

3. **List of new drugs approved in the year 2019 till date**

S.No.	Name of drug	indication	Date of issue
1	Fenspiride hydrochloride film coated extended release tablet 80 mg and fenspiride hydrochloride bulk	1. Acute Rhinosinusitis 2. moderate persistent asthma as an add- on therapy	04-02-2019
2	Bilastine tablets 20 mg and Bilastine bulk	For symptomatic treatment of allergic rhino-conjunctivitis (seasonal and perennial) and urticaria in adults	06-02-2019
3	Iguratimod film coated tablets 25 mg and iguratimod bulk	For the treatment of active rheumatoid arthritis symptoms	18-02-2019
4.	Fingolimod Capsules 0.5 mg and Fingolimod hydrochloride bulk	For the treatment of patients with relapsing forms of multiple sclerosis (MS) to reduce the frequency of clinical exacerbations and to delay the accumulation of physical disability.	25.03.2019
5	Remogliflozin etabonate bulk and Remogliflozin etabonate film coated tablets 100 mg	Indicated in adults aged 18 years and older with type 2 diabetes mellitus to improve glycemic control as:- <ul style="list-style-type: none"> • Monotherapy when diet and exercise alone do not provide adequate glyceamic control. • Add on therapy with metformin together with diet and exercise, when these do not provide adequate glyceamic control. 	26.04.2019

05	45	Application for registration of bioavailability and bioequivalence study centre	5,00,000
07	47	Reconsideration of application for Registration of bioavailability and bio-equivalence study centre	1,00,000
08	52	Application for permission to manufacture new drugs or investigational new drugs for clinical trial or bioavailability or bioequivalence study	5000 per product
09	53	Reconsideration of application to manufacture new drugs or investigational new drugs for clinical trial or bioavailability or bioequivalence study	2000 per product
10	59	Application for permission to manufacture unapproved active pharmaceutical ingredient for development of formulation for test or analysis or clinical trial or bioavailability or bioequivalence study	5000 per product

11	60	Reconsideration of permission to Manufacture unapproved active pharmaceutical ingredient for development of formulation for test or analysis or clinical trial or bioavailability or bioequivalence study	2000
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12	67	Application for import of new drugs or investigational new drugs for clinical trial or bioavailability or bioequivalence study or for examination, test and analysis	5000 per product
13	68	Reconsideration of application for Import of new drugs or investigational new drugs for clinical trial or bioavailability or bioequivalence study or for examination, test and analysis	1000
14	75	Application for permission to import new drug (Finished Formulation) for marketing	5,00,000
15		Application for permission to import new Drug (Finished Formulation) already approved in the country for marketing	2,00,000
16		Application for permission to import new drug (Active Pharmaceutical Ingredient) for marketing	5,00,000

17		Application for permission to import new drug (Active Pharmaceutical Ingredient) already approved in the country for marketing	2,00,000
18		Application for permission to import approved new drug for new claims, new indication or new dosage form or new route of administration or new strength for marketing	3,00,000
19		Application for permission to import fixed dose combination having one or more of the ingredients as unapproved new molecules for marketing	5,00,000
20		Application for permission to import fixed Dose combination having approved ingredients for marketing	4,00,000
21		Application for permission to import fixed dose combination already approved for marketing	2,00,000
22		Application for permission to import fixed dose combination for new claims, new indication or new dosage form or new route of administration or new strength for marketing	3,00,000
23	76	Reconsideration of application for permission to import new drug for marketing	50,000
24		Application for permission to manufacture new drug (Finished Formulation or Active Pharmaceutical Ingredient) for sale or distribution	5,00,000
25		Application for permission to manufacture new drug (Active Pharmaceutical Ingredient) already approved in the country for sale or distribution	2,00,000
26	80	Application for permission to manufacture new drug (Finished Formulation) for sale or distribution	5,00,000
27		Application for permission to manufacture new drug (Finished Formulation) already approved in the country for sale or distribution	2,00,000
28		Application for permission to manufacture new drug (Active Pharmaceutical Ingredient) for sale or distribution	5,00,000