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Ministry of Chemicals and Fertilizers (Department of Pharmaceuticals)

New Delhi

Dated 15th May 2013

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ORDER

S.O. 1221(E).– In exercise of the powers conferred by section 3 of the Essential Commodities Act, 1955, (10 of 1955), and supersession of the Drug (Prices Control) Order, 1995, except as respect to things done or omitted to be done before such supersession, the Central Government hereby makes the following Order, namely:-

1. Short title and commencement.- (1) This Order may be called the Drugs (Prices Control) Order, 2013.

(2) It shall come into force on the date of its publication in the Official Gazette.

2. Definitions.- (1) In this Order, unless the context otherwise requires,-

(a) "Act" means the Essential Commodity Act, 1955 (10 of 1955);

(b) "active pharmaceutical ingredients or bulk drug" means any pharmaceutical, chemical, biological or plant product including its salts, esters, isomers, analogues and derivatives, conforming to standards specified in the Drugs and Cosmetics Act, 1940 (23 of 1940) and which is used as such or as an ingredient in any formulation;

(c) **"brand"** means a name, term, design, symbol, trademark or any other feature that identifies one seller's drug as distinct from those of other sellers;

(d) "**ceiling price**" means a price fixed by the Government for Scheduled formulations in accordance with the provisions of this Order;

(e) "**dealer**" means a person carrying on the business of purchase or sale of drugs, whether as a wholesaler or retailer and includes his agent;

(f) "**distributor**" means a person engaged in the work of distribution of drugs and includes an agent or a stockist for stocking drugs for sale to a dealer;

(g) **"existing manufacturer"** means manufacturer existing on the date of publication of this order in the Official Gazette.

(h) "Form" means a form specified in the Second Schedule;

(i) "**formulation**" means a medicine processed out of or containing one or more drugs with or without use of any pharmaceutical aids, for internal or external use for or in the diagnosis, treatment, mitigation or prevention of disease and, but shall not include –

(i) any medicine included in any bonafide Ayurvedic (including Sidha) or Unani (Tibb) systems of medicines;

(ii) any medicine included in the Homeopathic system of medicine; and

(iii) any substance to which the provisions of the Drugs and Cosmetics Act, 1940 (23 of 1940) do not apply;

(j) **"generic version of a medicine"** means a formulation sold in pharmacopeial name or the name of the active pharmaceutical ingredient contained in the formulation, without any brand name;

(k) "Government" means the Central Government;

(I) "**import**" with its grammatical variations and cognate expressions means bringing a drug into India from a place outside India for its sale;

(m) **"local taxes"** means any tax or levy (except excise or import duty included in retail price) paid or payable to the Government or the State Government or any local body under any law for the time being in force by the manufacturer or his agent or dealer;

(n) "**manufacturer**" for the purpose of this Order means any person who manufactures, imports and markets drugs for distribution or sale in the country;

(o) **"market share"** means the ratio of domestic sales value (on the basis of moving annual turnover) of a brand or a generic version of a medicine and the sum of total domestic sales value of the all brands and generic versions of that medicine sold in the domestic market having same strength and dosage form;

(p) "margin to retailer" for the purposes of this Order shall mean a percentage of price to retailer;

(q) **"market based data"** means the data of sales related to a drug collected or obtained by the Government as deemed fit, from time to time;

(r) **"maximum retail price"** means the ceiling price or the retail price plus local taxes and duties as applicable, at which the drug shall be sold to the ultimate consumer and where such price is mentioned on the pack;

(s) **"moving annual turnover"** in a particular month means cumulative sales value for twelve months in domestic market, where the sales value of that month is added and the corresponding sales of the same month in the previous year are subtracted;

(t) **"National List of Essential Medicines"** means National List of Essential Medicines, 2011 published by the Ministry of Health and Family Welfare as updated or revised from time to

time and included in the first schedule of this order by the Government through a notification in the Official Gazette;

(u) "**new drug**" for the purposes of this Order shall mean a formulation launched by an existing manufacturer of a drug of specified dosages and strengths as listed in the National List of Essential Medicines by combining the drug with another drug either listed or not listed in the National List of Essential Medicines or a formulation launched by changing the strength or dosages or both of the same drug of specified dosages and strengths as listed in the National List of Essential Medicines.

(v) "non-scheduled formulation" means a formulation, the dosage and strengths of which are not specified in the First Schedule;

(w) **"pharmacoeconomics"** means a scientific discipline that compares the therapeutic value of one pharmaceutical drug or drug therapy to another;

(x) "**price list**" means a price list referred to in paragraphs 24 and 25 and includes a supplementary price list;

(y) **"price to retailer"** means the price of a drug at which it is sold to a retailer which includes duties and does not include local taxes;

(z) "retail price" means the price fixed by the Government for a new drug under paragraph5 ;

(za) "retailer" means a dealer carrying on the retail business of sale of drugs to customers;

(zb) **"scheduled formulation**" means any formulation, included in the First Schedule whether referred to by generic versions or brand name;

(zc) "schedule" means a Schedule appended to this Order;

(zd) "wholesaler" means a dealer or his agent or a stockist engaged in the sale of drugs to a retailer, hospital, dispensary, medical, educational or research institution or any other agency;

(ze) **"wholesale price index"** means annual wholesale price index of all commodities as announced by the Department of Industrial Policy and Promotion, Government of India, from time to time.

(2) All other words and expressions used herein and not defined but defined in the Act or the Drugs and Cosmetics Act, 1940 (23 of 1940) shall have the meanings respectively assigned to them in the said Acts.

.3. Directions to manufacturers of active pharmaceutical ingredients or bulk drugs or formulations.- The Government may, -

(i) with a view to achieve adequate availability and to regulate the distribution of drugs, in case of emergency or in circumstances of urgency or in case of non-commercial use in public interest, direct any manufacturer of any active pharmaceutical ingredient or bulk drug or formulation to increase the production and to sell such active pharmaceutical ingredient or bulk drug to such other manufacturer(s) of formulations and to direct formulators to sell the formulations to institutions, hospitals or any agency as the case may be;

(ii) for the purpose of giving any direction under sub-paragraph (i), call for such information from manufacturers of active pharmaceutical ingredients or bulk drugs or formulations, as it may consider necessary and such manufacturer shall furnish the required information within such time the Government may fix.

4. Calculation of ceiling price of a scheduled formulation.– (1) The ceiling price of a scheduled formulation of specified strengths and dosages as specified under the first schedule shall be calculated as under:

Step1. First the Average Price to Retailer of the scheduled formulation i.e. P(s) shall be calculated as below:

Average Price to Retailer, P(s) = (Sum of prices to retailer of all the brands and generic versions of the medicine having market share more than or equal to one percent of the total market turnover on the basis of moving annual turnover of that medicine) / (Total number of such brands and generic versions of the medicine having market share more than or equal to one percent of total market turnover on the basis of moving annual turnover.)

Step2. Thereafter, the ceiling price of the scheduled formulation i.e. P(c) shall be calculated as below:

P(c) = P(s).(1+M/100), where

- P(s) = Average Price to Retailer for the same strength and dosage of the medicine as calculated in step1 above.
- M = % Margin to retailer and its value =16

(2) The ceiling price calculated as per sub-paragraph (1) and notified by the Government shall be applicable to scheduled imported formulations also.

5. Calculation of retail price of a new drug for existing manufacturers of scheduled formulations.– (1) The retail price of the new drug available in domestic market shall be calculated as provided in sub-paragraph (1) of paragraph 4.

(2) (i) the price to retailer of a new drug, not available in domestic market, shall be fixed by the Government on the principles of "Pharmacoeconomics" of the new drug, on the recommendation of a Standing Committee of Experts formed under paragraph 15.

(ii) the retail price of such new drug shall be fixed by adding sixteen percent margin to retailer on the price to retailer as fixed in item (i):

6. Ceiling price of a scheduled formulation in case of no reduction in price due to absence of **competition.**– (1) where the average price to retailer of a scheduled formulation, arrived at as per the formula specified in sub-paragraph (1) of paragraph 4, has the effect of,-

(a) no reduction in average price to retailer with respect to the prices to retailer of the schedule formulation; and

(b) there are less than five manufacturers for that formulation having one percent or more market share,

the ceiling price shall be calculated as under:-

(i) in the event of other strengths or dosage forms of the same scheduled formulation is available in the list of scheduled formulation, the average price to retailer shall be calculated as under:

Step1: First the Average Price to Retailer of such scheduled formulation i.e. P(s) shall be calculated as under:

 $P(s) = P_m \{1 - (P_{i1} + P_{i2} + ...)/(N*100)\}$ Where,

 \mathbf{P}_{m} = Price to Retailer of highest priced scheduled formulation under consideration.

 $P_i = \%$ reduction in Average Price to Retailer of other strengths and dosage forms (calculated as in step1 of sub-paragraph (1) of paragraph 4) in the list of schedule formulations w.r.t the highest priced formulation taken for calculating the average price to retailer of such strengths and dosage forms.

N = Number of such other strengths or dosage forms or both in the list of schedule formulations

Step2. Thereafter, the ceiling price of the scheduled formulation i.e. P(c) shall be calculated as under:

P(c) = P(s).(1+M/100), where

P(s) = Average Price to Retailer of the scheduled formulation as calculated in step1 hereinabove and

M = % Margin to retailer and its value=16

(ii) in the event of other strengths or dosage forms of the scheduled formulation is not available in the schedule but there are other scheduled formulations in same subtherapeutic category as that of the scheduled formulation, then the Ceiling Price shall be calculated as under:

Step1: First the Average Price to Retailer of such scheduled formulation i.e. P(s) shall be calculated as under:

P(s) = P_m{1-(P_{i1}+P_{i2}+...)/(N*100)}, Where,

 P_m = Price of highest priced formulation taken for calculating the average price to retailer of the formulation under consideration..

 $P_i = \%$ reduction in Average Price to Retailer of other schedule formulations (calculated as in step1 of sub-paragraph (1) of paragraph 4) in same sub-therapeutic category as that of the scheduled formulation under consideration w.r.t the highest priced formulation taken for calculating the average price to retailer.

N = Number of such other schedule formulations in same sub-therapeutic category as that of the scheduled formulation under consideration.

Step2. Thereafter, the ceiling price of the scheduled formulation i.e. P(c) shall be calculated as under:

P(c) = P(s)*(1+M/100), where

P(s) = Average Price to Retailer of the scheduled formulation as calculated in step1 above and

M = % Margin to retailer and its value=16

Explanation.- where the scheduled formulation under consideration is coming under more than one sub-therapeutic category, the Average Price to Retailer of the scheduled formulation shall be calculated after taking into consideration the percentage reduction in Average Price to Retailer of other schedule formulations under all such sub-therapeutic categories and the lowest average price to retailer shall be taken for calculating the ceiling price of the scheduled formulation under consideration; (iii) in case the other strengths or dosage forms of the scheduled formulation are not available in the schedule and there is no sub therapeutic category of the scheduled under consideration, the ceiling price shall be calculated as under:

Step1: First the Average Price to Retailer of such scheduled formulation i.e. P(s) shall be calculated as under:

 $P(s) = P_m \{1 - (P_{i1} + P_{i2} + ...)/(N*100)\}$ Where,

 \mathbf{P}_{m} = Price of highest priced formulation taken for calculating the average price to retailer of the formulation under consideration.

 $P_i = \%$ reduction in Average Price to Retailer of other schedule formulations (calculated as in step1 sub-paragraph (1) of paragraph 4) in same therapeutic category as that of the scheduled formulation under consideration w.r.t the highest priced formulation taken for calculating the average price to retailer.

N = Number of such other schedule formulations in same therapeutic category as that of the scheduled formulation under consideration.

Step2. Thereafter, the ceiling price of the scheduled formulation i.e. P(c) shall be calculated as under:

P(c) = P(s).(1+M/100), where

P(s) = Average Price to Retailer of the scheduled formulation as calculated in step1 above and

M = % Margin to retailer and its value=16

Explanation.- where the scheduled formulation under consideration is coming under more than one therapeutic category, the Average Price to Retailer of the scheduled formulation shall be calculated after taking into consideration the percentage reduction in Average Price to Retailer of other schedule formulations under all such therapeutic categories and the lowest average price to retailer shall be taken for calculating the ceiling price of the scheduled formulation under consideration. (2) Notwithstanding anything contained in this paragraph, where the price has been fixed and notified by the Government under the Drugs (Prices Control) Order, 1995 the provisions of sub-paragraph (1) shall not apply.

7. Margin to retailer. – While fixing a ceiling price of scheduled formulations and retail prices of new drugs, sixteen percent of price to retailer as a margin to retailer shall be allowed.

8. Maximum retail price.– (1) The maximum retail price of scheduled formulations shall be fixed by the manufacturers on the basis of ceiling price notified by the Government plus local taxes wherever applicable, as under:

Maximum Retail Price = Ceiling price + Local Taxes as applicable

(2) The maximum retail price of a new drug shall be fixed by the manufacturers on the basis of retail price determined by the Government plus local taxes wherever applicable, as under:

Maximum Retail Price = Retail Price + Local Taxes as applicable

9. Reference data and source of market based data.– (1) Initially, the source of market based data shall be the data available with the pharmaceuticals market data specializing company – IMS Health (IMS) and if the Government deems necessary, it may validate such data by appropriate survey or evaluation.

(2) The Government may in the due course of time come out with other appropriate mechanism of collecting or obtaining the market based data related to drugs and the decision of Government with respect to collection or obtaining of data shall be final.

(3) The market based data, for fixing the ceiling price of scheduled formulations for the first time after the notification of this order, shall be the data of May, 2012.

(4) The market based data for fixing the retail price of new drugs available in the market, shall be the data available for the month ending immediately before six months of receipt of application for fixing the price of the new drug.

(5) The market based data for fixing the ceiling price of a scheduled formulation due to a revision in the first schedule shall be the data available for the month ending immediately before six month of notification of revision in the first schedule.

(6) Notwithstanding anything contained in this order, the reference date for the formulations which are part of the Drugs (Prices Control) Order, 1995 shall be as per the provisions of paragraph 10 of this Order.

10. Pricing of the formulations covered under Drugs (Prices Control) Order, 1995.– (1) The prices of scheduled formulations, which are also specified in the First Schedule to the Drugs (Prices Control) Order, 1995, fixed and notified under the provisions of the said order, up to 31st May, 2012, shall remain effective for further one year i.e. up to 30th May' 2013 and the manufacturers may revise the prices of such scheduled formulations as per the annual wholesale price index for the previous calendar year announced by Department of Industrial Promotion and Policy and thereafter the formula as in sub- paragraph (1) of paragraph 4 of this Order shall be applied for fixing the ceiling prices of such formulations.

(2) The prices of scheduled formulations, which are also specified in the First Schedule to the Drugs (Prices Control) Order, 1995, fixed and notified under the provisions of Drugs (Prices Control) Order, 1995 after 31st May, 2012, shall remain effective for one year from the date of notification of such prices under Drugs (Prices Control) Order, 1995 and immediately thereafter the manufacturers may revise the prices as per the annual wholesale price index for the previous calendar year announced by Department of Industrial Promotion and Policy and on the 1st April of succeeding financial year, the formula as in sub-paragraph (1) of paragraph 4 of this Order shall be applied for fixing the ceiling prices of such schedule formulations.

(3) The prices of scheduled formulations, which are specified in the Drugs (Prices Control) Order, 1995 but not specified in the First Schedule of this order, fixed and notified under the provisions of the said order, up to 31st May,2012, shall remain effective for further one year i.e. up to the 30thMay'2013 and thereafter prices of such formulations shall be regulated as in case of other non-scheduled formulations as stated in paragraph 20 of this Order.

(4) The prices of scheduled formulations, which are specified in the Drugs (Prices Control) Order, 1995 but not specified in the First Schedule of this order, fixed and notified under the provisions of the said order, after 31st May,2012, shall remain effective for one year from the date of notification of such prices and thereafter prices of such formulations shall be regulated as in case of other non-scheduled formulations as stated in paragraph 20 of this Order.

11. Ceiling price or retail price of a pack.– (1) The average price to retailer calculated as per the provisions in paragraphs 4, 5 and 6 shall be on the dosage basis, (per tablet, per capsule or injection in volume as listed in first schedule) and the ceiling price or retail price of a pack shall be reached by multiplying the same with the number or quantity in the pack as the case may be.

(2) In the event of the unit of the dosage for a scheduled formulation not available in the first schedule, the lowest pack size for that category of medicine, as specified in the Drugs and Cosmetics Act, 1940 (23 of 1940) and the rules thereunder, shall be taken as unit dosage for calculating the ceiling price or retail price as the case may be, for that scheduled formulation and this shall be applicable while calculating the per unit price of even non-scheduled medicines for arriving at the retail price in case of paragraph 5.

12. Price of formulations (branded or generic version) listed in the National List of Essential Medicines, launched by a manufacturer.– (1) A manufacturer, launching a scheduled formulation, shall be free to fix the price of the scheduled formulation equal to or below the ceiling price fixed for that schedule formulation by the Government.

(2) Where an existing brand is re-launched by another manufacturer the provisions of paragraph 13 shall be applicable.

13. Price of scheduled formulations for the existing manufacturers.– (1) All the existing manufactures of scheduled formulations, selling the branded or generic or both the versions of scheduled formulations at a price higher than the ceiling price (plus local taxes as applicable) so fixed and notified by the Government, shall revise the prices of all such formulations downward not exceeding the ceiling price (plus local taxes as applicable):

Provided, that in case of scheduled formulations produced or available in the market before the date of notification of ceiling price, the manufacturers shall ensure within a period of forty-five days of the date of such notification that the maximum retail price of such scheduled formulation does not exceed the ceiling price (plus local taxes as applicable).

(2) All the existing manufactures of scheduled formulations, selling the branded or generic or both the versions of scheduled formulations at a price lower than the ceiling price (plus local taxes as applicable) so fixed and notified by the Government shall maintain their existing maximum retail price.

(3) Annual increase in maximum retail price may be carried out as per the increase in the wholesale price index with respect to previous year as per the provision of sub-paragraphs (2) and (3) of paragraph 16.

Provided that in case of decline in wholesale price index, a corresponding reduction in the prices shall be made as per the provision of sub-paragraph (4) of paragraph 16.

14. Fixation of ceiling price of scheduled formulations.– (1) The Government shall fix and notify the ceiling prices of the scheduled formulations in accordance with the provisions of the paragraphs 4 and 6, as the case may be, and no manufacturer shall sell the scheduled

formulations at a price higher than the ceiling price (plus local taxes as applicable) so fixed and notified by the Government.

(2) Where any manufacturer sells a scheduled formulation at a price higher than the ceiling price (plus local taxes as applicable) fixed and notified by the Government, such manufacturers shall be liable to deposit the overcharged amount along with interest thereon from the date of such overcharging.

15. Fixation of retail price of a new drug for existing manufacturers of scheduled formulations.– (1) The Government shall form a Standing Committee of such Experts, as it may deem fit, within sixty days of notification of this order with a view to recommend the retail prices of new drugs on the principles of "Pharmacoeconomics".

(2) Where an existing manufacturer of a drug with dosages and strengths as specified in National List of Essential Medicines launches a new drug, such existing manufacturers shall apply for prior price approval of such new drug from the Government in Form-I specified under Schedule-II of this Order.

(3) On receipt of the application under sub-paragraph (2), in the event of the new drug available in domestic market, the Government shall fix the retail price of the new drug in accordance with the provision of sub-paragraph(1) of paragraph 5 and in the event of the new drug not available in domestic market, the Government shall forward the same to the Standing Committee of Experts who shall examine the application on the principles of "Pharmacoeconomics" and make recommendations of retail price of the new drug to the Government within thirty days of the receipt of application.

(4) The Government shall, on receipt of recommendation under sub-paragraph (3), within thirty days, fix the retail price of such new drug and such price shall be applicable to such applicant of such new drug.

(5) Where existing manufacturer of scheduled formulation fails to apply for prior approval of the price of the new drug in Form-I, such manufacturer shall be liable to deposit the overcharged amount over and above such price fixed and notified by the Government, if any, along with interest thereon from the date of launch of the new drug, in addition to the penalty.

(6) No existing manufacturer of a scheduled formulation shall sell such a new drug at a price higher than the retail price (plus local taxes as applicable) fixed by the Government for such new drug and in case such a manufacturer is found to sell such a new drug at a price higher than the retail price (plus local taxes as applicable) fixed by the Government, such manufacturer of the new drug shall be liable to deposit the overcharged amount along with interest from the date of overcharge, in addition to the penalty.

16. Revision of ceiling price of scheduled formulations.– (1) The Government shall revise the ceiling prices of scheduled formulations as per the annual wholesale price index (WPI) for preceding calendar year on or before 1^{st} April of every year and notify the same on the 1^{st} day of April every year.

(2) The manufacturers may increase the maximum retail price (MRP) of scheduled formulations once in a year, in the month of April, on the basis of the wholesale price index with respect to previous calendar year and no prior approval of the Government in this regard shall be required.

(3) Information about the revision, if carried out, shall be forwarded to the Government in either electronic or physical form in Form-II within a period of fifteen days of such revision and non-submission of information under this sub-paragraph shall be construed as non revision of maximum retail price (MRP) and the concerned manufacturer shall be liable to deposit the amount charged over and above the pre-revised maximum retail price (MRP), alongwith interest thereon from the date of overcharging.

(4) In case of decline in wholesale price index, there shall be a corresponding reduction in the maximum retail price and in case of scheduled formulations produced or available in the market before the date of notification of revised ceiling price, the manufacturers shall ensure within a period of forty-five days of the date of such notification that the maximum retail price (MRP) of such scheduled formulation does not exceed the revised ceiling price (plus local taxes as applicable) and information about the revision shall be sent to the Government in either electronic or physical form in Form-II within a period of fifteen days of such revision.

(5) Non-submission of information under the sub-paragraph (4) shall be construed as non reduction in maximum retail price (MRP) and the concerned manufacturer shall be liable to deposit the amount charged over and above the maximum retail price revised based on decline in wholesale price index, alongwith interest thereon as overcharged amount from the date of overcharging.

17. Amendment of the list of scheduled formulation.– (1) A decision to amend the first schedule, clearly stating the reasons thereof, shall be taken by the Government within sixty days of receipt of communication from the Ministry of Health and Family Welfare and the amendment(s) or revision, if required, in the first schedule shall be notified and thereafter, the ceiling price(s) for the medicine(s) added in the first schedule shall be fixed as per the provisions of this order within a period of sixty days from the date of the notification.

(2) The medicines omitted from the first schedule shall fall under the category of non-scheduled formulations.

18. Revision of ceiling price on the basis of moving annual turnover (MAT).- The revision of ceiling prices on the basis of moving annual turnover value shall be carried out,-

(i) as and when the National List of Essential Medicines is revised by the Ministry of Health and Family Welfare or five years from the date of fixing the ceiling price under this Order whichever is earlier;

(ii) when the number of manufacturers of a scheduled formulation, having price of a scheduled formulation more than or equal to seventy five percent of the ceiling price fixed and notified by the Government, has decreased by twenty five percent or more than the number of manufacturers as existing on the reference date;

(iii) when the number of manufacturers of a scheduled formulation, having prices of their scheduled formulation equal to or lower than twenty five percent of the ceiling price fixed by the Government, has increased by twenty five percent or more than the number of manufacturers as existing on the reference date.

Explanation.- For the purpose of items (ii) and (iii) the "reference date" shall be for first revision of ceiling price May, 2012 and for second or subsequent revision the date of previous revision of the ceiling price.

19. Fixation of ceiling price of a drug under certain circumstances.- Notwithstanding anything contained in this order, the Government may, in case of extra-ordinary circumstances, if it considers necessary so to do in public interest, fix the ceiling price or retail price of any Drug for such period, as it may deem fit and where the ceiling price or retail price of the drug is already fixed and notified, the Government may allow an increase or decrease in the ceiling price or the retail price, as the case may be, irrespective of annual wholesale price index for that year.

20. Monitoring the prices of non-scheduled formulations.– (1) The Government shall monitor the maximum retail prices (MRP) of all the drugs, including the non-scheduled formulations and ensure that no manufacturer increases the maximum retail price of a drug more than ten percent of maximum retail price during preceding twelve months and where the increase is beyond ten percent of maximum retail price, it shall reduce the same to the level of ten percent of maximum retail price for next twelve months.

(2) The manufacturer shall be liable to deposit the overcharged amount along with interest thereon from the date of increase in price in addition to the penalty.

21. Monitoring the availability of scheduled formulations.– (1) The Government shall monitor the production and availability of scheduled formulations and the active pharmaceutical ingredients contained in the scheduled formulation and the manufacturer of scheduled formulations and the active pharmaceutical ingredients contained in the scheduled formulation shall furnish the information as stated in Form-III of schedule-II of this Order quarterly.

(2) Any manufacturer of scheduled formulation, intending to discontinue any scheduled formulation from the market shall issue a public notice and also intimate the Government in Form-IV of schedule-II of this order in this regard at least six month prior to the intended date of discontinuation and the Government may, in public interest, direct the manufacturer of the scheduled formulation to continue with required level of production or import for a period not exceeding one year, from the intended date of such discontinuation within a period of sixty days of receipt of such intimation.

22. Recovery of dues accrued under the Drugs (Prices Control) Order, 1979 and to deposit the same into the Drugs Prices Equalisation Account.– (1) Notwithstanding anything contained in this order, the Government may by notice, require a manufacturer, importer or distributor as the case may be, to deposit the amount which has accrued under the provisions of the Drugs (Prices Control) Order, 1979 on or before the commencement of this order, into the Drugs Prices Equalisation Account and the manufacturer, importer or distributor, as the case may be, shall deposit the said amount into the said account within such time as the Government may specify in the said notice.

(2) The existing amount, if any, in the Drugs Prices Equalisation Account on or before the date of commencement of this Order, and the amount deposited under sub-paragraph (1) shall be utilised for;-

(a) paying to the manufacturer, importer or distributor, as the case may be, the shortfall between his retention price and the common selling price or, as the case may be, the pooled price for the purpose of increasing the production, or securing the equitable distribution and availability at fair prices, of drugs;

(b) meeting the expenses incurred by the Government in discharging the functions under this paragraph; and

(c) promoting higher education and research in Pharmaceutical Sciences and Technology and for the purposes incidental thereto.

23. Recovery of overcharged amount under Drugs Prices Control Orders 1987 and 1995.– Notwithstanding anything contained in this order, the Government shall by notice, require the manufacturers, importer or distributor or as the case may be, to deposit the amount accrued due to charging of prices higher than those fixed or notified by the Government under the provisions of Drugs (Prices Control) Order, 1987 and Drugs (Prices Control) Order, 1995 under the provisions of this Order.

24. Carrying into effect the price fixed or revised by the Government, its display and proof thereof.– (1) For all the scheduled formulations having maximum retail price (MRP) higher than ceiling price (plus local taxes as applicable), the manufactures shall revise the maximum retail price (MRP) not exceeding the ceiling price (plus local taxes as applicable):

Provided that in case of scheduled formulations produced or available in the market before the date of notification of ceiling price, the manufacturers shall ensure within a period of forty-five days of the date of the notification that the maximum retail price of such scheduled formulation does not exceed the ceiling price (plus local taxes as applicable).

(2) Every manufacturer of a schedule formulation intended for sale shall display in indelible print mark, on the label of container of the formulation and the minimum pack thereof offered for retail sale, the maximum retail price of that formulation based on the ceiling price notified in the Official Gazette or ordered by the Government in this behalf with the words "Maximum Retail Price" preceding it and the words 'inclusive of all taxes' succeeding it.

(3) Every manufacturer shall issue a price list and supplementary price list, if required, in Form V to the dealers, State Drugs Controllers and the Government indicating reference to such price fixation or revision as covered by the order or Gazette notification issued by the Government, from time to time.

(4) Every retailer and dealer shall display the price list and the supplementary price list, if any, as furnished by the manufacturer, on a conspicuous part of the premises where he carries on business in a manner so as to be easily accessible to any person wishing to consult the same.

25. Display of prices of non-scheduled formulations and price list thereof.– (1) Every manufacturer of a non-Scheduled formulation intended for sale shall display in indelible print mark, on the label of container of the formulation and the minimum pack thereof offered for retail sale, the maximum retail price of that formulation with the words "Maximum Retail Price" preceding it and the words 'inclusive of all taxes' succeeding it.

(2) Every manufacturer shall issue a price list and supplementary price list, if required, of the non-Scheduled formulations in Form-V to the dealers, State Drugs Controllers and the Government indicating changes, from time to time.

(3) Every retailer and dealer shall display the price list and the supplementary price list, if any, as furnished by the manufacturer or importer, on a conspicuous part of the premises where he carries on business in a manner so as to be easily accessible to any person wishing to consult the same.

26. Control of sale prices of formulations.- No person shall sell any formulation to any consumer at a price exceeding the price specified in the current price list or price indicated on the label of the container or pack thereof, whichever is less.

27. Sale of split quantities of formulations.- No dealer shall sell loose quantity of any formulation at a price which exceeds the pro-rata price of the formulation.

28. Manufacturer, distributor or dealer not to refuse sale of drug. – Subject to the provisions of the Drug and Cosmetics Act, 1940 (23 of 1940) and the rules made thereunder, -

(a) no manufacturer or distributor shall withhold from sale or refuse to sell to a dealer any drug without good and sufficient reasons;

(b) no dealer shall withhold from sale or refuse to sell any drug available with him to a customer intending to purchase such drug.

29. Maintenance of records and production thereof for inspection.— Every manufacturer shall maintain records relating to the sales of individual active pharmaceutical ingredients or bulk drugs manufactured or imported and marketed by him, as the case may be, and the sales of formulations units and packs and also such other records as may be directed from time to time by the Government and the Government shall have the power to call for any record and to inspect such records at the premises of the manufacturer.

30. Power of entry, search and seizure.– (1) Any Gazetted Officer of the Central Government or of a State Government, as the case may be, authorised by a general or special order by the Central Government or by the State Government, as the case may be, in this behalf may, with a view to securing compliance with this Order or to satisfy himself that the provision of this Order have been complied with–

(a) enter and search any place;

(b) seize any drug, alongwith the containers, packages or coverings in which the drug is found, in respect of which he suspects that any provision of this Order has been, is being, or is about to be contravened, and thereafter take all measures necessary for securing production of the drug, containers, packages or coverings, so seized, in a court of law and for their safe custody pending such production;

(c) seize any document, such as, cash memo or credit memo books, books of account and records of purchase and sale of the drugs in respect of which he suspects that any provision of this Order has been, is being, or is about to be contravened.

(2) The provisions of Code of Criminal Procedure, 1973 (2 of 1974), relating to search and seizure shall, so far as may be, apply to searches and seizures under this Order.

31. Power to review.– Any person aggrieved by any notification issued or order made under paragraphs 4, 5 and 6 of this Order, may apply to the Government for a review of the notification or order within a period of thirty days of the date of publication of the notification in the Official Gazette or the receipt of the order by him, as the case may be, and the Government may make such order on the application as it may deem proper:

Provided that pending a decision by the Government on the application submitted under the above paragraph, no manufacturer shall sell a scheduled formulation or a new drug, as the case may be, at a price exceeding the ceiling price or retail price, as the case may be, fixed by the Government of which a review has been applied for.

32. Non–application of the provisions of this order in certain cases.- The provisions of this order shall not apply to, -

(i) a manufacturer producing a new drug patented under the Indian Patent Act, 1970 (39 of 1970) (product patent) and not produced elsewhere, if developed through indigenous Research and Development, for a period of five years from the date of commencement of its commercial production in the country.

(ii) a manufacturer producing a new drug in the country by a new process developed through indigenous Research and Development and patented under the Indian Patent Act, 1970 (39 of 1970) (process patent) for a period of five years from the date of the commencement of its commercial production in the country.

(iii) a manufacturer producing a new drug involving a new delivery system developed through indigenous Research and Development for a period of five years from the date of its market approval in India:

Provided that the provision of this paragraph shall apply only when a document showing approval of such new drugs from Drugs Controller General (India) is produced before the Government.

Explanation.- Notwithstanding anything contained in this Order, for the purpose of this paragraph "new drug" shall have the same meaning as is assigned to under rule 122E of the Drugs and Cosmetics Rules, 1945;

File No.31011/17/2012-PI-II

(Shambhu Kallolikar) Joint Secretary to the Government of India

Schedule-I

(See Paragraphs-2(t),2(zb))

Symbols P, S and T appearing in NLEM 2011 denote essentiality at Primary, Secondary and Tertiary levels respectively.

NATIONAL LIST OF ESSENTIAL MEDICINES 2011						
	Section: 1 – Anesthesia					
1	.1 General An	esthetics and Oxygen				
Medicines	Category	Route of Administration	Strengths			
Ether	S, T	Inhalation				
Halothane with vaporizer	S, T	Inhalation				
Isoflurane	S, T	Inhalation				
Ketamine Hydrochloride	P, S, T	Injection	10 mg / ml, 50 mg / ml			
Nitrous Oxide	P, S, T	Inhalation				
Oxygen	P, S, T	Inhalation				
Thiopentone Sodium	S, T	Injection	0.5 g, 1 g powder			
	Adde	d Medicines				
Sevoflurane	т	Inhalation				
Propofol	P,S,T	Injection	1% oil suspension			
	1.2 Loc	al Anesthetics				
Medicines	Category	Route of Administration/ Dosage Form	Strengths			

Bupivacaine Hydrochloride	S, T	Injection	0.25%, 0.5%, 0.5% to be mixed with 7.5% glucose solution
Lignocaine Hydrochloride	P, S, T	Topical Forms, Injection, Spinal	2-5%, 1-2%, 5% +7.5% Glucose
Lignocaine Hydrochloride + Adrenaline	P, S, T	Injection	1%, 2% + Adrenaline1:200,0 00
	Adde	d Medicines	
EMLA cream	т	Cream	
1.3 Preoperative	Medication and	d Sedation for Short Ter	m Procedures
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Atropine Sulphate	P, S, T	Injection	0.6 mg / ml
Diazepam	P,S,T S, T	Tablets Injection, Syrup, Suppository	5 mg 5 mg / ml 2mg/5ml 5 mg
Midazolam	P, S, T	Injection	1 mg / ml 5 mg / ml
Morphine Sulphate	S, T	Injection	10 mg / ml
Promethazine	P, S, T	Syrup	5 mg / 5 ml

Section: 2 - Analgesics , Antipyretics, Nonsteroidal Antiinflammatory Medicines, Medicines used to treat Gout and Disease Modifying Agents used in Rheumatoid Disorders

.1: Non-Opioid A	nalgesics, A	Antipyretics and Nonstero Medicines	bidal Anti-inflammator
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Acetyl Salicylic Acid	P, S, T	Tablets	325, 350 mg
Diclofenac	т	Tablets	50 mg
	Т	Injection	25 mg / ml
lleuronefere	БОТ	Tablets	200 mg, 400 mg
lbuprofen	P, S, T	Syrup	100mg/5ml
	P, S, T	Injection	150 mg / ml
Deressional	P, S, T	Syrup	125 mg / 5ml
Paracetamol	P, S, T	Tablets	500 mg
	P, S, T	Suppository	80 mg, 170 mg
	2	.2 Opioid Analgesics	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Morphine	о т	Injection	10 mg / ml
Sulphate	S, T	Tablets	10 mg
		Added medicines	
Transis I.I	0 T	Injection	50 mg/ml
Tramadol	S,T	Сар	50 mg,100 mg
Fentanyl	S,T	Injection	50ug/ml 2ml ampou

2.3 Medicines used to treat Gout						
Allopurinol	S, T	Tablets	100 mg			
Colchicine	S, T	Tablets	0.5 mg			
2.4 Disea	se modifying	agents used in Rheumate	oid disorders			
Azathioprine	S, T	Tablets	50 mg			
Methotrexate	S,T	Tablets	5mg, 7.5mg, 10mg			
Sulfasalazine	S, T	Tablets	500 mg			
Added medicines						
Hydroxychloroquin e phosphate	S,T	Tablets	200 mg			
Leflunomide	S,T	Tablets	10mg, 20 mg tab			

Medicines	Category	Route of Administration/ Dosage Form	Strengths
Adrenaline Bitartrate	P, S, T	Injection	1 mg / ml
Chlorpheniramine Maleate	P, S, T	Tablets	4 mg
Dexchlorpheniramine Maleate	P, S, T	Syrup	0.5 mg / 5 ml
Dexamethasone	P, S, T	Tablets	0.5 mg
		Injection	4 mg / ml
Hydrocortisone Sodium Succinate	P, S, T	Injection	100 mg
Pheniramine Maleate	P, S, T	Injection	22.75 mg / ml
Prednisolone	P, S, T	Tablets	5 mg, 10 mg, 20 mg
Promethazine	P, S, T	Tablets	10 mg, 25 mg
	. ,	Syrup	5 mg / 5 ml
	Adde	d Medicines	
Cetrizine	P,S,T	Tablets	10mg
		Syrup	5 mg/ml

Section: 4 - Antidotes and Other Substances used in Poisonings				
	4.1: I	Nonspecific		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Activated Charcoal	P,S,T	Oral		
	4.2	: Specific		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Atropine Sulphate	P,S,T	Injection	1 mg/ml	
Specific Antisnake venom	P,S,T	Injection Polyvalent Solution/ Lyophilyzed Polyvalent Serum		
Calcium gluconate	P,S,T	Injection	100mg/ml	
Desferrioxamine mesylate	S, T	Injection	500mg	
Methylthioninium chloride (Methylene blue)	S, T	Injection	10 mg / ml	
Penicillamine	S, T	Tablets or Capsules	250 mg	
Dimercaprol	S, T	Injection in oil	50 mg / ml	
Flumazenil	Т	Injection	0.1 mg / ml	
Sodium Nitrite	S, T	Injection	30 mg / ml	
Sodium Thiosulphate	S, T	Injection	250 mg/ ml	
Naloxone	P,S,T	Injection	0.4mg/ml	
Pralidoxime Chloride(2- PAM)	P,S,T	Injection	25 mg/ml	
	Addee	d medicines:		
N-acetylcysteine	P,S,T	Injection	200 mg/ml(5 ml)	

Section: 5 – Anticonvulsants/ Antiepileptics					
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Carbamazepine	P, S, T	Tablets	100mg		
		Syrup	200mg		
			100 mg/5ml		
Diazepam	P,S,T	Injection	5 mg / ml		
Magnesium sulphate	S,T	Injection	500 mg /ml		
Phenobarbitone	P,S,T	Tablets	30 mg, 60 mg		
	ST	Injection	200 mg/ml		
	P,S,T	Syrup	20 mg/5ml		
Phenytoin Sodium	P,S,T	Capsules or Tablets	50 mg,		
		Syrup	100mg		
		Injection	25mg/ml		
			50 mg/ml		
Sodium Valproate	P,S,T	Tablets	200 mg,		
		Syrup	500mg		
			200 mg/5ml		
	т	Injection	100 mg/ml		
	Added Medicines				
Lorazepam	т	Injection	2mg/ml		

Section: 6 – Anti-infective Medicines					
	6.1 Anthelminthics				
	6.1.1 Intesti	nal Anthelminthics			
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Albendazole	P,S,T	Tablets Suspension	400 mg 200 mg/ 5 ml		
	Adde	d Medicines			
Piperazine	P,S,T	Tablets Solution	4.5 gm 750mg/5ml		
	6.1.2	Antifilarials	l		
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Diethylcarbamazine citrate	P,S,T	Tablets	50 mg		
6.1.3 Antisc	chistosomals	and Antitrematode Medicir	nes		
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Praziquantel	S, T	Tablets	600 mg		
	6.2 A	ntibacterials			
	6.2.1 Beta	lactam medicines			
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Amoxicillin	S,T	Powder for suspension	125 mg / 5 ml		
		Capsules	250 mg, 500 mg		
		Capsules	250 mg, 500 mg		
Ampicillin	P,S,T	Powder for suspension	125 mg / 5 ml		
		Injection	500 mg		

	1		
Benzathine Benzylpenicillin	P,S,T	Injection	6 lacs, 12 lacs units
Cefotaxime	S, T	Injection	125 mg, 250 mg 500 mg
Ceftazidime	S, T	Injection	250mg, 1g
Ceftriaxone	S, T	Injection	250 mg, 1 g
Cephalexin	P,S,T	Syrup Capsules	125 mg / 5 ml 250 mg, 500 mg
Cloxacillin	P,S,T	Capsules Injection Liquid	250 mg, 500 mg 250 mg 125mg/ 5 ml
	Adde	d Medicines	
Amoxicillin+Clavulinic acid	т	Tablets Powder for suspension Injection	625 mg 228.5mg/5ml 600mg, 1.2gm
Cefixime	Т	Tablet	100, 200mg
	6.2.2 Oth	er antibacterials	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Amikacin	S, T	Injection	250 mg / 2 ml
Azithromycin	S,T	Tablets Suspension Injection	100, 250,500mg 100mg/5ml 500mg
Ciprofloxacin Hydrochloride	P,S,T	Injection Tablets	200 mg /100 ml 250 mg, 500 mg
Co-Trimoxazole (Trimethoprim + Sulphamethoxazole)	P,S,T	Tablets Suspension	80 + 400 mg 160+800 mg 40 + 200 mg / 5 ml

Doxycycline	P,S,T	Tablets	100 mg
Erythromycin Estolate	P,S,T	Syrup Tablets	125 mg / 5 ml 250 mg, 500 mg
Gentamicin	P,S,T	Injection	10 mg/ml, 40 mg/ml
Metronidazole	P,S,T	Tablet Injection Syrup	200mg,400 mg 500mg/100 ml 100mg/5ml
Nitrofurantoin	P,S,T	Tablets	100 mg
Sulphadiazine	S, T	Tablets	500 mg
Vancomycin Hydrochloride	т	Injection	500 mg, 1 g
	6.2.3 Antile	eprosy medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Clofazimine	P,S, T	Capsules	50 mg, 100 mg
Dapsone	P,S, T	Tablets	50 mg, 100mg
Rifampicin	P,S, T	Capsules or Tablets	150 mg, 300 mg
	6.2.4 Antitube	erculosis medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Ethambutol	P,S,T	Tablets	200 mg, 400 mg, 600 mg, 800 mg
Isoniazid	P,S,T	Tablets Syrup	50 mg, 100 mg, 300 mg 100 mg/5ml
Ofloxacin	S, T	Tablets	100 mg, 200 mg

		Syrup	50 mg / 5 ml	
Pyrazinamide	P,S,T	Tablets	500 mg, 750 mg, 1000 mg, 1500 mg	
		Capsules/Tablets	50 mg, 150 mg, 300 mg,450 mg	
Rifampicin	P,S,T	Syrup	100 mg / 5 ml	
Streptomycin				
Sulphate	P,S,T	Injection	0.75 g, 1 g	
	6.3 Antifu	ingal medicines		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Amphotericin B	S, T	Injection	50 mg	
		Pessaries	100 mg, 200 mg,	
Clotrimazole	P,S,T	Gel	2%	
Fluconazole			50 mg, 100 mg, 150 mg,	
	S, T	Capsules or Tablets	200 mg	
Griseofulvin	P,S,T	Capsules or Tablets	125 mg, 250 mg	
		Tablets	500,000 IU	
Nystatin	P,S,T	Pessaries	100,000 IU	
	6.4 Antiv	viral medicines		
	6.4.1 Antih	erpes medicines		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Acyclovir	S, T	Tablets	200 mg, 400 mg	
1.59010111		Injection	250 mg, 500 mg	
		Suspension	400 mg / 5 ml	
	6.4.2 Antiretroviral medicines			
6.4.2.1 Nucleoside reverse transcriptase inhibitors				

Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Didanosine	S, T	Tablets	250 mg, 400 mg		
Lamivudin <i>e</i>	S, T	Tablets	150 mg		
Lamivudine + Nevirapine + Stavudine	S, T	Tablets	150 mg + 200 mg+ 30 mg		
Lamivudine + Zidovudine	S, T	Tablets	150 mg + 300 mg		
Stavudine	S, T	Capsules	15 mg, 30 mg, 40 mg		
Zidovudine	S, T	Tablets	100 mg, 300 mg		
	ADDEI				
Stavudine+ Lamivudine	S,T	Tablets	30mg+ 150mg		
Zidovudine+	S,T	Tablets	300mg+		
Lamivudine+			150mg+		
Nevirapine			200mg		
6.4.2.2 Non-	nucleoside r	everse transcriptase inhibit	ors		
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
			200 mg,		
Efavirenz	S, T	Capsules	600 mg		
		Capsules	200 mg		
Nevirapine	S, T	Suspension	50 mg / 5 ml		
6.4.2.3 Protease inhibitors					
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Indinavir	S, T	Capsules	200 mg, 400 mg		

		r				
S, T	Capsules	250 mg				
	Capsules	100 mg,				
S, T	Syrup	400 mg / 5ml				
S, T	Capsules	200 mg				
6.5 Antiprotozoal Medicines						
6.5.1 Antiamoebic and Antigiardiasis medicines						
Category	Route of Administration/ Dosage Form	Strengths				
P,S,T	Tablets	500 mg				
P,S,T	Tablets	200 mg, 400 mg				
	Injection	500 mg /100 ml				
6.5.2 Antileishmaniasis medicines						
Category	Route of Administration/ Dosage Form	Strengths				
S, T	Injection	50 mg				
S, T	Injection	200 mg				
S, T	Injection	100 mg / ml				
6.5.3 Antimalarial Medicines						
6.5.3.1 For	curative treatment					
Category	Route of Administration/ Dosage Form	Strengths				
P,S,T	Tablets	50 mg				
	Tablets	150 mg base				
	Injection	40 mg / ml				
P,S,T	Syrup	50 mg / 5 ml				
P,S,T	Tablets	2.5 mg, 7.5 mg				
	S, T S, T 6.5 Antipro tiamoebic ar Category P,S,T P,S,T 5.2 Antileish Category S, T S, T S, T 6.5.3 Antim 6.5.3 Antim 6.5.3 Antim 6.5.3 I For Category P,S,T	S, TCapsules SyrupS, TCapsules6.5 Antiprotocal Medicinestiamoebic attration/cospane FormRoute of Administration/ Dosage FormP,S,TTabletsP,S,TTabletsP,S,TTabletsS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TInjectionS,TSocage FormP,S,TTabletsP,S,TSyrup				

Pyrimethamine	P,S,T	Tablets	25 mg		
Quinine sulphate	P,S,T	Tablets	300 mg		
	ST	Injection	300 mg / ml		
Sulfadoxine + Pyrimethamine	P,S,T	Tablets	500 mg + 25 mg		
Medicines added					
Clindamycin	S,T	Tablet	150, 300mg		
6.5.3.2 For prophylaxis					
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Medicines added					
Mefloquine	S,T	Tablet	250 mg base		
6.5.4 Antipneumocystosis and Antitoxoplasmosis medicines					
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Co-Trimoxazole	P,S,T	Tablets	80 + 400 mg		
(Trimethoprim + Sulphamethoxazole)			160+800 mg		
		Suspension	40 + 200 mg / 5 ml		
Pentamidine Isothionate	S, T	Injection	200 mg		

Section: 7 – Antimigraine medicines					
7.1: For treatment of acute attack					
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Acetyl Salicylic Acid	P,S,T	Tablets	300 - 350 mg		
Dihydroergotamine	S, T	Tablets	1 mg		
Paracetamol	P,S,T	Tablets	500 mg		
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7.2: For Prophylaxis					
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Propranolol hydrochloride	P,S,T	Tablets	10 mg, 40 mg		

Section: 8 – Antineoplastic, immunosuppressives and medicines used in palliative care			
	8.1: Immunos	uppressive medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Azathioprine	Т	Tablets	50 mg
Cyclosporine	т	Capsules	10 mg, 25 mg, 50 mg, 100 mg
		Concentrate for Injection	100 mg/ml
	8.2: Cyto	otoxic medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Actinomycin D	Т	Injection	0.5 mg
Alpha Interferon	Т	Injection	3 million IU
Bleomycin	Т	Injection	15 mg
Busulphan	Т	Tablets	2 mg
Cisplatin	Т	Injection	10 mg / vial 50 mg / vial
Cyclophosphami de	Т	Tablets Injection	50 mg, 200 mg 500 mg
Cytosine	Т	Injection	100 mg/vial

arabinoside			500 mg/vial 1000 mg/vial
Danazol	Т	Capsules	50 mg, 100 mg
Doxorubicin	Т	Injection	10 mg, 50 mg
Etoposide	т	Capsules Injection	100 mg 100 mg/ 5 ml vial
Flutamide	Т	Tablet	250 mg
5-Fluorouracil	Т	Injection	250 mg / 5 ml
Folinic Acid	Т	Injection	3 mg / ml
Gemcitabine hydrochloride	т	Injection	200 mg 1 gm
L- Asparaginase	Т	Injection	5000 KU.
Melphalan	Т	Tablet	2 mg, 5 mg
Mercaptopurine	т	Tablet Injection	50 mg 100 mg / ml
Methotrexate	Т	Tablet Injection	2.5 mg 50 mg / ml
Mitomycin-C	Т	Injection	10 mg
Paclitaxel	Т	Injection	30 mg / 5 ml
Procarbazine	Т	Capsules	50 mg
Vinblastine sulphate	т	Injection	10 mg
Vincristine	Т	Injection	1 mg / ml
	Adde	ed medicines	
Carboplatin	Т	Injection	150 mg, 450 mg vial
Dacarbazine	Т	Injection	500 mg
Daunorubicin	Т	Injection	20 mg vial

Т	Injection	1 gm/2ml vial
Т	Injection	200 mg
Т	Injection	50 mg vial
Т	Tablets	100 mg, 400 mg
Т	Tablets	2 mg
8.3: Hormone	es and antihormones	
Category	Route of Administration/ Dosage Form	Strengths
	Tablets	5 mg
S, T	Injection	20 mg, 25 mg (as sodium phosphate or succinate)
Т	Tablets	60 mg
Т	Tablets	10 mg, 20 mg
8.4: Medicines	used in palliative care	
Category	Route of Administration/ Dosage Form	Strengths
Т	Tablets	10 mg
	Tablets	4 mg, 8 mg
S, T	Injection	2 mg/ml
	Syrup	2 mg/5 ml
Adde	ed Medicines	
Т	Injection	1 ml vial
Т	Tablets	100 mg
	T T T 8.3: Hormone Category S, T T S, T T 8.4: Medicines Category T S, T Adde T	T Injection T Injection T Tablets T Tablets T Tablets 8.3: Hormores and antihormones Administration/ Dosage Form Category Route of Administration/ Dosage Form S, T Injection T Tablets T Tablets S, T Injection T Tablets T Tablets T Tablets T Tablets Gategory Route of Administration/ Dosage Form T Tablets S, T Injection S, T Injection S, T Syrup T Syrup

Section: 9 – Antiparkinsonism medicines			
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Bromocriptine Mesylate	S, T	Tablets	1.25 mg, 2.5 mg
Levodopa+ Carbidopa	P,S,T	Tablets	100 mg+10 mg 250 mg+25 mg 100 mg+25 mg
Trihexyphenid yl Hydrochloride	P,S,T	Tablets	2 mg

Section: 10 – Medicines affecting the blood			
	10.1: A	ntianaemia medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Cyanocobalami n	P, S,T	Injection	1 mg/ml
Ferrous Sulphate/	P,S,T	Tablets	Tablets equivalent to 60 mg elemental iron
Fumrate		Oral solution	25mg elemental iron (as sulphate)/ml
Folic Acid	P,S,T	Tablets	1 mg , 5mg
Iron Dextran	S, T	Injection	50 mg iron/ml
Pyridoxine	P,S,T	Tablets	10 mg
	10.2: Medic	ines affecting coagulation	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Heparin Sodium	S, T	Injection	1000 IU/ml 5000 IU/ ml

Protamine Sulphate	S, T	Injection	10 mg/ml
Phytomenadion e	P, S, T	Injection	10 mg/ml
Warfarin sodium	S, T	Tablets	5 mg
Added Medicines			
Enoxaparin	Т	Injection	40mg, 60mg

Section: 11 –Blood products and Plasma substitutes			
	11.1:	Plasma Substitutes	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Dextran-40	P,S,T	Injection	10%
Dextran-70	P,S,T	Injection	6%
Fresh frozen plasma	т	Injection	
Hydroxyethyl Starch (Hetastarch)	S, T	Injection	6%
Polygeline	S, T	Injection	3.5%
	11.2: Plasm	a fractions for specific use	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Albumin	S, T	Injection	5%, 20 %
Cryoprecipitate	S, T	Injection	
Factor VIII Concentrate	S, T	Injection	Dried
Factor IX Complex (Coagulation Factors II,VII, IX,	S, T	Injection	Dried

X)			
Platelet Rich Plasma	S, T	Injection	

Section: 12 – Cardiovascular medicines			
	12.1: A	Intianginal medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Acetyl salicylic acid	P,S,T	Tablets	75mg, 100mg, 350 mg soluble/dispersible
Diltiazem	S, T	Tablets	30 mg, 60 mg
Glyceryl Trinitrate	P,S,T	Sublingual Tablets Injection	0.5 mg 5mg/ml
Isosorbide 5 Mononitrate/ Dinitrate	P,S,T	Tablets	10 mg, 20 mg
Metoprolol	P,S,T	Tablets Injection	25 mg, 50 mg 1mg/ml
	P	Added Medicines	
Clopidogrel	Т	Tablets	75 mg
	12.2: An	tiarrhythmic medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Adenosine	S,T	Injection	3 mg/ml
Amiodarone	S, T	Tablets Injection	100 mg, 200 mg 50 mg/ml (3 ml ampoule)

	1		
Diltiazem	S, T	Tablets	30 mg, 60 mg
	Т	Injection	5 mg/ ml
Esmolol	Т	Injection	10 mg / ml
Lignocaine Hydrochloride	S, T	Injection	1%, 2%
Procainamide Hydrochloride	Т	Tablets Injection	250 mg 100mg/ml
Verapamil	S, T	Tablets Injection	40 mg, 80 mg 2.5mg/ml
	12.3: Anti	hypertensive medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Amlodipine	P,S,T	Tablets	2.5 mg, 5 mg
Atenolol	P,S,T	Tablets	50 mg, 100 mg
Enalapril Maleate	P,S,T T	Tablets Injection	2.5 mg, 5mg 1.25mg/ml
Losartan Potassium	S, T	Tablets	25 mg, 50 mg
Methyldopa	P,S, T	Tablets	250 mg
Nifedipine	S, T	Capsules Tablets Sustained release tablets or capsules	5 mg, 10mg 10mg, 20mg 10mg, 20mg
Sodium Nitroprusside	Т	Injection	50 mg/ 5 ml

Added Medicines			
Hydrochlorthiazi	P,S,T	Tablets	12.5,
de	F,0,1	i ablets	25 mg

12.4: Medicines used in heart failure			
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Digoxin		Tablets	0.25 mg
	S, T	Injection	0.25 mg/ml
		Elixir	0.05 mg/ml
Dobutamine	S, T	Injection	50 mg / ml
Dopamine Hydrochloride	S,T	Injection	40 mg / ml
	12.5: An	tithrombotic medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Acetyl salicylic acid	P,S,T	Tablets	75mg, 100mg, 350 mg soluble/dispersible
Heparin Sodium	S, T	Injection	1000 IU /ml 5000 IU/ml
Streptokinase	S, T	Injection	750,000 IU 15,00,000 IU
Urokinase	Т	Injection	500,000

			IU/ml		
			10,00,000 IU/ml		
	New Category - ADDED				
	12.6 Hypolipidemic Medicines				
Atorvastatin	P,S,T	Tablets	5 mg, 10 mg		

Section: 13 – Dermatological medicines (Topical)			
	13.1: 4	Antifungal medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Miconazole	P,S,T	Ointment or Cream	2%
	13.2: Ai	ntiinfective medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Acyclovir	S, T	Cream	5%
Framycetin Sulphate	P,S,T	Cream	0.5%
Methylrosaniliniu m Chloride (Gentian Violet)	P,S,T	Aqueous solution	0.5%
Neomycin + Bacitracin	P,S,T	Ointment	5 mg + 500 IU / g
Povidone Iodine	P,S,T	Solution or Ointment	5%
Silver Sulphadiazine	P,S,T	Cream	1%
13.3: Antiinflammatory and antipruritic medicines			
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Betamethason	P,S,T	Cream / Ointment	0.05%

e Dipropionate				
Calamine	P,S,T	Lotion		
	13.4: A	Astringent Medicines		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Zinc Oxide	P,S,T	Dusting Powder		
13.5: M	edicines affecting	g skin differentiation and	proliferation	
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Coal Tar	P,S,T	Solution	5%	
Dithranol	Т	Ointment	0.1-2%	
Glycerin	P,S,T	Solution		
Salicylic Acid	P,S,T	Solution	5%	
	13.6: Scab	icides and Pediculicides		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Benzyl benzoate	P,S,T	Lotion	25 %	
Added Medicines				
Permethrin	S,T	Cream Lotion	5% 1%, 5%	

Section: 14 – Diagnostic agents				
	14.1: Ophthalmic medicines			
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Fluorescein	S, T	Eye drops	1%	
Lignocaine	S, T	Eye Drops	4%	

Tropicamide	S, T	Eye drops	1%
	14.2: Ra	adiocontrast media	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Barium Sulphate	S, T	Suspension	100% w/v, 250% w/v
Calcium Ipodate	S, T	Injection	3 g
Iopanoic Acid	S, T	Tablets	500 mg
Meglumine Iothalamate	S, T	Injection	60% w/v (iodine =280 mg / ml)
Meglumine lotroxate	S, T	Solution	5-8 g iodine in 100-250 ml
Propyliodone	S, T	Oily, suspension	500-600 mg / ml
Sodium Iothalamate	S, T	Injection	70% w/v(lodine =420 mg / ml)
Sodium Meglumine Diatrizoate	S, T	Injection	60% w/v(lodine conc. =292 mg / ml), 76% w/v(lodine conc. =370 mg / ml)

Section: 15 – Disinfectants and antiseptics				
15.1: Antiseptics Route of Medicines Category Administration/ Dosage Strengths				
Acriflavin+Glycerin	P, S, T	Form Solution		
Benzoin Compound	P, S, T	Tincture		
Cetrimide	P, S, T	Solution	20% (conc. for dilution)	

Chlorhexidine	P, S, T	Solution	5% (conc. for dilution)
Ethyl Alcohol 70%	P, S, T	Solution	
Gentian Violet	P, S, T	Paint	0.5%, 1%
Hydrogen Peroxide	P, S, T	Solution	6%
Povidone lodine	P, S, T	Solution	5%, 10%
	15.2	2: Disinfectants	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Bleaching Powder	P, S, T	Powder	Contains not less than 30 % w/w of available chlorine (as per I.P)
Formaldehyde Solution	P, S, T	Solution	Dilute 34 ml of formaldehyde solution with water to produce 100 ml (As per I.P)
Glutaraldehyde	S,T	Solution	2%
Potassium Permanganate	P, S, T	Crystals for solution	

Section: 16 –Diuretics			
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Furosemide	P,S,T	Injection Tablets	10 mg/ ml 40mg
Hydrochlorothiazid e	P,S,T	Tablets	25 mg, 50 mg
Mannitol	P,S,T	Injection	10%, 20%

Spironolactone	P,S,T	Tablets	25 mg

Section: 17 – Gastrointestinal medicines			
1	7.1: Antacids ar	nd other Antiulcer medicine	es
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Aluminium Hydroxide + Magnesium Hydroxide	P,S,T	Tablet Suspension	
Omeprazole	P,S,T	Capsules	10 mg, 20 mg, 40 mg
Ranitidine	P,S,T	Injection	25 mg / ml
	Ad	ded Medicines	
Pantoprazole	Т	Injection	40 mg
Famotidine	P,S,T	Tablets	20 mg
	17.	2: Antiemetics	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Domperidone	P,S,T	Tablets Syrup	10 mg 1 mg / ml
Metoclopramide	P,S,T	Tablets Syrup Injection	10 mg 5 mg / 5 ml 5 mg / ml
Promethazine	P,S,T	Tablets Elixir or Syrup Injection	10 mg, 25 mg 5 mg / 5 ml 25 mg / ml
	Ad	ded Medicines	

Ondansetron	S,T	Tablet Syrup Injection	4mg, 8 mg 2 mg/ml 2mg/ml		
	17.3: Antiin	flammatory Medicines			
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
	Ad	ded Medicines			
5-Amino salicylic Acid (5-ASA)	S,T	Tablets	400mg		
	17.4: Antis	spasmodic medicines			
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Dicyclomine Hydrochloride	P,S,T	Tablets Injection	10 mg 10 mg / ml		
Hyoscine Butyl Bromide	P,S,T	Tablets Injection	10 mg 20 mg / ml		
	17	7.5: Laxatives	I		
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Bisacodyl	P,S,T	Tablets, Suppository	5 mg		
Ispaghula	P,S,T	Granules			
17.6: Medicines used in diarrhorea					
	17.6.1 Or	al dehydration salts			
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Oral Rehydration	P,S,T	Powder for solution	Glucose: 13.5 g/L Sodium chloride:		

Salts			2.6
			g/L
			Potassium chloride:
			1.5 g/L
			Trisodium citrate
			dihydrate+: 2.9 g/L
			Powder for dilution
			in 200ml; 500 ml;
			1000ml. (As per I.P)
	17.6.2 Anti	diarrhoeal medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
	<u> </u>		<u> </u>
Medicines added			
Zinc Sulfate	P,S,T	Syrup	20 mg/5ml

Section: 18 –Hormones, other endocrine medicines and						
	contraceptives					
18.1: <i>A</i>	Adrenal hormon	es and synthetic substitut	es			
MedicinesCategoryRoute of Administration/StrengthsDosage Form						
Dexamethasone	S,T	Tablets Injection	0.5mg 4mg/ml			
Hydrocortisone Sodium Succinate	P, S,T	Injection	100 mg / ml			
Methyl Prednisolone	S,T	Injection	40 mg/ ml			
Prednisolone	P, S,T	Tablets	5mg, 10mg, 20mg			
18.2: Androgens						
Medicines Category Route of Strengths Administration/						

		Dosage Form	
Testosterone	P,S,T	Capsules Injection	40mg(as undecanoate) 25mg/ml(as propionate)
	18.3: Co	ontraceptives	
	18.3.1: Hormo	onal Contraceptives	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Ethinylestradiol + Levonorgesterol	P,S,T	Tablets	0.03 mg +0.15 mg
Ethinylestradiol + Norethisterone	P,S,T	Tablets	0.035 mg +1.0 mg
Hormone Releasing IUD	Т	Levonorgesterol Releasing	IUD
	18.3.2: Intr	auterine devices	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
IUD containing Copper	P,S,T		
	18.3.3: B	arrier Methods	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Condoms	P,S,T		
	18.4:	Estrogens	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Ethinylestradiol	P,S,T	Tablets	0.01mg 0.05mg
18	.5: Medicines u	sed in Diabetes mellitus	1

18.5	.1: Insulins and	other Antidiabetic agents	6
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Glibenclamide	P,S,T	Tablets	2.5 mg, 5mg
Insulin Injection (Soluble)	P,S,T	Injection	40 IU / ml
Intermediate Acting(Lente/NPH Insulin)	P,S,T	Injection	40 IU / ml
Metformin	P,S,T	Tablets	500mg
	Addeo	d medicines	
Premix Insulin 30:70 injection	P,S,T	Injection	40IU/ml
18.5	5.2 Medicines us	sed to treat hypoglycemia	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Glucagon	Т	Injection	1mg/ml
	Adde	d medicines	
25% Dextrose	P,S,T	Injection	100 ml
	18.6 Ovu	lation Inducers	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Clomiphene citrate	Т	Tablets	50mg, 100mg
	18.7 P	rogestogens	
Medicines	Category	Route of Administration/ Dosage Form	Strengths

	P,S,T	Injection	250 IU, 500 IU		
Polyvalent Antisnake Venom	P,S,T	Injection	10 ml		
Anti-D immunoglobin (human)	S, T	Injection	300 µg		
Drugs	Category	Route of Administration/ Dosage Form	Strengths		
	19.2: Sera a	nd immunoglobins			
Tuberculin, Purified Protein derivative	P,S,T	Injection	1 TU, 5 TU		
Drugs	Category	Route of Administration/ Dosage Form	Strengths		
	19.1: Dia	agnostic agents			
	Section: 19	Immunologicals			
lodine	S,T	Solution	8 mg / 5 ml		
Levothyroxine	P,S,T	Tablets	50µg, 100 µg		
			5040		
Carbimazole	P,S,T	Tablets	10mg		
<u> </u>			5mg,		
MedicinesCategoryRoute ofMedicinesCategoryAdministration /StrengtDosage FormStrengtStrengt					
1	18.8 Thyroid and	d antithyroid medicines			
Norethisterone	P,S,T	Tablets	5mg		
Progesterone Acetate	P,S,T	Tablets	10mg		
Medroxy	рст	Tablete	5mg,		

Antitetanus Human immunoglobin				
Diphtheria Antitoxin	S, T	Injection	10,000 IU	
Rabies immunoglobin	P,S,T	Injection	150 IU / ml	
	19.	3: Vaccines		
	19.3.1: For U	niversal Immunisation		
Drugs	Category	Route of Administration/ Dosage Form	Strengths	
B.C.G Vaccine	P,S,T	Injection		
D.P.T Vaccine	P,S,T	Injection		
Hepatitis B Vaccine	P,S,T	Injection		
Measles Vaccine	P,S,T	Injection		
Oral Poliomyelitis vaccine (LA)	P,S,T	Solution		
1	9.3.2: For Spec	ific Group of Individuals		
Drugs	Category	Route of Administration/ Dosage Form	Strengths	
Rabies Vaccine	P,S,T	Injection		
Tetanus Toxoid	P,S,T	Injection		
Section: 20	– Muscle Rel	axants (Peripherally a	cting) and	
Cholinesterase Inhibitors				
Drugs	Category	Route of Administration/ Dosage Form	Strengths	
Atracurium besylate	S, T	Injection	10 mg / ml	

Neostigmine	S,T	Tablets, Injection	15 mg, 0.5mg/ml	
Pyridostigmine	S, T	Tablets, Injection	60 mg, 1mg/ml	
Succinyl choline chloride	S,T	Injection	50 mg/ml	
Added drugs				
Vecuronium	P,S,T	Injection	2 mg/ml	

Section: 21 – Ophthalmological Preparations					
	21.1	: Anti-infective agents			
Medicines	es Category Route of Administration/ Dosage Stre				
Chloramph enicol	P,S,T	Drops/Ointment	0.4%, 1%		
Ciprofloxa cin Hydrochlor ide	P,S,T	Drops/Ointment	0.3%		
Gentamici n	P,S,T	Drops	0.3%		
Miconazol e	P,S,T	Drops	1%		
Povidone Iodine	S,T	Drops	0.6%		
Sulphacet amide Sodium	P,S,T	Drops	10%,20%		
21.2: Antiinflammaory agents					
Medicines	Category	Route of Administration/ Dosage Form	Strengths		
Prednisolo ne	P,S,T	Drops	0.1%		

Acetate			
Prednisolo ne Sodium Phosphate	P,S,T	Drops	1%
I	21.	3: Local Anaesthetics	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Tetracaine Hydrochlor ide	P,S,T	Drops	0.5%
	21.4: Miotic	s and Antiglucoma medicines	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Acetazola mide	S,T	Tablets	250 mg
Betaxolol Hydrochlor ide	Т	Drops	0.25%, 0.5%
Pilocarpin e	S,T	Drops	2%, 4%
Timolol Maleate	P, S, T	Drops	0.25%, 0.5%,
		21.5: Mydriatics	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Atropine Sulphate	P,S,T	Drops/Ointment	1%
Homatropi ne	P,S,T	Drops	2%
Phenyleph rine	P,S,T	Drops	5%

21.6: Ophthalmic Surgical Aids			
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Methyl Cellulose	т	Injection	2%

Section: 22 – Oxytocics and Antioxytocics				
	22.1: O	kytocics		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Methyl Ergometrine	P,S,T	Tablets Injection	0.125mg 0.2mg/ml	
Mifepristone	т	Tablets	200mg	
Oxytocin	S,T	Injection	5 IU/ ml, 10IU/ml	

Added medicines				
Misoprostol	Т	Tablets	100ug	
	22.2: Ant	ioxytocics		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Terbutaline Sulphate	S,T	Tablets Injection	2.5 mg 0.5 mg/ml	
Added Medicines				
Nifedipine	S,T	Tablets	10 mg	

Betamethason	e P,S,	T Injection	4 mg/ml						
	Section: 2	3 – Peritoneal Dialysis S	olution						
Medicines	Category	Route of Administration/ Dosage Form	Strengths						
Intraperitoneal Dialysis Solution	т		4Of approximate composition						
Section: 24 – Psychotherapeutic Medicines									
	24.1: Medicin	es used in Psychotic Disorde	ers						
Medicines	Category	Route of Administration/ Dosage Form	Strengths						
		Tablets	25 mg, 50mg, 100mg						
Chlorpromazine hydrochloride	P,S,T	Syrup	25mg/5ml						
,		Injection	25mg/ml						
Haloperidol	S, T	Injection	5mg/ml						
		Added medicines							
Olanzapine	Т	Tablets	5mg,10mg						
	24.2: Medic	ines used in mood disorders							
	24.2.1: Medicin	es used in Depressive disord	lers						
Medicines	Category	Route of Administration/ Dosage Form	Strengths						
Amitriptyline	P,S,T	Tablets	25 mg						
Fluoxetine hydrochloride	P,S,T	Capsules	20 mg						
Imipramine	Imipramine P,S,T Tablets		25 mg, 75 mg						
	24.2.2: Medic	ines used in Bipolar disorde	rs						
Medicines	Category	Route of Administration/ Dosage Form	Strengths						
Lithium Carbonate	Т	Tablets 300 mg							

			Added Me	edicines			
Sodium Valproate	Ρ,5	S,T		Tablets		200 mg, 500mg	
24.3: Me	edicines u	sed for	Generaliz	zed Anxiety and Sle	ep Disoro	ders	
Medicines Category Route of Administration/ Dosage Form						rengths	
Alprazolam	P,S	6,T		Tablets		.25 mg,).5 mg	
Diazepam	P,S	6,T		Tablets		2 mg, 5mg	
24.4: Medicir	nes used f	for obse	essive cor	mpulsive disorders	and panio	c attacks	
Medicines Category Route of Administration/ Dosage Form Strengths							
			Added Me	edicines			
Fluoxetine hydrochloride	P,S	,T		Capsules		20 mg	
Sectio	on: 25 –	Medic	ines act	ing on the respir	atory tr	act	
		25.1: A	Antiasthm	atic medicines			
Medicines	5	Cat	egory	Route of Adminis Dosage For		Strengths	
Beclomethas Dipropiona		F	P,S,T	Inhalatio	n	50 μg, 250μg/dose	
Hydrocortiso sodium succi		F	P,S,T	Injection		100 mg, 200mg, 400 mg	
Salbutam sulphate		Tablets Syrup Inhalatio	n	2mg, 4mg 2mg/5ml 100µg/dose			
			Added M	edicines		<u>I</u>	

lpratropium bromide	P,S,T		Inhalation	2	20µg/metere d dose	
	25.2: Ant	itus	ssives			
Medicines	Category	I	Route of Administratio Dosage Form	n/ Strengths		
Codeine phosphate	S,T		Tablets Syrup		10mg 15mg/ 5ml	
Dextromethorphan	P,S,T		Tablets		30mg	
Section: 26 – Soluti	ons correcting	g w	vater, electrolyte ar	nd ac	id-base	
	disturb	ban	ices			
	26.1:	Ora	al			
Medicines	Category		Route of Administrat Dosage Form	tion/	Strengths	
Oral Rehydration Salts	P, S, T	Powder for Solutio			As per IP	
	26.2: Pa	ren	teral			
Medicines	Category		Route of Administration/ Dosage Form	S	trengths	
Glucose	P, S, T		Injection		6 isotonic, 0%, 15%.	
Glucose with sodium chloride	P, S, T		Injection	5% + 0.9%		
Normal Saline	P, S, T		Injection	0.9%		
N/2 Saline	S, T		Injection			
N/5 Saline	S, T		Injection			
Potassium Chloride	P, S, T		Injection	11.2% Sol.		
Ringer Lactate	P, S, T		Injection	A	As per IP	
Sodium	P, S, T		Injection	A	s per IP	

Bicarbonate			
	26.3: Misc	cellaneous	
Medicines	Category	Route of Administration/ Dosage Form	Strengths
Water for Injection	P, S, T	Injection	2 ml, 5 ml, 10 ml

Secti	on: 27 – Vita	mins and Minerals		
Medicines	Category	Route of Administration/ Dosage Form	Strengths	
Ascorbic Acid	P,S,T	Tablets	100 mg, 500 mg	
Calcium carbonate	P,S,T	Tablets	250 mg, 500 mg	
Multivitamins (As per Schedule V of Drugs and Cosmetics Rules)	P,S,T	Tablets		
Nicotinamide	P,S,T	Tablets	50 mg	
Pyridoxine	P,S,T	Tablets	25 mg	
Riboflavin	P,S,T	Tablets	5 mg	
Thiamine	P,S,T	Tablets	100 mg	
Vitamin A	P,S,T	Tablets Capsules	5000 IU, 50000 IU, 100000 IU,	
		Injection	50000 IU/ml	
Vitamin D (Ergocalciferol)	P,S,T	Capsules	0.25 mg, 1 mg	
	Added N	ledicines		
Calcium gluconate	P,S,T	Injection	100mg/ml in 10 ml ampoule	

SCHEDULE-II

FORM - I

PROFORMA FOR APPLICATION FOR PRICE FIXATION / REVISION OF A NEW DRUG FORMULATION RELATED TO NLEM FORMULATION

(See paragraphs 2(u),5,7,8,9,15)

- 1. Name of the formulation:
- 2. Name and address of the manufacturer/importer :
- 3. Name of the Marketing Company, if any:
- 4. Composition as per label claimed and approved by Drug Control Authorities:
- 5. Drugs Control Authority Permission Number and Date (copy to be enclosed):
- 6. Date of commencement of production / import:
- 7. Type of formulation (Tablets/ Capsules/ Syrup/ Injection/ Ointment/ Powder etc.):
- 8. Size of packs (10's/ 100's/ 1 ml/ 2 ml/ 10 ml/ 5 gms/ 10 gms etc.)
- 9. Therapeutic category/ use of the formulation.
- 10. The Retail Price claimed for approval
- 11. Reason for submission of application for price fixation / revision.
- 12. Any other information relevant to product and its process of manufacturing/ packaging/ distribution.

The information furnished above is correct and true to the best of my knowledge and belief.

Place:

Authorized Signatory:

Name: Designation:

Date:

SCHEDULE-II FORM - II

PROFORMA FOR SUBMISSION OF REVISED-PRICES FOR SCHEDULED FORMULATIONS

(See paragraph 16)

- 1. Name and address of the manufacturer / importer / distributor.
- 2. Name and address of the marketing company, if any.

SI. No.	Name of the Product (Formulation and its dosage forms)	Composition of scheduled formulation/ new drug	Pack Size	WPI change w.r.t preceeding Year	Price to retailer (incl. of E.D.) (Rs.)		(Rs.) (Rs.)		Ceiling Price (Notified) (Rs.)	Effective Batch No. and date
				Tear	Pre-Revised	Revised	Pre-Revised	Revised		
(1)	(2)	(3)	(4)	(5)	(6)	(7)	(8)	(9)	(10)	(11)
	Scheduled Formulations									
	Own Manufactured Formulations									
	Purchased/Imported Formulations									

<u>Notes:-</u> In case of purchased formulation, name of the manufacturer shall be indicated.

The information furnished above is correct and true to the best of my knowledge and belief.

Place:

Date:

Authorised Signatory: Name: Designation:

SCHEDULE-II FORM – III

PROFORMA FOR QUARTERLY RETURN IN RESPECT OF PRODUCTION/IMPORT AND SALE OF NLEM DRUGS (See paragraphs 21(1))

- 1. Name and address of the manufacturer/importer:
- 2. Name and address of marketing company, if any:
- 3. Details of production/import and sale for the Quarter of a Year:

TABLE-A

	ne of the heduled	Composition /Strength	Dosage Form	Unit(No/ kg/Ltr)	Production/Import Level				Domestic S	Sale				
For	mulation				Previous		Curren	t Year		Previous		Curre	nt Year	
					Year	1 st	2 nd	3 rd	4 th	Year	1 st	2 nd	3 rd	4 th
						Quarter	Quarter	Quarter	Quarter		Quarter	Quarter	Quarter	Quarter
	(1)	(2)	(3)	(4)	(5)	(6)	(7)	(8)	(9)	(10)	(11)	(12)	(13)	(14)

TABLE-B

Name of the	Unit	Installed	Production/Import Level				Domestic S	estic Sale				
Bulk Drug/API	(Kg/	Capacity										
used in	Ltr)		Previous		Curren	t Year		Previous		Currer	nt Year	
Scheduled	-		Year	Year				Year				
Formulation				1 st	2 nd	3 rd	4 th		1 st Quarter	2 nd Quarter	3 rd Quarter	4 th Quarter
				Quarter	Quarter	Quarter	Quarter					
(1)	(2)	(3)	(4)	(5)	(6)	(7)	(8)	(9)	(10)	(11)	(12)	(13)

Constraints, if any:

Note: (1) Production outsourced / carried out on job work basis should also be included

The information furnished above is correct and true to the best of my knowledge and belief.

Place:

Date:

Authorised Signatory: Name: Designation:

SCHEDULE-II

FORM - IV

PROFORMA FOR SUBMISSION OF THE DETAILS IN RESPECT OF DISCONTINUATION OF THE PRODUCTION AND/ OR IMPORT OF SCHEDULED FORMULATION

(See paragraphs 21(2))

- 1. Name of the formulation:
- 2. Name and address of the manufacturer/importer :
- 3. Name of the Marketing Company, if any:
- 4. Composition as per label claimed and approved by Drug Control Authorities:
- 5. Drugs Control Authority Permission Number and Date (copy to be enclosed):
- 6. Celling Price and date of notification:
- 7. Existing maximum retail price (MRP) and its effective date:
- 8. Therapeutic category as per NLEM:
- 9. Date of commencement of production / import
- 10. Proposed date of discontinuation:
- 11. Reasons for discontinuation of production / import:
- 12. Year-wise Production/Import during the last 5 years including current year
- 13. Year-wise sale during the last 5 years including current year
- 14. Whether any new drug as defined under Proviso of Definition of "New Drug" under DPCO, 2013 has been launched or intended to be launched. If so, the details thereof:
- 15. Any other information relevant to discontinuation of scheduled formulation:

Authorized Signatory:

Name:

Designation:

Place:

Date:

SCHEDULE-II FORM - V

PROFORMA FOR PRICE LIST

(See paragraphs 2(x),24,25,26)

- 1. Name and address of the manufacturer / importer / distributor.
- 2. Name and address of the marketing company, if any.

TABLE-A

Sl. No.	Name of the Product (Formulation and its dosage forms)	Composition approved by Drug Control Authorities	Pack Size	Price to retailer (incl. of E.D.) (Rs.)	Maximum Retail Price (incl. of E.D.&Taxes) (Rs.)
(1)	(2)	(3)	(4)	(5)	(6)
	Scheduled Formulations				
	Own Manufactured Formulations				
	Purchased/Imported Formulations				

TABLE-B

Sl. No.	Name of the Product (Formulation and its dosage forms)	Composition approved by Drug Control Authorities	Pack Size	Price to retailer (incl. of E.D.) (Rs.)	Maximum Retail Price (incl. of E.D.&Taxes) (Rs.)
(1)	(2)	(3)	(4)	(1101 01 2.0.) (13.)	(6)
	Non-Scheduled Formulations	· · · ·			
	Own Manufactured Formulations				
	Purchased/Imported Formulations				

Notes:- In case of purchased formulation, name of the manufacturer shall be indicated.

The information furnished above is correct and true to the best of my knowledge and belief.

Authorised Signatory: Name: Designation:

Place: Date:

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