Irradiation combined with surgery for function preservation in the treatment of extramedullary plasmacytoma of the left labia majora during pregnancy: A case report

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Received October 11, 2014; Accepted July 28, 2015

DOI: 10.3892/ol.2015.3648

Abstract. Extramedullary plasmacytoma (EMP) in the vulva is extremely rare. The current study presents, for the first time, a case of EMP in the left labia majora in a 36-year-old patient during pregnancy. A painful 7x4x2-cm mass with ulceration, pus and blood scabs, previously misdiagnosed as vulvar ulcer in a local hospital, was proven to be an EMP by biopsy. Upon magnetic resonance imaging, the tumor was shown to occur in the left labia majora without lymphadenopathy. A complete multiple myeloma (MM) workup excluded coexistence with MM. The goal of treatment was to eradicate the tumor while synchronously preserving the vulva. Therefore, following the termination of the pregnancy, radiotherapy with a total dose of 4,500 cGy markedly reduced the size of the tumor. An extended local excision of the residual tumor, and anaplasty of the vulva preserved the appearance and function of the vulva to the utmost. No post-operative radiation was administered, as the resection margins were not microscopically involved. There was no relapse, metastasis or progression to MM in a 9-month post-operative follow-up period, but close follow-up and long-term surveillance are required.

Introduction

Solitary plasmacytoma (SP) is a rare plasma cell dyscrasia that is characterized by the presence of bone or extramedullary monoclonal plasma cell tumors, without evidence of multiple myeloma (MM) (1). According to the location, SP can be categorized into two groups: Solitary plasmacytoma of the bone (SPB) and extramedullary plasmacytoma (EMP) (1). EMP accounts for 3-5% of all plasma cell tumors, with an incidence of only 0.04 cases per 100,000 individuals (2). Approximately 80% of EMPs occur in the upper respiratory tract (3), but they can also occur in numerous other sites (3,4), including the gastrointestinal tract (5,6), brain (7), orbits (8), thyroid gland (9), breasts (10), lungs (11), pleura (12), kidneys (13), bladder (14), urethra (15), ovaries (16), testes (17) and skin (18). Usually, EMPs have no specific clinical manifestations. They typically present as well-localized submucosal masses or swellings in the fifth to seventh decades of the life, with a male to female ratio of 3:1 (3). As the majority of patients can be cured by local radiotherapy, EMP carries an optimistic prognosis, with only 5-20% recurrence (3,4,5,19).

EMP in the vulva is extremely rare (20) and to the best of our knowledge, no case involving the labia majora has previously been reported. The present study describes the first case of EMP occurring in the left labia majora in a young female during early pregnancy. The strengths of the case were the uncommon location and the function preservation treatment strategy. Written informed consent was obtained from the patient for inclusion in the present study.

Case report

In November 2013, a 36-year-old female (gravida 2, para 1) in the seventh week of pregnancy presented to the West China Hospital (Chengdu, China) with a gradually enlarging mass on the left labia majora, accompanied by ulceration and pain that had persisted for 2 months since August 2013. The mass presented as a small nodule ~2 cm in diameter at the onset, with a rough surface and tenderness, which quickly expanded to the whole left labia majora and ulcerated with a purulent exudate. The diagnosis of a vulvar ulcer was made previously in a local hospital in October 2013, therefore, anti-inflammation treatment with intravenous benzylpenicillin (3 million units every 8 h for 10 consecutive days) and debridement were performed. However, the symptoms were not alleviated and the patient was subsequently transferred to the West China Hospital.

Upon admission to the West China Hospital in November 2013, an irregular 7x4x2-cm mass was found in the...
left labia majora, with blood scabs and a purulent exudate. The mass was hard and tender on palpation. There were no palpable inguinal lymph nodes. The patient reported persistent spontaneous pain with a score of 6 in the numerical rating scale (21).

A biopsy was performed and the histopathology results revealed a diffuse infiltration of plasmacytoid cells in the dermis and subcutaneous tissue, a number of which invaded the nerves and vessels (Fig. 1). The cells were positive for cluster of differentiation (CD)79a and κ-light chain, partly positive for CD132, and negative for CD20, λ-light chain, CD23, CD5, B-cell lymphoma (Bcl)-2, Bcl-6 and CD10. Ki-67 staining revealed a high cell proliferation rate with ~50% immunoreactive cells. No monoclonal gene rearrangement of immunoglobulin (Ig)H or IgK was detected. Therefore, a diagnosis of EMP was proposed.

Subsequently, an extensive medical workup was performed to delineate the extent of the lesion and to rule out coexistence with MM. As the patient had decided to terminate the pregnancy, imaging examinations were conducted. An irregular mass with mixed signals and without lymphadenopathy was found on magnetic resonance imaging (MRI) (Fig. 2). No metastases were found on chest and abdominal computed tomography (CT) scans. A normal result with only 0.5% plasma cells was yielded upon bone marrow aspiration and biopsy. No bone lesions were found on bone single-photon emission CT. No anemia, hypercalcemia or renal impairment were detected. Serum electrophoresis did not show the M-band and Bence-Jones proteinuria was not detected. Serum albumin and Ig levels were normal, and serum and urinary β2 microglobulin levels were unremarkable. On the basis of these results, a definitive diagnosis of EMP was established.

Following termination of the pregnancy, pre-operative external irradiation with a 6-MV photon beam though an...
anterior portal was applied to the vulvar area and the inguino-femoral lymph nodes (Fig. 3) using daily fractionation of 200 cGy in five fractions weekly (21 fractions in total), except for the third fraction in which 500 cGy was adopted due to aggravated pain. Dynamic observations of the mass were made every week. No severe moist desquamation or maceration of the perineal skin occurred. The total 4,500-cGy treatment was completed over 32 days, and compared with the initial presentation (Fig. 4A), the tumor was satisfactorily reduced in size (Fig. 4B).

Subsequently, 40 days later, the mass had further decreased in size and only two small nodules existed (Fig. 4C). The mass was reevaluated and the surgical timing was appropriate. An extended local excision and anaplasty of the vulva was performed. As the post-operative pathology confirmed that no tumor cells were present, post-operative boost irradiation was not performed. The post-operative course was uneventful (Fig. 4D) and the patient was advised to attend regular follow-ups every 3 months. At 9 months post-surgery, there was no evidence of local recurrence, distant metastasis or progression to MM. The patient has continued to maintain a satisfactory sexual life.

**Discussion**

EMP of the labia majora is extremely rare. The current study presents, to the best of our knowledge, the first documented case of human EMP involving the labia majora and the first such case presenting during pregnancy. A thorough literature search in Pubmed found only one study reporting a case of simultaneous EMPs of the vagina and right labia minora, with a literature review of four cases invading the vagina and with a mention of three vulvar plasmacytomas (20). However, the present case is reported not only for its uncommon location, but also for its tortuous diagnosis, treatment strategy and close follow-up.

The differential diagnosis should be kept in mind and rare diseases should be considered when diagnosing all masses. Due to the rare occurrence and variety of clinical manifestations of EMP, an accurate diagnosis is often delayed. In the present case, the patient was initially misdiagnosed with a vulvar ulcer in a local hospital. In fact, the main feature of the disease was the progressively enlarging mass, while the ulceration was only an accompanying symptom. When treatment effectiveness cannot be achieved, rare diseases should be considered. Pathological biopsy, a golden criteria of diagnosis, is essential for improving diagnostic accuracy.

An EMP was proposed on the basis of diffuse monoclonal plasma cell infiltration in the biopsy tissue of the present patient. Once EMP is proposed by pathology, a medical workup should be arranged to exclude existence of MM. The recommended diagnostic criteria for EMP are: i) A single extramedullary mass of clonal plasma cells; ii) histologically normal bone marrow aspirate and trephine; iii) normal results on skeletal survey, including radiology of the long bones; iv) no anemia, hypercalcaemia or renal impairment due to plasma cell dyscrasia; and v) an absent or low serum or urinary level of monoclonal Ig (22). In the present case, all relevant tests had been conducted to exclude coexistence with MM.

Due to the small number of patients and lack of randomized controlled trials, there are no established criteria for the treatment of EMP. However, radiotherapy is considered as
the mainstay due to the high radiosensitivity of EMP, particularly in the head and neck area where resection is limited by the anatomical complexity. Several series have reported 80-100% local control rates (4,5,19,23-27). The optimal radiation dose recommended by the United Kingdom Myeloma Forum is in the range of 40-50 Gy (22,25). A dose of 40 Gy in 20 fractions can confer an excellent chance of local control in tumors that are ≤5 cm in diameter, whereas a higher dose of ~50 Gy in 25 fractions is required in tumors >5 cm due to a higher risk of local failure (27,28). The necessity of prophylactic irradiation of local lymph nodes is unclear, as excellent results could be achieved with elective inclusion of the draining lymph nodes (26) and inclusion of them only when clinically involved (27,29). Complete surgical excision can be considered at other sites outside the head and neck area if feasible. A combination of radiotherapy with surgery has been demonstrated to be less invasive and to produce better overall survival (30). The role of adjuvant chemotherapy is inconclusive and additional use of it may bring certain benefits to patients with large masses and high-grade histology (22).

For the present patient, radiotherapy dominated the treatment not only due to its effectiveness, but also due to the advantage of function preservation. If surgery had been adopted first, the destruction of the vulvar appearance or even function may have caused major psychosexual problems in this patient of reproductive age. A moderate dose of 4,500 cGy was considered enough due to its predetermined combination with the subsequent radical surgery. The inguinal lymph node region was prophylactically irradiated for safety. It is noteworthy that the temporary adjustment to 500 cGy irradiation for the third fraction contributed greatly to the pain relief without injury to the urethra. The external location of the tumor made direct observation, without the requirement for CT or MRI, easy. Therefore, the change in tumor size and appearance demonstrated vividly the radiosensitivity of EMP, which has been rarely reported in other cases. Surgical timing was determined to be appropriate ~1 month after radiotherapy to avoid acute edema immediately after radiotherapy and to make best use of the late effect of radiotherapy. Overall, as indicated in the present case, the primary treatment for the majority of patients will be radiotherapy, but surgery may also be required. Close communication between the hematologist, radiotherapist and surgeon is therefore vital for providing optimum care.

EMP has the best prognosis of all plasma cell tumors. The 5-year overall survival rate varies between 31 and 82% (5,25,29,31,32). However, unlike other tumors in which attention is focused more on relapse and distant metastasis during follow-up, EMP requires extra attention with regard to progression to MM, as the majority of mortalities associated with EMP are due to this conversion. Compared with SPB, progression of EMP to MM occurs less frequently. Approximately 60% of patients with SPB develop MM (27,28,33) while EMP progresses to MM in 5-44% of cases (3,5,23,28,29,33,34). Although progression to MM usually occurs within 2 years of the initial diagnosis, conversion has occurred up to 15 years later (3,5,19,23,30). Therefore, the long-term regular evaluation of patients with EMP is strongly recommended. A physical examination, imaging of the primary site, bone marrow aspiration and biopsy, radiographic studies of the skeleton, serum protein electrophoresis, free light chain assays and laboratory tests, including a complete blood count, renal function tests, and analyses of blood calcium, serum albumin and Ig levels, are required during follow-ups.

In conclusion, the present study, for the first time, reports a case of EMP involving the left labia majora in a pregnant woman. The combination of irradiation and surgery leads to excellent tumor control and function preservation. Regular follow-ups should be performed in such patients in case of relapse, metastasis or progression to MM.

References