Potential Leukocytic Toxicity Induce by Vinorelbine, Doxorubicin and Cisplatin in Human Patients of Breast, Cervix and Lung Cancer

Taha Nazir1,2, Nida Taha2, Muzhar Mustafa3, Azharul Islam4, Adeel Mahmood5 and Saeed ur Rasheed Nazir6

Abstract
Anticancer drugs are used extensively in chemotherapy. Bone marrow suppression is one of their major side effects. Therefore, we aimed this study to investigate the potential leukocytic toxicity induced by vinorelbine in cancer patients. The data of total 60 adult patients were selected and divided into two groups; Group-1 patients received Vinorelbine alone and group-2 Vinorelbine based combinations. The outcomes demonstrated no statistically important difference in the patients who were either on vinorelbine alone, vinorelbine plus cisplatin or vinorelbine plus doxorubicin combinations. On comparison of mean value SEM ± count (109) Per µL overtime pre & post chemotherapy, no statistical differences were observed in Basophil and Lymphocyte counts for Group-I; Eosinophil, Basophil, Monocyte, and Lymphocyte counts for Group-II. Comparison of mean values ± SEM (x109) per µL before therapy with that of at weeek-4 (pre and post chemotherapy) showed no significant difference in Eosinophil, Basophil and Lymphocyte counts for Group-I (P value 0.102, 0.221, 0.063); and TLC, Neutrophil, Eosinophil, Basophil, Monocyte and Lymphocyte counts (P value 0.148, 0.118, 0.665, 0.314, 0.083, 0.427) for Group-II. Hence, the overall Leukocytopenia, Neutropenia, Eosinopenia, Basopenia, Monocytopenia and Lymphocytopenia in both of the chemotherapy protocols allow the clinical oncologists and consultant physicians to select either of the chemotherapeutic agents. The therapeutic efficacy should constitute the intervening consideration in treating the patients of breast, cervix and lung cancer.

Keywords
Leukocytosis; Leucopenia; Vinorelbine; Cisplatin; Doxorubicin; Breast cancer; NSCLC

Introduction
Cancer is a fatal dilemma of human life and more mysterious of the major life threatening diseases. Despite of the scientific miracles, cancer is still a very real concern to public health, both in perception and reality. It is being treated stereoscopically with good or bad results. Cancer is still a very real concern to public health, both in perception and reality. It is being treated stereoscopically with good or bad results. There are several types and stages of chemotherapy used to treat cancer; induction therapy, consolidation therapy, intensification therapy, maintenance therapy, adjuvant therapy, palliative therapy and chemo preventive therapy [2]. We have focused over the chemotherapy of any type and stage of patient diagnosed as cancer; the leukocytic toxicities produced by Vinorelbine alone and combinations.

Materials and Methods
This retrospective study was conducted at Shaukat Khanum Memorial Cancer Hospital & Research Center (SKMCH&RC), M.A Johar town, Lahore, Pakistan. We have collected the information's of Non-small cell lung cancer, metastatic breast cancer, and cervix cancer patients, taken Vinorelbine alone, Vinorelbine/ Doxorubicin and Vinorelbine/Cisplatin treatment protocols. Our major goal was to investigate the changes in leukocytic laboratory profile of human cancer patients.

Study design
These patients were selected from outpatient department (OPD) of SKMCH&RC. The patients' selection criterion was including as under;
1. Should be differentially diagnosed the neoplasm.
2. Patients with Metastatic breast / NSCL/ Cervix Cancer
3. Patients took vinorelbine as part of their treatment in clinical setting.
4. Having no history of blood or liver disease
5. Either sex of male or female.

Similarly, an exclusion criterion was also designed to scrutinize the patient for this study. A total 60 cancer patients were divided into two groups; Group-1 comprising of patient received vinorelbine as single therapy and Group-2 having the cancer patients on treatment protocol of vinorelbine based combinations i.e. Vinorelbine/ Cisplatin or vinorelbine/Doxorubicin (Table 1).

Table 1: The chemotherapy protocols follow up schedule and cancer site of experimental patients.

<table>
<thead>
<tr>
<th>Group</th>
<th>Sample size</th>
<th>Chemotherapy protocol</th>
<th>Patient neoplasm type</th>
<th>Chemotherapy schedule (days)</th>
<th>Follow up schedule (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>G-I</td>
<td>45</td>
<td>Vinorelbine</td>
<td>Metastatic breast cancer</td>
<td>1, 7, 14, 21</td>
<td>6, 13, 20, 28</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>NSCL cancer</td>
<td>1, 7, 14, 21</td>
<td>6, 13, 20, 28</td>
</tr>
<tr>
<td>G-II</td>
<td>15</td>
<td>Vinorelbine/ Doxorubicin</td>
<td>Metastatic breast cancer</td>
<td>1, 8</td>
<td>7, 15</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>NSCL cancer</td>
<td>1, 8</td>
<td>7, 15</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Cervix Cancer</td>
<td>1, 8</td>
<td>7, 15</td>
</tr>
</tbody>
</table>
Preparations of standard regimen of chemotherapeutical agents

The standard treatment regimen for vinorelbine, cisplatin and doxorubicin is reported by Taha et al. [3,4]. The vinorelbine was administered 25 mg/ml on day 1, weekly 4, i/v, with 0.45% sodium chloride or 5% glucose solution as diluents and delivered over a period -15 to 20 minutes [6]. In combination therapy the dose of Vinorelbine was decreased and administer as 20 mg/ml on day 1, 8 I/V with diluents day 5½ normal saline and delivered over IVP. The Doxorubicin was given as 50 mg/m² on day 1 only [7]. Doxorubicin was administered slowly in to tubing of freely running infusion of Sodium Chloride 0.9% or Glucose 5% [8]. The Cisplatin was administered intra-venously as 40 mg/ml on day 1 only, with the diluent of day 5% NS and delivered over IVP.

Sample collection and neutrophils count

The 3 ml of blood samples were drawn from brachial veins in 5 cc disposal syringes and transferred to appropriately labeled (complete blood count (C.B.C) vials containing 20 w/v of EDTA as described by Taha et al. [3]. The neutrophils count was performed using a computerized auto-analyzer (Technicon 113, Bayer Laboratories USA) at the Pathology laboratory, SKMCH&RC.

Data analysis

The means of two groups were compared by student t-Test to avoid the consistent deviation of analytical results or systematic errors in the procedure. ANOVA used to identify any factor influencing the test results. The obtained p values of Group-I, Group-II and overall (60) patients were compared before, after and every week to identify potential leucocytopenia.

Result

Comparison of mean value SEM ± count (10³) Per uL overtime pre & post chemotherapy showed no statistical differences were observed in Basophil and Lymphocyte counts (P value 0.435, 0.64) for Group-I; Eosinophil, Basophil, Monocyte, and Lymphocyte counts (P value 0.759, 0.437, 0.08, 0.23) for Group-II (Table 2). On comparison of mean values ± SEM (x10³) µL before with that of a week-1,2,3,4 and 5 indicate no significant difference in TLC (P value 0.607, 0.944, 0.897); Neutrophil (P value 0.742, 0.208, 0.425, 0.048, 0.791); Eosinophil (P value 0.488, 0.145, 0.171, 0.738); Basophil (P value 0.517, 0.089, 0.434, 0.475, 0.270); Monocyte (P value 0.551, 0.112, 0.559, 0.372, 0.468); and Lymphocyte (P value 0.736, 0.555, 0.727) (Table 3). Comparison of mean values ± SEM (x10³) µL per before therapy with that of at week-4 (pre and post chemotherapy) showed no significant difference in Eosinophil, Basophil and Lymphocyte for Group-I (P value 0.102, 0.221, 0.063); and TLC, Neutrophil, Eosinophil, Basophil, Monocyte and Lymphocyte (P value 0.148, 0.118, 0.665, 0.314, 0.053, 0.427) for Group-II (Table 4).

The significant statistical differences were noted at every week pre and post chemotherapy in the mean (10³) µL counts of neutrophil, Eosinophil & Lymphocyt at week 2 (P value 0.014, 0.006 & 0.003); Lymphocyte at week 3 (P value 0.033); and Leukocyte & neutrophil at week 4 (P value 0.024 & 0.048) (Table 3).

In addition of that, on comparison of the mean SEM ± counts (10³) Per µL before therapy with that of at week-4, no significant statistical differences were observed in Eosinophil, Basophil, & Lymphocyte (P value 0.102, 0.221 & 0.063) in Group-I; and Leukemia, Neutropenia, Eosinopenia, Basopenia, Monocytopenia & Lymphocytopenia (P value 0.148, 0.118, 0.665, 0.314, 0.053 & 0.427) in Group-2 (Table 4).

Discussion

The findings under discussion are in line with the work Dorr et al. [5], who reported the dose limiting leucopenia of Vinorelbine. Marty et al. [9] reported the leucopenia as noncumulative and of short duration (<7 days). While Shamseddine et al. [8] reported the acceptable degree of leucocytopenia induced by Cisplatin and Vinorelbine. Misako et al. [10] reported doxorubicin used for the treatment of lung cancer in Japan. The effects of AMR (Amurubicin hydrochloride) investigated over cultured supernatant. The subcutaneously injected into rabbits introduced a significant decline in number of eosinophil around the injected site.

Moreover, Kharbangar et al. [11] and Cao et al. [12] reported the cisplatin mediated development of various hematological changes in mice bearing ascites (Dalton lymphoma tumor). Cisplatin treatment of tumor bearing mice reduces eosinophil, basophils and lymphocytes along with the development of various morphological abnormalities. However, combination treatment of cytome plus cisplatin resulted in lower the potential of hematological toxicities.

Faller et al. [13] reported the accepted standard of cisplatin adjuvant chemotherapy of non-small-cell lung cancer (NSCLC) patients. Cisplatin and vinorelbine administered for acute myelogenous leukemia and NSCLC.

In addition of that Sauer et al. [14] reported the incidences of...
leucopenia with different response rate induced by two vinorelbine combinations; Vinorelbine, rh-endostatin and Vinorelbine, cisplatin. The collected information’s from different databases were analyzed by Cochrane systematic review methods and the meta-analysis conducted through software RevMan 5.0. Brown et al. [15] and Oostendorp et al. [16] reported the potential therapeutically benefits of vinorelbine and doxorubicin in clinical practice. These were selected for breast cancer patient because of their wide range of anticancer benefits. The studies randomly compared the drugs in combination with targeted agents to provide reasonable scientific evidences for therapeutically usage in advanced breast cancer patients.

Conclusion

There were no significant differences in the overall hematological toxicities of both of the chemotherapy protocols. The consultant oncologist can select either of the protocol to provide maximum relief to patients by assuring successful treatment. Moreover the therapeutically efficacy should probably constitute the overall consideration while treating the particular neoplasm.

References


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