



GOVERNMENT OF ANDHRA PRADESH

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Web site: www.apecurement.gov.in

TENDER (e – Procurement platform)

For

Procurement of Procurement of General Medicines

(Rate Contract 2019-21)

FOR THE FINALIZATION OF RATE CONTRACT OF ANDHRA PRADESH STATE

TO SUPPLY PROCUREMENT OF GENERAL MEDICINES_TO
13 CENTRAL MEDICINE STORES (CMS)

(Validity of rate contract : 24 months from the date of finalization of Price bid)

Tender Notice No: 46(19-20)/APMSIDC/Medicine Wing/2019-21, Dt: 09.10.2019.

Implementing Agency:

ANDHRA PRADESH MEDICAL SERVICES & INFRASTRUCTURE DEVELOPMENT
CORPORATION (APMSIDC)

(Formerly APMHIDC)

(AN ENTERPRISE OF GOVT. OF A.P.)APMSIDC,

Plot No.9, Survey No.49, IT Park, Mangalagiri, Guntur District – 522 503

GENERAL CONDITIONS
For
PROCUREMENT OF GENERAL MEDICINES

Bids are invited on the e-procurement platform from the Primary manufacturers or direct importers to enter into Rate Contract (R.C) for a period of two years for supply of **Procurement of General Medicines** to 13 Central Medicine Stores located in the District Head Quarters of Andhra Pradesh State. The details of bidding conditions and other terms can be downloaded from the electronic procurement platform of Government of Andhra Pradesh i.e., <https://tender.apecurement.gov.in>

1. a) Only firms having own Manufacturing Units and having WHO-GMP (WHO - Good Manufacturing Practice) certificate issued by the licensing authority.

b) The average annual turnover during last three years i.e. 2015-16, 2016-17 and 2017-18 or 2016-17,2017-18 and 2018-19 should not be less than **Rs.5 crores** for MSMEs of AP (13 districts only) and not less than **Rs.20.00 Crores** for all the other bidders. The bidders should have turnover in each year. The evidence of turnover should be supported by a Certificate from Chartered Account.

2. The Tenderers who participate in the tender shall upload the following scanned Certificates Online.
 - i) Processing Fee: The bidder shall remit processing fee Rs. 5625/- online to the account of Managing Director, APMSIDC,Managalagiri (Account No **142410011000314, IFSC code : ANDB0000366**) and upload the original transaction slip with UTR number. Failure to pay the processing fee and submitting the slip in the aforesaid manner will lead to rejection of the bid .

 - ii) Valid manufacturing License issued by the concerned Drug control authority/
Valid Import License for Importers.

 - iii) Valid Product wise license for manufacturing the products issued by the concerned Drug Control authorities/Valid import license if the product is imported.
In addition to the state licenses, if any product is declared as new drug, licenses issued by DCGI need to be submitted.

 - iv) Annual Turnover certificate supported by a original Certificate from Chartered Accountant.

 - v) The Earnest Money Deposit (EMD) shall be Rs.3 lakhs to each firm. The Earnest Money Deposit shall be paid in the form of Bank Guarantee or online to the account of Managing Director, APMSIDC, Managalagiri (Account No **142410011000314, IFSC code : ANDB0000366**) and upload the original transaction slip with UTR number. Failure to pay the EMD and submitting the slip in the aforesaid manner will lead to rejection of the bid.

The bidders should note that the local MSME/SSI units are exempted from payment of E.M.D, subject to production of necessary documents to that extent by them.

- vi) Non Conviction Certificate issued by DCA authorities for manufacturers and for direct importers that the firm has not been convicted for the last 3 years, issued on or after 01.05.2019. If the firm stands L1 they have to submit NCC issued on or after 20.11.2019.

In addition to that a Notarized affidavit on Rs. 100/- Non Judicial Stamp paper from the firm's Proprietor / Managing Partner / Managing Director / Authorized Signatory of the firm as per Companies act declaring that the firm is not convicted during the period from 01.05.2019 to till the date of tender submission shall be submitted by the bidding firm.

In the same affidavit the firm has to declare that their firm was not blacklisted by any of the procuring agencies in terms of quality issues and submission of fabricated / forged documents/Default Supplies. If the declaration is proved as false, the firm shall pay Rs.1,00,000/-(Rupees One Lakh) as penalty to APMSIDC in addition to legal action for cheating / misleading the Corporation. If stood L1 and during the Rate Contract Period, the firm or any product of the firm is blacklisted by any of the other Procurement agencies, the same shall be intimated to APMSIDC immediately failing which the penal actions mentioned in this paragraph shall be binding.

In the same affidavit the firm has to declared that "we have approved qualified staff, machines & equipments along with capacity to manufacture above category of drugs and our unit have been issued WHO-GMP* by Licensing Authority vide letter No.....dated.....valid upto.....".

(The Notarized affidavit shall be uploaded on E Procurement in online)

vii) WHO-GMP (WHO-Good manufacturing practices Certificate) Certificate issued by the Licensing Authority. The WHO-GMP certificate must not be older than one year from the due date of Bid submission in the case where validity is not mentioned in the certificate. The WHO-GMP certificate of all the manufacturing plants, of which products have been quoted, should be submitted. The Importer should produce WHO-GMP/COPP of the manufacturing firm or a certificate which is at par with WHO-GMP issued by exporting countries like US-FDA approval, etc. In the case of imported drugs, labels and product literature of all quoted products must be submitted.

The Firm will continue to hold WHO-GMP certificate for the product during entire rate contract period of the product. If WHO-GMP certificate expires, it is firm's responsibility to inform APMSIDC about the same and not to accept any further purchase order till re-issue / renewal of WHO-GMP certificate. During the period of non validity of WHO-GMP certificate of the firm the rate contract will deemed to be suspended. If the firm fails to inform APMSIDC about the expiry of WHO-GMP certificate and accept purchase order of APMSIDC and

later on it comes to the knowledge of APMSIDC in this situation firm shall be liable for a panel action.

- viii) Authorization of a senior responsible Person of the company with Authority to transact business.
 - ix) All the documents submitted online should be serially numbered and the first page should contain an index of contents.
 - x) The details of GST i.e. Number etc, should be enclosed in Online.
 - xi) Declaration form with details of Firm contact number, email ID and address.
 - xii) Memorandum and articles of association of the company.
 - xiii) Self declaration of production capacity. In case of bulk quantity if L2 bidder matches with the L1 price, the ordered quantity will be divided in the ratio 70:30. If L2 and L3 also matches then 60:20:20.
 - xiv) Self declaration of batch size for each quoted product.
 - xv) Audited balance sheets of last three financial years, 2015-16, 2016-17 and 2017-18. or 2016-17,2017-18 and 2018-19.
3. The rate quoted per unit should be inclusive of all taxes, all other levies and duties etc., packing, forwarding to Free On Road (FOR) destination at various locations in A.P., India including Insurance, Storage, Transportation Loading, Unloading, License fee, Octroi, Road permits etc. The Basic Price Should include all levies and duties etc., packing, forwarding to FOR destination at various locations in A.P., India including Insurance, Storage, Transportation Loadings, Unloading, License fee, Octroi, Road permits etc. If there is any variation in GST during the contract period, the same will be taken into account and the rates will be revised accordingly.
4. The Participant Bidders have to submit their bids online at <https://tender.apecurement.gov.in>
5. The Bidders have to scan the above particulars and submit online at <https://tender.apecurement.gov.in> on **20.11.2019** by **5.00 P.M.** Indian Standard Time (IST).
6. All the bidders should quote their rate in Indian Currency only.
7. Details of the Tenders Scheduled are as follows:
- a. Downloading of tender document: From **02.11.2019** to **20.11.2019** upto 12.00 PM.
 - b. Bid submission closing date **20.11.2019** upto **5.00 P.M**
 - c. Time and date of opening of technical bids : **20.11.2019** at 5.01 P.M

d. Pre bid Meeting on **07.11.2019** at **11 AM** in the Conference Hall,2nd floor
Plot No.9, Survey No.49, IT Park, Mangalagiri, Guntur District – 522 503
(The Bidders shall submit their queries regarding Tender document on or before
5:00p.m. of **06.11.2019** to the Mail ID:**tenders.apmsidc@gmail.com**).

8. **Objections**

Any queries or objections on other participant documents shall be accepted
upto **5 pm** of **19.11.2019**. After that no queries will be accepted.

9. **Technical Evaluation:**

a) Technical evaluation is conducted with the DCA authorities.

10. **Time extension**

a) Tenders must be received on electronic platform not later than the time and
date specified in the invitation for Tenders. In the event of the specified date for
submission of Tenders being declared as a holiday for the purchaser the
Tenders will be received upto the appointed time on the next working day.

b) The purchaser may at its discretion extend this deadline for submission of
Tenders by amending the Tender document in which case all rights and
obligations of the purchaser and tenderers previously subject to the original
deadline will then be subject to be the new deadline.

THE TENDER DOCUMENT

CONTENT OF TENDER DOCUMENT

The goods required, tender procedures and contract terms are prescribed in the
tender documents.

- a. Instructions to bidders.
- b. Conditions of Contract.
- c. Price Schedule (Model Tender Format)

A) INSRUCTIONS TO TENDERER

1. The tenderer is expected to examine all the instructions, forms, terms and
Specifications in the tender documents. Failure to furnish all information
specified in the tender documents, or submission of tenders not substantially

responsible to the tender document in every respect will be at the Bidder's risk and may result in rejection of Tender.

2. Procedure for Evaluation of Tenders: - Evaluation of bids will be done in two stages:

- **Technical Evaluation.** Technical evaluation will be based on information furnished in the Technical bid document and the supporting documents. Technical evaluation will be done by Technical Evaluation Committee as per G.O.Rt.No.1357. To qualify in the Technical bids, a bidder has to satisfy all the conditions and furnish all supporting documents.
- **Financial Bid Evaluation.** Bids of the only those bidders who qualify in technical evaluation will be opened for evaluation of Financial bids. Financial bids will be evaluated based on prices of all inclusive prices quoted in the tender.

3. PREPARATION OF TENDERS

LANGUAGE OF THE TENDER

The tender prepared by the tenderer and all correspondence and documents relating to the tender exchanged by the tenderer and the purchaser, shall be written in English language, provided that any printed literature furnished by the tenderer may be written in another language so long as accompanied by an English translation of its pertinent passage in which case, for purpose of interpretation of tender, the English translation shall govern.

4. PRICE SCHEDULE/FINANCIAL BID

The tenderer shall complete the Price schedule on e-procurement Platform and should submit online only.

5. AWARD OF CONTRACT:

The Purchaser reserves the right to accept or reject any tender, and to annulment the tender process and reject all tenders at any time prior to award of contract, without thereby incurring any liability to the affected bidder or

bidders or any obligation to inform the affected bidder or bidders of the grounds for the purchaser's action.

The Managing Director, APMSIDC will be at liberty to terminate without assigning any reasons thereof the contract either wholly or in part on one month's notice. The tenderer will not be entitled for any compensation whatsoever in respect of such termination.

6. NOTIFICATION OF AWARD.

The Tenders shall be valid up to 90 days from the last date of receipt of tenders. Prior to the expiration of the period of tender validity, the purchaser will notify the successful tenderer in writing by registered letter, that its tender has been accepted. In exceptional circumstances, the purchaser may solicit the Tenderers consent to an extension of the period of the validity and in such case the responses there to shall be made in writing. The EMD provided shall also be suitably extended. Tenderer may refuse the request without forfeiting the EMD

7. ENTERING INTO AGREEMENT

Within 10 days of the receipt of the acceptance letter, the successful tenderer shall enter into agreement on Rs.100/- (Rupees one hundred only) worth Non-Judicial Stamp Paper. The specimen form of agreement will be supplied by APMSIDC. Failure of the successful tenderer to enter into constitute sufficient grounds for the annulment of the award, in which even the Purchaser may make the award to the next lowest evaluated tenderer or call for new tenders, duly forfeiting the E.M.D.

The purchase order will be issued from time to time for a specific quantity during the course of Rate Contract Period. The supplier should supply the goods at the rate for which the Agreement is concluded for the purchase orders placed.

8. CONDITIONS OF TENDER FOR THE SUPPLY OF DRUGS AND MEDICINES TO APMSIDC UP TO A PERIOD OF TWO YEARS FROM THE DATE OF AGREEMENT

I. APMSIDC reserves right to reject the tender of companies blacklisted by APMSIDC or any other State/National Organizations.

- II . The tenderers are requested to note that any taxes to be deducted at source at the rate fixed by the appropriate Govt. i.e. State / Central,
- III. The tenderer to whom supply contract is awarded shall not charge a higher price to APMSIDC than the price he quotes to any other Govt. organization or to a private Agency during the validity period of Rate Contract. If it is found that the firm has quoted a lower rate in another tender for another organization in the country and that it is not passing on the benefit of the lower rate to APMSIDC the differential amount will be deducted from the bills of the Firm. Such firms will be blacklisted for a period of 2 years.
- IV. The manufacturers who are awarded the supply contract are required to manufacture the drugs in their own units which are directly under the supervision of the board of Directors of the Company. Out sourcing of the supply from other units will not be permitted.

9. EARNEST MONEY DEPOSIT :

Every Tender should accompany an Earnest Money Deposit in the shape of Bank Guarantee or payment in online in favour of Managing Director, APMSIDC, Mangalagiri and this E.M.D. is refundable to the unsuccessful tenderers.

SSI firms located within the State of Andhra Pradesh (13Districts) holding Permanent Registration Certificate from the District Industries Centers of Department of Industries, Govt. of A.P. will be granted exemption from payment of Earnest Money Deposit.

10 . AGREEMENT

Each successful Bidder is directed to execute an agreement by furnishing Performance security and a non judicial stamp paper of value of Rs. 100/- (Stamp duty to be paid by the bidder) to the Managing Director, APMSIDC, within 10 days from the date of receipt of intimation that his bid has been accepted. The specimen form of agreement will be supplied by the Managing Director, APMSIDC.

11 . PERFORMANCE SECURITY DEPOSIT

- The successful tenderer shall be required to pay a Performance Security Deposit of 5% of the contract value subject to a maximum of Rs.10 lakhs

per product. The Performance Security Deposit should be paid in respect of each contract on or before the due date fixed, in the form of Bank Guarantee or Demand Draft drawn in favour of the Managing Director, Andhra Pradesh Medical Services & infrastructure Development Corporation, Mangalagiri, Guntur Dist.

- The Security Deposit furnished by such tenderer in respect of his tender will be returned to him upon complete fulfillment of the tender period or the extended period if any to the satisfaction of the Managing Director, Andhra Pradesh Medical Services & infrastructure Development Corporation, Mangalagiri, Guntur Dist.
- In the case of successful tenderers, the Earnest Money Deposit may, at the discretion of Managing Director, Andhra Pradesh Medical Services & infrastructure Development Corporation, Mangalagiri, Guntur Dist be adjusted towards the Security Deposit payable by him. Unsuccessful tenderers EMD will be released after signing of the agreement by successful tenderer or after expiry of tender validity period.

12. QUANTITIES, PRICES AND OTHER CONDITIONS

- The Quantity mentioned is only the probable requirement and may increase or decrease as per the decision of the Managing Director, APMSIDC. The rates should not vary with quantum of the order or the destination.
- Tenders have been called for on the generic names of drugs only. The bidders should quote the rates for the generic products and should supply generic drugs only. Imported drugs will be accepted with their original labels with which they were imported into India.
- If the DCA authorities not mentioned any particular packing while issue of license, all packing specifications/ volumes will be considered. But, if the DCA authorities issued license with particular packing specifications / volumes, then it will be treated as the license is issued for that particular specifications/ volumes only.
- Rates inclusive of all duties and taxes should be quoted for each of the required drugs, medicines etc., separately on FOR basis, in metric system units according to the unit asked for, together with manufacturer

name, license number under the Drugs and Cosmetics Act, 1940 (Central Act 23 of 1940), Composition Strength of the Drugs, medicines etc., offered. Tender for the supply of Drugs, Medicines, etc., with conditions and tender in which the rate is quoted for a unit other than the one asked for shall not be considered. No handling, clearing or transport charges etc will be paid. The deliveries should be made as stipulated in the purchase order placed with successful tenderers.

- Bidder has to quote the price online only in the price schedule before 5.00PM on **20.11.2019**.
- The total unit price should be inclusive of all taxes, packing and forwarding FOR destination etc.
- All the bidders shall quote price for each Bottle, Sachet, Injection and for the Tablets and capsules the price maybe quoted for 100 Tablets / Capsules. If any firm is having the special packing like 30,60 or 140 Tablets or Capsules they can quote their price for their packing description and shall be mentioned in remarks column.
- All rates quoted by the bidder should be valid for a period of two years from the date of acceptance of the lowest bid.

13. The price quoted by the tenderer shall not in any case, exceed the controlled price, if any fixed by Central / State Government and the maximum retail price [MRP]. The APMSIDC at its discretion, will exercise, the right to revising the price at any stage so as to confirm to the controlled price or MRP as the case may be. This discretion will be exercised without prejudice any other action that may be taken against the tenderer.

14. If the Supplier/Rate Contract firm has not started the supply to the point of delivery as noted in the schedule noted above, the purchase orders will be cancelled and the corporation is at liberty to make alternative purchase of the items of medicines for which the purchase orders have been placed from any other sources or from the open market or from any other tenderer who might have quoted higher rates at the risk and the cost of the supplier and in such

cases the corporation has every right to recover the cost and impose penalties as mentioned in other parts of this document.

Apart from risk purchase action, the tenderer shall also suffer forfeiture of the Security Deposit.

15. The rate quoted and accepted will be binding on the tenderer for the stipulated period and on no account will any variation in the price be entertained till the completion of this tender period.

16.1 The value of the NSQ drugs will be deducted from the bills amount payable to the firm or from the performance security deposit of the firm.

16.2 If the drug is not consumed prior to its expiry date i.e., six months before expiry, the supplier will notify about the short expiry drugs, upon receipt of such information the supplier should replace the short expiry/expired quantity with fresh stock of longer shelf life, otherwise the expired product will be returned to the supplier and the value equal to the cost of expired quantity will be deducted from the bills or any other amount payable to the firm.

17. NOTE:

- (a). No Tender shall be quoted for a product for which the company has been blacklisted either by A.P Rate Contract Committee or by any other state / Central Government organization
- (b). No Company which has been blacklisted either by AP Rate Contract Committee, APMSIDC or by any other State Government or Central Government Organizations shall participate in the bid during the period of blacklisting.
- (c). No Tenderer shall be allowed at any time on any ground what so ever to claim revision of or Modification in the rates quoted by him. Clerical error, typographical etc., Committed by the bidders in the bid forms shall not be considered after opening of the bids. Conditions such as “SUBJECT TO AVAILABILITY” “SUPPLIES WILL BE MADE AS AND WHEN SUPPLIES ARE

RECEIVED” etc., will not be considered under any circumstances and the bids of those who have given such conditions shall be treated incomplete and for that reason, shall be summarily rejected.

- d). The Tenderer should supply the items at the rate quoted by him during the contract period. If at any time the price of bidded items is reduced by any law or Act of Central or State Government. Or by the Bidder himself, the payment will be made at reduced rate.

18. LOGOGRAMS:

Bids for the supply of drugs and medicines etc., shall be considered only if the bidder gives an undertaking in his bid that the supply will be prepared and packed with the logogram either printed or embossed or affixed on tablets and capsules, bottles etc., as per the design enclosed in Annexure – I. All the tablets and capsules have to be supplied in standard packing of 10x10 or 5 x 10 x 10 in strip or blister packing with different colour of PVC on one side and with aluminum foil on other side with printed logogram and shall also conform to Schedule P1 of the Drugs and cosmetics rules wherever it applies. Affixing of stickers and rubber stamps shall not be accepted. Vials, ampoules and bottles containing items bided for should also carry the logogram. Bids of manufacturers who are not willing to agree to this condition will be summarily rejected. Failure to supply drugs etc., with the logogram will be treated as breach of the terms of agreement and render the bidder liable for forfeiture of the EMD and security deposit, in addition to recovery of any attributable loss incurred by the Andhra Pradesh Medical Services & infrastructure Development Corporation.

19) PACKAGING CONDITIONS

Drugs and medicines shall be supplied in the packaging specified for the drug and carry the logograms specified as per the Annexure I. The Drug and medicines shall be supplied with their coding.

Apart from the above, the packing in each carton shall be strictly as per specifications mentioned in **Annexure – IV**. Failure to comply with this shall lead to non-acceptance of the goods besides imposition of penalties.

- (a) The cap of bottles of preparations should not carry the name of the supplier.
- (b) The labels in the case of Injectables should clearly indicate whether the preparations are meant for IV, IM, ID & SC etc.,

Note :- If the supply is received in damaged condition in respect of primary packing, it shall not be accepted. In case of any acceptable deficiencies or damages in the secondary or tertiary packing, marking and documentation, the supply will be accepted only after levying penalties on the total value of supply to that destination. The penalties are Drug condition 2% , supplies in brand name 0.5%, non printing of AP GOVT logo 0.5% , MRP printing 0.5%)

20. QUALITY TESTING:

- Samples of supplies in each batch will be chosen at the point of supply of distribution/ storage points for testing. The samples will be Sent to different Laboratories approved by DCA for testing or as decided by the APMSIDC.
- The drugs shall have the active ingredients at the maximum permissible level throughout the shelf life period of the drug. The samples will be drawn periodically throughout the shelf life period.
- The supplies will be deemed to be completed only upon receipt of the Quality certificates from the laboratories. Samples, which do not meet quality requirements, shall render the relevant batches liable to be rejected. If the samples do not conform to statutory standards, the bidder will be liable for relevant action under the existing laws. APMSIDC has the right to destroy such substandard goods.
- The bidder should clearly understand that the decision of the Managing Director, Andhra Pradesh Medical Services & infrastructure Development Corporation or any officer authorized by him as to assess the quality of the supplied drugs, medicines etc., shall be final and binding.

- Each and every batch of drugs received from the firm will be subjected to quality test. As soon as the supply is made in the Central Drug Stores, the samples will be drawn by the quality control wing of APMSIDC and sent to the empanelled analytical testing laboratories for their quality testing as per pharmacopoeia and other standards.
- Whenever a particular Drug is declared as “Not of Standard Quality” (NSQ) by the above laboratories in assay , dissolution and for
- **Liquid Preparations** : Showing Presence of Fungus, Foreign matter, Non Dispersible Lump or Cake formation
- **Parental preparations:** Failing in Test for Sterility, Test for Pyrogen / Endotoxin or undue Toxicity.
- **Sera / Vaccine** : Failing in Test for Sterility, Toxicity , Moisture Content
- **Ophthalmic preparation:** Failing in test for Sterility, Fungal Growth, Foreign Matter.
- **Powders** : Fungal Growth

another sample of same batch will be sent to DCA / CDL on the request of the firm at their cost for second opinion. In case the drug is declared as NSQ in the second opinion, then the item of the firm will be blacklisted against the firm. The DCL/CDL declares the product as NSQ the item of the firm will be blacklisted.

- In case of other parameters, if 3 batches are declared as NSQ then only the item of the RC approved firm will be black listed against the firm.
- The amount of the NSQ batch shall be deducted from the amount payable to the firm or from the performance security deposit of the firm. No purchase orders will be placed for the black listed item of the firm.

- 1) In case a firm is supplying more than one product and one of the products is declared as NSQ, in such case, in addition to the measure suggested above, 25% of total bill amount submitted by the firm will be withheld for a period of four months and will be paid after monitoring satisfactory supply of all other products.

2) If two items of any firm are black listed then the entire firm will be black listed and it will not be allowed to participate in tender for 3 consecutive years from the date of blacklisting

- The supplier shall furnish to APMSIDC the evidence of Bio-availability and/or Bio-equivalence reports of the drugs upon demand for certain critical drugs.
- The supplier shall furnish the evidence of the basis for expiration dating and other stability data of the drugs on request by the Corporation.

21. ACCEPTANCE OF BIDS AND SUPPLY CONDITIONS

- The Managing Director, APMSIDC reserves the right to reject the bids or to accept the bids for the supply of all articles or for any one or more of the articles bided for in a bid without assigning any reason.
- The Managing Director, APMSIDC will be at liberty to terminate without assigning any reasons thereof the contract either wholly or in part on one month's notice. The bidder will not be entitled to any compensation whatsoever respect of such termination. The acceptance of the bids shall be communicated to the bidders in writing.
- The supply should be started within 45 days and should be completed within 60 days for Category A Products and 70 days for Category B from the date of receipt of purchase order in phased manner. If no supply is received even after 60/70 (based on the category) days of receipt of the purchase orders from the supplier, the MD, APMSIDC is authorized to impose a penalty at the rate of 0.5 % of the value of goods not supplied will be levied for each day delayed upto a maximum period of 20 days if Multiple bidders have been qualified and 30 days if single bidder is qualified , after this period the P.O. is deemed to be cancelled. Supply shall be

considered complete only upon receipt of at least 95% of the ordered quantity at each of the 13 District Warehouses and uploading of the required documents in eAushadhi software. The date of actual receipt of the goods or the date of uploading of the required documents whichever is later shall be considered as date of receipt of the material.

- For the drugs requiring the CDL Kasauli clearance for release of the batch, the supply period will be 100 days instead of 70 as these products need not have to undergo Quality analysis and can be issued from the date of receipt itself.
- If the bidder fails to execute the supply within the stipulated time, the APMSIDC is empowered to levy 10% penalty of the unexecuted value or differential cost incurred for the alternate purchase, whichever is higher.
- All the supplies will be scheduled for the period from the date of acceptance till the completion of the bid in installments, as may be stipulated in the supply order. The supplied medicines and drugs should have a minimum potency for the maximum period as prescribed in the Drugs and Cosmetics Act 1940 and rules there under on the date of supply.
- It shall be the responsibility of the bidder for any shortages, damages at the time of receipt in the Central drug Stores and APMSIDC is not responsible for receipt of drug for which no order is placed.

22. Inspection of the firms

Whenever corporation feels that it is necessary to inspect the firm and its manufacturing facility either by MD or any person/committee nominated by MD, the bidder shall provide all data, documentation and information without cost. If any adverse report is received in such inspection, APMSIDC will issue show cause notice to the firm. If the reply is found not satisfactory, the APMSIDC shall have the right to reject the bid or terminate/ cancel the orders already issued or not to issue any further order.

The tenderer, whose manufacturing unit is found to be not complying with GMP or WHO-GMP during inspection, will be levied with a fine of Rs.50,000/- or the expenditure incurred by the APMSIDC in conducting such inspection whichever

is higher. This fine amount shall be deducted from the EMD deposited by the bidder or from any other amount payable to them in any nature. The amount shall be deducted without any notice. In case of deficit, legal action will be taken against the bidder for recovery as per law.

23. PAYMENT PROVISIONS:

- a) No advance payments towards costs of GENERAL DRUGS & I.V. FLUIDS will be made to the bidder. However as far as possible Payment will be made within 30 days after receipt of material at Central Drug stores.
- b) Payments towards the supply of GENERAL DRUGS & I.V. FLUIDS Will be made strictly as per the rules of the Andhra Pradesh Health & Medical Housing & Infrastructure Development Corporation.
- c) No claims shall be allowed against the Andhra Pradesh Health & Medical Housing & Infrastructure Development Corporation in respect of interest on Earnest Money Deposit or on Security Deposit or late payments.
- d) In case of any enhancement in GST due to notification of the Government after the date of submission of tenders and during the tender period, the quantum of additional GST so levied will be allowed to be charged extra as a separate item Without any change in price structure of the GENERAL DRUGS & I.V. FLUIDS approved under the tender.
- e) Payments will be made after completion of 50% of supplies of order quantity and remaining will be paid after completion of 95% of supplies of order quantity at each of the 13 District warehouses.

24. PENALTIES :-

- If the successful bidder fails to execute the agreement and / or deposit the required security within the time specified or withdraws his bid after the intimation of the acceptance of his bid has been sent to him or other reasons, he is unable to undertake the contract, his contract will be cancelled and the Earnest Money Deposit shall stand forfeited to the Andhra Pradesh Medical Services & infrastructure Development Corporation and he will also be liable for all damages sustained by the Managing Director,

Andhra Pradesh Medical Services & infrastructure Development Corporation, by reasons of breach, such as failure to supply / delayed supply, including the liability to pay any difference between the prices accepted by him and those ultimately paid for the procurement of the articles concerned. Such damages shall be accessed by the Managing Director, Andhra Pradesh Medical Services & infrastructure Development Corporation whose decision is final in the matter.

- If any articles or things supplied by the bidder have been partially or wholly used or consumed after supply and are subsequently found to be in bad order, unsound, inferior in quality or description or are otherwise faulty or unfit for consumption, then the contract price or prices of such articles or things will be recovered from the bidder, if payment had already been made to him. Otherwise the bidder will not be entitled to any payment whatsoever for such article. For infringement of the stipulations of the contract or for other justifiable reasons, the contract may be terminated by the Managing Director, Andhra Pradesh Medical Services & infrastructure Development Corporation and the bidder shall be liable for all the losses sustained by the Andhra Pradesh Medical Services & infrastructure Development Corporation in consequence of the termination which may be recovered personally from the bidder from his properties, as per rules.
- In the event of supplies declared as NSQ, contract with the bidder will be suspended and purchases made from alternative suppliers. Such Firm may be black listed for three (3) years beginning from the date of blacklisting. The bidder shall also be liable for action under criminal law and the matter shall be notified to the concerned Licensing authority of the Drugs Control Administration.
- In all the above conditions, the decision of the Managing Director, Andhra Pradesh Medical Services & Infrastructure Development Corporation shall be final and binding.
- In the event of any dispute arising out of the bids such dispute would be subject to the jurisdiction of the Civil Courts within the city of Mangalagiri.

26. SAVING CLAUSE:-

- a.) No suit, prosecution or any legal proceedings shall lie against APMSIDC or any person for anything, which is done in good faith or intended to be done in pursuance of bid.
- b.) A supplier / firm whose products or the supplier/firm, itself have been blacklisted by the corporation which is displayed in the corporation website i.e. <http://msidc.ap.nic.in> may within 15 days from the date of display, appeal to the State Government i.e. Special Chief Secretary/ Principal Secretary to state Government who deals the subject of APMSIDC activities in HM&FW Department A.P.

27. Special Condition:-

- (a.) If any Company / Supplier quotes a lower rate to any other Agency / State in the country at a rate lower than the rate at which supplied to APMSIDC during the Rate Contract period, the differential amount is liable to be recovered. The Supplier/ Company should furnish undertaking (Annexure II) that they will remit the differential cost if they quote and supply at a lower rate to any other Agency / State etc. in the country than the rate quoted or at which supplied to the Andhra Pradesh Medical Services & Infrastructure Development Corporation during the rate contract period.
- (b.) Substantiated supply of any part of whole consignment without meeting the quality specification shall also entail blacklisting of the firm for a minimum period of three years for that particular product apart from recovery of loss and such other action as provided for under the Drug and Cosmetics Act 1940 by rules 1945 and conditions of the tender document.
- (c) If the product is in BP/USP, at the time of granting product license by the DCA and the product later comes under IP the product shall be supplied in IP only .
- (d) If the product is not in IP, the firms who are having product approval in BP/ USP are eligible to participate in the tenders

28) Penalty charges for delayed supply of drugs:

- 1) For Category A Products 60 days and for Category B products 70 days from the date of issue of PO. -- No penalty.

- 2) For the next 20 days i.e. 61st day to 80th day for Category A products and 71st day to 90th day for Category B Products -- 0.5% per day of the value of drugs received during this period. The Purchase Order will be cancelled beyond 20 days from the stipulated period of supply of drugs if the product has multiple bidders and if only single bidder is approved the Purchase Order will be cancelled beyond 30 days from the stipulated period of supply of drugs. If more than one Purchase Order is not supplied in the stipulated period of supply then the firm will be declared as undependable supplier for the products for which the supplies are not executed.
- 3) The corporation will be at liberty to procure the drugs from L2 firm (or) Other lowest bidder and even from open market as situation demands and recover the extra cost from the L1 firm in case the bidder fails to supply (the rate difference between Rate Contract rate and Procured rate) from the amount payable or from the performance Security deposit of the Rate Contract holder.

Note: a) Purchase orders will be placed based on necessity and requirements only.

b) List of Drugs are appended at Annexure VII . For clarity on Specifications please refer Annexure VII while quoting the rates.

29) Appeals

29.A). A supplier/firm who's product or the supplier/firm, itself have been blacklisted by the corporation which is displayed in the corporation website i.e.://msidc.ap.nic.in// may within 15 days from the date of display, may appeal to the Director General, Drug Control Administration, A.P

The Director General, Drug Control Administration, A.P., after such enquiry into the matter, as is considered necessary and after giving the said supplier an opportunity for representing his views, may pass such order in relation thereto as he thinks fit

29.B). If the firm is not satisfied with the outcome may appeal within 15 days to the Principle Secretary, Health, Medical & Family Welfare, A.P. for review.The State Government after such enquiry into the matter, as is

considered necessary and after giving the said supplier an opportunity for representing his views, may pass such order in relation thereto as it thinks fit.

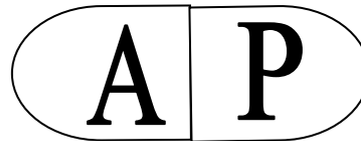
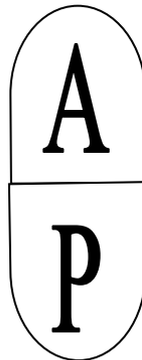
Managing Director

ANNEXURE -I

DESIGN FOR LOGOGRAMS

TABLET

CAPSULE



DESIGN FOR STRIP

REAR SIDE

AMPICILLIN 500mg	AMPICILLIN 500mg.	AMPICILLIN 500mg
	ANDHRA PRADESH GOVERNMENT SUPPLY NOT FOR SALE	
AMPICILLIN 500mg	AMPICILLIN 500mg.	AMPICILLIN 500mg
	ANDHRA PRADESH GOVERNMENT SUPPLY NOT FOR SALE	
AMPICILLIN 500mg	AMPICILLIN 500mg.	AMPICILLIN 500mg
	ANDHRA PRADESH GOVERNMENT SUPPLY NOT FOR SALE	
AMPICILLIN 500mg	AMPICILLIN 500mg.	AMPICILLIN 500mg
	ANDHRA PRADESH GOVERNMENT SUPPLY NOT FOR SALE	

MANUFACTURED BY

MFG. LICENCE NO:

BATCH NO:

DATE OF MANUFACTURE:

DATE OF EXPIRY:

SCHEDULE

NOTE:

BRAND NAME OF THE DRUG SHOULD NOT BE PRINTED ANY WHERE for indigenous products

Imported medicines accepted in brand name

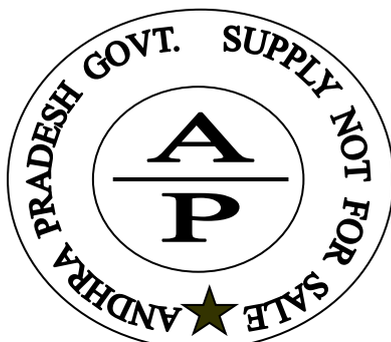
INJECTIONS

Injection in ampoule form should be supplied in Double constructed neck ampoules with the label bearing the words “Andhra Pradesh Govt. Supply – Not for sale” over printed and letter containing the logogram No. 1. Which will distinguish them from the normal rate packing.



Logogram No.1

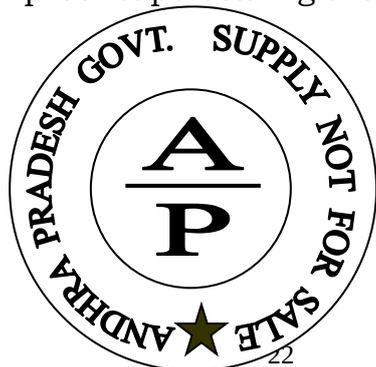
The vials should be supplied with aluminum seals containing the logogram.



In addition to the label bearing the logogram Andhra Pradesh Govt. Supply – Not for sale and the logogram No.1

LIQUIDS

Liquid preparations either be in HDPE bottles/ glass bottles depending on the nature of the product with pilfer-proof caps bearing the logograms:



On the top of the cap and the label to be affixed on the containers should bear a distinct colour different from the colour of the label of the trade packs and they should be over printed in red colour with the words Andhra Pradesh Government Supply -Not For Sale and the logogram above.

Top of the cap



SPECIMEN LABEL FOR OUTER CARTON

**A.P. GOVT.
SUPPLY
NOT FOR SALE**

(or)

**ఆంధ్రప్రదేశ్ ప్రభుత్వ
సరఫరా
అమ్మడానికి కాదు.**

DECLARATION

I do hereby declare that I will supply the drugs and medicines as per the above design.

Signature

ANNEXURE II

DECLARATION FORM

I/We _____ having our _____ Office at _____. The conditions of tender sent to me / us by the Managing Director, Andhra Pradesh Medical Services & Infrastructure Development Corporation, Mangalagiri for the tenders floated by him for the supply of drugs, medicines etc., for the tender for a period of two years from the date of acceptance and a Tender by all conditions set forth therein.

We hereby accept to supply the drugs at the accepted _____ (price) rates quoted by us in the tender document against the selected item or any matching price of Drug Price Control Organization (DPCO) as accepted by the department.

We will not quote & supply the drugs to the any agency / state in the country at the rate lower than the rate quoted in this tender.

If we quote lower rate than the rate quoted to the APMSIDC to any other agency / state in the country in future we will remit the differential cost to the APMSIDC.

I / We further declare that I / We posses valid Drug License bearing No.

valid upto

Signature :

Date :

Name of the
Firm and address :

Annexure-III

STATEMENT OF CAPACITY OF PRODUCTION

01. Name of the firm :

Address
Telephone
Telex

The installed capacity of this firm is as follows per shift

Tablets	Vials
Capsules	Dry syrups

Internal

(Liquids And colloids)

Syrups	Ampoules
--------	----------

External

Liquids

Signature of the tenderer: _____ Date _____

Full Name (IN BLOCK LETTERS) _____

NOTE :- Details are to be provided for two month's production capacity

Signature and seal of the Tenderer_____

ANNEXURE-IV

I. SCHEDULE FOR PACKING OF GENERAL DRUGS & I.V.FLUIDS

GENERAL SPECIFICATION :

- 1) No corrugated package should weight more than 15 kgs (ie. Product + inner carton + corrugated box). Except in the following cases :
 - a) Glutaraldehyde Solution
 - b) Conc. Heamodialysis Fluid
- 2) All Corrugated boxes should be of 'A' grade paper ie., Virgin.
- 3) All items should be packed only in first hand box only.

FLUTE :

- 4) The corrugated boxes should be of narrow flute.

JOINT :

- 5) Every box should be preferably single joint and not more than two joints.

STITCHING :

- 6) Every box should be stitched using pairs of metal pins with an interval of two inches between each pair. The boxes should be stitched and not jointed using calico at the corners.

FLAP :

- 7) The flaps should uniformly meet but should not over lap each other.

The flap when turned by 45-60° should not crack.

TAPE :

- 8) Every box should be sealed with gum tape running along the top and lower opening

CARRY STRAP:

- 9) Every box should be strapped with two parallel nylon carry straps. (They should not intersect).

LABEL:

- 10) Every corrugated box should carry a large label clearly indicating that the product is for “ A.P. Govt. Supply – Not for Sale”. The lower one third of the large label will indicate in bold the value of the product as depicted in Annexure III of this document.
- 11) The product label on the carton should be large atleast 15 CMS x 10 CMS dimension. It should carry the correct technical name, strength or the product, date of manufacturing, date of expiry, quantity packed and net weight of the box.

OTHERS:

- 12) No box should contain mixed products or mixed batches of the same product.
- 13) Every consignment of blood and blood related products should be certified as
 - a) AIDS Free
 - b) Hepatitis B Free

Strips of **aluminum foils** refer to gauge 04.

Aluminum foils as back material for blisters refer to gauge 025.

All hygroscopic drugs and sugar-coated tablets should be stripped in aluminum foil.

The rigid PVC used in blister packing should be of not less than 250 microns.

All glass bottles should be new, neutral glass and of USP type I glass.

I.V. Fluids bottles should be fitted with nipple caps.

Small tablets packed in blisters should be so packed to facilitate removal of the tablet without breaking / crushing.

Specifications of outer cartons are as given in the schedule. (annexure VII)

In case of any conflict between carton specifications and packets per carton specifications (Last column of this table), the specification of the carton / annexure VIII shall prevail.

All tablets should have score line.

All liquid orals should be provided with a measuring device

All plastic containers should be made of virgin grade plastic of HDPE/LDPE.

All plastic jars above 450 gms/ml should carry an inner plastic lid and should be of HDPE.

II SPECIFICATION FOR CORRUGATED BOXES HOLDING TABLETS/CAPSULES/PESSARIES

- 1) The box should not weigh more than 7-8 kgs. The grammage should be 120-150 gsm (outer paper should be 150 gsm and others may be 120 gsm).
- 2) The box should be of 5 ply with Bursting strength of 9 kg/Cm²

III SPECIFICATION FOR IV FLUIDS

- 1) Each corrugated box may carry a maximum of 50 bottles of 100 ml in 2 rows with individual sealed polythene cover center partition pad, top and bottom pads of 3 ply.
- 2) Grammage : 120-150 Gsm(outer paper 150 gsm; Others may be 120 gsm).
- 3) Ply : 5 or 7 ply.
- 4) Bursting Strength : Not less than 12 Kg/Cm²

IV SPECIFICATION FOR LIQUID ORALS: 50 ml to 120 ml bottles.

- 1) 100 bottles of 50 ml or 60 ml may be packed in a single corrugated in 2 rows with top, bottom and center pad of 3 ply.

50 bottles of 100 ml – 120 ml may be packed in a similar manner in a single corrugated box.
- 2) If the bottles are not packed in individual carton, 3 ply partition should be provided between each bottle. The measuring device should be packed individually.
- 3) Grammage : 120 – 150 Gsm (outer paper 150 gsm; others may be 120 gsm)
- 4) Ply : 7 Ply
- 5) Bursting Strength: Not less than 12Kg/Cm².
- 6) In case the box is heavier than 7 kg but less than 10 kg, the grammage may be 150gsm (outer paper 150gsm; others may be 120 gsm) 5 Ply and bursting strength should not be less than 9kg /Cm².

V. SPECIFICATION FOR INJECTABLES(IN VIALS)

- 1) Vials may be packed in corrugated boxes weighing upto 15 Kgs. Ampoules should be packed in C.B weighing not more than 8 Kgs.
- 2) CB for vials should be of 150 Gsm (outer paper 150 Gsm; others be 120 Gsm) and 3 ply, while CB for ampoules should be of 150 Gsm (outer 150 Gsm; others may be 120 Gsm) and 5 ply.
- 3) Bursting strength for CB boxes for
 - a) Vails : Not less than 12 Kg/Cm²
 - b) Amp : Not less than 9 Kg/Cm²
- 4) In the case of 10ml Ampoules 100 or 50 Ampoules may be packed in a Grey board box. Multiples of grey board boxes packed in CB. In case of ampoules larger than 10 ml only 25 ampoules may be packed in a Grey board box with partition.
- 5) If the vials are packed in individual carton, there is no necessity for grey board box packing. The individual carton may be packed as such in the CB with center pad.
- 6) In case of ampoules every grey board box should carry 5 ampoule Cutters and to be placed in a polythene bag.
- 7) Eye and ear drops Vials has to be packed in plastic bottles of HDPE/LDPE with HDPE cap and should be packed in an individual carton and they should be packed in 50's in a Grey board box.

VI. SPECIFICATIONS FOR OINTMENT / CREAM / GELS PACKED IN TUBES:

- 1) No corrugated box should weigh more than 7-9 kgs.
- 2) Every Ointment tube should be individually packed in a carton and then packed in 20's in a Grey board box, which may be packed in a corrugated box.
- 3) Grammage : 120-150 gsm (outer paper
150 gsm; others may be 120 gsm).
- 4) Ply : 5
- 5) Bursting Strength: Not less than 9 Kg/Cm²

VII. SPECIFICATIONS FOR ORS

- 1) THE SACHETS SHOULD BE OF Aluminum foil laminated with glassine or heat sealable plastic film, outer paper may contain label information.
- 2) 50 sachets may be packed in grey board boxes and 10 grey board boxes in a CB.

- 3) grammage : 120-150 gsm (outer paper
150 gsm; others may be 120 gsm)
- 4) Ply : 5
- 5) Bursting strength : Not less than 9 Kg/Cm²

**Annexure -V
DECLARATION**

I,.....S/o..... aged about..... Years Resident of.....do here by affirm on oath as under.

That I am Managing Director/Director/Partner/Proprietor of M/s.....on whose behalf an application for grant of License to manufacture Drugs has been made to the Licensing Authority.

That I am responsible for the day to day affairs and conduct of business of M/s..... for the purpose of Section 34 of the Drugs and Cosmetics Act, 1940 to which M/s..... and its Director/Partners etc., are held liable for any act of omission punishable under the Drugs and Cosmetics Act, 1940 and other enactment enforced by the Officers of Drugs Control Administration.

That in the event of any change in the constitution of the Company, I will inform the concerned licensing authority.The following are the Directors/Partners of the Company as on date and whose Names and permanent address are given below:

Name . S/O Age Residential Addresses .

- 1.
- 2.
- 3.

WITNESSES WITH FULL ADDRESS:

- 1.
- 2.

I, Srido hereby declare on oath that the above contents are true to the best of my knowledge and belief and nothing has been hidden.

DEPONENT.

Annexure –VI MODEL PRICE BID FORMAT



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Cost Component Details

Tender Details

Tender No : test/sai/aphrmhdc

Department : APHMHIDC-APHMHIDC

Tender Subject : test tender

Tender Type : Open

Default Cost Component

Item Name	Type	Value/Percentage	Remarks
Excise Duty	Exclusive (%)	2.00	
CST	Exclusive (%)	2.00	dsfsd
LST/ VAT	----select----		
Discount	Exclusive (%)	2.00	sdfsd
Packing Charges	Exclusive (Value)	2.00	dsfsdfs
Remarks	dsfsd		

Note: 1. Inclusive - Included in the Basic Price
 2. Exclusive - Separately Calculated from Basic Price
 3. Applicable for all Items.

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Price Bid

Aluminium Hydroxide Gel IP Aluminium Hydroxide Gel IP

UOM - bottles

Location	Total NIT Quantity	Brand/Make/Model	Basic Price Per Unit(INR) A	Basic Price per Unit Rate In Words	Cost Component	Total Cost component per Unit(INR) B	Total Landed Cost per Unit(INR) (A+B)
dsf(345)	345	LG	100.00	one hundred	Add	(8)	(108)

Note: 1. On Submission of PriceBid with Department Public Key the price data would be encrypted and will not be Visible.
 2. In Case of Edit/Update Price Bid after successful bid submission the last price bid submitted for the Item would be Superseded.

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Note: Price Bid should be submitted online only

Annexure VII List of items

S.No	Item Code	Item Name	Description	Strength	Unit	Probable required Tender Qty per Annum
Category A						
1	736 DG	Albendazole Tablets Chewable IP 400 mg	10 X 10 Or 50 X 10 Blister With Aluminum Foil Pack (Albendazole 400 Mg Tablets Chewable That Meets The New Amendment To I.P. -2018 i.e. Dissolution Test)	400 mg	100	40000000
2	1189 DG	Amikacin Sulphate Injection IP 500 mg / 2 ml	10X 10 / 5 X 10 , Type I glass	500 mg / 2 ml vial	1	2950700
3	1432 DG	Glimepride Tablets IP 1mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	1mg	100	196359000
Category B						
4	2200 DG	5-Fluorouracil Injection IP	1 X 10 Amples Type 1 Glass	500 mg / 10 ml	1	95040
5	1728 DG	Acenocoumarol Tablets IP 1 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	1mg	100	79500

6	1523 DG	ACT Kit for (9-14 years)	Per One Kit	9-14 Years:-Artesunate Tablets 50mg (3 tabs) + Artesunate Tablets 100mg (3 tabs), Pyrimethamine IP 25mg, Sulphadoxine IP 500mg (2tabs) -RED COLOR Blister.	1	6990
7	1522 DG	ACT Kit for Adults	Per One Kit	Adult:- Artesunate Tablets 200mg (3tab),Pyrimethamine IP 37.5mg, Sulphadoxine IP 750mg (2tabs) -WHITE COLOR Blister.	1	15050
8	2452 DG	Activated Charcoal Tablets IP 500mg Tablets	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	500mg	100	100000
9	1731 DG	Acyclovir Ophthalmic Ointment 3% I.P	1 X 25 (or) 1 X 50	5gm	1	3180
10	1614 DG	Ambroxal Syrup 30mg/5ml 60 ml bottle	60ml x 100 amber colour pet bottle	30mg/5ml 60 ml bottle	1	2016350

11	1615 DG	Amino acids essential IV 500ml as per IP	Each Glass Bottle	Each ml. Contains:L- Afgioine Hydrochloride / USP 2.7mg L- HistidineHydrochloride H2O / BP 1.4mg L- Isoleucine USP 1.8mg L- Leueine USP 4.1mg L- Lysine Hydrochloride USP 7.4mg L-Methionine USP 2.4mg L- Phenylalanine USP 2.9mg L- Threonine USP I.8mg L- Tryptophan	1	34100
12	1736 DG	Amiodarone Hcl Injection IP 150 mg /3 ml	10 x 10 / 5 x 10 type-1 glass	150 mg /3 ml	1	10000
13	1738 DG	Atamoxetine Capsules 10mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	10mg	100	525600
14	2211 DG	Atropine Sulphate eye ointment 1% 3.5gm	1 X 10 Tubes	1% 3.5gm	1	93010
15	2213 DG	Barium Sulfate Suspension 100% w/w 340 gms	1 X 24 Pack	100% w/w 340 gms	1	690
16	2456 DG	Benzathene Penicillin LA 12 lakh Injection 1 gm / vial	1 x 25	12 lakh Injection 1 gm / vial	1	10000

17	520 DG	Benzyll penicillin Injection IP 10 lakhs, Unit	With 10 ml Water For Injection	10 lakhs, Unit	1	5000
18	739 DG	Betamethasone Dipropionate Ointment USP 0.1% 10 gms tube	1X10 (or) 1X 25 Tetrapack	10 gms tube	1	1693530
19	1455 DG	Betaxolol eye Drops 0.25%, 5 ml	5ml X 5 X 5 OR 10 X5 FFS/BFS	5 ml	1	20630
20	58 DG	Bisacodyl Tablets IP 5 mg(Enteric Coated)	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5 mg	100	11084000
21	1746 DG	Bupropion Hcl Tab 150 mg USP	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	150 mg USP	100	50000
22	2223 DG	Calciferol oral Drops 75 mcg/ml 20 ml	1 X 25 Bottles	75 mcg/ml 20 ml	1	622130
23	6632 DG	CAPD (Continous Ambulatory Peritoneal Dialysis) Solution Set 4.25%, 2 Litres Bag With Integrated Asymmetric Y.Set	2 Ltrs bag with integtraed asymmetric Y set	CAPD (Continous Ambulatory Peritoneal Dialysis) Solution Set 4.25%, 2 Litres Bag With Integrated Asymmetric Y.Set	1	2000
24	2070 DG	Carbmazepine Tab 400mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	400mg	100	2498500
25	2232 DG	Carbolic Acid (Phenol) 500ml bottel	1 X 20 Bottles Type 1 Glass	500ml	1	29030

26	1755 DG	Chlorhexidine Gluconate+ Metronidazole Oral Gel	10 X 10	Chlorhexidine gluconate 0.25 % with Metronidazole 1 % w/w Oral gel 10 gm	1	102720
27	1295 DG	Chlorpromazine Tablets IP 100 mg,	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	100 mg	100	975500
28	123 DG	Ciprofloxacin Hcl Eye /Ear Drops IP 0.3% w/v in 5 ml	5ml x 5 x 5	0.3% w/v in 5 ml	1	3590940
29	2242 DG	Cisplatin Injection IP 50 mg / 10 ml vial	1X25 Vial	IP 50 mg / 10 ml vial	1	13500
30	2246 DG	Clonazepam 0. 5mg Tablet	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	0.5mg Tablet	100	1623700
31	2248 DG	Clopidogrel 150mg Tablet	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	150mg	100	1579800
32	1771 DG	Cyclophosphamide Injection IP vial 500mg	10 ml Water For Injection	500mg	1	5000
33	1770 DG	Cyclophosphamide Tablets IP 50mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	50mg	100	65400
34	1774 DG	Cyclosporine Capsules USP 25 mg	10 x 10 Blister With Aluminum Foil Pack	25mg	100	5000
35	2260 DG	Dexmedetomidine 100mcg Injection	1 X 25 Vials	100mcg	1	13000
36	2261 DG	Dextran 40% I.V 500 ml	1 X 24 Bottles	40% I.V 500 ml	1	8360
37	2265 DG	Dicyclomine drops 10mg/ml, 20 ml Bottle	1 X 50 Bottles	10mg/ml, 20 ml Bottle	1	38950

38	2267 DG	Diphtheria Anti Toxin 10000 IU/ 10 ml ampoule	1 X 10 Vials/ Ampoules	10000 IU/ 10 ml	1	1150
39	2449 DG	Doxophylline Tab 400mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	400mg	100	500000
40	2274 DG	Doxylamine succinate Tablet 25 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	25 mg	100	126240
41	2278 DG	Empagliflozin Tablet 10mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	10mg	100	162900
42	2450 DG	Ethamsylate Tab 250 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	250 mg	100	500000
43	1800 DG	Etoposide Capsules IP 100 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	100 mg	100	240
44	1799 DG	Etoposide Injection 100 mg / 5 ml vial	5 x 5 type-1 glass	100 mg / 5 ml vial	1	2000
45	1803 DG	Fenofibrate Capsules USP 160mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	160mg	100	201800
46	2285 DG	Fentanyl Transdermal Patch 50mcg/hr	1 X 10 Patches (or) 1 X 2 Patches	50mcg/hr	1	12320
47	2090 DG	Fluconazole For Oral suspension 50mg per 5ml (Pack Size : 35ml)	1x10x10 or 240	50mg per 5ml (Pack Size : 35ml)	1	1000
48	2291 DG	Fluconazole Injection 2 mg/ ml 100 ml bottle	1 X 24 Bottles	2 mg/ ml 100 ml bottel	1	16120
49	2294 DG	Flurometholone Eye Drops 0.1%, 5ml vial	1 X 50 Vials	0.1%, 5ml vial	1	13500

50	2299 DG	Gatifloxacin (0.3% w/w) + Dexamethasone (0.1% w/w) eye Drops in 5ml	1 X 25 Vials	(0.3% w/w) + (0.1% w/w) in 5ml	1	104100
51	303 DG	Gentamicin Ear Drops IP 0.3% in 5ml w/v	5ml X 5 X 5 OR 10 X5 FFS/BFS	0.3% in 5ml w/v	1	1759410
52	1688 DG	Glimepride Tablets IP 2mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	2mg	100	66368000
53	1819 DG	Homatropine eye Drops IP 2% 5 ml	5 X 5	2% 5 ml	1	2000
54	1824 DG	Ichthymol 1.5gm Glycerine upto 15ml NFL III	5ml X 5 X 5 FFS	Ichthymol 1.5gm Glycerine upto 15ml NFL III	1	4730
55	2311 DG	Inj. Glycopyrrolate (0.5mg) + Neostigmine (2.5mg) /5 ml	1 X 10 Amps	(0.5mg) + (2.5mg)/5 ml	1	68298
56	2149 DG	Isoniazide (INH) Tab. 300mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	300mg	100	1920
57	2185 DG	Isoniazide (INH) Tab.100mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	100mg	100	720
58	1843 DG	Isoprenaline Hcl Injection IP 200 micrograms / ml	2ml amp 10 x 10 / 5x10 type-1 glass	200 micrograms / ml	1	2000
59	1844 DG	Isosorbide Dinitrate Tablets 20mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	20 mg	100	1387600
60	2104 DG	Ivabradine HCL Tablets 5 mg	10 X 10 or 50X10 Blister With Aluminum Foil Pack (or) 14X4	5mg	100	42300

61	1847 DG	L- Asparaginase Injection vial 5000 IU	Each	5000 IU	1	2000
62	1853 DG	Letrozole Tablets 2.5mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	2.5mg	100	148500
63	1856 DG	Levodopa 100mg +Carbidopa 10mg Tablets I.P	30 Capsules in a HDPE White Colour Bottle	Levodopa 100mg +Carbidopa 10mg Tablets I.P	100	10000
64	2324 DG	Levofloxacin Injection 5 mg/ml 50 ml vial	1 X 10 Vials	5 mg/ml 50 ml vial	1	439290
65	1857 DG	Lignocaine Eye Drops BP 4 % in 5ml vial	5ml X 5 X 5 OR 10 X5 FFS/BFS	4 % in 5ml	1	15560
66	757 DG	Lignocaine Hcl Gel IP 2% in 30 gms tube	30 gm x 10x10 tubes	2% in 30 gms tube	1	229740
67	2329 DG	Lignocaine Hydrochloride Gel IP 10% 50 ml	1 X 10 Vials	(Lignocaine Gel, Lidocaine Hcl Gel, Lidocaine Gel): Strength: 1%> w/v and 2% w/v	1	5350
68	2331 DG	Linezolid Injection 2mg/ml, 300 ml vial	1 X10 Vial	2mg/ml, 300 ml vial	1	41060
69	2336 DG	Lorazepam Injection 2mg/ml, 2ml ampoules	1 X 50 Amps	2mg/ml, 2ml	1	211590
70	2337 DG	Lorazepam Tablet 2mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	2mg	100	1143900
71	2458 DG	Loteprednol eye drops 0.5%w/v	1 x 25	0.5%w/v	1	5000

72	1867 DG	Melphalan Tablets IP 5 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5 mg	100	600
73	1870 DG	Mercaptopurine Tablets IP 50 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	50 mg	100	2400
74	2455 DG	Meropenam Injection IP 1gm Vial	1 x 25	1gm Vial	1	100000
75	1873 DG	Methimazole Tablet 5mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5mg	100	38600
76	1875 DG	Methotrexate Injection IP 50 mg / ml in 2ml amp	2ml amp10 X 10 TYPE-1 GLASS OR 5X 10	50 mg / ml in 2ml amp	1	3650
77	1879 DG	Metolazone Tablets USP 5mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5mg	100	105700
78	1829 DG	Midazolam Inj 1mg/ml 10ml	10ml vial 10ml x 5 x10	Inj 1mg/ml 10ml	1	141820

79	1644 DG	Multivitamins Tablets (Therapeutic)	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	Formula: Methylcobalamin 500mcg+ Ascorbic acid 100mg+Niacinamide 50mg+Vit E25mg+ Calcium Pantothenate 12.5mg+ Vit B1 10mg + Riboflavin 10mg + Vit B6 3mg+ Vit A 5000IU+ Folic Acid 1mg + Zinc Oxide 15mg + Copper Gluconate 2.5mg + Manganese 1.4mg+ Chromium 65mcg+ Selenium 60mcg Tablets for Vitamin and mineral deficiency states in Adult patients.	100	50000000
80	2347 DG	Mupirocin Oint. 2% 15 gm	1 X10 Tubes	2% 15 gm	1	84740
81	2348 DG	Nacl 3% (Sodium Chloride) ,100ml bottle I.V	1X24 Bottles	100ml bottle I.V	1	139400
82	2457 DG	Natamycin eye drops 5%w/v	1 x 25	5%w/v	1	5000
83	1314 DG	Neostigmine Injection IP 0.5 mg/ ml in 5ml vial	5ml x 5 x 5 type- 1 glass vial / amp	0.5 mg/ ml in 5ml vial	1	289610
84	2359 DG	Nifedipine Retard Tablet 20mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	20mg	100	287000
85	1897 DG	Nitroglycerine (N.T.G.) Injection 5mg/ml in 5 ml	10 x 10 / 5 x 10 type-1 glass	5mg/ml in 5 ml	1	50000

86	1900 DG	Non Ionic Contrast Media (Iohexol) 400/50 (50ml)	Non Ionic Contrast Media (Iohexol) 400/50 (50ml) Mono Pack	400/50 (50ml)	1	260
87	1901 DG	Octreotide Injection 100mcg/ml	1ml AMP 10 X 10 OR 5 X 10 TYPE-1 GLASS	100mcg/ml	1	1000
88	1660 DG	Ofloxacin ophthalmic solution IP 0.3%w/v in 5ml vial	5ml X 5 X 5	0.3%w/v in 5ml vial	1	65680
89	1902 DG	Ofloxacin Tablets 100mg	5X10X10 BLISTER WITH ALUMINUM FOIL PAC	100mg	100	2863000
90	2369 DG	Olmesartan Medoxomil Tablets 40mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	40 mg	100	365900
91	2459 DG	Olopatadine eye drops 0.1%w/v	1 x 25	0.1%w/v	1	5000
92	1317 DG	Ondansetron Tablets IP 4 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	4 mg	100	5992000
93	2454 DG	Paracetamol Paediatric Oral Suspension IP 15ml,125mg/5ml	1x10	15ml,125mg/5ml with 1ml pillar	1	1300000
94	2453 DG	Paracetamol Tablets IP 650mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	650mg	100	50000000
95	2380 DG	Phenobarbitone Injection 200mg/ml,1ml Ampoules	1 X50 Amp	200mg/ml,1ml	1	92620
96	1447 DG	Phenylephrine HCL Eye Drops USP 5% 5 ml	5ml X 5 X 5 OR 10 X5 FFS/BFS	5% 5 ml	1	5000
97	1651 DG	Phenytoin sodium Syrup 25mg/5ml 60ml bottle	60ml X 100 Amber Colour Pet Bottle	25mg/5ml 60ml bottle	1	5000

98	1167 DG	Phenytoin sodium Tablets IP 100mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	100mg	100	18649700
99	2384 DG	Piracetam 800mg Tablet	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	800mg	100	213900
100	1839 DG	Piracetam Tablets 400mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	400mg	100	35400
101	104 DG	Povidone Iodine Scrub IP 7.5% W/V,500ml	500 ml Bottle Packing: 500 ml Bottle Carton: 20X 500ml	7.5% w/v,500ml	1	70400
102	1914 DG	Praziquantel Tablets IP 600 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	600 mg	100	11400
103	576 DG	Prednisolone Tablets IP 5mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5mg	100	17508000
104	1519 DG	Primaquine Tablets IP 7.5 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	7.5 mg	100	2061900
105	2391 DG	Probio sachet (Ig)	1 X 25 Sachets	Total Probiotic not less than 1.25 billion CFU (Lactobacillus acidophilus, Lactobacillus rhamnosus, Bifidobacterium longum, Saccharomyces boulardii)	1	204960

106	1925 DG	Pyridostigmine Tablets BP 60 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	60 mg	100	10000
107	2393 DG	Pyridoxine HCL CR Tab. 100mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	100mg	100	144000
108	2190 DG	Pyridoxine Hcl Tablets IP 25 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	25 mg	100	5986700
109	574 DG	Pyridoxine Hcl Tablets IP 5 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5 mg	100	200000
110	2396 DG	Quadrivalent Influenza Injection vaccine	10 Doses Per Vial (Vaccine in Vial)	1. An A/Michigan/45/2015(H1N1)pdm09-like virus; 2. an A/Singapore/INFIMH -16-0019/2016(H3N2)- like virus; and 3. aB/Phuket/3073/2013 -like virus (B/Yamagata/16/88 lineage). It is recommended that quadrivalent vaccines containing two influenza B viruses contain the above three viruses and B/Brisbane/60/2008- like virus (Victoria Lineage)	1	3750
111	2399 DG	Quinine Sulphate Injection 600mg/ 2ml Ampule	1 X 50 Amps	600mg/ 2ml	1	217280

112	2407 DG	Risperidone 0.5mg Tablet	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	0.5mg	100	325500
113	2409 DG	Rocuronium Bromide Injection 10mg/ml, 2.5ml Vials	1 X 25 Vials	10mg/ml, 2.5ml Vials	1	18760
114	2410 DG	Ropinirole ER Tablets 1 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	1 mg each film coated tab contains; Ropinirole Hcl equi. To Ropinirole 1 mg	100	147000
115	2414 DG	S. Metoprolol PR Tablet 25mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	25mg	100	525800
116	635 DG	Salbutamol Sulphate Tablets IP 2 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	2 mg	100	85410600
117	2421 DG	Sertaconazole Nitrate powder 2%w/w 100gram bottle	1 X 25 Bottles	2%w/w 100gram bottle	1	370370
118	1948 DG	Silicon Oil Injection 1000mm 2/5 (cst) Injection 10ml Vial	5 X 5 , TYPE-1 GLASS	1000mm 2/5 (cst) Injection 10ml Vial	1	100
119	2425 DG	Sodium Valporate Injection 500mg / 5ml Vial	1 X 25 Vials	500mg / 5ml Vial	1	59300
120	2426 DG	Sodium Valporate Syrup 200mg / 5 ml,100ml Syrup	1 X 50 Bottles	200mg / 5 ml,100ml Syrup	1	102830
121	2428 DG	Spiramycin Tablet 3.0 M.I.U	10 X 10 Aluminium Blister Packs	3.0 M.I.U	100	1800

122	1120 DG	Stable Bleaching Powder 25Kgs As per IS 1065-1989 Grade-I or equivalent 34%	Each	25Kgs As per IS 1065-1989 Grade-I or equivalent 34%	1	516550
123	611 DG	Streptomycin Sulphate Injection IP 0.75g	Streptomycin Sulphate Injection Ip 0.75G	0.75g	1	2000
124	1957 DG	Sumatriptan Succinate Tablets BP 50mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	50mg	100	10000
125	2431 DG	Suspension posaconazole oral 40 mg / ml bottle of 105 ml	1 X 25 Bottles	40 mg / ml 105 ml	1	300
126	1961 DG	Tacrolimus Capsules 5 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5 mg	100	18900
127	1962 DG	Tamoxifen Citrate Tablets IP 20 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	20 mg	100	162600
128	1963 DG	Tamsulosin HCL Capsules 400 mcg BP/USP	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	400 mcg	100	98100
129	2451 DG	Teneligliptin 20 mg	1 X 7 (or) 10 X 10 Blister With Aluminum Foil Pack	20 mg	100	250000
130	1275.01 DG	Tetanus Vaccine (Adsorbed) IP 0.5ml Ampoule	120Amps x 12	0.5ml Ampoule	1	400000
131	2433 DG	Thiamine Tablet 50mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	50mg	100	5359900
132	2436 DG	Tramadol Capsule 50 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	50 mg	100	3879500

133	1970 DG	Tranexamic acid Tablets BP 250mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	250mg	100	464200
134	678 DG	Trifluoperazine Tablets IP 5 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5 mg	100	200000
135	1373 DG	Tropicamide Eye drops BP 0.5% in 5ml vial	5ml x 5 x 5 or 10 x5 BFS / FFS	0.5% in 5ml vial	1	19290
136	1330 DG	Vecuronium bromide Injection 4 mg / 2 ml	2ml amp 10 X 10 OR 5 X 10 TYPE-1 GLASS	4 mg / 2 ml	1	401440
137	1974 DG	Vincristine Injection IP 1mg/ml in 1ml ampoule	1ml AMP 10 X 10 OR 5 X 10 TYPE-I GLASS	1mg/ml in 1ml ampoule	1	1000
138	1976 DG	Warfarin sodium Tablets IP 3 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	3 mg	100	33300
139	1975 DG	Warfarin sodium Tablets IP 5 mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	5 mg	100	5000
140	1718 DG	Zinc Syrup in 20 mg /5 ml in 100 ml bottle	100ml x 20 amber colour pet bottle in one box	Zinc Syrup in 20 mg /5 ml in 100 ml bottle	1	629500
141	2441 DG	Zolpidem tablets 10mg	10 X 10 (or) 50 X 10 Blister With Aluminum Foil Pack	10mg	100	155100
142	2443 DG	Zuclopenthixol (Acetate / Decanoate) Injection I.P 50 mg	1 X 25 Vials	50mg	1	10560

Annexure – VIII (Check list)

Sno	Name of the Document	Submission	Page no
1	Processing Fee: The bidder shall remit processing fee Rs. 5625/- online to the account of Managing Director, APMSIDC, Managalagiri (Account No 142410011000314, IFSC code : ANDB0000366) and upload the original transaction slip with UTR number. Failure to pay the processing fee and submitting the slip in the aforesaid manner will lead to rejection of the bid.	Online	
2	Valid manufacturing License issued by the concerned Drug control authority/ Valid Import License for Importers	Online	
3	Valid Product wise license for manufacturing the products issued by the concerned Drug Control authorities/Valid import license if the product is imported.	Online	
4	Annual Turnover certificate by Chartered Account.	Online	
5	The Earnest Money Deposit (EMD) shall be Rs.3 lakhs to each firm. The Earnest Money Deposit shall be paid in the form of Bank Guarantee or online to the account of Managing Director, APMSIDC, Managalagiri (Account No 142410011000314, IFSC code : ANDB0000366) and upload the original transaction slip with UTR number. Failure to pay the EMD and submitting the slip in the aforesaid manner will lead to rejection of the bid. The bidders should note that the local MSME/SSI units are exempted from payment of E.M.D, subject to production of necessary documents to that extent by them.	Online	
6	Non Conviction Certificate issued by DCA authorities for manufacturers and for direct importers that the firm has not been convicted for the last 3 years (2015-16, 2016-17 and 2017-18 or 2016-17,2017-18 and 2018-19) issued on or after 01.05.2019.	Online	
7	WHO-GMP (WHO-Good manufacturing practices Certificate) Certificate issued by the Licensing Authority. The WHO-GMP certificate must not be older than one year from the due date of Bid submission in the case where validity is not mentioned in the certificate. The WHO-GMP certificate of all the manufacturing plants, of which products have been quoted, should be submitted. The Importer should produce WHO-GMP /COPP of the manufacturing firm or a certificate which is at	Online	

	<p>par with WHO-GMP issued by exporting countries like US-FDA approval, etc. In the case of imported drugs, labels and product literature of all quoted products must be submitted.</p> <p>The Firm will continue to hold WHO-GMP certificate for the product during entire rate contract period of the product. If WHO-GMP certificate expires, it is firm's responsibility to inform APMSIDC about the same and not to accept any further purchase order till re-issue / renewal of WHO-GMP certificate. During the period of non validity of WHO-GMP certificate of the firm the rate contract will deemed to be suspended. If the firm fails to inform APMSIDC about the expiry of WHO-GMP certificate and accept purchase order of APMSIDC and later on it comes to the knowledge of APMSIDC in this situation firm shall be liable for a panel action.</p>		
8	Authorization of a senior responsible Person of the company with Authority to transact business.	Online	
9	Memorandum and articles of association of the company along with List of Directors downloaded from the website of ministry of company affairs, signed by MD/Director / Company secretary.	Online	
10	Self declaration of production capacity.	Online	
11.	Self declaration of batch size for each quoted product.	Online	
12.	Audited balance sheets of any of the three of last three years, 2015-16, 2016-17 and 2017-18 or 2016-17,2017-18 and 2018-19).	Online	
13	The details of GST i.e. Number etc, should be enclosed in Online	Online	
14	Declaration form with details of Firm contact number, email ID and address (as per Manufacturing license)	Online	

Managing Director