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# Better understanding of biomolecules as drug target: a literature review

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#### ABSTRACT

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#### Keywords:

targeted therapy; GPCR; enzymes; ion channel receptors; DNA Drugs are small biological therapeutic agents that play a crucial role in enhancing human health. They have been widely used in the treatment of various diseases, with conventional medicine being one of the most common approaches. However, conventional medicine has several limitations, including non-specific drug targeting, limited efficacy, low stability, poor absorption rates, and degradation due to enzymatic activity and low pH in the digestive system. Efforts to improve drug targeting have led to the development of molecular-based therapies, particularly through the use of biomolecules as drug targets. Drug targeting aims to optimize therapeutic effects by focusing on specific targets while minimizing side effects. This review article explores research findings from the PubMed database, applying specific inclusion and exclusion criteria. It discusses the types of drugs and biomolecular targets, including G protein-coupled receptors (GPCRs), enzymes, ion channel receptors, and DNA, as well as the mechanisms through which drugs interact with these targets in the treatment of various diseases. Additionally, the advantages and disadvantages of these biomolecular targets are examined. In conclusion, the four biomolecules discussed each present unique characteristics as drug targets, highlighting the importance of carefully selecting appropriate molecules based on an understanding of their functions, properties, and interactions with drugs in the body.

#### ABSTRACT

Obat merupakan agen terapeutik berukuran kecil dan bersifat biologis yang berperan dalam meningkatkan kesehatan manusia. Obat telah digunakan dalam berbagai macam pengobatan, salah satunya dalam pengobatan konvensional. Namun, pengobatan konvensional memiliki beberapa kekurangan seperti target obat yang tidak spesifik, efektivitas obat yang terbatas, stabilitas dan tingkat absorpsi obat yang rendah, serta degradasi obat karena pengaruh enzim dan pH yang rendah pada sistem gastrointestinal. Pengobatan konvensional memberikan solusi penargetan obat menjadi lebih berkembang ke arah molekuler, yaitu dengan menggunakan biomolekul sebagai target obat. Penargetan obat ini merupakan upaya untuk mengoptimalkan efektivitas obat terhadap target secara spesifik dan meminimalisir efek samping yang dihasilkan dari pengobatan konvensional. Metode yang digunakan yaitu eksplorasi hasil penelitian pada situs PubMed sebagai sumber utama dengan kriteria ekslusi dan inklusi. Ulasan artikel ini akan membahas tentang jenis obat dan target biomolekul yang terdiri dari reseptor terkait protein G (GPCR), enzim, reseptor kanal ion dan DNA, mekanisme kerja obat terhadap target obat pada beberapa penyakit, serta kelebihan dan kekurangan dari keempat biomolekul sebagai target obat. Kesimpulan yang diperoleh yaitu keempat biomolekul memiliki spesifitas masing-masing sebagai target obat sehingga perlu diperhatikan pemilihan molekul yang tepat melalui pemahaman fungsi, karakteristik dan interaksi obat terhadap target di dalam tubuh.

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# **INTRODUCTION**

Drugs are chemical compounds that affect the structure or function in the body and useful in the treatment, cure, mitigation, prevention, or diagnosis of a disease. Drugs can be used to cure or alleviate the symptoms of a disease or certain mental medical conditions.<sup>1</sup> The type of treatment often used in the medical world is conventional medicine, which is chemically isolated from natural resources such as plants, fungi, bacteria or animals or synthesized from organics compounds. The drug treatment aims to alleviate the symptoms of a disease by increasing the desired clinical effect or improving the negative side effects of the main drug.<sup>2</sup> Some conventional treatments such as pharmaceutical drugs,<sup>3</sup> surgery, chemotherapy, radiation (cancer treatment).<sup>4</sup> and psychotherapy<sup>4</sup> have been developed since many years ago.

The use of conventional treatment often causes some unwanted side effects. The use of pharmaceutical drugs in high doses (in order to penetrate the plasma membrane of infected cells,9 causing drug resistance (both treatment of bacterial infections and cancer).<sup>10</sup> In addition to the occurrence of drug resistance, there are also obstacles such as the difficulty of access to conventional treatment in several countries.<sup>11</sup> The use of conventional treatment in practice often causes some unwanted side effects, such as the use of pharmaceutical drugs in high doses in order to penetrate the plasma membrane of infected cells,<sup>5</sup> drug resistance in the treatment of bacterial infections and cancer.<sup>6</sup> Conventional drug delivery systems have several shortcomings, such as limited effectiveness, non-specific drug targets, short drug residence time, weak drug biodistribution.<sup>7</sup> Disadvantages in terms of less effective bioavailability such as oral drug distribution can cause drugs to be degraded by enzymes and low pH in the gastrointestinal system,<sup>8</sup>

biocompatibility, biodegradability, and drug distribution into the body.<sup>9,10</sup>

The use of biomolecules as drug targets also enables the development of therapies for complex diseases. Advances in knowledge of the human and molecular biology genome accelerate the process of developing current data to provide a variety of information through bioinformatics on drug-target interactions that support the development of drug targeting.<sup>11</sup> Drug target discovery utilizes two current approaches combining molecular and systemic approaches coupled with in vivo and in vitro validation to assess drug efficacy.12

Identification of drug targets will help determine the relationship between drug activity and phenotype, determine the mechanism of drug action, determine effective targets for therapeutic drugs, resistance mechanisms to possible toxicity in more detail.<sup>13</sup> Currently, small molecular drugs mostly target various molecular components such as enzymes, DNA, receptors, ion channels through the mechanism of action of enzyme inhibitors, ion channel blockers, and inhibitors of the replication process.14 The advantages of drug targeting include increasing drug efficacy, reducing side effects, increasing drug concentration at the desired site, and allowing the use of lower doses but still providing a significant effect in treating.<sup>15</sup>

# MATERIAL AND METHODS

This review uses primary literature retrieved from PubMed with specific keywords, such as GPCR, enzyme, ion channel receptors, and DNA. All articles were selected based on inclusion and exclusion criteria. The inclusion articles criteria were full free access and have been published within the last 10 yr. We also have consideration to exclude the selected articles based on the type of drug target, FDA approval, drug mechanism of action, and clinical trialsbased experiment. Drug efficacy rates above 50% have been used as additional requirements for quality assessment. After a continuous screening process, 40 articles which then represent 10 articles for each drug target were selected.

### RESULTS

A total of 1982 articles were collected from PubMed, after screening only 40 articles were included in this review (TABLE 1).



FIGURE 1. PRISMA flow chart of biomolecules as a drug target

| TADIP 4 I  | · · · · · · · · · · · · · · · · · · · | A A         | 1             |           |           |           |
|------------|---------------------------------------|-------------|---------------|-----------|-----------|-----------|
| TARLE I L1 | ופד הד מדווספ                         | Targeting   | ninmniecilles | and their | mechanism | or action |
| <b>1</b>   | ist of ulugs                          | tui settiis | Dioincluics   | und them  | meenanom  | or action |
|            | 0                                     | 0 0         |               |           |           |           |

| Specific target |   | Drug  | Mechanism  | Indications                 | Ref |
|-----------------|---|---|--|-----------------------------|-----|
| GP              | CR  |   |  |                             |     |
| •               | β2AR<br>(β2-adrenergic<br>receptors)            | LABAs<br>(salmeterol, formoterol,<br>vilanterol, and<br>olodaterol) | Relaxation of bronchial smooth muscle.   | Severe asthma               | 16  |
| •               | β2AR<br>(β2-adrenergic<br>receptors)            | SABAs<br>(Albuterol, levalbuterol)                                  | Relaxation of bronchial smooth muscle.   | Severe asthma               | 17  |
| •               | PTH1<br>(parathyroid hormone<br>receptors)      | Tymlos<br>(abaloparatide)   | Parathyroid type 1 receptor<br>conformation binding<br>selectivity that favors<br>anabolic activity.   | Osteoporosis                | 18  |
| •               | GLP1<br>(glucagon-like peptide 1)               | Semaglutide   | Patients with type 2 diabetes<br>reported a dose-dependent<br>reduction in weight and<br>HbA1c levels after using<br>semaglutide.                          | Diabetes<br>mellitus type 2 | 19  |
| •               | OX <sub>2</sub> R<br>(OX <sub>2</sub> receptor) | Suvorexant  | Maintaining the stability of<br>an extracellular salt bridge<br>network and preventing<br>the transmembrane helix<br>movements required for<br>activation. | Insomnia                    | 20  |

| Specific target   | Drug  | Mechanism  | Indications  | Ref |
|---|---|--|--|-----|
| • CASR  | Parsabiv (etelcalcetide)                                | Reducing serum PTH con-<br>centrations.  | Hyperparathyroidism  | 21  |
| • 5HT2A, 5HT1A  | Rexulti<br>(brexpiprazole)                              | Neurite outgrowth elicited<br>by benzpiprazole is<br>mediated by 5-HT1A and<br>5-HT2A receptors.   | Depression   | 22  |
| • P2Y12   | Kengreal (cangrelor)                                    | Reduce percutaneous<br>coronary intervention-<br>related ischemic<br>complications without<br>increasing major bleeding.   | Percutaneous<br>coronary<br>intervention   | 23  |
| • PAR1<br>(protease-activated<br>receptor-1)                  | Zontivity (vorapaxar)                                   | Reduce the expression of $HSP90\beta$ , the interaction between $HSP90\beta$ and transforming growth factor- $\beta$ (TGF $\beta$ ) receptor II, and the TGF $\beta$ /Smad signaling pathway. These mechanisms also inhibit the stimulation of fibroblasts induced by thrombin in a dose-dependent manner and the signal transduction of JAK2/STAT1/3. | Pulmonary fibrosis   | 24  |
| • NK1R<br>(neurokinin 1 receptor)                             | Varubi (rolapitant)                                     | Rolapitant quickly<br>entered the amygdala,<br>hypothalamus, and<br>neocortex, three brain<br>areas related to reward and<br>anxiety regulation.   | Alcohol self-<br>administration in<br>action, alcohol use<br>on a voluntary basis,<br>and the recurrence<br>of alcohol-seeking<br>behavior due to<br>stress. | 25  |
| Enzymes   |   |  |  |     |
| <ul> <li>Cyclooxygenase (COX)<br/>1 &amp; 2</li> </ul>        | Aspirin   | Inhibiting synthesis of the<br>procoagulant Thromboxane<br>A2  | Ischemic stroke  | 26  |
| • Hydroxymethylglutaryl-<br>coenzyme A (HMG-CoA)<br>reductase | Statin drugs (lovastatin,<br>simvastatin, atorvastatin) | Competitively inhibit<br>HMG-CoA reductase in<br>mevalonate synthesis<br>pathway thereby reducing<br>cholesterol synthesis in the<br>liver   | Dyslipidemia and<br>cardiovascular<br>disease  | 27  |
| • Angiotensin converting enzyme (ACE)                         | Captopril and lisinopril                                | ACE competitive inhibitor,<br>leads to blood pressure<br>reduction by inhibiting<br>conversion of angiotensin I<br>to angiotensin II   | Hypertension   | 28  |
| • Guanylate cyclase type C                                    | Linaclotide   | Guanylate cyclase type<br>C activator, results in an<br>elevation of cGMP levels   | Chronic idiopathic constipation  | 29  |
| • Xanthine oxidase (XO)                                       | Allopurinol and febuxostat                              | Competitive inhibitor<br>of xanthine oxidase<br>which decreases the<br>formation of uric acid by<br>inhibiting hydroxylation of<br>hypoxanthine to xanthine<br>and xanthine to uric acid   | Hyperuricemia and<br>gout  | 30  |

# Table 1 cont.

| Specific target  | Drug                                | Mechanism   | Indications   | Ref |
|--|-------------------------------------|---|---|-----|
| • Dihydropteroate synthase   | Sulfa drugs                         | Dihydropteroate synthase<br>inhibitor by directly<br>competing with the substrate<br><i>p</i> ABA results in disruption of<br>folate biosynthesis | Antibacterial   | 31  |
| • Dihydrofolate reductase<br>(DHFR)  | Methotrexate                        | Inhibits Dihydrofolate<br>Reductase, reducing the<br>amount of tetrahydrofolate<br>required for the synthesis of<br>nucleotide bases              | Lymphoma, acute<br>lymphoblastic<br>leukemia, and<br>osteosarcoma     | 32  |
| <ul> <li>Thyroid peroxidase<br/>(TPO)</li> </ul>                           | Methimazole and<br>Propylthiouracil | Thyroid Peroxidase inhibitor<br>by interfering oxidation and<br>organification of iodine  | Hyperthyroidism   | 33  |
| • 5'-Lipoxygenase  | Zileuton                            | Inhibitor of 5'-Lipoxygenase,<br>leads to synthesis disruption<br>of leukotriene B4 (LTB4) and<br>cysteinyl leukotrienes (CysLT)                  | Asthma  | 34  |
| <ul> <li>γ-Aminobutyric acid<br/>aminotransferase<br/>(GABA-AT)</li> </ul> | Vigabatrin (Sabril)                 | Active-site directed inhibitor<br>of GABA-AT by blocking the<br>degradation of brain GABA   | Epilepsy and drug<br>addiction  | 35  |
| Ion channel  |                                     |   |   |     |
| • Calcium channel  | Nimodipine                          | Lowering the CaV1.2 mRNA expression and block the calcium uptake  | Diabetic<br>encephalopathy  | 36  |
| • Calcium channel  | Nifedipine                          | Activating PPAR <b>γ</b> to<br>suppresses MCP-1 and ABCA1   | Atherosclerosis   | 37  |
| • Sodium channel   | Lamotrigine                         | Inhibit Na voltage-gated ion<br>channel through a dual-<br>pocket mechanism   | Epilepsy  | 38  |
| Sodium-glucose     cotransporter 2 (SGLT2)                                 | Empagliflozin                       | Direct inhibition of SGLT2<br>channel to reduce glycated<br>hemoglobin  | Diabetes mellitus type 2  | 39  |
| • Potassium channel  | Dalfampridine                       | Blocking the potassium<br>channel exposed through MS-<br>related demyelination.   | Multiple sclerosis  | 40  |
| • ATP-sensitive potassium<br>(K ATP)                                       | Carbamazepine                       | Inhibiting KATP channel<br>activity primarily by<br>abolishing the stimulatory<br>effect of MgADP   | Congenital<br>hyperinsulinism,<br>neonatal diabetes,<br>DEND syndrome | 41  |
| Ion channel  |                                     |   |   |     |
| • Kv1.3 potassium channel  | Dalazatide                          | Blocking the Kv1.3 potassium channel and inhibit the T cell migration   | Plaque psoriasis  | 42  |
| <ul> <li>Epithelial sodium<br/>channel</li> </ul>                          | Amiloride                           | Blockade renin-angiotensin-<br>aldosterone system (RAAS)  | Proteinuric kidney  | 43  |

# Table 1 cont.

| Specific target   | Drug   | Mechanism   | Indications   | Ref |
|---|--|---|---|-----|
| Ion channel   |  |   |   |     |
| • Transient receptor<br>potential cation channel<br>member A1 (TRPA1) | GDC-0034<br>(Proline sulfonamide)              | Bind to closed state of<br>TRPA1 and restrict the<br>conformational change<br>related with gating<br>mechanism  | Asthma, airway<br>inflammation  | 44  |
| • Calcium released-related calcium (CRAC) channel                     | Auroxa   | Inhibit calcium channel<br>and the release of<br>proinflammatory<br>cytokine  | Pneumonia in<br>COVID-19  | 45  |
| DNA   |  |   |   |     |
| • Breast cancer<br>susceptibility gene 1 and<br>2 (BRCA1 and 2)       | Olaparib, rucaparib,<br>niraparib              | Combat homologous<br>recombination-deficient<br>(HRD) cancers with Poly<br>(ADP-ribose) polymerase<br>inhibitor (PARPi)   | Ovarian cancer  | 46  |
| • DNA damage response<br>(DDR) genes                                  | Camonsertib                                    | Synthetically lethal by<br>loss-of-function (LOF)<br>ataxia telangiectasia-<br>mutated (ATM)  | Ovarian cancer  | 47  |
| • DNA polymerase theta  | RP-6685  | Against breast and  | Breast and Ovarian<br>cancer Tumors that<br>have Alterations in<br>DNA Repair Genes | 48  |
| (Pol0)  | Novobiocin                                     | harboring BRCA-mutant<br>alleles. Blocks the activity<br>of a protein called DNA<br>polymerase theta, which<br>helps repair DNA that<br>has become damaged as<br>cells grow and divide. |   |     |
| • BCL-2 (B-cell lymphoma 2) protein (BCL2 gene)                       | Venetoclax                                     | Inhibits BCL-2 (B-cell<br>lymphoma 2)   | Small lymphocytic<br>lymphoma (SLL)   | 49  |
| • Single strand DNA breaks  | Talazoparib                                    | Inhibits the<br>polyadenosine<br>5'-diphosphoribose<br>polymerase (PARP)<br>enzymes, which play a<br>critical role in repairing<br>DNA single-strand breaks                             | Early triple negative<br>breast cancer  | 50  |
| • <i>BDNF</i> gene Val66Met polymorphism                              | Antipsychotic drugs                            | BDNF modulates the<br>major neurotransmitter<br>systems including<br>the dopaminergic,<br>glutamatergic, and<br>serotonergic systems  | Schizophrenia   | 51  |
| • Trophoblast cell surface<br>antigen 2 (TROP2)                       | Datopotamab deruxtecan<br>(Dato-DXd, DS-1062a) | Internalized into<br>tumor cells followed by<br>intracellular trafficking<br>to lysosome and DXd<br>release, which induced<br>DNA damage and<br>apoptosis in TROP2                      | Negative breast<br>cancer and urothelial<br>carcinoma                               | 52  |

#### TABLE 1. Cont.

| Specific target  | Drug  | Mechanism   | Indications                                 | Ref |
|--|---|---|---|-----|
| • DNA damage response<br>(DDR)-related gene  | Toripalimab, lenvatinib,<br>and gemcitabine plus<br>oxaliplatin (GEMOX) | Inhibits PD-1 (programmed<br>cell death protein 1)<br>protein on T cells, inhibits<br>a number of enzymes<br>involved and inhibits DNA<br>synthesis | Intrahepatic<br>cholangiocarcinoma<br>(ICC) | 53  |
| • EGFR (epidermal growth factor receptor) protein  | Osimertinib   | Osimertinib inhibits EGFR,<br>which is essential for the<br>growth and proliferation of<br>lung cancer cells  | Non-small cell lung<br>cancer (NSCLC)       | 54  |
| <ul> <li>Cancer-associated of<br/>proliferating cell nuclear<br/>antigen (caPCNA)</li> </ul> | Dacarbazine,<br>cyclophosphamides, and<br>Busulfan                      | Exploit this genetic<br>instability by inducing<br>additional DNA damage<br>to overwhelm the repair<br>system in cancer cells                       | Cancer                                      | 55  |

TABLE 1. Cont.

#### DISCUSSION

Target identification during the early stage of the drug discovery process plays an important role in successfully developing novel therapeutic agents. A 'drug target' refers to biomolecules that directly bind to drugs through specific binding sites and alter signaling pathways, leading to physiological therapeutic response to achieve benefits.<sup>56</sup> Most drug targets are protein including receptor, enzyme, ion channel, and transporters. Yet, some drugs bind directly with DNA rather than protein.<sup>57</sup> According database to curation conducted by Santos *et al.*,<sup>58</sup> 893 human and pathogen-derived biomolecules are primarily involved as targets of 1,578 FDA-approved drugs. G-protein coupled receptor (GPCR) make up the largest group of drug targets (33%), accompanied by enzyme and other biomolecules (30%), ion channel includes both voltagegated and ligand-gated (18%), nuclear receptor (16%), and kinase (3%). FDAapproved small molecule and biologic drugs are most widely used for nervous system therapeutic area, involving 249 drugs commonly act as antiepileptics (N03A), antipsychotic (N05A), and antidepressants (N06B). Followed by antineoplastic and immunomodulating agents (209 drugs), cardiovascular (205), antiinfections (204), alimentary tract and metabolism system (190), sensory organs (154), and many more.

An extensive knowledge of a drug's mechanism of action is fundamental to completely understanding its biological effects on the organism. Despite that, mechanisms of several drugs remain unclear even if the targets have been recognized. Therefore. molecular biology approaches are needed to obtain comprehensive insight of the interaction between drug and drug targets for pharmacological achieving desired effect.

#### GPCRs as drug target

G-protein-coupled receptors are a superfamily of seven transmembranespanning proteins that generate in a majority of organs and are activated by a diverse array of extracellular ligands. Several biological processes, such as neurotransmission, chemotaxis, inflammation, cell proliferation, cardiac and smooth muscle contraction, as well as visual and chemosensory perception, are influenced by signaling through these receptors. A particular kind of membrane protein known as GPCR is necessary for cell signaling. It is an important focal point for therapeutic research and is involved in a number of physiological processes.<sup>59</sup>

The mechanism of GPCR as a drug target involves the binding of a drug to the GPCR, which causes a conformational change in the receptor and triggers downstream signaling cascades. This activation or inhibition of GPCR signaling can lead to therapeutic effects in various diseases. Throughout signal transduction, agonists and ligands bind to the extracellular area of GPCRs, inducing the protein to first modify shape and become activated. By separating Gα from Gβγ, this activated GPCR further converts the inactive G protein to active G protein complex.<sup>60</sup> Additionally, helix-helix interaction ensures the GPCRs maintain their functional tertiary structure, which is essential for ligand binding, receptor folding and stability, and ligand-induced conformational changes that lead to G protein coupling.61

According to their amino acid sequences, the human GPCR family is classified into classes A (rhodopsin), B (secretin and adhesion), C (glutamate), and F (Frizzled) subfamilies.<sup>62</sup> Class A GPCRs such as 5HTR2A and P2Y12 (TABLE 1), also known as rhodopsin-like receptors, play a significant role in human disease. They are involved in a wide range of physiological processes such as sensory perception, neurotransmission, and immune response. These GPCRs have been implicated in numerous diseases including cardiovascular disorders. metabolic disorders, neurological disorders, and cancer.<sup>63</sup>5HT2A is a specific subtype of serotonin receptor that has been implicated in the pathophysiology of depression. It is believed that alterations in the signaling and activity of 5HT2A receptors may contribute to the development and maintenance of depressive symptoms. Targeting 5HT2A receptors with drugs that modulate their activity has shown promise in the treatment of depression.<sup>64,65</sup> Meanwhile,

adenosine 5' diphosphate (ADP)-binding platelet P2Y12 receptor (P2Y12R) is necessary for hemostasis, thrombosis, and platelet function. Mild-to-moderate bleeding diathesis appears in patients with hereditary P2Y12R abnormalities.<sup>66</sup> Particularly in the case of cardiovascular disorders, these receptors are crucial in the development of numerous human diseases. In biomechanically the induced aggregation mechanism, P2Y12 can directly contribute to thrombus formation and can also activate PI3K kinase and Syk kinase to drive thrombus formation.67

Receptors for peptide hormones, such as PTH1 and GLP1, belong in the class B GPCRs of the secretin family (TABLE 1).<sup>68</sup> These types of receptors are significant drug targets in a variety of human diseases, such as diabetes, osteoporosis, cancer, neurodegeneration, cardiovascular disease, headaches, and psychiatric disorders.<sup>69</sup> PTH1, also referred to as the parathyroid hormone receptors, is involved in bone structure maintenance.<sup>70</sup> encouraging Bv osteoblast reducing growth and osteoblast cell death or apoptosis, PTH can increase the number of boneforming cells.<sup>71</sup> While glucagon works to maintain blood glucose levels through the conversion of glycogen to glucose, the incretins glucagon-like peptide 1 (GLP-1) and glucagon itself are members of the glucagon subfamily. The primary way that the incretins lower blood glucose is through the release of insulin.72

Moreover, class C GPCRs are a distinct family of receptors characterized by their unique structural features and functional roles in various physiological processes. This class includes important receptors such as metabotropic glutamate receptors (mGluRs), γ-aminobutyric acid B receptors (GABA-B), calciumsensing receptors (CaSR), and taste receptors. Class C GPCRs play critical roles in various physiological processes, including calcium homeostasis, neurotransmission, taste sensation, and signal modulation. The calcium-sensing receptor (CaSR) regulates calcium levels in the bloodstream by detecting extracellular calcium concentrations and modulating parathyroid hormone (PTH) secretion accordingly. Modulating CaSR (TABLE 1) activity presents potential therapeutic avenues for managing disorders related to calcium imbalance, such as osteoporosis or hyperparathyroidism.<sup>26</sup>

Last but not least, class F GPCRs, consisting of ten Frizzled (FZD) paralogs and smoothened (SMO), play crucial roles in embryonic development, stem regulation, and tumorigenesis. cell Specifically, FZDs mediate WNT signaling, SMO mediates Hedgehog whereas signaling. Despite lacking conserved motifs like those in class A GPCRs, such as the E/DRY motif, Class F receptors exhibit ligand-induced and constitutive activities involving the opening of the transmembrane bundle by a TM6 swing-out motion, akin to Class A and B GPCRs' activation mechanisms.<sup>73,74</sup> Drug mechanisms targeting Class F receptors aim to disrupt aberrant signaling pathways contributing to diseases like cancer. For instance, inhibitors designed to interfere with the GLI transcription factors downstream of SMO activation show promise in treating basal-cell carcinomas caused by aberrant Hedgehog signaling. Similarly, modulators of Wnt/Fzd interactions could offer therapeutic options for conditions influenced by dysregulation of this pathway. Further research will continue to uncover specific druggable sites within these receptors, enabling effective treatments tailored more towards disrupting pathologic signaling while preserving physiologic functions.<sup>74</sup>

G-protein coupled receptors are a widely studied class of cell surface receptors that play a crucial role in various physiological processes. Although GPCRs have been successful targets for drug development, there are several limitations and challenges associated with targeting them. Firstly, limitation is the potential for off-target effects, this means that drugs targeting a specific GPCR may also interact with other receptors, leading to unintended side effects.<sup>75</sup> Secondly, the complexity of GPCR signaling pathways. The signaling pathways involving GPCRs are highly intricate and interconnected, making it difficult to selectively modulate the desired pathway without affecting others. Thirdly, the limited structural information available for GPCRs, which hinders the rational design of drugs.<sup>76</sup> Moreover, GPCRs exhibit a high degree of structural flexibility, undergoing conformational changes upon ligand binding, which further complicates the rational design of drugs targeting GPCRs.77,78

Understanding the characteristics and functioning of GPCRs as a drug target is crucial in developing effective medications for several reasons. Firstly, GPCRs are one of the largest and most important protein families involved in cellular signaling and regulation. Secondly, they are involved in a wide range of physiological processes and making them pathways, attractive targets for drug intervention. Thirdly, GPCRs have been successfully targeted by numerous drugs and account for a significant portion of pharmaceuticals on the market. By knowing the functioning characteristics and of GPCRs as a drug target, researchers can better understand how to manipulate these receptors with medications. This knowledge can lead to the development of more targeted and effective drugs with fewer side effects. Additionally, understanding GPCRs allows researchers to identify the endogenous ligands that naturally bind to these receptors. This can provide insights into the normal physiological roles of GPCRs and potential therapeutic targets. Moreover, by understanding the structure and functioning of GPCRs, researchers can design drugs that specifically target these receptors and modulate their activity. Ultimately, this knowledge can drive the discovery of novel drugs that have better efficacy, improved safety profiles, and reduced side effects.<sup>79,80</sup>

# Enzyme as drug target

Enzymes are exceptionally powerful capable biocatalvsts of promoting chemical transformations that take place in any living cell of an organism including metabolism, cell growth, division, and motility, as well as cellular signaling with high efficiency and selectivity. The existence of substrate binding sites which can be exploited and modified are one of the considerations of targeting enzymes for drug development.<sup>81</sup> According to Agarwal *et al.*,<sup>82</sup> enzymes comprise 27% of FDA-approved oral drugs. Hydrolase are the most common class of enzymatic drug targets followed by oxidoreductases, transferase, isomerase, lyase, and ligase.<sup>83</sup> Drugs acting on enzymes can either inhibit or activate them. Inhibition of enzymes has become a recurrent strategy for drug design because of upregulating mechanisms being prevalent in numerous diseases.

Enzyme inhibitors can either decrease or totally inhibit enzyme catalytic activity through three common mechanisms called competitive, noncompetitive, and uncompetitive inhibition.<sup>84</sup> Competitive inhibitor drugs serve as substrate analog to occupy the active site of enzymes and prevent substrate attachment<sup>85</sup> (FIGURE 2b). A great number of therapeutic agents are developed through this regulation. The most prominent atherosclerotic cardiovascular disease drug named Statins has been declared as 3-hydroxy-3-methylglutarate CoA reductase competitive inhibitor. Randomized trials conducted by Hong et al.,<sup>86</sup> showed the reduction of low-density lipoprotein cholesterol (LDL-C) levels around 50-70 mg/dL after high-intensity statin therapy for around 3 yr. While some drugs act as competitive inhibitors, non-competitive inhibitors perform another binding property. In non-competitive inhibition, the inhibitor binds to the allosteric site of enzymes, leading to configurational and disrupting change enzyme's catalytic activity<sup>85</sup> (FIGURE 2c). Aspirin, cardiovascular drug, establishes а non-competitive mechanism а bv inhibiting synthesis of Thromboxane A2. Randomized trials elucidated that aspirin treatment reduced the 6 weeks risk of recurrent ischemic stroke by about 60% and disabling fatal ischemic stroke by about 70%.<sup>87</sup> A third category, uncompetitive inhibitor, also considered as an inhibition type which disturbs enzyme-substrate complexes, leading to dead end complex formation<sup>84</sup> (FIGURE 2d). Unfortunately, this phenomenon is unusual in the drug development process and excluded in this current review.



FIGURE 2. Three kind of enzyme inhibition mechanism; (a) Common binding of enzyme; (b) Competitive inhibitor act as substrate analog; (c) Non-competitive inhibitor bind to allosteric site leads to conformational change; and (d) Uncompetitive inhibitor bind to enzyme-substrate complex (https://BioRender.com)

Drugs that act as an enzyme activator can accelerate biochemical reaction through enzyme modification (ie. phosphorylation) or low molecular weight positive modulators.<sup>88</sup> There are particular conditions where enzyme activators comprise therapeutic benefits. For instance, activator of guanylyl cvclase С (linaclotide), increases phosphorylation rate resulting in an elevation of cGMP levels and induces several cellular responses related to chronic idiopathic constipation improvement.<sup>34</sup> The effectiveness and efficacy of linaclotide treatment against gastrointestinal disease have been studied by Peng et al.,<sup>89</sup> Group of irritable bowel syndrome (IBS) patients treated with linaclotide (290  $\mu$ g/d) showed a significantly greater improvement than placebo in all secondary endpoints from the first 2 wk.

Several pharmaceutical agents targeting enzymes have been evolved for a wide range of applications, including gastrointestinal, respiratory, cancer, cardiovascular, and infectious diseases. However, research related to enzymatic drug targets faces various challenges and limitations. Certain drugs are known to interact with others through drugdrug interactions, leading to unwanted side effects. Besides, developing and manufacturing enzymatic drug targets can be very expensive, making them cost-prohibitive under certain circumstances. Other challenges lie in the similarity of enzyme structure resulting in lack of drug specificity and increasing off-target probability. Enzymes are also known to have short in vivo half-life and in particular, several studies found that the immune system exhibits enzyme clearance. The future of enzyme-based drug expansion lies in the development of selective and potent drugs as well as the ability to target enzymes involved in multiple pathways. Combination of enzyme targeting drugs with other therapeutic strategies such as gene therapy also give promising approaches.<sup>90,91</sup>

# Ion channel as drug target

channel-targeting Ion drugs are novel but essential in modern pharmacology, significantly influencing numerous physiological processes.<sup>92,93</sup> It offers a precise and potent way to alter cellular signaling and function. Multiple sclerosis secondary-progressive multiarm randomization trial (MS-SMART) evaluated the neuroprotective has effect of amiloride among other agents in patients with secondary progressive multiple sclerosis. The study included 90 patients per treatment arm and assessed the percentage of brain volume change (PBVC) as the primary outcome. Results indicated a reduction in PBVC of 0.5% in the amiloride group compared to placebo over 96 wk.94 A consensus report from experts in China highlighted the safe and effective use of ion channel drugs for managing chronic pain, emphasizing the need for standardized practices based on clinical evidence.<sup>95</sup> That therapeutic approach spans a broad potential spectrum of medical conditions based on the 2 prominent families of ion channeltargeting drugs.

Voltage-gated ion channels are transmembrane proteins that open and close in response to changes in membrane potential.96 Thev control cellular excitability and signal transmission by regulating the flow of several ions, such as sodium, potassium, and calcium.97 Representing a primary target for pharmacological intervention due to its fundamentalroleinorchestratingcellular excitability and signaling. As mentioned in TABLE 1, the use of nimodipine, a voltage-gated ion channel-targeting drug, has also successfully normalized the activity of intracellular free calcium levels in diabetic animal models. In a study with randomized controlled trial involving 1,000 patients with trigeminal neuralgia, carbamazepine was found to reduce pain episodes by approximately 70% compared to placebo over 12 wk.98 A meta-analysis involving 38 studies with over 8,000 patients showed that gabapentin significantly reduced pain scores compared to placebo, with a mean difference of -1.42 (95% CI: -1.73 to -1.10) on a 0-10 scale, indicating substantial efficacy in managing neuropathic pain.99 Compared to ligand-gated, voltage-gated channels offer unique opportunities for precise modulation of neuronal activity, making them particularly attractive targets for therapeutic intervention in various neurological and cardiovascular disorders. This emphasis on voltagechannels underscores gated ion their significance in drug discovery and development efforts to address complex physiological and pathological conditions.

On the other hand, ligand-gated ion channels are responsible for opening or closing in response to the binding of ligand-specific molecules, such as neurotransmitters hormones.<sup>100</sup> or Ligand-gated ion channels are crucial for neurotransmission and involve several processes, such as synaptic plasticity and sensory perception.<sup>101</sup> A study involving 404 patients demonstrated that memantine reduced cognitive decline as measured by the Alzheimer's disease assessment scale-cognitive subscale (ADAS-Cog) by an average of 2.2 points compared to placebo over six months.<sup>102</sup> The evidence shows that ion channeltargeted drugs are not only responsible for synaptic diseases in particular but are also involved in other health problems.

However, the experiment regarding ion channel-targeting drugs faces significant challenges in present and future development. One of the current challenges lies in specificity and safety, such as off-target effects and potential adverse reactions that could happen. For example, while targeting sodium channels can relieve pain, it may also disrupt regular neuronal activity, causing side effects like dizziness or sedation.<sup>103</sup> This necessitates extensive safetv profiling in clinical trials. Furthermore, the ion channel structure complexity and their regulatory mechanism pose a pressing challenge in drug development and optimization, such as individual genetic differences that can affect patients' response to ion channel drugs, complicating treatment strategies. For instance, polymorphisms in P2X receptor genes have been associated with variations in blood pressure responses to treatments targeting these channels.<sup>104</sup> The present and future of this field will likely depend on overcoming these hurdles to enhance the efficacy and safety of new therapeutic agents.

# DNA as drug target

Nucleic acids are important for transferring the genetic code from DNA and translating it into specific proteins. Hybridization can be used to target the activity of specific genes and genes have potential use in drug development. For example, if a gene carried by a virus is silenced by a drug, the virus may not be able to survive inside the cells it has infected and will die. Another example is if cancer cells grow uncontrollably because a gene is permanently stuck in the active position, it can be stopped by suppressing its expression by administering drugs that target DNA.<sup>105</sup> RNA has also been shown to be closely involved in gene regulation, namely the process of turning genes on or off at specified times and locations in living cells. Drugs that target DNA work in different ways in gene therapy. Intercalation is the process of changing the shape of DNA so that it cannot be used by ribosomes, for example chloramphenicol changes the shape of DNA. Alkylation is the process by which a drug can change the shape of DNA so that it cannot be used by ribosomes. An

example of a drug that uses an alkylation mechanism is neomycin. Apart from that, there is also a mechanism by which this drug can damage DNA replication.<sup>106</sup> This has implications for diseases such as cancer, where many genes are deregulated, leading to uncontrolled cell growth. For example, combination of toripalimab, lenvatinib, and gemcitabine plus oxaliplatin (GEMOX) as the first therapy for advanced stage ICC, which is done by inhibiting the PD-1 (programmed cell death protein 1) protein on T cells, which is a mechanism used by cancer cells to avoids detection and destruction by the immune system, inhibits a number of enzymes involved and inhibits DNA synthesis.<sup>53</sup> According to Okajima *et* al.,<sup>52</sup> drugs that target DNA datopotamab deruxtecan can internalized into tumor cells followed by intracellular trafficking to lysosome and DXd release, which induced DNA damage and apoptosis in TROP2.

The drugs shown in TABLE 1 are intended to bind to and stop the expression of specific genes, either at the RNA level or directly on DNA. The mechanism by which drugs target DNA can have an effect on the properties of the nucleic acid molecule, as well as the way it hybridizes or binds to its target sequence meaning that it sometimes connects to other areas, causing offtarget effects such as immunogenicity. Immunogenicity refers to the ability of a molecule to enhance the immune system, which is an undesirable characteristic of drugs administered to humans. The main problem with nucleic acidbased drugs is their delivery to target tissues.<sup>107</sup> To target DNA as a target for a drug can be optimized by knowing that the selected target DNA is directly involved in treating the disease being targeted, ensuring that the target DNA can be accessed by the drug, the DNA to be targeted has high specificity thereby reducing the potential for mistargeting or targeting DNA. It is also important to test the right dose to achieve a therapeutic effect without excessive side effects, considering the possibility of resistance to the drug so that the drug remains effective in treating a disease. The DNA that will be used as a drug target should be stable so that there are no significant changes in the DNA structure that could reduce the action of the drug used.

# **CONCLUSION**

receptors. G-protein coupled one of the biomolecules involved in modulating receptor activity and various physiological processes, can be targeted by affecting relevant signaling pathways to provide therapeutic effects against certain diseases, such as osteoporosis and carcinoma cancer. Targeting biomolecules, like enzymes, has the potential to overcome diseases such as cancer and infections through the understanding of the enzyme inhibition or activation mechanism as well as the mechanism between and substrates. enzymes Targeting drugs at ion channels that regulate ion transmission has the potential to treat various neurological and cardiovascular disorders. A DNA and RNA can be targeted especially in the treatment of diseases such as cancer by altering gene expression or damaging DNA involved in disease progression. Further research needs to be done that focuses on understanding selective and specific drug delivery mechanisms, the structure and the complexity of each biomolecule, and the interactions between drugs and biomolecules to improve treatment effectiveness.

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# REFERENCES

- Karaman R. Commonly used drugs

   uses, side effects, bioavailability
   approaches to improve it. New
   York: Nova Science Publishers Inc.,
   2015; 1-293.
   https://doi.org/10.13140/
   RG.2.1.1444.4640
- Glynn J, Bhikha RAH. Herbal products and conventional drugs

   an uneasy alliance. Int J Human Health Sci (IJHHS) 2018; 2(4):193. https://doi.org/10.31344/ijhhs.v2i4.55
- 3. Karimi A, Majlesi M, Rafieian-Kopaei M. Herbal versus synthetic drugs; beliefs and facts. J Nephropharmacol 2015; 4(1):27-30.
- 4. Patel S, Sadhu P, Kumari M, Dash DK, Jain S, Sen AK. A review on anticancer agents: conventional drugs and novel target specific inhibitors. Orient J Chem 2023; 39(3):657-69.

https://doi.org/10.13005/ojc/390316

5. Nazli A, He DL, Liao D, Khan MZI, Huang C, He Y. Strategies and progresses for enhancing targeted antibiotic delivery. Adv Drug Deliv Rev 2022; 189:114502. https://doi.org/10.1016/j.

addr.2022.114502

- Vasan N, Baselga J, Hyman DM. A view on drug resistance in cancer. Nature 2019; 575(7782):299-309. https://doi.org/10.1038/s41586-019-1730-1
- 7. Aminu N, Bello I, Umar NM, Tanko N, Aminu A, Audu MM. The influence of nanoparticulate drug delivery systems in drug therapy. J Drug Deliv Sci Technol 2020; 60(Suppl 1):101961. https://doi.org/10.1016/j.

jddst.2020.101961

8. Bhutani U, Basu T, Majumdar S.

Oral drug delivery: conventional to long acting new-age designs. Eur J Pharm Biopharm 2021; 162:23-42. https://doi.org/10.1016/j. ejpb.2021.02.008

9. Sivamaruthi BS, Nallasamy PK, Sivakumar S, Kesika P, Chaiyasut C. Pharmaceutical and biomedical applications of starch-based drug delivery system: A review. J Drug Deliv Sci Technol 2022; 77(3):103890. https://doi.org/10.1016/j.

jddst.2022.103890

- 10. Farjadian F, Ghasemi A, Gohari O, Roointan A, Karimi M, Hamblin MR. Nanopharmaceuticals and nanomedicines currently on the market: challenges and opportunities. Nanomedicine (Lond) 2019; 14(1):93-126. https://doi.org/10.2217/nnm-2018-0120
- 11. Karel P. Molecular drug targets and drug delivery. Open Access J Cancer Oncol 2017; 1(2). h t t p s://doi.org/10.23880/OAJCO-16000107
- 12. Laurieri N, Delgoda R. Novel targets in drug discovery. Pharmacognosy 2017; 617-32. https://doi.org/10.1016/B978-0-12-802104-0.00031-7
- Gao D, Chen Q, Zeng Y, Jiang M, Zhang Y. Applications of machine learning in drug target discovery. Curr Drug Metab 2020; 21(10):790-803.

https://doi.org/10.2174/1567201817 999200728142023

14. Muhamadejevs R, Živković L, Dzintare M, Sjakste N. DNAbinding activities of compounds acting as enzyme inhibitors, ion channel blockers and receptor binders. Chem Biol Interact 2021; 348:109638.

https://doi.org/10.1016/j.cbi.2021.109638

15. Prabahar K, Alanazi Z, Qushawy M. Targeted drug delivery system: advantages, carriers and strategies. IJPER 2021; 55(2):346-53.

- 16. Walker JKL, Penn RB, Hanania NA, Dickey BF, Bond RA. New perspectives regarding  $\beta(2)$ adrenoceptor ligands in the treatment of asthma. Br J Pharmacol 2011; 163(1):18-28. https://doi.org/10.1111/j.1476-5381.2010.01178.x
- Pollock M, Sinha IP, Hartling L, Rowe BH, Schreiber S, Fernandes RM. Inhaled short-acting bronchodilators for managing emergency childhood asthma: an overview of reviews. Allergy 2017; 72(2):183-200.

https://doi.org/10.1111/all.13039

- 18. Miller PD, Hattersley G, Riis BJ, Williams GC, Lau E, Russo LA, *et al.* Effect of abaloparatide vs placebo on new vertebral fractures in postmenopausal women with osteoporosis: a randomized clinical trial. JAMA 2016; 316(7):722-33. https://doi.org/10.1001/jama.2016.11136
- 19. Nauck MA, Petrie JR, Sesti G, Mannucci E, Courrèges JP, Lindegaard ML, *et al.* A phase 2, randomized, dose-finding study of the novel once-weekly human glp-1 analog, semaglutide, compared with placebo and open-label liraglutide in patients with type 2 diabetes. Diabetes Care 2016; 39(2):231-41.

https://doi.org/10.2337/dc15-0165

20. Yin J, Mobarec JC, Kolb P, Rosenbaum DM. Crystal structure of the human OX2 orexin receptor bound to the insomnia drug suvorexant. Nature 2015; 519(7542):247-50.

https://doi.org/10.1038/nature14035

21. Block GA, Bushinsky DA, Cheng S, Cunningham J, Dehmel B, Drueke TB, *et al.* Effect of etelcalcetide vs cinacalcet on serum parathyroid hormone in patients receiving hemodialysis with secondary hyperparathyroidism: a randomized clinical trial. JAMA, 2017; 317(2):156-64.

https://doi.org/10.1001/jama.2016.19468

22. Ishima T, Futamura T, Ohgi Y, Yoshimi N, Kikuchi T, Hashimoto K. Potentiation of neurite outgrowth by brexpiprazole, a novel serotonin-dopamine activity modulator: a role for serotonin 5-HT1A and 5-HT2A receptors. Eur Neuropsychopharmacol 2015; 25(4):505-11.

https://doi.org/10.1016/j. euroneuro.2015.01.014

23. Rymer JA, Bhatt DL, Angiolillo DJ, Diaz M, Garratt KN, Waksman R, *et al.* Cangrelor use patterns and transition to oral P2Y12 inhibitors among patients with myocardial infarction: initial results from the cameo registry. J Am Heart Assoc 2022; 11(11):e024513.

# https://doi.org/10.1161/JAHA.121.024513

24. Xiao T, Ren S, Bao J, Gao D, Sun R, Gu X, et al. Vorapaxar proven to be a promising candidate for pulmonary fibrosis by intervening in the PAR1/JAK2/STAT1/3 signaling pathway-an experimental in vitro and vivo study. Eur J Pharmacol 2023; 943:175438. https://doi.org/10.1016/j.

ejphar.2022.175438

- 25. Kokhan VS, Anokhin PK, Abaimov DA, Shamakina IY, Soldatov VO, Deykin AV. Neurokinin-1 receptor antagonist rolapitant suppresses anxiety and alcohol intake produced by repeated withdrawal episodes. FEBS J 2022; 289(16):5021-9. https://doi.org/10.1111/febs.16400
- 26. Kamarova M, Baig S, Patel H, Monks K, Wasay M, Ali A, *et al.* Antiplatelet use in ischemic stroke. Ann Pharmacother 2022; 56(10):1159-73. h t t p s : / / d o i . org/10.1177/10600280211073009
- 27. Hirota T, Fujita Y, Ieiri I. An updated review of pharmacokinetic drug interactions and pharmacogenetics of statins. Expert Opin Drug Metab

Toxicol, 2020; 16(9):809-22. https://doi.org/10.1080/17425255.20 20.1801634

28. Cutrell S, Alhomoud IS, Mehta A, Talasaz AH, Van Tassell B, Dixon DL. Ace-inhibitors in hypertension: a historical perspective and current insights. Curr Hypertens Rep 2023; 25(9):243-50.

https://doi.org/10.1007/s11906-023-01248-2

29. Rao SS, Manabe N, Karasawa Y, Hasebe Y, Nozawa K, Nakajima A, *et al.* Comparative profiles of lubiprostone, linaclotide, and elobixibat for chronic constipation: a systematic literature review with meta-analysis and number needed to treat/harm. BMC Gastroenterol 2024; 24(1):12.

https://doi.org/10.1186/s12876-023-03104-8

30. White WB, Saag KG, Becker MA, Borer JS, Gorelick PB, Whelton A, *et al.* Cardiovascular safety of febuxostat or allopurinol in patients with gout. N Engl J Med 2018; 378(13):1200-10.

https://doi.org/10.1056/NEJMoa1710895

31. Griffith EC, Wallace MJ, Wu Y, Kumar G, Gajewski S, Jackson P, *et al*. The structural and functional basis for recurring sulfa drug resistance mutations in *Staphylococcus aureus* dihydropteroate synthase. Front Microbiol, 2018; 9:1369. https://doi.org/10.3389/

fmicb.2018.01369

32. Raimondi M, Randazzo O, La Franca M, Barone G, Vignoni E, Rossi D, *et al.* DHFR inhibitors: reading the past for discovering novel anticancer agents. Molecules 2019; 24(6):1140.

h t t p s://doi.org/10.3390/ molecules24061140

33. Yoshihara A, Luo Y, Ishido Y, Usukura K, Oda K, Sue M, *et al.* Inhibitory effects of methimazole and propylthiouracil on iodotyrosine deiodinase 1 in thyrocytes. Endocr J 2019; 66(4):349-57. https://doi.org/10.1507/endocrj. EI18-0380

34. Makullah M, Ellis DA, Jones M, Steele C. Hematopoietic 12/15-lipoxygenase activity negatively contributes to fungalassociated allergic asthma. Am J Physiol Lung Cell Mol Physiol 2023; 325(2):L104-13.

h t t p s : // d o i . o r g / 1 0 . 1 1 5 2 / ajplung.00090.2023

Silverman 35. RB. Design and Mechanism of GABA inactivators. aminotransferase Treatments for epilepsies and Chem Rev 2018; addictions. 118(7):4037-70. https://doi.org/10.1021/acs.

chemrev.8b00009

36. Singhal K, Sandhir R. L-type calcium channel blocker ameliorates diabetic encephalopathy by modulating dysregulated calcium homeostasis. J Neurosci Res 2015; 93(2):296-308. https://doi.org/10.1002/inr.22478

https://doi.org/10.1002/jnr.23478

- 37. Grosskopf I, Shaish A, Charach G, Harats D, Kamari Y. Nifedipine treatment for hypertension is associated with enhanced lipolytic activity and accelerated clearance of postprandial lipemia. Horm Metab Res 2016; 48(4):257-62. https://doi.org/10.1055/s-0035-1565180
- Huang J, Fan X, Jin X, Teng L, Yan N. Dual-pocket inhibition of Nav channels by the antiepileptic drug lamotrigine. Proc Natl Acad Sci USA 2023; 120(41):e2309773120. https://doi.org/10.1073/pnas.2309773120
- Zinman B, Wanner C, Lachin JM, Fitchett D, Bluhmki E, Hantel S, *et al.* Empagliflozin, cardiovascular outcomes, and mortality in type 2 diabetes. N Engl J Med, 2016; 373(22):2117-28.

https://doi.org/10.1056/NEJMoa1504720

40. Lo AC, Ruiz JA, Koenig CM,

Anderson BM, Olson KM, Triche EW. Effects of dalfampridine on multi-dimensional aspects of gait and dexterity in multiple sclerosis among timed walk responders and non-responders. J Neurol Sci 2015; 356(1-2):77-82.

https://doi.org/10.1016/j.jns.2015.06.008

41. Zhou Q, Chen PC, Devaraneni PK, Martin GM, Olson EM, Shyng SL. Carbamazepine inhibits ATPsensitive potassium channel activity by disrupting channel response to MgADP. Channels 2014; 8(4):376-82.

https://doi.org/10.4161/chan.29117

42. Tarcha EJ, Olsen CM, Probst P, Peckham D, Muñoz-Elías EJ, Kruger JG, *et al.* Safety and pharmacodynamics of dalazatide, a Kv1.3 channel inhibitor, in the treatment of plaque psoriasis: A randomized phase 1b trial. PloS One 2017; 12(7):e0180762. https://doi.org/10.1371/journal.

pone.0180762

- 43. Shen W, Alshehri M, Desale S, Wilcox C. The effect of amiloride on proteinuria in patients with proteinuric kidney disease. Am J Nephrol 2021; 52(5):368-77. https://doi.org/10.1159/000515809
- 44. Balestrini A, Joseph V, Dourado M, Reese RM, Shields SD, Rougé L, *et al.* A TRPA1 inhibitor suppresses neurogenic inflammation and airway contraction for asthma treatment. J Exp Med 2021; 218(4):e20201637.

https://doi.org/10.1084/jem.20201637

- 45. Bruen C, Al-Saadi M, Michelson EA, Tanios M, Mendoza-Ayala R, Miller J, *et al.* Auxora vs. placebo for the treatment of patients with severe COVID-19 pneumonia: a randomized-controlled clinical trial. Crit Care 2022; 26(1):101. https://doi.org/10.1186/s13054-022-03964-8
- 46. Lau CH, Seow KM, Chen KH. The

molecular mechanisms of actions, effects, and clinical implications of PARP inhibitors in epithelial ovarian cancers: a systematic review. Int J Mol Sci 2022; 23(15):8125.

https://doi.org/10.3390/ijms23158125

- 47. Yap TA, Fontana E, Lee EK, Spigel DR, Højgaard M, Lheureux S, et al. Camonsertib in DNA damage response-deficient advanced solid tumors: phase 1 trial results. Nat Med 2023; 29(6):1400-11. https://doi.org/10.1038/s41591-023-02399-0
- 48. Gu L, Hickey RJ, Malkas LH. Therapeutic targeting of DNA replication stress in cancer. Genes (Basel) 2023; 14(7):1346. https://doi.org/10.3390/genes14071346
- 49. Lachowiez C, DiNardo CD, Konopleva M. Venetoclax in acute myeloid leukemia - current and future directions. Leuk Lymphoma 2020; 61(6):1313-22. https://doi.org/10.1080/10428194.20 20.1719098
- 50. Hoy SM. Talazoparib: first global approval. Drugs 2018; 78(18):1939-46. https://doi.org/10.1007/s40265-018-

1026-z

51. Han M, Deng C. BDNF as a pharmacogenetic target for antipsychotic treatment of schizophrenia. Neurosci Lett 2020; 726:133870. https://doi.org/10.1016/j.

neulet.2018.10.015

52. Okajima D, Yasuda S, Maejima T, Karibe T, Sakurai K, Aida T, *et al.* Datopotamab deruxtecan, a novel TROP2-directed antibody-drug conjugate, demonstrates potent antitumor activity by efficient drug delivery to tumor cells. Mol Cancer Ther 2021; 20(12):2329-40. https://doi.org/10.1158/1535-7163

https://doi.org/10.1158/1535-7163

53. Shi GM, Huang XY, Wu D, Sun HC, Liang F, Ji Y, *et al.* Toripalimab combined with lenvatinib and GEMOX is a promising regimen as first-line treatment for advanced intrahepatic cholangiocarcinoma: a single-center, single-arm, phase 2 study. Signal Transduct Target Ther 2023; 8(1):106.

https://doi.org/10.1038/s41392-023-01317-7

54. Sakata Y, Sakata S, Oya Y, Tamiya M, Suzuki H, Shibaki R, *et al.* Osimertinib as first-line treatment for advanced epidermal growth factor receptor mutation-positive non-small-cell lung cancer in a real-world setting (OSI-FACT). Eur J Cancer 2021; 159:144-53.

https://doi.org/10.1016/j.ejca.2021.09.041

- 55. Gu L, Hickey RJ, Malkas LH. Therapeutic targeting of dna replication stress in cancer. Genes (Basel) 2023; 14(7):1346. https://doi.org/10.3390/genes14071346
- 56. Zhang X, Yu W, Li Y, Wang A, Cao H, Fu Y. Drug development advances in human genetics-based targets. Med Comm 2024; 5(2):e481. https://doi.org/10.1002/mco2.481
- 57. Ritter JM, Flower RJ, Henderson G, Loke YK, MacEwan D, Rang HP. Rang & Dale's Pharmacology. Elsevier Health Sciences; 2018.
- 58. Santos R, Ursu O, Gaulton A, Bento AP, Donadi RS, Bologa CG, *et al.* A comprehensive map of molecular drug targets. Nat Rev Drug Discov 2016; 16(1):19-34.

https://doi.org/10.1038/nrd.2016.230

- 59. Cacace A, Banks M, Spicer T, Civoli F, Watson J. An ultra-HTS process for the identification of small molecule modulators of orphan G-protein-coupled receptors. Drug Discov Today 2003; 8(17):785-92. https://doi.org/10.1016/s1359-6446(03)02809-5
- 60. Mansfield R, Able S, Griffin P, Irvine B, James I, Macartney M, Dorr P. CCR5 pharmacology methodologies and associated applications. Methods Enzymol 2009; 460:17-55.

https://doi.org/10.1016/s0076-6879(09)05202-1

- 61. Tuteja N. Signaling through G protein coupled receptors. Plant Signal Behav, 2009; 4(10):942-7. https://doi.org/10.4161/psb.4.10.9530
- 62. Baldwin JM. Structure and function of receptors coupled to G proteins. Curr Opin Cell Biol 1994; 6(2):180-90. https://doi:10.1016/0955-0674(94)90134-1
- 63. Hauser AS, Attwood MM, Rask-Andersen M, Schiöth HB, Gloriam DE. Trends in GPCR drug discovery: new agents, targets and indications. Nat Rev Drug Discov 2017; 16(12):829-42. https://doi.org/10.1038/nrd.2017.178
- 64. Cheng L, Xia F, Li Z, Shen C, Yang Z, Hou H, *et al.* Structure, function and drug discovery of GPCR signaling. Mol Biomed 2023; 4(1):46. https://doi.org/10.1186/s43556-023-00156-w
- 65. Wang L, Zhou C, Zhu D, Wang X, Fang L, Zhong J, *et al.* Serotonin-1A receptor alterations in depression: a meta-analysis of molecular imaging studies. BMC Psychiatry 2016; 16(1):319. https://doi.org/10.1186/s12888-016-1025-0
- 66. Zięba A, Stępnicki P, Matosiuk D, Kaczor AA. Overcoming depression with 5-HT2A receptor ligands. Int J Mol Sci 2021; 23(1):10. https://doi.org/10.3390/ijms23010010
- 67. Cattaneo M. P2Y12 receptors: structure and function. J Thromb Haemost 2015; 13 Suppl 1:S10-6. https://doi.org/10.1111/jth.12952
- 68. Wang L, Wang J, Xu J, Qin W, Wang Y, Luo S, *et al.* The role and molecular mechanism of P2Y12 receptors in the pathogenesis of atherosclerotic cardiovascular diseases. Appl Sci 2021; 11(19):9078.

https://doi.org/10.3390/app11199078

69. Alexander SPH, Benson HE, Faccenda E, Pawson AJ, Sharman IL, Spedding M, et al. The concise guide to PHARMACOLOGY 2013/14: G protein-couple receptors. Br J Pharmacol 2013; 170(8):1459-581. https://doi.org/10.1111/bph.12445

- Pal K, Melcher K, Xu HE. Structure 70. and mechanism for recognition of peptide hormones by Class B G-protein-coupled receptors. Acta Pharmacol Sin 2012; 33(3):300-11. https://doi.org/10.1038/aps.2011.170
- 71. Esbrit P, Alcaraz MJ. Current perspectives on parathyroid hormone (PTH) and PTH-related protein (PTHrP) as bone anabolic therapies. Biochem Pharmacol 2013; 85(10):1417-23.

https://doi.org/10.1016/j.bcp.2013.03.002

Canalis E, Giustina A, Bilezikian JP. 72. Mechanisms of anabolic therapies for osteoporosis. N Engl J Med 2007; 357(9):905-16.

https://doi.org/10.1056/NEJMra067395 Wright SC, Kozielewicz P, Kowalski-

- 73. Jahn M, Petersen J, Bowin CF, Slodkowicz G, et al. A conserved molecular switch in Class F receptors regulates receptor activation and pathway selection. Nat Commun 2019; 10(1):667. https://doi.org/10.1038/s41467-019-08630-2
- 74. Schulte G, Kozielewicz P. Structural insight into class F receptors - What have we learnt regarding agonistinduced activation? Basic Clin Pharmacol Toxicol 2020; 126(Suppl 6):17-24.

https://doi.org/10.1111/bcpt.13235

Cho YM, Merchant CE, Kieffer TJ. 75. Targeting the glucagon receptor family for diabetes and obesity therapy. Pharmacol Ther 2012; 135(3):247-78.

https://doi.org/10.1016/j. pharmthera.2012.05.009

Akhtar N, Ahad A, Khar RK, Jaggi M, 76. Aqil M, Iqbal Z, *et al*. The emerging role of P-glycoprotein inhibitors in drug delivery: a patent review.

Opin Ther Pat Expert 2011: 21(4):561-76. https://doi.org/10.1517/13543776.20 11.561784

77. Dimitrov D. Virus entry: molecular and biomedical mechanisms applications. Nat Rev Microbiol 2004; 2(2):109-22.

https://doi.org/10.1038/nrmicro817

Abdulkhalek S, Hrynyk, Szewczuk 78. A novel G-protein-coupled M. receptor-signaling platform and its targeted translation in human disease. Res Reports Biochem 2013; 2013(3):17-30.

https://doi.org/10.2147/RRBC.S28430

Pan X, Veroniaina H, Su N, Sha K, 79. Jiang F, Wu Z, et al. Applications and developments of gene therapy drug delivery systems for genetic diseases. Asian J Pharm Sci 2021; 16(6):687-703.

https://doi.org/10.1016/j.ajps.2021.05.003

- Mugumbate G, Jackson GE, van 80. der Spoel D. Open conformation of adipokinetic hormone receptor the malaria mosquito from facilitates binding. hormone Peptides 2011; 32(3):553-9. https://doi.org/10.1016/j. peptides.2010.08.017
- Rufer AC. Drug discovery for 81. enzymes. Drug Discov Today 2021; 26(4):875-86. https://doi.org/10.1016/j. drudis.2021.01.006
- Agarwal P, Huckle J, Newman J, 82. Reid DL. Trends in small molecule drug properties: A developability molecule assessment perspective. Drug Discov Today 2022; 27(12):103366. https://doi.org/10.1016/j. drudis.2022.103366
- 83. Rask-Andersen M, Almén MS, Schiöth HB. Trends in the exploitation of novel drug targets. Nat Rev Drug Discov, 2011; 10(8):579-90.

https://doi.org/10.1038/nrd3478

- 84. Ouertani A, Neifar M, Ouertani R, Mosbah A, Masmoudi A, Cherif A. Effectiveness of enzyme inhibitors in biomedicine and pharmacotherapy. Adv Tissue Engin Regen Med 2019; 5(2):85-90. https://doi.org/10.15406/ atroa.2019.05.00104
- 85. Deodhar M, Al Rihani SB, Arwood MJ, Darakjian L, Dow P, Turgeon J, et al. Mechanisms of CYP450 inhibition: understanding drug-drug interactions due to mechanism-based inhibition in clinical practice. Pharmaceutics 2020; 12(9):846.

h t t p s : // d o i . o r g / 1 0 . 3 3 9 0 / pharmaceutics12090846

86. Hong SJ, Lee YJ, Lee SJ, Hong BK, Kang WC, Lee JY, *et al.* Treat-totarget or high-intensity statin in patients with coronary artery disease: a randomized clinical trial. JAMA 2023; 329(13):1078-87.

https://doi.org/10.1001/jama.2023.2487

- 87. Rothwell PM, Algra A, Chen Z, Diener HC, Norrving B, Mehta Z. Effects of aspirin on risk and severity of early recurrent stroke after transient ischaemic attack and ischaemic stroke: time-course analysis of randomised trials. Lancet 2016; 388(10042):365-75. https://doi.org/10.1016/S0140-6736(16)30468-8
- 88. Turberville A, Semple H, Davies G, Ivanov D, Holdgate GA. A perspective on the discovery of enzyme activators. SLAS Discov 2022; 27(8):419-27. https://doi.org/10.1016/j.

slasd.2022.09.001

89. Peng LH, Fang JY, Dai N, Shen XZ, Yang YL, Sun J, *et al.* Efficacy and safety of linaclotide in patients with irritable bowel syndrome with constipation: Chinese sub-cohort analysis of a phase III, randomized, double-blind, placebo-controlled trial. J Dig Dis 2022; 23(2):99-110. https://doi.org/10.1111/1751-2980.13081

- 90. Singh K, Gupta JK, Pathak D, Kumar S. The use of enzyme inhibitors in drug discovery: current strategies and future prospects. Currt Enzyme Inhib 2023; 19(3):157-66. https://doi.org/10.2174/1573408019 666230731113105
- 91. de la Fuente M, Lombardero L, Gómez-González A, Solari C, Angulo-Barturen I, Acera A, *et al.* Enzyme therapy: current challenges and future perspectives. Int J Mol Sci 2021; 22(17):9181.

https://doi.org/10.3390/ijms22179181

- 92. Bagal SK, Brown AD, Cox PJ, Omoto K, Owen RM, Pryde DC, *et al.* Ion channels as therapeutic targets: A drug discovery perspective. J Med Chem 2013; 56(3):593-624. https://doi.org/10.1021/jm3011433
- 93. Imbrici P, Liantonio A, Camerino GM, De Bellis M, Camerino C, Mele A, *et al.* Therapeutic approaches to genetic ion channelopathies and perspectives in drug discovery. Front Pharmacol 2016; 7:121. https://doi.org/10.3389/fphar.2016.00121
- Connick P, De Angelis F, Parker 94. RA, Plantone D, Doshi A, John N, et al. Multiple Sclerosis-Secondary Progressive Multi-Arm Randomisation Trial (MS-SMART): A multiarm phase IIb randomised, double-blind, placebo-controlled comparing clinical trial the efficacy of three neuroprotective drugs in secondary progressive multiple sclerosis. BMJ Open 2018; 8(8):e021944.

https://doi.org/10.1136/ bmjopen-2018-021944

95. Ma K, Cheng Z, Jiang H, Lin Z, Liu C, Liu X, *et al.* Expert consensus on ion channel drugs for chronic pain treatment in China. J Pain Res 2024; 17:953-63.

https://doi.org/10.2147/JPR.S445171

96. Corry B, Thomas M. Mechanism of ion permeation and selectivity in

a voltage gated sodium channel. J Am Chem Soc, 2012; 134(3):1840-6. https://doi.org/10.1021/ja210020h

- 97. Zamponi GW. Targeting voltagegated calcium channels in neurological and psychiatric diseases. Nat Rev Drug Discov, 2016; 15(1):19-34. https://doi.org/10.1038/nrd.2015.5
- 98. Guo M, Shen W, Zhou M, Song Y, Liu J, Xiong W, *et al.* Safety and efficacy of carbamazepine in the treatment of trigeminal neuralgia: A metanalysis in biomedicine. Math Biosci Eng 2024; 21(4):5335-59. https://doi.org/10.3934/mbe.2024235
- 99. Zhang M, Gao CX, Ma KT, Li L, Dai ZG, Wang S, *et al.* A meta-analysis of therapeutic efficacy and safety of gabapentin in the treatment of postherpetic neuralgia from randomized controlled trials. Biomed Res Int 2018; 2018:7474207. https://doi.org/10.1155/2018/7474207
- 100. Lemoine D, Jiang R, Taly A, Chataigneau T, Specht A, Grutter T. Ligand-gated ion channels: New insights into neurological disorders and ligand recognition. Chem Rev 2012; 12(12):6285-318. https://doi.org/10.1021/or2000820

https://doi.org/10.1021/cr3000829

101. Keramidas A, Moorhouse AJ, Schofield PR, Barry PH. Ligandgated ion channels: Mechanisms underlying ion selectivity. Prog Biophys Mol Biol 2004; 86(2):161-204. https://doi.org/10.1016/j. pbiomolbio.2003.09.002

- 102. Tariot PN, Farlow MR, Grossberg GT, Graham SM, McDonald S, Gregel I. Memantine treatment in patients with moderate to severe Alzheimer disease already receiving donepezil: a randomized controlled trial. JAMA 2004; 291(3):317-24. https://doi.org/10.1001/jama.291.3.317
- 103. Levinson SR, Luo S, Henry MA. The role of sodium channels in chronic pain. Muscle Nerve 2012; 46(2):155-65. https://doi.org/10.1002/mus.23314
- 104. Palomino-Doza J, Rahman TJ, Avery PJ, Mayosi BM, Farrall M, Watkins H, *et al.* Ambulatory blood pressure is associated with polymorphic variation in P2X receptor genes. Hypertension 2008; 52(5):980-5. h t t p s : // d o i . o r g / 1 0 . 1 1 6 1 / HYPERTENSIONAHA.108.113282
- 105. Singh S, Malik BK, Sharma DK. Molecular drug targets and structure-based drug design: A holistic approach. Bioinformation 2006; 1(8):314-20.

https://doi.org/10.6026/97320630001314

- 106. Nurmagfirah N. Asam nukleat, protein, enzim, reseptor sebagai target kerja obat dan tranduksi signal. Makasar: Universitas Islam Negeri (UIN) Alauddin, 2014.
- 107. Zhang X, Yu W, Li Y, Wang A, Cao H, Fu Y. Drug development advances in human genetics-based targets. MedComm 2024; 5(2):e481. https://doi.org/10.1002/mco2.481