

e-TENDER NO:- BPPI/DRUG-054/2017

TENDER FOR SUPPLY OF DRUGS

TO

Bureau of Pharma Public Sector Undertakings of India (BPPI)

For the year 2017-19



BUREAU OF PHARMA PUBLIC SECTOR UNDERTAKINGS OF INDIA

(Set up under the Department of Pharmaceuticals, Govt. of India)

IDPL corporate office Complex, Old Delhi-Gurgaon Road, Dundahera, Gurgaon 122016

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BPPI/DRUG-054/2017

BUREAU OF PHARMA PUBLIC SECTOR UNDERTAKINGS OF INDIA (BPPI)

(SET UP UNDER THE DEPARTMENT OF PHARMACEUTICALS, GOVERNMENT OF INDIA)

Regd. Office: Core No. 6, First Floor, SCOPE Complex, Lodi Road, New Delhi-110003

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ONLINE TENDER FOR THE SUPPLY OF DRUGS TO BUREAU OF PHARMA PSU OF INDIA FOR THE YEAR 2017-2019

Tender Reference	BPPI/DRUG-054/2017 Dt. 31/08/2017
Date of availability of tender documents on website	31/08/2017 (Thursday)
Time and date and place pre-bid meeting	11:00 AM on 07/09/2017(Thursday)
	Bureau of Pharma PSUs of India, IDPL corporate office, IDPL Complex, Old Delhi-Gurgaon Road, Dundahera, Gurgaon- 122016 (Haryana)
Last date and time for submission of Online Bid i.e. Bid Submission End Date and time	21/09/2017 upto 11:00 A.M.
Last Date for submission of EMD in physical Form in office of Bureau of Pharma PSUsof India, IDPL corporate office Complex, Old Delhi-Gurgaon Road, Dundahera, Gurgaon- 122016 (Haryana)	22/09/2017
Time and date of opening of Technical Bid	11:30 AM on 22/09/2017 (Friday)

Place of opening of tender	Bureau of Pharma PSUs of India,			
	IDPL corporate office Complex, Old Delhi-Gurgaon Road, Dundahera, Gurgaon- 122016 (Haryana)			
Address for Communication	Bureau of Pharma Public Sector Undertakings of India,			
	IDPL corporate office Complex, Old Delhi-Gurgaon Road, Dundahera, Gurgaon- 122016 (Haryana)			
Cost of the Tender Document	Free of cost			
Contact Person for clarification if any	1. Sh. Mahadev Agarwal, Manager (Procurement) Phone:- 0124-4040756 Mob:- 9811780789 Email: mahadevpharm.bppi@gmail.com 2.Mr. Sandeep Sapan Patra, Sr. Executive (Procurement) Phone:- 0124-4556767 Mob:- 9090512532 Email:- proc5.bppi@gmail.com 3. Mr. Rupak Kumar, Executive (Procurement) Phone:- 0124-4556764/767 Mob:- 7291087675 Email:- proc3.bppi@gmail.com			

The tender document can be downloaded free of cost from the CPPP e-Procurement Portal https://eprocure.gov.in and from the website of BPPI: janaushadhi.gov.in.

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BUREAU OF PHARMA PUBLIC SECTOR UNDERTAKINGS OF INDIA

ONLINE TENDER FOR THE SUPPLY OF DRUGS TO <u>BUREAU OF PHARMA</u> <u>PUBLIC SECTOR UNDERTAKINGS OF INDIA</u>

FOR THE YEAR 2017-19

PRADHAN MANTRI BHARTRIYA JANAUSHADHI PARIYOJANA(PMBJP) is the initiative of Department of Pharmaceuticals, Ministry of Chemical and Fertilizer, Government of India launching with the noble objective of making quality generic medicines available at affordable prices for all, particularly the poor and disadvantaged, through specialized outlets called PRADHAN MANTRI BHARTRIYA JANAUSHADHI KENDRA (PMBJK). BPPI was established in December 2008 under the Department of Pharmaceuticals, Government of India, with the support of all the CPSUs, and identified as the executing agency for PMBJP.

The Bureau has been registered as an independent society under the Societies Registration Act, 1860, in April, 2010. BPPI follows the provisions of GFR 2017 as amended from time to time, the CVC guidelines, and instructions from the Department of Pharmaceuticals.

It aims to open more than 3000 stores during current financial year. It is proposed to channelize efforts to popularize PMBJP and ensure availability of the complete basket of medicines at affordable prices.

Tender Inviting Authority – C.E.O, Bureau of Pharma Public Sector Undertakings of India, IDPL Corporate Office, IDPL Complex, Old-Delhi-Gurgaon Road, Dundahera, Gurgaon - 122016 (Haryana) (hereinafter referred as **Tender Inviting Authority** unless the context otherwise requires).

Tender Accepting Authority – CEO, Bureau of Pharma Public Sector Undertakings of India,(hereinafter referred as **BPPI** unless the context otherwise requires).

Tender Inviting Authority invites Tender for the supply of Drugs to BUREAU OF PHARMA PUBLIC SECTOR UNDERTAKINGS OF INDIA, for the year 2017-2019.

1. LAST DATE AND TIME FOR SUBMISSION OF ONLINE TENDERS.

- (a) Online Bids [in two separate Cover {Technical bid ("Cover A") and price bid (Cover "B")}] will be submitted till **11.00 A.M. up to 21/09/2017(Thursday) on** CPP portal i.e. eprocure.gov.in.
- (b) The price bid shall be valid for a period of 120 days from the date of opening of Technical Bid. Prior to the expiry of the bid validity, the Tender Inviting Authority may request the Tenderers to extend the bid validity for further period as deemed fit on their original quoted prices and all terms &conditions. However, BPPI reserves the right to place purchase orders at the quoted rate till such period.

2. ELIGIBILITY CRITERIA

- (a) (i) Tenderer shall be a manufacturer having valid drug manufacturing unit duly licensed by licensing authorities. Loan licensee is also eligible.
- (ii) Tenderer shall be direct importer holding valid import license. The Importer should have valid sale license.
- (iii) Tenderer shall be a marketer of manufacturer who have exclusive rights to market the products and manufacturer do not market the products duly supported by valid agreement with the manufacturer and in that case BPPI shall sign tri party agreement for supply of drugs if they are eligible for award of contract. **Distributors/Suppliers/Agents are not eligible to participate in the Tenders.** The Marketer should have valid sale license.
- (b) (i) Manufacturer should have valid WHO-GMP (World Health Organisation-Good Manufacturing Practices) certificate issued by licensing authority.
 - (ii) Average Annual turnover of manufacturer in the last three years i.e.2013-14, 2014-15 and 2015-16 shall not be less than **Rs.10 Crores.** In case of loan licensees and Marketer, average annual turnover of manufacturer in the last three years i.e. 2013-14, 2014-15 and 2015-16 shall not be less than **Rs.10 Crores.**
- (c) Market Standing Certificate (MSC) issued by the state licensing authority under generic or brand name as a Manufacturer for each product quoted in the tender for a minimum 2 years.
- (d) A certificate from their C.A. (Chartered Accountant) or ICWA that they have manufactured & marketed at least 2 commercial batch in last three years.
- (e) Non-conviction Certificate not older than 6 month issued by the licensing authority of the State certifying that the firm/company has not been convicted.
- (f) A certificate from the C.A. (Chartered Accountant) or ICWA that the bidder has Production & financial capacity to manufacture and deliver the drugs quoted by the firm in the tender as per quantity mentioned in tender during contract period.
- (g) Tenderer should not be submitted for the product(s) for which the firm / company has been blacklisted/debarred/de-registered/banned by any State Government / Central Government / its Drug procurement agencies due to quality failure of the drugs *at the time of submission of online bid.*

- (h) The Tenderer should have not been blacklisted/debarred/de-registered/banned due to quality failure for the quoted product /firm by any State Government / Central Government / its Drug procurement agencies **at the time of submission of bid**. Further, quoted drugs have not been failed in house testing or testing by any State Government/Central Government / its Drug procurement agencies/BPPI during last two years.
- (i) During the validity of the tender if the firm / Company is blacklisted/debarred/deregistered/banned by any State Government / Central Government / its Drug procurement agencies / convicted by any Court of law in India, it shall be intimated to BPPI along with relevant authentic document by the tenderer firm/ company within one month.
- (j) The tenderer should confirm that they have read tender document including Amendment(s) to Tender document (if any) along with terms and condition and these terms and condition of tender document including Amendment(s) to Tender document (if any) are acceptable unconditionally to them.
- (k) Tenderer are required to incorporate bar codes as per GS1 standards at various packaging levels (primary, secondary and tertiary. (Annexure I)

3. GENERAL CONDITIONS.

- (i) The tender document shall be download from the websites janaushadhi.gov.in; and CPP portal i.e.eprocure.gov.in. Tender Document is free of cost. No tender cost is to be deposited.
- (ii) **EMD** (Earnest Money Deposit): EMD of Rs.1,00,,000/- (Rupees One Lakh only as specified in Clause 7 of the Tender document in the form of Bank Guarantee or Bankers Cheque or Demand Draft from nationalised/Scheduled Bank favouring "Bureau of Pharma Public Sector Undertakings of India ", payable at Gurgaon/Delhi which is to be delivered in original to BPPI, Gurgaon on or before the date stipulated against 'Bid opening Date'. Name & full address of the bidder may be written at the back of the Demand Draft/Pay Order. Signed and scanned soft copy of the EMD instrument must be uploaded (ANNEXURE IV) to the e-Procurement portal. EMD in any other form like *cheque/cash/postal order* etc. will not be accepted. The Bid (in case not exempted for EMD as mentioned in tender document) without EMD shall be summarily rejected.
- (iii) Tenders will be opened online. However, authorized representatives of bidder who like to attend online bid opening on the specified date and time should bring letter of authority authorising to attend online bid opening on the printed letter head of the company.
- (iv) (a) At any time prior to the last date of submission of online bid, Tender Inviting Authority may, for any reason, whether on own initiative or in response to a clarification requested by a prospective Tenderer, may modify the condition in Tender documents by an amendment uploading on website on **janaushadhi.gov.in**; and CPP portal i.e. **eprocure.gov.in** will be binding on them. In order to provide reasonable time to take the amendment into account in preparing their bid, Tender Inviting Authority may at discretion, extend the date and time for submission of online bid.

- (b) Any person who has downloaded the tender document should watch for amendment, if any, on the website **janaushadhi.gov.in**; and CPP portal i.e.**eprocure.gov.in** for which BPPI will not issue any separate communication to them.
- (v) Interested eligible Tenderers may obtain further information in this regard from the office of the Tender Inviting Authority on all working days between 10:00 AM and 5:00 PM.
- (vi) During tender or agreement period, if L1 bidder is debarred/deregistered/blacklisted/banned by any Central Government or state Government or its procurement agencies due to quality failure, BPPI may purchase the drugs from L2 bidder or may go for fresh tender as per discretion of BPPI.
- (vii) The BPPI reserves the right to purchase any drugs full or part quantity from PSU as per discretion of BPPI. In case of emergencies, BPPI may go to PSU and price will be as per negotiation and at the discretion of BPPI.

3.1 SPECIAL CONDITIONS.

- (i)Bids shall be submitted online only at CPPP website :https://eprocure.gov.in. Manual bids shall not be accepted except for the original documents/instruments as mentioned in tender document.
- (ii) Bidders are advised to follow the 'Special Instructions to the Contractors/Bidders for the e-submission of the bids online' available through the link 'Help for Contractors' at the e-Procurement Portal https://eprocure.gov.in.
- (iii) Bidder shall not modify the downloaded tender form including downloaded price Bid template in any manner. In case any tender form/Price bid template is found to be tampered with/modified in any manner, such bid will be summarily rejected, Bid Security would be forfeited and bidder is liable to be banned from doing business with BPPI.
- (iv)Bidders are advised to check the *website of BPPI: janaushadhi.gov.in* and CPPP website https://eprocure.gov.in at least 3 days prior to closing date of submission of tender for any corrigendum, addendum, or amendment to the tender document.

4. TECHNICAL BID - COVER "A"

- **4.1.** The Tenderer should upload the following documents in while submitting technical bid hereafter called <u>"Cover A"</u>. (Scanned copies of each page of all documents should be uploaded while submitting Technical bid).
- (a) (i) The marketer of manufacturer who have exclusive rights to market the products and manufacturer do not market the products should upload valid agreement with the manufacturer with technical bid (ANNEXURE II) and the original agreement should be submitted on or before the schedule date of technical bid opening. In case, bidder is a marketer of a manufacturer, scanned copy of valid sale license is required to be uploaded. The Importer is also

required to upload copy of valid sale license. In case bidder is Importer, it is not mandatory to submit ANNEXURE II but it is advisable to submit the same from their Manufacturer.

- (ii) The tenderers are required to upload scanned undertaking on stamp paper duly notarized by authorised signatory confirming they are holding the valid drug license, valid WHO- GMP certificate, 2 years market standing certificate for quoted products issued by licensing authority, a certificate for manufactured & marketed of two batches for quoted drugs within 3 years issued by CA or ICWA, valid Non conviction certificate not older than 6 months issued by licensing authority, valid import license, undertaking as per para 2(h) & (j), undertaking to supply the drug with bar code as per ANNEXURE I and as per Annexure XIV & XIV A, undertaking for Clause 7.2 and also enclosed all undertaking/declaration as per Annexure mentioned in the tender document. In case the bidder is Importer, they may strike the clause or part of clause not applicable in their case. On the basis of such undertaking, the price bid shall be opened within a week after opening of technical bid. However, the bidder is required to upload/submit all the documents along with the technical bid and in case any document is not complying as per undertaking, their contract/agreement shall be cancelled with forfeiture of EMD/Security Deposit/Bank guarantee. (ANNEXURE - III). The original ANNEXURE III should be submitted to BPPI, Gurgaon on or before the schedule date of technical bid opening.
- (b) Earnest Money Deposit as indicated in Clause 3(ii) and Clause 7. of the tender document shall be in the form of **Bank Guarantee or Bankers Cheque or Demand Draft** favouring "Bureau of Pharma Public Sector Undertakings of India " payable at Gurgaon/Delhi. Tender cost and EMD in any other form like *cheque/cash/postal order* etc. will not be accepted. Scanned soft copy of the EMD instrument must be uploaded (ANNEXURE IV) to the e-Procurement portal. and original EMD instrument should be submitted to BPPI, Gurgaon on or before the schedule date of technical bid opening.
- (c) (i) Documentary evidence for the constitution of the Company/Firm such as Memorandum and Articles of Association, Partnership deed, Permanent Registration Number etc. with details of the Name, Address, Telephone Number, Fax Number, e-mail address of the firm and of the Managing Director / Partners / Proprietor. The list of present Directors in the Board of the Company duly certified by a Company Secretary of the Company/Practicing Company Secretary / Chartered Accountant to be uploaded.
- (ii) In case the bidder is MSME, documentary evidence for valid MSME certificate for quoted drugs is required to upload with technical bid failing which the bidder shall be treated as non MSME. Further, MSME unit owned by SC/ST entrepreneurs if applicable, is also required to upload documentary evidence with technical bid, failing which such MSME bidder shall not be treated from SC/ST entrepreneurs.
- (d) The instruments such as power of attorney, resolution of board etc., authorizing an officer of the Tenderer as the Authorized signatory of the Company/Firm should be uploaded.
- (e) Authorization letter nominating an officer of the Tenderer on the printed letter head of the company to transact the business with the BPPI to be uploaded.

- (f) (i) A certificate from the C.A.(Chartered Accountant) or ICWA that the bidder has Production & financial capacity to manufacture (as per format ANNEXURE V) and deliver the drugs quoted by the firm in the tender as per quantity mentioned in tender during contract period. The certificate should be uploaded along with the technical bid. In case the bidder is Importer, a certificate from the C.A.(Chartered Accountant) or ICWA that the bidder has Financial capacity to manufacture (as per format ANNEXURE VA) and deliver the drugs quoted by the firm in the tender as per quantity mentioned in tender during contract period. The certificate should be uploaded along with the technical bid.
- (ii) Average Annual Turnover certificate from Chartered Accountant of manufacturer (in cluding manufacturer of loan licensees and Marketer) in the last three years i.e.2013-14, 2014-15 and 2015-16 certifying not be less than **Rs. 10 Crores is required to upload as per format(ANNEXURE-VI).** The certificate from C.A. (Chartered Accountant) is required to upload as per format(ANNEXURE-VI).
- (g) The Tenderer should upload Scanned copy of valid drug Manufacturing Licence for the product, duly approved by the Licensing Authority for each and every product quoted as per specification in the tender. The licence must have been duly renewed up to date and the items quoted shall be clearly highlighted in the licence. Original documents should be produced for verification when demanded. However, if renewal application for manufacturing licence has been filed, Scanned copy of same duly receipted by drug authorities must be uploaded along with the validity certificate from state licencing authority (SLA).
- (h) Scanned copy of import license (in Form 10 with Form 41), as per Rule 122A of the Drugs and Cosmetics Act 1940, if the product is imported should be uploaded. The licence must have been renewed up to date. A copy of a valid licence for the sale of Drugs imported by the firms issued by the State Licensing Authority shall be uploaded. Original documents should be produced for verification when demanded.
- (i) MARKET STANDING CERTIFICATE (MSC) ISSUED BY THE STATE LICENSING AUTHORITY UNDER generic or brand name as a Manufacturer for each product quoted in the tender for a minimum 2 years (Certificate should be uploaded with list of items). In case of direct importer, evidence for importing the said items such as bill of landing, bill of entry and certificate of analysis are to be uploaded. MSC issued under brand name or under generic name (by the state licensing authority) will also be accepted but supplies will be accepted as per packing and label by foreign manufacturer in their brand subject to affixing sticker for Logo as approved by BPPI & BPPI MRP. However, for those newly launched drugs whose first product permission to manufacture and sale has been issued within 2 years by the respective country's / state drug authority, MARKET STANDING CERTIFICATE (MSC) issued by the respective country's /STATE LICENSING AUTHORITY under generic or brand name as a Manufacturer for less than 2 years shall be acceptable to BPPI. In case the bidder is Importer, MARKET STANDING CERTIFICATE (MSC) ISSUED BY THE STATE LICENSING AUTHORITY shall not be applicable.
- (j)The bidder should upload a certificate from their C.A.(Chartered Accountant) or ICWA that they have manufactured & marketed at least 2 commercial batch in last three years. The details of commercial batch no., month of manufacture, batch size in last three years period duly certified by their C.A. or ICWA should be uploaded along with technical bid. In case the bidder is Importer also, a certificate from their C.A.(Chartered Accountant) or ICWA that they have manufactured & marketed at least 2 commercial batch in last three years. The details of

commercial batch no., month of manufacture, batch size in last three years period duly certified by their C.A. or ICWA should be uploaded along with technical bid.

- (k) The copies of relevant pages approved by drug authorities of concerned country for any quoted Drug/product offering CoPP certificate and quoted drugs/product approved by from US FDA, TG Australia, Health Canada, EU approval, MCC South Africa approval, Brazil Anvisa) should be uploaded with technical bid.
- (I) Scanned copy Non-conviction Certificate issued by the licensing authority of the State certifying that the firm/company has not been convicted should be uploaded. **The certificate should not be more than 6 months old at the time of submission of technical bid**.
- (m) Scanned copy of Valid WHO-GMP(World Health Organisation-Good Manufacturing Practices) Certificate (for manufacturer only) issued by the Licensing Authority should be uploaded. In case of Imported drugs , labels and product literature of all quoted product(s) must be uploaded COPP certificate as per WHO format of their Principal Manufacturing company/firm.
- (n) a. Scanned copy of GST registration certificate/Latest Sales Tax Clearance certificate/returns are to be uploaded.
- b. Scanned copy Latest Income tax assessment orders/returns filed are to be uploaded.

In case, bidder is a marketer of a manufacturer, the GST registration certificate/latest Sales Tax Clearance certificate/returns and the latest Income tax assessment orders/returns of bidder as well as of their manufacturer/manufacturers are to be uploaded.

- (o) Documents, if any, to show that the manufacturing unit/importer has been recognized by any other Indian / International Standard Organizations etc. as applicable.
- (p)The loan license bidder are required to upload scanned copies of all the documents as per tender requirements including manufacturing unit.
- (q)List of items quoted (The name & Drug code of the Items quoted as shown in the **ANNEXURE-VII** should be uploaded and **the rate of those items should not_be indicated in this list).**
- (r) A Checklist (**ANNEXURE-VIII**) shall be uploaded with technical bid. If a company/firm has two or more separate manufacturing units at different sites / States, which are not separate entities then the company will be allowed to submit only one tender for all units but necessary document regarding separate manufacturing units will uploaded as a separate set with the same tender. However, one bidder will be allowed to submit only one offer for one product.
- (s) All the documents uploaded should also be signed by the authorized official of the Tenderer.
- **4.2.** The all documents indicated above should be uploaded and shall be opened at the time of Technical bid opening.

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5. PRICE BID - COVER "B"

- **5.1.** Cover "B" contains the Price Bid of the Tenderer.
- (i) The Tenderer shall fill in the rate per unit size, % age rate of GST and total rate inclusive of GST in respective column of BOQ for the items quoted. In case, any bidder offers CoPP or offers product approved by US FDA, TG Australia, Health Canada, EU approval, MCC South Africa approval, Brazil ,Anvisa, copies of relevant pages of valid document approved by drug authorities of concerned country for imported drug should be uploaded on line with technical bid.

(ii) **Determination of L1 bidder:**

- (a) In determining the lowest evaluated price, the rate quoted per unit size inclusive if GST as indicated in column No. 8 of the **BOQ** shall be taken into consideration.
- (b) The Price preference of up to 5% over L1 bidder (if L1 bidder is not offering certificate of pharmaceutical product i.e. **CoPP** issued in the format recommended by the World Health Organization) shall be given to the bidder having CoPP for the particular drugs and shall be awarded contract. Scanned copy of Valid CoPP issued by the Licensing Authority must be uploaded.
- (c) The Price preference up to 10% over L1 bidder (if L1 is not offering product approved by from US FDA, TG Australia, Health Canada, EU approval, MCC South Africa approval, Brazil Anvisa) shall be given to the bidder having product approval from US FDA. TG Australia, Health Canada, EU approval, MCC South Africa approval, Brazil Anvisa.
- (d)(i) If the participating Micro and Small Enterprises (MSE) meets all the other eligibility criteria and their quoting price is within price band of L1+15 (fifteen) per cent shall also be allowed to supply a portion of requirement by bringing down their price to L1 price in a situation where L1 price is from someone other than a MSE and such MSE shall be allowed to supply up to 20 (twenty) per cent of total tendered value. The 20 (twenty) per cent quantity is to be distributed proportionately among these bidders, in case there are more than one MSMEs within such price band.
- (ii) Within this 20% (Twenty Percent) quantity, a purchase preference of four per cent (that is, 20 (twenty) per cent out of 20 (twenty) per cent) will be reserved for MSEs owned by Scheduled Caste (SC)/Scheduled Tribe (ST) entrepreneurs (if they participate in the tender process and match the L1 price). Provided that, in event of failure of such SC/ST MSE to participate in tender process or meet tender requirements and L1 price, four per cent sub-target shall be met from other MSE. MSEs would be treated as owned by SC/ST entrepreneurs: a) In case of proprietary MSE, proprietor(s) shall be SC/ST b) In case of partnership MSE, the SC/ST partners shall be holding at least 51% (fifty-one percent) shares in the unit c) In case of Private Limited Companies, at least 51% (fifty-one percent) share shall be held by SC/ST promoters.

Note 1:- (a) Price preference as in Clause 5.1 (ii) (c) will be get preference over the clause 5.1 (ii) (b).

Note 2:- Later on, if product does not comply CoPP and product having approval of the any agency like US FDA, TG Australia, Health Canada, EU, MCC South Africa approval,

Brazil Anvisa approved product as declared in tender, the extra price paid to the supplier shall be recovered in addition to other penal action.

- (iii) The rate quoted inclusive of GST in column 8 of **BOQ** should be for a unit and for the given specification. **The rates quoted should be in rupees and paisa up to 2 digits.** The Tenderer is not permitted to change/alter specification or unit size given in the **ANNEXURE-IX**
- (iv) GST (Goods and Services Tax)-<u>The tenderers must indicate the rate of GST applicable and payable by them. In case no information is given, it shall be presumed that rate are inclusive of GSAT and no GST shall be charged by them under any circumstances.</u>
- (v) The bidder is required to indicate GST in % only against the heading of column BOQ and not to indicate amount of GST in Rs. at particular cell of excel sheet of BOQ.

6. OPENING OF COVER "A" AND COVER "B" OF TENDER

- **6.1** Only authorized official as indicated in Clause 4.1. (e) are entitled to be present at the time of opening of Technical Bid Cover "A" of the tender submitted by them.
- **6.2** Tenderers, who are found eligible on satisfying the criteria for technical evaluation/based on undertakings & Declaration, will only be informed the time and date of opening of Price Bid Cover "B" of the tender.
- 6.3 In case, the date for opening of technical bid is declared holiday, the technical bid shall be opened on next working day at 11.30 A.M.

7. EARNEST MONEY DEPOSIT

- 7.1. The Earnest Money Deposit referred to under Clause 3(ii) & 4.1(a), shall be Rs. 1 lakh. The Earnest Money Deposit shall be paid in the form of Bank Guarantee or Bankers Cheque or Demand Draft in favour of BUREAU OF PHARMA PUBLIC SECTOR UNDERTAKINGS OF INDIA, payable at Gurgaon/Delhi. In case EMD in form of Bank Guarantee, Irrevocable Bank Guarantee in favour of Bureau of Pharma Public Sector Undertakings of India from any Nationalised/scheduled Bank should be valid for a period beyond 270 days/9 months from the date of tender opening. The format of Bank Guarantee is at ANNEXURE-X. BPPI will not pay interest on any deposit held in the form of Bankers Cheque or Demand Draft.
- **7.2**. (i) The tender submitted without sufficient EMD will be summarily rejected.
- (ii) The Earnest Money Deposit will be refunded to the successful bidders within 30 days from the date of signing the contract agreement and on the deposit of Security Deposit.

- (iii) The Earnest Money Deposit (EMD) of the unsuccessful bidders will be returned after finalization of tender/signing of agreement with eligible bidder.
- (iv) The Earnest Money Deposit (EMD) will be forfeited, if the tenderer withdraws his bid any time after opening of price bid / non execution of agreement /undertaking within the period prescribed.
- (v) The Earnest Money Deposit (EMD) will be forfeited, in case of the lowest bidder, fails to execute the contract agreement and / or deposit the security Deposit within the stipulated time. The EMD shall be forfeited if the undertaking as Annexure II is not found correct.
- (vi) Tenderer may be exempted from the payment of EMD, if valid **registration** certificate from NSIC/MSME is uploaded **for the product for which bidder has submitted quotation.**
- (vii) PSUs are exempted from the payment of EMD.

8. OTHER CONDITIONS

- **8.1**.(i) The details of the required drugs, medicines, etc., are shown in **ANNEXURE -IX**. *The tender quantity mentioned herein is not a fixed procurement quantity and it is only a tentative requirement and may be increased or decreased* by the BPPI, at its discretion, depending on it is actual need. Though the tentative quantity is indicated in the agreement, the BPPI, will confirm the actual requirement then / there through purchase order/orders. The tenderers shall supply the drugs only on the basis of the purchase order issued time to time within validity of contract period by the BPPI. Any supply without a valid purchase order will not be acceptable by BPPI and the BPPI shall not be responsible for any loss on this account.
- (ii) The Tenderer shall fill in manufacturing capacity per year in units, Shelf life in months and manufacturing batch size in units for each quoted drugs in required column of **ANNEXURE XI and upload along with technical bid.** In case the bidder is Importer, the importer is required to sign and upload ANNEXURE XI on behalf of the exporter which would be supported by documentary evidence provided by the manufacture
- (iii) However, once the purchase order/orders is/are issued by the BPPI, the tenderer shall not renege from the commitment of supplying the quantity mentioned in the agreement / undertaking.
- (iv) The rates quoted shall not be varied with the ordered quantity during the full contract period.
- **8.2.** Tender has been called for in the <u>Generic name of drugs</u>. The Tenderers should quote the rates for the generic products only. The composition, strength and packing of each product should be as per specifications given in **ANNEXURE-IX**. Any variation, if found, will result in rejection of the tender. However, the imported/combination drugs are allowed to quote in trade / brand name.
- **8.3.** Rates (inclusive of Customs duty, packing & forwarding charges, transportation, insurance and any incidental charges, but exclusive **GST** should be quoted for each of the required drugs, medicines etc., separately on door delivery basis according to the unit ordered.

Tender for the supply of drugs, medicines, etc. with cross conditions like "AT CURRENT MARKET RATES" shall not be accepted. Handling, clearing, transport charges etc., will not be paid separately. The delivery should be made as stipulated in the purchase order placed with Tenderers.

- **8.4.** Each bid must contain not only the unit rate but also the total value of each item quoted for supply in the respective columns. The aggregate value of all the items quoted in the tender shall also be furnished.
- **8.5.** (i) The price quoted by the tenderers shall not, in any case exceed the DPCO controlled price, if any, fixed by the Central/State Government, the Maximum Retail Price (MRP) and the selling price of the tenderer. Tender Inviting Authority at its discretion, may exercise, the right to revise the price at any stage so as to conform to the controlled price or MRP or the selling price of the tenderer as the case may be. This discretion will be exercised without prejudice to any other action that may be taken against the Tenderer. In case delivery of drugs is not made within delivery period mentioned in Purchase order, the supplier must confirm from BPPI whether BPPI MRP is to be reduced due to changes in DPCO ceiling rate after issue of purchase order.

(ii) FALL CLAUSE:

If at any time during the execution of the contract, the controlled price becomes lower or the supplier reduces the sale price or sells or offers to sell such stores, as are covered under the contract, to any person / organization including the purchaser or any department of Central government/state Govt. or its procurement agencies at a price lower than the price chargeable under the contract, he shall forthwith notify such reduction or sale or offer of sale to the purchaser and the price payable under the contract for the stores supplied after the date of coming into force of such reduction or sale or offer of sale shall stand correspondingly reduced.

- **8.6.** The rates quoted and accepted will be binding on the Tenderer for the full contract period of two years and any increase in the price will not be entertained till the completion of this contract period. Accordingly, this clause will be applicable for all orders placed during the contract period. However, agreement validity period may be extended for period up to further one year at same rate, terms & conditions with the consent of the supplier.
- **8.7.** No Tenderer shall be allowed at any time and on any ground, whatsoever it may be, to claim revision or modification in the rates quoted by them. Representation to make correction in the tender documents on the ground of Clerical error, typographical error, etc., committed by the Tenderers in the Bids shall not be entertained after submission of the tenders. Cross Conditions such as "SUBJECT TO AVAILABILITY", "SUPPLIES WILL BE MADE AS AND WHEN SUPPLIES ARE RECEIVED" etc., will not be entertained under any circumstances and the tenders of those who have mentioned such conditions shall be treated as incomplete and accordingly the Tender will be summarily rejected.
- **8.8.** Supplies should be made directly by the tenderer and not through any other Agency / Dealer / Distributors.

- **8.9.** The Tenderer shall allow inspection of the factory at any time after the opening of technical bid and during the entire contract period by a team of Experts/Officials nominated by the Tender Inviting Authority for the purpose. The Tenderer shall extend necessary cooperation to such team in inspection of the manufacturing process, quality control measures adopted etc., in the manufacture of the items quoted. If Company/Firm does not allow for any such inspection, their tenders will be rejected. If any such situation arises after placement of contract, the same shall be cancelled at the firm's risk cost.
- **8.10** "MRP inclusive of all taxes" is to be printed on each unit/label. MRP will be intimated to successful bidders at the time of placing purchase orders.

9. ACCEPTANCE OF TENDER

- **9.1.** (i.) Evaluation of the tender and determination of the L1 rate (Lowest rate) will be done based on rate per unit size inclusive of GST as mentioned in column 8 of **BOQ** considering price preference for <u>CoPP</u> and for product having approval of the any agency like US FDA, TG Australia, Health Canada, EU, MCC South Africa approval, Brazil Anvisa approved product. However, to have additional source of supply, the L1 bidder shall be awarded contract/agreement for 50% of tender quantity indicated in the tender document. Balance 50% of the tender quantity indicated in the tender document shall be awarded to **L2 bidder if they agree to supply the drugs at L1 rates.**
- (ii) In case, L2 bidder does not agree to match L1 rate, 100% tender quantity shall be awarded to L1 bidder. The purchase order shall be issued to L1 bidders and in case they fail to supply in stipulated time or due to quality failure, the purchase order shall be issued to L2 bidder. During the agreement period of 2 years, in case L1 bidder completes the supply of drugs for contracted quantity, next supply shall be taken from L2 bidder accordingly for contracted quantity.
- (iii). The issue of purchase orders in same manner as mentioned above. Negotiation if required will be done at our premises and the same will be done strictly as per Central Vigilance Commission guidelines.
- Note 1.:- No quantity distribution shall be applicable if L1 rates quoted by more than one bidder keeping in view of sharing of quantity as per clause no. 11(c) provided that no L1 bidder has <u>CoPP</u> certificate or drug/product approval of the any agency like US FDA, TG Australia, Health Canada, EU, MCC South Africa approval, Brazil Anvisa.
- Note 2. In case, MSME bidder is not eligible as per clause 5.1(ii)(d) and single bid is available after determination of L1 bidder considering price preference 5% for CoPP and 10% for product having approval of the any agency like US FDA, TG Australia, Health Canada, EU, MCC South Africa approval, Brazil Anvisa approved product, Such bidder shall be awarded 70% of quantity indicated in tender document and balance 30% quantity shall be awarded to lowest bidder at their quoted rates if applicable or L2 bidder at lower rate by 5% over L1 rate in case of CoPP bidder and by 10% over L1 rate in case product having approval of the any agency like US FDA, TG Australia, Health Canada, EU, MCC South Africa approval, Brazil Anvisa approved product,

- **9.2.** BPPI reserves the right to accept or reject the tender for the supply of all or any one or more items of the drugs tendered for in a tender without assigning any reason.
- **9.3.** BPPI or its authorized representative(s) has/have the right to inspect the manufacturing premises of Tenderers, before accepting the rate quoted by them or before releasing any purchase order(s) or at any point of time during the continuance of tender and also has the right to reject the tender or terminate/cancel the purchase orders issued and/or not to place further order, based on adverse reports brought out during such inspections.
- **9.4.** The acceptance of the tenders will be communicated to the Tenderers in writing.

10. SECURITY DEPOSIT AND AGREEMENT

10.1Security Deposit:

On being informed about the acceptance of the tender and at the time of signing the Agreement, the Tenderer shall pay the Security Deposit @Rs.50000/- per drug code accepted for award of contract/agreement subject to minimum Rs.100000/- and maximum Rs.20,00,000/- in the form of *Demand Draft or irrevocable Bank Guarantee* in favour of Bureau of Pharma Public Sector Undertakings of India from any scheduled Bank. In case the Security Deposit is paid in form of Bank Guarantee, the bank guarantee shall be valid for a period beyond **one year of the validity of the agreement**. The format of Bank Guarantee is at **ANNEXURE-XII**.

- **10.2.** The Tenderer shall execute an agreement on a non-judicial stamp paper of value of Rs.100/- (stamp duty to be paid by the Tenderer) within 15 days from the date of the intimation from BPPI informing that his tender has been accepted. The Specimen form of agreement is available in **ANNEXURE-XIII.**
- **10.3.** The Tenderer shall not, at any time, assign, sub-let or make over the contract or the benefit thereof or any part thereof to any person or persons what so ever.
- **10.4.** All notices or communications relating to and arising out of this agreement or any of the terms thereof shall be considered duly served on or given to the Tenderer if delivered to him or left at the premises, places of business or abode as provided by the tenderer.
- **10.5.** If the lowest selected Tenderer fails to execute the agreement and/or to deposit the required security deposit within the time specified or withdraws the tender, after the intimation of the acceptance of the tender or owing to any other reasons to undertake the contract, the contract will be cancelled and the Earnest Money Deposit deposited by the tenderer along with the tender shall stand forfeited by the BPPI and the firm will also be liable for all damages sustained by the BPPI apart from blacklisting and other penal actions. The security deposit shall be forfeited if the undertaking as Annexure II is not found correct.
- **10.6**. The security deposit of supplier will be returned by BPPI only after the supplier has given undertaking to replace such medicines and indemnify BPPI against any loses on account of quality parameters.

11. METHODOLOGY FOR PLACING ORDERS

For the above purpose the following procedures will be adopted

- (a) After the conclusion of Price Bid opening (Cover B), the rates offered by tenderers for each product are evaluated and lowest acceptable rate (L1 Rate) arrived at is declared and that tenderer is informed.
- (b) The Successful Tenderer is eligible for the placement of Purchase Orders only after depositing the required amount as Performance Security and on execution of the agreement.
- (c) If two or more than two Tenderer's are declared as lowest suppliers for the same item(s), such Tenderers shall execute necessary agreement as specified in the Tender Document on depositing the required amount as Performance Security and on execution of the agreement such Tenderers are eligible for the placement of Purchase Orders for such item(s) for which they are declared as lowest. Placement of order shall be shared equally amongst these bidders subject to their manufacturing capacity.
- (d) In the case of purchase of goods where the quantity offered at the lowest price is less than the total quantity required, the BPPI may, after placing orders with the lowest evaluated Tenderer for the entire quantity offered by such Tenderer subject to his ability to supply, require all the other eligible Tenderers who participated in the tender and offered a price higher than that offered by the lowest evaluated Tenderer, to submit sealed offers of the quantity they would be willing to supply at the price quoted by the lowest evaluated Tenderer, and thereafter place orders for the remaining required quantity with all those who match the lowest evaluated price such that those who bid lower prices in the original tender get a higher priority for supply.
- (e) If a supplier fails to execute supply order, the 5% value of supply order shall be recovered from pending bill or EMD/Bank Guarantee and their bad performance shall be kept in record of BPPI for future dealing as considered appropriate by BPPI.
- (f) Notwithstanding anything contained in para (e) above, the supplier, after committing the default in supply either partly or fully, can inform the BPPI about his willingness to execute the Purchase Order during the tender period. The BPPI at discretion may consider the willingness of the supplier on merit. However, such supplies will be subjected to the levy of Liquidated Damages, unexecuted fine and other penalties as stipulated in the tender document, agreement and purchase order.
- (g) The supplier shall start supply of the Drugs/Medicines required by BPPI at Central Ware House (CWH), Gurgaon or any other place decided by BPPI within the stipulated period.

- (h) The Drugs/Medicines supplied in excess of the ordered quantity shall not be accepted and the supplier shall take back the excess at their cost. BPPI will not be responsible for the loss to the supplier and will not entertain any demand/claim.
- (i) The supplier shall supply the Drugs/Medicines at the CWH, Gurgaon along with copy of Purchase order, copy of test reports and 3 original copies of Invoice. No payment will be processed without test reports.
- (j) The supplier shall take utmost care in supplying the quality Drugs/Medicines and ensure that the batch number mentioned in the packages of the Drugs/Medicines tally with the batch number mentioned in the Invoice produced to BPPI for payment. Also the supplier shall ensure the quantity relevant to the Batch Number of the Drugs/Medicines is mentioned in the invoice.
- (k) It is the duty of the supplier to supply Drugs/Medicines at the CWH Gurgaon or any other place decided by BPPI and supply shall conform to the conditions mentioned in the provisions of tender documents, viz., logo, nomenclature, specification etc.,
- (l) Subject to above, BPPI will process the invoices submitted by the supplier and the payments against supply will be made within 30 days from the date the Drugs/Medicines supplied has been declared of STANDARD QUALITY, by the Empanelled laboratory of BPPI subject to various terms and conditions of the tender.
- (m) Subject to the conditions mentioned in the Purchase Order, Tender Document, Agreement executed by the supplier and here under, the Supplier is entitled for the payment against supply. In case of any discrepancy in levy of LD, Penalty, Unexecuted Fine, Short Passing of Bills, such discrepancy shall be intimated within 30 days from the date of receipt of payment, failing which BPPI will not entertain any claim thereafter.
- (n) BPPI reserves the right to place upto 50% additional purchase order of the quantities as contracted within validity of contract.

12. SUPPLY CONDITIONS

- **12.1.** Purchase orders will be issued to the Tenderer(s) at the discretion of the BPPI as per actual requirements. All the supplies shall be received at the central warehouse at Gurgaon or any other place decided by BPPI.
- **12.2.** Within 3 days from the receipt of purchase orders the Tenderer should inform BPPI through fax and mail the confirmation for the receipt of the purchase order.

- **12.3.** The Tenderer should also fax and mail the details of supply dates as specified in Annexure, to BPPI within 7 days from the receipt of the purchase order. In case, the supply shall not be made by the date as conveyed by the supplier, supply order shall be cancelled at their risk and cost. If no response is received within 7 days from the supplier / tenderer about supply of drugs as per purchase order, it shall be presumed that the supplier/tenderer is not interested to supply the drugs ordered as per purchase order and BPPI shall purchase the drugs from alternative sources.
- **12.4**. (a) For the first purchase order, the supplier must supply the ordered quantity within 45 days from the date of Purchase Order.
- (b) For Subsequent purchase orders, the supplier shall complete the supply within 30 days from the date of purchase order at the destinations mentioned in the purchase order.
- (c) If the above day for 12.4 (a) &(b) above happened to be a holiday for BPPI, the supply should be completed by 5.00 PM on the next working day.
- (d) In case of Non- execution of the order, BPPI reserves the right to place purchase orders (partially/fully) on alternate source at the risk and cost of the default tenderer(s) without any notice/Information.
- (e) If the Tenderer fails to execute the supply within the stipulated time, the BPPI is at liberty to make alternative arrangement for purchase of the items for which the Purchase orders have been placed, from any other sources or in the open market or from any other Tenderer who might have quoted higher rates, at the risk and the cost of the defaulted supplier and in such cases the BPPI has every right to recover the cost and impose Liquidated Damages as mentioned in Clause 18.
- (f) The liquidated damages as specified in clause 18.1 and 18.2 of the tender conditions will be levied on the quantity supplied after the 45thday and 30th day for 12.4 (a) &(b) respectively. However, no supplies will be accepted after 75th days/ 60 days for 12.4 (a) &(b) respectively from the date of issue of purchase order and the purchase order shall be cancelled at the risk and cost of the supplier. However, the supplier must take prior approval from BPPI for supply of drugs beyond stipulated delivery period in Purchase order.
- 12.5. Supplier shall complete the earliest pending purchase order before commencing the supply of subsequent purchase orders. Further, supplies ordered against a purchase order are to made in minimum numbers of batches as far as possible and same batch should not be supplied in repeated consignment.
- 12.6. The supplied Drugs (covered in SCHEDULE "P" of Drugs and Cosmetics Act) should have the prescribed potency throughout the shelf life period as prescribed in the Drugs and Cosmetics Act 1940 and rules there under and in relevant Pharmacopoeias. However, in case of thermolabile drugs not covered in SCHEDULE "P" of Drugs and Cosmetics Act, the minimum shelf life should be 2 years from the date of manufacture.
- **12.7.** The Tenderer must submit an Analysis report for every batch of drug along with invoice. In case of failure on part of the supplier to furnish such report, the batch of drugs will be BPPI/DRUG-054/2017

returned to the suppliers and he is bound to replenish the same with Govt. approved lab test report. The Drugs supplied by the successful Tenderer shall be of the best quality and shall comply with the specifications, stipulations and conditions specified in the tender.

12.8. Tenderer should supply the product (a) within 2 months including month of manufacture of products having shelf life upto 2 years, (b) within 3 months including month of manufacture of products having shelf life more than 2 years & up to 3 years and (c) within 4 months including month of manufacture of products having shelf life more than 3 years. Products beyond the above mentioned period from the date of manufacture shall not be accepted. For example, product having manufacturing of November 2017 must be supplied before Dec 31, 2017 in case shelf life less than 2 months.

For imported products, 60% of shelf life should be available at time of supply.

- **12.9**. If at any time the Tenderer has, in the opinion of the BPPI delayed the supply of drugs due to one or more reasons related to Force Majeure events such as riots, mutinies, wars, fire, storm, tempest or other exceptional events at the manufacturing premises, the time for supplying the drugs may be extended by the BPPI at discretion for such period as may be considered reasonable. However, such extension shall be considered only if a specific written request is made by the Tenderer within 10 days from the date of occurrence of such event with necessary documentary evidence. The exceptional events does not include the Scarcity of raw material, Increase in the cost of raw material, Electricity failure, Labour disputes/Strikes, Insolvency, and Closure of the Factory/Manufacturing unit on any grounds etc.
- **12.10**. The supplier shall not be liable to pay LD and forfeiture of security deposit for the delay in executing the contract on account of the extension of supply period on the ground of force majeure events.

13. LOGOGRAMS

Logogram means, wherever the context occurs, the design as specified in ANNEXURE-XIV. The name of the drug shall be mentioned in English and Hindi.

- **13.1.** Tenders for the supply for Drugs etc., shall be considered only if the Tenderer gives an undertaking that the product(s) will be prepared as per the specifications such as name, strength, minimum size and packed with appropriate size of the strips/blisters/bottles/tubes etc as per the design enclosed as per **ANNEXURE –XIV &XIV-A**.
- **13.2.** All tablets and capsules have to be supplied in packing as specified in product list (**ANNEXURE IX**) and shall also conform to Schedule P1 of the Drugs & Cosmetics Act & Rules 1945, wherever it applies. Affixing of stickers and rubber stamps shall not be accepted and supplies will be returned back at supplier's cost.
- **13.3.** Vials, Ampoules (more or equal than 5 ml) and Bottles containing the items tendered for should also carry the printed Jan Aushadhi logogram of proportionate size.

13.4. Failure to supply Drugs etc., with the printed logogram of proportionate size will be treated as breach of the terms of agreement / violation of tender conditions. The purchase order shall be cancelled at the risk and cost of the supplier. However, if such failure continuous despite notice, will be viewed as a serious lapse and initiate blacklisting of the supplier.

Tenderers who are not willing to agree to conditions above will be summarily rejected.

13.5. For imported Drugs, the supplies will be accepted as per packing and label by foreign manufacturer in their brand subject to affixing sticker for Logo as approved by BPPI & BPPI MRP.

14. PACKING

- **14.1.** The drugs shall be supplied in the package specified in **ANNEXURE IX** and **ANNEXURE** -**XV** and the package shall carry the logograms of proportionate size specified in **ANNEXURE** -**XIV**, **XIV** -**A**. Non affixing of logograms will be treated as violation of tender conditions and fine will be deducted from the amount payable as per condition in Clause 18.5
- **14.2.** The minimum size of each tablet should be 6.4 mm in diameter and the minimum size of the blister packing/strip packing/Alu-alu packing should be 80mm x 35mm/50mm x 130 mm/45mm x 110mm respectively. The drugs in any dosage form to be supplied by the supplier should not be embossed indicating any code no./logo or name of the company. Failure to comply with this shall lead to non-acceptance of the goods besides imposition of penalties as per clause 18.5.
- **14.3.** The packing in each carton shall be strictly as per the specification mentioned in **Annexure-XV**. The outer carton/secondary packaging should be of pearl white duplex board (**off white/grey is not acceptable**) with a minimum of 350 GSM with **Gloss laminated** packing for the strips, blisters, ointments, creams etc. and for ampoules and vials should be with pearl white board of 350 GSM(**off white/grey is not acceptable**). **The material to be used for carton should be from virgin chemical pulp.** Failure to comply with this shall lead to non-acceptance of the goods besides imposition of penalties as per clause 18.5. Storage conditions must be indicated on outer label.
- **14.4.** The cap of bottle preparations should not carry the name of the supplier.
- **14.5.** The labels in the case of Injectable preparations should clearly indicate whether the preparations are meant for Intravenous (IV), Intra Muscular (IM), Intra Dermal (ID), Subcutaneous (SC) administration etc.
- **14.6.** It should be ensured that only first-hand virgin packaging material of uniform size, including bottle and vial, is used for packing.
- **14.7.** All primary packing containers should be strictly conforming to the specification included in the relevant pharmacopoeia.
- **14.8.** Packing should be able to prevent damage or deterioration during transit.

- **14.9.** In the event of items of drug supplied found to be **not as per specifications in respect of their packing and logogram**, the BPPI is at liberty to make alternative purchase of the items of drugs for which the Purchase orders have been placed from any other sources or in the open market or from any other Tenderer who might have quoted higher rates, at the risk and the cost of the supplier. In such cases the BPPI has every right to recover the cost and impose penalty as mentioned in Clause 18 & 19.
- **14.10.** Designs of packaging with the logograms shall be subject to approval by BPPI within 3 days of receipt of the same from the supplier, as per the specifications with soft copy of STP(standard testing procedure) for Non- Pharmacopoeil drugs as per clause 15.1.1. In case of failure of BPPI to do so, the supplier may go ahead with the design as per the specification in **ANNEXURE XIV and XV.**
- 14.11. The colour of the strength must be different from the colour of the generic name of the drug on primary and secondary packaging and the approval for the same should be taken from the quality/regulatory department while taking artwork approval. The printing ink used should be of good quality (clarity, brightness, contrast) which is easily readable.
- 14.12. WHO-GMP certified, Therapeutic code & NABL lab tested shall be indicated on the primary and secondary packaging and shall be incorporated as per the approval from the quality/regulatory department while taking artwork approval.

15. QUALITY TESTING

- **15.1.** Samples of supplies from each batch will be chosen at the point of despatch at supplier's site or receipt of supply or distribution/storage points for testing at discretion of BPPI. The samples will be sent to different laboratories including Government Drugs Testing Laboratory for testing as decided by the BPPI Handling and testing charges will be deducted by BPPI for the above purpose, as specified in Clause 17.
- **15.1.1** For Non- Pharmacopoeil drugs, supplier should send soft copy of STP(standard testing procedure) by mail to Quality and Regulatory officer of BPPI with art work approval for design of packaging with the logogram as per Clause 14.10.
- **15.2.** The Drugs shall have the active ingredients at the prescribed level as indicated in official compendiums throughout the shelf life period of the drug. The samples will be drawn periodically throughout the shelf life period and if found "Not of Standard Quality", the cost of entire batch paid will be recovered whether consumed fully/partially. Also action will be initiated for blacklisting as per clause No.19 irrespective of the period of supply. The supplies will be deemed to be completed only upon receipt of the quality certificates from the laboratories. Samples which do not meet quality requirement shall render the relevant batches liable to be rejected. If the sample is declared to be "Not of Standard Quality" or spurious or adulterated or misbranded, such batch/batches will be deemed to be rejected goods.
- **15.3.** In the event of the samples of Drugs supplied fails in quality tests or found to be not as per specifications, the BPPI is at liberty to make alternative purchase of the items of drugs for which the Purchase orders have been placed from any other sources or in 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 BPPI has every right to recover the cost and impose penalty as mentioned in Clause 19.

- **15.4.** The supplier shall furnish evidence of the basis for shelf life and other stability data concerning the commercial final package on request by the BPPI. In case of any complaint in the field, the B.M.R/B.P.R for the particular batch of the product(s) supplied shall be produced when demanded.
- **15.5.** The products should conform to the standards of IP/BP/USP/EP/JP as the case may be. In case the product is not included in the any of the said compendiums, the supplier, upon award of the contract, must provide the reference standards and testing protocols for quality control testing. For imported drugs, respective Country's Pharmacopoeial standards shall be acceptable (even if the product is official in IP).
- **15.6.** The case of admixture of drugs will be treated as a violation of tender conditions and fine will be levied as per clause 19. If such lapses happen more than twice in a tender period such cases will be treated as "Misbranded Drugs".

16. PAYMENT PROVISIONS

- **16.1.** No advance payments towards costs of drugs, medicines etc., will be made to the Tenderer.
- **16.2.** Payments towards the supply of drugs will be made within 60 days from the date of receipt of goods, strictly as per the tender terms and condition. The payment will be made either by means of a/c payee Cheque or through RTGS (Real Time Gross Settlement System)/Core Banking/NEFT. The Tenderer shall furnish the relevant details in original (ANNEXURE XVI) to make the payment through RTGS/Core Banking/NEFT.
- **16.3.** All bills/Invoices should be raised in triplicate and the bills should be drawn as per GST Rules in the name of Bureau of Pharma Public Sector Undertakings of India. IDPL Complex, Dundahera, Gurgaon 122016 or in the name of any other authority as may be designated.
- **16.4.** (i) Payments for supply will be considered only after supply of minimum 50% of Drugs ordered in the individual Purchase Order PROVIDED reports of Standard Quality on samples testing are received from Government Analyst or Approved Laboratories of BPPI.
- (ii) However, in case of cancellation of a particular purchase order due to failure in delivery, payment for part supplies less than 50% of the purchase order quantity on the date of cancellation of the purchase order may be considered for release of payment subject to the following:
- (a) If the Tenderer have supplied at least 50% of the quantity ordered in the subsequent purchase order within delivery period stipulated in purchase order from the issue of such purchase order.
- (b) If further purchase order is not placed with the supplier due to any reason, not attributable to the supplier, the amount eligible will be paid within 60 days from the date of last supply.

- (c) The payment for part supply as mentioned above will subject to the deduction of liquidated damages, penalty towards unexecuted quantity, risk and cost etc., as per the tender conditions.
- **16.5.** If at any time during the period of contract, the price of tendered items is reduced or brought down by any law or Act of the Central or State Government or by the Tenderer himself, the Tenderer shall be bound to inform the BPPI immediately about such reduction in the contracted prices. Tender Inviting Authority is empowered to unilaterally effect such reduction as is necessary in rates in case the Tenderer fails to notify or fails to agree for such reduction of rates.
- **16.6.** In case of any increase of decrease in the taxes/GST after the date of submission of tenders and during the tender period, such variation in the taxes/GST will be to the account of the BPPI. For claiming the additional cost on account of the increase in taxes/GST, the Tenderer should produce the proof of having paid additional amount on this account on the goods supplied to BPPI from the concerned authorities and also must claim the same in the invoice separately. However, the basic price structure and the price of the Drugs approved under the tender shall not be altered. Similarly, if there is any reduction in the taxes/GST and statutory levies as notified by the Govt., after the date of submission of tender, the Tenderer will be paid based on the unit rate worked out on the basis of the reduced taxes/GST/statutory levies without any change in the basic price or the price structure of the drugs approved under the tender. Any increase or decrease in taxes/GST and statutory levies will be considered based on the notification issued by the Government.

However, if the firm supplies after originally stipulated Delivery period, increase in taxes/GST due to statutory variation in taxes/GST shall be borne by the supplier. In case of decrease in taxes/GST due to statutory variation in taxes/GST, the same shall be passed on by the supplier to the BPPI.

17. HANDLING & TESTING CHARGES:

No handling & testing charges shall be applicable.

18. LIQUIDATED DAMAGES AND OTHER PENALTIES:

18.1. If the supply reaches the designated places or Central Warehouse after 5 PM of **45th day** from the date of issue of the Ist purchase order and after 5 PM of the **30th day** from the date of issue of the subsequent purchase order, a liquidated damages will be levied at 2% per week or part thereof, subject to maximum of 10% irrespective of the fact that whether the BPPI has suffered any damage/loss or not, on account of delay in effecting supply. If the 45th/30th day happens to be a holiday the supply will be accepted on the next working day without any penalty.

- **18.2.** If the supply is received in damaged condition, open delivery of the supplies shall be received, wherein it is possible to physically inspect the shipment. Damaged products shall not be accepted.
- **18.3.** All the Tenderers are required to supply the product(s) with printed logogram of appropriate size on the strips, blisters, vials, ampoules& bottles and with prescribed packing specification. If there are any deviation in these Tender conditions, action will be taken to blacklist the product and/or a separate damages will be levied @ 5% of value of the defaulted quantity irrespective of the Tender Inviting Authority having actually suffered any damage/loss or not, without prejudice the rights of alternative purchase specified in Clause No.14.11 and 13.4.

19. DEDUCTION & OTHER PENALTIES ON ACCOUNT OF QUALITY FAILURE:

- 19.1. If the samples do not conform to statutory standards, the Tenderer will be liable for relevant action under the existing laws and the entire stock in such batch has to be taken back by the Tenderer within a period of 30 days of the issue of the letter from the BPPI Such stock shall be taken back at the expense of the Tenderer. Further, actual handling and testing charges shall be paid to BPPI by the supplier otherwise these charges shall be recovered from their pending bill/EMD/security deposit. The BPPI has the right to destroy such "NOT OF STANDARD QUALITY DRUGS" if the Tenderer does not take back the goods within the stipulated time. The BPPI will arrange to destroy the "NOT OF STANDARD QUALITY DRUGS" after the expiry of 30 days mentioned above without further notice, and shall also collect demurrage charges calculated at the rate of 2% per week on the value of the drugs rejected till such time stipulated. Futher, the cost of disposal shall be recovered from the supplier.
- 19.2. If any items of Drugs/Medicines supplied by the Tenderer have been partially or wholly used or consumed after supply and are subsequently found to be in bad odour, unsound, inferior in quality or description(Adulterated/Spurious/Misbranded) or otherwise faulty or unfit for consumption, then the contract price or prices of total such batches supplied will be recovered from the Tenderer, if payment had already been made to him. In other words the Tenderer will not be entitled to any payment whatsoever for Items of drugs found to be of "NOT OF STANDARD QUALITY" whether consumed or not consumed and the Tender Inviting Authority is entitled to deduct the cost of such batch of drugs from any amount payable to the Tenderer. On the basis of the nature of failure, action will be initiated to blacklist the product/supplier.
- 19.3. For the supply of Adulterated/Spurious/Misbranded, as defined in the Drugs and Cosmetics Act, 1940, to BPPI, BPPI reserves the right to blacklist the supplier. No further supplies shall be accepted from the firm/company. If the tenderer is blacklisted, the tenderer shall also not be eligible to participate in tenders of Tender Inviting Authority of BPPI for supply of Drugs for a period of 5 years from the date of blacklisting. In case of supply of NOT OF STANDARD QUALITY drug(s) to BPPI, the product shall be blacklisted by BPPI and no further supplies shall be accepted for the particular drug(s). The Tenderer shall also not be eligible to participate in tenders of BPPI for supply of such Drugs for a period of 2 years from the date of blacklisting. In addition, the Director of Drugs Control of concerned State will be informed for initiating necessary action on the Tenderer in their state. Security deposit will also be forfeited without any intimation.

- **19.4.** The Tenderer shall furnish the source of procurement of raw material utilized in the formulations, if required by the BPPI. The BPPI reserves the right to cancel the purchase orders, if the source of supply is not furnished.
- **19.5.** The decision of the BPPI or any officer authorized by him, as to the quality of the supplied drugs, medicines etc., shall be final and binding. In such cases, the BPPI will be at liberty to terminate, the contract either wholly or in part on 30 days' notice. The Tenderer will not be entitled for any compensation whatsoever in respect of such termination besides forfeiture of Security deposit.
- **19.6.** For contravention of the stipulations of the contract or for other justifiable reasons, the contract may be terminated by the BPPI, and the Tenderer shall be liable to pay for all losses sustained by the BPPI in consequence of the termination which may be recovered from the Tenderer, as per rules besides forfeiture of Security deposit.
- **19.7.** Non-performance of any of the contract conditions and provisions will disqualify a firm from participating in the tender for the next 2 years besides forfeiture of Security deposit.
- **19.8.** In the event of making Alternative Purchase, as specified in Clause 12.4 (a), Clause 14.11 and in Clause 15.3 penalty will be imposed on the supplier. The excess expenditure over and above contracted prices incurred by the BPPI in making such purchases from any other sources or in the open market or from any other Tenderer who has quoted higher rates and other losses sustained in the process, shall be recovered from the Security Deposit or from any other money due and become due to the supplier and in the event of such amount being insufficient, the balance will be recovered personally from the supplier as per rules.
- **19.9.** In all the above conditions, the decision of the BPPI shall be final and binding.

20. BLACK LISTING IN THE EVENT OF WITHDRAWL FROM THE TENDER, AND NON-ADHERENCE TO THE QUALITY STANDARDS AND SUPPLY SCHEDULE

20.1. BLACKLISTING OF PRODUCT/TENDERER ON WITHDRAWAL OF TENDER

(a) If the Tenderer(s) fails to execute the agreement / to perform the obligations under the tender conditions / commits default in the performance of the contract, such Tenderers will be blacklisted for a period of 2 years by BPPI from the date of observing the defect besides forfeiture of Security deposit.

BLACKLISTING FOR QUALITY FAILURE

20.2.1. Quality Test by the Empanelled Laboratories of BPPI

a. Each batch of drugs/medicines shall be subjected to quality test by the Empanelled laboratories.

- b. The samples collected from each batch of supply of the each drugs will be sent to the empanelled testing laboratories for testing the quality of drugs. In addition to the above BPPI shall also draw the samples of products supplied in the market place and get the same tested, to make sure the products are conforming to quality requirements.
- c. If such sample passes quality test in all respects, BPPI will instruct its Warehouse to release such items of drugs.
- d. If the sample fails in quality test and report is received certifying that sample is "NOT OF STANDARD QUALITY" then supplies will be rejected & no further procurement of that drug from the supplier for two years from the date of sample being declared not of standard quality. If the supplier challenges and request for re-testing, the rejected supply shall be tested in two labs simultaneously at the cost of supplier. The cost testing shall be recovered from the supplier.
- (i) If such sample passes the quality test in both laboratories, the drugs representing the sample shall be qualified for issue to various Institutions.
- (ii) If the sample passes in one laboratory and fails in other laboratory or fails in both laboratories, the supply shall be rejected. No further procurement of said drug shall be made from such supplier.
- (iii) If **3** batches of item/drug supplied by the same supplier is reported to NOT OF STANDARD QUALITY in specification, then the firm shall be blacklisted for 2 years after observing procedure laid down in Para 20.2.3 besides forfeiture of Security Deposit.

20.2.2 Quality Test by Statutory Authorities:

- (a) If any drug is declared "NOT OF STANDARD QUALITY", by any government agencies or drug licensing authority, the issue of available stock of the particular item will be stopped. Further, the available stock of the product in hospitals/JAS will be retrieved.
- (b) If any batch of any product(s) supplied by the company/firm declared, NOT OF STANDARD QUALITY in specification as defined in the Drugs and Cosmetics Act, 1940, by the Government Authorities during the relevant tender period or during quality check within shelf life period, the company/firm shall be blacklisted for a period of 2 years from the date of blacklisting after observing procedure laid down in Para 20.2.3.

20.2.3 Procedure for Blacklisting:

(i) On receipt of complaint from Distributer/retailers/customers or report from Govt. Analyst/Drug Testing Laboratory indicating that a particular Item/Drug is "NOT OF STANDARD QUALITY/ ADULTERATED/ SPURIOUS/MISBRANDED" (As the case

may be), a show cause notice shall be issued to the supplier calling for explanation within 7 days from the date of notice. On receipt of explanation from the supplier, the CEO, BPPI may take appropriate action on merits of the case and impose penalty including the blacklisting of the item of the product/company or firm as deemed fit besides forfeiture of Security deposit

- (ii) If a particular item of the drug has been blacklisted according to the procedure stated above, the supplier is not eligible to participate in any of the tenders for that particular item floated by the BPPI until the period of blacklisting is over.
- (iii) If a supplier company/firm is blacklisted according to the procedure stated above, such supplier is not eligible to participate in any of the tenders floated by the BPPI until the period of blacklisting is over.

20.3 BLACKLISTING FOR NON-SUPPLY:

Due to non supply of item against any purchase order, 5 % value of purchase order shall be recovered from the supplier in addition of other penal like risk purchase. In case of repeated circumstances of non supply of items i.e. 3 times , the supplier may be blacklisted for 2 years in addition of forfeiture of security deposit/ EMD and other penal action.

21. SAVING CLAUSE

No suit, prosecution or any legal proceedings shall lie against the Tender Inviting Authority or any person for anything that is done in good faith or intended to be done in pursuance of the tender.

22. RESOLUTION OF DISPUTES

(i) The BPPI and the supplier shall make every effort to resolve, amicably by direct informal negotiation any disagreement or dispute arising between them under or in connection with the contract,

ARBITRATION AND JURISDICTION

Normally, there should not be any scope of dispute between the BPPI and the supplier after entering into a mutually agreed valid contract/agreement.

However, due to various unforeseen reasons, problems may arise during the progress of the contract/agreement leading to disagreement BPPI and the supplier shall first try to resolve the same amicably by mutual Consultation. If the parties fail to resolve the dispute by such mutual consultation within twenty-one days, then, depending on the position of the case, either the BPPI or the supplier shall give notice to other party of its intension to commence Arbitration procedure as per Indian Arbitration and Conciliation Act, 1996. Such disputes/differences shall be referred to Sole Arbitrator to be appointed by the President/ CEO of BPPI. The venue of Arbitration Shall be at New Delhi. The award published by the Arbitrator shall be final and binding on the parties.

23. APPEAL:

(i) Any Tenderer aggrieved by the order passed by the Tender Accepting Authority under section 10 of the said Act, may appeal to the Department of Pharmaceuticals, Ministry of Chemical and Fertilizer, Government of India within ten days from the date of receipt of order and the Department of Pharmaceuticals, Ministry of Chemical and Fertilizer, Government of India shall dispose the appeal within fifteen days from the date of receipt of such appeal.

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(ii) No Appeal shall be preferred while the tender is in process and until tender is finalized and Notification of award is issued by the BPPI.

24. CONTACTING THE BPPI BY THE BIDDER:

- (i) No bidder shall contact the *BPPI* on any matter relating to its bid, from the time of bid opening to the time the contract is awarded.
- (ii) Any effort by a bidder to influence the *BPPI* in the *Purchaser*'s bid evaluation, bid comparison or contract award decisions may result in rejection of the bidder's bid.
- (ii) The bidder shall not make any attempt to establish unsolicited and unauthorized contact with the Tender Accepting Authority, Tender Inviting Authority or Tender Scrutiny Committee after opening of the bids and prior to the notification of award and any attempt by any bidder to bring to bear extraneous pressures on the Tender Accepting Authority, Inviting Authority or Tender Scrutiny Committee, shall be sufficient reason to disqualify the bidder.
- (iv) Not withstanding anything contained in clause (iii) above the Tender Inviting Authority or the Tender Accepting Authority, may seek bonafide clarifications from bidders relating to the bids submitted by them during the evaluation of bids.

25. FRAUDULENT AND CORRUPT PRACTICES:

(1)For bidders:

It is purchaser's policy to ensure that suppliers and their authorized representatives/agents observe the highest standard of ethics during the procurement and execution of such contracts. (In this context, any action taken by a bidder, supplier, contractor, or by their authorized representatives/agent, to influence the procurement process or contract execution for undue advantage is improper) In pursuance of this policy, the purchaser;

- (a) defines, for the purposes of this provision, the terms set forth below as follows:
- (i) "corrupt practice" is the offering, giving, receiving or soliciting, directly or indirectly, of anything of value to influence improperly the actions of another party ("another party" refers to a public official acting in relation to the procurement process or contract execution]. In this context, "public official" includes staff and employees of other organizations taking or reviewing procurement decisions.
- (ii) "fraudulent practice" is any act or omission, including a misrepresentation, that knowingly or recklessly misleads, or attempts to mislead, a party to obtain a financial or other benefit or to avoid an obligation (a "party" refers to a public official; the terms "benefit" and "obligation" relate to the procurement process or contract execution; and the "act or omission" is intended to influence the procurement process or contract execution).

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- (iii) "collusive practice" is an arrangement between two or more parties designed to achieve an improper purpose, including to influence improperly the actions of another party ["parties" refers to participants in the procurement process (including public officials) attempting to establish bid prices at artificial, non competitive level].
- (iv) "coercive practice" is impairing or harming, or threatening to impair or harm, directly or indirectly, any party or the property of the party to influence improperly the actions of a party (a "party" refers to a participant in the procurement process or contract execution).
- (v) "obstructive practice" is (a) deliberately destroying, falsifying, altering or concealing of evidence material to the investigation or making false statements to investigators in order to materially impede a investigation into allegations of a corrupt, fraudulent, coercive or collusive practice; and/or threatening, harassing or intimidating any party to prevent it from disclosing its knowledge of matters relevant to the investigation or from pursuing the investigation; or acts intended to materially impede the exercise of the purchaser's inspection and audit rights provided for under sub-clause (e) below.
- (b) will reject a proposal for award if it determines that the bidder considered for award has, directly or through an agent, engaged in corrupt, fraudulent, collusive, coercive or obstructive practices in competing for the contract in question;
- (c) will cancel the contract if the purchaser determines at any time that the bidder, supplier and contractors and their sub contractors engaged in corrupt, fraudulent, collusive, or coercive practices.
- (d) will sanction a firm or individual, including declaring in eligible, either indefinitely or for a stated period of time, to be awarded a contract if it at any time determines that the firm has, directly or through an agent, engaged in corrupt, fraudulent, collusive, coercive or obstructive practices in competing for, or in executing, a contract; and
- (e) will have the right to inspect the accounts and records of the bidders, supplier, and contractors and their subcontractors/authorized representatives and to have them audited by auditors appointed by the purchaser.

(2) For suppliers:

If the BPPI determines that a Supplier has engaged in corrupt, fraudulent, collusive, coercive or obstructive practices, in competing for or in executing the Contract, then the BPPI may, after giving 7 days notice to the Supplier, terminate the Supplier's engagement under the Contract and cancel the contract, and the procurement will be made at the risk and cost of the supplier besides blacklisting the bidder for 5 years with forfeiture of Security Deposit apart from other penal actions.

- (a) For the purposes of this Sub-Clause:
- (i) "corrupt practice" is the offering, giving, receiving or soliciting, directly or indirectly, of anything of value to influence improperly the actions of another party;
- (ii) "fraudulent practice" is any act or omission, including a misrepresentation, that knowingly or recklessly misleads, or attempts to mislead, a party to obtain a financial or other benefit or to avoid an obligation;
- (iii) "collusive practice" is an arrangement between two or more parties designed to achieve an improper purpose, including to influence improperly the actions of another party;
- (iv) "coercive practice" is impairing or harming, or threatening to impair or harm, directly or indirectly, any party or the property of the party to influence improperly the actions of a party;

(v) "obstructive practice" is (aa) deliberately destroying, falsifying, altering or concealing of
evidence material to the investigation or making false statements to investigators in order to materially
impede a purchaser investigation into allegations of a corrupt, fraudulent, coercive or collusive practice;
and/or threatening, harassing or intimidating any party to prevent it from disclosing its knowledge of
matters relevant to the investigation or from pursuing the investigation; or (bb)acts intended to materially
impede the exercise of the purchaser's inspection and audit rights provided for.

26. JURISDICTION

In the event of any dispute arising out of the tender such dispute would subject to the jurisdiction of the Civil Court within the city of Delhi only.

ANNEXURE I

(BARCODE REQUIREMENTS)

Reference clause 2(k)

GS1 barcode requirements on Drugs procured by Bureau of Pharma Public Sector undertakings of India (BPPI)

These requirements cover medicines/drugs procured by Bureau of Pharma Public Sector Undertakings of India (BPPI), Gurgaon, meant for supply and distribution through BPPI regulated distribution channel.

Barcode based on GS1 identification standards are provided below at various levels of product packaging which includes primary, secondary and shipper/carton levels and need to be complied with while supplying medicines/drugs to BPPI.

GS1 India is unique identification & barcoding standards body setup by Ministry of Commerce & Industry, Govt. of India along with APEDA, BIS, Spices board, IIP and apex industry chambers like CII, FICCI, ASSOCHAM to assist India industry and govt. bodies on adoption of global standards.

Suppliers are also required to provide GS1 subscription validity certificate at the time of supply of medicines/drugs issued by GS1 India. For validity certificate suppliers can contact GS1 India at 011-4289-0890

Barcodes based on GS1 global standards are required to be printed on product packaging at primary, secondary and tertiary packaging levels **in addition** to other, existing statutory labelling & marking requirements.

Technical Specification for GS1 Standards

Tertiary Level Pack: Data attributes captured

- *a) Unique product identification code (GTIN Global Trade Identification Number)*
- *b)* Expiry date
- c) Batch no.
- d) Quantity
- e) Serial Shipping Container Code (SSCC)

e.g. 1st Barcode: (02)1 8901072 00253 4 (17) 180815 (10) RNBXY0514 (37)5000 2ndBarcode: (00) 1 8901072 001234567 6

Attribute	Description	Length	Nature	Data Type
(02)	Application Identifier to indicate GTIN-14 Brackets not encoded in the	2	Fixed	Numeric

	barcode			
1 8901072 00253 6	Unique Product Number-GTIN-14	14	Fixed	Numeric
(17)	Application Identifier to indicate Expiry date Brackets not encoded in the barcode	2	Fixed	Numeric
180815	Expiry Date in YYMMDD format	6	Fixed	Date
(10)	Application identifier to indicate Lot/batch number Brackets not encoded in the barcode	2	Fixed	Numeric
RNBXY0514	Batch No / Lot No	20	Variable	Alphanumeric
(37)	Application identifier to indicate Quantity in Outer Carton	2	Fixed	Numeric
5000	Quantity/no of units	Upto 8	Variable	Numeric
(00)	Application identifier to indicate the SSCC Brackets not encoded in the barcode	2	Fixed	Numeric
1 8901072 001234567 6	Unique number of the tertiary pack	18	Fixed	Numeric
Recommended Barcode – GS-128				

To,

Warehouse-BPPI, Gurgaon Haryana Mnfd By,

AAA Pharma Company 125, SEZ

Ahmedabad-382213 Gujrat

Drug Name: Dobucin 500 mg Exp Date: 15 Aug 2018 Batch No: RNBXY0514





Secondary Level Pack: Data Attributes Captured

- a. Unique product identification code (GTIN)
- b. Expiry date
- c. Batch No.
- d. Qty

e.g. Barcode - (02) 1 8901072 00253 6 (17) 180815 (10) RNBXY0514 (37) 500

Attribute	Description	Length	Nature	Data Type
(02)	Application Identifier to indicate GTIN-14. Brackets not encoded in the barcode	2	Fixed	Numeric
1 8901072 00253 6	GTIN-14- Unique product code with first digit being the packaging indicator	14	Fixed	Numeric
(17)	Application Identifier to	2	Fixed	Numeric

500	Quantity/Units in Secondary pack	Upto 8	Variable	Alphanumeric
(37)	Brackets not encoded in the barcode	2	<i>г</i>	Ivumeric
(27)	Application Identifier to indicate serial number	2	Fixed	Numeric
RNBXY0514	Batch No / Lot No	Upto 20	Variable	Alphanumeric
(10)	Application identifier to indicate Lot/batch Brackets not encoded in the barcode	2	Fixed	Numeric
180815	Expiry Date in YYMMDD format	6	Fixed	Date
	Brackets not encoded in the barcode			
	indicate Expiry date			

Recommended Barcode depending upon the space available – GS1 Data matrix

Or

GS1-128



(02) 1 8901072 00253 6 (17) 180815 (10) RNBXY0514 (37) 500

or



(02) 1 8901072 00253 6 (17) 180815 (10) RNBXY0514 (37) 50

Primary Level Pack: Data Attributes Captured

a. Unique product identification code (GTIN)

Barcode e.g. - (01) 1 8901072 00253 6

Attribute	Description	Length	Nature	Data Type
(01)	Application Identifier to indicate GTIN-14 Brackets not encoded in the barcode	2	Fixed	Numeric
1 8901072 00253 6	GTIN-14 with first digit being the packaging indicator	14	Fixed	Numeric
Recommended Barcode – GS1 Datamatrix,		(01) 1 89010	72 00253 6	

Please contact GS1 India office for any further assistance – GS1 India

(Under Min. of Commerce, Govt. of India) 330, 2nd Floor, 'C' Wing, August Kranti Bhawan, Bhikaji Cama Place, New Delhi - 110066

T +91-11-42890890, (D) +91-11-42890846

F +91-11-26168730

E ankit@gs1india.org

W http://www.gslindia.org

ANNEXURE-II

{ Clause 4.1(a)(i)}

MANUFACTURER'S AGREEMENT WITH MARKETER

To
The CEO
Bureau of Pharma PSUs of India,
IDPL corporate office Complex, Old Delhi-Gurgaon Road,
Dundahera, Gurgaon- 122016 (Haryana),

	orporate office Complex, Old Delhi-Gurgaon Road, hera, Gurgaon- 122016 (Haryana) ,
Dear S	Sir,
1.	We who are established and reputable manufacturer of Drugs having factory/factories atand hereby declare that we do not market our products. Therefore, we authorize M/S (Name and address) to bid, negotiate and contract with your tender No. BPPI/Drug-054/2017 dated 31/08/2017 for supply of drugs manufactured by us.
2.	No company of the firm or individual other than M/Sauthorised to bid, negotiate and conclude the contract in regard to this business against this specific tender as also for a business in the entire territory of India.
3.	This agreement is valid fromto(This period will be the date of opening tender till valid one year shelf life of the drugs or period of contract/price agreement whichever is more.
4.	The ex-factory cost of the Drugs being quoted will be provided by us whenever called for. We also undertake that we will not quote a price higher than supplied to any institute in last 6 months. In case our submission is found wrong, we undertake to be liable for punitive action in the form of recovery of excess amount/ withholding of payment/ any other action as deemed appropriate by department.
5.	An marketing commission of% is included in the gross ex-works price is applicable of
N	N/s

- 6. We hereby extend our full guarantee and warrantee as per relevant conditions of contract for the goods offered for supply against this invitation for bid by the above firm. In the event of failure by authorized marketer in honouring the contract, we undertake to provide remedial action at the earliest without any additional charges.
- 7. Our other responsibilities include.

 Provision of requisite inspection and testing fa 	cilities at our works in respect of
supply order placed on our agent.	
ii	
{Here specify in detail manufacturers responsibilities	es]
The services to be rendered by the marketer, M as under:-	/sare
i	
ii	
{Here specify the services to be rendered by the	ne agent}
8. We certify that neither we, nor our agent is blacklist by any Govt. agency.	ted/ debarred/de-registered/banned
9.We jointly agree to abide by the following clauses in	the contract:
a) Penalty for use of undue influence.b) Access to books of accounts.	
10. We undertake that GST registration numb (PAN), last three years Bank Statement of Accounts, agent/ distributor will be provided as and when demand	Banker details of our authorized
Yours faithfully,	
(Name, Signature & Stamp)	(Name, Signature & Stamp)
(Name of Authorized Marketer)	(Name of Manufacturer)
Date:	Date:
NOTE: THIS LETTER OF AUTHORIZATION SHOULD BE OMANUFACTURING CONCERN AND SHOULD BE INK MANUFACTURERE & THE AUTHORIZED MARKETER, BY A	SIGNED BY BOTH THE ORIGINAL

MANUFACTURING CONCERN AND SHOULD BE INK SIGNED BY BOTH THE ORIGINAL MANUFACTURERE & THE AUTHORIZED MARKETER, BY A PERSON WHO IS COMPETENT AND HAVING THE POWER OF ATTORNEY TO BID THE MANUFCTURER, A COPY OF NOTARIZED POWER OF ATTORNEY SHOULD ALSO BE FURNISHED, NAME, SIGNATURE AND OFFICAL STAMP OF MANUFACTURER AND MARKETER TO BE APPENDED.

ANNEXURE –III

(On nonjudicial Stamp Paper)

Ref. Clause No. 4.1(a)(ii)

DECLARATION

(NOTE:-In case Bid is submitted by the Marketer, this declaration is to be signed by Marketer as well as Manufacturer)

represented by its Proprietor/Managing Partner /Managing Director

I/We M/s.

having its registered office at	at
do hereby declare as under:-	
(I) that I/we have carefully read all the terms and conditions of tender in ref. no. BPPI/Dru 054/2017 dated 31/08/2017 including Amendment(s) to Tender document (if any) issued Bureau of pharma public sector undertakings of INDIA, GURGAON, 122016 and account unconditionally all terms and condition of tender document including Amendment(s) Tender document (if any).	by ept
(II) that I/We are holding and have uploaded (a) valid drug license for quoted drugs,(b) valid WHO-GMP certificate, (c) 2 years market standing certificate for quoted products issued licensing authority for quoted drugs, (d) a certificate manufactured & marketed two batch within 3 years issued by C.A. for quoted drugs, (e) valid non conviction certificate not of than 6 months,(f) Valid Import license (If applicable) and also enclosed undertaking/declaration as per Annexure mentioned in the tender document. On the basis such undertaking, the price bid shall be opened within a week after opening of technical However, any document uploaded with technical bid is not complying as per undertaking, contract/agreement shall be cancelled with forfeiture of EMD/SECURITY DEPOSIT/B guarantee against tender no. BPPI/Drug-054/2017 dated 31/08/2017 along with other action.	by thes all s of bid the
(III) I/We declare that we possess the valid drug manufacturing licence and WHO-GMP(Wo Health Organisation-Good Manufacturing Practices) Certificate issued by competent author and complies and continue to comply with the condition lied in schedule M of Drug & cosme act, 1940 the rules made there under.	ity

- I am / We are aware of the Tender inviting Authority's right to forfeit the Earnest Money Deposit and /or Security Deposit and blacklist me/us for a period of 5 years if,any information furnished by us proved to be false at time the of inspection and not complying the condition as per schedule M of the said Act for a period of five years.
- (IV) (a) I do hereby declare that I will supply the drug with bar code as per ANNEXURE I and as per the design as per enclosures to ANNEXURE XIV enclosed with tender document as well as other instruction given in this regard.
- (b) Further, I / we do hereby declare that I will supply the drugs by affixing logo on Primary/Secondary/ Tertiary packing for the imported items along with the generic name as per BPPI/DRUG-054/2017

the designs given in enclosures to Annexure XIV A as well as other instructions given in this regard.

(V) that in pursuant to the conditions in Clause No. 7.2 of the tender, the Earnest Money Deposit can be forfeited by the Tender Inviting Authority in case of violation of any of the conditions and non-performance of the obligation under tender document.

(VI) that our company/applied items have not been blacklisted/debarred/de-registered/banned due to quality failure of the drugs supplied either by any State government or Central Government Organization or its drug procurement agencies for the following products quoted in the tender at the time of submission of bid. Further, quoted drugs have not been failed in house testing or testing by any State Government/Central Government / its Drug procurement agencies/BPPI during last two years. We are eligible to participate in the tender ref. No. **BPPI/Drug-054/2017 dated 31/08/2017** for the following products:-

S. No.	Drug Code	Name of the Drug
		Signed
		Name
		Designation
		(Company Seal)
Witnes	s:-(1)	
	(2)	

To be attested by the Notary

ANNEXURE-IV

Ref. Clause No. 7.1 & 3(ii)

DETAILS OF E.M.D SUBMITTED UPLOAD THE SCANNED COPY OF DRAFT/ PAY ORDER/BANK GURANTEE

ANNEXURE-V

Ref. Clause No. 4.1 (f)(i)

(Format for a certificate from the C.A.(Chartered Accountant) or ICWA)

It is certified that M/s	has Financial capacity to
manufacture and deliver the drugs quoted by them in the tender as	per quantity & delivery
schedule mentioned in tender. This certificate is based on their Ma	anufacturing capacity,
inventory of raw Material and financial statement.	
Date (Name	e, Signature & Stamp) Registration no.

ANNEXURE-V A

Ref. Clause No. 4.1 (f)(i)

(Format for a certificate from the C.A.(Chartered Accountant) or ICWA)

It is certified that M/s	has Production and Financial
capacity to manufacture and deliver the drugs quoted by them in	the tender as per quantity &
delivery schedule mentioned in tender. This certificate is based	on their Manufacturing
capacity, inventory of raw Material and financial statement.	
Date (Nar	ne, Signature & Stamp) Registration no.

ANNEXURE -VI

Ref. Clause No. 4.1{f(ii)}

ANNUAL TURNOVER STATEMENT

The annual Turnover of M/s.for the past three years are given below

Rs.....Lakhs

Registration no.

Sl.No.	Financial Year	Turnover in Lakhs(Rs.)
1.	2013-14	
2.	2014-15	
3.	2015-16	
TOTAL		Rs Lakhs

and certified that the statement is true and correct.

Average Turnover per annual

Date: Signature of Auditor/Chartered Account
Seal: (Name in Capital)

ANNEXURE – VII

Ref. clause 4.1 (q)

LIST OF ITEMS QUOTED

Sl.No.		Detai	ils		
1.	Name of the firm and address				
	(As given in	n Drug licence)			
2.	Drug Licen	ce No. in form 2:	5 & 28		
	Or import L	icence No.			
3.	Date of issu	e & validity			
4.	WHO-GMF	(World Health (Organisation-Good		
	Manufactur	ing Practices) Co	ertificate obtained	on	
5.	Non-convic	tion Certificate (Obtained on		
6.	Market stan	ding Certificate	Obtained on		
7.	Details of E	Indorsement for a	all products quoted	l:	
Sl.No.	Drug	Drug Name	Specifications	Date of	Whether
	Code		IP/BP/USP	Endorsement	Endorsement
				obtained from	is in Generic
	the State Drugs or Trade				
				Controller Name	
1.					
2.					

Authorised signatory: Date:

ANNEXURE – VIII

Ref. Clause 4.1 (r)

CHECK-LIST(Whether Uploaded the documents)

COVER – A

S.No.	Check List	YES	NO
1.	Check list - ANNEXURE – VII		
2.	EMD Rs.100,000/- in the form of Bank Guarantee or Bankers		
	Cheque or Demand Draft uploaded as per ANNEXURE-IV DD		
	NoDatedissued by		
	(name of bank) and delivered to BPPI.		
	Uploaded NSIC certificate for exemption if any.		
3.	Documentary evidence for the constitutions of the		
	company / concern		
4.1	Documentary evidence for MSME if applicable		
4.2	In case of MSME, Documentary evidence that unit owned		
	by SC/ST entrepreneurs if applicable		
5.	Scanned copy of License for the Product duly approved by		
	the Licensing Authority for each and every product quoted		
6.	Scanned copy of Import License, if Imported and whole		
	sale Drug license		
7.	COPP certificate as per WHO format of their Principal		
	Manufacturing company including Imported drugs.		
8.	The instruments such as power of attorney, Resolution of		
	board etc.,		
9.	Authorization letter nominating a responsible Person of the		
	tenderer to transact the business with the Tender inviting		
	Authority.		
10.	Scanned copy of Market Standing Certificate issued by the		
	Licensing Authority		
11.	A certificate from their C.A.or ICWA that manufactured at		
	least 2 commercial batch in last three years.		
12.	Scanned copy of WHO-GMP(World Health Organisation-		
	Good Manufacturing Practices) Certificate.		
13.	Scanned copy of Non Conviction Certificate issued by the		
	licensing authority not older than 6 months.		
14.	Scanned copy of GST registration certificate/Latest Sales		
	Tax Clearance Certificate/returns filed.		
15.	Scanned copy of Latest income tax assessment		
10.	orders/returns filed.		
16.	Copies of approval of the any agency like US FDA, TG		
	Australia, Health Canada, EU, MCC South Africa		
	approval, Brazil Anvisa approved product , if any.		
17.	Scanned copy of ANNEXURE-II		
1,.	(Agreement with Manufacturer) if any , original Annexure I		
	delivered to BPPI.		
18.	Scanned copy of ANNEXURE –III (Declaration for eligibility in		
= -	participating the tender) original Annexure II delivered to		

	BPPI.	
18A	Scanned copy of ANNEXURE V{certificate from the	
	C.A.(Chartered Accountant) or ICWA that the bidder has	
	Production & financial capacity}.	
19.	Scanned copy of ANNEXURE V A in case of	
	importer{certificate from the C.A.(Chartered Accountant)	
	or ICWA that the bidder has financial capacity}.	
20.	Scanned copy of ANNEXURE -VI (A certificate from CA	
	for Annual Turnover Statement)	
21.	Scanned copy of ANNEXURE - VII(List of Items quoted without rates) .	
22.	Scanned copy of ANNEXURE-XI	
	(Details for Manufacturing Capacity & Batch Size)	
23.	Scanned copy of ANNEXURE—XV (Mandate form)	

NOTE:-EMD instrument, ANNEXURE II (if applicable) and ANNEXURE III are to be delivered in original to BPPI, Gurgaon on or before 'Bid opening date.

Name and signature of authorise	ed signatory (with company seal)	
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Annexure -IX

Clause 8.1 &8.2

Bureau of Pharma Public Sector Undertakings of India, Gurgaon Tender for supply of drugs (Tender No. BPPI/Drug-054/2017) dated 31/08/2017)

(1)	(2)	(3)	(4)	(5)	(6)	(7)
Sr. No.	Drug Code	Generic name of Medicines	Unit Size	Pack Size	Packing per Carton (Shipper Pack)	Tender quantity in unit size
1	8	Diclofenac Sodium + Serratiopeptidase (50mg + 10mg) Tablets	10's	10'sX10	(10's x10x10)x10	800000
2	9	Diclofenac Sodium (SR) Tablets IP 100 mg	10's	10's x10	(10's x10x10)x10	1200000
3	10	Diclofenac Sodium 25mg per ml Inj. IP	3 ml ampoul e	10X1's	100x10x1's	1200000
4	14	Ibuprofen 400mg + Paracetamol 325mg Tablets	15's	15's x 10	(15's x10x10)x 5	1500000
5	15	Ibuprofen film coated Tablets IP 200mg	10's	10's x10	(10's x10)x10x10	1000000
6	30	Amikacin Sulphate 50mg/ml inj. IP	2ml Vial	1's x 20	1'sx20x50	300000
7	36	Amoxycillin + Clavulanic acid (200 mg+28.5 mg /5ml) Dry Syrup	30 ml bottles	1's x 10	20 x (1's x 10)	300000
8	39	Amoxycillin + Clavulanic acid (500 mg + 125 mg) film coated Tablets IP	6's	6's x 10	(6's x 10 x 10) x10	1500000
9	43	Amoxycillin 125mg/ 5ml Powder for Suspension IP	60 ml bottle	1's x 10	(1's x 10 x 50)	500000
10	44	Amoxycillin 250 mg Caps IP	10's	10's x10	(10's x10x10)x10	400000
11	45	Amoxycillin 500 mg Caps IP	10's	10's x10	(10's x10x10)x10	400000
12	47	Azithromycin (100mg/ 5ml)	15 ml	1's x 10	(1's x 10 x 50)	300000

		Suspension IP	bottle			
13	49	Azithromycin 250 mg film coated Tablets IP	10's	10's x10	(10's x10x10)x10	300000
14	50	Azithromycin 500 mg film coated Tablets IP	10's	10's x10	(10's x10x10)x10	1000000
15	51	Cefadroxil film coated Tablets IP 250mg	10's	10's x10	(10's x10x10)x10	500000
16	55	Cefixime 200mg film coated Tablets IP	10's	10'sX10	(10's x10x10)x10	500000
17	59	Cefotaxime Sodium & Sulbactam Sodium (1g + 500 mg) Inj.	Vial & wfi	1's x10	1's x10 x 20	200000
18	60	CEFOTAXIME SODIUM 250 MG & SULBACTAM SODIUM 125 MG INJECTION	VIAL & WFI	1's x10	1's x10 x 20	200000
19	61	Cefotaxime Sodium & Sulbactam Sodium (500 mg + 250 mg) Inj.	Vial & wfi	1's x 10	30 x (1's x 10)	200000
20	62	Cefotaxime Sodium Injections IP 1000mg	Vial & wfi	1's x 10	(1's x 10) x 50	200000
21	63	Cefotaxime Sodium Injections IP 250mg	Vial & wfi	1's x 10	(1's x 10) x 50	200000
22	64	Cefotaxime Sodium Injections IP 500mg	2ml Vial & wfi	1's x 10	(1's x 10) x 50	200000
23	66	Cefpodoxime 200 mg film coated Tablets IP	10's	10's x10	(10's x10x10)x10	400000
24	68	Ceftazadime 250 mg Inj.	Vial & wfi	1's x 10	30 x (1's x 10)	200000
25	71	Ceftriaxone + Tazobactum 1000 mg + 125 mg Inj.	Vial & wfi	1's x 10	10 x (1's x 10)	300000
26	84	Ciprofloxacin + Tinidazole (500 mg + 600 mg) film coated Tablets	10's	10's X 10	(10's x10x10)x10	300000
27	85	Ciprofloxacin 250 mg film coated Tablets IP	10's	10's x10	(10's x10x10)x10	200000
28	88	Co-trimoxazole (Sulphamethoxazole 200mg + Trimethoprim 40mg / 5ml) Susp	50 ml bottle	1's x 10	(10X1's)x50	200000

29	89	Co-trimoxazole Tablets IP (160 MG + 800 MG)	10's	10's x10	(10 x 10 x 10) x 5	200000
30	91	Co-trimoxazole Tablets IP (80 mg + 400 mg)	10's	10's x10	(10's x10x10)x10	200000
31	98	NORFLOXACIN 400 mg + TINIDAZOLE 600 mg FILM COATED Tablets	10's	10'sX10	(10's x10x10)x10	500000
32	99	Norfloxacin 400 mg film coated Tablets	10's	10's x 10	50 x (10X10's)	300000
33	104	Roxithromycin (50 mg/ 5ml) Susp.	30 ml bottles	1's x 10	50 x (1's x 10)	200000
34	106	Roxithromycin 300mg film coated Tablets	10's	10's x 10	50 X (10X10's)	200000
35	107	Tinidazole 300mg film coated Tablets	10's	10's x10	100 X 10X10's	400000
36	111	Benzyl Benzoate Application IP 25% w/v Lotion	100ml bottles	1's x 10	25 x (1's x 10)	100000
37	119	Fluconazole 150 mg Tablets IP	10's	10's x10	(10's x10x10)x10	400000
38	124	Povidone Iodine 5% w/w Ointment USP	250 gm Jar	1's x 10	(1's x 10)x 5	50000
39	126	Povidone Iodine 10 % Solution IP	500 ml bottles	1's x 5	(1's x 5) x 2	100000
40	127	Povidone lodine 5% Solution 100ml	100ml bottle	1's x 6	(1's x 6)x3	156000
41	130	Chlorhexidine + Cetramide (1.5 % w/v + 3% w/v) Solution	100 ml bottles	1's x 10	1's x 10 x 10	100000
42	133	Glibenclamide 2.5 mg Tabs IP	10's	10's x10	(10's x10) x10 x10	1000000
43	141	Glipizide Tablets IP 5mg	10's	10's x10	(10's x10x10)x10	700000
44	142	INSULIN INJECTION IP 40 IU/ml (Insulin Human Recombinant)	10 ml Vial	1's X10	1's x 10 x 20	300000
45	153	Cisplatin 10 mg Injection	Vial	1'sx10	(1'sx10x10)x10	30000
46	158	Etoposide Inj. IP 100 mg/5ml	Vial & wfi	1's x 10	(1's x 10) x 50	30000

47	171	Mannitol 20% w/v Injection	350 ml	1'sx10	(1'sx10x5)x10	200000
48	172	METRONIDAZOLE INJECTION IP 5 mg / ml	100 ml	(1's x 10)	(10X1's) x10	300000
49	176	Sterile Water for Injection	5ml Amp	1'sx10	100x(10x1's)	300000
50	179	Albendazole 400mg Tabs IP	1's	1's x10	(1's x10x10)x100	5000000
51	197	Lactulose Syrup 10 gm/ 15 ml	100 ml bottle	1's X 6	1's X 6 X 5	300000
52	198	Aluminium Hydroxide + Magnesium Hydroxide (250+250mg / 5ml) Suspension	170 ml	1's X 10	1's x 10 X 5	500000
53	200	Metoclopramide Injections IP 5mg/ml	2ml vial	1's x 10	(1's x 10 x 10) x 10	400000
54	216	RANITIDINE INJECTION IP 50 MG/ 2ML	2 ML vial	1'sx10	(1'sX10)x25	1137000
55	244	Theophylline 25.3mg + Etophylline 84.7mg /2ml Injections	2ml ampoul e	1's x 10	(1'sx10x10) x 10	500000
56	245	Etophyllin +Theophylline (77 mg + 23 mg) Tablets	10's	10's x10	(10's x10x10)x10	1000000
57	256	SALBUTAMOL Tablets IP 2 MG	10's	10's x 10	(10's x10x10)x10	400000
58	257	Salbutamol 2.5 mg Respule	2.5 ml	1's x 20	(10's x10x10)x10	500000
59	259	Salbutamol Syrup IP 2mg/5ml	100ml bottle	1's x 10	(1's x 10) x 10	300000
60	260	Salbutamol Tablets IP 4mg	10's	10'sx10	(10's x10x10)x10	401000
61	274	Dopamine HCl 200 mg/5ml Inj. IP	5 ml ampoul e	1's x 5	(1's x 5 x 10) x 10	200000
62	279	Furosemide 40mg Tabs	10's	10'sX10	(10's x10x10)x10	700000
63	280	Heparin Sodium 1000iu/ml Inj. IP	5 ml vial	1's x 5	(1's x 10 x 10) x 10	100000

64	290	Metoprolol 25 mg Tabs	10's	10'sX10	(10's x10x10)x10	700000
65	298	Telmisartan + Hydrochlorthiazide (40 mg + 12.5 mg) Tabs	10's	10'sX10	(10's x10x10)x10	2500000
66	300	Telmisartan 40 mg Tabs	10's	10'sX10	(10's x10x10)x10	3000000
67	305	Chloroquine Phosphate 250 mg film coated Tablets	10's	10'sX10	(10's x10x10)x10	200000
68	312	Oral Rehydration Salts Citrate IP 20.5 gm (WHO Formula) Sachet	1's	1'sx20	(1'sx20x10)x10	1000000
69	316	Betahistine 8 mg Tabs	10's	10's x10	(10's x10x10)x10	1200000
70	318	Carbamazepine 200mg Tabs IP	10's	10's x10	(10's x10x10)x10	700000
71	327	PHENYTOIN Tablets IP 100 mg	100's in Bottle	1's x 10	1's x 10 x 10	400000
72	331	Thyroxine Sodium 50 mcg Tabs	100's in A Bottle	1's x 10	1's x 10 x 10	1500000
73	333	DEXAMETHASONE 0.5 MG TABS IP	10's	10's x10	(10's x10x10)x20	600000
74	364	GLIMEPIRIDE 2 MG + METFORMIN HYDROCHLORIDE 500 MG TABLETS SR	15's	15'sx10	(15'sx10)x10x10	4000000
75	374	ARTEMETHER 80 MG + LUMEFANTRINE 480 MG SR TABLETS	6's	6's X 10	(6's X 10)X 10x10	100000
76	377	CLINDAMYCIN CAPSULES IP 300 MG	10's	10'sx10	(10'sx10x10)x20	400000
77	382	LINEZOLID TABLETS IP 600 MG	10's	10'sX10	(10's x 10x 10)x10	300000
78	392	GRISEOFULVIN TABLETS IP 250 MG IP	10's	10's x10	(10's x10x10)x10	100000
79	394	PYRANTEL PAMOATE ORAL SUSPENSION IP 250 MG/5 ml IP	10 ml bottle	1'sx10	(1'sx10x10)x10	300000
80	397	Oxytetracycline Cap I.P 250mg	10's	10'sx10	(10'sx10x10)x20	100000

81	407	IVERMECTIN TABLETS 12 MG	10's	10'sx10	(10's x 10x 10)x10	200000
82	420	ATORVASTATIN 10 MG+ CLOPIDOGREL 75 MG CAPSULES	10's	10'sx10	(10's x 10x 10)x10	2000000
83	421	NEBIVOLOL TABLETS IP 5 MG	10's	10's x10	(10's x10x10)x20	700000
84	424	CARVEDILOL TABLETS IP 3.125 MG	10's	10'sx10	(10's x 10x 10)x10	700000
85	438	INDAPAMIDE TABLETS IP 1.5 MG	10's	10'sx10	(10's x 10x 10)x10	300000
86	452	WARFARIN TABLETS IP 5 MG	10's	10'sx10	(10'sx10x10)x20	700000
87	461	BETAMETHASONE VALERAT 0.1 % w/w + NEOMYCIN SULFATE 0.5 % w/w CREAM	20 gm tube	1'sx20	1'sx20x10	500000
88	466	URSODEOXYCHOLIC ACID Tablets IP 300 mg	10's	10'sx10	(10'sx10x10)x10	700000
89	472	DOMPERIDONE 30 MG+ ESOMEPRAZOLE 40 MG CAPSULE	10's	10'sx10	(10's x10x10)x10	700000
90	476	LIQUID PARAFFIN 1.25 ML+ MILK OF MAGNESIA 3.75ML+ SODIUM PICOSULPHATE 3.33MG /5ML EMULSION	170 ml bottle	1'sX10	1'sx10x5	250000
91	480	LEVOSULPIRIDE (SR) 75 MG+ ESOMEPRAZOLE 40 MG (EC) CAPSULES	10's	10's x10	(10's x10x10)x10	700000
92	487	DICYCLOMINE 10 MG + DIMETHICONE 40 MG /5 ML SUSPENSION	30 ml bottle	1'sx10	1'sx10x50	200000
93	491	ITOPRIDE TABLET 50 MG	10's	10'sx10	(10'sx10x10)x10	500000
94	492	Sulfasalazine Tablets EC BP 500 MG	10's	10's x10	(10's x10x10)x10	200000
95	500	LEVO-THYROXINE TABLETS IP 100 MCG	100's in A Bottle	100 ML X6	(1'sX 10)X200	700000
96	506	LEVO-THYROXINE TABLETS IP 50 MCG IP	100's in Bottle	1'sX 10	(1'sX 10)X10x10	700000

97	517	THIOCOLCHICOSIDE 4 MG+ ACECLOFENAC 100 MG TABLETS	10's	10'sx10	(10'sx10x10)x20	400000
98	531	GUAIFENESIN 100 mg+ TERBUTALINE 2.5 mg+ BROMHEXINE 8 mg /10 ml SYRUP	100 ml bottle	1's X 6	(1's X 6) X20	500000
99	555	DOXOFYLLINE TABLETS IP 400 MG	10's	10'sx10	(10's x10x10)x10	200000
100	567	SALBUTAMOL 100 MCG + IPRATROPIUM 20 MCG /PUFF INHALER	200 MDI	1's x 10	(1's x 10)x10	50000
101	585	VITAMIN-D (CHOLECALCIFEROL) 60000 IU /1 GM SACHET	1 SACHE T	1's X 25	(1'sX25)x25	700000
102	592	L-LYSINE + MULTIVITAMINS (VIT- B1,B2,B3,B5,B6) SYRUP	200 ML bottle	1'sX10	1'sx10x5	500000
103	603	Cetirizine Dihydrochloride IP 5mg, Phenylephrine Hydrochloride IP 10mg., Paracetamol IP 325 mg Tablets	10's	10's x10	(10's x10x10)x10	500000
104	607	Beclomethasone Dipropionate 0.025% w/w, Neomycin Sulphate 0.5% w/w (3500 Unit /G) Chlorocresol 0.1% w/w Cream	15gm tube	1's x 20	(1's x 20) x 25	300000
105	608	Betamethasone 0.05% w/w + Salicylic acid 3% w/w cream	20 Gm	1'sx20	(1'sx20x10)x10	400000
106	630	laxative Suspension Liqid Paraffin 3.75ml+Milk of Magnesia 11.25ml	170ml Bottle	1'sX10	1'sx10x5	200000
107	634	Clobetasol Propionate BP0.05 % w/w, Neomycin Sulphate IP0.50 % w/w., Miconazole Nitrate IP2.00 % w/w,Chlorocresol IP(as preservative) 0.10 % w/w Cream/Ointment	20gm tube	1's x 20	(1's x 20) x 25	500000
108	643	Paracetamol 125mg+ CPM 1 mg + Sodium Citrate 60mg in a flavour syrup base	60 ml Bottle	60ml X 10	60ml X 10 X10	300000
109	647	Diclofenac 1.16%w/w+ Menthol 5%+ Oleum 3% + Methyl Salicylate 10% Gel	20gm tube	1's x 20	(1's x 20) x 25	1000000

110	648	Diclofenac Diethylamine BP 1.16 %, Linseed Oil BP 3 % w/w, Methyl Salicylate IP 10 % w/w, Menthol IP 5 % w/w, Excipients and Propellant q.s. to 100 % w/w Spray	35 gms.	1'sx10	1'sX10)x25	500000
111	649	Dicyclomine 10mg + Act. Dimethicone 40mg per ml	10ml Bottle	1's X 10	1's X 10 X 20	200000
112	652	Dicyclomine 10mg + Mefenamic 250 mg Tablets	10's	10'sx10	(10's x10x10)x10	200000
113	657	Hydroquinone 2.0% w/w + Tretinoin 0.025% w/w + Mometasone Furoate 0.1% w/w in a cream base q.s	15gm tube	1's x 20	(1's x 20) x 25	200000
114	671	Diacerein 50 mg + Glucosamine Sulphate 500 mg Tablet	10's	10'sx10	(10's x10x10)x10	700000
115	676	Triamcinolone Acetonide 0.1 % Mouth Ulcer gel	10gm tube	1's X 20	1's X 20X 20	50000
116	678	levodopa 250mg, Carbidopa 25mg Tablets IP	10's	10's x10	(10's x10x10)x10	200000
117	687	Lactulose 10gm/15 ml syrup	200ml bottle	1'sx6	1'sx6x10	360000
118	691	Ofloxacin Ophthalmic Solution IP 0.3% w/v	10 ml	10ml x 10	(10ml x 10) x 50	300000
119	692	Olopattadine 0.1% w/v Eye Drops	10 ml Vial	10ml X 10	10ml X 10 X 20	200000
120	701	Pilocarpine Eye drop IP 2 %	10 ml Vial	10ml x 10	10ml x 10) x 50	200000
121	713	Glibenclamide 5mg + MetforminHcl 500 mg tab	10's	10's x10	(10 x 10 x 10) x 5	700000
122	723	Water for Injection amp polypack	10ml	10ml X 10	10ml X 10 X 20	1000000
123	732	Sodium Chloride Injection IP 0.9%w/v (Normal Saline (NS) 0.9% w/v)	100ml IV fluid plastic contain er	1's	1's x 20	50000

			using FFS/BF S technol			
124	735	Misoprostol 25mcg Tablets	4's	4's x10	(4's x10x10)x20	200000
125	736	Megeestrol Acetate Tablets IP 40mg	10's	10's x10	(10's x10x10)x20	200000
126	741	Cefpodoxime Proxetil dispersible tablet 50 mg	10's	10's x10	(10's x10x10)x20	200000
127	742	Cefaclor Tablet I.P 250 mg	10's	10's x10	(10's x10x10)x10	200000
128	746	Valganciclovir Hydrochloride USP 450 mg tablet	10's	10's x 10	(10's x10x10)x10	100000
129	750	HYOSCINE BUTYLBROMIDE INJECTION 20 mg/1 mL	1ml ampoul e	1's x 10	(1's x 10 x 10) x 10	100000
130	752	Clotrimazole 1%w/w 100 gm Dusting Powder	100 gm Powde r	1's x 10	(1's x 10) x 10	300000
131	753	Clotrimazole 1% w/w, Beclometasone Dipropionate 0.025% w/w 15 ml lotion	15 ml Lotion in Bottle	1's x 10	1's x 10 x 30	400000
132	755	Povidone-lodine 10% medicated paint	50 ML	1's x 10	(1's x 10) x 10	100000
133	757	Cefuroxime Injection IP 1500 mg	10 ml Vial & wfi	1's x 10	(1's x 10 x 10)x10	200000
134	762	Nortriptyline Tablet 25 mg Tablet	10's	10's x10	(10's x10x10)x20	500000
135	766	L-methylfolate calcium 7.5mg Tablet	10's	10'sx10	(10'sx10x10)x10	500000
136	772	Adenosine Injection IP 3mg/ml	2ml ampoul e	(1's x10)	(1'sx10x10)x10	100000
137	784	Amisulpride Tablets I.P 50mg	10's	10'sx10	(10'sx10x10)x20	200000

138	785	Amitriptyline hydrochloride 25mg Tablets I.P	15's	15'sx10	(15'sx10x10)x20	300000
139	786	Amitriptyline hydrochloride 10mg Tablets I.P	10's	10'sx10	(10'sx10x10)x20	400000
140	788	Anastrozole FC Tablets IP 1mg	10's	10'sx10	(10'sx10x10)x20	30000
141	793	Atenolol Tablets 25 mg	14's	14'sx10	(14'sx10x10)x10	1000000
142	796	Atorvastatin I.P 10mg, Aspirin I.P (EC) 75mg Capsules	10's	10'sx10	(10'sx10x10)x20	1000000
143	797	Atracurium Besilate Injection I.P 25mg/2.5ml	2.5ml	(2.5ml x 10)	(2.5mlx10x10)x1 0	100000
144	800	Bacitracin Zinc 250 lu Neomycin 5 Mg, Sulphacetamide Sodium 60 Mg Per 1gm Dusting Powder	10gm Powde r	1's x10	(1'sx10)x100	200000
145	804	Betamethasone Inj. I.P 4 mg/ml	1 ML ampoul e	(1's x10)	(1'sx10x10)x10	500000
146	809	Bortezomib Injection IP 3.5mg	Vial	1'sx10	(1'sx10)x100	10000
147	811	Bromfenac Sodium Ophthlamic solution IP 0.09%w/v	5ml	1'sx10	(1'sx 10)x100	200000
148	812	Bromocriptine Mesylate Tablets I.P 2.5mg	10's	10'sx10	(10'sx10x10)x20	200000
149	819	Capecitabine FC Tablets I.P 500mg	10's	10'sx10	(10'sx10x10)x10	10000
150	820	Carboprost Tromethamine Injection IP 250 mcg/ml	1ml ampoul e	(1's x10)	(1'sx10x10)x10	100000
151	822	Cefazolin Sodium Injection IP 500mg	Vial&W FI	1'sx10	(1'sx10)x100	100000
152	829	Chloramphenicol Eye Ointment IP 1%w/w	5 gm	1'sx10	(1'sx10x10)x20	100000
153	831	Chlorpromazine Tablets IP 50mg	10's	10'sx10	(10'sx10x10)x20	200000
154	832	Chlorthalidone Tablets IP 12.5mg	10's	10'sx10	(10'sx10x10)x20	200000
155	837	Cilnidipine Tablets 20mg	10's	10'sx10	(10'sx10x10)x20	400000

156	840	Citicoline Tablets IP 500mg	10's	10'sx10	(10'sx10x10)x10	300000
157	844	Clonazepam Tablets IP 1mg	10's	10'sx10	(10'sx10x10)x20	300000
158	860	Dextromethorphan HBr Syrup IP 13.5mg/5ml	50ml bottle	1's x10	(1's x10)x10	300000
159	865	Diacerein Capsules IP 50mg	10's	10'sx10	(10'sx10x10)x20	500000
160	868	Dicyclomine HCl (Dicycloverine) Injection IP 10mg/ml	2ml	(2ml x10)	(2mlx10x10)x10	300000
161	878	Drotaverine HCl 80mg, Mefenamic Acid 250mg Tablets	10's	10'sx10	(10'sx10x10)x20	300000
162	879	Drotaverine HCl FC Tablets IP 40mg	10's	10'sx10	(10'sx10x10)x20	400000
163	882	Efavirenz Tablets IP 600mg	30's	30's x 10	(30's x10)x25	50000
164	888	Febuxostat Tablets 40mg	10's	10'sx10	(10'sx10x10)x20	1200000
165	889	Febuxostat Tablets 80mg	10's	10'sx10	(10'sx10x10)x20	1200000
166	893	Filgrastim 300mcg/1ml Prefilled Syringe	1's	1 x 10	(1x10x10)x10	100000
167	901	Gabapentin Capsules IP 300mg	10's	10'sx10	(10'sx10x10)x20	300000
168	904	Glimepiride 1mg Metformin SR 500mg Tablets	10's	10'sx10	(10'sx10x10)x10	700000
169	905	Glimepiride 2mg, Metformin Hydrochloride 1g Tablets	10's	10'sx10	(10'sx10x10)x5	1500000
170	910	Human Chorionic Gonadotrophin 5000 IU Powder For Inj. With solvent	Vial and solvent	1'sx10	(1'sx10)x100	20000
171	911	Human Menopausal Gonadotrophin Injection 75 IU with solvent	Vial and solvent	1'sx10	(1'sx10)x100	20000
172	913	Hydrocortisone Sodium Succinate Injection IP 100mg	Vial & WFI	1x10	(1x10)x100	300000
173	916	Imatinib mesylate FC Tablets IP 400mg	10's	10'sx10	(10'sx10x10)x10	50000
174	923	Isosorbidemononitrate Tablets IP 20mg	10's	10'sx10	(10'sx10x10)x20	400000

175	926	Ketoconazole Cream 2%w/w	15gm Tube	1 x 25	(1 x 25) x 20	400000
176	935	Letrozole FC Tablets USP 2.5mg	10's	10'sx10	(10'sx10x10)x20	50000
177	936	Leuprolide Acetate Injections BP 3.75mg	1's	1's x 10	(1'sx10x10)x10	50000
178	937	Levetiracetam Syrup100 Mg/5ml	100ml	1's x10	(1's x10)x5	400000
179	938	Levocarnitine Injections 1gm	5ml vial	1's x 10	(1's x 10) x 100	200000
180	945	Levosulpiride Tablets 25mg	10's	10'sx10	(10'sx10x10)x20	400000
181	947	Lithium Carboinate PR Tablets IP 450mg	10's	10'sx10	(10'sx10x10)x10	400000
182	948	Lorazepam Tablets IP 1mg	10's	10'sx10	(10'sx10x10)x20	500000
183	949	Lorazepam Tablets IP 2mg	10's	10'sx10	(10'sx10x10)x20	500000
184	963	Methyldopa Tablets IP 250mg	10's	10'sx10	(10'sx10x10)x20	500000
185	973	Naloxone Injection I.P 400mcg	1ml ampoul e	(1's x10)	(1'sx10x10)x10	100000
186	975	Nebivolol 5 mg, Hydrochlorothiazide 12.5 mg Tab.	10's	10'sx10	(10'sx10x10)x20	700000
187	976	Nebivolol Tablets IP 2.5mg	10's	10'sx10	(10'sx10x10)x20	700000
188	978	Nepafenac Ophthalmic Solution 0.1% w/v	5ml	1 x 25	(1x25)x20	100000
189	987	Nitrofurantoin Tablets I.P 100mg	10's	10'sx10	(10'sx10x10)x20	400000
190	990	Olanzapine Tablets I.P 10mg	10's	10'sx10	(10'sx10x10)x20	300000
191	991	Olanzapine Tablets I.P 5mg	10's	10'sx10	(10'sx10x10)x20	300000
192	993	Ondansetron Oral Solution I.P 2 mg/5ml	30ml bottle	1's x10	(1's x10)x20	500000
193	994	Oxaliplatin Injections 50mg USP	Vial	1x10	(1x10)x100	50000
194	996	Oxcarbazepine Tablets I.P 300mg	10's	10'sx10	(10'sx10x10)x20	300000

195	997	Paclitaxel Injection IP 100mg (6mg/ml)	16.7ML Vial	1x10	(1x10)x100	30000
196	1003	Permethrin Cream 5% w/w	30gm tube	1's x10	(1's x10)x25	400000
197	1007	Phenylephrine Injection I.P 10mg/ml	1ml ampoul e	(1's x5)	(1'sx10x10)x10	200000
198	1011	Piracetam Syrup 500mg/5ml	100ml bottle	1's x10	(1's x10)x5	300000
199	1012	Piracetam Tablets 400mg	10's	10'sx10	(10'sx10x10)x10	300000
200	1013	Potassium Chloride 500mg/5ml Syrup	200ml bottle	1's x10	(1's x10)x5	200000
201	1032	Quetiapine Tablets I.P 100mg	10's	10'sx10	(10'sx10x10)x20	500000
202	1042	Risperidone Tablets 4mg	10's	10'sx10	(10'sx10x10)x20	500000
203	1050	Sertraline HCl Tablets I.P 100mg	10's	10'sx10	(10'sx10x10)x20	700000
204	1066	Spironolactone Tablets I.P 100mg	15's	15'sx10	(15'sx10x10)x10	100000
205	1067	Streptomycin Sulphate Injection I.P 1000mg	Vial	1'sx10	(1'sx10)x50	50000
206	1068	Sucralfate 1gm, Oxetacaine 10mg/10ml Suspension USP	100ml bottle	1's x10	(1's x10)x5	100000
207	1071	Tacrolimus Capsules 1mg	10's	10'sx10	(10'sx10x10)x20	500000
208	1072	Tamsulosin Modified-Release Capsules 0.4 mg	10's	10'sx10	(10'sx10x10)x20	1000000
209	1074	Telmisartan 80mg, Hydroclorthiazide 12.5mg Tablets	10's	10'sx10	(10'sx10x10)x20	1200000
210	1098	Voglibose 0.2mg, Metformin 500mg Tablets SR	10's	10'sx10	(10'sx10x10)x10	1200000
211	1099	Voglibose 0.3 mg, Metformin 500mg Tablets SR	10's	10'sx10	(10'sx10x10)x10	1200000
212	1105	Zolpidem Tablets I.P 10mg	10's	10'sx10	(10'sx10x10)x20	500000
213	1107	Pregabalin Capsules 75mg	10's	10'sx10	10'sx10x100	1000000

214	1109	Prazosin Tablets 2.5mg	30's	30'sx10	30'sx10x10x20	200000
215	1116	Ciprofloxacin Hydrochloride Sterile Ophthalmic ointment 0.3%	3gm tube	1'sx10	(1'sx10x10)x10	200000
216	1124	Fluvoxamine Maleate Tablets 100mg	10's	10'sx10	(10'sx10x10)x10	300000
217	1126	Mecobalamin 1500mcg, Alpha lipoic acid 100mg, Benfotiamine 100mg, Vitamin B6 3mg, Folic acid 1.5mg Capsules	10's	10'sx10	(10'sx10x10)x20	500000
218	1138	Desvenlafaxine Tablet 50mg	10's	10'sx10	(10'sx10x10)x10	200000
219	1142	Venlafaxine Capsule 75mg	10's	10'sx10	(10'sx10x10)x10	200000
220	1145	Venlafaxine Capsule 37.5mg	10's	10'sx10	(10'sx10x10)x10	200000
221	1151	Vitamin C Tablet 500mg	100's in a Bottle	1'sx10	(1'sx10x10)x10	500000

ANNEXURE -X(Ref:-Clause 7.1)

MODEL BANK GUARANTEE FORMAT FOR FURNISHING EMD

Whereas	(hereinafter called the
"tenderer") has submitted their offer supply	dated for the
BBPI/DRUG-054/2017 KNOW ALL	·
	having our registered office at are bound unto Bureau of Pharma Public Sector
Undertakings of India Gurgaon/Delhi(he lakh only for which payment will and tru	reinafter called the "Purchaser) in the sum of Rs. One ally to be made to the said Purchaser, the Bank binds a presents. Sealed with the Common Seal of the said
THE CONDITIONS OF THIS OBLIGATION A	RE:
(1) If the tenderer withdraws or amends within the period of validity of this tender	, impairs or derogates from the tender in any respect
(2) If the tenderer having been notified during the period of its validity:-	of the acceptance of his tender by the Purchaser
a) If the tenderer fails to furnish the contract.	the Performance Security for the due performance of
b) Fails or refuses to accept/execu	te the contract.
demand, without the Purchaser having demand the Purchaser will note that the	to the above amount upon receipt of its first written to substantiate its demand, provided that in its he amount claimed by it is due to it owing to the ions, specifying the occurred condition or conditions.
This guarantee will remain in force up should reach the Bank not later than the	to 30.06.2018 and any demand in respect thereof above date.
(Signature of the authorized officer of	the Bank)
Name and designation of the officer	
Seal, name & address of the Bank and add	dress of the Branch

Annexure – XI

{Ref:- clause 8.1(ii)}

(1)	(2)	(3)	(4)	(5)	(6)	(7)
Sr.	Drug			Shelf	Manufacturing	Manufacturi
No.	Code	Generic name of Drug	Unit Size	Life in	Capacity per	ng Batch Size
				months	year in Units	in Units
		Diclofenac Sodium +	4.01			
1	8	Serratiopeptidase (50mg +	10's			
		10mg) Tablets Diclofenac Sodium (SR)				
2	9	Tablets IP 100 mg	10's			
_	10	Diclofenac Sodium 25mg per	3 ml			
3	10	ml Inj. IP	ampoule			
4	14	Ibuprofen 400mg +	15's			
		Paracetamol 325mg Tablets				
5	15	Ibuprofen film coated Tablets IP 200mg	10's			
		Amikacin Sulphate 50mg/ml				
6	30	inj. IP	2ml Vial			
		Amoxycillin + Clavulanic acid	30 ml			
7	36	(200 mg+28.5 mg /5ml) Dry	bottles			
		Syrup	bottles			
	20	Amoxycillin + Clavulanic acid	CI.			
8	39	(500 mg + 125 mg) film coated Tablets IP	6's			
		Amoxycillin 125mg/ 5ml	60 ml			
9	43	Powder for Suspension IP	bottle			
10	44	Amoxycillin 250 mg Caps IP	10's			
11	45	Amoxycillin 500 mg Caps IP	10's			
12	47	Azithromycin (100mg/5ml)	15 ml			
	.,	Suspension IP	bottle			
13	49	Azithromycin 250 mg film coated Tablets IP	10's			
		Azithromycin 500 mg film				
14	50	coated Tablets IP	10's			
15	F4	Cefadroxil film coated	1010			
15	51	Tablets IP 250mg	10's			
16	55	Cefixime 200mg film coated	10's			
		Tablets IP				
17	E0	Cefotaxime Sodium &	Vial & wfi			
17	59	Sulbactam Sodium (1g + 500 mg) Inj.	Vidi & Wil			
		CEFOTAXIME SODIUM 250				
18	60	MG & SULBACTAM SODIUM	VIAL &			
		125 MG INJECTION	WFI			
		Cefotaxime Sodium &				
19	61	Sulbactam Sodium (500 mg	Vial & wfi			
20	62	+ 250 mg) Inj. Cefotaxime Sodium	Vial & wfi			
20	UZ	Cerotaxiiile Souluiii	viai & Wil			

		Injections IP 1000mg			
21	62	Cefotaxime Sodium	Vial 9 wfi		
21	63	Injections IP 250mg	Vial & wfi		
22	64	Cefotaxime Sodium	2ml Vial &		
22	04	Injections IP 500mg	wfi		
23	66	Cefpodoxime 200 mg film	10's		
		coated Tablets IP			
24	68	Ceftazadime 250 mg Inj.	Vial & wfi		
25	71	Ceftriaxone + Tazobactum 1000 mg + 125 mg Inj.	Vial & wfi		
26	84	Ciprofloxacin + Tinidazole (500 mg + 600 mg) film coated Tablets	10's		
27	85	Ciprofloxacin 250 mg film coated Tablets IP	10's		
28	88	Co-trimoxazole (Sulphamethoxazole 200mg + Trimethoprim 40mg / 5ml) Susp	50 ml bottle		
29	89	Co-trimoxazole Tablets IP (160 MG + 800 MG)	10's		
30	91	Co-trimoxazole Tablets IP (80 mg + 400 mg)	10's		
31	98	NORFLOXACIN 400 mg + TINIDAZOLE 600 mg FILM COATED Tablets	10's		
32	99	Norfloxacin 400 mg film coated Tablets	10's		
33	104	Roxithromycin (50 mg/ 5ml) Susp.	30 ml bottles		
34	106	Roxithromycin 300mg film coated Tablets	10's		
35	107	Tinidazole 300mg film coated Tablets	10's		
36	111	Benzyl Benzoate Application IP 25% w/v Lotion	100ml bottles		
37	119	Fluconazole 150 mg Tablets IP	10's		
38	124	Povidone Iodine 5% w/w Ointment USP	250 gm Jar		
39	126	Povidone Iodine 10 % Solution IP	500 ml bottles		
40	127	Povidone lodine 5% Solution 100ml	100ml bottle		
41	130	Chlorhexidine + Cetramide (1.5 % w/v + 3% w/v) Solution	100 ml bottles		
42	133	Glibenclamide 2.5 mg Tabs IP	10's		
43	141	Glipizide Tablets IP 5mg	10's	 	

	1	INSULIN INJECTION IP 40	İ		
44	142	IU/ml (Insulin Human	10 ml Vial		
1	172	Recombinant)	10 mm viai		
45	153	Cisplatin 10 mg Injection	Vial		
		Etoposide Inj. IP 100			
46	158	mg/5ml	Vial & wfi		
47	171	Mannitol 20% w/v Injection	350 ml		
48	172	METRONIDAZOLE INJECTION	100 ml		
	1/2	IP 5 mg / ml	100 1111		
49	176	Sterile Water for Injection	5ml Amp		
50	179	Albendazole 400mg Tabs IP	1's		
51	197	Lactulose Syrup 10 gm/ 15	100 ml		
	137	ml	bottle		
		Aluminium Hydroxide +			
52	198	Magnesium Hydroxide	170 ml		
		(250+250mg / 5ml)			
		Suspension			
53	200	Metoclopramide Injections	2ml vial		
		IP 5mg/ml			
54	216	RANITIDINE INJECTION IP 50	2 ML vial		
		MG/ 2ML Theophylline 25.3mg +			
55	244	Etophylline 84.7mg /2ml	2ml		
33	244	Injections	ampoule		
		Etophyllin +Theophylline (77			
56	245	mg + 23 mg) Tablets	10's		
		SALBUTAMOL Tablets IP 2			
57	256	MG	10's		
58	257	Salbutamol 2.5 mg Respule	2.5 ml		
	250	Salbutamol Syrup IP	100ml		
59	259	2mg/5ml	bottle		
60	260	Salbutamol Tablets IP 4mg	10's		
61	274	Dopamine HCl 200 mg/5ml	5 ml		
01	2/4	Inj. IP	ampoule		
62	279	Furosemide 40mg Tabs	10's		
63	280	Heparin Sodium 1000iu/ml	5 ml vial		
		Inj. IP			
64	290	Metoprolol 25 mg Tabs	10's		
		Telmisartan +			
65	298	Hydrochlorthiazide (40 mg +	10's		
		12.5 mg) Tabs			
66	300	Telmisartan 40 mg Tabs	10's		
67	305	Chloroquine Phosphate 250	10's		
		mg film coated Tablets			
66	242	Oral Rehydration Salts	415		
68	312	Citrate IP 20.5 gm (WHO	1's		
60	216	Formula) Sachet	10'0		
69	316	Betahistine 8 mg Tabs	10's		
70	318	Carbamazepine 200mg Tabs	10's		
		IP			

71	327	PHENYTOIN Tablets IP 100 mg	100's in Bottle		
72	331	Thyroxine Sodium 50 mcg Tabs	100's in A Bottle		
73	333	DEXAMETHASONE 0.5 MG TABS IP	10's		
74	364	GLIMEPIRIDE 2 MG + METFORMIN HYDROCHLORIDE 500 MG TABLETS SR	15's		
75	374	ARTEMETHER 80 MG + LUMEFANTRINE 480 MG SR TABLETS	6's		
76	377	CLINDAMYCIN CAPSULES IP 300 MG	10's		
77	382	LINEZOLID TABLETS IP 600 MG	10's		
78	392	GRISEOFULVIN TABLETS IP 250 MG IP	10's		
79	394	PYRANTEL PAMOATE ORAL SUSPENSION IP 250 MG/5 ml IP	10 ml bottle		
80	397	Oxytetracycline Cap I.P 250mg	10's		
81	407	IVERMECTIN TABLETS 12 MG	10's		
82	420	ATORVASTATIN 10 MG+ CLOPIDOGREL 75 MG CAPSULES	10's		
83	421	NEBIVOLOL TABLETS IP 5 MG	10's		
84	424	CARVEDILOL TABLETS IP 3.125 MG	10's		
85	438	INDAPAMIDE TABLETS IP 1.5 MG	10's		
86	452	WARFARIN TABLETS IP 5 MG	10's		
87	461	BETAMETHASONE VALERAT 0.1 % w/w + NEOMYCIN SULFATE 0.5 % w/w CREAM	20 gm tube		
88	466	URSODEOXYCHOLIC ACID Tablets IP 300 mg	10's		
89	472	DOMPERIDONE 30 MG+ ESOMEPRAZOLE 40 MG CAPSULE	10's		
90	476 480	LIQUID PARAFFIN 1.25 ML+ MILK OF MAGNESIA 3.75ML+ SODIUM PICOSULPHATE 3.33MG /5ML EMULSION LEVOSULPIRIDE (SR) 75 MG+	170 ml bottle		
			103		

		ESOMEPRAZOLE 40 MG (EC)				
		CAPSULES				
0.0		DICYCLOMINE 10 MG +	30 ml			
92	487	DIMETHICONE 40 MG /5 ML	bottle			
02	404	SUSPENSION	401-			
93	491	ITOPRIDE TABLET 50 MG	10's			
94	492	Sulfasalazine Tablets EC BP 500 MG	10's			
95	500	LEVO-THYROXINE TABLETS	100's in A			
	300	IP 100 MCG	Bottle			
96	506	LEVO-THYROXINE TABLETS	100's in			
		IP 50 MCG IP	Bottle			
		THIOCOLCHICOSIDE 4 MG+				
97	517	ACECLOFENAC 100 MG	10's			
		TABLETS				
		GUAIFENESIN 100 mg+	400			
98	531	TERBUTALINE 2.5 mg+	100 ml			
		BROMHEXINE 8 mg /10 ml SYRUP	bottle			
		DOXOFYLLINE TABLETS IP				
99	555	400 MG	10's			
		SALBUTAMOL 100 MCG +				
100	567	IPRATROPIUM 20 MCG	200 MDI			
100	307	/PUFF INHALER	200 10101			
		VITAMIN-D				
101	585	(CHOLECALCIFEROL) 60000	1 SACHET			
101	233	IU /1 GM SACHET	2 07 (0.12)			
		L-LYSINE + MULTIVITAMINS	200 ML			
102	592	(VIT-B1,B2,B3,B5,B6) SYRUP	bottle			
		Cetirizine Dihydrochloride IP				
		5mg, Phenylephrine				
103	603	Hydrochloride IP 10mg.,	10's			
		Paracetamol IP 325 mg				
		Tablets				
		Beclomethasone				
		Dipropionate 0.025% w/w,				
104	607	Neomycin Sulphate 0.5%	15gm			
101	007	w/w (3500 Unit /G)	tube			
		Chlorocresol 0.1% w/w				
		Cream				
46-	000	Betamethasone 0.05% w/w				
105	608	+ Salicylic acid 3% w/w	20 Gm			
		cream				
100	630	laxative Suspension Liqid	170ml			
106	630	Paraffin 3.75ml+Milk of	Bottle			
		Magnesia 11.25ml				
		Clobetasol Propionate BP0.05 % w/w, Neomycin				
107	634	Sulphate IP0.50 % w/w.,	20gm			
107	034	Miconazole Nitrate IP2.00	tube			
		% w/w,Chlorocresol IP(as				
		70 W/ W, CITIOTOCIESOTIT (ds	<u> </u>	<u> </u>		

		preservative) 0.10 % w/w			
		Cream/Ointment			
		Paracetamol 125mg+ CPM	60 ml		
108	643	1 mg + Sodium Citrate 60mg	Bottle		
		in a flavour syrup base			
109	647	Diclofenac 1.16%w/w+ Menthol 5%+ Oleum 3% +	20gm		
103	047	Methyl Salicylate 10% Gel	tube		
		Diclofenac Diethylamine BP			
		1.16 %, Linseed Oil BP 3 %			
		w/w, Methyl Salicylate IP 10			
110	648	% w/w, Menthol IP 5 %	35 gms.		
		w/w, Excipients and			
		Propellant q.s. to 100 % w/w Spray			
		Dicyclomine 10mg + Act.	10ml		
111	649	Dimethicone 40mg per ml	Bottle		
112	652	Dicyclomine 10mg +	10's		
112	032	Mefenamic 250 mg Tablets	10.5		
		Hydroquinone 2.0% w/w +			
113	657	Tretinoin 0.025% w/w + Mometasone Furoate 0.1%	15gm tube		
		w/w in a cream base q.s	tube		
		Diacerein 50 mg +			
114	671	Glucosamine Sulphate 500	10's		
		mg Tablet			
115	676	Triamcinolone Acetonide 0.1	10gm		
	070	% Mouth Ulcer gel	tube		
116	678	levodopa 250mg, Carbidopa	10's		
		25mg Tablets IP	200ml		
117	687	Lactulose 10gm/15 ml syrup	bottle		
118	691	Ofloxacin Ophthalmic	10 ml		
110	031	Solution IP 0.3% w/v	101111		
119	692	Olopattadine 0.1% w/v Eye	10 ml Vial		
120	701	Drops	10 mal \/iml		
120	701	Pilocarpine Eye drop IP 2 % Glibenclamide 5mg +	10 ml Vial		
121	713	MetforminHcl 500 mg tab	10's		
122	723	Water for Injection amp	10ml		
122	/23	polypack			
			100ml IV		
		Codium Chlorida Inication ID	fluid		
		Sodium Chloride Injection IP 0.9%w/v	plastic container		
123	732	(Normal Saline (NS) 0.9%	using		
		w/v)	FFS/BFS		
		,	technolog		
			у		
124	735	Misoprostol 25mcg Tablets	4's		
125	736	Megeestrol Acetate Tablets	10's		

		IP 40mg			
120	741	Cefpodoxime Proxetil	10's		
126	741	dispersible tablet 50 mg	10 \$		
127	742	Cefaclor Tablet I.P 250 mg	10's		
128	746	Valganciclovir Hydrochloride USP 450 mg tablet	10's		
129	750	HYOSCINE BUTYLBROMIDE INJECTION 20 mg/1 mL	1ml ampoule		
130	752	Clotrimazole 1%w/w 100 gm Dusting Powder	100 gm Powder		
131	753	Clotrimazole 1% w/w, Beclometasone Dipropionate 0.025% w/w 15 ml lotion	15 ml Lotion in Bottle		
132	755	Povidone-lodine 10% medicated paint	50 ML		
133	757	Cefuroxime Injection IP 1500 mg	10 ml Vial & wfi		
134	762	Nortriptyline Tablet 25 mg Tablet	10's		
135	766	L-methylfolate calcium 7.5mg Tablet	10's		
136	772	Adenosine Injection IP 3mg/ml	2ml ampoule		
137	784	Amisulpride Tablets I.P 50mg	10's		
138	785	Amitriptyline hydrochloride 25mg Tablets I.P	15's		
139	786	Amitriptyline hydrochloride 10mg Tablets I.P	10's		
140	788	Anastrozole FC Tablets IP 1mg	10's		
141	793	Atenolol Tablets 25 mg	14's		
142	796	Atorvastatin I.P 10mg, Aspirin I.P (EC) 75mg Capsules	10's		
143	797	Atracurium Besilate Injection I.P 25mg/2.5ml	2.5ml		
144	800	Bacitracin Zinc 250 lu Neomycin 5 Mg, Sulphacetamide Sodium 60 Mg Per 1gm Dusting Powder	10gm Powder		
145	804	Betamethasone Inj. I.P 4 mg/ml	1 ML ampoule		
146	809	Bortezomib Injection IP 3.5mg	Vial		
147	811	Bromfenac Sodium Ophthlamic solution IP 0.09%w/v	5ml		
148	812	Bromocriptine Mesylate	10's		

		Tablets I.P 2.5mg		
149	819	Capecitabine FC Tablets I.P	10's	
149	019	500mg		
150	820	Carboprost Tromethamine	1ml	
150	020	Injection IP 250 mcg/ml	ampoule	
151	822	Cefazolin Sodium Injection	Vial&WFI	
		IP 500mg		
152	829	Chloramphenicol Eye	5 gm	
		Ointment IP 1%w/w Chlorpromazine Tablets IP		
153	831	50mg	10's	
		Chlorthalidone Tablets IP		
154	832	12.5mg	10's	
155	837	Cilnidipine Tablets 20mg	10's	
156	840	Citicoline Tablets IP 500mg	10's	
157	844	Clonazepam Tablets IP 1mg	10's	
150	960	Dextromethorphan HBr	50ml	
158	860	Syrup IP 13.5mg/5ml	bottle	
159	865	Diacerein Capsules IP 50mg	10's	
		Dicyclomine HCl		
160	868	(Dicycloverine) Injection IP	2ml	
		10mg/ml		
4.54	070	Drotaverine HCl 80mg,	401	
161	878	Mefenamic Acid 250mg	10's	
		Tablets Drotaverine HCl FC Tablets		
162	879	IP 40mg	10's	
163	882	Efavirenz Tablets IP 600mg	30's	
164	888	Febuxostat Tablets 40mg	10's	
165	889	Febuxostat Tablets 80mg	10's	
		Filgrastim 300mcg/1ml		
166	893	Prefilled Syringe	1's	
167	001	Gabapentin Capsules IP	10's	
167	901	300mg	10 5	
168	904	Glimepiride 1mg Metformin	10's	
100	304	SR 500mg Tablets	103	
169	905	Glimepiride 2mg, Metformin	10's	
		Hydrochloride 1g Tablets		
170	010	Human Chorionic	Vial and	
170	910	Gonadotrophin 5000 IU Powder For Inj. With solvent	solvent	
		Human Menopausal		
171	911	Gonadotrophin Injection 75	Vial and	
1,1	311	IU with solvent	solvent	
472	042	Hydrocortisone Sodium	\/:- 0 \\/(5)	
172	913	Succinate Injection IP 100mg	Vial & WFI	
173	916	Imatinib mesylate FC Tablets	10's	
1/3	210	IP 400mg	10.3	
174	923	Isosorbidemononitrate	10's	
_,.		Tablets IP 20mg		

175	926	Ketoconazole Cream 2%w/w	15gm Tube	
176	935	Letrozole FC Tablets USP 2.5mg	10's	
177	936	Leuprolide Acetate Injections BP 3.75mg	1's	
178	937	Levetiracetam Syrup100 Mg/5ml	100ml	
179	938	Levocarnitine Injections 1gm	5ml vial	
180	945	Levosulpiride Tablets 25mg	10's	
181	947	Lithium Carboinate PR Tablets IP 450mg	10's	
182	948	Lorazepam Tablets IP 1mg	10's	
183	949	Lorazepam Tablets IP 2mg	10's	
184	963	Methyldopa Tablets IP 250mg	10's	
185	973	Naloxone Injection I.P 400mcg	1ml ampoule	
186	975	Nebivolol 5 mg, Hydrochlorothiazide 12.5 mg Tab.	10's	
187	976	Nebivolol Tablets IP 2.5mg	10's	
188	978	Nepafenac Ophthalmic Solution 0.1% w/v	5ml	
189	987	Nitrofurantoin Tablets I.P 100mg	10's	
190	990	Olanzapine Tablets I.P 10mg	10's	
191	991	Olanzapine Tablets I.P 5mg	10's	
192	993	Ondansetron Oral Solution I.P 2 mg/5ml	30ml bottle	
193	994	Oxaliplatin Injections 50mg USP	Vial	
194	996	Oxcarbazepine Tablets I.P 300mg	10's	
195	997	Paclitaxel Injection IP 100mg (6mg/ml)	16.7ML Vial	
196	1003	Permethrin Cream 5% w/w	30gm tube	
197	1007	Phenylephrine Injection I.P 10mg/ml	1ml ampoule	
198	1011	Piracetam Syrup 500mg/5ml	100ml bottle	
199	1012	Piracetam Tablets 400mg	10's	
200	1013	Potassium Chloride 500mg/5ml Syrup	200ml bottle	
201	1032	Quetiapine Tablets I.P 100mg	10's	
202	1042	Risperidone Tablets 4mg	10's	
203	1050	Sertraline HCl Tablets I.P 100mg	10's	

204	1066	Spironolactone Tablets I.P 100mg	15's		
205	1067	Streptomycin Sulphate Injection I.P 1000mg	Vial		
206	1068	Sucralfate 1gm, Oxetacaine 10mg/10ml Suspension USP	100ml bottle		
207	1071	Tacrolimus Capsules 1mg	10's		
208	1072	Tamsulosin Modified- Release Capsules 0.4 mg	10's		
209	1074	Telmisartan 80mg, Hydroclorthiazide 12.5mg Tablets	10's		
210	1098	Voglibose 0.2mg, Metformin 500mg Tablets SR	10's		
211	1099	Voglibose 0.3 mg, Metformin 500mg Tablets SR	10's		
212	1105	Zolpidem Tablets I.P 10mg	10's		
213	1107	Pregabalin Capsules 75mg	10's		
214	1109	Prazosin Tablets 2.5mg	30's		
215	1116	Ciprofloxacin Hydrochloride Sterile Ophthalmic ointment 0.3%	3gm tube		
216	1124	Fluvoxamine Maleate Tablets 100mg	10's		
217	1126	Mecobalamin 1500mcg, Alpha lipoic acid 100mg, Benfotiamine 100mg, Vitamin B6 3mg, Folic acid 1.5mg Capsules	10's		
218	1138	Desvenlafaxine Tablet 50mg	10's		
219	1142	Venlafaxine Capsule 75mg	10's		
220	1145	Venlafaxine Capsule 37.5mg	10's		
221	1151	Vitamin C Tablet 500mg	100's in a Bottle		

ANNEXURE -XII

Ref. Clause No.10.1

Performance Security Bank Guarantee

(unconditional)

To:	Bureau	of	Pharma	Public	Sector	Undertakings	of	India,	(Name	of	purchaser)	IDPL
Con	nplex, Ol	ld-D	Delhi-Gur	gaon Ro	oad, Dur	ndehera, Gurga	on ?	122016	(Haryar	na)		

Supplier) l	nerein called	"the Suppl	ier" has undertake oly of Drugs for the Contract".	n, in purs	uance of Te	ender BPF	-	
shall furni	sh you with	a Bank (een stipulated by your Guarantee for the formance obligation	sum spec	cified therei	n as sec	urity	
AN	D WHERE	AS we have	agreed to give the S	Supplier a (Guarantee			
behalf	of	the	affirm that we are Supplier,	upto	a	total		of
the Supplie sums with Words and	er to be in d in the limit	efault under of aforesaid, w	ertake to pay you, to the Contract and ithout your needing the therein.	without ca	avil or argur (Amount of	ment., any the Gua	y sum rantee	n or e in
This guara	ntee is valid ı	ıntil the	day of.		2019.			
				Signa	ture and Sea	l of Guara	antors	;
				Date			2	016
				Addı	ress			

ANNEXURE-XIII

Ref. Clause No.10.2

(NOTE:-In case Bid is submitted by the Marketer, the Agreement is be signed by Marketer as well Manufacturer)

AGREEMENT

THIS AGREEMENT made theday of2017 Between
Bureau of Pharma Public Sector Undertakings of India, IDPL Complex, Old-Delhi Gurgaon
Road, Dundahera, Gurgaon 122016 (Haryana)
(Name of purchaser) of (Country of Purchaser) (here in after "the Purchaser") of the one
part and(Name of Supplier) of(City and
Country of Supplier) (herein after called "the Supplier") of the other part :
WHEREAS the Purchaser is desirous that certain Goods and ancillary services viz;
Supply of Drugs in the tender Reference No. BPPI/DRUG-054/2017 (Brief Description of
Goods and Services) and has accepted a bid by the Supplier for the supply of those goods and

NOW THIS AGREEMENT WITNESSETH AS FOLLOWS:

(hereinafter called "the Contract Price").

1. In this Agreement words and expressions shall have the same meanings as are respectively assigned to them in the Conditions of Contract referred to, and they shall be deemed to form and be read and construed as part of this agreement.

services for the sum of(Contract Price in Words and Figures)

- 2. The following documents shall be deemed to form and be read and construed as part of this Agreement, viz
 - a. The Letter of Acceptance issued by the purchaser.
 - b. The Notice Inviting Tender
 - c. The supplier's bid including enclosures, annexures, etc.
 - d. The Terms and Conditions of the Contract
 - e. The Schedule of Requirement
 - f. The Technical Specification
 - g. Any other document listed in the supplier's bid and replies to queries, clarifications issued by the purchaser, such confirmations given by the bidder which are acceptable to the purchaser and the entire Addendum issued as forming part of the Contract.

- 3. In consideration of the payments to be made by the Purchaser to the Supplier as hereinafter mentioned, the Supplier hereby covenants with the Purchaser to provide, the goods and services and to remedy defects therein in conformity in all respects with the provisions of the Contract.
- 4. The purchaser hereby covenants to pay the Supplier in consideration of the provision of the goods and services and the remedying of defects therein, the Contract Price or such other sum as may become payable under the provisions of the Contract at the times and in the manner prescribed by the Contract.

Brief particulars of the goods and services which shall be supplied / provided by the Supplier are as under:

S.No.	Drug	Name of	UNIT	Tender Qty in	Unit Price (in	Rate of	Total value
	Code	Product	SIZE	Unit Size*	Rs.)exclusive	GST in %	inclusive of
					of GST		GST in Rs.

* Tender quantity indicated here is tentative and may vary subjected to various terms and conditions of the tender.

DELIVERY SCHEDULE

Supply shall all complete within 45th day from the date of issue of Ist purchase order and within 30th day from the date of issue of subsequent purchase order.

Dispute Resolution

Th	is agreement	shal	l be	deemed	l to	have	been	made/	executed	at	Del	hi	for	all	purp	ose.
----	--------------	------	------	--------	------	------	------	-------	----------	----	-----	----	-----	-----	------	------

Normally, there should not be any scope of dispute between the BPPI and the _____ after entering into a mutually agreed valid contract/agreement.

However, due to various unforeseen reasons, problems may arise during the progress of the								
contract/agreement leading to disagreement BPPI and the shall first try to resolv								
the same amicably by mutual Consultation. If the parties fail to resolve the dispute by sucl								
mutual consultation within twenty-one days, then, depending on the position of the case, eithe								
the BPPI or the shall give notice to other party of its intension to commence								
Arbitration procedure as per Indian Arbitration and Conciliation Act, 1996. Such								
disputes/differences shall be referred to Sole Arbitrator to be appointed by the President/ CEC								

of BPPI. The venue of Arbitration Shall be at New Delhi. The award published by the Arbitrator shall be full and final which shall be binding on both the parties.

Governing Law/Jurisdiction

BPPI/DRUG-054/2017

The applicable law governing this agreement shall be the laws of India and the court of Delhi shall have the exclusive jurisdiction to try any dispute arising out of the violation of any terms and condition of this agreement.

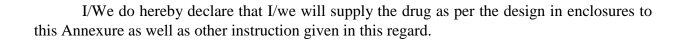
IN WITNESS where of the parties here to have caused this Agreement to be executed in accordance with their respective laws the day and year first above written.

Signed, Sealed and Delivered by the
Said(For the Purchaser)
Name –
Address- IDPL Complex, Old-Delhi-Gurgaon Road, Dundehera, Gurgaon 122016 (Haryana)
Designation -
In the presence of witness
Signature
Name
Address- IDPL Complex, Old-Delhi-Gurgaon Road, Dundehera, Gurgaon 122016 (Haryana)
Designation – Executive (Procurement)
Signed, Sealed and Delivered by the
Said(For the Supplier)
Name
Address
Designation
In the presence of witness
Signature
Name
Address Designation

ANNEXURE -XIV

Ref. Clause no 13

DECLARATION



Signature of the Tenderer

Name

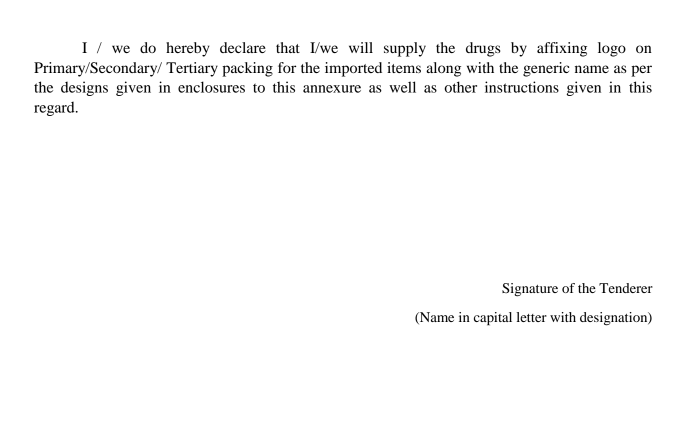
Designation

(Company Seal)

ANNEXURE -XIV(A)

Ref. Clause No. 13

UNDERTAKING



Enclosure–1 to <u>ANNEXURE</u> -XIV AND XIVA)

Ref. Clause No. 13

DESIGN FOR: Foil / blister of tablet and capsule

- 1. Text Matter Printing on Foil /Blister should be in minimum two colour i.e. Black & red. However, colour and design of PMBJP(Pradhan Mantri Bhartiya Janaushadhi Pariyojana) logogram in standard colour from at as per approval at the time of ART WORK approval before supply should be as given below.
- 2. PMBJP Logogram should be placed along with the address as given below
- 3. BPPI helpline number 1800 180 8080 should be printed
- 4. Font type should in CALIBIRI format for any type of title name of generic medicines
- 5. Title name of generic medicine should be **bold** in minimum 12 font size & the strength corresponding to it must be **bold** in minimum 14 font size and it may increase respectively according to size of label & the rest text matter should be in minimum 8 font size.
- 6. The stereo printing of batch no./manufacturing /expiry date & other details shouldn't overlap the text matter.
- 7. "Bureau of Pharma PSUs of India" should be running text only and should not be prominent.

Manufactured for:



Bureau of Pharma PSUs of India

IDPL Plant complex, Dundahera, Gurgaon 122016 (Haryana)

BPPI helpline number 1800 180 8080

Or

Manufactured for:



Bureau of Pharma PSUs of India

IDPL Plant complex, Dundahera, Gurgaon 122016 (Haryana)

BPPI helpline number 1800 180 8080

1. Pradhan Mantri Bharitya Janaushadhi Priyojana should be printed in Hindi at side of strips.

Enclosure – 2 to <u>ANNEXURE</u> –XIV & <u>ANNEXURE</u> –XIV(A)

Ref. Clause No. 13

1. Design for injection for primary packing

- a) Vial (5ml or more) should be supplied with the following PMBJP logogram **as per approval at the time of ART WORK approval before supply** as under:
- b) BPPI helpline number 1800 180 8080 should be printed
- c) Font type should in CALIBIRI format for any type of title name of generic medicines
- d) Title name of generic medicine should be minimum 12 font size and it may increase respectively according to size of label.
- e) "Bureau of Pharma PSUs of India" should be running text only and should not be prominent.



Manufactured for:

Bureau of Pharma PSUs of India

IDPL Plant complex, Dundahera, Gurgaon 122016 (Haryana)

BPPI helpline number 1800 180 8080

b) Ampoules or Vials less than 5 ml for primary packing

- (i) Injection in ampoule or vial (less than 5 ml) should be supplied with PMBJP logogram as per approval at the time of ART WORK approval before supply as under (colour should be black)
- (ii) BPPI helpline number 1800 180 8080 should be printed
- (iii) Font type should in CALIBIRI format for any type of title name of generic medicines
- (iv) Title name of generic medicine should be minimum 12 font size and it may increase respectively according to size of label.
- (v) "Bureau of Pharma PSUs of India" should be running text only and should not be prominent.

Manufactured for:



Bureau of Pharma PSUs of India

IDPL Plant complex, Dundahera, Gurgaon 122016 (Haryana)

BPPI helpline number 1800 180 8080

(vi) LIQUID:

- a) Liquid preparation should by supply with pilfer proof ROPP cap.
- b) Bottle cap should not bear the manufacturer's logogram
- c) Bottle label should bear PMBJP logogram as per approval at the time of ART WORK approval before supply as below:
- d) BPPI helpline number 1800 180 8080 should be printed
- e) "Bureau of Pharma PSUs of India" should be running text only and should not be prominent
- f) Font type should in CALIBIRI format for any type of title name of generic medicines
- g) Title name of generic medicine should be minimum 12 font size and it may increase respectively according to size of label.



Manufactured for:

Bureau of Pharma PSUs of India

IDPL Plant complex, Dundahera, Gurgaon 122016 (Haryana)

BPPI helpline number 1800 180 8080

3. OINTMENTS / CREAMS

a) Ointment / Cream /Gel /Glass Jar should bear JANASHADHI or PMJAY logogram as per approval at the time of ART WORK approval before supply as below:

Manufactured for:



Bureau of Pharma PSUs of India

IDPL Plant complex, Dundahera, Gurgaon 122016 (Haryana)

BPPI helpline number 1800 180 8080

- b) BPPI helpline number 1800 180 8080 should be printed
- c) Ointment / cream tube should be packed in mono carton (secondary packing) with PMBJP logogram as per approval at the time of ART WORK approval before supply as given below.
- d) "Bureau of Pharma PSUs of India" should be running text only and should not be prominent
- e) Font type should in CALIBIRI format for any type of title name of generic medicines
- f) Title name of generic medicine should be minimum 12 font size and it may increase respectively according to size of label.

Enclosure 3 to <u>ANNEXURE</u> –XIV(A)

SPECIMEN LABEL FOR MONO CARTON (Secondary Packing)

Rx 10 X 10's Tablets

Generic Name of Product



Manufactured for:

Bureau of Pharma PSUs of India

IDPL Plant complex, Dundahera, Gurgaon 122016 (Haryana)

BPPI helpline number 1800 180 8080

For Ampoules/vials :- All secondary packing box/carton should be supplied with printed text matter as per guidelines.

Note: An additional to statuary requirement under Drug & Cosmetic Act 1940 and rules 1945

ANNEXURE-XV

Ref. Clause No.14.1

SCHEDULE FOR PACKAGING OF DRUGS

GENERAL SPECIFICATIONS

- 1. Strips of Aluminum foils should be gauge 04.
- 2. Aluminum foils s back material for blisters should be gauge 025.
- 3. The rigid PVC used in blister packing should be of not less than 250 micron
- 4. All glass bottles should be new neutral glass. Pet bottles so accepted as per drug laws stipulation.
- 5. Ointments should be packed in lacquerized Aluminum Tubes or Lami tubes.
- 6. Small Tablets packed in blisters should be packed to facilitate easy removal of the tablet without breaking / crushing.
- 7. Specification of outer cartons are as given in this Schedule.
- 8. In case of any conflict between Carton specifications and packets per carton specification the specification of the packets / carton shall prevail.
- 9. All plastic containers should be made of virgin grade plastics
- 10. Injection in vials should have a flip-off seals.
- 11. The strips shall be aluminum strip / blisters with aluminum foil back.
- 12. The minimum diameters of each tablets should be of 6.4mm
- 13. The outer carton/secondary packaging should be of pearl white duplex board (off white/grey is not acceptable) with a minimum of 350 GSM with Gloss laminated packing for the strips, blisters, ointments, creams etc. and for ampoules and vials should be with pearl white board of 350 GSM(off white/grey is not acceptable). The material to be used for carton should be from virgin chemical pulp.
- 14. All liquid oral preparations to be provided with a measuring plastic cup, fitted over the cap of the bottle.
- 15. All primary/secondary/tertiary packaging should have JA/PMJAY logo.
- 16. Two Horizontal/vertical/standing lines in two different colour will be there on Primary and secondary packaging, so as to differentiate therapy groups. The colours of lines will be intimated during Artwork approval.

17. The primary packing should be decided by the party depending on the drug category as per D&C act. For e.g if drug is hygroscopic then tablet should be packed in Alu/Alu blister or if it is light sensitive then to be packed in Amber colour PVC e.t.c.

(Schedule)

1.	CORRUGATED BOXES(Liquid)						
	1. No corrugate package should weigh more than 15 kgs (i.e. product + inner						
	carton + corrugated box).						
	2. All Corrugated boxes should be of 'A' grade paper i.e. Virgin and 7 Ply.						
	3. All items should be packed only in first hand boxes only.						
2.	FLUTE						
	The corrugated boxes should be of narrow flute.						
3.	JOINT						
	Every box should be preferably single joint and not more than two joints.						
4.	STITCHING						
	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 joined using calico at the						
	corners.						
5.	FLAP						
	The flaps should uniformly meet but should not over lap each other. The flap when turned						
	by 45 - 60° should not crack.						
6.	TAPE						
	Every box should be sealed with gum tape running along the top and lower opening.						
7.	CARRYSTRAP:						
	Every box should be strapped with two parallel nylon carry straps (they should intersect).						
8.	LABEL						
	The product label on the carton should be large at least 15 cms x 10 cms dimension.						
	It should carry the correct technical name, strength of the product, date of						
	manufacturing, date of expiry, quantity packed and net weight of the box.						
9.	OTHERS						
	No box should contain mixed products or mixed batches of the same product.						

II. SPECIFICATION OF CORRUGATED BOXES HOLDING TABLETS / CAPSULES / PESSARIES

- (1) The box should not weigh more than 7-8 kgs. The grammage of outer box should be 150 gsm and inside partition / lining should be 120 gsm.
- (2) The box should be of 7 ply with bursting strength of 9 Kg / Cm2

III. SPECIFICATIONS OF CORRUGATED BOXES FOR OINTMENT / CREAM / GELS PACKED IN TUBES:

- (1) No corrugate box should weigh more than 7-8 Kgs.
- (2) Every Ointment tube should be individually packed in carton and then packed in 20's in a white board box, which may be packed in a corrugated box.
- (3) Grammage: Outer box should be 150 gsm inside partition /

Lining should be 120gsm.

IV. SPECIFICATIONS OF CORRUGATED BOXESFOR INJECTABLE (IN VIALS AND AMPOULES)

- (1) Vials may be packed in corrugated boxes weighing up to 15 Kgs. Ampoules should be packed in C.B weighing not more than 8 kgs.
- (2) C.B. for vials should be of 150 gsm (outer box should be 150 gsm and inside partition / lining should be 120 gsm) and 7 ply, while C.B. for ampoules should be of 150 gsm (outer box should be 150 gsm and inside partition / lining should be 120 gsm) and 7 ply.
- (3) Bursting strength for CB boxes for

i. Vials : Note less than 13 Kg/Cm2

ii. Amp : Note less than 9 Kg/Cm2

- (4) In the case of 10 ml Ampoules 100 or 50 ampoules may be packed in a grey board box. Multiples of white 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 vial is 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 centre pad.
- (6) In case of ampoules every white board box should carry 5 amps. Cutters placed in a polythene bag.
- (7) Vials of eye, ear drops and nasal drops should be packed in an individual carton with a dispensing device. If the vial is of FFS/BFS technology, they should be packed in 50's in a white board box.

ANNEXURE -XVI

MANDATE FORM

Ref. clause 16.2

Sl.No.	Details Required	
1.	Company Name	
	PAN Number	
	TIN Number	
	Date of Inception	
	Licence No. & Date	
	Issued By	
	Valid Upto	
2.	Postal Address of the Company	
	Telephone No.	
	Fax No.	
	E-mail ID	
	Alternate E-mail ID	
3.	Name of the Managing Director /	
	Director / Manager	
	Mobile No. / Phone No	
	E-mail ID	
4.	Name and Designation of the	Name:
	authorized company official	Designation:
	Mobile No.	
	E-mail ID	
5.	Bank Details	
	a) Name of the Bank	
	b) Branch Name & address	
	c) Branch Code No.	
	d) Branch Manager Mobile No.	
	e) Branch Telephone no	
	f) Branch E-mail ID	
	g) 9 digit MICR code number of	
	the bank and branch appearing on the MICR	
	appearing on the MICR cheque issued by the bank	
	h) IFSC Code of the Branch	
	i) Type of Account (Current /	
	Savings)	
	j) Account Number (as appear	
	in cheque book)	

(In lieu of the bank certificate to be obtained, please <u>upload the original cancelled cheque</u> issued by your bank for verification of the above particulars).					
transaction is delayed would not hold Bure have read the cond	d or not effected at all cau of Pharma Public itions of the tender	culars given above are correct and complete. If the I the reasons of incomplete or incorrect information, I Sector Undertakings of India (BPPI) responsible. I / agreement entered and agree to discharge the ompany as a tenderer / successful tenderer.			
Date:	Company Seal	Signature			
Place:		(Name of the person signing & designation)			
CERTIFIED THAT CORRECT AS PER		S FURNISHED ABOVE BY THE COMPANY ARE			
Bank Seal with addre	ess:	Signature of the authorized official of the bank			