Role of the Clinical Pharmacist in Detection of Drug Therapy Problems in Critically Inpatients: Experience Report

Gabrielle Mari Rosetti Alves¹, Fabiana Rossi Varallo¹, Rosa Camila Lucchetta² and Patricia de Carvalho Mastroianni³

¹Drugs and Medicines Department of the School of Pharmaceutical Sciences of Universidade Estadual Paulista “Júlio de Mesquita Filho” (UNESP), Brazil
²Pharmacy Department of Universidade Federal do Paraná (UFPR), Brazil
³Corresponding author: Patricia de Carvalho Mastroianni, Faculdade de Ciências Farmacêuticas – UNESP, Rodovia Araraquara - Jaú, km 1 - CEP: 14801-902, Araraquara – SP, Brazil, Tel: +55 16 3301-6977; Fax: +55 16 3322-0073, +55 16 3301-6960; E-mail: pmastro@fcfar.unesp.br

Received date: July 21, 2014, Accepted date: July 25, 2014, Published date: August 1, 2014

Abstract

This is an experience report on clinical pharmacy in New York, United States of America, in a teaching hospital, describing the results of drug therapy monitoring in critically ill patients, as well as interventions to solve or prevent identified drug therapy problems. The cross-sectional study was conducted by the clinical staff at the Surgical Intensive Care Unit during August 20th to 24th, 2012. Blood counts, serum levels of certain antibiotics, microbiological cultures and their antibiotic susceptibility, possible drug interactions, dosage of each drug prescribed and the compatibility between the route of administration and pharmaceutical form were assessed daily through review of electronic medical records. Twenty seven patients were followed up and 16 drug therapy problems were identified: Unnecessary drug therapy (seven), adverse drug reaction (four), needs additional drug therapy (two), noncompliance (two) and dosage too low (one). After evaluation, the drug therapy problems and their pharmaceutical interventions were reported to clinical pharmacist responsible for the Surgical ICU, as well as the multidisciplinary team. Further, the clinical outcomes were monitored and interventions were classified as to its acceptance. Data demonstrate that clinical pharmacists can contribute to the security and proper use of medications, as the trigger tools for intensive monitoring helps in early detection of drug therapy problems and patient safety.

Keywords: Patient safety; Pharmacovigilance; Drug-related side effects and adverse reactions; Intensive Care Units

Introduction

Pharmacovigilance is a cross practice to the role of clinical pharmacy, since drug therapy monitoring helps to detect problems related to safety, effectiveness and quality of drugs [1].

According to Cipolle [2], the drug therapy problems are events that prevent or delay patient recovery. The processes for identifying, resolving and preventing drug therapy problems is a mainstay of pharmacotherapy management, also being the scope of post-marketing surveillance of drugs [3].

Studies show that the inclusion of the clinical pharmacist in the intensive care unit reduces mortality and adverse events [4-7], improves clinical outcomes and reduces hospital costs [6-8].

Considering that polypharmacy and the high proportion of drug administration by intravenous contribute to increase the risks associated with the use of medications, drug therapy monitoring in critically ill patients is important [9]. Therefore, this study aimed to describe the results of drug therapy monitoring of critically ill patients, as well as interventions to prevent or resolve identified drug therapy problems.

Materials and Methods

This is an experience report on clinical pharmacy in New York, United States of America, in a teaching hospital, member of the New York Presbyterian Health System. The hospital has 651 beds, of those, 27 are for Surgical Intensive Care Unit (ICU), where was developed this report experience. Surgical ICU is intended for all adult patients, in preoperative and postoperative period. Inclusion criteria were all patients in the unit during the study period.

The hospital is certified by the Joint Commission on Accreditation of Healthcare Organizations (JCAHO) and fulfills the Health Insurance Portability and Accountability Act of 1996 (HIPAA), which protects patient identification, preventing their names and personal data of been published. Therefore, in this study patients will be identified by the Medical Record Number (MR #).

At the Surgical ICU the clinical pharmacy services are developed through reviewing electronic medical records of all patients. For this analysis, we considered all medical records during August 20th to 24th, 2012.

Electronic charts were daily monitored to detect possible drug therapy problems. Trigger tools (blood count, serum levels of certain antibiotics, microbiological cultures and their antibiotic susceptibility) were followed up. The staff also evaluated the occurrence of possible drug interactions, dosage of each drug prescribed and the compatibility between the route of administration and pharmaceutical form used.

After reviewing the medical records, clinical rounds were initiated in order to verify the presence of mechanical ventilation, enteral or parenteral nutrition and ability to ambulate.
Drug therapy problems identified were classified according to their etiology [2] and seriousness [10]. Thus, according to the etiology, drug therapy problems can be classified into:

1. Unnecessary drug therapy: The drug therapy is unnecessary because the patient does not have a clinical indication at this time.

2. Needs additional drug therapy: Additional drug therapy is required to treat or prevent a medical condition.

3. Ineffective Drug: The drug product is not effective at producing the desired response.

4. Dosage too low: The dosage is too low to produce the desired response.

5. Adverse drug reaction: The drug is causing an adverse reaction.

6. Dosage too high: The dosage is too high resulting in undesirable effects.

7. Noncompliance: The patient is not able or willing to take the drug regimen appropriately.

According to the National Council for Medication Error Reporting and Prevention (NCC MERP), an organization that aims to develop strategies to assist health professionals to avoid possible medication errors, provides that when the performance of a health professional could lead to inappropriate medication use or harm the user, these events can be classified according to the potential medication errors:

1. Category A: The situation has the ability to lead to a medication error.

2. Category B: A medication error occurred, but was previously identified to the application of the medication.

3. Category C: A medication error occurred, but did not cause harm to the patient.

4. Category D: A medication error occurred, resulting in the need for patient monitoring.

5. Category E: A medication error occurred and contributed temporarily to clinical worsening of the patient. Intervention was required.

6. Category F: A medication error occurred and contributed temporarily to clinical worsening of the patient, resulting in hospitalization or prolongation of hospitalization.

7. Category G: A medication error occurred and may have resulted in permanent damage to the patient.

8. Category H: A medication error occurred causing the need for intervention to sustain life.

9. Category I: A medication error occurred that may have contributed to the patient’s death.

After assessment, drug therapy problems and their pharmaceutical interventions were reported to clinical pharmacist responsible for Surgical ICU, as well as the multidisciplinary team. Further, the clinical outcomes were monitored and interventions were classified as to its acceptance.

Accepted interventions were considered those in which there was a change of prescription medication in the system. Not accepted interventions were considered those which there were not properly completed or have not been implemented due to the discharge of the patient.

Results

During the period, 27 patients were followed, whom seven had drug therapy problems and 16 interventions were implemented. The drug therapy problems were classified into unnecessary drug therapy (seven), adverse drug reaction (four), needs additional drug therapy (two), noncompliance (two) and dosage too low (one). The situations identified were classified in the categories A (seven), B (seven) and C (two) from NCC MERP classification (Table 1).

The overall prevalence demonstrates that clinical pharmacy services permeated 26% of patients in the surgical ICU, in other words, in a proportion of four patients, one required of the clinical pharmacist activity, and could have had clinical complications if there was not a clinical pharmacist seeking improvements to their drug therapies.

Moreover, the prevalence of drug therapy problems demonstrates that for each patient treated by clinical pharmacist were found at least two drug therapy problems liable to intervention.

Discussion

By convention the JCAHO 2001, patients should not feel pain [11]. However, from this convention, it was observed the increasing use of opioids and also the increased frequency of adverse reactions. Subsequent studies have concerns, security should not restrict the effectiveness of pain management, so therapeutic options with opioids should be available [12].

According to Honiden [13], the pain goals should be individualized to meet the needs of each patient, and for being subjective, the most reliable evaluations require the active participation of the patient. Among patients who cannot communicate, the pain can be inferred from observable behavior through the aid of instruments such as ‘Behavior Pain Scale’ or ‘Critical Care Pain Observation Tool’, but there are limitations due to interpretation.

Most surgical ICU patients reported pain or makes use of opioids during surgery. In these cases, because of published studies warning about the possible adverse reactions, the JCAHO recommends monitoring of them [14-16]. Therefore, in order to monitor the occurrence of adverse reactions caused by opioids, symptoms such as nausea, vomiting, dizziness and sedation should be reported to the multidisciplinary team for treatment.

As the hospital is accredited by the JCAHO, should follow these recommendations, and therefore, patients MR #4021893, MR #742896 and MR #4924432 received pharmaceutical interventions to prevent drug related problems that are common to these drugs and encourage communication between patients, doctors and nurses.

The MR #4021893 patient received intervention, in order to prevent receive the medicine that was prescribed only for the day of surgery (morphine, fentanyl and ondansetron).

The MR #742896 patient was receiving two drugs of similar action mechanism, metoprolol and labetalol, which could accentuate the adverse effects of these drugs, in addition to increasing the likelihood of a hypotensive crisis. The perception by the clinical pharmacist and the acceptance of pharmaceutical intervention on the same day may have avoided such drug therapy problems [17-19]. This same patient also had other preventive intervention in order to avoid
administration of hydromorphone medication, scheduled only for the surgery day. With the discontinuation of this medication, potentially serious drug related problems may have been avoided, such as hypotension, bradycardia, bronchospasm, among other potential adverse reactions [20].

<table>
<thead>
<tr>
<th>Drug therapy problems etiology</th>
<th>Drug therapy problems</th>
<th>NCCMERP</th>
<th>Pharmaceutical intervention</th>
<th>Acceptance (reason for not acceptance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unnecessary drug therapy</td>
<td>Ketorolac scheduled</td>
<td>A</td>
<td>Ask the necessity of NSAIDs. It is necessary only if the patient feels pain. Change the status to PRN</td>
<td>Not accepted (discharged)</td>
</tr>
<tr>
<td>Unnecessary drug therapy</td>
<td>Morphine, fentanyl and ondansetron scheduled. It should be prescribed only for the day of surgery</td>
<td>B</td>
<td>Discontinue morphine, fentanyl and ondansetron</td>
<td>Not accepted (discharged)</td>
</tr>
</tbody>
</table>

**Patient Medical Record 4021893**

<table>
<thead>
<tr>
<th>Drug therapy problems etiology</th>
<th>Drug therapy problems</th>
<th>NCCMERP</th>
<th>Pharmaceutical intervention</th>
<th>Acceptance (reason for not acceptance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unnecessary drug therapy</td>
<td>Droperidol scheduled</td>
<td>A</td>
<td>Change the status to PRN.</td>
<td>Not accepted (discontinued)</td>
</tr>
<tr>
<td>Unnecessary drug therapy</td>
<td>Metoprolol and labetalol scheduled</td>
<td>B</td>
<td>Discontinue labetalol</td>
<td>Accepted</td>
</tr>
<tr>
<td>Unnecessary drug therapy</td>
<td>Hydromorphone scheduled when it should have been stopped the day before</td>
<td>B</td>
<td>Discontinue hydromorphone</td>
<td>Accepted</td>
</tr>
</tbody>
</table>

**Patient Medical Record 742896**

<table>
<thead>
<tr>
<th>Drug therapy problems etiology</th>
<th>Drug therapy problems</th>
<th>NCCMERP</th>
<th>Pharmaceutical intervention</th>
<th>Acceptance (reason for not acceptance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Noncompliance</td>
<td>Metoprolol bid</td>
<td>B</td>
<td>Change the status to q12h</td>
<td>Accepted</td>
</tr>
</tbody>
</table>

**Patient Medical Record 4924432**

<table>
<thead>
<tr>
<th>Drug therapy problems etiology</th>
<th>Drug therapy problems</th>
<th>NCCMERP</th>
<th>Pharmaceutical intervention</th>
<th>Acceptance (reason for not acceptance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unnecessary drug therapy</td>
<td>Metoclopramide scheduled</td>
<td>A</td>
<td>Change the status to PRN or discontinue it</td>
<td>Not accepted (discharged)</td>
</tr>
</tbody>
</table>

**Patient Medical Record 4348201**

<table>
<thead>
<tr>
<th>Drug therapy problems etiology</th>
<th>Drug therapy problems</th>
<th>NCCMERP</th>
<th>Pharmaceutical intervention</th>
<th>Acceptance (reason for not acceptance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Needs additional drug therapy</td>
<td>No stress ulceration prophylaxis</td>
<td>A</td>
<td>Add proton pump inhibitor</td>
<td>Accepted</td>
</tr>
</tbody>
</table>

**Patient Medical Record 4918096**

<table>
<thead>
<tr>
<th>Drug therapy problems etiology</th>
<th>Drug therapy problems</th>
<th>NCCMERP</th>
<th>Pharmaceutical intervention</th>
<th>Acceptance (reason for not acceptance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Needs additional drug therapy</td>
<td>PRN metoprolol</td>
<td>C</td>
<td>Change the metoprolol status to scheduled</td>
<td>Not accepted (medical decision)</td>
</tr>
</tbody>
</table>

**Patient Medical Record 4923104**

<table>
<thead>
<tr>
<th>Drug therapy problems etiology</th>
<th>Drug therapy problems</th>
<th>NCCMERP</th>
<th>Pharmaceutical intervention</th>
<th>Acceptance (reason for not acceptance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dosage too low</td>
<td>Glucose: 174 mg / dL</td>
<td>B</td>
<td>Increasing the frequency of aspartate 100 U insulin / mL to q8h</td>
<td>Accepted</td>
</tr>
</tbody>
</table>

**Patient Medical Record 9797720**

<table>
<thead>
<tr>
<th>Drug therapy problems etiology</th>
<th>Drug therapy problems</th>
<th>NCCMERP</th>
<th>Pharmaceutical intervention</th>
<th>Acceptance (reason for not acceptance)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adverse drug reaction</td>
<td>Absence of therapeutic drug monitoring collection after the 5th dose of vancomycin</td>
<td>B</td>
<td>Collect blood sample in vancomycin trough (30 minutes before the next dose)</td>
<td>Not accepted (collection was not performed on vancomycin trough)</td>
</tr>
</tbody>
</table>

Table 1: Description of drug therapy problems identified per patient, according to their etiology, description of pharmaceutical interventions and clinical outcomes observed during August 20th to 24th, 2012, at the Surgical Intensive Care Unit.
To ensure that the patient MR #742896 receive your medicine at optimal dose spacing, it was important that the scheduling of the drug to be q12h instead of bid, because if the patient is in a procedure, and metoprolol scheduled q12h, as soon as the patient returns from the procedure will receive a dose of medicine, and this will not be ignored by the nursing staff. The intervention was accepted.

Studies showed that the rate related to stress-induced ulceration mortality reaches 50% of ICU patients [21]. This type of ulcer lead to clinically significant bleeding in 3-6% of patients, clinically characterized, 24 hours after the onset of ulceration, by decrease equal or greater than 20 mmHg in systolic blood pressure when the patient is bedridden; increased heart rate by 20 beats/min and decreased systolic blood pressure by 10 mmHg when the patient is upright; and also, a decrease in hemoglobin of at least 2 g/dL [22].

Bleeding occurs because at the time of ICU admission, the patient feels stressed, cornered and there is activation of the sympathetic nervous system, with increased release of catecholamines and pro-inflammatory cytokines, which in turn will leads to displacement of blood from the gastrointestinal (GI) tract to organs related with escape, such as muscles and brain. When prolonged, this gastric hypoperfusion and consequent reduction in oxygen supply decrease the secretion of bicarbonate and mucus in the stomach, besides reducing gastric motility. These effects of the GI tract, make it more susceptible to the corrosive action of pepsin and other bile acids, leading to ulceration and constant degradation of formed clots [22].

Once the ulcer is formed in a patient at ICU, the probability of sepsis or multiple organ failure is imminent, may lead him to death. Therefore, to prevent the worsening of a clinical picture to this point, prophylaxis is the most effective way.

In these cases, cimetidine remains the only H2-receptor agonist approved by the Food and Drug Administration (FDA) for the prevention of gastrointestinal bleeding in critical care patients. The scheme approved by the FDA is 50 mg/h IV continuous infusion, however, in practice, intermittent bolus infusion is more used [22]. A comparative study of continuous IV cimetidine versus oral omeprazole resulted in a significant reduction of gastrointestinal bleeding with omeprazole compared with cimetidine [23]. Therefore, prevention is done with the use of proton pump inhibitors.

MR #4924432 and MR #4923104 patients were able to receive oral medication, but were still receiving proton pump inhibitor, pantoprazole, intravenously. For economic reasons and to prevent possible infections and formation of biofilms [24], it is practice prioritize oral instead of parenteral, when it is possible. However, patients were discharged and would not need prophylaxis any more.

The MR #4348201 patient’s metoprolol status was scheduled and PRN. In this case, the scheduled medicine should be evaluated as to dosage, or even combined another antihypertensive drug (different mechanism of action), if it was not controlling blood pressure effectively [25,26]. But it should not be in PRN in cases of hypertension, because the likelihood of drug therapy problems would be increased.

Medicines for blood pressure control are extremely important to the health and welfare of the patient, it must always be on scheduled status. The metoprolol prescription of patient MR #4918096, was PRN, which could lead to drug therapy problems such as rebound hypertension due to the inconsistent use of the drug [25].

Drugs that have an irritant effect on the stomach, such as acetylsalicylic acid can be coated with a substance that will dissolve only in the small intestine, when crushed to be able to pass through the nasogastric tube of patient MR #4918096, lose this property, thus having a different effect than expected [27]. The suggestion of the exchange for chewable presentation, demonstrates how important it is the presence of the clinical pharmacist in the round, along with other professionals in the multidisciplinary team, because it was possible that this patient had worsening of their clinical condition by ineffectiveness of the drug chosen or adverse drug effects. Furthermore, Viktil8 demonstrated that the presence of clinical pharmacist during rounds, gives greater confidence to other professionals of the multidisciplinary team, increases the percentage of acceptance of suggested interventions.

The MR #4923104 patient had diabetes mellitus and insulin use was for glycemic control and prevention of infections [28]. But the insulin was not controlling glycaemia as required; a greater amount of insulin was necessary, thus the suggestion was to increase the administration dose to q6h. Glucose decreased from 174 mg/dL to 153 mg/dL.

Patient safety is a priority for all employees, especially for pharmacists, since despite all precautions, some medications may be unsafe for certain patients, as was the case of the patient MR #979720, with severe infection, requiring, therefore, vancomycin, which could be diluted in 5% dextrose. Apparently the diluted could be raising the patient’s blood sugar level, corresponding to a therapeutic problem. The change of blood glucose level by diluent is still a controversial topic, but has been reported in hospitalized patients [29].

The drugs used in the ICU or operating room may also induce hyperglycemia such as exaggerated administration of epinephrine and norepinephrine, resulting in insulin resistance by increasing gluconeogenesis. Hyperglycemia has been hypothesized that prevent the normal physiological responses against infections: mechanisms including inactivation of the complement system, uneven adhesion of granulocytes, impaired phagocytosis (occurs at even lower levels than 200 mg/dL), delayed and decreased Oxidative “Burt” chemotaxis [28].

Hyperglycemia is common in ICU patients, regardless of whether these patients have diabetes mellitus, and is associated with increased morbidity and mortality [28]. Therefore, for the case of a patient who underwent surgery, and presents a serious infection, the blood glucose level should be controlled. It was suggested that the amount of glucose delivered to the patient MR #979720 was reduced and the solvent was replaced for normal saline, given its physical and chemical compatibility [29]. After the intervention, the patient’s blood glucose decreased from 188 mg/dL to 118 mg/dL.

The main antibiotics used at U.S. that requires therapeutic drug monitoring are vancomycin, when administered as IV, and gentamicin. These drugs, as are broad coverage antibiotics, are in general the first medical choice, while they wait the results of susceptibility testing. According to the pharmacokinetic parameters of these drugs, the measurement of trough phase is the easiest way to know the effectiveness of therapy compared to assessment by the ratio of the area under the curve on minimum inhibitory concentration (AUC/MIC) clinical form, as in this case many samples of the patient’s blood is required. Thus, determine the concentration in the trough phase is the closest parameter to ensure that the patient is receiving the minimum appropriate amount of antibiotic to fight infection, and reduce the phenomenon of bacterial resistance [30].
The suitable serum levels at trough phase are the minimum antibiotics amount to combat infecting bacteria and depend on the antibiotic and the severity of the infection. For vancomycin serum trough levels recommended range between 15-20 mg/mL for serious infections (bacteremia, endocarditis, osteomyelitis and meningitis and pneumonia) [30-32].

There is evidence that, for patients in the ICU, is economically viable monitoring of vancomycin levels in an attempt to avoid nephrotoxicity, particularly those who are also receiving other potentially nephrotoxic drugs, as well as cancer patients. This fact demonstrates that the presence of a clinical pharmacist brings benefits not only to the health of patients, but also for the health service by reducing new clinical problems, becoming an economic benefit [30].

The MR #979720 patient has not had vancomycin levels dosed since the fourth dose of the antibiotic. However, the intervention was applied incorrectly, because to obtain the measurement at trough phase is necessary that the blood collection be made 30 minutes before the next dose of vancomycin [30].

There was a predominance of drug therapy problems related to unnecessary pharmacotherapy, different from what was found by Reis [33] in Brazil, where the use of unnecessary or inappropriate medication is the second most common error, being dose errors the first.

The emergence of a new drug therapy problem related to the use of unnecessary medication can prolong the stay of these patients in the ICU, a fact that is financially inefficient for the hospital, to the patient, the health insurance of the patient and/or the government [8]. Although no situation of category F (NCCMERP) have been identified, the presence of the pharmacist in the multidisciplinary team allows identification and interception of medication errors.

Conclusion

The prevalence drug therapy problems in critically ill patients demonstrated that to each patient, at least two drug therapy problem can be prevented and/or remedied by clinical pharmacists. Data demonstrate that this professional can contribute to the security and proper use of medications, as the trigger for intensive monitoring helps in early detection of drug therapy problems, therefore patient safety.

References

15. The Joint Commission Sentinel Event Alert. JCAHO.
