A Case Report of Bloom Syndrome Complicated by Colonic Cancer Due to Polyposis Degeneration: The Importance of Colorectal Cancer Screening

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Abstract

Bloom syndrome is a rare autosomal recessive disorder that attributes a chromosomal instability, described at the first time in 1954 by David Bloom, a dermatologist [1]. The Bloom’s syndrome registry (http://weill.cornell.edu/bsr/) reports 265 patients (140 male and 125 female) worldwide of whom about 26% are Ashkenazi Jewish ancestry, reaching a frequency of approximately 1 in 48000 [2]. Bloom syndrome is associated with the early occurrence of various cancer [3]. These patients develop the same tumour observed in the general population. However, they occur at an earlier age. Colon cancer is one of the most common tumours reported [4]. We report a case of Bloom syndrome in a 30-year-old man with colonic cancer developed due to polyposis degeneration to suggest the importance of colon and rectal cancer (CRC) screening for these patients.

Keywords: Colonic cancer; Bloom syndrome; Colonic polyposis; Screening

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Case Report

A 30-year-old, non-Ashkenazi Jewish man, with BS and hypothyroidism operated for ectopic testicular in 2011, developed in July 2013 an abdominal pain relieved by recurrent diarrhea specially located in the right lumbar and iliac regions, slightly abdominal distension, asthenia, loss of appetite and weight loss. Which occurred intermittently over at least a 3-month-period before he sought medical attention. There was no family history of the genetic disorder, personal or family history of colorectal carcinoma. Manifestations of BS in this patient included a severe growth deficiency even though the proportions of the body are normal with a slightly small cranium, a generalized facial erythema with facial telangiectatic erythema and repeated respiratory infections. The patient underwent a total colectomy. The patient recovered without complication and is currently being followed with endoscopic surveillance of the rectal stump. We present a case report about CRC diagnosed in a Bloom syndrome patient at an early age to suggest the necessity of CRC screening for these patients. Then, we recommend the necessity of doing at least annual examinations and colonoscopy as well as a rapid and thorough investigation of any new symptoms.

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Discussion

We present this clinical case on the diagnosis of CRC in a patient with Bloom syndrome at an early age to suggest the importance of CRC screening in this population. Bloom syndrome belongs to a group of "chromosomal breakage syndromes" which are transmitted in an autosomal recessive mode. The commonly acknowledged chromosomal breakage syndrome are Fanconi anaemia, ataxia telangiectasia, xeroderma pigmentosum and Bloom syndrome. However, genetically confirmation is necessary to eliminate a differential diagnosis. Growth deficiency is frequently the first manifestation that causes parents to seek medical attention. Patients with Bloom syndrome have a photosensitive lupus-like rash on the face, areas of café-au-lait spots [5]. The decrease of immunoglobulin level leads to recurrent respiratory and gastrointestinal tract infections [6]. A cytogenetic analysis looking for increased sister chromatid exchanges can then confirm or refute the diagnosis. Making the diagnosis has important management implications due to the increased risk of malignancy, probably hypersensitivity to chemotherapy and radiotherapy, and also the possibility that patients with Bloom syndrome are more likely to suffer from a range of complications of all treatments due to the other features of the condition immunodeficiency [7]. Then we should take extreme caution in using chemotherapy in patients with this disorder. Several authors reported similar toxicities in patients affected by Bloom syndrome associated with cancers (epipharyngeal B-cell-type lymphoma, nasopharyngeal B-cell-type malignant lymphoma, Burkitt lymphoma, hepatocellular carcinoma, Wilms tumour) [8,9]. As in the normal population, the colon is a common site of malignancy in persons with BS. The incidence of colonic adenomas among BS patients is unknown. The increased risk of malignancy in patients with colonic adenomas and BS is well established. Among the 210 individuals followed in The Bloom's Syndrome Registry, patients with colonic adenomas and BS is well established. Among BS patients is unknown. The increased risk of malignancy in persons with BS seems to have more right-sided and transverse colon tumours [10]. To make a definitive conclusion is impossible due to the small number of cases. The treatment of carcinomas by surgical resection at an early stage is the best curative option for the adult patient. Then, we recommend the necessity of CRC screening in this population by doing at least annual examinations as well as rapid and thorough investigation of any new symptoms.

Conclusion

The genetic disorder responsible for BS is uncovered. Genetic testing is currently available for diagnostics and to identify asymptomatic persons. BS is a worldwide, even in Tunisia, a rare genetic disorder. Only two cases are registered in Bloom syndrome register from North Africa. It might be more common. This population is exposed to develop cancer at a young age, particularly breast and colon carcinomas. Then a screening program should be offered to diagnose and treat these two cancers at an early stage avoid severe side effects. The possibility of antenatal diagnosis is possible when the risk of BS transmission is well evaluated.

Author Disclosures

Authors have no conflicts of interest or financial ties to disclose.

References