Role of oncology clinical pharmacist: a case of life-saving interventions

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INTRODUCTION
CP role is emerging as vital addition to the health care team resulting in improved patient care. Interacting with health care professionals on rounds, patient interview, medication reconciliation, and providing patient discharge counseling and follow-up all resulted in improved outcomes. In oncology setting, CP has a major role in the adjustment of medications and being part of multidisciplinary team is recommended to prevent and manage medications errors.

CP has to follow and evaluate the chemotherapy protocols regarding appropriate doses, potential drug interactions and side effects management to achieve safe and effective drug therapy and to prevent medication errors. CP has potential role in symptoms management and was effective in optimizing the use of antiemetic for chemotherapy induced nausea and vomiting. Pain management is another field were clinical pharmacist interventions were effective in improving safety, efficacy and patient satisfaction. Also CP participation in patient care is important to improve patient compliance to the prescribed medications and avoidance of cost-ineffective drug therapy.

CASE REPORT

Case summary
We report a case of 42 years old male patient with Non-Hodgkin Lymphoma (DLBCL), treated with high dose methotrexate (MTX) as prophylaxis for central nervous system (CNS) lymphoma, and followed up by CP regarding medications related issues. Role of CP is mainly to identify, prevent and manage any drug related problem including drug choice, dosage, interactions, administration and side effects. Oncology CP was very effective in optimizing medication use and has a promising role through providing clinically important interventions regarding medication use. We present this educational case report to show the vital role of CP in patient care by providing important interventions that are effective and sometimes life-saving.

Keywords: Clinical pharmacy, Methotrexate, Oncology, Chemotherapy, Pharmaceutical care
axillary lymph nodes, mild heart burn, constipation and moderate ascites.

**Relevant lab tests**

Albumin 1.48 g/dl, Liver Enzymes (ALT 45 IU/L, AST 32 IU/L), Total Bilirubin: 0.53 mg/dl, Phosphate 7.7 mg/dl, Potassium 3.6 mEq/L, Uric Acid 8.8 mg/dl, Magnesium 2.3 mEq/L, Sodium 138 mEq/L, Chloride 98 mEq/L. Neutrophils count 1100/μL, Platelets 170,000/μL, White Blood Cells 1600 μL, Serum Creatinine 0.89 mg/dl, Calcium 7.5 mg/dl, Lactate Dehydrogenase 620 IU/L.

- Calculated Creatinine Clearance using Cockcroft and Gault formula: 104 ml/min.
- Calculated body surface area (BSA) using Mosteller Method: 2 m².

**Current medications**

<table>
<thead>
<tr>
<th>Medication</th>
<th>Dose &amp; Frequency</th>
<th>Route</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Omeprazole</td>
<td>20 mg once daily</td>
<td>Oral</td>
<td>Stress ulcer prophylaxis</td>
</tr>
<tr>
<td>TMP-SMX*</td>
<td>960 mg every 48 hours</td>
<td>Oral</td>
<td>PCP** prophylaxis</td>
</tr>
<tr>
<td>Calcium Carbonate</td>
<td>500 mg 3 times daily</td>
<td>Oral</td>
<td>Hypocalcemia</td>
</tr>
<tr>
<td>Senna</td>
<td>24 mg once daily</td>
<td>Oral</td>
<td>Constipation</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>400 mg 3 times daily</td>
<td>Oral</td>
<td>Osteoarthritis</td>
</tr>
</tbody>
</table>

*Trimethoprim-Sulfamethoxazole, **PCP: Pneumocystis Carinii Pneumonia

**Plan**

Therapeutic plan was to give intravenous (I.V) high dose MTX for CNS lymphoma prophylaxis according to the following protocol (body surface area= 2 m²):

**Urine Alkalization:**

Start oral sodium bicarbonate 1.5 g every 6 hours. Dipstick urine every 2 hours to check PH. If pH <7 give additional bicarbonate.

**Hydration:**

Start IV hydration 12 hours before administration of MTX with Sodium chloride 0.45% with sodium bicarbonate 50 mEq per liter at rate of 150 ml/hour.

Continue hydration throughout and post MTX administration until MTX level is less than 0.05 mcg/ml.

**Premedications:**

Ondansetron 8 mg I.V. once pre MTX.

Dexamethasone 8 mg I.V. once pre MTX.

**Chemotherapy:**

Methotrexate (3.5 gm/m²) = 7 grams in Dextrose 5% in water 1000 mL I.V over 6 hours on day 1.

Leucovorin (25 mg/m²) = 50mg IV beginning 24 hours after the start of MTX and continue every 6 hours until the serum MTX level is <0.05mcg/ml.

* Check urine pH every 8 hours. If less than 7, give sodium bicarbonate 50 mEq IV.

* Therapeutic drug monitoring (MTX post-dose level):

MTX levels every morning at 24, 48 and 72 hours after the completion of MTX and continue daily until serum MTX level is <0.05mcg/ml.

**Clinical pharmacist interventions**

1. To stop TMP-SMX one week before chemotherapy, since it associated with serious and fatal interaction that increases risk of MTX toxicity.

2. Drainage of the ascites should be done before initiation of MTX; consider adding Albumin (20%) 100 ml every 12 hours and Furosemide 20 mg every 12 hours to resolve ascites before MTX therapy to prevent MTX toxicity.

3. To replace Omeprazole 20 mg with oral Ranitidine 150 mg twice daily to avoid the interaction between proton pump inhibitors (PPIs) and MTX.

4. To hold Ibuprofen and resume (if needed) after MTX level decreased to <0.05 mcg/ml since non-steroidal anti-inflammatory drugs (NSAIDs) interfere with the excretion of MTX and may potentiate toxicity; consider oral Paracetamol 1000 mg 4 times daily as alternative.

5. To add oral Allopurinol 300 mg once daily and oral Sevelamer 800 mg three times daily to treat hyperuricemia and hyperphosphatemia, respectively.

6. Consider stopping Calcium Carbonate replacement if there is no other indication.

**Follow up**

- Pre-hydration was given as recommended by the protocol (150 ml/hr). CP found that post-hydration was given as 15 ml/hr for 6 hours (instead of 150 ml/hr) due to nurse mistake, assigned consultant was informed and
the decision was to hold the post-hydration and to give 1 liter of sodium chloride 0.45% over 2 hours, then post-hydration was resumed as 150 ml/hr again as it should be.

- After 24 hours:

MTX level was 57 mcg/ml, urine PH was 7, serum creatinine increased from 0.89 mg/dl to 2.1 mg/dl, Total Bilirubin 0.5 mg/dl and Liver enzymes elevated (ALT 152 IU/L & AST 121 IU/L). Patient developed mild dermatitis on the back, nausea and vomiting. Also the patient complained of diarrhea for which Loperamide was prescribed. No pulmonary or neurological changes were observed.

The CP recommended the following actions:

7- Continue intravenous hydration with addition of Furosemide 20 mg twice daily to maintain urine adequate output.

8- Increase Leucovorin dose to be 100 mg every 6 hours and continue until MTX level <0.05 mcg/ml.

9- Continue oral Sodium Bicarbonate as 1.5 g every 6 hours to maintain urine PH >7, preferably >8.

- After 48 hours:

MTX level has dropped to 1.88 mcg/ml, urine PH was 8, serum creatinine decreased to 1.42 mg/dl with continuous hydration, Total Bilirubin 0.52 mg/dl, Liver Enzymes decreased (ALT 73 IU/L, AST 68 IU/L) and patient is still suffering from the symptoms of diarrhea and dermatitis.

- After 72 hours:

MTX level was decreased to 0.08 mcg/ml and the patient was stable, symptoms of diarrhea and dermatitis improved and patient was discharged 5 days later.

DISCUSSION

DLBCL, the most common histological type of non-Hodgkin's lymphoma, is an aggressive disease associated with CNS secondary involvement in 3-5% of cases and mainly treated with the R-CHOP chemotherapy protocol. Rate of CNS involvement is dramatically increased in patients with certain risk factors at the time of diagnosis.6

Elevated LDH, age >60 years, involvement of >1 extranodal site, hypoalbuminemia and high International Prognostic Index (IPI) score are common risk factors for secondary CNS lymphoma. Patients with ≥2 of these risk factors should receive prophylaxis either with intrathecal or high-dose systemic chemotherapy.8 Our patient had ≥2 risk factors, so the decision was to give high dose MTX as CNS prophylaxis. The patient was taking TMP-SMX for PCP prophylaxis. CP recommendation was to stop this medication 1 week before starting MTX therapy because there is a well-documented fatal drug interaction between MTX and TMP-SMX causing increased risk of MTX toxicity like serious pancytopenia9, so this intervention by CP is very important and considered life-saving.

The use of high dose MTX is contraindicated in the presence of a third space fluid like pleural effusions or ascites. Presence of ascites will prolong MTX plasma half life and increase the risk of toxicity. It is recommended to drain the accumulated fluid before MTX therapy.10

Using PPIs concurrently with MTX is considered major risk factor for delayed MTX elimination and increased toxicity, so PPIs should not be used during MTX therapy.11 Thus, the CP recommendation was to discontinue Omeprazole and use Ranitidine 150 mg twice daily as safe alternative. Similarly, MTX elimination is also impaired in patients receiving concurrent NSAID therapy like Ibuprofen. So it is recommended to avoid such combination.12

Patients treated with chemotherapy for DLBCL are at high risk for tumor lysis syndrome (TLS). Allopurinol for hyperuricemia and Sevelamer for hyperphosphatemia were recommended to this patient as treatment and also as prophylaxis for TLS.13 There is no clear indication for Calcium Carbonate replacement in this patient. Calcium level is 7.5 mg/dl and albumin level is also low, but corrected calcium level is 9.52 mg/dl which is normal.

When MTX toxicity is suspected based on high MTX level and symptoms, maintaining urine output, urinary alkalinization, monitoring serum creatinine, electrolytes and plasma MTX concentrations, and pharmacokinetically-guided leucovorin rescue, are the cornerstones of management for patients who develop early signs of renal dysfunction and delayed MTX elimination.14 CP recommendations were mainly directed to adjust Leucovorin dose, maintain urine PH >7 and continuous hydration.

This case report clearly shows the important role of the oncology CP as a member of the oncology multidisciplinary team, in optimizing medication use to achieve safe, effective and cost-effective drug therapy. By providing therapeutic interventions, CP can monitor, detect, prevent, and manage many serious and sometimes fatal drug-related problems.

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REFERENCES
