INTRODUCTION

Drugs are capable of causing immense comfort as well as irreparable harm to man. Harms caused by medicines are often described as adverse drug events and these can be due to either medication errors or adverse drug reactions [1]. While medication error is a broad term and points at an error which is in control of the health care professional or the patient, the World Health Organization (WHO) defines an adverse drug reaction (ADR) as "a reaction which is noxious and unintended, and which occurs in doses normally used in human for prophylaxis, diagnosis, or therapy of disease, or for the modification of physiological functions [2]."

The Government of India established National Pharmacovigilance Program in November 2004, after which the Pharmacovigilance program of India (PvPI) was launched in 2010. This programme aims at establishing the habit of ADR notification by health care workers all over India.

The purpose of ADR monitoring is to help ensure that patients are entitled to safe and effective medicines and also to generate ADR data which is crucial for understanding the purpose of pharmacovigilance and for educating the health care providers about this discipline.

The criteria for serious adverse drug reactions (serious ADRs) have been specified by the WHO and includes any untoward medical occurrence at any dose that results in death, life-threatening, requires or prolongs hospitalization, or results in persistent or significant disability or incapacity [3].

The overall incidence of serious ADRs in the US between 1966 to 1996 was 6.7% and a number of fatal medical errors were 0.32% of hospitalized patients. Serious ADRs occurred in 10–20% of hospitalized patients making these reactions between the fourth and sixth leading cause of death [4].

In this study we attempt to analyze the serious ADRs in a tertiary care centre and assess their seriousness, outcome, causality and severity.

In a resource limited country like India, the concept of ADR monitoring and the practice of ADR reporting is yet to catch up. Our study emphasizes on increasing numbers of serious ADRs and the need to improve models for patient care benefit in intensive care units by training pharmacovigilance cell members to collaborate with hospital consultants and help reduce mortality due to serious ADRs.

MATERIALS AND METHODS

A prospective observational study was carried out at ADR monitoring centre of Bhaskar General Hospital from July 2016 to Dec 2016 after obtaining Permission from Institutional Ethical Committee of the hospital.

64 suspected ADRs were collected using the ADR notification form (yellow form) and CDSCO form (red form). These were analysed for seriousness using Hardwick Siegal scale and causality assessment was done using WHO-UMC scale.17 out of 64 ADRs were reported to be serious.

ADR s were diagnosed by the treating consultants, and relevant details of each ADR were collected in spontaneous ADR reporting form(yellow form) The details were then fed into the CDSCO form(red form) and sent to the National Coordinating Centre via Vigiflow software.

ADR s were reported from Departments of Dermatology, Gynaecology, Haematology, Medicine, Ophthalmology, Pediatrics, Psychiatry, and Pulmonology.

The collected information included patient’s initial, age, gender, reporting department of the hospital, description of the reaction,
duration of the reaction, the name of the suspected drug causing the reaction, concomitant drug if any, and outcomes.

Serious ADRs were identified as per the WHO-UMC criteria and analyzed to find the time relationship with the initiation of drug treatment, causal drug group, and body system as per system organ class (SOC). Causality assessment was done using the WHO-UMC scale [5]. While Severity was assessed using modified Hartwig and Siegel scale [6].

The outcome of the patients with ADR was recorded as fatal, fully recovered (patient fully recovered during the study period), recovering (patient recovering, but not fully recovered during the study period) and unknown (insufficient information and not documented).

Type of study
Prospective observational study

Study design
The data will be collected from the prescriptions of patients suffering with ADRs by the respective consultants in the ADR reporting form and will include Patient particulars, history, diagnosis, drugs-the dosage, frequency, and duration of treatment, comorbid conditions, suspected drug that caused ADR, description of the ADRs, details on hospital stay, concomitant drugs, generic or brand prescription information. The CDSCO form is then duly filled at the ADR monitoring centre and with access to the yellow form, the data is will be analysed, tabulated, and statistical analysis will be carried out.

Selection criteria

Inclusion criteria
All the suspected ADRs that may be due to the medications, both prescribed and over the counter, taken by patients either as inpatients or outpatients, that were ultimately documented.

Exclusion criteria
The use of the alternative system of medicines such as Ayurveda, Homeopathy, Unani, etc. was excluded.

All mentally retarded, drug addicted, and unconscious patients were also excluded from the study.

Patients admitted due to alcohol or drug abuse, a suicide attempt or incoherent patients with unreliable data were excluded.

Statistical analysis
Data entry and analysis was done using Microsoft Excel 2010 version. Data was presented in percentages and proportions. Bars diagrams and pie charts were used to depict percentages.

RESULTS
64 patients were reported to experience ADR at BGH during the study period. Of these, 17 (26.5%) were serious and 47 (73.4%) were non-serious as per the Hartwig Seigal scale

Fig. 2 shows the distribution of ADRs in different age groups, both male and female. The mean age of patients with serious ADRs was 49 y. There were no severe ADRs in the pediatric age group.

Fig. 2: The severe ADRs, 9 patients (52.9%) were females and 8(47%) were males with a male: female ratio of 1:0.88

Most of the severe reactions were among the age group 40-60 y and it was observed that Polypharmacy is ubiquitous for the development of ADRs.

Fig. 3 shows the percentage of cases reported to be possible, probable and certain. The causality assessment revealed that most of the ADRs belonged to "probable" category (14, 82.3%) followed by "certain" category (2,11.7%).

Fig. 3: Causality Assessment

Fig. 4: Shows the various ADRs reported system wise
Cutaneous manifestations which included rash, urticaria, dermatitis, Steven Johnson syndrome, Toxic epidermal necrolysis etc were most common ADRs with an incidence of 58.8% and these were most commonly due to antiepileptic agents like Phenytoin and Carbamazepine.

The next organ system commonly involved was a gastrointestinal system which included gastritis, nausea, vomiting and dyspepsia etc accounting for 17.6% of total ADRs. 

23.53% of the severe ADRs were due to DOTS-ATT specifically Rifampicin, 23% due to analgesics with paracetamol one of the active ingredients and 17.65% were due to anti epileptic agents like Phenytoin and CBZ.

Table 1: Clinical pattern of serious ADRs

<table>
<thead>
<tr>
<th>Clinical presentation</th>
<th>Suspected drug</th>
<th>Number of severe ADRs (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Steven Johnsons Syndrome</td>
<td>Carbamazepine</td>
<td>1(5.8)</td>
</tr>
<tr>
<td></td>
<td>Phenytoin</td>
<td>1(5.8)</td>
</tr>
<tr>
<td>Severe Urticaria</td>
<td>Phenytoin</td>
<td>1(5.8)</td>
</tr>
<tr>
<td></td>
<td>DOTS</td>
<td>2(11.7)</td>
</tr>
<tr>
<td></td>
<td>Paracetamol</td>
<td>1(5.8)</td>
</tr>
<tr>
<td></td>
<td>Amoxicillin+Clavulanic acid</td>
<td>1(11.4)</td>
</tr>
<tr>
<td></td>
<td>Ceftriaxone</td>
<td>1(5.8)</td>
</tr>
<tr>
<td></td>
<td>Olmesartan</td>
<td>1(5.8)</td>
</tr>
<tr>
<td>Gastritis, Nausea, Vomiting, Syncope</td>
<td>Rifampicin</td>
<td>2(11.7)</td>
</tr>
<tr>
<td></td>
<td>Paracetamol</td>
<td>2(11.7)</td>
</tr>
<tr>
<td></td>
<td>Acedofenac+Paracetamol</td>
<td>1(5.8)</td>
</tr>
<tr>
<td>Generalised Edema and shock</td>
<td>Azithromycin</td>
<td>1(5.8)</td>
</tr>
<tr>
<td>Fever with chills, falling BP progressing to Shock</td>
<td>Inj. Iron-Sucrose for anemia</td>
<td>1(5.8)</td>
</tr>
</tbody>
</table>

A maximum number of Severe ADRs were reported from the Dept of General Medicine, followed by Departments of Pulmonology and Skin. All cases were admitted in the ICU of BGH and treated until recovery.

DISCUSSION

According to a meta-analysis review of 39 prospective studies carried out in the US from 1966 to 1996, the overall incidence of serious ADRs was 6.7% and of fatal ADRs was 0.32% of hospitalized patients [4]. Similarly, it has been estimated that 5% of all hospital admissions in the EU are caused by ADRs, 5% of hospitalized patients will experience an ADR during their hospital stay, and that ADRs cause 197,000 deaths annually throughout the EU [7]. A study by Ramesh et al. in India carried out in a tertiary referral centre in India showed that admissions due to ADRs accounted for 0.7% of total admissions and deaths due to ADRs accounted for 18% of total ADRs [8].

In this study, spontaneously reported adverse drug reactions were evaluated for a period of 6 mo, with emphasis on serious ADRs. Of the total adverse drug reactions reported, 17 (26.5%) were serious. This shows that severe ADRs are a significant proportion of the ADR burden which is supported by a study conducted by Lukshmy M Hettihewa et al. [9] who reported in their study that severe adverse drug reactions were 31% of all ADRs reported.

It was observed that adverse drug reactions were more commonly reported in elderly (above 45 y) due to polypharmacy, as more drugs were being used in this age to address other comorbid conditions.

The male and female ratio was found to be 1:0.88, with the nearly equal incidence of males and females. This is in contrast with the study by Sarminder Kaur et al. [10], in which preponderance of ADRs was seen in female subjects as compared with males. 27.9% of females enrolled in their study suffered from ADR as compared with 15.8% of enrolled males.

Systems most commonly affected were dermatological in 58.9% of patients, gastrointestinal in 17.6% of patients. The results were comparable with an international study conducted by Suh et al. [11], which revealed that the system most badly affected were the dermatological and gastrointestinal systems.

The drug class mostly associated with ADR were anti tubercular antibiotics and NSAIDs, both accounting for 23% of cases each, followed by anti epileptic drugs in 17.6% of patients in the present study. Murphy and Frigo developed and implemented an ADR reporting program in Loyola University Medical Center, Chicago and their study revealed that the most common adverse reactions were rash; and antibiotics were the most commonly implicated drug class [12]. Our results were also comparable with other studies like one done by Glassen et according to whom NSAIDs are implicated in the majority of ADRs [13].

CONCLUSION

We conclude that Anticonvulsants, Analgesics and Antimicrobials are responsible for most of the ADRs in this study.

In view of the increasing percentage of severe ADRs being reported, it would be helpful if ADR monitoring centre members are given basic training in monitoring and treating severe ADRs.

ACKNOWLEDGEMENT

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CONFLICT OF INTERESTS

Declare none

REFERENCES


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