Role of Pharmacovigilance in Drug Development

Lekha Saha
Assistant Professor, Department of Pharmacology,
Post Graduate Institute of Medical Education & Research (PGIMER), Chandigarh-160012, INDIA

Corresponding author: Dr. Lekha Saha, Assistant Professor, Department of Pharmacology, Post Graduate Institute of Medical Education & Research (PGIMER), Chandigarh-160012, INDIA, Tel: +91-0172-2755253, +91-9463503752 ; E-mail: lekhasaha@rediffmail.com
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Pharmacovigilance (PV) is the science and activities relating to the detection, assessment, understanding and prevention of adverse effects or any other drug related problem. PV collects, records, codes ADEs/ADRs analyses and assesses the reports, promotes the safe use of drugs, creates appropriate structures and means of communication needed to perform its tasks. The aims of PV are to improve patient care and safety, improve public health and safety, contribute to the assessment of benefit, harm, effectiveness and risk of medicines, promote education and clinical training, promote effective communication to public and promote rational and safe use of medicines.

The process of drug discovery and development is a long-term, competitive, expensive and complicated process. Bringing the drugs from the bench to the market, that is, from screening and identification of the drug as a compound to its introduction to the market, takes several years of efforts [1]. The process of discovering and developing a new drug involves an intricate interaction between investors, industry, academia, patent laws, regulatory authorities, marketing and the necessity to balance confidentiality with communication [2]. The complete process of presenting a drug to the patients involves four stages- 1. Drug discovery, 2. Drug development, 3. Regulatory review and approval, and 4. Marketing [3].

Drug Discovery

In the process of drug discovery, the first step is to identify an appropriate ‘drug’ target [4], which can be a biomolecule or a protein receptor that is explicitly associated with a disease condition or pathology. After the target has been identified, the next step involves target validation and the confirmation of its role in the disease progression. This is followed by testing of the target against different small molecule compounds to identify lead compounds which can interact with the target biomolecule and display the potential therapeutic activity either by nullify or slow the disease development. The lead compounds can be identified by screening a library of compounds through various methods, such as high-throughput screening [5], de novo synthesis [6], and isolation from the natural products [7].

Drug Development

The drug development phase involves stringent testing and optimization of the selected compounds to identify the ‘drug candidate’ which might be most effective in terms of safety, toxicity [6], dosage, and efficacy. For this purpose, the selected lead compounds are tested in cells (in vitro) and in animals (in vivo) to study their pharmacodynamic and pharmacokinetic properties, which include Absorption, Distribution, Metabolism, Excretion and Toxicity (ADME/Tox) properties. The successful lead candidate must be non-toxic and should have good bioavailability, can be distributed to the drug target in the body, and can be metabolized efficiently and effectively as well as successfully excreted from the body. This part of the development process is referred to as the ‘preclinical phase’ in which the drug candidate is meticulously examined, optimized, and prepared for testing in humans. This phase is followed by the ‘clinical phase’ of development, in which the efficacy and safety of a drug candidate is scrutinized in patients. This ‘clinical trial’ has 3 phases: Phase 1 perform initial human testing in a small group of healthy volunteers to demonstrate the safety and pharmacokinetics, phase 2 involves testing in a small group of patients to demonstrate the safety, efficacy and pharmacokinetics and phase 3 includes testing a large group of patients to show safety and efficacy of the drug candidate in them since the healthy and sick people have potentially different metabolic patterns for the drugs.

Regulatory Review and Approval

The outcome of the ‘clinical trial’ decides whether the drug candidate is safer and effective enough in treating the disease. At this point, new drug applications (NDA) with all the essential evidence, including quality, preclinical and clinical data collected during development of the drug candidate, are submitted to the relevant regulatory authorities, e.g., the United States Food and Drug Administration (USFDA), which oversees the development, approval, and marketing of drugs [8]. They need to approve the drug applications so that the company can commercialize the drug in their jurisdictions (e.g., New Drug Application (NDA) in USA, and Marketing Authorization Application (MAA) in Europe).
Marketing (Commercialization)

This is the last phase of drug development process. Once the drug has been approved, it is marketed or commercialized. The drug manufacturers need to submit marketing authorization applications in every country in which they want to sell the drug. Following this, post-marketing surveillance is conducted by the manufacturer to continue evaluating the safety and efficacy of the marketed drug, and to better design its further development. These studies are considered as phase 4 clinical trials and are compulsory in some countries, e.g., Japan and Philippines. As the drug is typically targeted to a very large number of patients, the manufacturer is expected to monitor this stage cautiously and submit reports to the FDA. The reports include evidence for medicine-related problems, e.g., treatment failure, adverse reaction, counterfeit/poor quality medicines, drug interactions, or incorrect use. These reports are significant in terms of generating proof of efficacy that will inspire public confidence and trust.

The entire process of drug discovery and development costs around $0.8 - 1.0 billion per drug [11,12] and takes about 10-15 years and even then, there is a very high attrition rate for new drug candidates in clinical stage. To deal with the issues involved in failure of new drugs, USFDA has launched the “Critical Path Initiative and the Critical Path Opportunity” program in 2004 to guide the development process of new drug [13].

Introducing a new medicine to the patients is not only a highly time-consuming and expensive process but also requires an extremely essential and strict vigilance on the safety and efficacy of the drug. Therefore pharmacovigilance plays a pivotal role in drug discovery and development. The drugs may appear to be safe and well-tolerated in preclinical and clinical testing, but their safety in the ‘real world’ may not be distinct, after-effects of the drug when used frequently or in combination with other drugs are generally unknown, safety in vulnerable groups with different metabolic profiles (e.g. pregnant women and breastfeeding mother, elderly person, young children) can be uncertain, and rumors and myths can destroy the integrity, adherence to, and success of a treatment.

Pharmacovigilance is the pharmacological science which deals with drug safety including accumulation, detection, assessment, monitoring, and prevention of adverse effects of the drugs. It is an iterative process focusing on detection of unidentified safety issues, identification of risk factors, quantifying risks and preventing patients from being adversely affected unnecessarily. Pharmacovigilance plays a critical role at various stages of drug discovery and development process; for example, in clinical research, pharmacovigilance requires submission of the reports on adverse events during clinical trials to regulatory authorities within a specified time frame, notification of such events to all investigators and ethics committees, and a safety review by independent Drug Safety Monitoring Boards (DSMB). Annual reports, a summary and analysis of all the serious adverse events, new safety findings from animal studies, and evaluations of benefit and risk are also required. Pharmacovigilance also plays a significant role when the drug is commercialized. Reporting the safety reviews is mandatory for companies in a marketing phase. These safety reviews include Risk Management Plan (RMP), Periodic Benefit Risk Evaluation Report (PBPER), the Development Safety Update Report (DSUR) [14], Periodic Safety Updates Report (PSUR) [15], phase 4 studies (post-marketing surveillance), clinical trials (intervene disease management), and pharmacoepidemiological studies (non-interventional or observational).

Pharmacovigilance is a very critical and inevitable part of the drug discovery and development process. It will require comprehensive documentation and severe monitoring at every phase of drug development including pharmacovigilance inspection and audit, risk management, and reporting of ADR medicinal drugs, periodic safety update report, post-authorization safety studies, additional monitoring, and safety communication. Therefore, it is absolutely essential to establish good pharmacovigilance practices for improving the understanding of the drug safety issues during the drug development and its post-approval so that the attrition rates can be reduced and the patients can be provided with safe and efficacious innovative medicines to meet their prerequisite medical needs.

References