Case Report

Synchronous colonic malignancy

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ABSTRACT

Synchronous colorectal neoplasias, defined as 2 or more primary tumors identified in the same patient and at the same time, are caused by common genetic and environmental factors. Since intraoperative palpation can miss up to 69% of the SN, currently, synchronous neoplastic lesions are usually diagnosed at a preoperative staging by colonoscopy or virtual colonoscopy; according to data from literature, 3% of the patients with SN are affected by different types of malignant lesions while 33-55% shows villous adenomas. Literature also confirms the presence of primitive synchronous cancers; malignant synchronous lesions are very rare, showing the following incidence: between 0.17% and 0.69% in case of 2-3 synchronous lesions, 0.19% in case of 4-5 synchronous lesions. The most voluminous synchronous cancer is called "first primitive" or "index" cancer. When the index cancer is located in the caecum, the incidence of left colon synchronous cancers is higher than when the index cancer is located in the left colon. Colorectal adenomas standard treatment is usually represented by endoscopic polypectomy; indeed only 5% of synchronous colorectal lesions require a surgical treatment.

Keywords: Abdominal discomfort, Caecum and descending colon, Bowel thickening, Adenocarcinoma, Chemotherapy

INTRODUCTION

Colorectal cancer is the third most common malignancy worldwide and the fourth most common in the United States, with estimated 146,970 new cases diagnosed in 2009. In the United States, approximately 49,920 cancer-related deaths were attributed to a colorectal malignancy in 2009, making it second only to mortality from lung cancer.3

Despite the high incidence of colorectal carcinoma that is diagnosed yearly, the majority of lesions are resected with curative intent (70%-80%) and colorectal cancer-related deaths account for 20%-30% of those diagnosed and treated surgically, making it a highly curable malignancy if identified in its early-stages. Approximately, 55% of colorectal cancers diagnosed on screening are found to present with Stage I or II disease.2,3 Patients with Stage I or Stage II colon cancer have a greater than 60% 5-year survival, and for patients with Stage I or Stage II rectal cancer, there is a greater than 50% 5-year survival.4,5

Because of this, colorectal screening exams in asymptomatic patients have become the recommended standard of care for average risk patients at the age of 50 and at the age of 40 or younger for those in the moderate and high risk groups.

Approximately, 70% of colorectal lesions occur distal to the splenic flexure, and looking at colon cancer alone, approximately 25% are found in the sigmoid, 10% at the rectosigmoid junction, and 4%-6% located in the descending colon.5 Anatomically, the left colon has a
smaller diameter than the right, and as a result, left-sided carcinomas can cause varying degrees of intraluminal occlusion and patients more frequently present with obstructive symptoms. Large bowel obstructions present a challenging clinical scenario for the physician in the diagnosis, operative management, and the timing of colonic surveillance. Patients presenting with advanced lesions causing partial or high-grade large bowel obstruction commonly have a distant history or no history of previous colonic surveillance.

This fact is of clinical importance because the diagnosis of a primary colorectal malignancy is accompanied by an overall incidence of synchronous colorectal cancers between 2% and 10% and a frequency of synchronous adenomatous polyps ranging from 15% to 50%. Interestingly, when compared by stage, no statistically significant difference has been found in the overall and disease-free survival between synchronous and primary cancers.

**CASE REPORT**

47 years old male residing at Chandur Bazar farmer by occupation was presented with chief complaints of vague abdominal discomfort since 12 months; incomplete bowel evacuation since 12 months; decreased appetite since 12 months; and loss of weight since 12 months.

There was no history of vomiting, hematemesis, malena, bleeding per rectum, fever, breathlessness, hemoptysis, chest pain, giddiness, headache, backache or urinary complaints. With the past history of pulmonary tuberculosis for which CAT 1 AKT was taken 15 years ago. General conditions was satisfactory as afebrile, with pulse rate of 96/min; blood pressure 130/80 mm Hg and paleness was present, and no signs of lymphadenopathy, icterus, oedema, and signs of dehydration.

**Laboratory investigations**

Hb- 11.2 gm%; TLC- 13800 cmm; Platelets- 193000 cmm; ESR-14 mm; LFT, KFT- within normal limits; Mauntoux test – negative.

**Colonoscopy**

Colonoscopy could not be passed beyond distal part of descending colon s/o (?) extrinsic mass on descending colon.

**Surgical procedure**

Under spinal plus epidural anesthesia midline incision taken and peritoneum opened; evidence of grossly distended plastered large bowel with strictures near caecum and near splenic flexure of colon; No evidence of growth on liver surface; No evidence of LN involvement; Caecum, ascending colon, tranverse colon, descending colon resected and ileo-sigmoid Anastomosis done; Resected segment sent for HPE; Two ROMO ADK drains (no.32) were kept; Right drain in pelvic region and left drain in left paracolic gutter.

**Post-operative orders (for first 24 hours)**

Patient was kept nil by mouth and i.v. fluids-2 RL, 2 DNS, 1 NS; Injection meropenam 1 gm iv 8 hourly; Injection amikacin 500 mg iv 12 hourly; Injection metro 100 ml iv 8 hourly; Injection pantop 40 mg iv OD; Injection tramadol 1cc iv 8 hourly; Injection ondem 4mg iv sos, if vomiting.

**On post op day 1**

Rt drain 120 ml (serous); Lt drain 100 ml (serous); 1 PCV given. Epidural catheter was removed on day 3, bowel sounds were returned on day 4; liquids started on day 5; soft diet on day 7; right drain removed on day 7 and left drain removed on day 13.

**Histopathology report**

s/o well differentiated adenocarcinoma of caecum and descending colon with infiltration of ¾ th thickness of muscle wall; surgical margins, mesocolic tissue and mesocolic LN are free from tumor cells. Suture removal done and patient discharged on post op day 18; As per HPE report patient was referred to oncologist for chemotherapy. 6 cycles of chemotherapy was given for
every 21 days. Patient was advised to follow up after completion of chemotherapy schedule. CECT abdo/pelvis done s/o no locoregional residual tumour, no evidence of distal metastasis or lymphadenopathy.

CONCLUSION

With the development of early diagnostic technologies, more SCRCs have been diagnosed. Compared with the solitary CRC, SCRCs possess distinctive features. These should be fully investigated in preoperative clinical workups to avoid the possibility of accessory lesions being overlooked and not resected. Older male patients with concurrent adenomas in the colon are at a particularly high risk of SCRCs and close attention should be paid to them.

The extent of surgical resection relies on the location and the number of lesions. The resected region should include enough intestinal length and regional lymph nodes to capture any local spread, thereby guarding against recurrence or subsequent malignant change. Of course, the mechanisms regarding the formation of multicentricity need to be fully elucidated in order to gain insight into SCRC genesis and development, which will ultimately benefit molecular therapies and improve patient survival.

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REFERENCES


Management

- The choice of type of the operative procedure for synchronous colorectal carcinomas remains controversial.
- Some authors proposed radical operations such as subtotal colectomy to remove enough length of intestines and numbers of local lymph nodes to prevent future development of metachronous tumors.
- Some experts recommended a more conservative approach for older patients and radical procedures for younger patients with regionally confined non-metastatic disease.
- We agree with the second opinion because major surgery such as subtotal colectomy could increase morbidity in older patients.(17% in patients with age >70 years compared to 3.4% in patients with age <50 years)

Prognosis

- In many studies, there was no difference in survival between the patients with primary multiple and single colorectal cancers and there were only minor differences in stage distribution.
- Postoperative survival in synchronous tumor cases was not worse than that in single tumor cases, and was mainly dependent on the pathological stage and curability of the index lesions.

Follow up

As total or subtotal colectomy performed for these patients, therefore the follow up will be by tumour markers, CT scan, PET scan and MRI.

Figure 2: Barium enema showing apple core appearance near caecum and near splenic flexure of colon.