ADVERSE DRUG REACTION

1*Soheil A Pathan. 2 Dr. A M Shaikh.
AAEMF’s Delight College of pharmacy, Pimple jagtap road,Koregaon Bhima, Maharashtra 412216

ABSTRACT: Drug instability (Adverse Drug Reaction) is a major problem for modern medicine, mainly due to the complexity of treatment, the aging process and the growing number of diseases. This article addresses some of the key issues related to ADR, as well as issues related to their prevention, diagnosis, reporting, factor affecting and current treatment.

KEYWORDS: Adverse drug, WHO, Side effects, Pharmacodynamics, Etiology, Occurrence, Magnitude of ADRs.

I. INTRODUCTION:

HIGHLIGHTS:
• Adverse drug reactions.
• Medical pharmacology.
• Adverse drug reactions and adverse reactions.
• Factors affecting on ADRs.
• Pharmacovigilance reporting system.

ABOUT ADVERSE DRUG REACTIONS:

DEFINATION: According to the world health organization (WHO) An ADR is defined as “a response to a drug which is noxious and undesired and which occurs at normal doses when used in man for prophylaxis, diagnosis or therapy of disease or for the modification of physiologic function.” Thus an ADR is a type of ADE who cause can be directly related to a drug and its physiological properties.

Adverse drug reaction (ADR) is an unwanted, undesirable, effect of a medicine that occurs during usual clinical application. It occurs almost daily in health care institutions and can adversely effect a patients quality of life. Significant research in the United States and the United Kingdom from the late 20th to the early 21st century shows that medical side effects are common, including the causes of unexpected hospitalizations that occur in the hospital and appear after treatment. NPD levels do not change over time and, despite various precautions, 5% to 10% of patients suffer from side effects after taking, taking, or leaving home. The extent of incidents depends on the way these incidents are detected, and most NPDs do not cause violent incidents. However, this high risk is associated with morbidity and mortality, is costly and can adversely affect the doctor-patient relationship and should be carefully considered.

Hospital medications, especially those containing ADR, include antiplatelet agents, anticoagulants, anticancer drugs, immunosuppressants, diuretics, antidiabetic drugs, and antibiotics. If there are fatal side effects, it is often associated with bleeding and is suspected to be the most common cause of oral anticoagulants / anticoagulants and NSAIDs.

A RASH DUE TO A DRUG REACTION
KEY ALIMENTS:
Adverse drug reactions (ADRs) - unexpected, drug-related side effects - occur as the cause and timing of a significant proportion of unplanned hospitalizations. Careful medical history can help the patient understand previous experiences with the drug, especially previous side effects that preclude re-exposure. Prevention of ADRs depends on the refusal of treatment in cohorts of hypersensitive patients or the provision of treatment as part of a treatment plan that reduces the risk of side effects (eg, concomitant use of other drugs, monitoring of blood test results). Spontaneous reports based on ADR suspicion (using the yellow card scheme in the UK) are an important part of pharmacovigilance, but in general, health facilities and sectors are poorly informed about ADRs. If in doubt, it is better to report.

BACKGROUND:
Drug-related adverse reactions (ADRs). According to WHO (World Health Organization) is defined as "a drug reaction that harms people and is deliberately used for treatment." Various factors can result in unforeseen errors, such as medication or incompatibility. Uncontrolled outcomes can have a profound effect on a patient's quality of life and health. Different outcomes are the leading cause of illness and death worldwide, and will continue to be a serious problem for human health and the complexity of medications to treat various diseases in the elderly. A recent study found that in addition to about 3.5% of hospital cases, catastrophic effects in Europe cost up to £197,000 annually. The causes of drug-related side effects are very complex and varied. Adverse events can be divided into the following categories: Diet / Exercise Disorders or Mental Health. Side effects of medications are usually dietary, but sometimes unavoidable. It depends on the patient's healing difficulties and the combination of medications used. There are usually no side effects, but they are common. Infectious diseases occur when a patient does not respond well to medication that can prevent dehydration immediately, including skin care and prompt good advice. The effects of the drug are not well understood and are not very serious. Few people get sick, the consequences can be managed.

The risk of patients taking certain medications, such as adults, is a major problem. The study found that 75% of all STIs received some form of treatment after being transferred from hospital to primary care. Many of the side effects of drug use are caused by common medical problems. According to a large previous study, the error rate in the English system is 5%. With the introduction of technology into medical systems, the implementation of high-level error-based computer support systems can lead to poor or difficult outcomes. Currently, there are limited data and evidence on the prevalence of adverse reactions. Following the previous literature review (Cochrane, JBI data on performance reviews and performance reporting. Ovid MEDLINE), there was no systematic review or assessment that provided an overview of drug activity. Most available studies are small and are usually limited to one group. Obviously, the new study focuses on specific measures and medications to reduce treatment failures and errors. While definitions of drugs exist that focus on the consequences of medical errors and the effectiveness of procedures, they do not explain the types of side effects. A 2017 study by Khalil’s colleagues tested the effectiveness of various treatments for reducing mortality, emergency care, and hospitalization. The authors found little evidence to support the professional and institutional benefits of resolving treatment errors. Assiri et al., Investigated 2018 cases of drug failure and related effects and risk factors. Controversy has been raised over the definition of drug risks, the methods used to assess side effects, and the various levels of risk.

MECHANISM:
ADR can be described as unacceptable and dangerous due to drug intervention. The aetiology of adverse reactions can be divided into studies of direct toxicity and hypersensitivity reaction, which occur due to changes in the pharmacokinetics and pharmacodynamics of the drug product. Specific toxic effects may result from toxic effects of compounds or metabolic processes seen in various organs, as well as from harmful chemicals, physiological dysfunctions, DNA damage, or damage to structures and tissues of our cells. On the other hand, the effects of hypersensitivity can be explained after a person’s immune system shows a drastic change in medication or metabolism, which includes allergies and anaphylaxis. This process is the result of direct cytotoxicity and an overactive immune system seen in various organs, such as the skin, liver, lungs, brain, and kidneys.

MAGNITUDE OF ADRs:
Side effects are one of the leading causes of disease and mortality in health. The Institute of Medicine in the USA (2000) reports between 44,000 and 98,000 deaths annually caused by ADRs. The total number, an estimated 7,000 deaths occur from side effects. 39 studies examining the US pharmacy system over four decades showed that in 1994, 106,000 people died due to side effects. More than 2 million people have been affected by serious side effects (Pomeranz and Bruce, 1998). These figures showed that the trend of death and injury increased the side effects. Side effects are the fourth leading cause of death in the United States after heart disease, cancer, and stroke (Jemal et al. 2005). In another review conducted by the American Society of Pharmacists, Byrne et al. (2006) found that 85% of patients who answered the questions expressed concern about at least one drug-related network, such as taking borrowed drugs, having a harmful pharmacovigilance effect, or receiving the illegal drug. Side effects are a significant public health problem in the world. Side effects not only cause deaths and injuries, but also affect the length of stay in hospital, which in turn increases healthcare costs and reduces patient productivity. Moura et al. (2009) The frequency of adverse reactions in intensive care units decreased and its effect on the duration of care assessed and found after adverse reaction presented by a patient increased by 2.38 days in the intensive care unit. In research conducted at the University of Liverpool, an estimated 18,820 patients. It turned out that the total amount of 1225 side effects was reported, giving a prevalence of 6.5%. The average stay was 8 days, which was calculated for 4%
of the hospital bed capacity (Nainggolan, 2004). Another prospective cohort study was conducted to evaluate outpatient prescriptions for more than 1,200 patients, to map and review over a period of 4 weeks. The researchers found that 25% of patients with side effects had experienced selective serotonin reuptake inhibitors, beta blockers, angiotensin-converting enzyme inhibitors and non-steroidal anti-inflammatory drug classes. The incidence of adverse reactions has reached 27 per 100 patients (Gandhi et al., 2005). The ADR report is not yet fully developed. "The need for increased awareness of the importance of withdrawing ADR in Malaysia (Aziz et al. 2007)."

DATA SOURCES:
A systematic review of the literature between search in the period 1991-2012 was based on PubMed, Cochrane database of systematic reviews, EMBASE and IDIS. The article included original studies, WHO, FDA, reports and conventional medical records. Keywords used are: medical error, medication, side effect, iatrogenic disease, other factors, outpatient care, primary care, side effects and emergency care. The research was done at the time of publication.

FACTOR AFFECTING THE OCCURRENCE OF ADRs:
Kitteringham et al. (1994) recommend that for many adverse effects, especially idiosyncratic responses to drugs, there may be a complex pattern involving not only genetic defects in many areas but also environmental factors such as coexisting infections or drug use in various diseases. Many of these side effects are due to the broad pharmacological action of the desired drugs, often due to the wide variability in pharmacokinetics and pharmacodynamics observed in patients. Pharmacological, immunological and genetic factors involved in the pathogenesis of ADR. Factors that predispose to pharmacological side effects are dosage, drug composition, pharmacokinetic or pharmacodynamic abnormalities, and drug interactions. The conversion of metabolic drugs to metabolism has now been recognized as a requirement for a great number of specific drug responses (Masubuchi et al., 2007). The correlation between high levels of drug metabolism, decreased detoxification, or decreased cellular defense against reactive drugs appears to be an important trigger (Guengerich and MacDonald, 2007). Immunological and genetic factors can play a role in how the body responds to the drugs it is administering. Rusticitas et al. (2004) also suggested that ethnic diversity plays an important role in the development of RAM. Evans (2005) found that some risk factors include side effects and several classes of therapeutic drugs, among other specific classes. Carefully recommend high-risk factors based on patient characteristics (gender, age, weight, creatinine, diabetes, and number of comorbidities) and drug (dose, route of administration, number of concomitant medications). Factors that may increase the likelihood of an effect are: extreme age, gender, use of multiple medications, medical condition, history of side effects or allergy, genetics, high doses, etc. - Drug withdrawal or dose adjustment may be important in the development of side effects in some people, especially the elderly. The Health Research Agency (2001) suggested that another possible cause of potentially harmful side effects for physicians was the reluctance to seek treatment with appropriatedoses of drugs for fear of causing toxicity. Side-effects can be caused by mistakes in preparing, administering, prescribing, administering or administering drugs. Eighteen percent of drug-related adverse events identified in the Harvard Medical Studies Study define neglect as failure to follow a medical procedure reasonably expected by a qualified physician for patient care (Lea et al., 1991). Bates et al., 2003a, Bates et al., 2003b suggested that adverse effects often occur during the observation period and can be avoided or corrected with a few simple methods. Factors influencing the occurrence of ADR were divided into five groups. Patient factors, social factors, medication factors, disease factors, and adverse event factors.

A) PATIENT RELATED FACTOR;

1) AGE:
All these drugs can cause side effects, but not all patients will experience side effects equally and generally. Age is very important because it affects the outcome of side effects. Elderly patients with multiple health conditions, having multiple medications, having adverse effects, and having reduced ability to clear drugs are at high risk of adverse effects. A study by Debellis et al. (2003) From the incidence and prevention of adverse ambulatory effects in elderly men, they concluded that adverse effects are common and often avoidable in elderly individuals in the ambulatory clinical setting. More serious side effects can be avoided. Prevention strategies should target prescriptions and pursue levels of pharmacy care. The intervention was intended to promote the patient's adherence to the prescribed regimen and to monitor prescribed medications. Elderly and pediatric patients are particularly at risk for side effects, as these drugs are less likely to be studied in extreme age groups, and medication absorption and metabolism are more variable in these two groups and less foreseen. Efforts are needed to prevent and prevent side effects in children, infants and infants at risk of developing side effects, because the ability to metabolize the drug has not been fully estimated. Certain factors affecting the development of side effects in newborns (Clavenna & Bonati, 2008)
- Having immature renal tubular functions, when under 8 weeks old, digoxin, aminoglycosides, ACE inhibitors, NSAIDs are a must.
- Hypoalbuminemia affects physiological drug dosage in newborns. Caution is advised when dealing with important protein binding drugs such as NSAIDs.
- Children born with low body fat they might be affected by unbound fatty drugs.
- Anesthetic effects increased due to a delayed premature cerebral hemorrhage less than 8 weeks of age.
- Predisposition to hypotension due to poor cardiac compliance and premature baroreceptors.
People are at high risk of developing side effects for a number of reasons. People tend to have many health issues and hence have many prescriptions and over the counter medications. As the elderly lose their ability to metabolize liver drugs. Even older adults are guaranteed side effects more than twice as severe as younger ones. As people age, the amount of water in the body decreases and the amount of fatty tissue increases relative to water. As in adults, drugs that are dissolved in water reach a higher concentration because there is less water to dissolve and drugs discontinue more in fat, which is to accumulate fat in the substance bodily. Also, as people age, the kidneys are less able to eliminate drugs in the urine, and the liver is less able to metabolize many drugs. Jimmy and Padma (2006) in their study concluded that the incidence of ADRs in adults and the elderly was significantly higher than other age groups. Although gender-related adverse reactions developed and differed between age groups, type A reactions in the elderly (85.9%) were the most common and type B reactions in adults (35.9%, %) were more frequent at other ages. Due to all the age-related changes, many drugs tend to stay in the larger body rather than the smaller body, prolonging the effects of the drugs and increasing the risk of trauma.

2 GENDER:
Biological differences affect the activity of multiple drugs in men and women. Anatomical and physiological differences are body weight, body composition, factors of gastrointestinal tract, liver metabolism, and kidney function. Females compared to men have lower body weight and organ size, more body fat, lower gastric motion and glomerular filtration rate. These differences can affect the process of treating drugs by changing the pharmacokinetics and pharmacodynamics of drugs, including drug absorption, distribution, metabolism and elimination. The genus produces effects with adverse effects. The study of gender differences in the effects of antiretroviral drugs indicates potential gender differences in the frequency and severity of adverse effects of antiretroviral drugs. Liver CYP3A4 kinase is more active in women than in men, leading to various effects on the metabolism of drugs. Also, women are more prone to advancing abdominal pains than men in the management of narcotics that spend much of cardiac reactivation. Females due to acute and chronic health problems limit their activity to about 25% more days per year than men, spending about 40% more days in bed each year than men. Women aged 17 to 44 have twice as many doctors visits and hospitalizations as men. With the exception of six generative and other specialized conditions, the difference in hospital rooms is virtually nil, but the difference in outpatient care is still around 30%. After 45 years, excluding six specific conditions, women continue to have more than 10–20% of medical visits, compared with men who have a greater frequency of hospitalization (Ensom, 2000).

One of the most consistent health studies is that women report more symptoms of the disease than men. It is unclear whether this is due to the severe or severe clinical severity of the disease, or the following variance: Behavioral disorder - women are more likely to define discomfort as a symptom than men; Symptom view - women's anxiety about physical discomfort increases the symptoms of symptoms and assess their symptoms as a disease; o Educational reports: Women have the opportunity to remember and report symptoms.

3 MATERNITY STATUS:
Implications for the pharmacological treatment of pregnancy. Not only women are affected by the drug, but the fetus will also be exposed to adverse drugs. There are some physiological changes that occur during pregnancy that can affect the pharmacokinetics and pharmacodynamics of drugs: these are changes; Total blood volume increased by 30-40% (1500-1800 ml), increased extravascular volume during the second and third trimesters, which caused a decrease in the plasma iron revision process, and some drugs improved kidney function with increased renal plasma. hemorrhage by 30 and GFR increased by 50%, serum protein 1-1.5 lower; As a result, drugs for the kidneys have more rate of elimination. Cardiovascular changes are noted, an increase in heart rate of about 32%, an increase in heart rate (10-15 bpm) and an increase in stroke volume, and blood pressure is relatively constant. The movement, acidity, and sound of the GIT decrease during pregnancy and this may impede absorption or excretion of a drug, and ultimately drug metabolism may be affected by some stages of pregnancy. Drugs during pregnancy can affect the mother or the embryo or both. The impact of drugs on fetal organogenesis is crucial, as it can cause teratogenicity and dysmorphogenesis. Many medications, for example, antihypertensive drugs such as angiotensin kinase (ACE) inhibitors and angiotensin II receptor blockers, pose a risk to the normal health and development of the fetus.

4 ALLERGY:
They can induce sensitive drug independent cross-reactive antigens, which can manifest as allergy drugs. The existence of such cross-reactivity is supported by the medical literature. After the first sensitization to the causative drug, the second exposure causes T cells and antibodies to enter the evoked phase, which responds to specific immune reactions (Gell and Coombs Classification). Most allergy medications were observed with type I or IV reactions; type II and III reactions were only infrequently encountered. The formation of immune complexes, a common event in the normal immune response, usually occurs without symptomatology. Rarely do immune complexons bind to endothelial cells and induce immune complex deposition with complementary activation in the blood vessels. Type III clinical signs include serum sickness reactions (e.g., β-lactams), mediation induced lupus erythematosus (e.g., quinidine), and vasculitis (e.g., minocyclina). Drug-mediated T-cell hypersensitivity may have a variety of clinical manifestations, ranging from skin-only onset to systemic diseases such as lightning. Often the drugs that involve are sulfa antibiotics and β-lactams.

5 CREATININE CLEARANCE CATEGORY:
Creatinine clearance is important for renal function, which is responsible for the clearance of various medicines. Any change in the drug's renal toxicity profile increases or decreases the therapeutic effects. Kidney disease affects drug clearance and metabolism. Veniz (2000) concluded that chronic renal failure affects both the eliminated drugs and the liver, as well as the metabolic effects of uremia drugs on the liver during renal failure. This, in turn, may affect the regulation of metabolic drugs through changes in plasma protein binding and hepatic metabolism. Sol et al. (2006) stated that drug carriers, as well as metabolic enzymes, can reduce the clearance of drugs responsible for renal deficiencies in patients with renal impairment. This reduced activity of metabolic enzymes may affect medication clearance (Naud et al., 2008). The effect of kidney disease on drug excretion.
does not lead to any renal disease associated with renal failure affected by this deficiency and increases the capacity of specific ADRs in that patient.

6 FETAL DEVELOPMENT:
The fetus, which is exposed to drugs in the maternal bloodstream, is very sensitive to the effects of drugs, because it is small, low in blood plasma protein and has a weak ability to metabolize and get rid of drugs. When drug molecules reach the fetus they may cause teratogenicity (anatomical malformations) or other ADRs can cause it. The gestation period is divided into three quarters; first, second and third trimesters. The effect of drugs in each trimester varies according to the degree of development of the fetus. The drug is likely to be teratogenic, especially when taken during the first trimester of pregnancy when the fetal organs are formed. Drugs taken during the second and third trimesters, ADRs in the newborn (birth to month) or baby (month to year) tend to show delay, respiratory problems, infection or bleeding. In general, the effects are mainly determined by the type and quantity of drugs, the duration of exposure, and the degree of growth and development of the fetus with drug exposure. Both therapeutic and non-therapeutic drugs can affect the fetus.

B) SOCIAL RELATED FACTORS:

1) ALCOHOL CONSUMPTION:
Alcohol affects the metabolism of many drugs and contributes to side effects. The interaction between drugs and alcohol is related to the possibility that alcohol can alter the intensity of adverse events, which more or less harms the patient, either in a pharmacokinetic or pharmacodynamic manner. Concomitant alcohol use with certain drugs can cause several side effects, such as nausea, vomiting, headache, drowsiness, poor coordination, hypotension, and many other side effects. Heavy internal bleeding can occur when alcohol is used in combination with non-steroidal anti-inflammatory drugs in patients with peptic ulcer or gastritis. Chronic alcohol consumption activates enzymes that convert some drugs into toxic chemicals that can damage the liver and organs in the body. Alcohol can also activate sedative effects and medications to increase its effects on the brain. Alcohol can affect hepatitis cirrhosis and hepatitis, which affects the body’s ability to use drugs for metabolisation, especially hepatic and metabolic drugs that support initial metabolism. For example, the toxicity of beta blockers increases liver problems. This will ensure that physicians and pharmacists are informed of the health risks associated with alcohol and patients with pulmonary tuberculosis. Alcohol-related drugs can be more severe than mixed-aged patients, as alcohol can cause many age and health problems.

2) SMOKING RELATED FACTORS:
Smoking is a risk factor for many diseases, such as peptic ulcer, cancer and heart disease. It also affects metabolic processes involving liver enzymes, which trigger the activation of hepatic cytochrome P-450 (CYP) isoenzymes 1A1, 1A2, and possibly 2E1. Many antibiotics that replace CYP1A2 can stimulate their metabolism in smokers, thereby reducing the necessary pathogenesis. This nicotine drug is not made entirely from tobacco. Because nicotine stimulates the body’s metabolism, it can attenuate the side effects of many medications. Numerous studies around the world have shown that the interaction between smoking and theophyll, flecainidum, insulin, mouthwash, beta-blockers, thiotyoxene and H2 drugs are the factors that can affect the response to smoking. Medical research has shown that diabetics need 15-20% more insulin than non-smokers and that up to by 30% when they smoke heavily. Smoking increases the clearance of heparin, probably due to the intensity of tobacco smoke, with increased binding of heparin to antithrombin III. Nicotine-induced skin vasoconstriction may reduce the rate of insulin secretion after physical activity (Schwing et al., 1999). Smoking reduces the effect of beta blockers on the blood and heart.

C) DRUG RELATED FACTORS:

1) POLYPHARMACY:
Taking multiple medications, both prescription and over-the-counter, contributes to the risk of ADR. The number and severity of ADRs increase disproportionally as the number of drugs taken increases. There are many definitions are applied for polypharmacy. This involves prescribing too many medications for a particular patient, and may increase the risk of ADR. The more drugs are prescribed, the larger the range of polypharmacies, this does not necessarily mean that patients do not have many drugs. Polypharmacy results from many conditions; Patients suffer from more than one disease, especially in the elderly. Patients may seek more than one prescription at a time in several diseases, or acute or chronic conditions. ADRs can be caused by drug interaction, synergism, duplication, additive effect, discontinuation of therapies, changes in dosing to save money, omitting certain medications, and physiological difficulties. One important aspect of the development of ADRs by polypharma is the inability of some patients, especially the elderly, to keep track of their drug use regardless of how well these drugs may work if given alone. There is also a cascade of prescription effects from polypharmacies in which certain medicines are used to treat the side effects of other medicines. This is the potential for unlimited medications used by the patient. It is possible to ignore the signs and signs of polypharmacies by confusing them with signs of aging or the disease itself. But this happens with many medicines to the sick. Constipation, diarrhea, fatigue, lameness, skin slip, catastrophe, anxiety and many other symptoms can be caused by both disease and polypharmacy.

2) DRUG DOSE AND FREQUENCY:
Drug dosing influences the development of ADR in various ways; for example. some medicines should be given in the morning, others in the evening, and others at bedtime. Taking bisphosphonates in bed may temporarily lead to the esophagus, the anti-platelet effect of aspirin in the evening may be preferable in the morning. Dosage should be considered as a factor that may have an effect on the development of ADR.
### D) DISEASE RELATED FACTORS:
Concomitant disease may also affect the patient's susceptibility to ADRs. For example; the increasing frequency of idiosyncratic poisoning with anti-infective drugs such as trimethoprim-sulfamethoxazole. Many diseases make patients more vulnerable to ADRs due to the presence of many diseases and the use of advanced medicines. If hypertension is accompanied by other diseases, these diseases can affect the body’s response to antihypertensive drugs, as it will worsen the body's metabolic processes. In patients with renal impairment, the effect of these drugs is reduced due to loss of local renal activity. This leads to an increase in dosage, which in turn induces further ADRs. The same problem occurs in patients with peptic ulcer, many drugs, which are prescribed NSAIDs, can lead to serious drug problems. Multiple disease factors are important for drug-disease interactions and ADRs. Drugs are beneficial in one disease and harmful in another. For example, some beta blockers for heart disease or high blood pressure can make asthma worse and make it harder for people with diabetes to report when their blood sugar is too low.

### ETIOLOGY OF ADVERSE DRUG REACTION:
Most medications-related adverse reactions are dosage-related; others allergic or idiosyncratic. Dose-related ADRs are generally predictable; Non-dose-related ADRs are generally unpredictable. Dosage-related ADRs are of particular concern when drugs have a limited therapeutic index (e.g., bleeding due to an oral anticoagulant). Adverse effects may result from decreased drug clearance in patients with renal or hepatic impairment, or from drug interactions.

Allergic ADRs are not dose-related and require prior exposure. An allergy develops when a drug acts as an antigen or allergen. Once the patient is experienced, subsequent exposure to the drug produces one of several types of allergic reaction. A proper medical history and skin care exams can sometimes help predict allergic side effects.

Idiosyncratic ADRs are unexpected side effects that are not dose-related or allergic side effects. They occur in a small percentage of patients taking the medicine. Idiosyncrasy is a definite term that is similarly defined as an abnormal response to a drug, but not all idiosyncratic reactions have a pharmacogenetic cause. A deprecated name can be created for device-specific ADRs.

### CLASSIFICATION OF ADRs:
Drug induced ADRs are classified on the basis of:

1. Pharmacological mechanism, and
2. Their preventability

1) Pharmacological mechanism:

Traditionally, ADRs have been classified into two types:

Type A reactions - sometimes called augmented reactions - are "dose dependent" and can be predicted based on the pharmacology of the medicine.

Type B reactions - strange reactions - which are unpredictable and unpredictable based on pharmacology.

Although widely mentioned, this initial classification does not work for all ADRs, such as chronic side effects associated with cumulative drug exposure (such as osteoporosis with long-term corticosteroid therapy) or withdrawal reactions (e.g., blood reflected by stopping the concentrated antihypertensive effect). An alternative and perhaps more comprehensive classification scheme is DoTS, which classifies reactions based on drug dose, response period, and related predisposing factors (such as other genetic, pathological, and biological differences). Benefits that help to consider the diagnosis and prevention of ADR in practice.

Predictable ADR can be classified on the basis of:

- Over doses,
- Adverse effect
- Drug-drug interaction, and
- Drug disease interaction.

Unpredictable ADRs may be happen due to:

- Intolerance,
- Idiosyncracy,
- Pseudo-allergy, and
- Allergy.
PREVENTION OF SIDE EFFECTS OF THE DRUGS:
While some side effects are unpredictable, such as anaphylaxis after exposure to penicillin-containing antibiotics, many can be avoided with appropriate predictability and control. Prevention (or contraception) usually refers to cases where the drug is inconsistent with current practice based on facts or is unrealistic due to known circumstances. It can be prevented that it is much easier to diagnose later. However, measures to reduce the risk of developing ADHD can be a way to reduce the risk of injury to the patient.

There are two main steps you can take to avoid ADR;

1. Determine the proportion of patients who are most likely to have side effects, and adjust treatment options accordingly.
2. Make sure your treatment plan minimizes possible side effects.

SUSCEPTIBILITY IDENTIFY:
Understanding patient susceptibility can provide a basis for defining prescribing and reduce the risk of ADR. The patient's history identifies previous side effects and, as a whole, prevents any new drug exposure. In other cases, susceptibility factors such as age, gender, stage of shipment and ethnicity may indicate a risk of using AMR. For example, the National Institutes of Health Mitigation and Excellence Guideline suggests that patients of African or Caribbean descent should take an angiotensin-II receptor blocker in the presence of an angiotensin enzyme (ACE) inhibitor for hypertension caused by hypotension. Inhibitor-induced angioedema. Pharmacogenetics is often used to design more personalized drug options that are more appropriate for the use of specific ADRs.

DIAGNOSIS:
Recognition of undesired delivery reactions; ADRs are just great healthcare mimics, often mimicking "traditional diseases" and manifesting their use in every body system. Drug use-related problems in patients with severe renal impairment encompass a variety of modalities, including biochemical and blood abnormalities (including acute renal, gastrointestinal, or health-related disorders). Persistent eruptions, drug-induced eosinophilia, or juvenile edema will certainly experience a history that serves to link the symptom or late detection to ADRs, and to avoid future ADRs that may help attribute the cause to specific drugs.

As soon as a drug is taken, these are easily referred to drug use. However, the diagnosis of chronic symptoms due to the use of drugs requires a significant degree of uncertainty and is often complicated. A suppressing drug is sometimes necessary, but an essential drug is difficult if it is not suitable. When testing the relationship between drugs and symptoms, a challenge should be considered, in addition to cases of severe allergic reactions. Therefore two actions should be performed in the diagnosis:

1. Consideration of rechallenge.
2. Reporting of suspected ADRs to MedWatch.

Doctors must report suspicious drug reactions to MedWatch (ADR Food and Drug Administration Monitoring Program), which is an early warning system. Unexpected reports such as ADRs can only be identified and investigated. MedWatch also monitors the nature and frequency of ADRs changes. Forms for information about ADRs reporting are available on the Medical Report Desk and the FDA's Drug News Daily Bulletin. Nurses, pharmacists and other health care professionals should also report ADRs.

TREATMENTS:
A careful and safe prescription is necessary to reduce the errors that can contribute to ADRs. Treatment plans must be treated and mitigated for all adverse effects. For example, co-prescription ofolic acid with methotrexate will reduce the incidence of adverse effects Combined with folate deficiency; and monitoring electrolyte and renal function in the treatment of renal active drugs or diuretics. All these examples can prevent adverse effects of treatment, although they may be limited, recommendations that appear unsatisfactory, often inadequate or ambiguous. It is important to remember that an accurate prescription may also completely prevent the use of drugs and treatment choices without pharmacology or preservatives should always be considered.

Dosage-related adverse reactions, dosage modifications or elimination or reduction factors are generally effective; however, an increase in the excretion of drugs is rarely required. Generally medications should be removed for allergic and idiomatic side effects. Because you don't have what you can pay. In the case of allergic side effects, and sometimes in the case of dose-dependent side effects, it is usually unsuccessful to prescribe other medications.

PREVENTIBILITY FOR ADRS:
Knowledge of drugs and their potential mechanisms is important to prevent drug side effects. To investigate potential drug interactions, data analysis can be used when a drug is modified or added to a test. For traditional medicine, the initial dose is carefully selected. Adverse reactions should always be taken before starting symptomatic treatment if the patient has no specific symptoms.
PHARMACOVIGILANCE:
Pharmacovigilance is defined as ‘the science and activities relating to the detection, assessment, understanding and prevention of adverse events or any other drug-related problem’.  
New legislation was introduced in the European Union in 2012 to ensure good vigilance practice for pharmaceutical companies and the medicines regulators. This new guidance clearly identifies the roles and responsibilities of relevant stakeholders in terms of drug safety. Notably, the guidance has introduced a programme of more intensive surveillance for new pharmacological agents and biological agents with black triangle status (i.e. those requiring additional monitoring). One of the guiding principles is that the pro-active strategies of the risk management policy replace the previous reactive strategies.

REPORT SIDE EFFECTS OF ADRs:
Potential ADR exposure has been based on recall systems for over half a century, such as the UK Yellow Card, managed by the Medicines Agency (MHRA) and the Human Health Commission (CHM). This project was created in 1964, shortly after the Thalidomide disaster in the 1950s. Through a voluntary report, the system collects information about suspected adverse reactions to all authors, including licensed drugs and vaccines, including more than counter-prescriptions and drugs that are available. In order to receive an effective report, only four parts of the information you desire: patient identification, response, suspected narcotics identification, and report identification. He urged the notary to provide as much information as possible, i.e. The UK Council reports that it still receives about 25,000 a year, allowing health officials to view the ADR results. Unfortunately, cancellation remains an important problem, with about 5% of all ADRs used. This limits the system’s ability to supply accurate flash memory information. In 2014, the UK NHS and MHRA published a joint recommendations: To improve reporting and search for therapeutic errors. In this regard, drug-related adverse events will be referred to the Yellow Card system under the National Report and Training System (NRLS).

Patients are increasingly involved in their therapeutic management, and as major patient evaluation procedures reveal the effectiveness of the Yellow Card, all patients are now actively encouraged to report adverse reactions. Paper (original yellow paper) has been largely replaced by the use of Yellow Card reporting systems or applications. Treatment email addresses commonly used in some hospitals and clinics can also include integrated messages that send ADR data directly to a central agency before it can be entered into national and international databases for processing.

Although outside the scope of this article, the modern generation can detect signal potential for early signs of harm and alert clinicians to potential new therapeutic risks. For detecting such standards, statistical algorithm extractions are run on the basis of a great service, but generally require additional estimations before the operation is completed. The potential for undesirable and possible adverse effects in databases, such as the Clinical Practice Research Datalink (CPRD), a database of anonymised UK primary care boards, can support or disprove potential symptoms.

Pharmacovigilance uses many other methods data systems, including official pharmaceutical surveillance studies, published data, pharmaceutical company data on periodic safety update reports (PSURs), and internationally distributed data. However, regulators and researchers are also looking at the potential of other “big data” sources, such as social media, to detect early signs. This is still an interesting, mostly unexplored field of research.

CAUSES OF ADR REPORTED:
The causes of side effects vary from study to study. However, most patients have adverse effects due to old age, poor patient readiness, and discomfort. Many studies have cited drug-related causes, such as misdiagnosis, conflicting counselling, incomplete physical examinations, misconceptions, and lack of vigilance.

Three studies also reported drug-specific factors such as drug administration, recurrence errors, drug effects, and similar drugs. Studies by Shehab and others have reported allergic reactions to adverse effects. The reasons for the opinion were also reported in a study. The two searches reportedly had no adverse effects.

DETECTION AND REPORTING:
The health care professional should be aware of ADR reports that ADR reports are, for the most part, just associations that suspect the drug has caused some adverse event. The ADR report does not imply an accidental link between an adverse drug reaction. But in case of doubt, it is better to report than not to report.

WHAT TO REPORT:
When there is suspicion of undesirable adverse events associated with the use of drugs, biologicals, herbal medicines, cosmetics or medical devices, it must be reported.

The reports should include the following information;

1. All ADRs are as a result of prescription and non-prescription.
2. Whether there is any increase in frequency of a given reaction.
3. Whether a serious reaction was expected or not.
4. ADRs occurring from overdose or medication errors.
5. Unusual lack of efficacy or when suspected pharmaceutical defects are observed.
6. Whether or not the drug was used in accordance with the drug information provided by the company marketing the drug.
7. ADRs in special fields of interest such as drug abuse and drug use in pregnancy and during lactation.
The minimal information to be provided for proper assessment of the ADR case reports is given below:

1. Patient information.
2. Description of adverse reactions.
3. Information related to the suspected drugs.
4. Information about how the adverse reactions had been managed.
5. Information about the reporter.

Reports are advised to study carefully and adhere to the guidelines on how to complete an ADR Reporting from for proper interpretation.

A) PATIENT INFORMATION:

1. Patient identity: The initials or number of the patient in a hospital, name of the medical institution, dispensary, clinic or pharmacy are to be indicated or recorded in the report.
2. Birth dates or age: The date, month and year of birth of each patient are to be recorded.
3. Sex: Male or Female 4. Weight: Should be in kilograms

B) ADVERSE REACTION:

1. Brief description of the ADR(s) indicates that the adverse reaction(s) is reported by marking X in the appropriate box. As clearly as possible the nature of the adverse reaction being reported including the site of the body and severity are to be mentioned briefly.
2. Time/date of onset of the adverse reactions: The time of onset or the occurrence of the adverse reaction in relation to the administration of the drug need to be recorded. The date of onset should be indicated in the following order: day, month and year. For example, the drug reactions appear immediately after drug administration or there was temporal or spatial correlation with administration.
3. Other related information:
   a) Patient's medical history or laboratory data including dates if available should be entered. If the ADR(s) being reported is considered relevant to the case.
   b) Appropriate laboratory tests are to be done on the patient and to confirm the adverse reaction the results are to be mentioned.
   c) This should be stated briefly but clearly.

C) SUSPECTED DRUG:

1. Name of the suspected drug(s): Trade name should preferably be used, if trade name is not available, generic name may be used. Strength of the drug(s) should also be stated.
2. Therapy date: The dates of starting and end of the administration of each drug should be started, and preferably recorded as follows: Date, Month and year. When the dates are not available, the duration of treatment is to be recorded. If drug administration has not been terminated at the time of reporting, it is to be recorded as ‘continuing.’
3. Batch no and expiry date: If available are to be recorded.
4. Reason for use: The indication or condition (reason)
5. The relevant information on medical devices are to be provided.
6. Dosage, frequency and route of administration should be clearly mentioned. For example:
   a) Dosage: The dosage form such as tablet, capsules, syrup, injection, cream, eye drops, etc. including total amount of drugs are to be specified.
   b) Frequency: Unit of dose i.e. mg, ml, mg/kg or per given or taken such as twice, thrice or 4 times daily are to be specified.
   c) Route of administration by which the drug was administered: This should be entered in full term or abbreviated using the World Health Organization (WHO) codes as follows:
Osteoporosis is a metabolic bone disease characterized by low bone density and deterioration of bone architecture that increase the risk of fractures. Osteoporosis-related fractures can increase pain, disability, nursing home placement, total health care costs, and mortality. The diagnosis of osteoporosis is primarily determined by measuring bone mineral density (BMD) using noninvasive dual-energy x-ray absorptiometry. Osteoporosis medications include bisphosphonates, receptor activator of nuclear factor kappa-B ligand inhibitors, estrogen agonists/antagonists, parathyroid hormone analogues, and calcitonin. Emerging therapies utilizing novel mechanisms include a cathepsin K inhibitor and a monoclonal antibody against sclerostin.

While professional organizations have compiled recommendations for the management of osteoporosis in various populations, a consensus has yet to develop as to which is the g