


Chapter 9

Role of Artificial Intelligence and Machine Learning in Drug Discovery and Drug Repurposing

Sameer Quazi

 <https://orcid.org/0000-0002-1258-4088>
GenLab Biosolutions Private Limited, India

Zarish Fatima

University College Lahore, Pakistan

ABSTRACT

Drug designing and repurposing is the most important field in the pharmaceutical industries and biomedical sciences. Because the challenges caused by drug such as low retention time, sensitivity can affect the efficacy of developmental process. As AI or ML has proven to be a potential activity in the health and biomedical sciences and from previous research it has found that AI can learn new data and transform it into the useful knowledge. So, in field of pharmacology, the aim is to design more efficient and novel vaccines using this method which is also cost effective. The underlying fact is to predict the molecular mechanism and structure for increased likelihood of developing new drugs. Clinical, electronic, and high-resolution imaging datasets can be used as inputs to aid the drug development niche. Moreover, the use of comprehensive target activity has been performed for repurposing a drug molecule by extending target profiles of drugs which also include off targets with therapeutic potential providing a new indication.

DOI: 10.4018/978-1-6684-6523-3.ch009

INTRODUCTION

Over the last few years, an immense progress has made in the fields of artificial intelligence, machine learning and bioinformatics and more research is needed for understand the data in biological science and related problems. Bioinformatics is a subdivision of science that involves the analysis of biological data using mathematical principles, statistical tools and certain algorithms in addition to computational approaches (Ezziane, 2006). Artificial intelligence is the ability to solve various problems related to human intelligence and in turn the simulation of these intelligence processes using computer systems or software. It involves machine learning that allows one to perform all of these tasks based on its training (Narayanan et al., 2002). Basic or structural bioinformatics tools use artificial intelligence and machine intelligence for the design of drugs and repurposing various novel compounds against many diseases such as cancer, neural inflammation and others using the silica approach by applying certain tools with principles of artificial intelligence and machine learning. Bioinformatics has been used to analyze data and logical conclusions. The huge amount of data obtained from whole genome sequencing projects and bioinformatics is used for the annotation of biological data in meaningful ways (Nicolas, 2020). Similarly a large collection of problems has been solved by combining the knowledge and abilities of artificial intelligence with bioinformatics approaches for the prediction of genes, studies involving the study of protein interactions, computational systems for drug design, repurposing drugs for better efficiency, next generation sequencing and development of other software. Therefore, both artificial intelligence and machine learning have useful applications in the field of bioinformatics. The proficiency of artificial intelligence can be changed by varying input data. Artificial learning is further classified into generalized and applied branches. Both of them are totally different where applied is involved in the use of machines and algorithms while applied to stimulate the data into expressions similar to the thoughts of humans automatically (Bülow, 2021). Current machine learning has overcome some of the barriers and has predicted a huge amount of data, increased computational power and the revival of neural networks. These algorithms can be trained for exploit further data and thus don't require human labor or programmers for data prediction (Mitra, 2005).

If safety testing has been performed then it will display compatibility of dosage with the new indications. Drug repurposing has been discovered by chance on the basis of random testing and exploration. For example in the market, sildenafil citrate was discovered as a hypertensive drug and then repurposed with Pfizer resulting in the formation of a new drug molecule Viagra for the treatment of erectile dysfunction proved by clinical studies directly from its formulation which in turn provided massive sales in addition to additional health benefits. From the last few

14 more pages are available in the full version of this document, which may be purchased using the "Add to Cart" button on the publisher's webpage: www.igi-global.com/chapter/role-of-artificial-intelligence-and-machine-learning-in-drug-discovery-and-drug-repurposing/318557

Related Content

P Colonies of Capacity One and Modularity

Ludk Cienciala, Lucie Ciencialová and Miroslav Langer (2014). *Natural Computing for Simulation and Knowledge Discovery* (pp. 122-138).

www.irma-international.org/chapter/p-colonies-of-capacity-one-and-modularity/80060

Lifelike Self-Replicators

Eleonora Bilotta and Pietro Pantano (2010). *Cellular Automata and Complex Systems: Methods for Modeling Biological Phenomena* (pp. 210-247).

www.irma-international.org/chapter/lifelike-self-replicators/43222

Diversity Conserved Chaotic Artificial Bee Colony Algorithm based Brightness Preserved Histogram Equalization and Contrast Stretching Method

Krishna Gopal Dhal and Sanjoy Das (2015). *International Journal of Natural Computing Research* (pp. 45-73).

www.irma-international.org/article/diversity-conserved-chaotic-artificial-bee-colony-algorithm-based-brightness-preserved-histogram-equalization-and-contrast-stretching-method/164541

Dynamic Modeling and Parameter Identification for Biological Networks: Application to the DNA Damage and Repair Processes

Fortunato Bianconi, Gabriele Lillacci and Paolo Valigi (2011). *Handbook of Research on Computational and Systems Biology: Interdisciplinary Applications* (pp. 478-510).

www.irma-international.org/chapter/dynamic-modeling-parameter-identification-biological/52329

Cell Motility Viewed as Softness

Koji Sawa, Igor Balaž and Tomohiro Shirakawa (2012). *International Journal of Artificial Life Research* (pp. 1-9).

www.irma-international.org/article/cell-motility-viewed-softness/65071