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**Schistosomal (bilharzial) polyps: Travel through the colon and beyond**

Emara MH *et al*. Schistosomal polyps

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**Abstract**

Schistosomiasis (bilharziasis) is a major neglected tropical disease. It is endemic in many tropical and subtropical communities. Schistosomal polyps (S. polyps) are not uncommon presentation of this infection. Although the colon is the most commonly affected organ, many other organs are affected. S. polyps are associated with a variable range of morbidity independent of the Schistosomal infection. S. polyps are frequently described in endemic areas and increasingly reported in non-endemic areas mainly among immigrants and visitors to the endemic areas.This review aimed to increase awareness of practitioners, especially gastroenterologists, for this peculiar type of polyps caused by this neglected infection hence improving patient outcomes.Web-based search of different databases was conducted for the literature focusing the development of S. polyps in the colon and other organs with analysis of the clinical manifestations, diagnosis and treatment. The following key words were used in the search, “Schistosomiasis” OR “Bilharziasis” AND “Polyps” OR “Polyp” AND “ Colon” OR “Small intestine” OR “ Duodenum” OR “ Stomach” OR “Esophagus” OR ” Gallbladder” OR” Pharynx” OR “Larynx” OR “Trachea” OR ”Urinary bladder” OR “ Ureter” OR “Renal Pelvis” OR “Urethra”. All publication types including case reports, case series, original research, and review articles were retrieved and analyzed. S. polyps are not infrequent presentation of acute or chronic Schistosomal infection. S. polyps are described in many organs including the bowel, genitourinary tract, skin, gallbladder and the larynx. Presentation of S. polyps is variable and depends on the site, number as well as the polyp size. The relationship of *S. polyps* to malignant transformation is a matter of discussion. Presence of S. polyps is sometimes the only manifestation of Schistosomiasis. Small polyps can be treated medically with praziquantel, while large accessible polyps are amendable for endoscopic excision through different polyp resection techniques. However, huge, complicated, non-accessible and suspicious polyps are indicated for surgical management or advanced endoscopic resection when appropriate. Clinicians and endoscopists should be aware about these facts when treating patients living in, immigrated from or visiting endemic areas.

**Key Words:** Schistosomiasis; Bilharziasis; *Schistosomal polyps*; Colon; Praziquantel

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**Core Tip:** Schistosomiasis is associated with a wide range of pathological lesions including development of polyps. Colon is the commonest site for polyp development, however polyps are reported in many organs including urinary bladder, ureters, larynx, duodenum, small intestine, gallbladder, anus, uterine cervix and external genitalia. Schistosomal polyps are associated with a wide range of morbidity according to the polyp site, size and number. The malignant potential of these polyps is a hot point of discussion. Although small sized polyps can regress with medical therapy using praziquantel, large accessible polyps can be retrieved endoscopically. Complicated, huge and inaccessible polyps can be treated surgically.

**INTRODUCTION**

Schistosomiasis is a major neglected tropical disease. It is endemic in many geographical regions mainly in Africa, Latin America and Asia**.** Many data have been published from endemic areas and also from areas where the infection is not likely to occur with diverse clinical presentations. In non-endemic areas the infection is reported among immigrants or visitors to endemic areas[1,2].

This parasitic infestation was described in early human history, with evidence of infection reported in the ancient Egyptian papers and probably other old civilizations. The infection was characterized in the 19th century by the German pathologist Theodor Bilharz; that is why it is named after him as “Bilharziasis”. There are many species of the parasite. Human infection is likely caused by five species, namely *Schistosoma Mansoni, S. Haematobium, S. Japonicum, S. Intercalatum and S. Mekongi*[1]. It is obvious that, the infection is linked to water supplies because of the snail intermediate host settles in the water canals and hence agricultural communities are the ultimate victims of the infestations, although persons who came in contact with infected water are prone also to catch the infection.

The clinical manifestation of this infection is either acute or chronic. The acute manifestations are related to the invasion of the human body by the cercarial invading stage through the skin, migration within the body and the early stage of ovi-position. Chronic manifestations are related to the establishment of adult worms and trapping of the deposited ova within the tissues and consequently granuloma formation[1]. The acute presentations include constitutional manifestations with fever, myalgia, urticarial rashes and with ovi-position the manifestations will change to hematuria (urinary Schistosomiasis), diarrhea with blood (intestinal Schistosomiasis), and chronic blood loss; manifested as anemia[1,3]. In chronic cases, development of fibrosis is the hallmark of the disease and this results in devastating sequelae including portal hypertension, hepatic peri-portal fibrosis, and splenomegaly for intestinal Schistosomiasis, while urinary Schistosomiasis is associated with obstructive lesions, increased incidence of stone formation, chronic urinary tract infections, and urinary bladder malignancy[4].

The current review aimed to increase awareness of practitioners, especially gastroenterologists, for this peculiar type of polyps caused by this neglected infection and hence improve patient outcomes.

**LITERATURE SEARCH**

Web-based search of different databases was conducted, including PubMed/MEDLINE, Cochrane library, Web of Science, Ovid, Science Direct, Scopus, Directory of Open Access Journals, EBSCO HOST, ProQuest, Institute for Scientific Information, EBESCO, Egyptian knowledge bank, Google scholar, *Reference Citation Analysis* (https://www.referencecitationanalysis.com/) and the Research Gate, for relevant articles focusing on the development of Schistosomal polyps (S. polyps) in the colon and other organs with analysis of the clinical manifestations, diagnosis and treatment. The following key words were used in the search, “Schistosomiasis” OR “Bilharziasis” AND “Polyps” OR “Polyp” AND “ Colon” OR “Small intestine” OR “ Duodenum” OR “ Stomach” OR “Esophagus” OR ” Gallbladder” OR” Pharynx” OR “Larynx” OR “Trachea” OR ”Urinary bladder” OR “ Ureter” OR “Renal Pelvis” OR “Urethra”. All publication types including case reports, case series, original research, and review articles were retrieved and analyzed.

**PATHOGENESIS OF S. POLYPS**

The development of polyps in general is linked to hollow organs or organs with contact to space; this enables external growth. Consequently, polyps are frequently reported in the bowel rather than other organs. The final shelters for adults Schistosomes are the intestinal and pelvic venous plexuses whatever the species are and hence the development of S. polyps in the hollow organs drained by this plexuses is not strange, *e.g.,* colon and urinary bladder, while development of polyps in other organs, *e.g.,* skin is considered ectopic. Furthermore, S. polyps may develop in atypical sites, *e.g.,* duodenum, that are not drained by the intestinal or pelvic veins.

***The development of polyps in different stages***

**Stage of ova deposition and trapping:** The eggs laid by the adult parasite had a characteristic spine that helps it to dig and find way out, *e.g.,* through the colon mucosa to be released with the stool. The initial step in polyp formation starts when Bilharzial eggs are deposited and trapped within the superficial layers of submucosa. The submucosal connective tissue is delicate and not superficially bound by firmer tissue. Consequently, accumulation of large amounts of eggs, reactive cellular debris and vascular granulation tissue do occur. In the submucosa, the trapped ova produce a TH2 cell-mediated inflammatory response with many cellular and chemical elements being recruited to the area and granuloma formation begins. The hallmark of the granuloma is infiltration by eosinophils (eosinophilic granuloma)[1], however, during this exudative stage of granuloma formation, other inflammatory cells are seen including the epithelioid macrophages and mononuclear cells[1,5].

**Stage of proliferation:** Within the granuloma necrosis occur[5]. Healing of the developed necrotic foci is associated with fibrous connective tissue formation. Furthermore, the adjacent muscularis mucosa becomes hypertrophied[1,5]. The fibrous tissue in the submucosa and the hypertrophied muscularis mucosa form a barrier hindering the ova transit from the veins to the lumen, hence more ova are entrapped. Sometimes, the adult worms either alone or matted are seen lodged and obstructing small venules within the polyp[5].

**Stage of growth and protrusion:** The granuloma formation is the basic pathology unit of chronic Schistosomiasis. Ova entrapment induces a foreign body reaction with progressive inflammation and fibrosis (chronic granuloma). As this process continues, a nodule is formed that elevates the hypertrophied muscularis mucosa and mucosa to form the polyp[6]. The polyp’s mucosa harbors many goblet cells which secrete large amounts of mucus and this matches the abdominal pain and passage of mucus with stool seen among those patients. Furthermore, the delicate, and highly vascular nature of the polyp facilitates bleeding with the passage of stools[1,5-7].

This previously described mechanism of fibrosis and ova entrapment explain why the Schistosomal eggs are concentrated within the polyps more frequently than in the surrounding mucosa and submucosa[8]. Furthermore, non-inflammatory polyps associated with Schistosomiasis including adenomatous, hamartomatous, and hyperplastic polyps have been reported[1,9].

**Stage of fibrosis:** When the infection became chronic, the developed granulomas are infiltrated with fibroblasts, and fibrosis does occur[2,10]. Consequently, the polyps may become fibrotic. Small polyps especially when medically treated with praziquantel (PZQ) can then regress in size[11].

**SCHISTOSOMAL COLONIC POLYPS**

***Site***

The most common site for the development of S. polyps is the left colon, mainly recto-sigmoid region[3,12] because mesenteric (especially inferior mesenteric) and pelvic venous plexuses are the final habitat of the adult parasite. However, no part of the colon is immune against development of S. polyps. The polyps were described in all parts of the colon mainly in the rectosigmoid[12], but also were described in the caecum[13], ileocaecal valve[14], appendix[15], ascending colon[16], transverse colon[17] as well as the descending colon[12], some reports documented the distribution of S. polyps all through the large bowel starting from the caecum up to the anus during the course of heavy infestations in endemic areas[18].

***Number***

Schistosomiasis in the endemic areas, is associated with development of multiple polyps and multiplicity correlates with the density of infection[3,18]. However, there are many reports that Schistosomiasis can manifest by single colon polyp discovered either incidentally during colonoscopy or complicated[13,16,19] even without any clinical manifestations suggestive of Schistosomal infection.

***Size***

Schistosomiasis is usually associated with small-sized polyps[1,3,12]. However, large-sized S. polyps have been described in many case reports[12,16,18,19]. We reported earlier[20] that, no part of the colon was immune against development of large S. polyps. Many cases presented with solitary S. polyps[13,16,19,21] even in absence of any Schistosomiasis-related colon inflammation[9,13,16,19,21].

***Atypical presentations and complications***

Schistosomal colonic polyps had a wide range of atypical presentations. Elbatee *et al*[19], Alyhari *et al*[7], and Al-Zubaidi *et al*[22], described colo-rectal cancer (CRC) like presentation due to huge polyp size, abnormal polyp morphology or both, respectively. Furthermore, Smith *et al*[23], described diffuse colonic polypoid masses with dysensteric features, a picture endoscopically indistinguishable from familial polyposis and severe ulcerative colitis. Polyp-like lesion in the appendix manifestated as acute appendicitis was reported in a Chinese woman by Zhu *et al*[15], while pan-colonic inflammation and polyposis were described in Kenia by Bosire *et al*[18].

**S. POLYPS BEYOND THE COLON**

Urinary bladder polyps due to *S. Haematobium* have long been described and development of polyps correlates with heaviness of the infection in endemic areas[23]. Urinary bladder S. polyps were also reported among visitors to endemic areas[24].Urinary S. polyps are not limited to urinary bladder, they were described within the ureters with a prevalence of 5.9% (30/511) in a large Egyptian study[25]. Due to the anatomical constraints, ureteric S. polyps tend to be small[25].

The atypical sites of S. polyps (Table 1) were described not infrequently in the literature. As early as in 1951, [Gilges](https://www.cabdirect.org/cabdirect/search/?q=au%3a%22Gilges%2c+W.%22)[26], described a skin polyp in the clitoris area of an African child. Furthermore, Schistosomal vulvual polyps have been described as vulvual swelling in 9- and 11-year-old girls from endemic areas in Senegal[27] and Nigera[28] respectively.

An ectopic cervical polyp was reported in Puerto Rico by File *et al*[29], despite the light Schistosomal infection. While Eladl *et al*[30] reported an endocervical polyp containing granulomas rich in viable eggs of *S. Hematobium* in a 43-year-old Egyptian woman who manifested with vaginal bleeding.

A slowly growing anal polyp containing Schistosomal ova as well as adult worms was described in a young Brazilian adult by Raso *et al*[5], the parasites probably migrated to the anus through the veins of hemorrhoidal plexus and it was associated with troublesome mass lesion with intense pruritus.

Duodenal polyps due to schistosomiasis have been reported in the literature although not common. In the duodenum it is discovered during endoscopic evaluation for obscure anemia and/or abdominal pain. The described duodenal S. polyps are either small, multiple and sessile[31]or solitary and large[32]. One case of dull abdominal pain was diagnosed in the United States with Schistosomiasis complicating a huge duodenal Peutz-Jeghers hamartomatous polyp[9].

The small intestine is infrequently affected with S. polyps and due to its narrow lumen, which is usually characterized by obstructive manifestations and diagnosis is established either at laparotomy or more frequently retrograde through histopathologic assessment. Small bowel obstruction due to huge polyp/polypoid mass at the ileocaecl valave[14] was described among visitors as well as residents of endemic areas[33]and small bowel obstruction is the usual presentation of such cases.

Gallbladder S. polyps are infrequently reported in the literature, the developed polyps are usually small in size and grew in the fundus, hence obstructive presentation is not likely and it is usually manifest (if any) as right upper quadrant pain. Both urinary (*S. Haematobium*) and intestinal Schistosomiasis (*S. Mansoni* and *S. Japonicum*) have been associated with gallbladder Schistosomiasis[34,35].

Larynx is less commonly affected during the course of Schistosomiasis. Toppozada described S. polyp involving the right vocal cord of a 25-year-old Egyptian male presented with 5-mo progressive hoarseness of voice, surgical microscopic excision and pathological examination showed granuloma with terminal sine ova[36].

**S. POLYPS AND MALIGNANT TRANSFORMATION**

The malignant potential of S. polyps is debatable. Although the relationship of urinary Schistosomiasis to malignancy is well established, the direct tumorigenic impact of intestinal S. polyps is questionable.

The bowel S. polyps have been implicated in the development of CRC and liver (HCC) cancers. The evidence supporting the malignant potential of intestinal Schistosomiasis is in favor of *S. Japonicum* rather than *S. Mansoni*. The chronic inflammatory state induced by ovi-position in the submucosa and other tissues have been proposed as the potential mechanism linking *S. Japonicum* in the Far East to the development of CRC and liver cancer[37].

So far the link between *S. Mansoni* and CRC has been viewed as no more than an epidemiological association and most published literatures deny the precancerous potential of the *S. Mansoni*-associated lesions including the polyps[38]. However, the mood of clinicians is spoiled by the emerging evidence incriminating *S. Mansoni* as a potential carcinogen. A large Egyptian study proposed an association between *S. Mansoni* infection and CRC relying on the increased levels of carcinoembryonic antigen within Bilharzial polyps’ tissue[39]. In addition, one case report[40] retrieved Schistosome ova from the CRC specimens during histologic examination. Parasitism is associated with DNA repair defects with resultant genome instability, a commonly reported anomaly in CRC[38]. Furthermore, an emerging evidence through biomolecular mechanisms suggests an association between *S. Mansoni* and human carcinogenesis mainly for HCC and CRC and this is likely through many egg-related agents and Th2-immune mechanisms[41]. A single recent report described for the first time a concomitant infection with *S. Mekongi* and rectal cancer[42].

The association of *S. Haematobium* and the urinary bladder squamous cell cancer is well established[43] and that is why *S. Haematobium* is classified as group 1 definitive biological carcinogenic. This association is probable with all forms of urinary bladder- induced Schistosomal pathology including chronic bladder wall inflammation and irritation by the deposited ova, the induced ulcers, sandy patches and also to the developed S. polyps[44]. There is no clear evidence in the literature linking intestinal Schistosomiasis to bladder tumors and vice versa[9].

**DIFFERENTIAL DIAGNOSIS**

The severe forms of intestinal Schistosomiasis are characterized by the presence of multiple variable sized polyps against a background of diffuse mucosal affection showing mucosal erythema, and ulcerations, a picture mistaken with inflammatory bowel disease[3]. S. polyps of the colon are usually located in the recto-sigmoid region[12] and tend to be small[3,12] (Figure 1A) and dark grey in color with surface ulceration, however other conditions with multiple colonic polyps should be considered in the differential diagnosis.

The solitary forms of S. polyps should be differentiated from other polyps according to the location. In fact, there is no characteristic morphology for S. polyps[45]. The huge polyps are commonly mistaken as malignant polyps (Figure 1B) and in such cases diagnosis is achieved through histopathology either by biopsy or after excision[19,21,22].

**DIAGNOSIS**

Diagnosis of polyps of the Schistosomal origin is sometimes challenging. There are no peculiar endoscopic morphologic features of S. polyps in comparison to other types of polyps and high index of suspicion is required especially in endemic areas. However, most cases are diagnosed either through endoscopic biopsy from the polyps or retrograde upon histopathology examination of removed polyps where the characteristic eosinophilic granulomas with the pathogenomic ova (Figure 2) of the parasite are seen [12-19,22] and infrequently the mature parasites may also be seen embedded within the background tissue of the excised polyps[22,29] or is seen lodged within thrombosed veins inside the polyp[5].

Linking the developed polyps, whatever its location, with active intestinal or urinary Schistosomiasis is not always achievable. In many cases described in the literature, identification of Bilharzial ova in fresh stool or urine samples taken from the same cases was not possible[9,16,31]. Furthermore, S. polyps may be the only clue of a Bilharzial infection[19,31].

Barium enema was used frequently to diagnose Bilharzial polyps of the bowel[46] in endemic areas before the era of endoscopy. However, currently the role of contrast study in diagnosis of intestinal or urinary Schistosomiasis is not likely to replace the standard methods for direct ova detection in stool or urine samples or detection of the ova with its characteristic granuloma on endoscopic or surgical specimens due to low sensitivity and specificity of these contrast studies[47].

Upon histologic examination, the polyp had a stalk of fibrous connective tissue that project from the submucosa into the lumen and is covered with mucosa. The overlying mucosa harbor distorted glands showing variable degrees of mucoid activity (Figure 1C), mucinous degeneration, and adenomatous hyperplasia. The covering mucosa frequently had focal areas of ulceration. Larger areas of ulceration may be replaced by granulation tissue. Mononuclear cells, eosinophils, and few polymorphonuclear leucocytes infiltrate the mucosa[48].

The peculiar granuloma had a centrally located ova (viable and/or nonviable) surrounded by cellular infiltrate characteristically with eosinophils, mononuclear cells and multinucleated histiocytes (Figure 2)[1,5], chronic granulomas are surrounded by fibroblasts (Figure 3). The supporting tissue is composed of fibrous connective tissue and muscle derived from the muscularis mucosa. Blood vessels may be present in large numbers but diminish with progression of fibrosis[48].

**TREATMENT**

***Medical treatment***

Currently, PZQ is the recommended first-line medical therapy for Schistosomiasis. The drug is active and effective in treatment of all Schistosome species with no reports of resistance[49]. Small-sized polyps can be cured with PZQ therapy alone. The dose is 40-60 mg/kg, it is given as a single dose after a meal[1] and can be divided over one day[3,9], and can be repeated[7]. The side effects associated with PZQ are usually mild and include nausea, vomiting, abdominal cramps and allergic reactions. S. polyps are exceptional types of polyps that can regress with medical treatment[11] especially when they are small and are not associated with manifestations necessitating excision. All cases associated with large or ectopic polyps after being excised either endoscopically or surgically were followed by treatment with PZQ irrespective of the patients' active intestinal or urinary Schistosomiasis state[9,13,19,34].

***Endoscopic treatment***

Large polyps are indicated for excision. Endoscopic excision forS. polyps have been reported thoroughly in the literature as early as the 80s of the last century[50,51] with the early time of endoscopy. S. polyps were excised with different techniques including cold snare for small polyps, standard direct hot polypectomy for larger polyps whatever the site is the duodenum[9], or the colon[3].Hot polypectomy after application of endoloop[52], or clips[13] has been associated with good outcomes. While advanced endoscopic resection techniques including endoscopic mucosal resection[21,22]and endoscopic submucosal dissection[53]were also used for endoscopic removal of large S. polyps. All endoscopic resection techniques were used with accepted safety profile. The adverse events reported were not different from those reported with other types of polyps[3,9,13,16,22]. Resection of S. polyps combined with medical treatment using PZQ was successful [3,9,13,16,22,48]. Endoscopic resection of urinary S. polyps is less frequently performed in comparison to bowel and was associated with acceptable success rates[24].

***Surgical treatment***

Surgical excision is not the standard of care in treatment of S. polyps. However, surgery is usually the ultimate solution for challenging situations including cases presented with bowel obstruction[14,33], or when the lesion is morphologically or radiologically suspected to be malignant[7,12,19] and also for lesions in ectopic areas including the anus[5], cervix[30] and the external genitalia[27,28]. Atypical sites of S. polyps may require special surgical approaches, *e.g.,* in the larynx, the surgical microscope is essential to excise the polyp[36], while in other situations, *e.g.,* gallbladder, the organ is excised with the polyp en bloc[34]. All cases managed by surgery, similar to endoscopic management, should be followed by PZQ therapy.

***Outcomes of treatment***

The outcome of S. polyps achieved by the above mentioned treatments depend on many factors. First, the number of lesions detected. For single lesions excision (either endoscopic or surgical) is usually curative, while multiple lesions may require more rounds of excision[3]. Second, patients' residency; patients residing in endemic areas are susceptible to re-infection and hence should follow the general preventive measures together with PZQ therapy that sometimes given as chemoprophylaxis or as mass treatment campaigns even without testing for the presence of the parasite[54], while patients out of the endemic areas a single course of PZQ is usually sufficient for treatment. Third, the location of the polyps, some polyps are accessible for endoscopic treatment, such as polyps in the colon, and urinary bladder, while other lesions, *e.g.,* gallbladder are not. Last, presentation of the lesions, such as lesions presented with acute manifestations including bowel obstruction, surgery usually the corner stone of treatment and in such cases diagnosis is often achieved retrograde[14,33].

**CONCLUSION**

Schistosomiasis is a neglected tropical disease and its prevalence is no more limited to endemic areas. S. polyps are not infrequent presentation of acute or chronic Schistosomal infection. S. polyps are described in many organs including the bowel, genitourinary tract, skin, gallbladder and the larynx. Presentation of S. polyps is variable and depends on the site, number as well as the polyp size. The relationship of S. polyps to malignant transformation is a matter of discussion. Small polyps can be treated medically with PZQ, while large accessible polyps are amendable for endoscopic excision through different polyp resection techniques. However, huge, complicated and suspicious polyps are indicated for surgical management or advanced endoscopic resection if appropriate. Clinicians and endoscopists should be aware about these facts when treating patients living in, immigrated from or visiting endemic areas.

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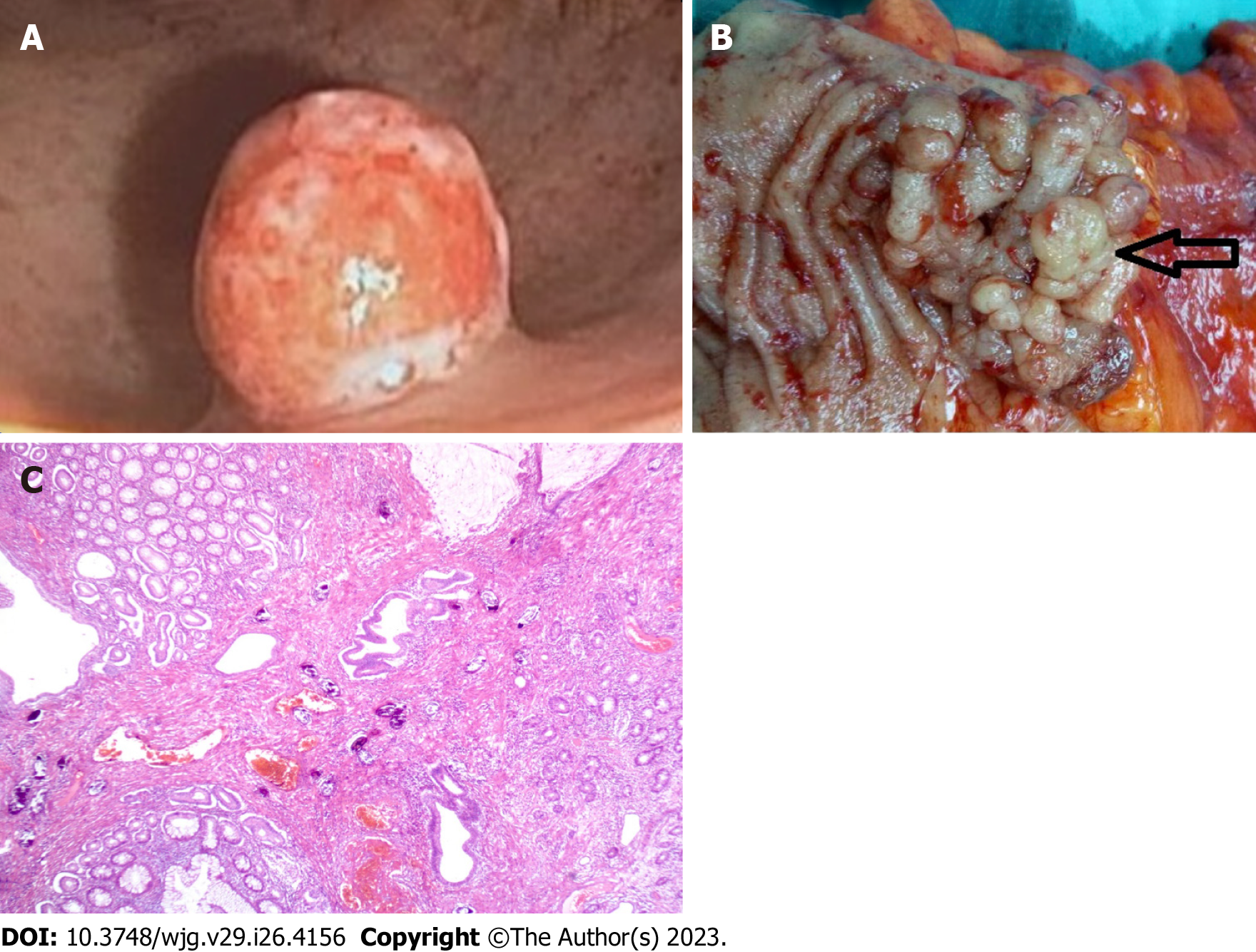
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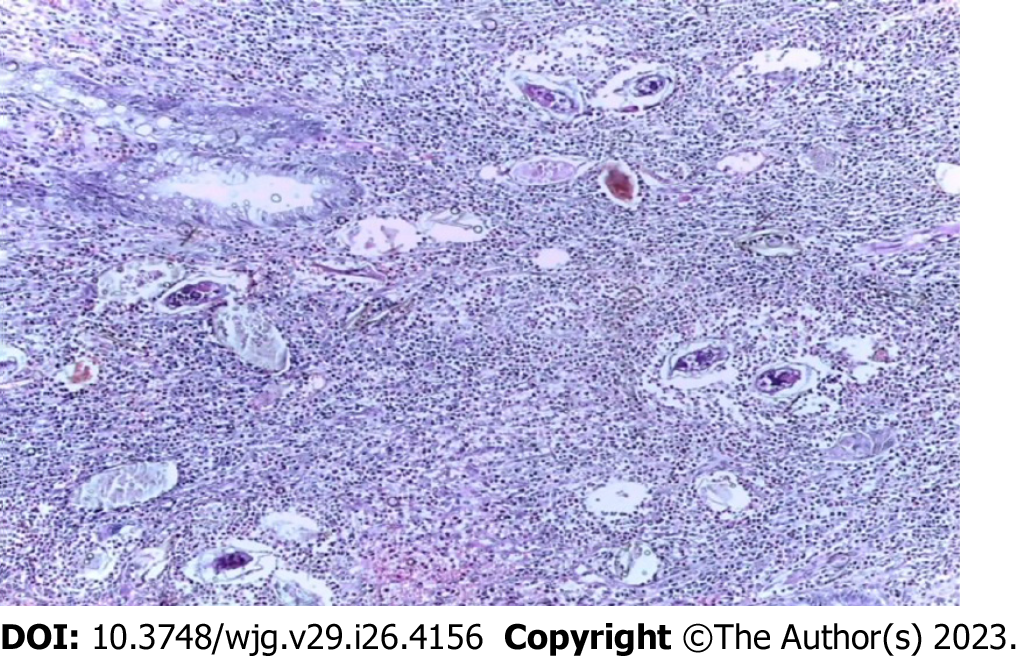
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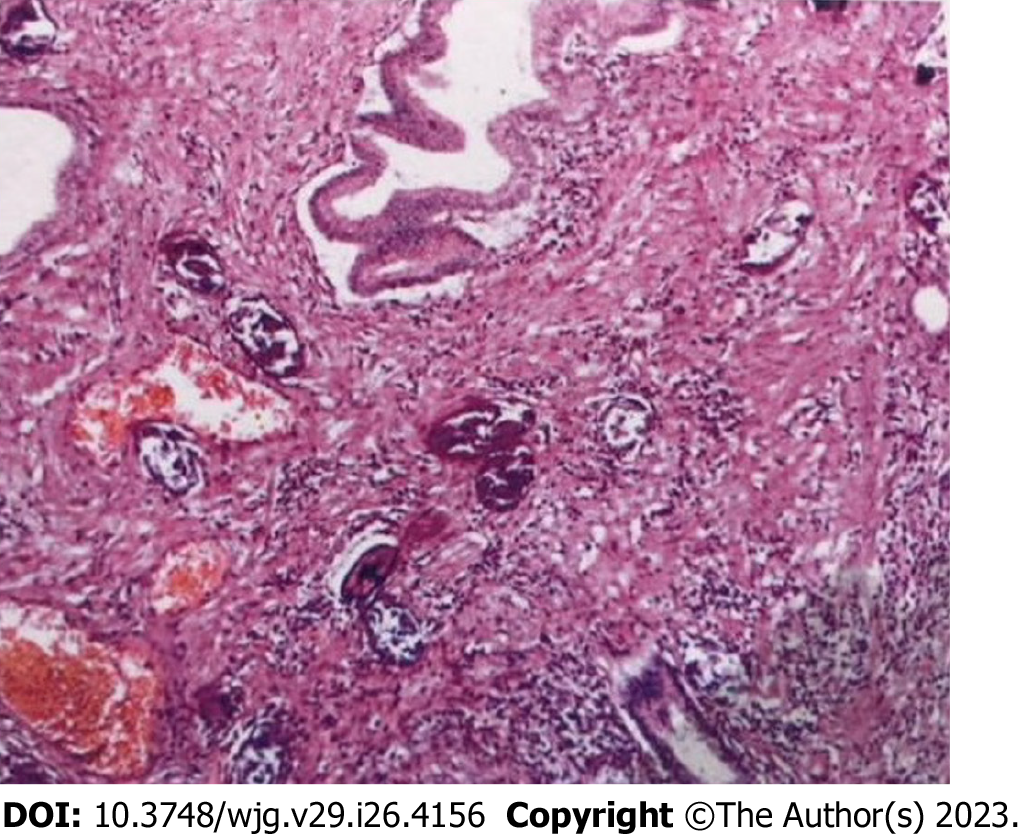
**Figure Legends**



**Figure 1 Schistosomal polyp.** A: Small Schistosomal polyp located in the rectum. Note its dark red color with minute surface ulcerations; B: Surgical specimen of huge lobulated Schistosomal polyp morphologically and radiological confused with colo-rectal cancer; C: Schistosomal polyp with noticeable mucoid hyperplasia.



**Figure 2 Active schistosomal granuloma; multiple schistosomal eggs in the submucosa surrounded by cellular infiltrate composed of eosinophils, lymphocytes, plasma cells and macrophages.**



**Figure 3 Schistosomal granuloma infiltrated by fibroblasts with dense submucosal fibrosis.**

**Table 1 Summary of atypical and ectopic schistosomal polyps**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Ref.** | **Site** | **Size** | **Presentation** | **Source** |
| Raso *et al*[5], 2013 | Anus | Large (2.5 cm) | Swelling with pain and intense pruritus | Endemic area |
| Gilges[26], 1951 | External genitalia (clitoris) | Not mentioned | Mass like | Report from endemic area |
| Eladl *et al*[30], 2012 | Cervix | Large sized (3 mc) | Vaginal bleeding | Endemic area |
| File *et al*[29], 1998 | Cervix | Large | Gynecologic | Common area of spread. Adult worms within the polyp |
| Dioussé *et al*[27], 2016 | Vulva | Large | Mass like | Endemic area |
| Sahabi and Rabiu[28], 2017 | Vulva | Large | Mass like | Endemic area |
| Gonzalez *et al*[9], 2021 | Duodenum | Huge (≥ 4 cm) | Dull abdominal pain | Resident of non-endemic area with a history of travel to endemic areas |
| Altonbary *et al*[31], 2014 | Duodenum | Diminutive (≤ 4 mm) | Abdominal pain and generalized lymphadenopathy | Endemic area |
| Thatcher *et al*[32], 1984 | Duodenum | Large | Isolated polyp | Non-endemic area |
| Lamyman *et al*[14], 2006 | Ileocaecal valve area | Large | Intestinal obstruction | Visitor to endemic areas |
| Ali *et al*[34], 2021 | Gallbladder fundus | Small 6 mm | Vague Abdominal pain | Patient from endemic area |
| Ghimire *et al*[35], 2020 | Gallbladder fundus | Diminutive 3 mm | Right hypochndrial pain | Non-endemic areas |
| Toppozada[36], 1985 | Larynx, right vocal cord | Large | Hoarseness of voice | Endemic area |



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