Introduction

Rhabdomyosarcoma of uterus is an unusual sarcomatous tumor in gynecologic practice; and when seen it is associated as a component of carcinosarcoma; considered now as a classical example of epithelial mesenchymal transition tumor in human.

Oncogenesis

As a distinct tumor, this is recognized long back in 1968 by Enterline and Horn [1]. Later was analyzed in detail a large survey of 112 cases by Enzinger and Shiraki [2]. The oldest patient in their series was 58 years. Parham and Barr [3] succinctly defined it as cellular sarcomatous neoplasm; containing a monomorphous population of primitive cells and showing skeletal muscle differentiation.

Case Presentation

The main complaints of this 70 year old lady was foul smelling vaginal discharge for few months and post menopausal bleeding for a month. Speculum examination revealed a huge polyp occupying the cervix and extending further downwards. On operation, it turned out to be an endometrial polyp arising from the fundus. Histologically the polyp showed classical features of alveolar rhabdomyosarcoma; confirmed immunochemically.

Discussion

Endometrial polyps at this age are unusual; most often are due to Malignant Mixed Mullerian Tumor (MMMT) or rarely due to adenocarcinoma arising from EIN in endometrial polyp. The rarity of this tumor is well illustrated in the IRS study dealing with ARMS; in which only eight
such cases were of genitourinary system [4]. The histological hallmark of this neoplasm of pseudo-alveolar spaces amidst solidly arranged tumor cells; after downgrading its extent from 75% to 50% helped to diagnose this condition better and realistically. Another histological feature like detecting many forms of rhabdomyoblasts amidst nondescript cells arranged solidly as well as in pseudo alveolar spaces helped to diagnose this condition accurately and rule out other possibilities like small cell synovial sarcoma, malignant lymphoma, small blue cell tumors and solid carcinomas of epithelial origin.

After the advent of immunochemistry, the diagnosis can be firmly established by employing myogenic transcription factors; myogenin and Myo-D1. They are transcriptional regulatory proteins expressed early in muscle differentiation [5] and display a nuclear staining pattern. In a recent review of 956 cases, myogenin and MyoD1 showed a sensitivity of 97% and specificity of 90% and 95% respectively [6]. Myogenin is considered technically superior to other markers [7]. Though Ober long ago recognized malignant mesenchymal tumors of uterus as homologous or hetero logos or mixed, the number of case reported as mixed is still rare; particularly collision tumor as described in this patient.

Reported cases of ARMS involving uterus are rare and when compared with the case under discussion differed markedly in presentation. Fukanya [8] reported a case of ARMS of uterine corpus in a 70 year old woman with the tumor replacing the wall of the body of uterus by multiple friable nodules. She developed extensive metastases and died 12 months after surgery. The second such case [9] was in a 21yr old lady; unique for uterine inversion secondary to ARMS. The last such case was seen in a 44 year old African woman [10]; presenting with widespread disseminated tumor and involving all abdominal viscera. It is clear the case details recorded in our patient differed vastly from other reported cases and appears unique. We feel from our experience myogenin immunochemistry in future will help to discover many more cases and to plan for effective therapy in such patients.

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