A Collision Tumor With Features of Breast Cancer and Plasma Cell Myeloma as Primary Tumors

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Abstract

Collision tumors, consisting of two different histological types of cancer, are rare but may have important implications, both for the patient and possibly in terms of cancer risks and predisposition. This patient has had prior invasive ductal breast cancer treated with surgery and chemotherapy 5 years earlier, but she developed a pathological hip fracture and anemia. The workup uncovered a plasma cell dyscrasia and the bone biopsy from the hip showed the collision tumor with an interface between breast cancer and plasma cell cancer. Such combination tumors may be diagnostic and therapeutic challenges, but they also raise significant questions regarding pathogenesis. Local factors such as neuro-immune crosstalk or cytokine perturbations may be at play. Alternatively, or in addition, oncogenic growth factors or the effects of prior chemotherapy may be significant.

Keywords: Collision tumor; Histopathology; Monoclonal gammopathy; Plasma cell dyscrasia; Lytic/blastic bone lesions; Genetic predisposition

Introduction

Collision tumors are defined by the presence of cancers of different histopathological origin at the same anatomical organ or site. The diagnosis of this very rare type of tumor depends upon histopathology and also needs careful review of morphology, immunohistochemistry staining, and specific cytogenetics. While collision tumors of other histopathologies have been described in multiple locations, including the gastric cardia, uterine cervix, renal pelvis, bladder, liver, lung, oral cavity, thyroid, ovary and bile ducts, this is, to our knowledge, the first reported case of a collision tumor consisting of both breast cancer and multiple myeloma found in bone [1-11]. The pathogenesis of such combination tumors is unclear; they may be simply unfortunate and rare dual afflictions, but may alternatively be due to genetic, cellular, cytokinetic, or other perturbations, local or systemic. The diagnosis and treatment of such dual tumors may be more complex and may require careful planning and timing of multiple anti-cancer modalities to maximize their effects and minimize complications.

Case Report

This 79-year-old woman had a history of right breast invasive ductal carcinoma stage IIB, estrogen receptor (ER) and progesterone receptor (PR) positive, human epidermal growth factor receptor 2 (HER2/neu) negative by FISH/IHC and, was originally treated with modified radical right mastectomy followed by adjuvant chemotherapy and adjuvant aromatase inhibitor. She failed regular follow-up with her oncologist but was living independently and had full functional status prior to this hospitalization. Five years after her original diagnosis, she presented to the emergency department with excruciating right hip pain after sustaining a ground level fall. Computed tomography (CT) of the pelvis revealed a compound fracture of the right femoral neck and a displaced fracture of the left sacral ala suggestive for pathological fracture (Fig. 1). She had a microcytic, hypochromic anemia and elevated total protein and serum globulins on admission (Table 1). She underwent right hip open reduction and internal fixation and a bone biopsy obtained from the area during surgery showed metastatic breast carcinoma which was positive for GATA3 (Fig. 2), strongly positive for ER and PR (Fig. 3) but HER2/neu negative. In addition to metastatic breast carcinoma, the adjacent bone marrow showed an abnormal population of plasma cells positive for CD138 (Fig. 4) and representing approximately 80% of the total cellularity of the marrow. Serum and urine electrophoresis, serum and urine immunofixation, and free light chain assays were done as part of the plasma cell dyscrasia workup. The results revealed an abnormal IgG kappa monoclonal protein elevated to 5,300.0 mg/dL with elevated kappa-lambda light chain ratio (K/L) 2,406.90 (Table 2). Additional immunohistochemical stains were performed and showed negative CD56 and clonality positive for kappa but no staining for lambda light chains. These findings were all consistent with involvement of bone by malignant clonal plasma cells as well. Given this dual diagnosis of metastatic...
breast cancer and multiple myeloma, additional workup was initiated to determine the stages of these diseases. She underwent a Tc 99m nuclear medicine bone scan as a part of staging for breast cancer which revealed increased radiotracer activity in the left frontal skull, the posterior aspect of the left 10th rib at the costovertebral junction, the thoracic (T3) vertebral body, and also left sacroiliac joint. CT of the chest and abdomen revealed enlargement of the pericarinal, subcarinal, and right hilar nodes as well as multiple pleural-based masses in the right lung consistent with metastatic disease. No evidence of metastatic disease was detected in the liver, spleen or adrenal glands. Bone marrow biopsy was deferred given the above results of the surgical specimen.

Treatment

The findings were compatible with a collision tumor of breast cancer (stage IV) and multiple myeloma (stage II by the international staging system) of the right hip (Fig. 2); both were felt to be radiosensitive malignancies. Palliative radiotherapy followed by chemotherapy with an aromatase inhibitor initiated for metastatic breast carcinoma and first cycle of induction chemotherapy with the combination of a proteasome inhibitor, cyclophosphamide and dexamethasone (CyborD) was started for the newly diagnosed multiple myeloma. It was felt that the alkylating agents such as cyclophosphamide would also target the metastatic breast cancer. The patient tolerated the first cycle of chemotherapy well without significant complications and was discharged home on continued anastrozole with plans for a second cycle as an out-patient.

Discussion

Combination tumors of different histology are divided into two pathological groups: collision and composite tumors. Collision tumors have two or more juxtaposed masses and each mass has a distinct boundary, whereas composite tumors have intermingled cell types. Collision of two or even more malignancies can occur between cancers originating in the same organ or with metastases from other sites [1]. Bone is a well-known sanctuary site for metastatic breast cancer and is a primary site for multiple myeloma involvement. Both can certainly result in skeletal-related events such as the pathological fractures experienced by our patient.

The cause of collision tumors such as that in our patient would

Table 1. Laboratory Results on Admission

| CBC | White blood cell count 6.3
| Neutrophils 66
| Lymphocytes 21
| Monocytes 12 (H)
| Red blood cell count 3.09 (L)
| Hemoglobin 8.1 (L)
| Hematocrit 25.6 (L)
| MCV 83.0
| RDW 19.3 (H)
| Platelets 187
| LFT and coagulation factors
| AKP 68
| ALT 10
| AST 53 (H)
| Bilirubin 0.5
| ALB 2.5 (L)
| Protein 9.8 (H)
| Globulin 7.3 (H)
| PT 15.5 (H)
| APTT 28.8
| INR 1.23 (H)
| BMP
| Glucose 100 (H)
| BUN 20
| Creatinine 0.65
| Sodium 130 (L)
| Chloride 100
| Potassium 4.4
| Calcium 9.3
| CO₂ 22
| Tumor marker
| CEA < 1

CBC: complete blood count; MCV: mean corpuscular volume; RDW: red blood cell distribution; AKP: alkaline phosphatase; ALT: alanine amino transferase; AST: aspartate aminotransferase; ALB: albumin; PT: prothrombin time; APTT: activated partial thromboplastin time; INR: international normalized ratio; LFT: liver function test; BMP: basic metabolic panel; CEA: carcinoembryonic antigen; L: low; H: high.
is unclear. Of course, coincidence is a possibility with the unfortunate meeting of two coexisting neoplasms developing independently and finally colliding [12]. This theory does not provide any particular explanation of the colliding pattern and makes no differentiation between composite and collision neoplasms [12]. Other possibilities seem more intriguing.

Our patient was previously treated with systemic chemotherapy and local breast radiation therapy 5 years prior to presenting with her multiple myeloma; thus, an explanation may be that chemotherapeutic medications used to treat a previous cancer may trigger the development of a second primary cancer. A second potential explanation is the “seeds and soil” theory in which cancer cells are the “seeds” and the specific organ microenvironment is the “soil”. Interaction between the “seeds” and the “soil” may determine the formation of a secondary tumor [13]. The presence of metastatic breast cancer and its interaction on cytokines or its elaboration of growth factors may create a fertile microenvironment for second malignancies. Crosstalk between cancer cells is another possible explanation. Cancer cells can influence their microenvironment and bi-directionally communicate with other systems including the immune and nervous systems. While the immune system is well-recognized for its cancer regulatory function, the nervous system may also play a fundamental role in influencing immune responses to a range of disease states including cancer development and progression [14]. Finally, another possible explanation is that genetic predisposition or alteration may increase the likelihood of the development of two different cancers.

Conclusions

Our case is that of an unusual collision tumor with breast can-
Table 2. More Diagnostic Data Suggestive for Plasma Cell Dyscrasia

<table>
<thead>
<tr>
<th>SPEP</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Total protein</td>
<td>10.3 (H)</td>
</tr>
<tr>
<td>Albumin</td>
<td>3.2</td>
</tr>
<tr>
<td>Alpha 1</td>
<td>0.5 (H)</td>
</tr>
<tr>
<td>Alpha 2</td>
<td>0.9</td>
</tr>
<tr>
<td>Beta</td>
<td>0.9</td>
</tr>
<tr>
<td>Gamma</td>
<td>4.7</td>
</tr>
<tr>
<td>B2 microglobulin</td>
<td>3.5</td>
</tr>
<tr>
<td>Free light chain</td>
<td></td>
</tr>
<tr>
<td>Kappa</td>
<td>698.0 (H)</td>
</tr>
<tr>
<td>Lambda</td>
<td>0.29 (L)</td>
</tr>
<tr>
<td>K/L ratio</td>
<td>2,406.90</td>
</tr>
<tr>
<td>Immunofixation</td>
<td></td>
</tr>
<tr>
<td>IgA</td>
<td>&lt; 19</td>
</tr>
<tr>
<td>IgG</td>
<td>5,300 (H)</td>
</tr>
<tr>
<td>IgM</td>
<td>&lt; 8</td>
</tr>
</tbody>
</table>

SPEP: serum protein electrophoresis; IgA: immunoglobulin A; IgG: immunoglobulin G; IgM: immunoglobulin M; Alpha 1: alpha-1 globulin; Alpha 2: alpha-2 globulin; Beta: beta globulin; Gamma: gamma globulin.

Conflict of Interest

The authors declare no conflict of interest in preparing this article.

Disclosures

The above case has been presented as poster presentation at the American College of Physicians Southern California Regional Scientific Meeting in September 2017.

References