“Study And Assessment Of Drug-Drug Interactions In Hospitalised Patients In Quaternary Care Hospital”

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Abstract: Drug-Drug Interactions (DDI) in patients receiving multi-drug therapy are of wide spread. Such interactions are one of the important cause of adverse drug reactions (ADRs) and may lead to increased risk of mortality and morbidity. Hence, health cost will increase. This study estimates to find out the potential DDIs among the hospitalized patients. The final results of the study concluded with a major preponderance towards ATORVASTATIN and ASPIRIN in which the interaction of ATORVASTATIN and PANTOPRAZOLE can lead to (insert what will happen if HMG coa increases) and interaction between ASPIRIN and HEPARIN can lead to (insert what will happen if the interactions occur) further intervention by clinical pharmacists to reduce such adversities have to be implemented.

Keywords: drug-drug interaction, adverse drug reaction, patient medical record.

I. Introduction

An interaction is said to occur when the effects of one drug are changed by the presence of another drug, herbal medicine, and food drink or by some environmental chemical agent. Drug-drug interaction is a modification of the effect of a drug when administered with another drug. The effect may be an increase or decrease in the action of either substance, or it may be an adverse effect that is not normally associated with either drug. This has become an important issue in the health care.1

The majority of the patients are treated with more than one drug simultaneously. Reasons for the treatment with multiple drugs include the treatment of multiple ailments in the same patient and the use of multiple drugs for the same ailment. With the increasing median age of the population, and the now known effectiveness of multiple therapy regimens for viral diseases, cancer, cardiovascular diseases and infectious diseases, so exposure of a patient to multiple drugs is a common rather than occurrence.2

The outcome can be harmful if the interaction causes an increase in the toxicity of the drug. For example, there is a considerable increase in risk of severe muscle damage if patients on statins start taking azole antifungals. A reduction in efficacy due to an interaction can sometimes be just as harmful as an increase: patients taking warfarin who are given rifampicin need more warfarin to maintain adequate and protective anticoagulation. These unwanted and unsought-for interactions are adverse and undesirable but there are other interactions that can be beneficial and valuable, such as the deliberate co-prescription of antihypertensive drugs and diuretics in order to achieve antihypertensive effects possibly not obtainable with either drug alone. The mechanisms of both types of interaction, whether adverse or beneficial, are often very similar.1

The basic and clinical scientific issues underlying pharmacokinetic drug interactions are becoming increasingly complex as poly pharmacy becomes more common and more drugs with enzyme-inducing or inhibiting properties are introduced into clinical practice. A well-planned, integrated approach is needed to address the clinical problems. Ideally, the approach should incorporate the collaborative participation of individuals with expertise in molecular pharmacology, cytochrome biochemistry, in vitro metabolism, clinical pharmacokinetics-pharmacodynamics, and clinical therapeutics. The ultimate goal should be the informed and safe use of drug combinations in clinical practice. Drug interactions are said to occur when the pharmacological activity of a drug is altered by the concomitant use of another drug or by any pharmacologically active substance. The drug whose activity is affected is affected by such an interaction is called as object drug and the agent which precipitates such an interaction is referred to as the precipitant. The net effect of a drug interaction is Increased or decreased effect (quantitative) Rapid or slow effect (seldom qualitative) Precipitation of newer or increased adverse effects. Most interactions are undesirable. Rarely desirable (beneficial) example:- enhancement of activity of penicillin’s when administered with probenecid.3
II. Methodology

Data was retrieved from patient case file and Patient Medical Record (PMR) and stored in a data collection form for a period of 6 months. DDIs were identified using Lexi-Comp, database system. The drug-drug interactions were classified as major, moderate and minor. Inclusion criteria were only Hospitalized patients and Exclusion criteria were Prescription containing less than three drugs.

2.1 DATA COLLECTION FORM:
The data collection form was developed by referring available literatures. It includes patient demographics, laboratory results, final diagnosis and medication chart.

2.2 DOCUMENTATION:
The data collected from the patients was documented for further analysis. Microsoft excel software was used for statistical analysis.

III. Results

This was a prospective observational study conducted for 6 months. During the study total of 202 prescriptions were collected and analysed. Out of 202 prescriptions 131 were male (64.8%), 71 were females (35.1%).

Out of 202 prescriptions, 12 prescriptions were of age <20 years (6%), 35 were between 20-35 years (17%), 34 prescriptions were between 36-50 years (17%), 68 prescription were between 51-65 years (34%), 47 prescription were between 66-80 years (23%), 6 prescription were >80 years (3%).
Table 1: Prevalence of DDIs in study population

<table>
<thead>
<tr>
<th>Type of case</th>
<th>Number of prescriptions</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cases with DDIs</td>
<td>181</td>
<td>10.3</td>
</tr>
<tr>
<td>Cases with out DDIs</td>
<td>21</td>
<td>89.6</td>
</tr>
<tr>
<td>Total No. of cases</td>
<td>202</td>
<td>100</td>
</tr>
</tbody>
</table>

The identified DDIs were analysed and classified based on severity as major, moderate and minor. A total of 1238 DDIs were identified of which 583 combinations were obtained from 181 prescription. From these 314(25.36%) were of major, 821(66.31%) were of moderate and 103(8.31%) were of minor.

The drugs are classified according to the Lexi-Comp, Inc. Version: 2.7.5 Copyright 2016, Lexi-Comp, Inc. database where the drugs are classified as X(avoid combination), D(consider therapy modification), C(monitor therapy), B(no action needed), A(no known interaction). Out of 1238 DDIs, 53 were X(4.28%), 177 were D(14.29%), 826 were C(66.72%), 166 were B(13.40%), 16 were A(1.29%).

The most prevalent DDIs were identified. The most frequent interacting combinations were Atorvastatin+pantoprazole, clopidogrel+aspirin, clopidogrel+atorvastatin, aspirin+heparin, ondansetron+tramadol.
IV. Discussion

4.1. RISK FACTORS CONTRIBUTING TO DRUG INTERACTIONS:

Multiple drug therapy: This is very common in most acute and chronic care settings, for e.g., therapy in patient suffering from hypertension and congestive heart failure includes antihypertensives as well as digitals which together may lead to abnormal heart rhythms. Concurrent use of non-prescription drugs, for e.g. aspirin as well as herbal medications also lead to drug interactions. Theoretically, the possibility for drug interactions to occur is over 50% when a patient is receiving five medications, and the probability increases to 100% when seven drugs are used.

Multiple prescribers: some individuals go to more than one physician, and it is common for a patient to be treated by one more specialist in addition to a family doctor. E.g: one doctor may prescribe an anxiolytic for a patient while another prescribes an antihistamine having sedative properties with a possible consequence of an excessive depressant effect.

Multiple diseases: some patient take several drugs owing to their suffering from more than one disease e.g. a patient with both diabetes and hypertension takes hypoglycemics and beta blockers which result in decreased response to anti-diabetic drug resulting in elevated blood sugar level. Advanced age of patients: increased tendency of drug interactions in elderly due to decrease in liver function. Drug related factors: clinically significant interactions are most likely to occur between drugs that have potent effects and narrow therapeutic index. Poor patient compliance: multiple drugs, inadequate information by pharmacist or doctor, confusion regarding taking medicines which may lead to under dosing or overdosing and consequent drug interactions.

In this study total of 202 prescriptions were collected. According to gender classification there were 131(64.8%) males and 71(35.1%) females. Then according to age classification out of 202 prescriptions ,12 prescriptions were of age <20years(5.9%),35 prescriptions were between 20-35years(17.32%), 34 prescriptions were between 36-50years(16.38%),68 prescriptions were between 51-65years(33.6) prescriptions were between 66-80years(23.2%) and 6 prescriptions were >80years(2.9%). Based on severity out of 202 prescriptions, 181(10.3%) prescriptions were classified as DDIs and 21(9.6%) were without DDIs. From these total 181 DDIs, 314(25.36%) were major, 821(66.31%) were of moderate and 103(8.31%) were of minor. A total of 1238 DDIs were identified of which 583 different combinations were obtained from 181 prescriptions.

The drugs are classified according to the Lexi-Comp,IncVersion: 2.7.5 Copyright 2016 database where the drugs are classified as X(avoid combination), D(consider therapy modification), C(monitor therapy), B(no action needed), A(no known interaction). Out of 1238 DDIs, 53 were X(4.28%), 177 were D(14.29%), 826 were C(66.72%), 166 were B(13.40%), 16 were A(1.29%).

Classification based on number of drugs prescribed, out of 202 prescriptions 39 patients were prescribed with 3-6 drugs in which 71 DDIs were found (5.46%), 90 patients were prescribed with 7-10 drugs in which 431 DDIs were identified (33.17%), 41 patients were prescribed 11-14 drugs in which 363 DDIs were identified (27.94%), 26 patients were prescribed with 15-18 drugs in which 346 DDIs were identified (26.6%), 6 patients were prescribed with greater than 19 drugs in which 88 DDIs were identified (6.77%). The most frequent interacting combinations were Atorvastatin+ Pantoprazole, Clopidogrel+ Aspirin, Clopidogrel+ Atorvastatin , Aspirin+ Heparin , Ondansetron+ tramadol. Similar study was conducted by O Jimmy Devi, Blessy KG, Hephizibah MC, Vinay DM, Sushilkumar PL at tertiary care Teaching Hospital, Davangere. It also shows that the most prevalent DDI are with Aspirin+Clopidogrel.

V. Conclusion

The Drug-Drug Interactions are frequent among the hospitalized patients who were prescribed with more than three drugs. The drugs that are significantly found having major preponderance are “ATORVASTATIN+PANTOPRAZOLE and ASPIRIN+HEPARIN”. Prevalence of drug interactions increases by a linear mode according to numbers of drugs prescribed, number of therapeutic drug classes, patient’s gender and age. The results were obtained according to the Theoretical hypothesis obtained by us in hospitalised set up. So these interactions has been monitored by us to minimize the potential adverse drug reactions and intervention as appropriate.

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References


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