

Bid Document

Bid Details	
Bid End Date/Time	12-12-2022 14:00:00
Bid Opening Date/Time	12-12-2022 14:30:00
Bid Offer Validity (From End Date)	90 (Days)
Ministry/State Name	Ministry Of Health And Family Welfare
Department Name	Department Of Health And Family Welfare
Organisation Name	North Eastern Indira Gandhi Regional Institute Of Health And Medical Sciences (neigrihms)
Office Name	Neigrihms, Shillong
Total Quantity	1500
Item Category	Blood Bags-IS:15102 (Q2)
Minimum Average Annual Turnover of the bidder (For 3 Years)	3 Lakh (s)
OEM Average Turnover (Last 3 Years)	3 Lakh (s)
Years of Past Experience Required for same/similar service	3 Year (s)
MSE Exemption for Years Of Experience and Turnover	Yes
Startup Exemption for Years Of Experience and Turnover	Yes
Document required from seller	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) *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
Past Performance	30 %
Bid to RA enabled	No
Type of Bid	Two Packet Bid
Time allowed for Technical Clarifications during technical evaluation	5 Days
Estimated Bid Value	500000
Evaluation Method	Total value wise evaluation

EMD Detail

Required	No
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ePBG Detail

Required	No
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Splitting

Bid splitting not applied.

MII Purchase Preference

MII Purchase Preference	Yes
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MSE Purchase Preference

MSE Purchase Preference	Yes
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1. If the bidder is a Micro or Small Enterprise as per latest definitions under MSME rules, the bidder shall be exempted from the requirement of "Bidder Turnover" criteria and "Experience Criteria" subject to meeting of quality and technical specifications. If the bidder is OEM of the offered products, it would be exempted from the "OEM Average Turnover" criteria also subject to meeting of quality and technical specifications. In case any bidder is seeking exemption from Turnover / Experience Criteria, the supporting documents to prove his eligibility for exemption must be uploaded for evaluation by the buyer.
2. If the bidder is a Startup, the bidder shall be exempted from the requirement of "Bidder Turnover" criteria and "Experience Criteria" subject to their meeting of quality and technical specifications. If the bidder is OEM of the offered products, it would be exempted from the "OEM Average Turnover" criteria also subject to meeting of quality and technical specifications. In case any bidder is seeking exemption from Turnover / Experience Criteria, the supporting documents to prove his eligibility for exemption must be uploaded for evaluation by the buyer.
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 {themselves or through reseller(s)} should have regularly, manufactured and supplied same or similar Category Products to any Central / State Govt Organization / PSU / Public Listed Company for number of Financial years as indicated above in the bid document before the bid opening date. Copies of relevant contracts 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 .In case Buyer has selected Purchase preference to Micro and Small Enterprises clause in the bid, the same will get precedence over this clause.

7. Purchase preference to Micro and Small Enterprises (MSEs): Purchase preference will be given to MSEs 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 the Purchase preference, the bidder must be the manufacturer of the offered product in case of bid for supply of goods. Traders are excluded from the purview of Public Procurement Policy for Micro and Small Enterprises. 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. 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 Seller shall be given opportunity to match L-1 price and contract will be awarded for 25%(selected by Buyer) percentage of total QUANTITY.

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 30% 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 / Public Listed Company. 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.

Blood Bags-IS:15102 (1500 pieces)

(Minimum 50% and 20% Local Content required for qualifying as Class 1 and Class 2 Local Supplier respectively)

Brand Type	Registered Brand
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Technical Specifications

* As per GeM Category Specification

Specification	Specification Name	Bid Requirement (Allowed Values)
GENERAL FEATURES	Clinical Purpose	Collection, processing and storage of whole blood and blood components
	Disposable	Yes
PRODUCT INFORMATION	Conformity to standard for Blood Bag	ISO 3826/IS 15102: Latest Revision
	Type of blood bag	Quadruple (top and bottom)
	Capacity of blood Bag	350 ml
	Material of Bag (Medical grade)	DEHP Plasticized PVC

Specification	Specification Name	Bid Requirement (Allowed Values)
	Blood Collection Bags should be collapsible non vented sterile containers complete with collecting tube for completely closed system to avoid the chances of contamination	Yes
	Flexible pre-sterilized and pyrogen free	Yes
	Non toxic, non haemolytic, biocompatible material	Yes
	There should be no risk of contamination and air embolism (closed system) with all leak proof seals (Disposable bags)	Yes
	Slit on both sides of the bags should be enough to accommodate 5 -10 ml volume test tubes	Yes
	The capacity of the bag should be enough to prevent any ballooning/rupture of the abg from the seam when it is filled up with the requisite volume of blood	Yes
TUBING OF BAG	Flexible kink resistant tubing	Yes
	Non sticking	Yes
	Transparent	Yes
	Leak Proof	Yes
	Length of tubing from primary bag to needle	≥ 80 Cm
	The tubing should have same ID/segment number as that on the bag	Yes
	The tubes should have multiple printed ID/segment numbers	Yes
	Clamp provided for closed system	Yes
NEEDLE	Needle Size	16 G
	Ultra thin walled and straight to reduce penetration force and enable painless vein puncture	Yes
	Sharp, regular and smooth margins and bevelled tip	Yes
	Rust proof	Yes

Specification	Specification Name	Bid Requirement (Allowed Values)
	Tightly fixed with hub covered with sterile guard	Yes
	Hermetically sealed	Yes
	The needle should not separate from the tube at any point of time, especially while removing it from the vein for donor safety	Yes
	The needle must confirm to ISO 1135-3 standard	Yes
EXTERNAL PORT	Tamper proof and should not be re-capped	Yes
	Easily accessible	Yes
ANTICAOGULANT AND PRESERVATIVE SOLUTION	Type of anticoagulant present	CPDA-1, CPD
	Quantity of anticoagulant solution	14 ml per 100 ml of blood
	Solution should be clear and colorless	Yes
	There should be no discoloration of solution on storage at room temperature	Yes
	Additive solution present	Yes
	Type of additive solution	SAGM
	Quantity of Additive solution(ml)	100, NA
	Anticoagulant and/or additive solution should be sterile and pyrogen free	Yes
LABEL	Availability of anticoagulant/additive quality check certificate from manufacture (proof of same to be submitted to buyer)	Yes
	Non-peel off	Yes
	Heat sealed/ Pressure embossed label	Yes
	The label should remain attached between room temperature to - 80°C with a transparent adhesive	Yes
	Date of manufacturing, date of expiry and batch number must be mentioned on each bag	Yes

Specification	Specification Name	Bid Requirement (Allowed Values)
RESISTANCE TO DISTORTION	Bag (Fiiled to normal capacity) shall withstand a acceleration of 5000 g for 30 min at temperature 4°C to 24°C without becoming permanently distorted	Yes
	Bag (Fiiled to normal capacity) should be able to withstand temperature upto - 80°C without breakage	Yes
PACKAGING	Individual bag packed in plastic pack and multiple bags packed in moisture proof aluminum foil (Protective dual packaging) eliminating microbial contamination on surface maintaining the contents of the bag	Yes
	The supplier should ensure proper transportation of the consignment of blood bags in temperature controlled conditions (Storage temperature should not exceed 30°C)	Yes
CERTIFICATIONS & REPORTS	Availability of valid drug license issued from competent authority defined under Drugs and Cosmetics Act, 1940 (Proof of the same to be submitted to buyer on demand)	For Manufacture, For Sale
	Manufacturer certifications (Proof of the same to be submitted to the buyer on demand)	ISO 13485, GMP, WHO GMP
	Product Certifications (Proof of the same to be submitted to buyer on demand)	ISO, EU-CE
	Four digit number of notified body If product is EU-CE certified	7556
	Each batch supplied should be accompanied with quality assurance test report from NABL approved lab/any lab approved from govt of India as well as in house lab	Yes

Specification	Specification Name	Bid Requirement (Allowed Values)
	Biocompatibility of the material of the product must certified by the manufacturer and be supported by the test reports of cell culture cytotoxicity, hemolysis, systemic infections, sensitization, Intra-cutaneous injection, pyrogen test and Sterility	Yes
	Submission of manufacturer's documented evidence of biochemical parameters of blood stored in CPDA/CPDA-1/CPD-SAGM containing DEHP plasticized PVC blood bags manufactured by the company on 28th/35th/42nd day of storage	Yes
SHELF LIFE	Shelf Life from the date of manufacture (in months)	24
	Stability report from a recognized laboratory must be submitted to the buyer at the time of supply	Yes
	The product should have atleast 3/4 of the total shelf life at the time of dispatch to the consignee	Yes

Consignees/Reporting Officer and Quantity

S.No.	Consignee/Reporting Officer	Address	Quantity	Delivery Days
1	W.Shanborlang Swer	793018,P.O. NEIGRIHMS, Mawdiangdiang, Shillong	1500	30

Special terms and conditions-Version:2 effective from 20-10-2022 for category Blood Bags-IS:15102

1. All Provisions of Drugs and Cosmetics Act, 1940 and Rules made there under as amended till date will always be applicable. This will include all notifications issued by *Central Drugs Standard Control Organisation (CDSCO)*, Ministry of Health & Family Welfare (MoHFW) and Department of Pharmaceuticals (DOP), Ministry of Chemicals & Fertilizers time to time in this regard.
2. The sellers are registered on GeM based on self-declaration of valid Drug License, product certification, test reports etc. However, buyers must check and validate the details at their end for all applicable licenses and certifications e.g., validity and authenticity/genuineness of drug license, product certification, manufacturer certification/licenses, test reports etc.

3. The price offered by the seller shall not, in any case exceed the DPCO controlled price, if any, fixed by the Central/State Government, the Maximum Retail Price (MRP) and the selling price. The seller must reduce the prices if there is any reduction in DPCO ceiling price, if any.
4. Any other Terms and Conditions which is not included or at variance with the conditions specified in STC/GTC, may be added by the buyer through Additional Terms and Conditions in the bid to ensure items are procured from authentic/validated source with appropriate and applicable quality. The above terms and conditions are in reverse order of precedence i.e., ATC shall supersede specific STC which shall supersede General Terms and Conditions ("GTC"), whenever there are any conflicting provisions.

Buyer Added Bid Specific Terms and Conditions

1. Experience Certificate for the supply of the same to any Govt/ PSU/ any renowned private organisation along with Supply/ Purchase Order.
2. If the agency is registered under MSME or NSIC, then EMD exemption certificate needs to be enclosed.
3. Make in india specific authorisation certificate needs to be enclosed.
4. **Generic**

OPTION CLAUSE: The Purchaser reserves the right to increase or decrease the quantity to be ordered up to 50 percent of bid quantity at the time of placement of contract. The purchaser also reserves the right to increase the ordered quantity by up to 50% of the contracted quantity during the currency of the contract at the contracted rates. Bidders are bound to accept the orders accordingly.

5. Buyer Added Bid Specific ATC

Buyer Added text based ATC clauses

Samples for this Bid is mandatory, Samples are to be submitted at Central Store NEIGRIHMS before Bid opening Date, failing which Bid participation will be rejected

Disclaimer

The additional terms and conditions have been incorporated by the Buyer after approval of the Competent Authority in Buyer Organization. Buyer organization is solely responsible for the impact of these clauses on the bidding process, its outcome, and consequences thereof including any eccentricity/restriction arising in the bidding process due to these ATCs and due to modification of technical specifications and/or terms and conditions governing the bid. Any clause incorporated by the Buyer such as demanding Tender Sample, incorporating any clause against the MSME policy and Preference to make in India Policy, mandating any Brand names or Foreign Certification, changing the default time period for Acceptance of material or payment timeline governed by OM of Department of Expenditure shall be null and void and would not be considered part of bid. Further any reference of conditions published on any external site or reference to external documents/clauses shall also be null and void. If any seller has any objection/grievance against these additional clauses or otherwise on any aspect of this bid, they can raise their representation against the same by using the Representation window provided in the bid details field in Seller dashboard after logging in as a seller within 4 days of bid publication on GeM. Buyer is duty bound to reply to all such representations and would not be allowed to open bids if he fails to reply to such representations. Also, GeM does not permit collection of Tender fee / Auction fee in case of Bids / Forward Auction as the case may be. Any stipulation by the Buyer seeking payment of Tender Fee / Auction fee through ATC clauses would be treated as null and void.

[This Bid is also governed by the General Terms and Conditions](#)

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.

---Thank You---