Case Report

Management of Para-Aortic Lymph Node Metastasis in Rectal Squamous Carcinoma

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Background

Rectal squamous cell carcinoma (SCC) of the rectum is a rare malignancy of the gastrointestinal (GI) tract without a clarified pathogenesis mechanism and a rather aggressive clinical behaviour [1]. The incidence of this malignancy has been reported to be 0.25 to 1 per 1000 colorectal carcinomas [2]. The SCC of the rectum mainly affects individuals between the age of 39 and 93 years and is more frequently found in women [3].

The SCC tumors are associated with a worse overall 5-years survival than adenocarcinomas. Interestingly, squamous cell carcinomas presenting with more advanced stage disease showed a similar survival rate to adenocarcinomas of the same stage [4]. Retroperitoneal nodal metastasis (RNM) represent 1-2% of metastasis from colorectal cancer (CRC) and radical retroperitoneal lymphadenectomy (RRL) in this setting remains controversial; however, there is no mention in the literature on how to treat SCC retroperitoneal lymph node metastasis.

Case Presentation: The patient was a 47-years-old woman with SCC of the rectum. Concomitant chemoradiotherapy (CRT) followed by surgery was planned. After 8 weeks, CT of the abdomen for restaging showed a 6 cm para-aortic mass involving and obstructing the medium third of the left ureter. The patient was submitted to abdominoperineal rectal resection with left nephrectomy and en-bloc resection of para-aortic tumor (nodal bulking).

Conclusion: After a thorough review of the literature regarding RNM for rectal SCC, we found no mention of how to treat RNM for rectal SCC. The purpose of this paper is to report a very rare case of RNM for rectal SCC and to discuss the surgical approach. Surgery seems to be the best option for local control and improvement in overall survival (OS).
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The patient was submitted 14 weeks after neoadjuvant CRT to abdominoperineal rectal resection with left nephrectomy (renal exclusion), en-bloc resection of the para-aortic tumor and retroperitoneal lymphadenectomy (Figure 3). Histopathological examination showed a complete pathological response in the rectum, and squamous carcinoma metastasis in the para-aortic mass (Figure 4), with R0 resection. The patient had no postoperative morbidity, with complete relief of symptoms, and was discharged from hospital 6 days after surgery.

Discussion

SCC is one of the rarest forms of rectal cancer. Pure SCC is a very rare histological diagnosis and clinical occurrence [6]. Furthermore, many supposed rectal SCC may be an extension of anal or gynecological carcinoma. For that reason, Williams et al. established guidelines before a definitive diagnosis of SCC can be made [7]. These criteria include:

- Exclusion of primary SCC from distant sources;
- Exclusion of a squamous lined fistula from bowel tumor;
- Tumor must originate from the rectum and not be an extension of anal SCC;
- Confirmation of SCC by histology.

To fulfill the above criteria, we decided to proceed with magnetic resonance, which confirmed its origin from the rectal wall with enlarged perirectal lymph node and no local spread to adjacent pelvic organs. Due to its rarity, the natural history and progression of SCC are not well established.

The treatment of rectal SCC has traditionally involved surgery, in some cases preceded or followed by adjuvant radiotherapy or chemotherapy. The definitive therapy is surgical resection of the affected rectum, either by low anterior resection or by abdominoperineal resection, depending on the location of the tumor [8]. However, in the last decade, there has been increasing interest in the response of rectal SCC to definitive chemoradiotherapy, with very encouraging results [9]. Clinical examination is not a reliable predictor of tumor response after treatment. Our decision on surgical rescue was based on the persistence of symptoms and significant rectal thickening 12 weeks after the end of CRT.

Several studies for colorectal cancer have suggested that surgery for retroperitoneal nodal metastasis is the only potentially curative treatment and can achieve long-term overall survival in selected patients with acceptable postoperative morbidity. Median OS in operated patients was 34-46 months, compared with 3-33 months in those who did not have surgery, and five-year OS was 15-53% versus 0-12% respectively [10].
Conclusion

In summary, we described a very rare case of RNM for SCC rectal cancer, and the surgical approach seems to be the best option for local control and improvement in OS. Further research, including prospective trials and large observational studies, is needed to better define the role of surgery in the management of these patients.

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Author Contributions

Cecília Araújo Carneiro Lima and Mário Rino Martins drafted the manuscript; Juliana Karine Ferreira Santos Lessa provided the original pictures; and Rogério Luiz dos Santos and Luciana Mata da Silva reviewed the manuscript. All authors read and approved the final manuscript.

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Availability of Data and Materials

The datasets supporting the conclusions of this article are included within the article and its additional files.

Ethics Approval and Consent to Participate

Not applicable.

Consent

Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the editor-in-chief of this journal.

Competing Interests

None.

Abbreviations

SCC: Squamous Cell Carcinoma

RNM: Retroperitoneal Nodal Metastasis
CRC: Colorectal Cancer
RRL: Radical Retroperitoneal Lymphadenectomy
GI: Gastrointestinal
HPE: Histopathological Examination
CT: Computed Tomography
CRT: Chemo-Radiotherapy

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