A Case of More Abundant and Dysplastic Adenomas in the Interposed Colon than in the Native Colon

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INTRODUCTION

Colon graft has been a favored surgical technique for benign esophageal stricture and esophageal malignancy. Early post-operative complications of this procedure are common and include anastomosis leaks, fistula formation, colon graft edema, ischemia, and stricture. Late complications include peptic ulcer, fistula formation, reflux colitis, atonia, outlet obstruction in stomach, and rarely malignancy. We report a rare case of colonic adenoma with carcinomatous transformation arising in the interposed colon.

CASE REPORT

A 60-year-old woman, who had undergone esophagectomy with colon interposition, due to esophageal lye-structure 40 years ago, was admitted to hospital with hematemesis and heartburn sensation. Laboratory evaluation including hematological and biochemistry studies were unremarkable except for mild anemia. Endoscopic examination of the colonic graft could not find the focus of the bleeding but six polyps of 3 to 5 mm in size were found incidentally. Colonoscopy of the native colon revealed two polyps of similar size, one in the ascending colon, and the other in the cecum. Consequently, endoscopic and colonoscopic polypectomy was carried out. Histologically, all resected specimens were adenomatous polyps and only two polyps from interposed colon showed high grade dysplasia, and others showed no dysplasia.

8 months later, the patient was admitted due to melena. Endoscopic examination of the colon graft revealed a 1.0 cm sized, active staged ulceration at cologastric anastomosis. In addition, two flat to slightly elevated lesions of 12 and 20 mm in size which might have been missed at the previous examination, were noted at the anastomosis site and 4 cm proximal to the anastomosis site (Fig. 1 A, B). As there was no evidence of active bleeding, endoscopic biopsies were performed in ulcer margin while endoscopic mucosal resection of colonic polyps were carried out (Fig. 1C, D). Pathology of the resented 20 mm sized adenoma disclosed tubulo-villous adenoma with high grade dysplasia and focus of intramucosal well-differen-
tiated adenocarcinoma (Fig. 2A, B). We performed the test for microsatellite instability (MSI)-testing of the lesion. The DNA from the tumor was amplified by polymerase-chain-reaction with the microsatellite markers and genotyped after fluorescence labeling, and the result of MSI-testing was stable (Fig. 3). The patient discharged without further bleeding episode and was treated with H₂ receptor antagonist.

DISCUSSION

Colonic interposition for esophageal reconstruction was described first in 1911. This operation has been generally performed for malignant disease of esophagus and cardia, but also has been largely applied to the benign esophageal disease, such as lye-induced strictures. Because the patients who underwent colonic interposition for benign disease may have normal life-expectancy, late morbidity associated with this operation is particularly important. Malignancy arising in interposed colon is a rare complication of colonic interposition.

In general, the etiology of colon cancer is multifactorial comprising environmental factors, especially diet, and genetic factors. The diet with high animal fat and low fiber has been reported to be associated with carcinomatous transformation of colon. The currently accepted hypothesis regarding animal fat is that the ingestion of animal fat leads to an increased proportion of anaerobes in the gut microflora, resulting in the conversion of normal bile acids into carcinogen. The theory related to dietary fiber is that the fiber accelerates intestinal transit time, reducing the...
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exposure time of colonic mucosa to potential carcinogen and diluting these carcinogens. These studies have been performed in the view of indirect effect of diet.

However, interposed colonic segment would have a marked change of location and functional demand. Such changes include direct exposure of colonic mucosa to environmental substances with potential toxicity or carcinogenicity to colonic mucosa, and to different microflora. The colonic mucosa encounters these factors right after the change of location, and also affected by gastric juice and other physiologic factors involved in alimentary tract function. Altorjay et al.\(^6\) reported that the muscle contraction of interposed colon was segmental without spreading waves on manometric examination. The passage time of food contents in grafted colon is longer than in normal esophagus, and the grafted colon may well be more vulnerable to the refluxates from stomach due to absence of sphincter. Therefore, the duration of exposure to noxious agent and carcinogen is longer in the grafted colon than the normal esophagus.\(^7\) Interestingly, Lindahl et al.\(^8\) reported that interposed colonic mucosa tended to undergo frequent premalignant transformation such as fundic type gastric metaplasia of interposed colon, colonic dysplasia, and increased aneuploid cell population, in a study with histologic and flow cytometric examination of interposed colonic mucosa.

To date, 10 cases\(^6,9,17\) of primary adenocarcinoma arising from interposed colon have been reported worldwide to our knowledge, including the report of Fritscher-Ravens et al.\(^12\) in which the tumor in the interposed colon was diagnosed 7 months after esophagectomy, probably developing from undiagnosed polyps in the transposed colon (Table 1). In the current case, synchronous adenomas were found in the native colon, as well. However, number of adenomas found in the interposed colon was larger than that of the native colon, and carcinomatous transformation was found only in the interposed colon. Of course, the left colon, which is mostly used to replace the esophagus is a common site of cancer over the lifespan. Even in the unoperated patient the finding of dysplasia and cancer is more common in the left colon than the right colon. Adenocarcinoma in the interposed colon may only be the result of the natural history of the dysplasia-carcinoma sequence outside the native anatomical location. However, we confirmed that she had undergone transverse colectomy on colonoscopic

Table 1. Review of Literatures of the Cases of Primary Adenocarcinoma Arising in Interposed Colon

| Name          | Age | Sex | Reason for esophagectomy | Time since graft (yrs) | Detail                                      |
|---------------|-----|-----|--------------------------|------------------------|---------------------------------------------|
| Goldsmith et al.,\(^9\) 1968 | 48  | F   | Esophageal cancer (scc)  | 2                      | Adenocarcinoma invading muscle              |
| Licata et al.,\(^10\) 1978 | 51  | M   | Esophageal stricture     | 11                     | Metastatic adenocarcinoma                   |
| Haerr et al.,\(^11\) 1987 | 72  | M   | Esophageal cancer (scc)  | 7                      | Colonic adenocarcinoma, unsectable         |
| Houghton et al.,\(^12\) 1989 | 64  | M   | Esophageal stricture     | 20                     | Dukes A carcinoma arising in a colonic adenoma |
| Theile et al.,\(^13\) 1992 | 55  | M   | Carcinoma at GE junction| 12                     | Colonic adenocarcinoma, T3N0M0              |
| Lee et al.,\(^14\) 1994 | 75  | F   | Postcricoid carcinoma(scc)| 20                     | Dukes A carcinoma                           |
| Altorjay et al.,\(^6\) 1995 | 70  | M   | Grave esophagitis       | 5                      | Colonic adenocarcinoma                      |
| Fritscher-Ravens et al.,\(^15\) 1999 | 62  | F   | Esophageal carcinoma (scc)| 0.6                    | Tubulovillous adenoma with adenocarcinoma   |
| Goyal et al.,\(^16\) 2000 | 78  | M   | Carcinoma of gastric cardia| 8                      | Colonic adenocarcinoma, T3N0M0              |
| Liau et al.,\(^17\) 2004 | 78  | M   | Esophageal cancer        | 30                     | Colonic adenocarcinoma                      |

scc, squamous cell carcinoma; GE, gastro-esophageal.
Microsatellite instability is a well-recognized feature of non-sporadic colorectal cancer, especially in the hereditary non-polyposis colorectal cancer. However, about 10% of sporadic colorectal cancers have microsatellite instability.\textsuperscript{18,19} We were concerned about the result of microsatellite instability in the sporadic colorectal tumors under mechanical stress and carcinogens. Our MSI-testing showed the absence of instability further emphasizing the importance of environmental exposures in this case.

We anticipate the average span of patient’s life with interposed colon would be long enough in whom esophageal colon graft has been performed for benign diseases of esophagus and stomach. The early and regular screening of the interposed colonic graft to detect and treat malignancy may be demanded because malignancy arising from interposed colon is asymptomatic, but fatal, and re-operation is difficult.

REFERENCES

1. Agha FP, Orringer MB. Colonic interposition: radiographic evaluation. AJR Am J Roentgenol 1984;142:703-8.
2. Larson TC 3rd, Shuman LS, Libshitz HI, McMurtrey MJ. Complications of colonic interposition. Cancer 1985;56:681-90.
3. Lundell L, Olbe L. Colonic interposition for reconstruction after resection of cancer in the esophagus and gastroesophageal junction. Eur J Surg 1991;157:189-92.
4. Briel JW, Tamhankar AP, Hagen JA, DeMeester SR, Johansson J, Choustoulakis E, et al. Prevalence and risk factors for ischemia, leak, and stricture of esophageal anastomosis: gastric pull-up versus colon interposition. J Am Coll Surg 2004;198:536-42.
5. Neagoe A, Molnar AM, Acalovschi M, Seicean A, Serban A. Risk factors for colorectal cancer: an epidemiologic descriptive study of a series of 333 patients. Rom J Gastroenterol 2004;13:187-93.
6. Altorjay A, Kiss J, Vörös A, Szanto I, Bohák A. Malignant tumor developed in colon-esophagus. Hepatogastroenterology 1995;42:797-9.
7. Bucknell A, Clark C. An experimental method for recording the behaviour of human isolated colonic segments. Gut 1967;8:569-73.
8. Lindahl H, Rintala R, Sariola H, Louhimori L. Long-term endoscopic and flow cytometric follow-up of colon interposition. J Pediatr Surg 1992;27:859-61.
9. Goldsmith HS, Beattie EJ Jr. Malignant villous tumor in a colon bypass. Ann Surg 1968;167:98-100.
10. Licata AA, Fecanin P, Glowitz R. Metastatic adenocarcinoma from oesophageal colonic interposition. Lancet 1978;1:285.
11. Haerr RW, Higgins EM, Seymore CH, el-Mahdi AM. Adenocarcinoma arising in a colonic interposition following resection of squamous cell esophageal cancer. Cancer 1987;60:2304-7.
12. Houghton AD, Jourdan M, McColl I. Dukes A carcinoma after colonic interposition for esophageal stricture. Gut 1989;30:880-1.
13. Theile DE, Smithers M, Strong RW, Windsor CJ. Primary adenocarcinoma in a colonic ‘oesophageal’ segment. Aust N Z J Surg 1992;62:158-60.
14. Lee SJ, Koay CB, Thompson H, Nicolaides AR, Das Gupta AR. Adenocarcinoma arising in an oesophageal colonic interposition graft. J Laryngol Otol 1994;108:80-3.
15. Fritscher-Ravens A, Sriram PV, Thonke F, Jaeckle S, Maydeo A, Soehendra N. Synchronous adenocarcinoma in the transposed colonic conduit after esophagectomy for squamous cell cancer: endoscopic palliative resection while awaiting surgery. Gastrointest Endosc 1999;50:852-4.
16. Goyal M, Bang DH, Cohen LE. Adenocarcinoma arising in an oesophageal colonic interposition graft. Dis Colon Rectum 2000;43:555-8.
17. Liu CT, Ho S, Yeow KM. Primary adenocarcinoma arising in oesophageal colon interposition: report of a case. Hepatogastroenterology 2004;51:748-9.
18. Jass JR. HNPCC and sporadic MSI-H colorectal cancer: a review of the morphological similarities and differences. Fam Cancer 2004;3:93-100.
19. de la Chapelle A. Microsatellite instability. N Engl J Med 2003;349:209-10.