Synchronous caecal small-cell neuroendocrine carcinoma and adenocarcinoma of the rectum

Sinkroni neuroendokrini karcinom malih stanica cekuma i adenokarcinom rektuma

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Summary

Small-cell neuroendocrine colon carcinoma is a rare entity with a usually poor prognosis. An 80-year-old female had colon cancer surgery due to synchronous tumour of the rectum and caecum. Pathohistological analysis of the caecal tumour showed trabecular and solid clusters, relatively uniformed small to middle sized epithelial cells, deficient cytoplasm and there were a great number of mitosis with larger areas of necrosis in the connective tissue. The immunohistochemistry was positive for chromogranin A. The caecal tumour was diagnosed as a small-cell neuroendocrine carcinoma. In addition, the rectal tumour showed microscopic findings consistent with stage IIA adenocarcinoma. The immunohistochemical panel showed that the tumour was negative for neuroendocrine markers. There were no clinical findings suggestive of hormone hypersecretion. Cancer metastases were not found. Postoperative chemotherapy was applied. The patient is still alive, in good general condition and with no signs of tumour progression.

Key words: neuroendocrine, small-cell carcinoma, caecum, immunohistochemistry

Sažetak


Ključne riječi: neuroendokrini, karcinom malih stanica, cekum, immunohistokemiija

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Introduction

Neuroendocrine carcinomas are malignant tumours that show histopathological and immunohistochemical evidence of neuroendocrine differentiation. Neuroendocrine tumours can develop at many different sites of the body, but colon is a very rare site.¹ Most neuroendocrine tumours are carcinoids and they have a better prognosis than conventional adenocarcinomas. The original term “carcinoid” equates to neuroendocrine tumours that are both grade 1 and 2 (well differentiated), whereas grade 3 neuroendocrine tumours are mostly small-cell neuroendocrine

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carcinomas (SCNEC) with a minority component being large-cell type (poorly differentiated). Small-cell neoplasia most commonly originates in the lung but also other sites such as skin, thymus, kidney, breast, ovary, uterus, urinary bladder, hepatobiliary tree, pancreas and salivary glands. More specifically, the colorectal SCNEC accounts for 0.2% to 1.5% of all colonic cancers and is considered to be an aggressive neoplasm with rapid growth and early spread carrying poor prognosis.

**Case report**

An 80-year-old female was admitted with leading symptom of abdominal discomfort. Patient's past medical history was unremarkable. In the last four months the patient is periodically experiencing upper abdominal pain and loss of appetite, denying nausea and vomiting. In the same period of time she lost 20 kg of weight. In the last four months patient's stool is irregular (absent from time to time in periods of 8 days) with occasional diarrhea without tinge of blood or melena. The laboratory results verify microcytic anemia.

Colonoscopy has showed a tumorous mass in the mid third of rectum. Irigography has verified a lack in filling measured 40 mm in diametar, in the area of rectosigmoid crossing, which is considerably reducing the lumen of the bowel and where the barium enema isn't passing through. Computed tomography (CT) of the abdomen and pelvis has shown a thickening of the bowel wall which was highly suspect of an infiltration of the rectal ampulle on the left side just above the anocutaneous border, and just above the same rectosigmoid area near the back bowel wall has shown a round tumorous mass 5 cm in diameter (Picture 1). The wall of the caecum was slightly thicker (Picture 2). The pelvic CT showed no enlarged lymphatic nodes.

On the third day of patient's hospitalization, based upon diagnostic criteria, we decide to perform a medial laparotomy which has shown a smaller tumour measured 5 cm in diameter in the mid third of rectum. Right hemicolecotomy with the resection of infiltrated ileum was performed and the Hartmann procedure in the rectosigmoidal area. The recovery was unremarkable. The patient was discharged 12 days after the surgical procedure.
The macroscopic examination has shown, in the area of the Bauchin's valve, caecum and ascendant colon on a length of 8.5 cm, a yellowish white tumorous tissue that engages the whole circumference of the bowel (Picture 3). This tumour involved the pericolic adipose tissue. The rectum margin was free from tumour. Metastasis was found in neither of eleven dissected regional lymph nodes. The histological examination has shown that the tissue in question was a cluster of trabecular and solid, relatively uniformed small to middle sized epithelial cells with deficient cytoplasm and in the connective tissue were a lot of mitosis (up to 100 per 10 high power field) with larger areas of necrosis. An immunohistochemical panel was performed, and the tumour was found to be positive for chromogranin A. The synaptophysin and CD56 was negative. Therefore, our diagnosis of the tumour was a giant small-cell neuroendocrine carcinoma (Picture 4).

The macroscopic examination of the second sample, a part of rectosigmoid colon, has shown an exophytic tumour sized 4.5 x 2 cm, which engages the whole circumference of the bowel (Picture 5). All sampled lymph nodes and surgical ends were free of tumour involvement. The tumour is histologically made of atypical glandular formations coated with malignant epithelium partly making a cribriform form. The tumorous tissue infiltrates the whole thickness of the bowel wall and spreading to the under laying fat tissue. Diagnosis was stage IIA adenocarcinoma. On the immunohistochemical staining panel, the tumour was negative for neuroendocrine markers (chromogranin A, synaptophysin and CD56). Thus, we concluded that the patient had synchronous a primary adenocarcinoma of the rectum and a primary giant SCNEC in the caecum. The patient received postoperative chemotherapy with 5-fluorouracil. Five cycles of chemotherapy were performed. The patient is still alive and with no sign of tumour progression or distant metastasis 6 month after the treatment. In the follow up of patient we determined carcinoembryonic antigen (CEA), performed liver ultrasound and control thorax-abdomen-pelvis CT.

Picture 3 Giant yellowish white neuroendocrine carcinoma in the caecum (arrow)
*Slika 3. Veliki žuto-bijeli neuroendokrini karcinom u cekumu (strelica)*

Picture 4 Microscopic finding. Immunohistochemical staining revealed positive findings for chromogranin A.
*Slika 4. Mikroskopski nalaz. Imunohistokemijsko bojenje pokazalo je pozitivan nalaz na kromogranin A.*
Discussion

This case has two aspects of clinical interest. First, synchronous adenocarcinoma and SCNEC is the rare appearance in clinical practice. Second, there are no clear guidelines for appropriate treatment and follow up of SCNEC with a synchronous secondary malignancy; therefore, our experience may help in the consideration of treatment for similar patients.

SCNEC are rare in the large bowel, but more common in this location than elsewhere in the intestines. Neuroendocrine carcinomas including SCNECs comprise approximately 0.6% of all carcinomas of the large bowel. The average age of patients with neuroendocrine carcinomas of the large bowel is 61.5 years and the male-to-female ratio is essentially 1 : 1, similar to the values for large-bowel adenocarcinomas. Neuroendocrine carcinomas of the colorectum are grossly similar to conventional adenocarcinomas. If an associated adenoma is present, they may be polypoid; if not, they are typically ulcerated neoplasms with a raised border, and on cut section they are infiltrative. SCNEC is a neuroendocrine tumour that has features in common with neuroendocrine-differentiated tumours as well as specific cytological features: small cells with minimal cytoplasm and fusiform nuclei and have very high mitotic rate, averaging 65 per 10 high power field. Small cell carcinomas typically express chromogranin A, synaptophysin and CD56. In the report by Vilallonga et al., on neuroendocrine tumours, excluding carcinoid tumours, most patients presented with a paraneoplastic or carcinoid syndrome. However, our patient had no such symptoms. However, we cannot rule out the possibility of secreted hormones in the plasma at very low concentrations not associated with a clinical syndrome, or if there were secreted peptides they were not associated with clinical effects. The precise pathogenesis of secondary cancers with neuroendocrine carcinoma remains unclear. Some reports have suggested that neuroendocrine tumours occur as secondary primary carcinoma. According to one previous study, gastrin and cholecystokinin were associated with neuroendocrine carcinoma; this resulted in tissue growth in the gastrointestinal tract and carcinogenesis leading to colorectal and gastric cancer. In addition, Prommeger et al. reported that neuroendocrine tumours are associated with a high risk of secondary gastrointestinal malignancy. They studied 96 patients with neuroendocrine tumours and found that 14 patients had a neuroendocrine tumours...
and a second primary malignancy. Five patients had a synchronous second primary malignancy. The locations were two colon cancers with one double colon cancer, one gastric cancer, one bladder cancer, and one ovarian cancer. In recent study Kato et al. suggested a possible link between colorectal neuroendocrine carcinoma and adenocarcinoma. These reports are consistent with the findings in our case where the patient had an advanced stage of neuroendocrine carcinoma in the caecum and a synchronous adenocarcinoma in the rectum. It is possible that the neuroendocrine caecum was associated with the development of the adenocarcinoma of the rectum. However, further confirmation is needed to support this.

For colonic neuroendocrine tumours (excluding appendical tumours), 5-year survival rates are 76.0% for localized disease, 71.6% for regional disease, and 30.0% for tumours with distant metastases. Overall, the 5-year survival rate is 61.8%. If the neuroendocrine carcinoma is completely resected, the 5-year survival rate has been reported to be 61%. However, most patients are not candidates for curative resection.

After completely resection both of the tumours, our patient received systemic chemotherapy with 5-fluorouracil. 5-fluorouracil is the chemotherapeutic agent of first choice in the treatment of patients with colorectal cancer. Five cycles of systemic chemotherapy were provided. The patient is still alive with good performance status and no sign of tumour progression.

In conclusion, we report a SCNEC of the caecum that coexisted with an adenocarcinoma, suggesting possible association between colorectal neuroendocrine carcinoma and adenocarcinoma. Further investigations are required to determine the pathogenesis of these synchronous tumours.

References