Corrigendum

AIIMS Rishikesh tender Enquiry No. T.No 24/RC/Pharmacy/Medicines/ 612/2021-RISH(Admn) "Rate Contract document for procurement of Medicine (Musculoskeletal System/ Analgesics And Gastro-Intestinal Medicines) for Central Pharmacy AIIMS, Rishikesh"

S. No.	Original point	Revised (To be read as)		
1.	Instruction to Bidders; Point1. c. No third-party manufacturers are allowed to quote for any drug.	Bidders with products manufactured on P2P basis from third party but having quality control of such products by in-house Quality Assurance Department are eligible to participate in tender.		
2.	Technical Evaluation Point d: Bidder must have recent experience, within last two years, for supplying salt, for which bid submitted, to developed countries	Bidder must have recent experience, within last two years, for supplying salt, for which bid submitted, to developed country(ies). For imported drug manufacturer must be Supplying/selling the product to/ in developed country(ies).		
3.	Technical evaluation; Point e. Bidder(s) shall submit a Manufacturing & Market Standing Certificate / Experience Certificate issued by the Centre / State Drug Licensing Authority of the respective state that quoted product is manufactured and marketed by them since last 05 (Five) years.	Bidder(s) shall submit a Manufacturing & Market Standing Certificate / Experience Certificate issued by the Centre / State Drug Licensing Authority of the respective state that quoted product is manufactured and marketed by them since last 03 (Three) years. Any firm who took over another manufacturer portfolio in past 3 years from date of publishing of this tender shall submit Manufacturing & Market Standing Certificate / Experience Certificate issued by the Centre / State Drug Licensing Authority stating time duration for which quoted product(s) was manufactured by either manufacturer totaling for 03 (Three) years. In such scenario, current manufacturer must fulfil other eligibility and technical criteria of tender document.		
4.	Technical Evaluation; Point h: Quality Related Action taken by drug regulatory authority of India Bidder has to provide certificate issued from regulatory authority, India that no warning has been issued or action has been taken in last 5years, in terms of quality of production and produced batches.	Bidder has to provide certificate issued from regulatory authority, India that no warning has been issued or action has been in last 5 years or from date of approval to market, whichever is latest, in terms of quality of production and produced batches.		
5.	Technical Evaluation; Point i: Projection Capacity Assessment Certificate: Manufacturing firm must enclose certificate issued by Chartered Accountant/ concerned State Drug Controller indicating actual production detail of a particular molecule, batchwise, for items quoted.	Capacity Assessment Certificate: Manufacturing firm must enclose certificate as per Annexure B (Attached below) *If any bidder already got prepared the format as per earlier clause with all required information for last three financial years then it will also be accepted.		
6.	Financial Evaluation; Point c: Supplier has to submit a notarized affidavit on Indian Non Judicial Stamp Paper of Rs.10/- that bidder has not quoted price higher than previously supplied to any government Institute/ Organization/reputed Private Organization or DGS&D rate in recent past.	Supplier has to submit a notarized affidavit on Indian Non Judicial Stamp Paper of Rs.10/- that bidder has not quoted price higher than previously supplied to any government Institute/ Organization/ reputed Private Organization or DGS&D rate in <i>last three years or since the last change in M.R.P. whichever is latest.</i>		

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Therefore, if at any stage it is found that Therefore, if at any stage it is found that supplier has supplier has quoted lower rates than those quoted lower rates than those quoted in this tender; quoted in this tender; Institute (purchaser) Institute (purchaser) would be given benefit of lower would be given benefit of lower rates by rates by Supplier and Supplier and any excess payment if any, will become immediately any excess payment if any, will become payable to AIIMS, Rishikesh. If such affidavit is not submitted, tender will be summarily rejected. (Part of immediately payable to AIIMS, Rishikesh. If such affidavit is not submitted, tender will be technical bid) summarily rejected. (Part of technical bid) 7. (B) Special Conditions of Contract (SCC) / Contract terms; Point 3. Pharmacopoeia Specification; Point b): It should be ensured that ISI Code No. is Point b). is omitted. indicated on the packing and at the time of Point c). to be read as b). supplies has ISI Mark as well as Code No. as is the statutory requirement of the Bureau of Indian Standards. 8. Special Conditions of Contract (SCC) / Contract terms; Labelling; Point 5a. Before making supply, approved rate contract holder Before making supply, approved rate contract should ensure that all primary, secondary and tertiary holder should ensure that all primary packaging packaging labels (i.e. Ampoules, vials, bottles, jars, tubes labels (i.e. Ampoules, vials, bottles, jars, tubes etc.) should be embossed, imprinted, stamped etc.) should be embossed/ imprinted with barcode carrying information about batch number, date of with letters, or requirements like "AIIMS-RISHIKESH manufacture, expiry date etc. Packing should have labels mentioning "AIIMS-RISHIKESH SUPPLY NOT FOR SUPPLY NOT FOR SALE" stamp with permanent ink on each item/strip up to primary SALE" in permanent ink on each item/strip up to primary level. level. 9. Eligibility criteria cum Administrative cum Technical Evaluation Checklist (Annexure I): 12. Please provide a notarized affidavit on Non 12. Please provide a notarized affidavit on Non Judicial Judicial stamp paper of Rs. 10/- in format given stamp paper of Rs. 10/- in format given in Annexure-V in Annexure-V that you have not quoted price that you have not quoted price higher than previously higher than previously supplied to any supplied to any government Institute/reputed Private government Institute/I/reputed Private I or Institute or DGS&D/GeM rate in last three years or since DGS&D/GeM rate in last three years. If you the last change in M.R.P. whichever is latest. If you don't don't fulfil this criterion, your tender will be out fulfil this criterion, your tender will be out rightly rejected. * If the quoted product M.R.P. was changed within rightly rejected. preceding three years of date of tender publish, then bidder has to submit documents as evidence of approval of such change from drug regulatory authority. 21. Details of DD No of Tender fee: Date: 21. Invoice against claim of supply/ sell of quoted Payable at: molecules to developed country (ies) 22. In-house quality assurance lab with accreditation certificate in name of bidder in reference to Instruction to Bidders; revised Point1. c. 23. Details of DD No of Tender fee: Date: Payable at: 10. Annexure III: I agree to maintain the following information in I agree to maintain the following information in the the Company's record for a period of 8 years Company's record for a period till 01 (one) year after all and shall make this available for verification to contractual obligations are over and shall make this available for verification to any statutory authorities any statutory authorities

Corrigendum regarding Bill of Quantities (BOQ)/ Price schedule:

Column number 05 with heading "Units":

In BOQ unit for quoting price against tablet or capsules is **to be read as one tab./cap.** instead of Stp. (strip)

Annexure B

CAPACITY ASSESSMENT CERTIFICATION

No.:							
Telephone	e firm:						
India, is in go) is properly regist od legal and statu manufacturer of th	tory standing w	rith the respo	onsible health	authorities ar	Il devices in nd is licensed	
3.The produc follows:	tion capacities for			(n	ame of Manu	facturer) is as	
S.No. (As per tender	Medicine Name	Formulation	Strength	Production Capacity in Financial Year			
document)	Name			2017-18	2018-19	2019-20	
					•	ed Accountant eal with date)	
quality standa 5. We hereby	tches and quality ards during the pas certify that the ab	st three years, a	ults, has der and will exhi n is true and	bit these on re accurate to th	mpliance with equest. ne best of our	n the M GMP	
Signature & S Date:	Seal of the Authorit	ty of company __			<u> </u>		
	inted)						
Position	•			<u> </u>			