turnover	Yes   Partial   Turn over value - 0 (in lakhs)
टर्नओवर के लिए स्टार्टअप को छूट प्राप्त है / Startup Relaxation for Turnover	Yes   Partial   Turn over value - 0 (in lakhs)

<b>बिड विवरण/Bid Details</b>	
<b>विक्रेता से मांगे गए दस्तावेज़/Document required from seller</b>	Experience Criteria,Past Performance,Bidder Turnover,Certificate (Requested in ATC),OEM Authorization Certificate,OEM Annual Turnover,Additional Doc 1 (Requested in ATC),Additional Doc 2 (Requested in ATC),Additional Doc 3 (Requested in ATC),Additional Doc 4 (Requested in ATC),Compliance of BoQ specification and supporting document *In case any bidder is seeking exemption from Experience / Turnover Criteria, the supporting documents to prove his eligibility for exemption must be uploaded for evaluation by the buyer
<b>क्या आप निविदाकारों द्वारा अपलोड किए गए दस्तावेज़ों को निविदा में भाग लेने वाले सभी निविदाकारों को दिखाना चाहते हैं? संदर्भ मेनू है/Do you want to show documents uploaded by bidders to all bidders participated in bid?</b>	Yes (Documents submitted as part of a clarification or representation during the tender/bid process will also be displayed to other participated bidders after log in)
<b>बिड लगाने की समय सीमा स्वतः नहीं बढ़ाने के लिए आवश्यक बिड की संख्या। / Minimum number of bids required to disable automatic bid extension</b>	1
<b>दिनों की संख्या, जिनके लिए बिड लगाने की समय-सीमा बढ़ाई जाएगी। / Number of days for which Bid would be auto-extended</b>	4
<b>ऑटो एक्सटेंशन अधिकतम कितनी बार किया जाना है। / Number of Auto Extension count</b>	1
<b>विगत प्रदर्शन /Past Performance</b>	40 %
<b>बिड से रिवर्स नीलामी सक्रिय किया/Bid to RA enabled</b>	No
<b>बिड का प्रकार/Type of Bid</b>	Two Packet Bid
<b>प्राथमिक उत्पाद श्रेणी/Primary product category</b>	Condoms (Free Supply) for Family Planning Programme
<b>तकनीकी मूल्यांकन के दौरान तकनीकी स्पष्टीकरण हेतु अनुमत समय /Time allowed for Technical Clarifications during technical evaluation</b>	2 Days
<b>निरीक्षण आवश्यक (सूचीबद्ध निरीक्षण प्राधिकरण /जेम के साथ पूर्व पंजीकृत एजेंसियों द्वारा)/Inspection Required (By Empanelled Inspection Authority / Agencies pre-registered with GeM)</b>	No
<b>क्या पार्ट क्वांटिटी बोली लगाने की अनुमति है? / Is Part Quantity Bidding Allowed?</b>	Yes
<b>मूल्यांकन पद्धति/Evaluation Method</b>	Item wise evaluation
<b>मध्यस्थता खंड/Arbitration Clause</b>	No
<b>सुलह खंड/Mediation Clause</b>	No

#### ईएमडी विवरण/EMD Detail

एडवाइजरी बैंक/Advisory Bank	State Bank of India
Schedule 1 ईएमडी राशि/EMD Amount (In INR)	1473141
Schedule 2 ईएमडी राशि/EMD Amount (In INR)	201104
Schedule 3 ईएमडी राशि/EMD Amount (In INR)	1646039
Schedule 4 ईएमडी राशि/EMD Amount (In INR)	314880
Schedule 5 ईएमडी राशि/EMD Amount (In INR)	344655
Schedule 6 ईएमडी राशि/EMD Amount (In INR)	126108

### ईपीबीजी विवरण /ePBG Detail

आवश्यकता/Required	No
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(a). जेम की शर्तों के अनुसार ईएमडी छूट के इच्छुक बिडर को संबंधित कटेगरी के लिए बिड के साथ वैध समर्थित दस्तावेज प्रस्तुत करने है। एमएसई कटेगरी के अंतर्गत केवल वस्तुओं के लिए विनिर्माता तथा सेवाओं के लिए सेवा प्रदाता ईएमडी से छूट के पात्र हैं। व्यापारियों को इस नीति के दायरे से बाहर रखा गया है।/EMD EXEMPTION: The bidder seeking EMD exemption, must submit the valid supporting document for the relevant category as per GeM GTC with the bid. Under MSE category, only manufacturers for goods and Service Providers for Services are eligible for exemption from EMD. Traders are excluded from the purview of this Policy.

(b). The EMD Amount will be applicable for each schedule/group selected during Bid creation.

(c). ईएमडी और संपादन जमानत राशि, जहां यह लागू होती है, लाभार्थी के पक्ष में होनी चाहिए। / EMD & Performance security should be in favour of Beneficiary, wherever it is applicable.

### लाभार्थी /Beneficiary :

DG&CEO

Central Medical Services Society, Department of Health and Family Welfare, Central Medical Services Society (CMSS), Ministry of Health and Family Welfare (Cmss New Delhi)

### विभाजन/Splitting

विभाजन/Splitting Applied	Yes
बोलीदाताओं की अधिकतम संख्या, जिनके बीच ऑर्डर विभाजित किया जा सकता है। / Maximum No. Of Bidders Amongst Which Order May Be Split	2
विभाजन मानदंड इस बात पर आधारित है कि कौन सी क्वांटिटी को वितरित किया जाएगा / Split Criteria based on which quantity will be distributed	70:30

### एमआईआई खरीद वरीयता / MII Purchase Preference

एमआईआई खरीद वरीयता / MII Purchase Preference	Yes
मेक इन इंडिया विक्रेताओं को खरीद में प्राथमिकता, यदि उनका मूल्य L1+X% तक की सीमा में है / Purchase Preference to MII sellers available upto price within L1+X%	20
मेक इन इंडिया खरीद में प्राथमिकता के लिए बिड की मात्रा का अधिकतम प्रतिशत / Maximum Percentage of Bid quantity for MII purchase preference	50
सार्वजनिक खरीद (मेक-इन-इंडिया को प्राथमिकता) आदेश 2017 के अनुसार केवल क्लास 1/क्लास 2 के स्थानीय आपूर्तिकर्ताओं को ही भागीदारी की अनुमति है दिनांक 16.09.2020 (समय-समय पर संशोधित एवं लागू) / Allow participation only from Class 1/Class 2 local suppliers as per the Public procurement(Preference to Make-in-india) order 2017 date 16.09.2020(as amended and applicable time to time)	Yes, in compliance with the MII ORDER : DPIIT Order(as amended and applicable time to time)

#### एमएसई खरीद वरीयता/MSE Purchase Preference

एमएसई खरीद वरीयता/MSE Purchase Preference	Yes
सूक्ष्म और लघु उद्यम मूल उपकरण निर्माताओं को खरीद में प्राथमिकता, यदि उनका मूल्य L1+X% तक की सीमा में हो / Purchase Preference to MSE OEMs available upto price within L1+X%	15
सूक्ष्म और लघु उद्यम को खरीद में प्राथमिकता के लिए बिड की मात्रा का अधिकतम प्रतिशत / Maximum Percentage of Bid quantity for MSE purchase preference	25

1. If the bidder is a Micro or Small Enterprise (MSE) as per latest orders issued by Ministry of MSME, the bidder shall be relaxed from the eligibility criteria of "Bidder Turnover" as defined above subject to meeting of quality and technical specifications. If the bidder itself is MSE OEM of the offered products, it would be relaxed from the "OEM Average Turnover" criteria also subject to meeting of quality and technical specifications. The bidder seeking Relaxation from Turnover, shall upload the supporting documents to prove his eligibility for Relaxation.
2. If the bidder is a DPIIT registered Startup, the bidder shall be relaxed from the the eligibility criteria of "Bidder Turnover" as defined above subject to their meeting of quality and technical specifications. If the bidder is DPIIT Registered OEM of the offered products, it would be relaxed from the "OEM Average Turnover" criteria also subject to meeting of quality and technical specifications. The bidder seeking Relaxation from Turnover shall upload the supporting documents to prove his eligibility for Relaxation.
3. The minimum average annual financial turnover of the bidder during the last three years, ending on 31st March of the previous financial year, should be as indicated above in the bid document. Documentary evidence in the form of certified Audited Balance Sheets of relevant periods or a certificate from the Chartered Accountant / Cost Accountant indicating the turnover details for the relevant period shall be uploaded with the bid. In case the date of constitution / incorporation of the bidder is less than 3-year-old, the average turnover in respect of the completed financial years after the date of constitution shall be taken into account for this criteria.
4. Experience Criteria: In respect of the filter applied for experience criteria, the Bidder or its OEM of the product offered in the bid {themselves or through reseller(s)} should have regularly, manufactured and supplied same or similar Category Products to any Central / State Govt Organization / PSU for number of Financial years as indicated above in the bid document before the bid opening date. Copies of relevant contracts and delivery acceptance certificates like CRAC to be submitted along with bid in support of having supplied some quantity

during each of the Financial year. In case of bunch bids, the category of primary product having highest value should meet this criterion.

5. OEM Turn Over Criteria: The minimum average annual financial turnover of the OEM of the offered product during the last three years, ending on 31st March of the previous financial year, should be as indicated in the bid document. Documentary evidence in the form of certified Audited Balance Sheets of relevant periods or a certificate from the Chartered Accountant / Cost Accountant indicating the turnover details for the relevant period shall be uploaded with the bid. In case the date of constitution / incorporation of the OEM is less than 3 year old, the average turnover in respect of the completed financial years after the date of constitution shall be taken into account for this criteria.

6. Preference to Make In India products (For bids < 200 Crore): Preference shall be given to Class 1 local supplier as defined in public procurement (Preference to Make in India), Order 2017 as amended from time to time and its subsequent Orders/Notifications issued by concerned Nodal Ministry for specific Goods/Products. The minimum local content to qualify as a Class 1 local supplier is denoted in the bid document. If the bidder wants to avail the Purchase preference, the bidder must upload a certificate from the OEM regarding the percentage of the local content and the details of locations at which the local value addition is made along with their bid, failing which no purchase preference shall be granted. In case the bid value is more than Rs 10 Crore, the declaration relating to percentage of local content shall be certified by the statutory auditor or cost auditor, if the OEM is a company and by a practicing cost accountant or a chartered accountant for OEMs other than companies as per the Public Procurement (preference to Make-in -India) order 2017 dated 04.06.2020. Only Class-I and Class-II Local suppliers as per MII order dated 4.6.2020 will be eligible to bid. Non - Local suppliers as per MII order dated 04.06.2020 are not eligible to participate. However, eligible micro and small enterprises will be allowed to participate .The buyers are advised to refer the OM No.F.1/4/2021-PPD dated 18.05.2023.

[OM No.1 4 2021 PPD dated 18.05.2023](#) for compliance of Concurrent application of Public Procurement Policy for Micro and Small Enterprises Order, 2012 and Public Procurement (Preference to Make in India) Order, 2017.

7. Purchase preference will be given to MSEs having valid Udyam Registration and whose credentials are validated online through Udyam Registration portal as defined in Public Procurement Policy for Micro and Small Enterprises (MSEs) Order, 2012 dated 23.03.2012 issued by Ministry of Micro, Small and Medium Enterprises and its subsequent Orders/Notifications issued by concerned Ministry. If the bidder wants to avail themselves of the Purchase preference, 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 for any purchase preference. 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 for purchase preference based on documentary evidence submitted, while evaluating the bid. If L-1 is not an MSE and MSE Seller (s) has / have quoted price within L-1+ 15% (Selected by Buyer) of margin of purchase preference /price band defined in relevant policy, such MSE Seller shall be given opportunity to match L-1 price and contract will be awarded for 25% (selected by Buyer) percentage of total quantity. The buyers are advised to refer the OM No. F.1/4/2021-PPD dated 18.05.2023 [OM No.1 4 2021 PPD dated 18.05.2023](#) for compliance of Concurrent application of Public Procurement Policy for Micro and Small Enterprises Order, 2012 and Public Procurement (Preference to Make in India) Order, 2017. 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.

## पार्ट क्वांटिटी बोली लगाना / Part Quantity Bidding

Buyer has allowed part quantity bidding, bidders can offer maximum quantity that they can deliver keeping in mind their capacity and delivery period requirements. The offer quantity has to be more than minimum bid quantity as specified by the Buyer in the bid. Offers with quantity less than Minimum are liable to be rejected. It may however be noted that there is no guarantee that full offer quantity will be ordered by the buyer. Quantity to be ordered by the buyer will depend on various factors including the Ranking of the bidder, Offered quantity, Splitting criteria indicated by the buyer in the bid and the requirement of the buyer to have multiple sources of supply for ensuring supply chain etc. Sellers would be notified about likely order quantity or range of possible order quantity at the time of price match request made by the buyer. ward of contract will be subject to acceptance of price match request along with min / max offer quantity as decided by the Buyer.

8. Estimated Bid Value indicated above is being declared solely for the purpose of guidance on EMD amount and for determining the Eligibility Criteria related to Turn Over, Past Performance and Project / Past Experience etc. This has no relevance or bearing on the price to be quoted by the bidders and is also not going to have any impact on bid participation. Also this is not going to be used as a criteria in determining reasonableness of quoted prices which would be determined by the buyer based on its own assessment of reasonableness and

based on competitive prices received in Bid / RA process.

9. Past Performance: The Bidder or its OEM {themselves or through re-seller(s)} should have supplied same or similar Category Products for 40% of bid quantity, in at least one of the last three Financial years before the bid opening date to any Central / State Govt Organization / PSU. Copies of relevant contracts (proving supply of cumulative order quantity in any one financial year) to be submitted along with bid in support of quantity supplied in the relevant Financial year. In case of bunch bids, the category related to primary product having highest bid value should meet this criterion.

#### Pre Bid Detail(s)

<b>मूल्य भिन्नता खंड दस्तावेज/Pre-Bid Date and Time</b>	<b>प्री-बिड स्थान/Pre-Bid Venue</b>
11-05-2026 11:00:00	CMSS Conference Hall

#### मूल्यांकन विधि(मदवार मूल्यांकन विधि) / Evaluation Method ( Item Wise Evaluation Method )

Contract will be awarded schedulewise and the determination of L1 will be done separately for each schedule. The details of item-consignee combination covered under each schedule are as under:

मूल्यांकन अनुसूचियां / Evaluation Schedules	वस्तु/श्रेणी / Item/Category	मात्रा / Quantity
Schedule 1	Condoms (free Supply) For Family Planning Programme	80062000
Schedule 2	Emergency Contraceptives Pills (ecp) For Family Planning Programme	7476000
Schedule 3	Injectable Contraceptive (antara) For Family Planning Programme	4663000
Schedule 4	Iucd 380a For Family Planning Programme	820000
Schedule 5	Iucd 375 For Family Planning Programme	999000
Schedule 6	Tubal Rings For Family Planning Programme	558000

#### विक्रेता से आवश्यक मदवार न्यूनतम क्षमता / Itemwise Minimum Capacity Required From Seller

S.N o.	Schedule Name	Item Category	Item Quantity	Minimum Capacity
1	Schedule 1	Condoms (Free Supply) for Family Planning Programme	80062000	40031000
2	Schedule 2	Emergency Contraceptives Pills (ECP) for Family Planning Programme	7476000	3738000
3	Schedule 3	Injectable Contraceptive (Antara) for Family Planning Programme	4663000	2331500
4	Schedule 4	IUCD 380A for Family Planning Programme	820000	410000
5	Schedule 5	IUCD 375 for Family Planning Programme	999000	499500
6	Schedule 6	Tubal Rings for Family Planning Programme	558000	279000

#### Condoms (Free Supply) For Family Planning Programme ( 80062000 pieces )

(क्रमशः श्रेणी 1 और श्रेणी 2 के स्थानीय आपूर्तिकर्ता के रूप में अर्हता प्राप्त करने के लिए आवश्यक/Minimum 50% and 26% Local Content required for qualifying as Class 1 and Class 2 Local Supplier respectively)

#### 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)
PRODUCT INFORMATION	<b>Product Description</b>	Condom (Free Supply)
	<b>Conformity to technical specifications</b>	As per detailed technical specification document attached/uploaded
	Attached/uploaded technical specification has been seen, read and understood	Yes
CERTIFICATIONS	Compliance to Medical Device Rules (MDR) 2017 as amended till date	Yes
	Availability of valid medical device manufacturing license for the product issued from the competent authority defined under Drugs and Cosmetic Act 1940 and Rules made there under as amended till date	Yes
	Manufacturing Facility Certification	ISO:13485 (Latest)
	Availability of Test Report for each supplied batch/product as per Medical Device Rules (MDR) 2017 as amended till date	Yes
	Submission of all necessary certifications, licenses and test reports to the buyer at the time of bid submission and/or along with supplies as per buyer requirement	Yes

अतिरिक्त विशिष्टि दस्तावेज़ /Additional Specification Documents

Applicable Specification Document	<a href="#">View</a>
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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

क्र.सं./S.No.	परेषिती/रिपोर्टिंग अधिकारी /Consignee Reporting/Officer	पता/Address	मात्रा /Quantity	डिलीवरी के दिन/Delivery Days
1	Anurodh Singh	110021,Second Floor , Viswa Yuvak Kendra , Chanakyapuri New Delhi -110021	80062000	180

### Emergency Contraceptives Pills (ECP) For Family Planning Programme ( 7476000 packet )

(क्रमशः श्रेणी 1 और श्रेणी 2 के स्थानीय आपूर्तिकर्ता के रूप में अर्हता प्राप्त करने के लिए आवश्यक/Minimum 80% and 51% Local Content required for qualifying as Class 1 and Class 2 Local Supplier respectively)

#### 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)
PRODUCT INFORMATION	<b>Product Description</b>	Emergency Contraceptive Pills (ECP)
	<b>Conformity to technical specifications</b>	As per detailed technical specification document attached/uploaded
	Attached/uploaded technical specification has been seen, read and understood	Yes
CERTIFICATIONS	Availability of valid drug manufacturing license issued from the competent authority defined under Drugs and Cosmetic Act and Rules there under as amended till date	Yes
	Submission of all necessary certifications, licenses and test reports to the buyer at the time of bid submission and/or along with supplies as per buyer requirement	Yes

#### अतिरिक्त विशिष्टि दस्तावेज़ /Additional Specification Documents

Applicable Specification Document	<a href="#">View</a>
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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	Anurodh Singh	110021,Second Floor , Viswa Yuvak Kendra , Chanakyapuri New Delhi -110021	7476000	179

### Injectable Contraceptive (Antara) For Family Planning Programme ( 4663000 dose(s) )

(क्रमशः श्रेणी 1 और श्रेणी 2 के स्थानीय आपूर्तिकर्ता के रूप में अर्हता प्राप्त करने के लिए आवश्यक/Minimum 80% and 51% Local Content required for qualifying as Class 1 and Class 2 Local Supplier respectively)

#### 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)
PRODUCT INFORMATION	<b>Product Description</b>	Injectable Contraceptive (Medroxyprogesterone Acetate)
	<b>Conformity to technical specifications</b>	As per detailed technical specification document attached/uploaded
	Attached/uploaded technical specification has been seen, read and understood	Yes
CERTIFICATIONS	Availability of valid drug manufacturing license issued from the competent authority defined under Drugs and Cosmetic Act and Rules there under as amended till date	Yes
	Submission of all necessary certifications, licenses and test reports to the buyer at the time of bid submission and/or along with supplies as per buyer requirement	Yes

#### अतिरिक्त विशिष्टि दस्तावेज़ /Additional Specification Documents

Applicable Specification Document	<a href="#">View</a>
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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	Anurodh Singh	110021,Second Floor , Viswa Yuvak Kendra , Chanakyapuri New Delhi -110021	4663000	179

### IUCD 380A For Family Planning Programme ( 820000 pieces )

(क्रमशः श्रेणी 1 और श्रेणी 2 के स्थानीय आपूर्तिकर्ता के रूप में अर्हता प्राप्त करने के लिए आवश्यक/Minimum 50% and 26% Local Content required for qualifying as Class 1 and Class 2 Local Supplier respectively)

#### 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)
PRODUCT INFORMATION	<b>Product Description</b>	Copper-T 380A
	<b>Conformity to technical specifications</b>	As per detailed technical specification document attached/uploaded
	Attached/uploaded technical specification has been seen, read and understood	Yes
CERTIFICATIONS	Compliance to Medical Device Rules (MDR) 2017 as amended till date	Yes
	Availability of valid medical device manufacturing license for the product issued from the competent authority defined under Drugs and Cosmetic Act 1940 and Rules made there under as amended till date	Yes
	Manufacturing Facility Certification	ISO:13485 (Latest)
	Availability of Test Report for each supplied batch/product as per Medical Device Rules (MDR) 2017 as amended till date	Yes
	Submission of all necessary certifications, licenses and test reports to the buyer at the time of bid submission and/or along with supplies as per buyer requirement	Yes

**अतिरिक्त विशिष्टि दस्तावेज़ /Additional Specification Documents**

Applicable Specification Document	<a href="#">View</a>
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**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	Anurodh Singh	110021,Second Floor , Viswa Yuvak Kendra , Chanakyapuri New Delhi -110021	820000	178

**IUCD 375 For Family Planning Programme ( 999000 pieces )**

(क्रमशः श्रेणी 1 और श्रेणी 2 के स्थानीय आपूर्तिकर्ता के रूप में अर्हता प्राप्त करने के लिए आवश्यक/Minimum 50% and 26% Local Content required for qualifying as Class 1 and Class 2 Local Supplier respectively)

**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)
PRODUCT INFORMATION	<b>Product Description</b>	Copper Intra Uterine Contraceptive Device 375
	Conformity to technical specifications	As per detailed technical specification document attached/uploaded
	Attached/uploaded technical specification has been seen, read and understood	Yes
CERTIFICATIONS	Compliance to Medical Device Rules (MDR) 2017 as amended till date	Yes
	Availability of valid medical device manufacturing license for the product issued from the competent authority defined under Drugs and Cosmetic Act 1940 and Rules made there under as amended till date	Yes

**अतिरिक्त विशिष्टि दस्तावेज़ /Additional Specification Documents**

Applicable Specification Document	<a href="#">View</a>
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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	Anurodh Singh	110021,Second Floor , Viswa Yuvak Kendra , Chanakyapuri New Delhi -110021	999000	179

**Tubal Rings For Family Planning Programme ( 558000 pairs )**

(क्रमशः श्रेणी 1 और श्रेणी 2 के स्थानीय आपूर्तिकर्ता के रूप में अर्हता प्राप्त करने के लिए आवश्यक/Minimum 50% and 26% Local Content required for qualifying as Class 1 and Class 2 Local Supplier respectively)

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)
PRODUCT INFORMATION	<b>Product Description</b>	Tubal Rings
	<b>Conformity to technical specifications</b>	As per detailed technical specification document attached/uploaded
	Attached/uploaded technical specification has been seen, read and understood	Yes
CERTIFICATIONS	Compliance to Medical Device Rules (MDR) 2017 as amended till date	Yes
	Availability of valid medical device manufacturing license for the product issued from the competent authority defined under Drugs and Cosmetic Act 1940 and Rules made there under as amended till date	Yes
	Manufacturing Facility Certification	ISO:13485 (Latest)
	Availability of Test Report for each supplied batch/product as per Medical Device Rules (MDR) 2017 as amended till date	Yes
	Submission of all necessary certifications, licenses and test reports to the buyer at the time of bid submission and/or along with supplies as per buyer requirement	Yes

## अतिरिक्त विशिष्टि दस्तावेज़ /Additional Specification Documents

Applicable Specification Document	<a href="#">View</a>
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### 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	Anurodh Singh	110021,Second Floor , Viswa Yuvak Kendra , Chanakyapuri New Delhi -110021	558000	180

### Special terms and conditions-Version:1 effective from 29-07-2024 for category Condoms (Free Supply) for Family Planning Programme

1. All Provisions of Drugs and Cosmetics Act, 1940 and Rules (including Medical Device Rule 2017) made there under as amended till date will always be applicable. This will include all notifications issued by Central Drugs Standard Control Organization (CDSCO), Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers time to time in this regard.
2. The sellers are registered on GeM based on self-declaration of valid Medical Device License, product certification, test reports etc. However, buyers must check and validate the details at their end for all applicable licenses and certifications e.g., validity and authenticity/genuineness of license, product certification, manufacturer certification/licenses, test reports etc.
3. In case of authorized resellers/distributors, it will be the legal & regulatory liability of the manufacturer to ensure that their resellers/distributors are operating in compliance with all relevant laws and regulations and are properly licensed to sell the manufacturer's products, including verifying the validity and authenticity of Medical Device License license held by them.
4. The price offered by the seller/bidder shall not, in any case exceed the DPCO/NPPA controlled price or price fixed by State Government, if any. The seller must reduce the prices if there is any reduction in DPCO/NPPA ceiling price or price fixed by State Government, if any.
5. Any other Terms and Conditions which is not included or at variance with the conditions specified in STC/GTC, may be added by the buyer through Additional Terms and Conditions (ATC) in the bid to ensure items are procured from authentic/validated source with appropriate and applicable quality. The above terms and conditions are in reverse order of precedence i.e. ATC shall supersede specific STC which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

### Special terms and conditions-Version:1 effective from 29-07-2024 for category Emergency Contraceptives Pills (ECP) for Family Planning Programme

1. The sellers are registered on GeM and exempted from the Vendor Assessment process based on the undertaking & submitted copy of a valid Manufacturing Drug License 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.
2. The Buyer shall ask the seller to submit the "Notarized Undertaking" in the mentioned below format (scanned copy and hard copy). Details of the same may be verified by the buyer at their end.

#### UNDERTAKING

**(to be on non-judicial stamp paper of Rs 10 and notarized)**

I, \_\_\_\_\_, s/o / d/o / w/o \_\_\_\_\_, aged about \_\_\_\_ resident of \_\_\_\_\_, do hereby declare and undertake that;

1. I am the partner / proprietor / director of \_\_\_\_\_ (*name of entity*) and duly authorized to sign this undertaking on behalf of \_\_\_\_\_. (*Name of entity*)
2. We are the manufacturers of the drug/medicine \_\_\_\_\_ (“Product”) and intend to offer the same for sale through the GeM portal.
3. We state that the license for the Product has been granted/obtained by us as per the provisions of the Drug & Cosmetics Act, 1940 and rules framed there under as amended till date.
4. We further state that the details regarding the Product/licenses have been uploaded by us on the online ‘SUGAM’ portal of CDSCO as per rule 84AB of the Drugs and Cosmetics Rules, 1945 as amended till date. Reference no. for SUGAM portal is \_\_\_\_\_.
5. We undertake that all the information provided above is true and complete in all respect. We understand that in the event any false information/declaration is provided by us, suitable legal action/action as per Drugs and Cosmetics Act, 1940 as amended till date and rules made there under will be initiated.

Place:

Date:

.....

*Signature, Name, Designation & Seal*

*on behalf of the Manufacturer*

3. All Provisions of Drugs and Cosmetics Act, 1940 and Rules made there under as amended till date will always be applicable. This will include all notifications issued by *Central Drugs Standard Control Organization (CDSCO)*, Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers time to time in this regard.
4. All provisions of Narcotic Drugs & Psychotropic Substances Act, 1985 as amended till date will also be applicable in case of Narcotic Drugs & Psychotropic Substances.
5. The purchase shall be made through Bidding/RA only irrespective of the value.
6. 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. The Drug/medicine quoted should be clearly marked and highlighted in Drug Manufacturing License. The valid own manufacturing license shall be submitted to the buyer at the time of bid submission/delivery as per buyer requirement.

In case of authorized resellers/distributors, it will be the legal & regulatory liability of the manufacturer to ensure that their authorized resellers/distributors are operating in compliance with all relevant laws and regulations and are properly licensed to sell the manufacturer's products.

Manufacturer shall be responsible for verifying the validity and authenticity of drug license held by their authorized resellers/distributors.

*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.*

7. Bidder/Seller shall submit the valid GMP/WHO-GMP Certificate of the manufacturing site as per revised Schedule-‘M’ for the quoted drugs/medicines issued by the Concerned Drug Licensing Authority to the buyer at the time of bid submission.
8. Bidder/Seller shall submit a valid **non-Conviction** certificate for last two (2) consecutive years issued by the Concerned Drug Licensing authority to the buyer at the time of bid submission. The certificate must have been issued within 12 months from the date of technical opening of the bid.
9. Bidder/Seller shall submit **Manufacturing & Market Standing certificate** (in India) issued by the concerned drug licensing authority for at least latest 2 consecutive years for the drug/medicine quoted to the buyer at the time of bid submission. The drug/medicine quoted should be clearly marked and highlighted.

*This would not apply to drugs, which were introduced in India less than 2 years ago. A certificate from the Drug Controller General (India) shall be required for all new drug formulations to this effect.*

10. If a company/firm has two or more separate manufacturing units at different sites / States/region, which are not separate entities then the company will be allowed to submit only one bid for all units but necessary document regarding separate manufacturing units will be submitted to the buyer. However, one bidder will be allowed to submit only one offer for one product.
11. The manufacturer shall have in house testing facilities and valid Good Laboratory Practice (GLP Certificate) as per Schedule "L1" of the Drugs & Cosmetics Act and Rules made thereunder as amended up to date issued by Central / State Drug Controller / FDA for the quoted drugs/medicines (as applicable).
12. Bidder/Seller shall have Maximum Production Capacity Certificate (section wise) issued by concerned drug licensing authority highlighting the quoted product.
13. STP (Standard Testing Procedure) along with the required reference standards for non-Pharmacopoeia drugs/medicines are required to be submitted by the bidder/seller at the time of submission of the bid.
14. The bidder/seller shall submit complete stability data (long term stability studies and accelerated stability studies) for all drugs/medicines in specified packing for at least 3 batches whenever required by the buyer. For New drugs/medicines, complete stability data of 6 months period shall be acceptable. (If manufacturer has licensed a formula from another company and such licensed formula is used for the product, then the stability data of the licensor should be submitted along with licensing agreement.)
15. The bidder/seller should have not been blacklisted/debarred/de-registered/banned for the quoted product by any State Government / Central Government / Central or State Government's Drug procurement agencies at the time of submission of bid. Further, quoted drugs / medicines have not been failed in house testing or testing by any State Government / Central Government / its Drug procurement agencies during last two years. If any bidder/seller has been blacklisted / debarred / de-registered/banned due to quality failure, such bidder/seller or their Partner / Director / Owner shall not be permitted to participate in the bid.
16. During the validity of the bid if the firm/Company is blacklisted/debarred/de-registered/banned by any State Government / Central Government/Central or State Government's Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to the buyer along with relevant authentic document by the bidder/seller firm/ company within one month.
17. During Contract period, if the supplier is debarred/deregistered /blacklisted/ banned by any Central Government or State Government or its procurement agencies due to quality failure, buyer may cancel the contract and go for fresh bid as per discretion of the buyer.
18. The firm/company/ corporation and any of its director/proprietors/ partners/ Authorized signatories should not be convicted/ or a criminal case filed against or pending in any court of India by any department of Govt. under prevention of Corruption Act or for cheating/ defrauding Government/ embezzlement of Government fund or any criminal conspiracy in the said matter at the time of submission of bid.
19. Bidder/seller should submit a notarized undertaking on an affidavit of Rs. 100/- (Rupees One Hundred only) duly attested by the Notary stating that:

*They will comply with all the statues & legislation regarding manufacturing, import, sale, and supply of drugs in India and in particular the following Acts/Enactments viz., The Drugs and Cosmetics Act, 1940, The Drugs and Cosmetics Rules, 1945 (as amended), The Legal Metrology Act, 2009, The Drugs (Control) Act, 1950, The Indian Statistical Institute Act, 1959, GST Act.*

*To supply drugs of standard quality as prescribed under the provisions of Drug and Cosmetic Act, 1940 (as amended). The bidder shall also undertake not to supply items/drugs "not of standard", "Grossly sub-standard" and "Spurious and adulterated drugs" as per the latest guidelines issued by the Drug Controller of India from time to time.*

20. The price offered by the seller/bidder shall not, in any case, exceed the DPCO/NPPA controlled price or price fixed by the State Government, if any. The seller must reduce the prices if there is any reduction in DPCO/NPPA ceiling price or price fixed by State Government, if any.
21. Guidelines of Department of Pharmaceuticals applicable as nodal ministry for implementing the provision of Public Procurement (Preference to Make in India) order (PPO) 2017-revision as amended to date, related to procurement of Goods & Services in Pharmaceutical Formulations, will be applicable.
22. **Fall Clause:** Provision of fall clause will not be applicable on the sale of drugs which have an expiry date as per Ministry of Chemical and Fertilizer OM No 31026/1/2019-Policy dated 12-9-2020.
23. **Shelf Life:** Shelf life of each quoted drugs/medicines shall be in accordance with Schedule P of

Drugs and Cosmetics Rules, 1945 as amended up to date.

In case pre-dispatch inspection is not applicable, the life of the drugs/medicines shall not have passed more than one sixth (1/6th) of the total shelf life of the drug/medicine at the time of delivery to the consignee.

In case of pre-dispatch inspection, at the time when the stores are offered for inspection, the life of the drugs/medicines shall not have passed more than one sixth (1/6th) of the total shelf life of the drugs/medicines.

#### 24. **Recalls**

If products are recalled because of problems with product quality or adverse reaction to the pharmaceutical, the supplier will be obliged to notify the buyer, providing full details about the reason leading to the recall, and shall take steps to replace the product in question at suppliers own cost at the ultimate destination with a fresh batch of acceptable pharmaceuticals or withdraw and give a full refund of the unconsumed quantity if the product has been taken off the market due to safety problems.

#### 25. **Inspection, Testing and Quality Control**

- All the batches of the drugs/medicines supplied shall be accompanied with in-house **Test Report/Certificate of Analysis** from the manufacturer's own Quality Control Lab. The Test Report/Certificate of Analysis shall include:
  1. Generic name of the product
  2. Batch No.
  3. Pharmacopoeia Reference and/ or In-house method
  4. Batch quantity
  5. Date of manufacture
  6. Expiry date
  7. Date of test
  8. Description (clarity, color etc)
  9. All identity, potency, purity, sterility, pyrogen and all other test required by the specified pharmacopoeia and/or In-house method. Both the actual results and the limits for the individual tests should be given
  10. Conclusion
  11. Qualified Person's signature

The above-mentioned batch shall be manufactured in accordance with the applicable GMP regulations.

- Buyer will embark on stringent quality checks to ensure that drugs/medicines/goods meet required standards throughout specified shelf life. The buyer reserves the right to carry necessary inspections/tests from NABL Accredited/Government approved laboratories at any of, or any combination of or/ all following stages:

##### **a) At Pre-Dispatch stage**

**b) At Delivery Stage:** Inspection done once the drugs/medicines/goods reach at consignee location and before taking over supplied goods in inventory.

**c) Post Delivery Surveillance:** The Drugs/Medicines/goods shall have the active ingredients and all other parameters at the prescribed level as indicated in official compendiums or technical specifications throughout the shelf-life period of the drugs/medicines/ goods. Quality Monitoring Activities may also be organized by the buyer post-delivery.

- The Buyer may engage the services of a Quality Control Agent & Quality Control Testing Laboratories for the purpose of Inspection & Quality Control. The sampling quantities shall be borne by the supplier.
- The buyer's right to inspect, test and, where necessary reject the drugs/medicines/goods after the drugs/medicines/goods arrival at the final destination shall in no way be limited or waived by reason of the goods having previously been inspected, tested and passed prior to the goods dispatch from the place of manufacture.
- **Inspection Methodology:** At pre-dispatch and/or delivery stage, samples of supplies in each batch will be chosen for testing. The samples will be collected and sent to designated laboratories (NABL Accredited/Government approved laboratories) for testing as decided by the buyer.

**At post-delivery surveillance** - The samples will be collected from the warehouse of buyer/or final consignee in States/UTs and sent to designated Quality Control Labs in respect of supplied drugs/medicines/goods at any point during specified shelf life as per decision of the buyer.

*Handling and testing charges will be borne by the buyer for the above purpose.*

- In case of failure of batches during or at any stage (indicated above), the testing charges would be claimed for the defaulting vendor.
- The supplies will be deemed to be completed only upon receipt of the quality certificates from the laboratories.

“Not of Standard Quality” or spurious or adulterated or misbranded, such batch/ batches will be deemed to be rejected goods.”

- **At any of testing stage**, Samples which do not meet quality requirement shall render the relevant batches liable to be rejected. If the sample is declared to be “Not of Standard Quality” or spurious or adulterated or misbranded, such batch/batches will be deemed to be rejected drugs/medicines/goods and the cost of entire batch paid will be recovered from the supplier whether consumed fully/partially.
- If any inspected or tested drugs/medicines/goods fails to conform to the specifications or fails in quality tests, the buyer may reject them, and the supplier will be responsible to take back the rejected drugs/medicines/goods from the depots/consignee place at their own cost and replace with fresh stock duly inspected and tested within 45 days from the date of intimation from the buyer to the supplier or as specified by the buyer. The buyer has the right to destroy such rejected drugs/medicines/goods if the supplier does not take back the drugs/medicines/goods within the stipulated time. The buyer will arrange to destroy the “NOT OF STANDARD QUALITY ITEMS” after the expiry of 45 days.

Action may also be initiated by the buyer for debarring/blacklisting against the supplier for suitable period. The FDA/ Drugs Control Authority of concerned State will also be informed by the buyer for initiating necessary action on the supplier in their state. In addition, Security deposit will also be forfeited without any intimation.

The decision of the buyer or any officer authorized by buyer, as to the quality of the supplied drugs, medicines etc., shall be final and binding.

- In case any drug/medicine is found substandard either any of testing stage or during the shelf life of the item, the report of the NABL/Government approved laboratory shall be accepted by the supplier/seller. If the same is disputed by the supplier/seller giving the reasons, the sample will be sent to Central Drug Laboratory, Kolkata, and the report of CDL will only be accepted as final and conclusive report. However, the same should be submitted within three months, from the date of communication of the disputed test report to the supplier/seller. For this, supplier/seller should approach the concerned Drug Control Authorities for getting the drugs/medicines tested, as per procedure, from the Appellate Laboratory at their own cost.

The de-registration / debarment action will be taken by the buyer against the manufacturing unit and contract holding firms (both) according to the category-A and category-B defects as per guidelines issued by the Ministry of Health & Family Welfare, Government of India and same will be communicated to GeM.

- The supplier shall furnish evidence of the basis for shelf life and other stability data concerning the commercial final package on request by the buyer. In case of any complaint in the field, the B.M.R/ B.P.R for the particular batch of the product(s) supplied shall be produced when demanded. For New drugs/medicines, complete stability data of 6 months period shall be acceptable.
- The case of admixture of drugs will be treated as a violation of terms and conditions and will not be accepted.
- Statutory provisions on manufacture, distribution, storage and quality issues of drugs/medicines under the Drugs & Cosmetic Act, 1940 as amended up to date is vested with the DCGI (CDSCO)/ MoH& FW, including its Central/ Zonal/ Regional Drug Testing Labs. Similarly, the authority for confiscation, sealing or prosecution with relation to drugs/medicines under the said Act is also with the DCGI/ State Drug Control Authorities.
- In accordance with the provisions of Sec 22 & 31 of the Drugs and Cosmetic Act, 1940, as amended up to date, the DCGI/ CDSCO/ State Drug Controller/ Drug Inspector may exercise their powers as an Inspecting Agency.

26. **Deduction, Blacklisting, and other penalties on account of Quality failure**

The suitable conditions may be added by the buyer in the bid through Additional Term & Conditions (ATC) as applicable.

27. **Quality Test by Statutory Authorities:**

If any batch of any product(s) supplied by the supplier is declared "NOT OF STANDARD QUALITY", by any government agencies or drug licensing authority, the supplier shall inform the same immediately to the buyer so that the use of the available stock of the particular item will be stopped. Further, the available stock of the product with all consignee/users will be retrieved.

28. **Termination for Default**

The buyer may without prejudice to any other remedy for breach of contract, by written notice of default sent to the supplier, terminate the contract in whole or in part If the supplier fails to promptly replace any drug/medicine/goods rejected submitted for testing or subject to recall ordered by the applicable Regulatory Authority in the country of manufacture due to unacceptable quality or reports of adverse drugs reaction after giving prompt notice of the recall.

29. **Warranty**

- Supplies must fully comply in all respect with the Technical specifications and conditions laid down in the contract and in accordance with the Pharmacopoeia standards.
- Each supply should be accompanied with a "Warranty Certificate" duly signed by the Bidder as under:

"The Supplier/Seller hereby declares that the stores as detailed below sold to the buyer under this contract shall be of the best quality and workmanship and shall be strictly in accordance with the specifications and particulars mentioned and the supplier/seller hereby guarantees that the stores would continue to conform to the description of and quality aforesaid for a period of useful life of minimum of five sixth (5/6th) of the specified shelf life from the date of delivery of the said stores to the buyer, have overages within the ranges set forth in the technical specification and are not subject to recall by the applicable Regulatory Authority due to unacceptable quality or adverse Pharmaceuticals reaction. Notwithstanding the above, the fact that the said stores fail to conform to the description and quality aforesaid or have deteriorated and the decision of the buyer in that behalf is final and conclusive, the buyer will be entitled to reject the said stores, or such part thereof as may be discovered not to conform to the said description and quality. Losses due to premature deterioration due to biological and other activities during life potency will be made good and supplied by the firm at its own cost at consignee's site.

On such rejection, the stores will be at the seller's risk and all provisions herein contained relating to the rejection of stores shall apply. The supplier/Seller shall if so called upon to do so by the buyer in writing, replace the stores free of cost at the ultimate destination within a period of forty five days or such further period as may be extended from time to time by the buyer at his discretion, on application made there under by the supplier/seller after the stores or such portion of the stores thereof as is rejected by the buyer and in such an event the above mentioned warranty period shall apply to the stores replaced from the date of the replacement thereof otherwise the supplier/seller shall pay to the buyer such damage as may arise by reason of the breach of the conditions. Nothing herein contained shall prejudice any other rights of the buyer in that behalf under this contract or otherwise".

- Sl. No. & Date
- Nomenclature & Specification
- Name & Address of Manufacturing Unit
- Batch No.
- DOM & DOE
- Qty. of each batch
- Remarks

designation and date with rubber stamp

Signature name &

- If the supplier, having been notified, fails to replace within the period specified above, the buyer may proceed to take such remedial action(s) as may be necessary/deemed fit by the

buyer, at the suppliers' risk and expense and without prejudice to other rights which the buyer may have against the supplier under the contract.

30. **Packaging, Labelling and Marking Requirements**

Packaging, Labelling and Marking shall be as per the provisions contained in the Drugs and Cosmetics Act, 1940 and the Rules made there under as amended up-to-date, other particulars of packaging, labelling & marking, if any, prescribed by the buyer in the bid through Additional Terms and Conditions (ATC) shall be complied with.

31. **Bar Coding**

All drugs/medicines supplied should incorporate GS1 barcodes standards at various packaging levels (primary, secondary, and tertiary level packaging) and should encode the information within the barcodes as mentioned by the buyers in addition to other existing statutory labelling and marking requirements. Details of bar-coding will be given by the buyer through Additional Terms and Conditions (ATC) in the bid.

32. **Delivery Period**

- Minimum delivery period will be of 45 days from the date of issuing of the purchase
- The supplier should maintain the recommended temperature of the drug/medicine (wherever indicated) during transit, otherwise if on checking it is found that temperature has not been maintained, supply against the said order is liable to be rejected and cancelled.
- The items requiring special cold storage conditions shall be supplied with cold chain transporting system under cold chain norms from the manufacturing unit to the warehouses/consignee location.

33. Any specific requirements for the packaging, labelling, logograms, printing, artwork, bar coding or any other requirement mentioned by the buyer through Additional Terms and Conditions (ATC) in the bid will be applicable.

34. 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:1 effective from 29-07-2024 for category Injectable Contraceptive (Antara) for Family Planning Programme**

1. The sellers are registered on GeM and exempted from the Vendor Assessment process based on the undertaking & submitted copy of a valid Manufacturing Drug License 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.
2. The Buyer shall ask the seller to submit the "Notarized Undertaking" in the mentioned below format (scanned copy and hard copy). Details of the same may be verified by the buyer at their end.

**UNDERTAKING**

***(to be on non-judicial stamp paper of Rs 10 and notarized)***

I, \_\_\_\_\_, s/o / d/o / w/o \_\_\_\_\_, aged about \_\_\_\_ resident of \_\_\_\_\_, do hereby declare and undertake that;

1. I am the partner / proprietor / director of \_\_\_\_\_ (name of entity) and duly authorized to sign this undertaking on behalf of \_\_\_\_\_. (Name of entity)
2. We are the manufacturers of the drug/medicine \_\_\_\_\_ ("Product") and intend to offer the same for sale through the GeM portal.
3. We state that the license for the Product has been granted/obtained by us as per the provisions of the Drug & Cosmetics Act, 1940 and rules framed there under as amended till date.
4. We further state that the details regarding the Product/licenses have been uploaded by us on the online 'SUGAM' portal of CDSCO as per rule 84AB of the Drugs and Cosmetics Rules, 1945

- as amended till date. Reference no. for SUGAM portal is \_\_\_\_\_.
5. We undertake that all the information provided above is true and complete in all respect. We understand that in the event any false information/declaration is provided by us, suitable legal action/action as per Drugs and Cosmetics Act, 1940 as amended till date and rules made there under will be initiated.

Place:

Date:

.....

*Signature, Name, Designation & Seal*

*on behalf of the Manufacturer*

3. All Provisions of Drugs and Cosmetics Act, 1940 and Rules made there under as amended till date will always be applicable. This will include all notifications issued by *Central Drugs Standard Control Organization (CDSCO)*, Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers time to time in this regard.
4. All provisions of Narcotic Drugs & Psychotropic Substances Act, 1985 as amended till date will also be applicable in case of Narcotic Drugs & Psychotropic Substances.
5. The purchase shall be made through Bidding/RA only irrespective of the value.
6. 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. The Drug/medicine quoted should be clearly marked and highlighted in Drug Manufacturing License. The valid own manufacturing license shall be submitted to the buyer at the time of bid submission/delivery as per buyer requirement.

In case of authorized resellers/distributors, it will be the legal & regulatory liability of the manufacturer to ensure that their authorized resellers/distributors are operating in compliance with all relevant laws and regulations and are properly licensed to sell the manufacturer's products.

Manufacturer shall be responsible for verifying the validity and authenticity of drug license held by their authorized resellers/distributors.

*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.*

7. Bidder/Seller shall submit the valid GMP/WHO-GMP Certificate of the manufacturing site as per revised Schedule-'M' for the quoted drugs/medicines issued by the Concerned Drug Licensing Authority to the buyer at the time of bid submission.
8. Bidder/Seller shall submit a valid **non-Conviction** certificate for last two (2) consecutive years issued by the Concerned Drug Licensing authority to the buyer at the time of bid submission. The certificate must have been issued within 12 months from the date of technical opening of the bid.
9. Bidder/Seller shall submit **Manufacturing & Market Standing certificate** (in India) issued by the concerned drug licensing authority for at least latest 2 consecutive years for the drug/medicine quoted to the buyer at the time of bid submission. The drug/medicine quoted should be clearly marked and highlighted.

*This would not apply to drugs, which were introduced in India less than 2 years ago. A certificate from the Drug Controller General (India) shall be required for all new drug formulations to this effect.*

10. If a company/firm has two or more separate manufacturing units at different sites / States/region, which are not separate entities then the company will be allowed to submit only one bid for all units but necessary document regarding separate manufacturing units will be submitted to the buyer. However, one bidder will be allowed to submit only one offer for one product.
11. The manufacturer shall have in house testing facilities and valid Good Laboratory Practice (GLP Certificate) as per Schedule "L1" of the Drugs & Cosmetics Act and Rules made thereunder as amended up to date issued by Central / State Drug Controller / FDA for the quoted drugs/medicines (as applicable).
12. Bidder/Seller shall have Maximum Production Capacity Certificate (section wise) issued by

concerned drug licensing authority highlighting the quoted product.

13. STP (Standard Testing Procedure) along with the required reference standards for non-Pharmacopoeia drugs/medicines are required to be submitted by the bidder/seller at the time of submission of the bid.
14. The bidder/seller shall submit complete stability data (long term stability studies and accelerated stability studies) for all drugs/medicines in specified packing for at least 3 batches whenever required by the buyer. For New drugs/medicines, complete stability data of 6 months period shall be acceptable. (If manufacturer has licensed a formula from another company and such licensed formula is used for the product, then the stability data of the licensor should be submitted along with licensing agreement.)
15. The bidder/seller should have not been blacklisted/debarred/de-registered/banned for the quoted product by any State Government / Central Government / Central or State Government's Drug procurement agencies at the time of submission of bid. Further, quoted drugs / medicines have not been failed in house testing or testing by any State Government / Central Government / its Drug procurement agencies during last two years. If any bidder/seller has been blacklisted / debarred / de-registered/banned due to quality failure, such bidder/seller or their Partner / Director / Owner shall not be permitted to participate in the bid.
16. During the validity of the bid if the firm/Company is blacklisted/debarred/de-registered/banned by any State Government / Central Government/Central or State Government's Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to the buyer along with relevant authentic document by the bidder/seller firm/ company within one month.
17. During Contract period, if the supplier is debarred/deregistered /blacklisted/ banned by any Central Government or State Government or its procurement agencies due to quality failure, buyer may cancel the contract and go for fresh bid as per discretion of the buyer.
18. The firm/company/ corporation and any of its director/proprietors/ partners/ Authorized signatories should not be convicted/ or a criminal case filed against or pending in any court of India by any department of Govt. under prevention of Corruption Act or for cheating/ defrauding Government/ embezzlement of Government fund or any criminal conspiracy in the said matter at the time of submission of bid.
19. Bidder/seller should submit a notarized undertaking on an affidavit of Rs. 100/- (Rupees One Hundred only) duly attested by the Notary stating that:

*They will comply with all the statues & legislation regarding manufacturing, import, sale, and supply of drugs in India and in particular the following Acts/Enactments viz., The Drugs and Cosmetics Act, 1940, The Drugs and Cosmetics Rules, 1945 (as amended), The Legal Metrology Act, 2009, The Drugs (Control) Act, 1950, The Indian Statistical Institute Act, 1959, GST Act.*

*To supply drugs of standard quality as prescribed under the provisions of Drug and Cosmetic Act, 1940 (as amended). The bidder shall also undertake not to supply items/drugs "not of standard", "Grossly sub-standard" and "Spurious and adulterated drugs" as per the latest guidelines issued by the Drug Controller of India from time to time.*

20. The price offered by the seller/bidder shall not, in any case, exceed the DPCO/NPPA controlled price or price fixed by the State Government, if any. The seller must reduce the prices if there is any reduction in DPCO/NPPA ceiling price or price fixed by State Government, if any.
21. Guidelines of Department of Pharmaceuticals applicable as nodal ministry for implementing the provision of Public Procurement (Preference to Make in India) order (PPO) 2017-revision as amended to date, related to procurement of Goods & Services in Pharmaceutical Formulations, will be applicable.
22. **Fall Clause:** Provision of fall clause will not be applicable on the sale of drugs which have an expiry date as per Ministry of Chemical and Fertilizer OM No 31026/1/2019-Policy dated 12-9-2020.
23. **Shelf Life:** Shelf life of each quoted drugs/medicines shall be in accordance with Schedule P of Drugs and Cosmetics Rules, 1945 as amended up to date.

In case pre-dispatch inspection is not applicable, the life of the drugs/medicines shall not have passed more than one sixth (1/6th) of the total shelf life of the drug/medicine at the time of delivery to the consignee.

In case of pre-dispatch inspection, at the time when the stores are offered for inspection, the life of the drugs/medicines shall not have passed more than one sixth (1/6th) of the total shelf life of the drugs/medicines.

24. **Recalls**

If products are recalled because of problems with product quality or adverse reaction to the pharmaceutical, the supplier will be obliged to notify the buyer, providing full details about the

reason leading to the recall, and shall take steps to replace the product in question at suppliers own cost at the ultimate destination with a fresh batch of acceptable pharmaceuticals or withdraw and give a full refund of the unconsumed quantity if the product has been taken off the market due to safety problems.

25. **Inspection, Testing and Quality Control**

- All the batches of the drugs/medicines supplied shall be accompanied with in-house **Test Report/Certificate of Analysis** from the manufacturer's own Quality Control Lab. The Test Report/Certificate of Analysis shall include:
  1. Generic name of the product
  2. Batch No.
  3. Pharmacopoeia Reference and/ or In-house method
  4. Batch quantity
  5. Date of manufacture
  6. Expiry date
  7. Date of test
  8. Description (clarity, color etc)
  9. All identity, potency, purity, sterility, pyrogen and all other test required by the specified pharmacopoeia and/or In-house method. Both the actual results and the limits for the individual tests should be given
  10. Conclusion
  11. Qualified Person's signature

The above-mentioned batch shall be manufactured in accordance with the applicable GMP regulations.

- Buyer will embark on stringent quality checks to ensure that drugs/medicines/goods meet required standards throughout specified shelf life. The buyer reserves the right to carry necessary inspections/tests from NABL Accredited/Government approved laboratories at any of, or any combination of or/ all following stages:

**a) At Pre-Dispatch stage**

**b) At Delivery Stage:** Inspection done once the drugs/medicines/goods reach at consignee location and before taking over supplied goods in inventory.

**c) Post Delivery Surveillance:** The Drugs/Medicines/goods shall have the active ingredients and all other parameters at the prescribed level as indicated in official compendiums or technical specifications throughout the shelf-life period of the drugs/medicines/ goods. Quality Monitoring Activities may also be organized by the buyer post-delivery.

- The Buyer may engage the services of a Quality Control Agent & Quality Control Testing Laboratories for the purpose of Inspection & Quality Control. The sampling quantities shall be borne by the supplier.
- The buyer's right to inspect, test and, where necessary reject the drugs/medicines/goods after the drugs/medicines/goods arrival at the final destination shall in no way be limited or waived by reason of the goods having previously been inspected, tested and passed prior to the goods dispatch from the place of manufacture.
- **Inspection Methodology:** At pre-dispatch and/or delivery stage, samples of supplies in each batch will be chosen for testing. The samples will be collected and sent to designated laboratories (NABL Accredited/Government approved laboratories) for testing as decided by the buyer.

**At post-delivery surveillance** - The samples will be collected from the warehouse of buyer/or final consignee in States/UTs and sent to designated Quality Control Labs in respect of supplied drugs/medicines/goods at any point during specified shelf life as per decision of the buyer.

*Handling and testing charges will be borne by the buyer for the above purpose.*

- In case of failure of batches during or at any stage (indicated above), the testing charges would be claimed for the defaulting vendor.
- The supplies will be deemed to be completed only upon receipt of the quality certificates from the laboratories.

"Not of Standard Quality" or spurious or adulterated or misbranded, such batch/ batches will

be deemed to be rejected goods.”

- **At any of testing stage**, Samples which do not meet quality requirement shall render the relevant batches liable to be rejected. If the sample is declared to be “Not of Standard Quality” or spurious or adulterated or misbranded, such batch/batches will be deemed to be rejected drugs/medicines/goods and the cost of entire batch paid will be recovered from the supplier whether consumed fully/partially.
- If any inspected or tested drugs/medicines/goods fails to conform to the specifications or fails in quality tests, the buyer may reject them, and the supplier will be responsible to take back the rejected drugs/medicines/goods from the depots/consignee place at their own cost and replace with fresh stock duly inspected and tested within 45 days from the date of intimation from the buyer to the supplier or as specified by the buyer. The buyer has the right to destroy such rejected drugs/medicines/goods if the supplier does not take back the drugs/medicines/goods within the stipulated time. The buyer will arrange to destroy the “NOT OF STANDARD QUALITY ITEMS” after the expiry of 45 days.

Action may also be initiated by the buyer for debarring/blacklisting against the supplier for suitable period. The FDA/ Drugs Control Authority of concerned State will also be informed by the buyer for initiating necessary action on the supplier in their state. In addition, Security deposit will also be forfeited without any intimation.

The decision of the buyer or any officer authorized by buyer, as to the quality of the supplied drugs, medicines etc., shall be final and binding.

- In case any drug/medicine is found substandard either any of testing stage or during the shelf life of the item, the report of the NABL/Government approved laboratory shall be accepted by the supplier/seller. If the same is disputed by the supplier/seller giving the reasons, the sample will be sent to Central Drug Laboratory, Kolkata, and the report of CDL will only be accepted as final and conclusive report. However, the same should be submitted within three months, from the date of communication of the disputed test report to the supplier/seller. For this, supplier/seller should approach the concerned Drug Control Authorities for getting the drugs/medicines tested, as per procedure, from the Appellate Laboratory at their own cost.

The de-registration / debarment action will be taken by the buyer against the manufacturing unit and contract holding firms (both) according to the category-A and category-B defects as per guidelines issued by the Ministry of Health & Family Welfare, Government of India and same will be communicated to GeM.

- The supplier shall furnish evidence of the basis for shelf life and other stability data concerning the commercial final package on request by the buyer. In case of any complaint in the field, the B.M.R/ B.P.R for the particular batch of the product(s) supplied shall be produced when demanded. For New drugs/medicines, complete stability data of 6 months period shall be acceptable.
- The case of admixture of drugs will be treated as a violation of terms and conditions and will not be accepted.
- Statutory provisions on manufacture, distribution, storage and quality issues of drugs/medicines under the Drugs & Cosmetic Act, 1940 as amended up to date is vested with the DCGI (CDSCO)/ MoH& FW, including its Central/ Zonal/ Regional Drug Testing Labs. Similarly, the authority for confiscation, sealing or prosecution with relation to drugs/medicines under the said Act is also with the DCGI/ State Drug Control Authorities.
- In accordance with the provisions of Sec 22 & 31 of the Drugs and Cosmetic Act, 1940, as amended up to date, the DCGI/ CDSCO/ State Drug Controller/ Drug Inspector may exercise their powers as an Inspecting Agency.

#### 26. **Deduction, Blacklisting, and other penalties on account of Quality failure**

The suitable conditions may be added by the buyer in the bid through Additional Term & Conditions (ATC) as applicable.

#### 27. **Quality Test by Statutory Authorities:**

If any batch of any product(s) supplied by the supplier is declared “NOT OF STANDARD QUALITY”, by any government agencies or drug licensing authority, the supplier shall inform the same immediately to the buyer so that the use of the available stock of the particular item will be stopped. Further, the available stock of the product with all consignee/users will be retrieved.

#### 28. **Termination for Default**

The buyer may without prejudice to any other remedy for breach of contract, by written notice of default sent to the supplier, terminate the contract in whole or in part If the supplier fails to promptly replace any drug/medicine/goods rejected submitted for testing or subject to recall ordered by the applicable Regulatory Authority in the country of manufacture due to unacceptable quality or reports of adverse drugs reaction after giving prompt notice of the recall.

29. **Warranty**

- Supplies must fully comply in all respect with the Technical specifications and conditions laid down in the contract and in accordance with the Pharmacopoeia standards.
- Each supply should be accompanied with a "Warranty Certificate" duly signed by the Bidder as under:

“The Supplier/Seller hereby declares that the stores as detailed below sold to the buyer under this contract shall be of the best quality and workmanship and shall be strictly in accordance with the specifications and particulars mentioned and the supplier/seller hereby guarantees that the stores would continue to conform to the description of and quality aforesaid for a period of useful life of minimum of five sixth (5/6th) of the specified shelf life from the date of delivery of the said stores to the buyer, have overages within the ranges set forth in the technical specification and are not subject to recall by the applicable Regulatory Authority due to unacceptable quality or adverse Pharmaceuticals reaction. Notwithstanding the above, the fact that the said stores fail to conform to the description and quality aforesaid or have deteriorated and the decision of the buyer in that behalf is final and conclusive, the buyer will be entitled to reject the said stores, or such part thereof as may be discovered not to conform to the said description and quality. Losses due to premature deterioration due to biological and other activities during life potency will be made good and supplied by the firm at its own cost at consignee’s site.

On such rejection, the stores will be at the seller's risk and all provisions herein contained relating to the rejection of stores shall apply. The supplier/Seller shall if so called upon to do so by the buyer in writing, replace the stores free of cost at the ultimate destination within a period of forty five days or such further period as may be extended from time to time by the buyer at his discretion, on application made there under by the supplier/seller after the stores or such portion of the stores thereof as is rejected by the buyer and in such an event the above mentioned warranty period shall apply to the stores replaced from the date of the replacement thereof otherwise the supplier/seller shall pay to the buyer such damage as may arise by reason of the breach of the conditions. Nothing herein contained shall prejudice any other rights of the buyer in that behalf under this contract or otherwise".

- Sl. No. & Date
- Nomenclature & Specification
- Name & Address of Manufacturing Unit
- Batch No.
- DOM & DOE
- Qty. of each batch
- Remarks

Signature name &  
designaion and date with rubber stamp

- If the supplier, having been notified, fails to replace within the period specified above, the buyer may proceed to take such remedial action(s) as may be necessary/deemed fit by the buyer, at the suppliers' risk and expense and without prejudice to other rights which the buyer may have against the supplier under the contract.

30. **Packaging, Labelling and Marking Requirements**

Packaging, Labelling and Marking shall be as per the provisions contained in the Drugs and Cosmetics Act, 1940 and the Rules made there under as amended up-to-date, other particulars of packaging, labelling & marking, if any, prescribed by the buyer in the bid through Additional Terms and Conditions (ATC) shall be complied with.

31. **Bar Coding**

All drugs/medicines supplied should incorporate GS1 barcodes standards at various packaging levels (primary, secondary, and tertiary level packaging) and should encode the information within

the barcodes as mentioned by the buyers in addition to other existing statutory labelling and marking requirements. Details of bar-coding will be given by the buyer through Additional Terms and Conditions (ATC) in the bid.

### 32. **Delivery Period**

- Minimum delivery period will be of 45 days from the date of issuing of the purchase
  - The supplier should maintain the recommended temperature of the drug/medicine (wherever indicated) during transit, otherwise if on checking it is found that temperature has not been maintained, supply against the said order is liable to be rejected and cancelled.
  - The items requiring special cold storage conditions shall be supplied with cold chain transporting system under cold chain norms from the manufacturing unit to the warehouses/consignee location.
33. Any specific requirements for the packaging, labelling, logograms, printing, artwork, bar coding or any other requirement mentioned by the buyer through Additional Terms and Conditions (ATC) in the bid will be applicable.
34. 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:1 effective from 29-07-2024 for category IUCD 380A for Family Planning Programme**

1. 1. All Provisions of Drugs and Cosmetics Act, 1940 and Rules (including Medical Device Rule 2017) made there under as amended till date will always be applicable. This will include all notifications issued by Central Drugs Standard Control Organization (CDSCO), Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers time to time in this regard.
2. The sellers are registered on GeM based on self-declaration of valid Medical Device License, product certification, test reports etc. However, buyers must check and validate the details at their end for all applicable licenses and certifications e.g., validity and authenticity/genuineness of license, product certification, manufacturer certification/licenses, test reports etc.
3. In case of authorized resellers/distributors, it will be the legal & regulatory liability of the manufacturer to ensure that their resellers/distributors are operating in compliance with all relevant laws and regulations and are properly licensed to sell the manufacturer's products, including verifying the validity and authenticity of Medical Device License license held by them.
4. The price offered by the seller/bidder shall not, in any case exceed the DPCO/NPPA controlled price or price fixed by State Government, if any. The seller must reduce the prices if there is any reduction in DPCO/NPPA ceiling price or price fixed by State Government, if any.
5. Any other Terms and Conditions which is not included or at variance with the conditions specified in STC/GTC, may be added by the buyer through Additional Terms and Conditions (ATC) in the bid to ensure items are procured from authentic/validated source with appropriate and applicable quality. The above terms and conditions are in reverse order of precedence i.e. ATC shall supersede specific STC which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

### **Special terms and conditions-Version:2 effective from 27-03-2026 for category IUCD 375 for Family Planning Programme**

1. 1. All Provisions of Drugs and Cosmetics Act, 1940 and Rules (including Medical Device Rule 2017) made there under and all subsequent amendments till date will always be applicable. This will include all notifications issued by Central Drugs Standard Control Organization (CDSCO), Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers in this regard.
2. The seller's registration and product upload on GeM is based on their declaration of possessing a valid Medical Device License for the product, certifications, test reports as per MDR (2017), latest amended. However, buyers must check, validate and verify the details at their end for all applicable licenses and certifications e.g., validity and authenticity/genuineness of medical device license (or registration), product certifications, manufacturer's certifications/ licenses, test reports at the time of supply.
3. In case of authorized resellers/distributors, it will be the legal and regulatory liability of the manufacturer to ensure that their resellers/distributors are operating in compliance with all relevant

laws and regulations, and are licensed to sell the manufacturer's products, including verifying the validity and authenticity of license held by them.

4. Any other Terms and Conditions which is not included or at variance with the conditions specified in STC/GTC, may be added by the buyer through Additional Terms and Conditions (ATC) in the bid to ensure items are procured from authentic/validated source with appropriate and applicable quality. The above terms and conditions are in reverse order of precedence i.e. BID ATC shall supersede specific STC which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.
5. **Packing and Marking:** Should be as per MDR.

### **Special terms and conditions-Version:1 effective from 29-07-2024 for category Tubal Rings for Family Planning Programme**

1. All Provisions of Drugs and Cosmetics Act, 1940 and Rules (including Medical Device Rule 2017) made there under as amended till date will always be applicable. This will include all notifications issued by Central Drugs Standard Control Organization (CDSCO), Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers time to time in this regard.
2. The sellers are registered on GeM based on self-declaration of valid Medical Device License, product certification, test reports etc. However, buyers must check and validate the details at their end for all applicable licenses and certifications e.g., validity and authenticity/genuineness of license, product certification, manufacturer certification/licenses, test reports etc.
3. In case of authorized resellers/distributors, it will be the legal & regulatory liability of the manufacturer to ensure that their resellers/distributors are operating in compliance with all relevant laws and regulations and are properly licensed to sell the manufacturer's products, including verifying the validity and authenticity of Medical Device License license held by them.
4. The price offered by the seller/bidder shall not, in any case exceed the DPCO/NPPA controlled price or price fixed by State Government, if any. The seller must reduce the prices if there is any reduction in DPCO/NPPA ceiling price or price fixed by State Government, if any.
5. Any other Terms and Conditions which is not included or at variance with the conditions specified in STC/GTC, may be added by the buyer through Additional Terms and Conditions (ATC) in the bid to ensure items are procured from authentic/validated source with appropriate and applicable quality. The above terms and conditions are in reverse order of precedence i.e. ATC shall supersede specific STC which shall supersede General Terms and Conditions (GTC), whenever there are any conflicting provisions.

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

#### **1. Buyer Added Bid Specific ATC**

Buyer uploaded ATC document [Click here to view the file.](#)

### **अस्वीकरण/Disclaimer**

The Additional Terms and Conditions (ATC) have been incorporated by the Buyer after approval of their Competent Authority. The Buyer ,is solely responsible for the impact of these clauses on the bidding process, its outcome, and consequences thereof including any restriction arising in the bidding process due to these ATCs and including the modification of technical specifications and / or terms and conditions governing the bid.All representations / grievances pertaining to the ATC clauses shall be raised with the buyer organization directly and not with GeM.If any of the clause(s) is/are incorporated by the Buyer regarding the following, the bid & resultant contract shall be treated as null & void. Further, GeM reserves the right, at its sole discretion, to cancel the bid forthwith, without issuance of any prior notice or intimation :-

1. Publishing Custom / BOQ bids for items for which regular GeM categories are available (unless such Custom / BOQ item is bunched with the major regular product Category Item).
2. Mandating procurement of / from specific Brand / Make / Model / Manufacturer / Dealer except in case of Single Bid / Proprietary Article Certificate (PAC) Buying.
3. Inclusion of disqualification criteria related to suspension of seller / service provider, where such suspension period has already expired.

4. Mandating submission of documents in physical form as a pre-requisite to qualify bidders.
5. Publishing bids on GeM for procurement of works.
6. Procurement of Goods by creating a Service bid on GeM & vice-versa.
7. Seeking sample with bid or approval of samples during bid evaluation process. However, trial / sample, as the case may be, shall be permitted in cases where trial / sample are allowed as per approved and published procurement policy of the Buyers' controlling Ministry / Department / State / Public Sector Enterprises Headquarters. If there is any violation of trial / sample clause with regard to approved policy of the Buyers' Ministry / Department / State / Public Sector Enterprises Headquarters, then this is to be determined and redressed by the concerned Buyer Organisation only.
8. Seeking experience from specific organization / department / institute only or from foreign / export experience.
9. Creating bid for items from incorrect categories.
10. Reference of conditions published on any external site or reference to external documents/clauses.
11. Asking for any Tender fee / Bid Participation fee, as the case may be.
12. Buyer added ATC Clauses which are in contravention of clauses defined in bid detail section, including specifications, EMD Detail, ePBG Detail and MII and MSE Purchase Preference sections of the bid, unless otherwise allowed by the applicable GeM GTC.
13. Any ATC clause in contravention with GeM GTC Clause 4 (xiii) (h) will be invalid. In case of multiple L1 bidders against a service bid, the buyer shall place the Contract by selection of a bidder amongst the L-1 bidders through a Random Algorithm executed by GeM system.
14. 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 catalogues 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. 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.**

**This Bid is governed by the General Terms and Conditions, conditions stipulated in Bid and Service Level Agreement specific to the Service, as the case may be, as provided in the Marketplace.**

**However, in case of Service, if any condition specified in General Terms and Conditions is contradicted by the conditions stipulated in Service Level Agreement specific to said Service, then it will over-ride the conditions in the General Terms and Conditions.**

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---**