# **RESEARCH ARTICLE**

# A study of new drug approval pattern of a Southeast Asian developed country from 2017 to 2021

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# ABSTRACT

**Background:** The pattern of new drug approval is changing across the world as shown by the study using Center for Drug Evaluation and Research and European Medicines Agency data in US and UK with more drug approval for anti-cancer and immunomodulator drugs. There is a need to generate similar database for developed South East Asian countries too. **Aims and Objectives:** This study was conducted for one such country- Singapore for the new drug approval pattern of last 5 years (2017–2021). **Materials and Methods:** This was a pharmacoepidemiological study, in which government drug regulatory website data available in public domain was searched. The new drug approval data were classified according to active ingredient, drug approval date, new drug application category, indication of drugs, and World Health Organization Anatomic Thoracic Classification. **Results:** In this study, 418 new drug approvals were found in last 5 years in Singapore. From this maximum, drug approvals were given to anti-neoplastic and immunomodulator category drugs. In anti-neoplastic category new drugs approval few examples were Trastuzumab deruxtecan and Tucatinib for breast cancer therapy and Tepotinib and Capmatinib for non-small cell lung cancer therapy. **Conclusion:** This study shows that drug development in anti-cancer drug and immunomodulator is significant in Singapore. This trend is quite matching with other country such as US and UK.

KEY WORDS: New Drug Approval; Singapore; Health Science Authority

#### INTRODUCTION

New drug development is crucial for scientific progress of any country. Different countries have different unmet needs for new drug development that should be fulfilled by pharmaceutical industry. According to New drug and clinical trial rules 2019, new drug is defined as any active pharmaceutical ingredient or phytopharmaceutical drug which is not approved in that country or any approved drug having modified or new claim for indication, route of administration,

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dosage, and dosage form. As well as new drug is any new fixed dose combination of two approved drug or modified or new claim for indication, route of administration, dosage, and dosage form. It also includes a modified or sustained release form of a drug or novel drug delivery system of any approved drug. As well as a vaccine, r-DNA derived product, living modified organism, monoclonal anti-body, stem cell derived product, gene therapeutic product or xenografts, intended to be used as drug.<sup>[1]</sup> The Singapore have Clinical Trial Guidance material that provides guidance to manufacturers, importers, and suppliers on the regulatory requirements relating to the import and supply of clinical research materials.<sup>[2]</sup>

Every country has the regulatory authority to check the authentic information regarding new drug approval and such information is available in public domain, for example, for US authentic information is available on Center for Drug

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Evaluation and Research (CDER) website, established in 1987 under the US Food and Drug Administration (FDA).<sup>[3]</sup> Similarly for European union authentic data available on European medical agency (EMA) website<sup>[4]</sup> and for India it is Central drug standard control organization (CDSCO).<sup>[5]</sup>

The pharmaceutical sector and research and development remain strong pillar in Singapore economy. It contributes 5% of Singapore Gross Domestic Product (GDP). In 2021, pharmaceutical industry had contributed about 10.5 billion Singapore dollars (SGD) to the GDP which is significant increase from the previous year, that is, 8.77 billion SGD. The government of Singapore launches biomedical science initiative in 2000, that has amplified the pharmaceutical industry in the country.<sup>[6]</sup> Health science authority (HSA) is government authority for new drug approvals in Singapore. HSA had been formed on April 1, 2001, under the ministry of health and government of Singapore. This agency comprises three professional groups: the Health Products Regulation Group; Blood Services Group, and Applied Sciences Group. The Health Products Regulation Group ensures that medicines, innovative therapeutics, medical devices, and health-related products are wisely regulated and meet appropriate safety, quality, and efficacy standards. Its international partners include agencies such as the US FDA, Health Canada, Swissmedic, the Australian Therapeutic Goods Administration and China FDA.<sup>[7]</sup>

As a medical practitioner, it is important to know new drug approval pattern of the country so they can prescribe the relevant new medicine to the patient. As per a study, the drug approval process in India has more lag time as compared to developed countries such as US and UK which could lead to delay of few more years for the new drug to come out in market for the beneficiaries.<sup>[8]</sup> Study by Konwar et al. mentioned that absolute drug lag of CDSCO was 19 and 18 relative to the US FDA and Japan Pharmaceuticals and Medical Devices Agency (PMDA), respectively. The relative drug lag for the CDSCO vis a vis the US FDA, EMA, and PMDA was 43.2, 25.6, and 30.3 months, respectively.<sup>[9]</sup> For a Pharmaceutical industry, it is important to know new drug approval pattern of that particular country to remain relevant with the current trend in the pharmaceutical market. So study of drug approval pattern is important for various stakeholders of health-care delivery. Different studies conducted across the world have shown changing pattern of new drug approval with more number of anti-cancer and biologics being approved.<sup>[10]</sup>

Such studies have mainly been done in US FDA using CDER data, Japan using PMDA, European countries using EMA website data, and India using CDSCO. But similar baseline study is lacking in south east Asian developed country such as Singapore which formed the basis for our study. With Singapore becoming one of the fast growing countries in pharmaceutical engineering and export output of pharmaceutical products, it becomes imperative that a database is generated for its new drug approval pattern.

#### MATERIALS AND METHODS

This was a pharmacoepidemiological retrospective observational type of study in which drug approval pattern of new drugs in Singapore was studied. Data available on official website of Singapore government health authority were searched.<sup>[7]</sup> The data were compiled for the duration of January 2017–December2021.

Data on the website was available in form of monthly approval which was then entered to Libre office calc spread sheet. The new approved drugs were classified based on different parameters such as active ingredient, drug approval date, new drug application (NDA) category, Indication of drugs, and the World Health Organization (WHO) Anatomic Thoracic Classification (ATC).

In Singapore any new drug or biologics approval is given through NDA. There are three categories in NDA.

- 1. NDA category one for first strength of new chemical or biological entity which is not previously registered in Singapore
- 2. NDA category two applicable to first strength of product previously registered in Singapore having new combination, new dosage, new formulation, new presentation, new route of administration, new indication, new dosage recommendation, and new patient population included
- 3. NDA category three for any new subsequent strength of chemical or biological entity that was previously registered in Singapore under NDA 1 or 2.

A biosimilar product applies to a biological product demonstrated to be similar in physicochemical characteristics, biological activity, safety, and efficacy to an existing registered biological product. Biosimilar products are required to be submitted via NDA-2 or NDA-3. Here NDA 2 is for the first strength of a bio similar product with the same dosage form and route of administration as the Singapore reference biological product. And NDA 3 is for subsequent strengths of a biosimilar product that has been registered or has been submitted as an NDA-2. The product name, dosage form, indication, dosing regimen, and patient population should be the same as that for the NDA-2 submission. Hence, the data were classified in three types of NDA categories as described above.

In the WHO ATC classification, the active ingredients were classified in fourteen main anatomical or pharmacological groups. Active ingredient is also classified according to hierarchy in five different levels - from which we have used level one classification with fourteen main groups in this study. The descriptive analysis of data was done using Libre office calc. As the new drug approval data are freely available in public domain of Government website, so ethical permission is not required.

# RESULTS

There were total 418 new drugs which were approved in Singapore during period 2017–2021, with an average of 83.6 new drugs approved annually in last 5 years and 29 annual new drug approval in NDA 1 category [Figure 1].

On classification of all the drugs according to the WHO ATC classification, it was found that maximum number of new drugs approvals 129 (30.86%) were from anti-neoplastic and immunomodulating agent category which was followed by 47 (11.24%) new drug approvals in anti-infective category and 44 (10.53%) new drug approvals in nervous system category [Figure 2].

The data were also classified according to NDA category and it was found that maximum number of new drugs belonged to NDA-2 category- 178 (42.58%) new drug approval followed by NDA-1- 145 (34.69%) new drug approval and NDA-3 category- 95(22.73%) [Figure 3]. From these new approved drugs, 27 were new biological entity and 91 were new chemical entity.

There were 11 biosimilars approved in last 5 years. There were 47 new dosage forms and 36 new combinations approved of already approved drugs in Singapore under NDA 2 category. There were 95 approvals given to new subsequent strength of already approved new chemical/biological/Biosmilar entity under NDA 3 category.

We had classified new chemical entity/new biological entity approval in Singapore in last 5 years. Few example of new drug approval were – (a) breast cancer (Trastuzumab deruxtecan, Tucatinib), (b) non-small cell lung cancer (NSCLC) (Tepotinib, Capmatinib), (c) SARS-CoV-2 (Tozinameran, Remdesivir), (d) HIV 1 (Doravirine), (e) Complecated Intraabdominal infection (Eravacycline, Avibactam sodium), (e) Depression (Esketamine, Brexpiprazole), (f) Asthma (Erdosteine, Benralizumab), and (g) Pulmonary Arterial Hypertension (Selexipag) [Table 1].

#### DISCUSSION

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This retrospective observational study was conducted to know new drug approval pattern in Singapore in last 5 years. In our study, 83.6 is average annual new drug approval found in Singapore and 29 annual new drug approvals in NDA 1 category. In comparison CDER, USA approved 50 new drug in 2021 and 53 new drug in 2020 with last 5 year average approval of 51 new drugs per year.<sup>[11]</sup>



Figure 1: Annual new drug approval in Singapore in the last 5 years



**Figure 2:** ATC Classification of new drug approval in the last 5 years. ATC Group: (a) Alimentary tract and metabolism; (b) Blood and blood forming organs; (c) Cardiovascular system; (d) Dermatologicals; (g) Genitourinary system and sex hormones; (h) Systemic hormonal preparations, excluding sex hormones and insulins; (j) Anti-infective for systemic use; (l) Anti-neoplastic and immunomodulating agents; (m) Musculo-skeletal system; (n) Nervous system; (p) Anti-parasitic products, insecticides and repellents; (r) Respiratory system; (s) Sensory organs; (v) Various



Figure 3: Classification of new drug approval according to NDA category

We found upwards trend of annual new drug approval with each year in this study except during COVID-19 pandemic which

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Cytomegalo virus infection (CMV) Letermovir	vtomegalo virus infection (CMV)
SARS-CoV-2 Tozinameran, Remdesivir	ARS-CoV-2
HZ, PHN [Antigen] Recombinant Varicella Zoster Virus glycoprotein E (gE)	Z, PHN
Influenza Pandemic Influenza Virus Type A, inactivated and disrupted , Baloxavir marboxil	nfluenza
Botulism Botulism antitoxin (equine) serotypes A, B, C, D, E, F, G	otulism
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Depression Esketamine. Brexpiprazole	Pepression
Schizophrenia Brexpiprazole. Cariprazine	chizophrenia
Tardive Diskinesia Valbenazine ditosvlate	ardive Diskinesia
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Table 1: (Continued)		
Few common conditions/disease	New drug approval in last 5 years	
Parkinson's disease (PD)	Rasagiline	
Common respiratory system disease		
Chronic obstructive pulmonary disease (COPD)	Revefenacin, Erdosteine, Aclidinium	
Asthma	Erdosteine, Benralizumab, Mepolizumab	
Common CVS diseases		
Dyslipidemia	Inclisiran sodium, Pitavastatin, Evolocumab, Alirocumab	
CHF	Vericiguat, Finerenone	
Myocardial Infarction	Finerenone	
Essential Hypertension	Fimasartan Potassium	
Pulmonary Arterial Hypertension (PAH)	Selexipag	
Amyloidosis	Tafamidis meglumine	
Erectile dysfunction	Avafanil	
Hyperkalemia	Sodium zirconium cyclosilicate	
Hyprnatremia	Tolvaptan	
Common Blood diseases		
Methemoglobinemia	Methylthioninium chloride trihydrate (methylene blue)	
Hemophilia A	Turoctocog Alfa, Rurioctocog alfa pegol [Antihemophilic Factor (Recombinant), PEGylated], Lonoctocog alfa , Emicizumab	
Non valvular atrial fibrillation (NVAF), Deep vein thrombosis (DVT), Pulmonary embolism (PE)	Edoxaban Tosilate	
Febrile neutropenia	Lipegfilgrastim	
Congenital hypo- or afibrinogenaemia	Human Fibrinogen	
Iron deficiency anemia	Ferric Derisomaltose 417 mg/ml (Elementary Iron 100 mg/ml)	
Hemophilia B	Albutrepenonacog alfa	
Common Hormonal diseases		
Diabetes mellitus	Semaglutide, Ertugliflozin, Insulin Degludec	
Osteoporosis	Romosozumab	
X- Linked hypophosphetemia (XLH)	Burosumab	
Medical termination of pregnancy (MTP)	Mifepristone	
Vulvular vaginal atrophy (VVA)	Ospemifene	
Controlled ovarian stimulation	Follitropin delta	
Secondary Hyperparathyroidism (SHPT)	Etelcalcetide	
Common Opthalmic disease		
Glaucoma	Latanoprostene Bunod, Omidenepag Isopropyl, Ripasudil Hydrochloride Hydrate,	
Common skin disease		
Plaque psoriasis	Risankizumab, Ixekizumab, Guselkumab, Apremilast	
Atopic dermatitis	Dupilimumab	
Common GIT Disease		
Hepatic encephalopathy	L-ornithine-L-aspartate	
Peptic ulcer, GERD	Vonoprazan	
Constipation	Lubiprostone	
Morning sickness	Doxylamine Succinate+ Pyridoxine Hydrochloride	
Motion sickness	Cyclizine Hydrochloride	
Common Inflammatory condition		
Rheumatoid arthritis	Upadacitinib, Sarilumab, Baricitinib	
Common diagnostic agents		
Scintigraphy	Sodium pyrophosphate decahydrate	
Radiopharmceuticals	Lutetium (177Lu) chloride, Lutetium (177Lu) oxodotreotide, Ioflupane [123I],	

Table 1: (Continued)		
Few common conditions/disease	New drug approval in last 5 years	
PET imaging	Flutemetamol[18F]	
Other conditions		
Fabry's disease	Agalsidase alfa	
Ttype 1 Gaucher's disease	Velaglucerase alfa	
Submental Fat	Deoxycholic acid	

NSCLC: Non-small cell lung cancer, SHPT: Secondary Hyperparathyroidism, VVA: Vulvular vaginal atrophy, MTP: Medical termination of pregnancy, XLH: X- Linked hypophosphatemia, NVAF: Non valvular atrial fibrillation, DVT Deep vein thrombosis, PE: Pulmonary embolism, PAH: Pulmonary Arterial Hypertension, COPD: Chronic obstructive pulmonary disease, RCC: Renal cell carcinoma, SMA: Spinal Muscular atrophy NMOSD :Neuromyelitis optica spectrum disorders

has affected new drug approval all over the world significantly. During the pandemic period, majority countries including Singapore had given conditional Emergency Use Authorization to drugs which were useful in COVID management.

Observations from this study showed that maximum numbers of drugs (30.86%) were approved in ATC Class L which contains Anti neoplastic and immunomodulating agents. In one study by Kataria *et al.* (2016) from India showed anti-neoplastic and immunomodulating drug approval rate of 36.26%.<sup>[12]</sup> Similar other studies mentioned higher anticancer drug approvals recently.<sup>[13-16]</sup>

Maximum anti-cancer drugs approval in last 5 years in Singapore were for breast cancer and NSCLC. Tastuzumab and Tucatinib for breast cancer and Tepotinib and Capmatinib for NSCLC were few example of new anti-caner drug approvals.

Now a day, number of new drugs approval is increasing with each year.<sup>[11]</sup> Reasons cited by few studies for this trend were:

- a. Increase in number of NDA by pharmaceutical companies
- b. Significant growth in anti-cancer drugs development in the past decade
- c. Faster drug approval by few countries like US, UK in the form of expedited drug review program (fast track review/break through therapy review/priority review/ accelerated approval review)<sup>[17]</sup>
- d. Incentive to Orphan drug development by many countries.<sup>[11]</sup>

Our study is first study of new drug approval pattern in Singapore and generated baseline data. This study analyzed data of last 5 years. Therefore, to know the trend of new drug approval over larger period, further future study with longer duration can be planned. Drug recalls have not been considered for analysis in this study.

#### CONCLUSION

This was the first study on new drug approval pattern in developed South East Asian country Singapore in last 5 year. In this study, it was found that in Singapore maximum number of new drug approvals was in anti-neoplastic and immunomodulator category. This trend is quite matching with other country such as US and UK. In last 5 years, number of drug approval given for breast cancer, NSCLC was maximum. That shows drug development in anti-cancer drug and immunomodulator is significant. This study provides baseline data for future larger scale studies.

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