ANAND M INGALE, PRATIBHA NADIG*, ANANYA CHAKRABORTY  
Department of Pharmacology, Vydehi Institute of Medical Sciences and Research Centre, Bengaluru - 560 066, Karnataka, India. 
Email: drpratibhanadig@yahoo.co.in  
Received: 23 March 2018, Revised and Accepted: 08 May 2018

ABSTRACT

Objectives: The objectives of the study were to analyze the various adverse drug reactions (ADRs) collected in the Pharmacovigilance Unit of Vydehi Institute of Medical Sciences and Research Centre with respect to their causality, severity, and preventability and also to identify the various risk factors, concomitant medications, and comorbid conditions with the occurrence of these events.

Methods: A prospective, non-interventional, observational, and cross-sectional study was carried out in the various clinical departments of Vydehi Institute of Medical Sciences and Research Centre from June 2014 to May 2015. The Classes of drugs, Organ system involved, Comorbid conditions associated and Concomitant drugs involved in causing ADRs were looked into. The assessment for causality and severity was determined by Naranjo and Modified Hartwig and Siegel scales, respectively. The data were compiled and subjected to descriptive statistical analysis.

Results: A total of 433 patients developing ADR reports were analyzed during our study period. Of these, 53.59% were females. 75% of them were of adult age group. Antimicrobials and chemotherapy group showed the maximum ADRs. The skin and appendages (27.6%) were the most affected organ system followed by the gastrointestinal system (22.8%). Comorbid conditions were found in 76 (20.1%) reports, of which diabetes (28.9%) and hypertension (26.3%) were maximum. 74 were serious reports. Maximum reports were probable and of mild severity.

Conclusion: Through active surveillance of the ADRs helps in early detection and prevention of all the possible adverse events associated with the usage of drugs and thereby provides a better health-care treatment to the patients.

Keywords: Adverse drug reaction, Causality, Serious reaction, Naranjo, Hartwig, Pharmacovigilance.

INTRODUCTION

Adverse drug reactions (ADRs) form an important compounding factor in the management of any clinical conditions, constituting an important cause for the morbidity and mortality [1]. ADRs are the fourth leading cause of death [2]. They occur in 10–20% of hospitalized patients [3]. The incidence of ADRs varies from 0.15% to 30% [4]. Serious ADRs account for 6–7% of all hospital admissions [2]. A study in South India showed that ADRs accounted for 0.7% of total admissions, and 1.8% of ADRs resulted in death [5].

As most of these ADRs are preventable, a thorough knowledge about them helps in analyzing the pattern and severity of them in various clinical conditions, which, in turn, helps in reduction of the health-care cost [6].

Spontaneous reporting, a part of active surveillance, though theoretically the best method of ADR assessment, has not been an effective method due to a large percentage of under-reporting. Reporting of ADRs is only 3% of the global ADR occurrence [7]. Passive or stimulated reporting is by and large the most common method of ADR reporting. It is recommended by Pharmacovigilance Program of India. However, active surveillance through direct interaction with patients has been reported to be the most effective means to assess the ADRs prevailing in the society [8]. Our previous experience of active surveillance in a single department has clearly shown that active surveillance also improves spontaneous reporting [5].

Our review of literature has shown numerous studies with active surveillance. However, very few studies have been carried out to assess the association of factors such as comorbid conditions and concomitant medications with the occurrence of the ADRs, which may be playing a key role in their occurrences. The knowledge of the same will help in making the prescriber aware of the various possible adverse events associated with the use of drugs.

Hence, the present study was undertaken in various clinical departments of Vydehi Institute of Medical Sciences and Research Centre (VIMS and RC) through active surveillance to analyze the different ADRs occurring in them.

Research objectives

The objectives are as follows:
1. To analyze the various ADRs collected in the Pharmacovigilance Unit of Vydehi Institute of Medical Sciences and Research Centre with respect to their causality, severity, and preventability.
2. To identify the various risk factors, concomitant medications and comorbid conditions with the occurrence of these events.

METHODS

A prospective, non-interventional, observational, and cross-sectional study for a period of 1 year from June 2014 to May 2015 was conducted at the various clinical departments of VIMS and RC after obtaining the approval of the Institutional Ethics Committee (VIEC/2016/APP/028).

Regular visits to the outpatient department and the inpatients of various clinical departments were carried out. Standard Central Drugs Standard Control Organization form [9] was used for reporting and analyzing the ADRs.

Each patient was interrogated only after obtaining their verbal consent. A detailed survey of the patient's complaints, treatment history and the
various events that occurred after administering the drugs was done and documented. Any new and unusual events occurring after the drug administration were also interrogated. The collected reports were subjected to analysis and later submitted to our Pharmacovigilance Unit.

The activity was divided among two groups to reduce any chances of bias. One group was involved in collecting the data and the other in analyzing the collected data.

The analysis of data was carried out as follows

Validity of the reports
ADR report with an identifiable patient, identifiable drug, indication for the use of the drug, and proper dates were considered valid and was included in our study [10].

Demography
The age of the patient was recorded and was grouped accordingly as pediatric, adult, or geriatric. The sex of the patient was noted.

Drug Group and its classes
Drug group along with its class involved in causing the reaction was recorded and categorized [11].

Organ system involvement
The organ system being affected in the reaction was recorded and categorized accordingly [12].

Number of drugs in therapy
Each of the reaction was looked in for the number of drugs prescribed for therapy and was categorized as reaction with monotherapy in case of reactions involving only one drug prescription and reactions with more than one drug prescribed were categorized as polytherapy reactions.

Comorbid conditions
Details of the various comorbid conditions (diabetes mellitus, hypertension, thyroid diseases, and so on) present in the patient likely to influence the occurrence of ADRs were surveyed and tabulated.

Measures undertaken and the outcome measures
Measures undertaken and outcome measures seen in the ADR were looked into and tabulated [13].

Seriousness of the reactions
Seriousness of the reactions was looked into and the reactions were classified as serious or non-serious as per the World Health Organization guidelines [14]. In addition to this, the reasons for the seriousness of the reactions were also surveyed and tabulated.

Expectedness of the reaction
The expectedness of the reaction was analyzed by doing a literature search on the possible ADRs due to the causative drug and also from the summary of the product characteristics. The outcome of the reaction was cross-checked with all the possible outcomes that could occur and it was analyzed as to whether the outcome occurring in the reaction is an expected reaction or not [15].

Risk factors associated with ADRs
The cases were thoroughly looked into the possible risk factors such as smoking, alcohol consumption, history of allergy, age, and concomitant medications.

Causality assessment
Causality assessment was done by Naranjo et al. [16] scale and the reactions were categorized as probable, certain, possible, or unlikely depending on the scores obtained.

Severity assessment
Severity assessment was done by Modified Hartwig et al. [17] scale and the reactions were categorized as probable, certain, possible, or unlikely based on their scores.

Statistical analysis
The data were analyzed through descriptive statistics, and the values were expressed in numbers and percentages.

RESULTS
Validity of the reports
We encountered 433 patients developing ADRs. Out of these, 377 (85.15%) were valid reports and were included in our study.

Demography
Of the 377 valid reports, 175 (46.41%) were of males and 202 (53.59%) were of females. The age group of the patients is depicted in Fig. 1. The adult group possessed the maximum ADRs.

Drug groups and classes
Antimicrobial and chemotherapy group of drugs accounted for the maximum ADRs as depicted in Fig. 2.

Organ system involvement
The most affected organ system was the skin and appendages (27.6%) as depicted in Fig. 3.

Number of drugs in therapy
The monotherapy and polytherapy reports were 188 and 189, respectively, in our study.

Comorbid conditions associated with ADRs
On analyzing the comorbid conditions associated with the ADRs, we found that they were present in 76 (20.1%) reports, of which diabetes (28.9%) and hypertension (26.3%) were the most common ones as depicted in Table 1.

Measures undertaken and the outcome measures
The most common measure undertaken was stoppage of the drug (n=276; 73.2%) whereas drug was continued in 71 (18.8%) patients and 70 (18.5%) cases, it was unknown.

With respect to the outcome measures, maximum were recovered (n=212; 56.2%) whereas drug was continued in 71 (18.8%) patients and 70 (18.5%) cases, it was unknown.

Seriousness
We encountered 74 (19.62%) serious reports. The most common reason for seriousness was hospitalization-initial or prolonged as depicted in Fig. 4.

Expectedness
The literature search for all the possible ADRs possible with each drug was done and analyzed as to whether the particular ADR manifested is expected or not. We found that 290 (76.92%) reports were expected outcomes whereas 97 (25.72%) were unexpected.

![Fig. 1: Age groups of patients involved in adverse drug reactions](image-url)
Risk factors associated with ADRs
We encountered 83 (22.01%) cases having associated risk factors. Of these, 30 (36.1%) were alcoholic and 20 (24%) were smokers as depicted in Table 2.

Causality
The causality assessment showed that majority of the cases was probable as depicted in Table 3.

Severity
The severity of the reactions was classified as mild, moderate, and severe as per modified Hartwig et al. scale as depicted in Table 4.

DISCUSSION
Females had a higher incidence of ADRs, which are in accordance with many proven previous studies [18], wherein female sex was considered to be a risk factor for the development of ADRs. Women in comparison to men having lower bodyweight and organ size, more body fat, different gastric motility, and lower glomerular filtration rate are attributed by Alomar [19] for having higher ADRs.

Adult group showed a higher percentage of ADRs. This was similar to Venkatesan et al. [20] who also found maximum ADRs among the adult age group. The mean age of the patients developing ADRs was 45.95+17.93 in their study. However, studies on spontaneous reporting have almost always shown higher incidences in the elderly group [21].

Table 1: Comorbid conditions associated with ADRs

<table>
<thead>
<tr>
<th>Comorbid conditions</th>
<th>Number of ADRs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diabetes</td>
<td>22</td>
</tr>
<tr>
<td>Hypertension</td>
<td>20</td>
</tr>
<tr>
<td>Dysthymia</td>
<td>5</td>
</tr>
<tr>
<td>Lymphadenitis</td>
<td>4</td>
</tr>
<tr>
<td>Pregnancy</td>
<td>3</td>
</tr>
<tr>
<td>CRF</td>
<td>2</td>
</tr>
<tr>
<td>S. aureus positive infection</td>
<td>2</td>
</tr>
<tr>
<td>Anemia</td>
<td>2</td>
</tr>
<tr>
<td>Cushing’s syndrome</td>
<td>2</td>
</tr>
<tr>
<td>Gastritis</td>
<td>1</td>
</tr>
<tr>
<td>Constipation</td>
<td>1</td>
</tr>
<tr>
<td>Vestibular lesions</td>
<td>1</td>
</tr>
<tr>
<td>Dengue</td>
<td>1</td>
</tr>
<tr>
<td>Hypothyroidism</td>
<td>1</td>
</tr>
<tr>
<td>Urinary tract infection</td>
<td>1</td>
</tr>
<tr>
<td>Pain abdomen</td>
<td>1</td>
</tr>
<tr>
<td>Cholelithiasis</td>
<td>1</td>
</tr>
<tr>
<td>Fatty liver</td>
<td>1</td>
</tr>
<tr>
<td>Adult polycystic kidney and liver disease</td>
<td>1</td>
</tr>
<tr>
<td>Diabetic nephropathy</td>
<td>1</td>
</tr>
<tr>
<td>Neurocysticercosis</td>
<td>1</td>
</tr>
<tr>
<td>Osteosarcoma of the femur with</td>
<td>1</td>
</tr>
<tr>
<td>lung metastasis</td>
<td>1</td>
</tr>
<tr>
<td>Pleural effusion</td>
<td>1</td>
</tr>
</tbody>
</table>

*S. aureus: Staphylococcus aureus. ADRs: Adverse drug reactions*
It appears that in voluntary reporting system the ADRs of the adult group may undergo unnoticed and hence unreported.

Antimicrobials (n=90) and anti-cancer group of drugs (n=75) accounted for the higher incidence of ADRs, which are in accordance with the study done by Shah and Sattigeri [22].

The most commonly involved organ was the skin and its appendages, which is in accordance to that reported Bhabhor et al. [23] who reported 52.25% of skin and appendages disorders in their study. Similar results were observed by Siddiqui et al. [24]. Since these reactions are easily detectable, they seem to be prominent in any reports on ADRs.

Although polytherapy is supposed to be associated with the increased occurrence of ADRs due to drug interactions, our study had an almost similar number of ADRs with monotherapy as well as with polytherapy.

We encountered 76 reports possessing comorbid conditions. Diabetes and hypertension were the most common comorbid conditions associated. This is found to be associated with higher incidences of ADRs as there is a possibility of drug-drug interactions [25]. It is also observed that adequate control of these conditions will reduce the ADRs of other drugs.

As far as the remedial measures undertaken were concerned, the most common was to stop the offending drug (n=276) and the most common outcome seen was recovery of the patient (n=212). Stopping the offending drug usually helped in the recovery of the patient making the diagnosis of ADRs easier.

The causality assessment of suspected drugs to reactions shown more than half of reactions belonged to probable category, and only a few percentage reactions belonged to certain category, which is in accordance to Kumari et al. [26] and contrast to the previous studies on active surveillance showing maximum of possible causality [5].

With respect to the severity assessment, most of the reactions encountered were of mild grade (n=224) followed by moderate (n=123) and severe (n=30) grades.

Our study had 56 (14.85%) invalid reports. Validity of the reports could have been increased by collecting and documenting all the details relating to the ADRs. A more complete ADR form would have given higher validity rates.

Through our study, we encountered 97 unexpected reactions. This was possible mainly because of adopting active surveillance as the investigators were aware of the expected reaction. Our study, hence, shows that active surveillance has a better effect on the quality of reporting. Thoroug h and accurate knowledge of the drug profile on the part of the reporter helps in predicting the various possible ADRs with the usage of the drugs. The detection of the unexpected ADRs will aid in exploring the undetected ADRs that are all possible with the usage of the drugs and add to the existing knowledge about the drugs.

Limitations of our study
The results of our study are based only the data procured through a smaller population, which, need not depict the results of the general population. Studies on a larger population can reveal a better result.

CONCLUSION
Spontaneous reporting is the major modality of ADR reporting in our country. However, active surveillance seems to be a better modality to identify the risk factors and detects more unexpected reactions. More awareness about the usefulness and importance about ADR reporting will help in increasing the reporting rate among the physicians and also thus help in better patient management.

CONFLICTS OF INTEREST
The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

AUTHOR’S CONTRIBUTION
All the authors have equally contributed in the idea of the research; planning and execution of the study protocol; collection, compiling and analysis of the data, and preparing the manuscript.

REFERENCES


