Multiple Small Intestinal Stromal Tumors Associated with Neurofibromatosis-1

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Gastrointestinal stromal tumors (GISTs) are rarely noted in association with neurofibromatosis-1 (NF-1, von Recklinghausen disease) as an individual gastrointestinal manifestation. We report here a case of multiple GISTs with an abundant skeinoid fiber in the jejunum of a 43-year-old woman diagnosed as NF-1. Histologically, the tumors were composed of uniform spindle-shaped cells with a fascicular pattern, almost indistinguishable from the histology characteristic of usual GISTs. However, multiple synchronous tumor occurrence, abundant skeinoid fiber, and presence of microscopic miniatures of stromal tumors are additional characteristic features of this case.

Key Words: Gastrointestinal stromal tumor (GIST), Neurofibromatosis 1 (von Recklinghausen disease), multiple.

INTRODUCTION

Gastrointestinal manifestations of neurofibromatosis-1 (NF-1) can be divided into four main groups: neuronal dysplasia; gastrointestinal stromal tumors (GISTs); endocrine cell tumors of the duodenum and periampullary region; and a miscellaneous group of other tumors.¹ GIST is referred as the majority of all gastrointestinal mesenchymal tumors, excluding tumors showing a definite differentiation toward the smooth muscle or schwann cells.² The majority of GISTs are isolated neoplasm and they are sporadically noted in non-NF-1 individuals without histological features that can help reliably distinguish NF-1-associated cases from sporadic cases. However, previous reports pointed out that tumors associated with NF-1 frequently showed multiplicity, jejunal location, and abundant skeinoid fibers.³ Here, we report a case of multiple small intestinal stromal tumors with abundant skeinoid fibers and spindle cell hyperplasia of the myenteric plexus, presenting like miniatures of stromal tumors, in the jejunum of a 43-year-old woman diagnosed as NF-1.

CASE REPORT

Clinical summary

A 43-year-old woman, who had no family history of NF-1, was admitted with abdominal pain and unexplained anemia. The physical examination revealed numerous café-au-lait patches and multiple cutaneous neurofibromas on the upper extremities and trunk. Esophagogastroduodenoscopy showed mild erosive gastritis, and the small bowel study showed an intraluminal protruding mass in the jejunum (Fig. 1). The abdominal exploration revealed multiple solid nodules in the jejunum, located from 30cm to 75cm distally from the Treitz ligament, and the segmental resection of the jejunum was performed.

Pathological findings

The resected segment of the small intestine showed nine subserosal solid masses (Fig. 2 and 3). The largest mass measured 3×2.5×2cm, showing a short stalk and hemorrhagic appearance. The second in size mass had a dumbbell shape, with a diffuse mucosal ulceration. The remaining
seven tumors were small in size, and ranged from 4 to 10 mm. Histologically, the tumors were composed of interlacing fascicles of the uniform spindle cells with elongated cytoplasm (Fig. 4). The tumor cells lacked pleomorphism, and mitotic figures were extremely rare. The results of the immunohistochemical stainings revealed that the tumor cells were diffusely positive for CD117, CD34, neuron specific enolase (NSE), and vimentin; and negative for smooth muscle actin, S100 protein, and desmin. Prominent skeinoid fibers, which were amorphous and eosinophilic materials in Hematoxylin-Eosin stain and positively stained with Periodic Acid Schiff (PAS) stain, were noted in the tumorous stroma (Fig. 5). Some of the tumors were directly continuous to the adjacent hyperplastic myenteric plexuses, and nodular proliferations of the spindle cells, like miniatures of the stromal tumors, were also noted in the region of the myenteric plexus (Fig. 6), and showed the same immunoreactivity patterns as the main tumors (Fig. 7).

**DISCUSSION**

Neurofibromatosis type-1 (von Recklinghausen
disease) is an autosomal dominant hereditary disease that may affect the gastrointestinal tract in up to 25% of the patients. Stromal tumors have been reported to be the most common lesion of the gastrointestinal tract, but the symptomatic cases account for less than 5% of patients. These tumors may cause obstruction, volvulus, intussusception, ulceration, bleeding or perforation. In our patient, the presumable cause of anemia was the overlying mucosal ulceration and bleeding of the tumor.

GISTs also occur sporadically in non-NF-1 individuals, and there is no histological difference between the NF-1-associated cases and the sporadic cases. However, tumors associated with NF-1 frequently show multiplicity, jejunal location, and abundant skeinoid fibers. We found nine cases of GISTs with abundant skeinoid fibers in patients with NF-1 in the literature in English, and all of these tumors were noted in the small intestine, predominantly in the jejunum (8 of 9 cases). Five of the nine cases were multiple; the number of tumors in each case was 2, 4, 7, over 50, and about 100, respectively. Our case seems to be a typical presentation of NF-1 associated GISTs, demonstrating the entire

Fig. 5. Prominent skeinoid fibers were noted in the tumorous stroma.

Fig. 6. A minute nodular proliferation of the spindle cells is noted between the inner circular and outer longitudinal muscle layers of the macroscopically normal intestinal wall.

Fig. 7. Immunohistochemistry showing strong and diffuse CD117 positivity in neoplastic cells (A), and proliferative spindle cells within hyperplastic myenteric plexus (B & C).
spectrum of features mentioned in previous reports.

Up to the present time, 6 gastrointestinal tumors associated with NF-1 have been reported in Korean literature: two types of synchronous biliary tract cancers in a patient with solitary, incidentally found jejunal stromal tumors; one case of cholangiocarcinoma; and one case of metachronous tumors (breast cancer, small bowel sarcoma) in the same patient. The features of one out of the two cases reported by Park et al. were very similar to those noted in the present case: multiple stromal tumors were present in the small intestine, the largest one measured 11 cm and the remaining about 0.5 cm. They described these tumors as neurofibromas arising in the Auerbach's myenteric plexus; however, no additional immunohistochemical or ultrastructural data was presented.

In this case, some of tumors were continuous to hyperplastic myenteric plexus, and nodular proliferations of the spindle cells, like miniatures of the stromal tumors, were noted in the region of the myenteric plexus, apart from the main masses. Walsh et al. previously reported such lesions and suggested a possible explanation that the Auerbach's myenteric plexus may be the site of the origin for GISTs in NF-1. The miniatures of the stromal tumors noted in this case can be an additional evidence of this suggestion. However, Handra-Luca et al. reported similar findings in their case of multiple familial gastrointestinal stromal tumors. Thus, such findings might help to understand the histogenesis of multiple GISTs, though they are not limited to the cases associated with NF-1.

REFERENCES