Transanal endoscopic microsurgery: The first attempt in treatment of rectal amyloidoma

Richa Sharma, Virgilio V George

Abstract
Localized amyloidosis is characterized by amyloid protein deposition restricted to one organ or tissue without systemic involvement. Gastrointestinal manifestations of localized amyloidoma are unusual, which makes amyloidoma restricted to the rectum a very rare diagnosis requiring a high index of suspicion. We present a rare account for rectal amyloidoma with an unusual presentation of obstructive symptoms and its treatment using a sophisticated surgical modality, transanal endoscopic microsurgery (TEM), which resulted in complete excision of the lesion without hospitalization and complications. The successful treatment for this rectal amyloidoma using TEM emphasizes the need to broaden its application in the treatment of various rectal lesions while preserving organ function and decreasing recurrence.

Key words: Transanal endoscopic microsurgery; Transanal endoscopic microsurgery; Amyloidoma; Localized amyloidosis; Rectal amyloidoma

INTRODUCTION
Amyloidosis is a rare group of disorders with an annual incidence of eight patients per million and is characterized by pathological deposition of fibrillar protein named amyloid, which disrupts organ struc-
The diagnosis of amyloidosis requires a high index of suspicion since the symptoms are nonspecific and may involve a single or multiple organ systems. Additionally, slow progression of the disease often delays diagnosis causing limited or palliative treatments. The manifestation of amyloidosis can be classified as primary, secondary, localized, or familial with prognosis varying in regards to the specific type of disease process\(^2,3\).

Localized amyloidosis, an extremely rare condition, is limited to a single organ. Furthermore, systemic features such as urinary and serum monoclonal proteins and/or clonal plasma cells in the bone marrow are absent making diagnosis difficult. Common sites of organ-restricted deposition include respiratory\(^4,5\), genitourinary tract\(^6,7\), in addition to, skin and soft tissue\(^8,9\). The mechanism underlying the formation of a localized amyloidoma remains poorly understood. De novo amyloid production\(^10\), in addition to, local plasmacytosis in chronic inflammatory diseases\(^6,11\) have been proposed for tissue restricted amyloidoma. Additionally, localized amyloid has been found to arise from certain fibrillar proteins produced by neoplasms such as calcitonin in medullary thyroid carcinoma\(^12,13\), amylin in insulinoma\(^14\) or prolactin in prolactinoma\(^15,16\).

Localized amyloidoma of the gastrointestinal tract is extremely unusual. All reported cases affecting the large bowel presented clinically with lower gastrointestinal bleeding\(^17-20\). Rarely amyloid of the colon produces a mass lesion causing obstructive symptoms\(^21\). Only two cases have been described in literature for rectal amyloidoma\(^22,23\). This case study is a rare account of the presentation of rectal amyloidoma and its surgical resection using an older but sophisticated surgical device, transanal endoscopic microsurgery (TEM).

**CASE REPORT**

A 66-year-old Caucasian male with past medical history of diabetes mellitus type II, hypertension, hyperlipidemia, and obstructive sleep apnea presented with unusual lower abdominal pain in September 2013. On digital rectal examination, the lesion was easily graspable and characterized as a hard, lobulated mass at the dentate line. Computed tomography scan was obtained, which showed a thickening of the left lower rectal wall with adjacent free gas due to the local perforation of the amyloidoma (Figure 1). A follow up colonoscopy found an irregular, poorly defined, semi circumferential rectal mass occupying about 50% of the rectal lumen located 2 cm from the dentate line and biopsy of the mass was taken. Pathology evaluation demonstrated abnormal deposit, which was found to be positive for congo red stain and was confirmed to be a rectal amyloidoma. No personal or family history of amyloid diseases or chronic inflammatory diseases was reported.

Proper workup was performed to assess for systemic amyloidosis. Serum blood count, comprehensive metabolic panel, coagulation studies, and liver and kidney function tests were all within standard limits. Serum total protein, albumin, alpha-1 globulin, alpha-2 globulin, beta globulin, and gamma globulin were normal. Urine analysis revealed a low level of monoclonal peak is present on electrophoreogram but was too small to quantitate and immunofixation yielded no abnormal bands. Furthermore, serum free kappa and free lambda protein levels were within regular limits resulting in a normal free kappa: lambda ratio of 1.02 (0.26-1.65). Chest X-ray, electrocardiogram, and echocardiogram showed no findings. Conclusively, work up was negative for systemic amyloidosis.

Amyloidoma of the rectum is an extremely rare rectal tumor. Lesions this low and of this size are more commonly treated by radical surgical intervention (low anterior resection or abdominal perineal resection) to achieve negative margins and evaluate the lymphovascular system for invasion. Due to the benign behavior of localized amyloid tumors, we offered the patient a local excision with a TEM.

**Surgical technique and post-operative course**

TEM includes an adjustable, multi-port proctoscope combined with CO\(_2\) mediated insufflation for optimal exposure of the rectum. This closed in-vivo system is then connected to a stereoscopic angulated optical system, which allows visualization and projection on a screen with higher resolution capabilities.

Briefly, for this case, the patient was placed in a lithotomy position. After examination of the rectum with a rigid, beveled, proctoscope, patient was positioned appropriately to localize the lesion inferiorly. The TEM device was connected to the anesthesia table. The TEM proctoscope was advanced to visualize the lesion in the left posterior aspect of the rectum at the inferior part of the TEM.
proctoscope (Figure 2A). Insufflation of CO₂ was started. The diameter of the patient’s rectum was too large to keep the lesion in the center, which led to mobilization of the lesion 1 cm distally and using this edge as a handle in order to allow optimal exposure for full thickness excision of the rectal wall using the monopolar cautery. After establishing the posterior dissection, we advanced in the mesorectal fat where the mass was divided full-thickness circumferentially from left and right and lastly, proximal aspect (Figure 2B). Small bleeding was controlled with the cautery device. The mass was completely excised, margins were obtained, and specimen was sent for pathology (Figure 3A). The defect was left open due to the large size and the lack of bleeding after cautery, in addition to, decreasing the likelihood of complications such as abscess formation and increased pain that have occurred in previous TEM cases when closing the defect.

Post-operatively, the patient did not have any bleeding from the surgical site and was able to go home the same day of surgery after recovering from anesthesia. Patient was gas incontinent for 2 wk post-operatively but resumed normal bowel habits afterwards without any local recurrence thus far. Findings of the histopathological report revealed acellular, homogenous, eosinophilic material underlying benign colonic mucosa positive for congo red stain and identified as local rectal amyloidoma (Figure 3B). Small fibers of the internal sphincter were also noted within the specimen (Figure not shown).

**DISCUSSION**

Many classifications have emanated for Amyloidosis over the years due to the advancement in knowledge for the disease process. Amyloidosis can be classified into four major categories. Primary amyloidosis, also known as immunoglobulin light chain amyloidosis, is diagnosed in individuals without previous diseases or coexisting conditions except multiple myeloma. Secondary or reactive amyloidosis is associated with chronic inflammatory conditions resulting in accumulation of hazardous byproducts. Familial amyloidosis includes variety of heritable mutations of proteins, such as transthyretin (TTR - most common form), apolipoprotein A-I, fibrinogen, cystatin and gelsolin[2,3]. Furthermore, the central nervous system and its’ exclusive environment is susceptible to amyloid deposition, which manifests as various forms of familial dementias[24]. The gastrointestinal system is a common site of amyloid deposition in patients with primary amyloidosis (70%) and secondary amyloidosis (50%) and the colon is frequently involved within the multi-systemic disease process[25]. However, single organ or localized form of amyloidosis, without systemic involvement, is rarely found in the colon[26-28].

Localized amyloidomas of the gastrointestinal
Localized amyloidoma is a benign, pathological deposition of fibrillar protein named amyloid, which can disrupt organ structure and function.

**References**

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