Primary Squamous Cell Carcinoma of Stomach: A Rare Entity – Case Report and Brief Review of Literature

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ABSTRACT

Very few case reports of pure squamous cell carcinoma (SCC) of stomach are available in the world literature. The exact pathology of this uncommon carcinoma in stomach remains unknown. This is an additional case report of SCC in an elderly female arising in the gastric antrum. She underwent distal gastrectomy, gastrojejunostomy and jejunojejunostomy. The histopathology was reported as SCC of stomach without any adenocarcinomatous component. She had no other source of extra gastric primary SCC. After surgery, the patient was advised adjuvant chemotherapy. This is an additional case of primary SCC of stomach.

Key words: Primary, squamous cell carcinoma, stomach

INTRODUCTION

Primary squamous cell carcinoma (SCC) of stomach is extremely rare.[1] Very few cases of pure, primary SCC of the stomach are reported in world literature. Their number is approximately <100.[2] The exact pathogenesis of primary SCC is not known.[1] Various theories have been suggested to explain the occurrence of SCC in stomach.

CASE REPORT

An 80-year-old postmenopausal female presented with complaint of epigastric pain for 2 weeks. The pain was of burning type. It was not associated with vomiting, fever or jaundice. She also complained of loss of appetite and weight. She gave a history of on and off fever 2 months back, which was diagnosed as typhoid and she received treatment for the same. Per abdominal examination revealed a swelling of the size 4 cm × 5 cm in the right hypochondrium and epigastric region.

Her routine investigations were within normal limit. Ultrasonography examination revealed irregular hypoechoic wall thickening in the duodenum. Computed tomography scan of the abdomen revealed the evidence of concentric thickening of pylorus of the stomach.

Endoscopy showed an ulcer-infiltrative growth in antrum extending onto the anterior wall of first part of duodenum [Figure 1]. It also revealed a normal esophagus [Figure 2]. Intra-operatively the tumor was present in the distal part of stomach. In the rest of the viscera no abnormality was noted.

Following this she underwent distal gastrectomy, gastrojejunostomy and jejunojejunostomy.

Gross examination of the distal gastrectomy specimen revealed an ulceroproliferative growth measuring 8 cm × 5 cm present in the antrum [Figure 3].

Histopathological examination of the distal gastrectomy specimen revealed the lesional component. The tumor comprised of polygonal cells with hyperchromatic nuclei and abundant amount of eosinophilic cytoplasm. Mitotic figures were frequently noted. The tumor cells were arranged in the form of sheets and nests. Individual cell keratinization and small areas of keratin formation is also noted. The tumor was seen infiltrating all the layers of the stomach. Multiple sections did not reveal any glandular component or mucin. Squamous metaplasia foci were also not seen. There was no evidence of Helicobacter, fungal, or parasitic infection [Figures 4 and 5].

The proximal (gastric) and distal (duodenal) resected margins were free from involvement by tumor. The isolated perigastric lymph nodes were reactive in nature.
Her postoperative period was uneventful. Because of her old age and poor performance status, she was managed symptomatically. No other source of extra gastric primary SCC was noted. Adjuvant chemotherapy was advised, but the patient refused further treatment.

**DISCUSSION**

Pure SCC of the stomach is an extremely rare entity. Its worldwide incidence is around 0.04–0.07%. Very few cases are reported in the world literature [Table 1]. These reports show a peak incidence in the sixth decade of life and male predominance.

According to Parks for a diagnosis of primary SCC of the stomach the criteria required are absence of tumor in the cardia, no tumor extension to the esophagus and no evidence of SCC in any other part of the body. The present case fulfilled the above mentioned criteria.

Boswell and Helwing opined that in histopathology of SCC of the stomach, keratinizing cell masses with pearl...
Bar r virus (EBV) infection, in the pathogenesis of SCC of the stomach. [8]

One theory also suggests damage and squamous differentiation of preexisting glandular epithelium secondary to chronic mucosal ectopic squamous cell nests, squamous metaplasia of the presence of totipotential (stem) cells, an area of cell arrangement with sharp borders, intercellular bridges and a high concentration of sulfhydryl or disulfide groups, indicating the presence of keratin, are characteristics of SCC of the stomach. [8]

The exact pathogenesis of primary SCC is not known. [1] Several theories have been proposed regarding the origin of pure SCC of the stomach. [3] They include the presence of totipotential (stem) cells, an area of ectopic squamous cell nests, squamous metaplasia of glandular epithelium secondary to chronic mucosal damage and squamous differentiation of preexisting adenocarcinoma. [3,5,6] One theory also suggests Epstein-Barr virus (EBV) infection, in the pathogenesis of SCC arising in the stomach. [1]

In the present case there were no foci of squamous metaplasia in the adjacent gastric mucosa. We have not done studies to detect the presence of EBV in our case.

Mori et al. in their study reexamined histopathological sections of three primary SCC of the stomach. They noted revealed minute areas of adenocarcinoma in addition to large areas of SCC. Hence, they opined that some gastric SCCs may be adenosquamous carcinomas with a predominant squamous component. [9]

In the present case, multiple sections did not reveal presence of any adenocarcinomatous component or mucin.

Due to the advanced stage at the time of diagnosis in most of these cases, the prognosis is generally poor. [14,6]

Schmidt et al., have described that adjuvant radio and chemotherapy have resulted in survival rates of more than 3 years in a patient of SCC of stomach. [4]

In one case reported by Yildirim et al., complete remission was achieved with a combination of cisplatin and 5-fluorouracil, in spite of presence of extensive disease. [3]

Though our patient was advised further adjuvant chemotherapy, she refused the same and was subsequently lost to follow-up.

This adds to the existing literature regarding SCC of stomach.

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REFERENCES


Table 1: Primary SCC of stomach reported in literature

<table>
<thead>
<tr>
<th>Authors, year</th>
<th>Age/sex</th>
<th>Location</th>
<th>Morphology</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Muto et al., [6] 1999</td>
<td>75/male</td>
<td>Stomach</td>
<td>Squamous</td>
<td>Surgery</td>
</tr>
<tr>
<td>Schmidt et al., [8] 2001</td>
<td>83/male</td>
<td>Antrum of stomach</td>
<td>7 cm×5 cm, squamous</td>
<td>Surgery+polychemotherapy</td>
</tr>
<tr>
<td>Callacondo et al., [9] 2009</td>
<td>62/male</td>
<td>Antrum of stomach</td>
<td>15 cm×9 cm×5 cm, squamous</td>
<td>Surgery+combination chemotherapy</td>
</tr>
<tr>
<td>Yildirim et al., [10] 2005</td>
<td>80/female</td>
<td>Antrum of stomach</td>
<td>8 cm×5 cm, squamous</td>
<td>Surgery</td>
</tr>
</tbody>
</table>

SCC: Squamous cell carcinoma

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