Challenges and Future Consideration for Pharmacovigilance

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Abstract
Pharmacovigilance is studying the safety and effectiveness of drugs and practice to minimize the possible risk associated with the use of drugs. It has developed significantly and will continue to develop in response to the new challenges it faces. With the rapid spread of information across the globe, increased communication across borders, easy access to variety of medicinal products and increasing public expectation of safety, there is a need for careful strategic planning adopting a holistic approach to meet the additional challenges. We need continuing and dynamic progress of all aspects of pharmacovigilance to improve public health and safety. There is a need for new global network to share information and intelligence about benefit and risk of medicinal products.

Keywords: Pharmacovigilance; Medicinal products; World Health Organization

Introduction
There is always a trade-off involved between the advantages and the possible risk with the use of pharmaceuticals. The risk is minimized when medicines of good quality, safety and efficacy are used rationally by an informed health professional and by patients. Pharmacovigilance helps in reducing the risk of harm by ensuring good quality medicines are used appropriately and expectations and concerns of patients are considered when therapeutic decisions are made. It ensures that health care providers and consumers have the information they need to make their decision about treatment. Pharmacovigilance has emerged as science to examine the safety and efficacy of drugs and other therapeutic products. The World Health Organization (WHO) defines pharmacovigilance as discipline to identify, review and understand drug adverse effects and taking required precautionary actions to curtail these drug linked problems [1]. Its importance has been critical to public health and has become an integral part of effective clinical practice. The first systemic international efforts to address drug safety were initiated in 1961, following the thalidomide tragedy [2,3]. When thalidomide was introduced, the pharmaceutical industry comprised of number of emerging competing companies and many of them were not dedicated to drug development. Also there were very few or no laws regulating drug development and marketing. Thalidomide use by pregnant mothers resulted in birth of thousands of congenitally deformed infants [2]. In response, efforts were initiated to address the drug safety issues by developing a system for detecting previously unknown or poorly understood adverse effects of medicines. Guidelines were developed to monitor drugs, foods and environmental contaminants for adverse reactions and toxicity. In the beginning these guidelines were to meet local and national needs. With globalization of world economy, free trade, increased communication across borders and increasing use of internet resulted in recognized need of a system, applicable internationally, to ensure safety of medicinal products. These efforts and recognition of system to monitor and promote the safety and effectiveness medicines lead to emergence of practice and science of pharmacovigilance.

Originally, pharmacovigilance purpose was to improve scientific understanding of safety profile of drugs and advise regulatory authorities. It has been limited largely to detection of unidentified adverse drug effects and support public health by providing reliable scientific information. However, there has been a growing awareness and realization that pharmacovigilance has the potential to progress beyond its confined limits and the scope of pharmacovigilance should be expanded to include new kind of safety concerns such as illegal sale and drug abuse over internet, increasing self-medication practices, widespread manufacture and sale of counterfeit medicines [1,4,5]. In recent times, the range of pharmacovigilance has been expanded to include, herbal medicines, blood products, biologics, medical devices and vaccines [5-7]. Additionally, there has been increasing use of traditional/herbal medicines outside the boundaries of the traditional culture and with other medicines that can cause the potential adverse interaction [7,8]. The pharmacovigilance faces new challenges by increasing demands and availability of range of new products. This article briefly describes some of the serious challenges pharmacovigilance faces and their potential implication.

Drug Regulation
There is a need for strong links within drug regulators to ensure that authorities are well briefed on everyday practice safety issue. There has been a substantial increase in number of clinical trials around the world. There is also an increasing partnership between academia and drug industries. This gives rise to widespread concern over ethical and scientific issues such as potential conflict of interest, unethical patient recruitment process, lack of competence for clinical trial monitoring, unethical clinical practice and poor reporting of adverse events [9]. Therefore drug regulators have increasing responsibility to ensure that right and health of patients and their commitments are protected. The growing intricacies of clinical trials also present further challenges to regulators [10]. Since clinical trials are usually carried out at multiple locations in different countries, local regulators are incognizant of investigator’s proceedings at other international locations. Regulators often do not have capacity to efficiently ensure proper conduct of clinical trials. There is a need for standardized reporting systems for safety concerns arising during clinical trials and better coordination between ethics committees and investigators.
The process of new drug safety evaluation continues post marketing stage and regulators makes the final judgment regarding drug release considering detailed pharmacovigilance study. This is also important in introduction of generic medicines or safety profile of older medicines if there are new safety concerns. Some additional drug safety aspects such as excipients input to the safety profile, comparing safety profile of similar medicines and assessment of the adverse effects on human health of drug residues in animal needed to be included in long term effect study of medicines.

Globalization

The globalization of world economy has resulted in worldwide drug distribution and the increased exposure of massive populations to large volumes of medicines. The free trade and communication across borders and increasing use of internet have resulted in rapid spread of drug information and access to all medicinal products. These changes have resulted in new kind of safety concerns such as illegal sale of drug over internet, increases self-medication, public exposure to substandard products etc. The growing public awareness with global exposure has also heightened public expectation of safety and national pharmacovigilance centers are in further pressure to address all safety concerns [11]. There is a recognized need for competent and rapid communication between national drug regulatory centers as well as to other countries.

Web-Based Sales and Promotional Activities

Public health programs and media coverage aimed at increasing public awareness have made public in many countries to increasing influence health professional’s prescribing and pattern of drug use. The Internet has encouraged the unrestrained sale of medicines across borders. Drug advertisement with unreliable claims is distributed internationally through the web. It ranges from prescription medicines, unregistered drugs and controlled substances to herbal medicines with suspicious quality and safety. Aggressive promotion by drug companies often results in unnecessary and irrational use of medicines by consumers [12]. Direct advertising to the consumer of prescription medicine has also become common in many countries. With this information patients feel competent to take greater responsibility for their own therapeutic decision without proper assistance from physician or pharmacist. It is welcome development in some situations; however available information is not always reliable and scientifically valid and produces serious health risks. This results in increasing self-medication, illicit availability of medicines over internet, and also physician overprescribing succumbing to patient demand [12].

There are increasing public concern that close collaborations between academia and the pharmaceutical industries may unfavorably influence medical practice and clinical research. A disturbing development is “direct to consumer” advertising by drug makers, sellers and promoters with vested interests. The lack of reliability and accuracy of information provided to consumers may compromise patient care and safety. Furthermore, the internet provides a medium that makes communication possible across the borders even when direct advertising of prescription medicine to consumer is illegal. This makes national regulators irrelevant regarding advertising. These issues put forward the need for detailed monitoring of drug safety and scrutiny of advertising. It is necessary to ensure that promotional material provide truthful and unbiased information.

Drug Utilization Pattern

While drugs are a prominent item in health budget, the cost of disease to society and money spent on detecting; preventing and managing drug’s adverse effects are also included in cost consideration. Many studies have suggested the huge impact that poor product quality, adverse drug reactions and practice errors have on health care and patient health [13].

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These changes in drug use and practice have likely to have important consequences on public health and safety. Thus there is a need to ensure that existing centers are efficient and effective, their benefit on public health are evident.

Biologics and Biosimilars

Biologics have specific characteristics, which complicate their safety assessment and require modified systems of safety monitoring. Biologics are often prescribed for rare diseases, which make it difficult to include ample number of patients in the pre-approval clinical trials. Moreover, biologics are frequently used in multiple indications with different dosage regimens, which further complicate the exposure assessment of biological. They are often prescribed to patient with severe disease as second- or third-line therapy after the failure of standard treatment. Thus, pharmacovigilance for biologics involves additional challenges compared to small molecules.

Additional challenges are met for biosimilars, as they cannot be approved based on the same requirement that apply to generics. Biosimilars are made by a complicated production and purification
process, which differs from one manufacturer to another [17]. Thus, it is very unlikely that manufacturer of the biosimilar will be able to precisely replicate the reference protein product as the original method is proprietary information of the owner of the reference product. It is recognized that small changes in the production and purification process of biologicals alter their safety profile as an adverse event can be related to a specific biological product [18,19]. Thus, regardless of the fact that the biosimilar and reference drug can show similar efficacy, the biosimilar can show different safety profile, which may influence the benefit-risk profile. It is essential to supervise the clinical safety and risk-benefit evaluation of biosimilars on a regular basis during the post-approval phase [20]. The safety profile of biosimilars should be established by identifying immunogenicity risk and appropriate post marketing supervision.

Herbal and Traditional Medicines

There is a common belief that “herbal” means “safe” and long-term use of traditional medicine assures its efficacy and safety. However, it is a misconception and irrational use of herbal and traditional medicines raises serious concerns about safety. Recently, traditional medicines have moved local boundaries and being manufactured for global use and there has been increasing number of cases of traditional and herbal medicines being adulterated or contaminated with other medicines [7]. Further, self-medication by consumers heightens the risk. The use of traditional and herbal medicine in combination with other medicine can cause very serious adverse drug interactions [21]. Therefore, these products should be included into regulatory framework like other pharmaceutical products. There should be standard of safety, quality and efficacy governing these products similar to other products intended for human use. But, lack of clear boundaries between foods and medicines get in the way in development of regulatory guidelines. The regulatory status of herbal products differs from country to country. These regulatory discrepancies between countries have serious implication for international availability of such products. If one country restricts the availability of an herbal product by prescription only, it may be obtained from health food shop or by mail order or internet from another country.

Conclusion

Pharmacovigilance has emerged over time as an essential part of public health program but encounters some crucial challenges. Its range is increasing with number and diversity of pharmaceutical products rising. The globalization of world economy and increased public exposure to large number of medicines, calls for a better coordination between drug regulators. There is a need for better integration of pharmacovigilance into clinical practice and public policy. A complicated and essential relationship exists between numbers of partners in drug safety monitoring. Sustained partnership and commitments are necessary between these partners to meet the future challenges in pharmacovigilance. The present pharmacovigilance systems need to be reviewed and evolved further to address future challenges.

References


