

File No.- BMSIC/40030/375-2024/7760

दिनांक:-29/11/2024

बिहार चिकित्सा सेवाएं एवं आधारभूत संरचना निगम लि०, बिहार पटना

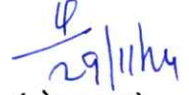
BMSICL की निविदा संख्या-BMSIC/MEDICAL DEVICES/CONSUMABLES/24-11 से संबंधित प्राप्त ई-निविदाओं के तकनीकी मूल्यांकन विवरणी पर साक्ष्य सहित दावा/आपत्ति की माँग संबंधी सूचना

बिहार चिकित्सा सेवाएं एवं आधारभूत संरचना निगम लि०, द्वारा Medical Devices/Consumables की अधिप्राप्ति हेतु ई-निविदा (निविदा संख्या-BMSIC/MEDICAL DEVICES/CONSUMABLES/24-11) आमंत्रित की गई थी, जिसके आलोक में निर्धारित समय सीमा के अन्तर्गत कुल 08 (आठ) ई-निविदा प्राप्त हुई। उक्त 08 (आठ) ई-निविदा के दस्तावेजों के प्राथमिक तकनीकी मूल्यांकन के उपरान्त तैयार की गई तकनीकी मूल्यांकन विवरणी के दावा/आपत्ति के आमंत्रण हेतु BMSICL के website (www.bmsicl.gov.in) एवं e-proc 2.0 के website (www.eproc2.bihar.gov.in) पर अपलोड की गई है।

निविदा में भाग लेने वाले निर्माता संस्थान अथवा अन्य आपूर्तिकर्ता अपलोड किये गए उपर्युक्त तकनीकी मूल्यांकन प्रपत्र में किसी प्रकार की विसंगति पाये जाने की स्थिति में, साक्ष्य सहित अपना दावा/आपत्ति दिनांक 06/12/2024 के अपराह्न 05:00 बजे तक केवल e-proc 2.0 के website (www.eproc2.bihar.gov.in) के माध्यम से कर सकते हैं।

e-proc 2.0 के website (www.eproc2.bihar.gov.in) के अलावा अन्य किसी माध्यम से प्राप्त किसी प्रकार के दावा/आपत्ति पर कोई विचार नहीं किया जायेगा।

अनुलग्नक- यथोक्त


(हरेन्द्र राम)

महाप्रबंधक (अधिप्राप्ति)

ज्ञापांक:- BMSIC/40030/375-2024/7760

दिनांक:-29/11/2024

प्रतिलिपि :- प्रबंधक (सिस्टम) एवं प्रबंधक (औषधि) को सूचनार्थ प्रेषित। निदेश है कि सूचना का अनुलग्नक सहित अविलंब BMSICL के website (www.bmsicl.gov.in) एवं e-proc2 (www.eproc2.bihar.gov.in) पर अपलोड करना सुनिश्चित करें।


(हरेन्द्र राम)

महाप्रबंधक (अधिप्राप्ति)



TENDER NO. BMSIC/MEDICAL DEVICES/CONSUMABLE/24-11

Check list - I: Evaluation sheet to be used for preliminary evaluation of technical bids

Company Name:.....LEELA MEDLIFE PRIVATE LIMITED..... Total Number of Pages Submitted in bid documents:...01 TO 78.....

Sl. No	SBD clause serial No	Description of the Clause/ Technical Eligibility Criteria as per SBD as mentioned in the Check List	Document/ fees submission status (Yes/ No)	Page number	Do the provided documents meet the eligibility criteria ? (Yes/ No)	Remarks
1	3. (a)	Tender Fee (Non-Refundable) of Rs 10,000/- + GST @18%, Total Rs 11,800 shall be submitted online (On E-Proc-2). This fee is payable only once for one tender irrespective of items contained therein.	Yes	78	Yes	-
2	3.(b)	EMD shall be submitted ONLY in form of Bank Guarantee (Offline mode), drawn in favor of Managing Director, Bihar Medical Services and Infrastructure Corporation Limited from any Scheduled/Nationalized bank payable at Patna, as mentioned in Clause 2, as per following table:- S.N. No. of MEDICAL DEVICES/CONSUMABLE quoted EMD Amount 1. Upto 5 MEDICAL DEVICES/CONSUMABLE Rs 50,000/- (Fifty thousand only) 2. 6 to 10 MEDICAL DEVICES/CONSUMABLE Rs 1,00,000 (One Lakhs only) 3. 11 to 20 MEDICAL DEVICES/CONSUMABLE Rs 1,50,000 (One Lakhs fifty thousand only) 4. More than 20 MEDICAL DEVICES/CONSUMABLE Rs 2,50,000 (Two Lakhs fifty thousand only)	Yes	27-30	Yes	AS Per Bid Document clause 3(b) BG Submitted for EMD must be schedule/Nationalized Bank Payable at Patna But firm has not submitted the same.
3	3.(c)	Documentary evidence of the constitution of the company/firm/Proprietorship such as Memorandum and Articles of Association, Partnership Deed etc. should be submitted with details of the Name, Address, Telephone Number, Fax Number, e-mail address of the firm and of the Managing Director / Partners / Proprietor should be submitted.	Yes	43-70	Yes	-
4	3.(k)	Copies of the Audited Balance Sheet and Profit and Loss statement showing details of their annual average turnover not less than 5 Crores for any three of the last four consecutive financial years (Auditor/CA certificate of turnover will not be accepted). Self-attested copies are to be submitted.	Yes	31-36	No	UDIN Number is Submitted for Only for F.Y-2021-2022.
5	3.(l)	Copy of Income Tax Return for any three of last four consecutive Assessment years should be submitted (Self Attested).	Yes	19-21	Yes	-
6	3.(o)	Copy of PAN Card of the bidder company should be submitted (self-attested).	Yes	22	Yes	-
7	3.(p)	Copy of certificate of valid GST registration of the bidder company should be submitted (self-attested).	Yes	23-25	Yes	-

Ankhanand Singh

[Signature]

TENDER NO. BMSIC/MEDICAL DEVICES/CONSUMABLE/24-11

Check list - II: Evaluation sheet to be used for preliminary evaluation of technical bids

Company Name: - M/s Leela Medlife Pvt. Ltd. Total Number of Pages Submitted in bid documents:- 77

Sl. No	SBD Clause serial No	Description of the Clause/ Technical Eligibility Criteria as per SBD, as mentioned in Check List	Document/ fees submission status (Yes/ No)	Page Number	Do the provided documents meet the eligibility criteria ? (Yes/ No)	Remarks
1	3.(d)	The details of Bidder Name, Address, Telephone Number, Fax Number, e-mail address of the bidder and of the Managing Director / Partners / Proprietor should be submitted in Annexure-V.	No	No	No	Not submitted
2	3.(e)	Power of Attorney or Resolution of the Board by which the authorized signatory has been authorized by the bidder firm to sign the documents should be submitted.	Yes	1	Yes	
3	3.(f)	(I) Minimum three years old valid Manufacturing License of the quoted product.	NA	NA	NA	
4		In case of manufacturer, the bidder firm must have a valid manufacturing license or duly acknowledged renewal application with old license issued by the state licensing authority/Central licensing approving authority under Medical Devices Rules 2017 where ever applicable.	NA	NA	NA	
5		Approved product list as per the license issued for quoted MEDICAL DEVICES/CONSUMABLE as Per Medical Devices Rules 2017.	NA	NA	NA	
6		Manufacturing License along with approved product list must be valid till the last date of the submission of tender.	NA	NA	NA	
7		Valid Pollution Control Clearance Certificate in accordance with Water [Prevention and control of Pollution] Act, 1974 & Air [Prevention and control of Pollution] Act, 1981 and Hazardous Wastes (Management, Handling & Trans Boundary Movement) Rules 2008 (Self Attested Copy of Certificate to be enclosed) (as applicable).	No	No	No	Not submitted
8		Bidders shall submit self-attested copies of required manufacturing license and approved product list in support of above-mentioned condition and they are required to specify the quoted product in their approved product list by highlighting it.	Yes	2	No	Highlighted product list & license not submitted.

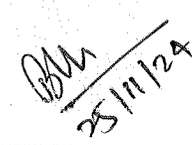
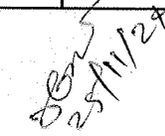
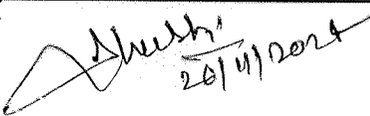
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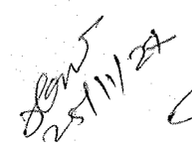
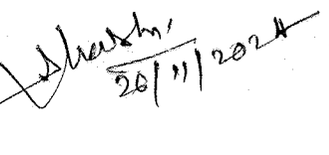
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9		(II) In case of Importer, the bidder (importer) firm must have a valid import License with product registration certificate issued by the Drugs controller General of India as per Medical Device Rule 2017. All Quoted products should be accompanied by their invoices, statement and import License showing that the quoted product are being imported and sold in India by the bidder (Importer) firm minimum for Last three years. Import license must be valid on the last date of submission of tender.	Yes	1-2	No	(i) Only IEC certificate submitted. (ii) Product list/ approvals per IEC/ MDR 2017 not submitted. (iii) Manufacturer/Import License as per MDR 2017 not submitted. (iv) Last 3 years import invoices not submitted.
10		(III) In case of Non-drug item(s) where neither the D & C Act 1945 nor the Medical Device Rule 2017 is applicable the bidder must have a manufacturing license/import export certificate (IEC) with an undertaking/Self declaration in his letter pas as per Annexure-VII that the item(s) quoted by the bidder is/are non-drug item(s) i.e. neither covered under D & C Act nor Under Medical Device Rule 2017.	Yes	38-39	No	License under MDR 2017 must be provided for the products falling under Medical Device Rules.
11	3.(g)	Bidder must have Market Standing Certificate of minimum three years issued by the concerned Licensing Authority from Drugs Control Department/Concerned Government Department for the quoted product. Self-attested copies are to be submitted.	Yes	8	No	MSC not issued by concerned department.
12	3.(h)	Non-Conviction Certificate (NCC) issued by the concerned Licensing Authority from Drugs Control Administration of the state for last three years should be submitted (wherever applicable). It should be not more than one year old. Self-attested copies are to be submitted.	Yes	9-10	No	NCC not issued by concerned department.
13	3.(i)	WHO-GMP/GMP (Good Manufacturing Practice) as per revised Schedule- 'M'/COPP Certificate of the manufacturing unit issued by the Licensing Authority/Drugs Control Department, (wherever applicable). The GMP certificate must not be older than one year from the date of publication of tender. Self-attested copies are to be submitted. Explanation- Generally the GMP Certificate is issued for one-year validity. Hence the provision that it should not be older than one year from the date of publication of tender implies mutatis mutandis that the GMP certificate should remain valid till the date of submission of tender.	NA	NA	NA	ISO 13485: 2016 certificate submitted valid upto 05.05.2024 (page no. 11)
14	3.(j)	Maximum Production Capacity Certificate (Section wise) issued by concerned Licensing Authority from Drugs Control Department highlighting the quoted product section (wherever applicable). Self-attested copies are to be submitted. In case of Importer, an affidavit (with Stamp) sworn before first class magistrate/ Notary stating the batch production capacity of the firm and also that said production (Importing) capacity shall be adequate for requirement laid in NIT. Importer will have also to submit Invoices/evidence of import in items of said product with quantity details.	Yes	15-16	No	MPCC not issued by concerned department.

15	3.(m)	The tenderer should give an affidavit sworn before first class magistrate / Notary stating that the firm & its quoted product is not black listed/Debarred currently (as on the date of submission of the tender) by Central Government / Central Government agencies/any state government or any of the state government agencies / or any Drug procurement agencies or by BMSICL as per Annexure-II .	Yes	6-7	Yes	
16	3.(n)	List of items quoted in prescribed format as per Annexure-III duly signed.	Yes	71-74	Yes	
17	5.(k)	Affidavit declaration regarding acceptance of tender conditions to be submitted by the bidding firm as per Annexure-IV .	Yes	26	Yes	
18	5.(l)	Filled check list as per given Annexure-VI to be submitted at the time of uploading the bid.	Yes	40-42	Yes	
19	7.(c)	Power of Attorney or Resolution of the Board by which the authorized signatory has been authorized by the bidder firm should sign the documents in cases where person other than the Managing Director/Managing Partner or sole Proprietor signs the document.	Yes	1	Yes	
20	10.(c)	If a particular document/Certificate to be uploaded as specified in bid, is not applicable for a bidder, the bidder shall attach a scanned copy of declaration in the letter head stating that the specific document is not applicable/exempted for the bidder in connection to this tender.	NA	NA	NA	
21	2 (d)	The price quoted by the bidders must not exceed the ceiling price as fixed by NPPA (National Pharmaceutical Pricing Authority) as per the provision of "Drugs Price Control Order (wherever applicable)" and the quoted rate should be at least 20% less than its MRP. In extraordinary cases the Managing Director, BMSICL has discretion to take decision). Explanation- In order to ensure procurement of the tendered drugs at the most appropriate and competitive rate, the bidders are directed to quote their lowest price as compared to the Rate provided to their respective Distributors/ Dealers/Wholesalers/ Carrying and Forwarding Agents/Authorized depot sales point in the State of Bihar. A Notarized affidavit to this effect on a Rs 100/- Non-judicial Stamp Paper should be submitted with the Bid in the prescribed format as mentioned in Annexure-X .	Yes	37-39	Yes	
22	-	Performance Statement- Annexure- IX	No	No	No	Not submitted
23	-	AFFIDAVIT FOR NON DRUG ITEM(S) ANNEXURE-VII	Yes	38-39	No	License under MDR 2017 must be provided for the products falling under Medical Device Rules.

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TENDER NO. BMSIC/MEDICAL DEVICES/CONSUMABLE/24-11

Check list - III: Evaluation sheet to be used for preliminary evaluation of technical bids

Company Name: - M/s Leela Medlife Pvt. Ltd. Total Number of Pages Submitted in bid documents:- 77

Sheet to be used for verification of product approval and market standing

Sl. No	NIT Sl. No	Name of the Quoted MEDICAL DEVICES/CONSUMABLE		Specification		Pack Size		Product category (Medical Device/Non Drug)	Approval Details			Last 3yrs market standing certificate/ New Drug
		As per NIT	As per Approval	As per NIT	As per Approval	As per NIT	As per Approval		Product Approval	Approved Upto	Approved in Brand /Generic Name	
1	1	Cryovials	Product list/ approval not submitted	(2ml) 1. Polypropylene Colourless sterile tubes to store human or animal cell 2. Temperatures as low as -190°C to -196°C. 3. It should be RNase or Dnase free. 4. ISO certified.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Product list/ approval not submitted	Product list/ approval not submitted	Not submitted for concerned department.
2	2	Cryobox	Product list/ approval not submitted	(100 set) 1. Box for storage of cryovials at -80°C 2. Solid polypropylene base with Clear polycarbonate cover for easy identification. 3. Temperature Range: -Withstand cryogenic storage conditions from -135 to 100 C Alphanumeric grid for easy sample identification 4. Withstand sterilization by autoclave, gas and disinfectant 5. Manufacturer should be ISO certified.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Product list/ approval not submitted	Product list/ approval not submitted	Not submitted for concerned department.
3	3	Pipette Aid	Product list/ approval not submitted	SPECIFICATION OF SEROLOGICAL PIPETTE CONTROLLER (PIPETTE AID) 0.5-10µL 1. Suitable smooth setting of pumping efficiency with high aspirating speed (20-25 ml in 5 second). 2. One finger control for flow and blow out. Single hand operation, must have delivery mode (Gravity delivery blowout) and adjustable motor speed range with thumb. 3. Cordless work for minimum upto 5 hours. LED light battery indicator that alert before 2 hours. Short charging time for battery to full capacity and at least 8-10 hours working capacity. 4. Continuing directly after recharging. Autoclavable pipette adaptors and filter holders. 5. Space saving wall holder and calibration certificates from CE/ISO certified agencies.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Product list/ approval not submitted	Product list/ approval not submitted	Not submitted for concerned department.
4	4	Pipette Aid	Product list/ approval not submitted	SPECIFICATION OF SEROLOGICAL PIPETTE CONTROLLER (PIPETTE AID) 10-20µl 1. Suitable smooth setting of pumping efficiency with high aspirating speed (20-25 ml in 5 second). 2. One finger control for flow and blow out. Single hand operation, must have delivery mode (Gravity delivery blowout) and adjustable motor speed range with thumb. 3. Cordless work for minimum upto 5 hours. LED light battery indicator that alert before 2 hours. Short charging time for battery to full capacity and at least 8-10 hours working capacity. 4. Continuing directly after recharging. Autoclavable pipette adaptors and filter holders. 5. Space saving wall holder and calibration certificates from CE/ISO certified agencies.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Product list/ approval not submitted	Product list/ approval not submitted	Not submitted for concerned department.
5	5	Pipette Aid	Product list/ approval not submitted	SPECIFICATION OF SEROLOGICAL PIPETTE CONTROLLER (PIPETTE AID) 20-200µl 1. Suitable smooth setting of pumping efficiency with high aspirating speed (20-25 ml in 5 second). 2. One finger control for flow and blow out. Single hand operation, must have delivery mode (Gravity delivery blowout) and adjustable motor speed range with thumb. 3. Cordless work for minimum upto 5 hours. LED light battery indicator that alert before 2 hours. Short charging time for battery to full capacity and at least 8-10 hours working capacity. 4. Continuing directly after recharging. Autoclavable pipette adaptors and filter holders. 5. Space saving wall holder and calibration certificates from CE/ISO certified agencies.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Product list/ approval not submitted	Product list/ approval not submitted	Not submitted for concerned department.

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6	6	Pipette Aid	Product list/ approval not submitted	<p>SPECIFICATION OF SEROLOGICAL PIPETTE CONTROLLER (PIPETTE AID) 100-1000µl</p> <ol style="list-style-type: none"> 1. Suitable smooth setting of pumping efficiency with high aspirating speed (20-25 ml in 5 second). 2. One finger control for flow and blow out. Single hand operation, must have delivery mode (Gravity delivery blowout) and adjustable motor speed range with thumb. 3. Cordless work for minimum upto 5 hours. LED light battery indicator that alert before 2 hours. Short charging time for battery to full capacity and at least 8-10 hours working capacity. 4. Continuing directly after recharging. Autoclavable pipette adaptors and filter holders. 5. Space saving wall holder and calibration certificates from CE/ISO certified agencies. 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
7	7	Forceps	Product list/ approval not submitted	<p>SPECIFICATION OF FORCEPS</p> <ol style="list-style-type: none"> 1. Must be made of Graded stainless steel conforming to IS 410 & 420 2. ISO 7151 Certified 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
8	8	Discarding jars	Product list/ approval not submitted	<p>SPECIFICATION OF DISCARDING JARS</p> <ol style="list-style-type: none"> 1. Plastic Container Made of HDPE – SPI Resin ID code 2 with Biohazard Sign 2. Puncture, Leak & Tamper Proof 3. Conform to ASTM Standard D1922 and D1909 or IS 14995 :2001 4. Able to Withstand 135°C 5. Size- 5 litre 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
9	9	Biomedical waste disposal (BMW)bags	Product list/ approval not submitted	<p>SPECIFICATION OF BIOMEDICAL WASTE DISPOSAL (BMW) BAGS, (with ties for sealing, preferably autoclavable) (Red, Black, Yellow).</p> <ol style="list-style-type: none"> 1. Plastic Bags of HDPL, LLPE, DP (Biohazard, ISO Certificate & Hospital Name should be Printed) should be biodegradable not reused plastic, bags should be superior quality of thickness of 50 micron or more thickness, non-chlorinated plastic with ties for sealing. 2. Toxicity Test Certification approved by FDA. 3. Bags should be Autoclavable & Puncture proof. 4. Should not emit thick Black smoke in incinerator. 5. Should burn without leaving traces into ashes. 6. Produced by IS/ISO certified under 17088-2008 test complies Indian Standard. 7. Aerobic Biodegradation in presence of Municipal Sewage Sludge Certification. 8. Anaerobic Biodegradation land fill conditions & All climate Condition certification. 9. The firm should be registered under Pollution Control Board as biodegradable Manufacturer. 10. Sizes :- 14 x 18 inches/ 18 x 22 inches 11. Colour : Red, Black and Yellow. 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
10	10	Biomedical waste disposal Bins	Product list/ approval not submitted	<p>SPECIFICATION OF BIOMEDICAL WASTE DISPOSAL (BMW) BINS</p> <ol style="list-style-type: none"> 1. Plastic bins LLDPE made from virgin polymer material. Thickness of bins shall be minimum 3mm (+/-0.2mm) 2. Foot operated lid and handle for lifting. Lid mechanism shall be of SS material only. 3. Able to Withstand 135°C 4. LLDPE bucket of Circular top with holding lid support stopper 5. Sizes: 10 litre/20litre 6. SS parts shall be smooth finished. Proper rubber studs shall be provided on the paddle and both ends of 20mm SS Square hollow pipe. 7. Lid of the bin shall be foot operated and SS bottom rod of foot operated mechanism 8. Bin shall be printed as per requirement of Bio Medical Waste Management Committee. 9. Dimensional accuracy ready to use. 10. Produced by an ISO 9001:2015 & 14001:2015 Certified unit. 11. Manufacturer should be CE certified 12. Demonstration of product during technical bid. 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				

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11	11	Iceboxes with gel packs	Product list/ approval not submitted	<p>SPECIFICATION OF ICEBOXES WITH GEL PACKS</p> <ol style="list-style-type: none"> The cool box should be durable construction, compact design and made from FDA approved grades of polyethylene material from food grade virgin double walled body with all round EPS/PUF filled insulation, inbuilt handle, heavy base, BIS marked. Excellent insulation maintains uniform temperature for storage & transportation. Ice Box manufacturer conforming to BIS/ISO 9001:2015 or exporter of EPS/PUF insulated Ice Box having Coolant Packs (Ice - Gel-Packs). Tight fittings to keep ice melting rate in line with the best in the world with rubber clamps for locking lid [to prevent corrosion]. Tapered inner bottom for maximum drainage with drainage plugs at bottom is preferable. Lid with suitable gasket for better temperature retention. 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
12	12	Tube rack	Product list/ approval not submitted	<p>SPECIFICATION OF TUBE RACK (FOR 15 ML TUBES)</p> <ol style="list-style-type: none"> Easy storage and access of test tubes in vertical position for 18/48/72 tubes Made of virgin polypropylene Should be unbreakable, autoclavable and able to withstand subfreezing temperatures Able to protect test tubes from minor shocks and damage during transportation. 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
13	13	Tube rack	Product list/ approval not submitted	<p>SPECIFICATION OF TUBE RACK (FOR 20 ML TUBES)</p> <ol style="list-style-type: none"> Easy storage and access of test tubes in vertical position for 18/48/72 tubes Made of virgin polypropylene Should be unbreakable, autoclavable and able to withstand subfreezing temperatures Able to protect test tubes from minor shocks and damage during transportation. 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
14	14	Permanent Marker pen	Product list/ approval not submitted	<p>SPECIFICATION OF SPECIFICATION OF MARKER PEN</p> <ol style="list-style-type: none"> Permanent marker pen. Colour : Black/blue Precise, narrowed tip for extreme control (0.3 to 0.5mm) Permanent on most surfaces, fade and water resistant. Quick-drying ink with non-toxic formula ISO Certified 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
15	15	Cellotape	Product list/ approval not submitted	<p>SPECIFICATION OF CELLO TAPE</p> <ol style="list-style-type: none"> High quality strong adhesive. 40 microns transparent film easily tearable by hands 1/2 inch width size of tape ISO Certified Length : 60 meters 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
16	16	Printer with label markers (for label print outs)	Product list/ approval not submitted	<p>SPECIFICATION OF PRINTER WITH LABEL MARKERS (FOR LABEL PRINT OUTS)</p> <ol style="list-style-type: none"> As per approved License 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
17	17	1.7 ml Eppendorf Tubes	Product list/ approval not submitted	<p>SPECIFICATION OF MICROCENTRIFUGE TUBES (EPPENDORF TUBES) 1.7 ML</p> <ol style="list-style-type: none"> Clear and transparent tubes FDA approved virgin polypropylene eliminating any trace of heavy metals Leak proof seal for prevention against accidental spillage Withstands temperatures from -80° to +121° C Clear and large graduation on tube body Choice of selecting low retention tube to increase recovery of the sample Compatible with all standard and high capacity centrifuge rotors Non cytotoxic & biologically inert Options for regular white tube, assorted color tube and in amber color for light sensitive reagents 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				

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18	18	Cryovial/ Eppendorf rack	Product list/ approval not submitted	SPECIFICATION OF MICROCENTRIFUGE TUBES RACK (EPPENDORF RACK) 1. Used for holding microtubes of 1.7ml capacity 2. Made of polypropylene, autoclavable 3. Withstand temperatures from 0° C to 135°C 4. Withstand sterilization by gas, autoclaving and disinfectant 5. Rigid container with mesh type gaps for holding 84 cryovials of 1.7ml capacity 6. ISO Certified	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.			
19	19	Micropipettes-	Product list/ approval not submitted	SPECIFICATION OF MICROPIPETTE (100 µL -1000 µL.) 1. Accuracy of 0.5%, CV 0.2% 2. Fully autoclavable 3. With calibration and accuracy certificate 4. ISO Certified 5. Easy to handle, grip and dispense and aspirate fluids. 6. Should not cause repeated stress injury to the user.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.			
20	20	Filter Barrier Tips:	Product list/ approval not submitted	SPECIFICATION OF FILTER BARRIER TIPS (200 µL) 1. Compatible with international standard pipettes attachable by firm grip. 2. Available in packs of 1000 tips 3. Non sticky in nature with Smooth internal surface 4. Tips should be sharp and pointed 5. Should be Less than 0.5% wastage due to any manufacturing defects 6. Should not be made of reused plastic 7. Manufacturer should be ISO certified	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.			
21	21	Filter Barrier Tips:	Product list/ approval not submitted	SPECIFICATION OF FILTER BARRIER TIPS (1000 µL) 1. Compatible with international standard pipettes attachable by firm grip. 2. Available in packs of 1000 tips 3. Non sticky in nature with Smooth internal surface 4. Tips should be sharp and pointed 5. Should be Less than 0.5% wastage due to any manufacturing defects 6. Should not be made of reused plastic 7. Manufacturer should be ISO certified	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.			
22	22	Tissue rolls	Product list/ approval not submitted	SPECIFICATION OF TISSUE ROLL 1. Tissue Roll white colour plain Paper of 1-2 mm thickness of GSM 50-80. 2. ISO 12625-1:2011 certified	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.			
23	23	PCR tubes/ PCR plates	Product list/ approval not submitted	SPECIFICATION OF PCR TUBES/PCR PLATES (200 µL) 1. 96 wells, raised rims for effective sealing, individually wrapped, D Nase, R Nase, Pyrogen free.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.			
24	24	PCR plates adhesive seals & plate sealer	Product list/ approval not submitted	SPECIFICATION OF PCR PLATES ADHESIVE SEALS & PLATE SEALER 1. Peel adhesive should withstand Temp range -20°C to 120° 2. Should be Colour clear.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.			
25	25	Micropipettes-	Product list/ approval not submitted	SPECIFICATION OF MICROPIPETTE (0.5 µl -10 µl) 1. Accuracy of 0.5%, CV 0.2% 2. Fully autoclavable 3. With calibration and accuracy certificate 4. ISO Certified 5. Easy to handle, grip and dispense and aspirate fluids. 6. Should not cause repeated stress injury to the user.	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.			

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26	26	Micropipettes-	Product list/ approval not submitted	<p>SPECIFICATION OF MICROPIPETTE (2 µl -20 µl)</p> <ol style="list-style-type: none"> 1. Accuracy of 0.5%, CV 0.2% 2. Fully autoclavable 3. With calibration and accuracy certificate 4. ISO Certified 5. Easy to handle, grip and dispense and aspirate fluids 6. Should not cause repeated stress injury to the user. 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
27	27	Micropipettes-	Product list/ approval not submitted	<p>SPECIFICATION OF MICROPIPETTE (20 µl -200 µl)</p> <ol style="list-style-type: none"> 1. Accuracy of 0.5%, CV 0.2% 2. Fully autoclavable 3. With calibration and accuracy certificate 4. ISO Certified 5. Easy to handle, grip and dispense and aspirate fluids. 6. Should not cause repeated stress injury to the user. 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
28	28	Filter Barrier Tips:	Product list/ approval not submitted	<p>SPECIFICATION OF FILTER BARRIER TIPS (10 µL)</p> <ol style="list-style-type: none"> 1. Compatible with international standard pipettes attachable by firm grip. 2. Available in packs of 1000 tips 3. Non sticky in nature with Smooth internal surface 4. Tips should be sharp and pointed 5. Should be Less than 0.5% wastage due to any manufacturing defects 6. Should not be made of reused plastic 7. Manufacturer should be ISO certified 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
29	29	Filter Barrier Tips:	Product list/ approval not submitted	<p>SPECIFICATION OF FILTER BARRIER TIPS (20 µL)</p> <ol style="list-style-type: none"> 1. Compatible with international standard pipettes attachable by firm grip. 2. Available in packs of 1000 tips 3. Non sticky in nature with Smooth internal surface 4. Tips should be sharp and pointed 5. Should be Less than 0.5% wastage due to any manufacturing defects 6. Should not be made of reused plastic 7. ISO certified 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
30	30	Nuclease free water for PCR	Product list/ approval not submitted	<p>SPECIFICATION OF NUCLEASE (RNASE) FREE WATER FOR PCR</p> <ol style="list-style-type: none"> 1. Clear and Colourless water. 2. Should be RNase & DNase free. 3. Should be in a pack of 100 ml 4. ISO/GMP certified 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				
31	31	RNase P	Product list/ approval not submitted	<p>SPECIFICATION OF RNASE P</p> <ol style="list-style-type: none"> 1. As per approved License 	Product list/ approval not submitted	Each Piece	Product list/ approval not submitted	MD	Product list/ approval not submitted	Not submitted for concerned department.				

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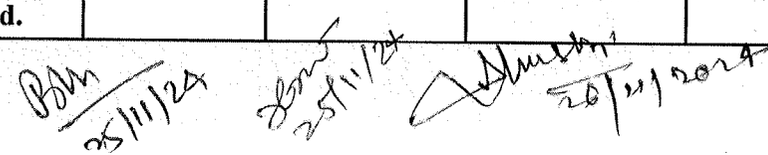
TENDER NO. BMSIC/MEDICAL DEVICES/CONSUMABLE/24-11

Check list - IV: Evaluation sheet to be used for preliminary evaluation of technical bids

Company Name: - M/s Leela Medlife Pvt. Ltd. Total Number of Pages Submitted in bid documents:- 77

Sheet for verification of licence details

Sl. No	NIT Sl. No	Name of the MEDICAL DEVICES/CONSUMABLE as per NIT	Manufacturer		Importer		Validity		GMP/ WHO GMP/ COPP (Import)/BIS/ISI/QMS	
			Forms Number	Manufacturing license Number	Forms Number	Import license Number	From	To	From	To
1	1	Cryovials	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
2	2	Cryobox	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
3	3	Pipette Aid	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
4	4	Pipette Aid	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
5	5	Pipette Aid	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
6	6	Pipette Aid	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024



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7	7	Forceps	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
8	8	Discarding jars	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
9	9	Biomedical waste disposal (BMW)bags	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
10	10	Biomedical waste disposal Bins	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
11	11	Iceboxes with gel packs	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
12	12	Tube rack	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
13	13	Tube rack	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
14	14	Permanent Marker pen	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
15	15	Cellotape	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024

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16	16	Printer with label markers (for label print outs)	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
17	17	1.7 ml Eppendorf Tubes	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
18	18	Cryovial/ Eppendorf rack	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
19	19	Micropipettes-	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
20	20	Filter Barrier Tips:	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
21	21	Filter Barrier Tips:	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
22	22	Tissue rolls	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
23	23	PCR tubes/ PCR plates	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
24	24	PCR plates adhesive seals & plate sealer	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024

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25	25	Micropipettes-	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
26	26	Micropipettes-	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
27	27	Micropipettes-	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
28	28	Filter Barrier Tips:	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
29	29	Filter Barrier Tips:	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
30	30	Nuclease free water for PCR	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024
31	31	RNase P	NA	NA	IEC	AAECL0809R License as per MDR 2017 not submitted.	02.05.2020	Not mentioned	ISO 13485: 2016 05.05.2023	Valid upto 05.05.2024

Note:- Assisted in technical evaluation in reference to letter no. BMSIC/40030/375-2024/7424 dt 21.11.2024 on the basis of documents provided by BMSICL as check list II, III & IV. Provided checklist compiled with due diligence and care. In spite, some inadvertent discrepancies could have been crept in. Humble request to all concerned to bring to the notice in due time if any discrepancies are observed for rectification.

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