SECONDARY HODGKIN LYMPHOMA AND MYELODYSPLASTIC SYNDROME (MDS) AFTER PACLITAXEL-CARBOPLATIN TREATMENT IN A PATIENT WITH SMALL CELL LUNG CANCER

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ABSTRACT

Herein synchronous occurrence of Hodgkin lymphoma and secondary myelodysplastic syndrome in a 60 year old male patient with small cell lung cancer treated with combined chemotherapy (carboplatin and paclitaxel) and radiotherapy is presented. The objective of this report is to stress the importance of documenting and monitoring adverse drug reactions that arise from chemotherapy. After four years of treatment with the combined chemotherapy, the patient presented inguinal lymphadenopathy and enlarged lymph nodes and histopathology rapport was suggestive for plasmacytoid variant of Castleman disease. Three years later, biopsy of lymph node was performed and diagnosis of Hodgkin lymphoma – mixed cellularity has been established. Molecular analyses revealed presence of dominant monoclonal population of the immunoglobulin genes in the oligo/monoclonal background. Bone marrow biopsy findings suggested secondary myelodysplasia and revealed signs of hematopoietic cells dismaturation with signs of megaloblastic maturation of the erythropoietic lineage, appearance of ALIP (abnormal localization of immature precursors) in the myeloid lineage and dysplastic megakaryocytes. In addition, an increased level of polyclonal plasmacytes (lambda vs kappa was 60%:40%) was found. Hodgkin lymphoma and MDS occurring after 4 years of carboplatin/paclitaxel therapy might be contributed to the accumulation of alkylator-related DNA damage. This emphasize the need of outlining a monitoring plan regarding development of secondary leukemia and other malignant hematological proliferations should be outlined in the protocols.

Keywords: Hodgkin lymphoma, MDS, Castleman disease paclitaxel, carboplatin

INTRODUCTION

Paclitaxel is a tubulin-binding agent that is widely used for the treatment of small cell lung cancer (SCLC), pancreatic, ovarian, breast and other cancers. The combination of paclitaxel and a platinum compound is an approved regimen for the treatment of advanced SCLC [1-3]. The rapid development of new therapies that improve patients’ quality of life and treat various diseases has led to occurrence of more complex adverse drug reactions. This is obvious in cancer therapy where various adverse events are more often neglected by the responsible clinicians because of the primary goal of cancer therapy [4-6]. Hence, documenting and monitoring adverse drug
reactions that arise from chemotherapy should be done more precisely especially in patients who need therapy longer duration and have cumulative dose.

A recent case report has documented a patient that developed secondary acute myeloid leukemia after several months of treatment with cyclophosphamide, alkylating agent used to treat lymphomas and various cancers [4]. Additionally, Gajendra et al, reported a rare case of Hodgkin lymphoma in a patient with chronic myeloid leukemia treated with imatinib, a tyrosine kinase inhibitor [7].

Herein, we report a patient with SCLC treated with combination of paclitaxel/carboplatin and radiotherapy who developed secondary Hodgkin Lymphoma mixed cellularity subtype accompanied with secondary myelodysplastic syndrome.

**CASE DESCRIPTION**

A sixty years old male has been admitted with previously diagnosed SCLC based on native chest X-ray (Figure 1), computer tomography (CT) (Figure 2), and bronchial biopsy rapport (Figure 3). His past medical history was insignificant. He started with first line of chemotherapy and sequential locoregional radiotherapy. The chemotherapy protocol has been consisted from 6 combined cycles of paclitaxel/carboplatin and locoregional radiotherapy with 44Gy. Soon after treatment, he recovered on clinical radiographic evaluation with complete remission of the disease, without signs of infiltration of the lung with malignant disease.

On periodical follow-ups on every six months, patients’ laboratory analyses from the peripheral blood, lung, liver and kidney function tests were within normal range. After four years, the patient presented with inguinal lymphadenopathy and enlarged lymph nodes in the retroperitoneal region on CT. CT also revealed lung fibrosis without enlarged mediastinal lymph nodes. Lymphadenectomy from the inguinal region has been done and the results from the histopathology report were suggestive for plasmacytoid variant of Castleman disease (Figure 4). Axillary lymphadenectomy followed descriptive histological finding for re-
active lesion. Patient was treated with antibiotics and there was improvement of the clinical picture with regression of the enlarged lymph nodes.

Three years later, the patient was admitted at the University Clinic of Hematology, with complaints of fatigue, prostration and increased sweating.

Physical and ultrasound examination revealed cervical, supraclavicular, axillary and inguinal lymphadenopathy with no hepatosplenomegaly. Mediastinal lymphadenopathy was found on CT scan. Abdominal CT scan examination revealed regular liver architecture, slightly enlarged spleen with hypodense region measuring 17 mm, combined with retroperitoneal and mesenterial lymphadenopathy.

Patients’ pertinent hematology and biochemistry laboratory investigation are shown in Table 1. Peripheral blood smear was normal. Lymph node biopsy from the axillary region has been done. A diagnosis of Hodgkin’s lymphoma – mixed cellularity has been established (Figure 5). Molecular analyses revealed presence of dominant monoclonal population of the immunoglobulin

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Results prior diagnosis of Hodgkin`s lymphoma</th>
<th>Follow-up results after ABVD protocol</th>
<th>Reference range</th>
</tr>
</thead>
<tbody>
<tr>
<td>WBC</td>
<td>8.17 x 10^3/µL</td>
<td>2.32 x 10^3/µL</td>
<td>4.00-10.50 x 10^3/µL</td>
</tr>
<tr>
<td>Neutrophil</td>
<td>6.19 x 10^3/µL</td>
<td>1.41 x 10^3/µL</td>
<td>1.40-6.50 x 10^3/µL</td>
</tr>
<tr>
<td>Lymphocyte</td>
<td>1.19 x 10^3/µL</td>
<td>0.57 x 10^3/µL</td>
<td>1.20-3.40 x 10^3/µL</td>
</tr>
<tr>
<td>Monocyte</td>
<td>0.40 x 10^3/µL</td>
<td>0.24 x 10^3/µL</td>
<td>0.00-0.60 x 10^3/µL</td>
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<tr>
<td>Eosinophil</td>
<td>0.38 x 10^3/µL</td>
<td>0.10 x 10^3/µL</td>
<td>0.00-0.70 x 10^3/µL</td>
</tr>
<tr>
<td>Basophil</td>
<td>0.01 x 10^3/µL</td>
<td>0.00 x 10^3/µL</td>
<td>0.00-0.20 x 10^3/µL</td>
</tr>
<tr>
<td>RBC</td>
<td>3.02 x 10^3/µL</td>
<td>2.57 x 10^3/µL</td>
<td>4.00-6.00 x 10^3/µL</td>
</tr>
<tr>
<td>HGB</td>
<td>8.3 g/dL</td>
<td>7.2 g/dL</td>
<td>11.0 – 18.0 g/dL</td>
</tr>
<tr>
<td>HCT</td>
<td>26.3%</td>
<td>23.3%</td>
<td>35.0 – 60.0%</td>
</tr>
<tr>
<td>PLT</td>
<td>78 x 10^3/µL</td>
<td>99 x 10^3/µL</td>
<td>150-450/µL</td>
</tr>
<tr>
<td>Glucose</td>
<td>7.13 mmol/L</td>
<td>-</td>
<td>3.89-6.40 mmol/L</td>
</tr>
<tr>
<td>Urea</td>
<td>6.2 mmol/L</td>
<td>-</td>
<td>3.0 – 9.2</td>
</tr>
<tr>
<td>Creatinine</td>
<td>89.5 µmol/L</td>
<td>-</td>
<td>63.6 – 110.5 µmol/L</td>
</tr>
<tr>
<td>AST</td>
<td>11 U/L</td>
<td>-</td>
<td>5-34 U/L</td>
</tr>
<tr>
<td>ALT</td>
<td>7 U/L</td>
<td>-</td>
<td>0-55 U/L</td>
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<tr>
<td>Alkaline phosphatase</td>
<td>67 U/L</td>
<td>-</td>
<td>40-150 U/L</td>
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<tr>
<td>Bilirubin</td>
<td>31.7 µmol/L</td>
<td>-</td>
<td>3.4 – 20.5 µmol/L</td>
</tr>
<tr>
<td>Albumin</td>
<td>31 g/L</td>
<td>-</td>
<td>35-52 g/L</td>
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<tr>
<td>Fe</td>
<td>6.2 µmol/L</td>
<td>-</td>
<td>11.6 – 31.3 µmol/L</td>
</tr>
</tbody>
</table>
genes in the oligo/monoclonal background. After echocardiographic evaluation of the cardiac function and spirometry for the respiratory capacity that revealed signs of borderline pulmonary arterial hypertension, the patient was further treated for secondary malignant disease Hodgkin lymphoma by ABVD protocol. The hematology results and differential leukocyte count, after applied two cycles of treatment with ABVD D1C1 and ABL D1D2 are shown in Table 1.

Bone marrow biopsy has been done with findings of secondary myelodysplasia (Figure 6). Histological analysis of the bone marrow biopsy revealed signs of hematopoietic cells dysmatura-

tion with signs of megaloblastic maturation of the erythropoietic lineage, appearance of ALIP in the myeloid lineage and dysplastic megakaryocytes. In addition, it was found an increased level of polyclonal plasmacytes (lambda vs kappa was 60%:40%).

**DISCUSSION**

Herein, we report the synchronous occurrence of Hodgkin lymphoma and secondary MDS in a patient with SCLC treated with combined chemotherapy (carboplatin and paclitaxel) and radiotherapy currently on chemotherapy. It is possible that carboplatin and/or paclitaxel contributed likely to the accumulation of alkylator-related DNA damage, due to a predisposition to the myelotoxic effects of chemotherapy [2-6].

Secondary myelodysplastic syndrome or secondary leukemia to chemotherapy usually develops four to seven years after the initial exposure and is often seen in young patients [8-10]. The most common abnormalities are related to chromosomes 5 and 7. Alkylating agents have been considered the most common drugs associated with MDS [6, 11, 12].

Although the potential contribution of paclitaxel and carboplatin to the induction of Hodgkin lymphoma and MDS is controversial, only few cases of the secondary diseases, particularly MDS and leukemia are reported in the literature where only carboplatin and paclitaxel were used as primary chemotherapy, but no radiotherapy [2, 12, 13]. To the best of our knowledge, there are not available data for the synchronous appearance of myelodysplasia and Hodgkin lymphoma. The individual impact of each treatment is uncertain in the presented case, even though the exposure to carboplatin and/or paclitaxel in combination with radiotherapy might have played a role in the development of therapy-related MDS. That might be the first hit in the process of development of the Hodgkin disease due to the escape of the immune surveillance in the condition of dysfunctional immune cells. This hypothesis is supported also by the appearance of the Castelman disease previously in the patient. There have been described therapy-related MDS/AML following cytotoxic chemotherapy of malignant tumors with classic alkylating agents, including nitrosoureas or procarbazin, radiation therapy alone, or combined chemotherapy and radiotherapy [14, 15]. The risk is higher in elderly patients, and there are no known factors other than age and duration of therapy to predict which patients might be at higher risk of the secondary diseases.

In a review including 28 cases of the secondary diseases, the median latency between the start of therapy and the diagnosis of therapy-related MDS/AML was 2.5 years [15, 16]. Our patient with SCLC developed Castelman disease three years after combined chemotherapy and radiotherapy. However, he was in good condition in the next three years when he manifested symptoms of Hodgkin disease and immediately symptoms of pancytopenia during the first two cycles of the ABVD-based chemotherapy. The development of MDS is related to the specific DNA-damaging agents, dose, therapy duration, and patient age [6, 11, 17]. The mechanism of the secondary disease occurrence related to carboplatin and paclitaxel remains to be determined. It is unlikely that this is a direct effect of carboplatin and paclitaxel only, since our patient has received also radiation thera-
py. It has been reported that the risk of developing secondary MDS and AML depends on the total exposure of chemotherapy and radiotherapy [14, 18, 20]. Hence, patients that are under long-term treatment must be followed-up carefully. This risk of developing secondary disease also depends on the age of the patient which must be considered by clinicians when developing treatment plan for cancer patients.

**CONCLUSION**

A monitoring plan regarding development of secondary leukemia and other malignant hematological proliferations should be outlined in the oncology protocols. Within the context of multi-agent chemotherapy regimens, in our case paclitaxel and carboplatin together with radiotherapy, we underline the importance of following the symptoms during and after the treatment, especially in patients that undergo longer chemotherapy and are exposed to high cumulative dose. To the best of our knowledge Hodgkin lymphoma in a case of SCLC has not been yet reported. Additionally, further studies are also needed to explain and understood the pathogenic relationship between the two entities and to assess the risk of Hodgkin lymphoma in SCLC patients treated with combination of chemotherapy (paclitaxel and carboplatin) and radiotherapy.

**REFERENCES**

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Abbreviations
SCLC - small cell lung cancer
ABVD - adriamycin, bleomycin, vinblastine, dacarbazine
RBC – red blood cell
WBC – white blood cell
HGB - hemoglobin
HCT - hematocrit
PLT - platelets
AST – aspartate aminotransferase
ALT – alanine transaminase
MDS - myelodysplastic syndrome
ALIP - abnormal localization of immature precursors
Резиме

РАЗВОЈ НА СЕКУНДАРЕН ХОЧКИНОВ ЛИМФОМ И МИЕЛОДИСПЛАСТИЧЕН СИНДРОМ (МДС) ПО ТРЕТМАН СО ПАКЛИТАКСЕЛ-КАРБОПЛАТИН КАЈ ПАЦИЕНТ СО СИТНОКЛЕТОЧЕН БЕЛОДРОБЕН КАРЦИНОМ

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Презентираме синхронизирана појава на Хочкинов лимфом и секундарен миелодиспластичен синдром кај 60-годишен пациент со ситноклеточен рак на белите дробови, третиран со комбинација на хемотерапија (карбоплатин и паклитаксел) и радиотерапија. Целта на овој случај е да се нагласи важноста за документација и следење на несаканите реакции што се појавуваат од хемотерапијата.

По четири години од третманот со комбинирана хемотерапија, пациентот презентирал ингвинална лимфаденопатија и зголемени лимфни жлезди со патохистолошки наод сугестивен за плазмацитоидна варијација на Кастелмановата болест. Три години подоцна, со изведба на биопсија на лимфен јазол, воспоставена е дијагноза на Хочкинов лимфом. Со користење на молекуларни анализи потврдено е присуство на доминантна моноклонална популација на имуноглобулинските гени во олиго/моноклонална заднина. Биопсијата на коскената срцевина сугерираше присуство на секундарна миелодисплазија и потврди знаци на незрелост на хематопоетичните клетки со знаци на металобластна матурација на еритопоетичната линија, појава на ALIP (абнормална локализација на незрели прекурсори) во миелодната линија и присуство на диспластични мегакариоцити. Исто така, најдено беше и покачано ниво на поликлонални плазмацити (lambda vs kappa 60% : 40%).

Сметаме дека карбоплатинот и/или паклитакселот придонесуваат за акумулација на алкилаторските оштетувања на ДНК поради предиспозицијата на миелотоксичниот ефект од хемотерапијата. Сметаме дека е неопходно да ја потенцираме потребата за поставување план за мониторинг во однос на потенцијалниот развој на секундарна леукемија и друга малигна хематолошка пролиферација во онколошките протоколи.

Ключни зборови: Хочкинов лимфом, миелодиспластичен синдром, паклитаксел, карбоплатин