Common Inflammatory Disorders and Neoplasia of the Ileal Pouch: A Review of Histopathology

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Abstract

Ileal pouch-anal anastomosis (IPAA) is the standard restorative procedure after proctocolectomy in patients with ulcerative colitis (UC) who require colectomy. The ileal pouch is susceptible to a variety of insults including mechanical injury, ischemia, fecal stasis, and infectious agents. In addition, the development of recurrent and idiopathic inflammatory bowel disease and neoplasia may occur in the ileal pouch. Although clinical, endoscopic, and radiographic examination can diagnose many ileal pouch diseases, histologic examination plays an essential role in diagnosis and management, particularly in cases with antibiotic refractory chronic pouchitis and pouch neoplasia.

Keywords: Inflammatory disorder; Neoplasia; Ileal pouch; Diagnosis; Pouchitis; Histopathology; Ulcerative colitis

Introduction

Over the last several decades, ileal pouch-anal anastomosis (IPAA) has evolved into the standard restorative procedure after proctocolectomy in ulcerative colitis (UC) and familial adenomatous polyposis syndrome (FAP). It is also used in patients with indeterminate colitis and in selected cases of Crohn’s disease (CD) without small bowel involvement. A variety of diseases can affect ileal pouch and correct diagnosis of these diseases requires a synthesis of clinical, endoscopic, radiographic, and pathologic information [1, 2]. Our knowledge of pouch diseases has been greatly expanded in recent years and it forms the basis of this review.

Pouchitis

Pouchitis is defined as a non-specific inflammation of the ileal reservoir. Pouchitis is the most common complication of IPAA in patients with UC as a result of dysbiosis. It also occurs in FAP patients with IPAA but at a much lower rate [2]. It has been reported that the frequency of pouchitis is related to years of follow-up, occurring in up to 50% of patients 10 years after IPAA in large series from major referral centers [2-5]. It is most frequently diagnosed within the first year after ileostomy closure. Symptoms of pouchitis are non-specific and may include increased stool frequency and liquidity, abdominal cramping, urgency, tenesmus, and pelvic discomfort. In addition, rectal bleeding, fever, or extraintestinal manifestations may also occur in some patients. Patients suspected of having pouchitis should undergo pouchoscopy to document the presence of epithelial injury and inflammation and to obtain mucosal biopsies for histologic confirmation. For optimal histologic evaluation, biopsies should be taken from different locations including the neo-terminal ileum, pouch body, and rectal cuff. Specimens should be submitted separately either fresh in saline or fixed in 10% neutral buffered formalin.

The 18-point pouchitis disease activity index (PDAI) was proposed by Sandborn et al in 1994 [6] to diagnose pouchitis; it consists of three components: clinical symptoms, endoscopic findings, and pathologic findings. Each component has a maximal score of 6 points. The pathologic findings are based on the degree of neutrophilic inflammation and ulceration. The histologic score is assigned as follows: neutrophilic infiltration (1, 2, or 3 for mild, moderate + crypt abscess, and severe + crypt abscesses, respectively) and mean ulceration per low-power field (1, 2, or 3 for < 25%, 25-50%, and > 50%, respectively). Although the clinical use of the 18-point PDAI was later validated by other groups [1], some researchers have suggested that a combination of symptomatology and endoscopic findings may be adequate for diagnosing pouchitis [7]. However, symptomatology per se is not sufficient for diagnosing pouchitis in up to 25% of cases [1]. In cases with symptoms, a PDAI of < 7 may suggest a diagnosis of irritable pouch syndrome in patients without cuffitis [8].

As with symptomatology or endoscopic findings, histological findings of pouchitis are non-specific. Histologic findings include acute inflammation with neutrophilic infiltration, crypt abscesses, erosion and ulceration, in association with a chronic inflammatory infiltrate [9, 10] (Fig. 1). The presence of inflammation alone in mucosal biopsies from an asymptomatic patient should not prompt the diagnosis of pouchitis and treatment with antibiotics.

Based on the clinical onset and duration of symptoms, pouchitis is classified into acute and chronic. The latter is fur-
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ther divided into antibiotic-responsive, -dependent, and -refractory depending upon the response to antibiotic treatment [11]. In one large series of IPAA, chronic pouchitis (defined as more than three attacks of pouchitis in a 12-month period) occurred in 15.9% of patients [2]. For patients with presumed chronic pouchitis who have persistent symptoms, alternative diagnoses should be considered along with histologic evaluation of biopsies to help narrow the differential diagnosis. If no specific etiologies are identified, the patients are considered to have chronic pouchitis and are usually treated with a long course of antibiotics.

Secondary pouchitis with identifiable infectious agents

Inflammatory bowel disease (IBD) is treated by immunosuppressive and biologic therapy creating an environment for opportunistic infections. Histologic examination of mucosal biopsies plays an important role in the diagnosis of certain types of infectious pouchitis.

C. difficile pouchitis

Ileal pouch C. difficile infection was initially reported by Mann et al (2003) [12]. Additional studies have revealed that C. difficile infection of the pouch is associated with male gender, pre-surgery left-sided colitis, pre-surgery antibiotic use, previous hospitalization, low serum immunoglobulin level and the presence of post-surgery mechanical intestinal complication [13, 14]. Clinically, C. difficile infection of the pouch can be refractory or recurrent. The diagnosis mainly relies on symptomatology, endoscopic examination and detection of C. difficile toxins A and B by enzyme immunoassay or C. difficile toxin B gene by PCR assay in stools. Histologic evaluation can be helpful in identifying cases without typical endoscopic features. For example, some mild cases may only demonstrate subtle abnormalities such as edema, loss of vascularity and friability while some severe cases may only show ulceration without obvious pseudomembranes. Histologic findings in pouch C. difficile infection are non-specific and include acute inflammation and villous blunting/atrophy [15].

Cytomegaloviral pouchitis

Cytomegalovirus (CMV) infection in patients with IPAA has been recognized as an etiology of chronic pouchitis and one of the treatable causes of secondary pouchitis [16, 17]. Clinical presentation of CMV pouchitis is non-specific and may include abdominal pain, increased stool frequency, and fecal incontinence, mimicking idiopathic/conventional pouchitis. However, patients with CMV pouchitis often have fever, which is not common in conventional pouchitis [18]. Endoscopic features of CMV pouchitis are non-specific and indistinguishable from conventional pouchitis. The diagnosis relies on the detection of CMV or its antigen. CMV can infect endothelium, stromal cells, and epithelial cells. CMV-infected cells often show typical features on routine hematoxylin and eosin (H&E) stained sections; the cells are typically enlarged, two to fourfold larger than the surrounding normal cells, and contain eosinophilic intranuclear inclusions. The intranuclear inclusions are often eccentrically placed and surrounded by a clear halo, rendering the characteristic “owl’s eye” appearance. Also commonly, those infected cells contain variably sized, eosinophilic, intracytoplasmic viral inclusions. Identification of typical CMV inclusions confirms the diagnosis of CMV pouchitis. In cases with possible CMV-infected cells, immunohistochemical staining for CMV antigen should be performed to confirm the diagnosis. However, caution should be taken when interpreting as some plasma cells may also show dense cytoplasmic staining. Only cells with dense, dark, nuclear staining should be interpreted as CMV-infected cells. Often, examination of serially sectioned levels will differentiate a CMV-infected cell from a plasma cell with dense cytoplasmic staining. The diagnosis of CMV pouchitis can also be made by other methods such as PCR analysis of CMV DNA in formalin-fixed and paraffin-embedded pouch biopsy material [19]. Complimentarily or alternatively, blood can be analyzed for CMV viremia in patients with pouchitis [18] before the initiation of anti-CMV treatment.

Figure 1. Histology of normal pouch (a) and pouchitis (b, c). (a) This pouch biopsy shows normal histology (H&E stain, × 20). The small bowel mucosa demonstrates preserved and slender villous projection into the lumen. There are a few mononuclear inflammatory cells in the lamina propria but without expansion of the lamina propria. There is no basal lymphoplasmacytosis, neutrophilic inflammation, epithelial injury, erosion or ulceration. (b, c) This pouch specimen shows small bowel mucosa with villous blunting, erosion, chronic and active inflammation (b, H&E, × 100; c, H&E stain, × 200), features of chronic pouchitis.
Fungal pouchitis

Candida pouchitis

Rare cases of Candida pouchitis in patients with chronic antibiotic-refractory pouchitis have been reported [18], but Candida pouchitis as a separate disease entity has not been substantiated. In those reported cases, the presence of Candida species was limited to the surface of the epithelial layer in the pouch biopsy, without true tissue invasion. The use of special stains, such as Gomori methenamine silver (GMS) may help identify these organisms in the mucosal biopsy.

Histoplasma capsulatum infection

H. capsulatum infection of the ileal pouch has been reported in a patient with refractory CD of the pouch after undergoing a total colectomy for indeterminate colitis [20]. Pouch biopsy from this case showed patchy chronic active inflammation and numerous non-caseating epithelioid granulomas on H&E stain. GMS stain revealed rare 2 - 5 µm intracellular yeast forms with morphology consistent with H. capsulatum. The patient also had positive urine and blood antigen tests for H. capsulatum antigen at the time of presentation.

Infectious pouchitis by multiple identifiable agents

Rare cases of pouchitis may be induced by multiple infections. For example, synchronous CMV and C. difficile infection of the pouch has been reported as a trigger for chronic pouchitis [21].

Other secondary pouchitis

Ischemic pouchitis

Ischemia may play an important role in the pathogenesis of pouchitis as well as distal ileitis or CD of the pouch. One early study has shown that pouch hypoperfusion is associated with pouch inflammation and local septic complications [22]. One small series proposed that endoscopic asymmetric, well demarcated, inflammation of the pouch appeared to represent ischemia-associated pouchitis [23]. Biopsies from ischemic pouchitis had higher frequency of extracellular hemosiderin or hematoidin deposits. The latter are characterized by extracellular golden yellow to yellow-brown crystalline pigment deposits on H&E stained sections and are often present in the exudates or adjacent debris. Hematoidin deposits are not stained by Prussian blue, in contrast to hemosiderin deposits which are more granular and darker brown in color [23]. Of note, hemorrhagic necrosis, crypt atrophy, and lamina propria hyalinization, typical features of ischemic enteritis/colitis, are not present in ischemic pouchitis. Patients with ischemic pouchitis have some unique clinical features. For example, they are more likely to have subsequent abdominal surgeries after IPAA, more likely to have postoperative portal vein thrombi and less likely to respond to conventional antibiotic therapy. Although a diagnosis of ischemic pouchitis requires clinicopathologic correlation, identification of hemosiderin and/or hematoidin pigments may provide histologic clues to the diagnosis.

Medication-associated pouchitis

A variety of drugs can cause injury to the gastrointestinal tract, with non-steroidal anti-inflammatory drugs (NSAIDs) being the most notable. One previous report showed that the withdrawal of NSAID use alone resulted in a significant reduction in the PDAI score, PDAI endoscopy subscore and PDAI histology subscore in patients with a variety of ileal pouch disorders including chronic refractory pouchitis, acute pouchitis, cuffitis, and irritable pouch syndrome [24], suggesting that NSAIDs may induce pouchitis. This is further supported by one article reporting that sulindac use induced ulcerative pouchitis in one patient with FAP [25].

Autoimmune pouchitis

A subset of patients with idiopathic antibiotic-refractory chronic pouchitis may have no apparent etiology, respond only to corticosteroids, immunomodulators, or biologics, and demonstrate positive serum autoantibodies and concurrent immune-mediated disorders [26]. The term autoimmune pouchitis has been proposed for the chronic pouchitis in this subset of patients. Histologically, in comparison to antibiotic-responsive pouchitis and normal pouches, this proposed entity has a significant increase in deep crypt apoptosis. With the use of an apoptotic score of 6 or greater (number of apoptotic cells per 10 high-power fields), this feature had a high specificity of 98.6% and a high positive predictive value of 90.9% [26]. Other features such as villous blunting, crypt distortion, ulceration, chronic inflammation, neutrophilic inflammation, deep crypt intraepithelial lymphocytosis, pyloric gland metaplasia (PGM), and goblet cell dropout are not diagnostically useful.

Pouch Prolapse

Prolapse of the ileal pouch is a rarely reported complication of IPAA with an overall incidence of 0.3% [27]. Prolapse of the pouch can be full thickness or superficial mucosal prolapse, and can be flat or polypoid [28]. Patients with pouch prolapse often experience external protrusion of tissue, sense of obstructed defecation, seepage, and pain [29]. The diagnosis of pouch prolapse often requires correlation with symptomatology and pouchoscopic examination. Biopsy may show intercrypt smooth muscle fibers, disrupted and thickened muscularis mucosae, and diamond shaped crypts, with or without mild diffuse chronic inflammation in the lamina propria [28]. In severe cases, erosion/ulceration can be seen.
CD of the Ileal Pouch

CD of the pouch is a relatively newly described entity. There is no consensus on terminology for CD that occurs in patients with IPAA. Multiple terms such as “CD of the pouch”, “CD in IPAA”, “Crohn’s pouchitis”, and “CD-like condition in ileal pouches” have been used in the literature to describe the scenario of an IPAA patient who has “characteristic” features of CD, such as non-caseating, non-rupture-associated granulomas on histology, proximal small bowel involvement, and/or late-onset stricture or fistula. CD can occur in the ileal pouch constructed after proctocolectomy for presumed UC and indeterminate colitis in up to 7% of cases [30]. In contrast, patients with preoperative CD had up to 41-64% risk of developing recurrent CD in the afferent limb and/or the pouch [31, 32] and pouch failure rate was reported to be 13.3% in one large study [2].

Based on the background idiopathic IBD, two types of pouch CD have been recognized. CD of the pouch diagnosed in CD patients with presumed or inadvertently diagnosed as UC or indeterminate colitis (IC) upon re-reviewing of the colectomy specimens may be classified as early-onset CD of the pouch. In contrast, CD of the pouch developing de novo in patients after IPAA for UC and with a confirmed lack of evidence of CD upon histopathologic reassessment of the proctocolectomy specimens may be classified as late-onset CD of the pouch or de novo CD of the pouch [33, 34]. Although outcomes in patients with delayed diagnosis are worse, there is still a 50% pouch retention rate at 10 years with good functional status of the pouch [33].

The diagnosis of pouch CD may be solely based on clinical features which include ulceration of the small bowel including the afferent limb without diffuse pouchitis persisting after at least 4 weeks of antibiotics. Stricture formation in the small bowel (including afferent limb), the pouch or pouch inlet combined with ulceration or inflammation in patients not taking NSAIDs was also regarded as CD. CD was also diagnosed in patients with fistulation following ileostomy closure. Of note, ulceration along staple lines should not be considered as CD. In addition, patients developing a complication that might be indicative of CD (for example, fistula formation) occurring < 3 months after ileostomy reversal should not be considered as CD on this basis alone [30]. Based on clinical features, CD of the pouch may be subclassified as inflammatory phenotype, fibrostenotic phenotype, and fistulizing phenotype [35].

Mucosal biopsy with histologic evaluation of the pouch only has a limited value in diagnosing CD of the pouch. The presence of non-caseating epithelioid granulomas, either in the lamina propria, submucosa, or in lymphoid aggregates, suggests a diagnosis of CD (Fig. 2). Unfortunately, only 10-12% of patients with known CD of the pouch have granulomas on mucosal biopsy [35]. Granulomas secondary to crypt rupture or giant cell response to foreign material in biopsies taken from the suture line should not be used as evidence of CD. Further it is not clear whether isolated or “incidental” granulomas on biopsy should be treated or be offered pharmaceutical prophylaxis [34]. Other than granulomas, CD of the pouch does not exhibit specific histologic features. In two small studies, the presence of PGM was suggested to be a potential histologic marker for diagnosing CD of the pouch [36, 37]; however, one other study showed that PGM can be found in both chronic antibiotic-refractory pouchitis and CD of the pouch and thus it cannot be used to diagnose CD of the pouch [38]. In the setting of IPAA, CD can also occur in any part of the gastrointestinal (GI) tract in addition to the distal afferent limb, pouch body, anal-transitional zone (ATZ), or perianal region, thus performing esophagogastroduodenoscopy with duodenal and/or stomach biopsy may yield additional information that supports the diagnosis [39].

Features of the common inflammatory disorders of the ileal pouch are summarized in Table 1.

Pouch Neoplasia

Pouch neoplasia includes a spectrum of neoplastic changes from dysplasia to adenocarcinoma. The risk of neoplasia in patients with UC and IPAA is small with a cumulative incidence at 5, 10, 15, 20, and 25 years of 0.9%, 1.3%, 1.9%, 4.2%, and 5.1% and this risk is not eliminated by mucosectomy [40]. In patients with IPAA after proctocolectomy for IBD, preoperative colorectal neoplasia increases risk for pouch neoplasia [40]. Currently, there is no recommendation regarding the surveillance for neoplasia in the pouch. In some institutes, the majority of patients undergo surveillance endoscopy every 1 - 3 years at the discretion of the treating IBD specialist or colorectal surgeon. The endoscopic procedure includes inspection of the cuff or anal transitional zone (ATZ), pouch body, and afferent limb. During a typical diagnostic and/or surveillance pouch endoscopy, 2 - 6 pieces of biopsy are taken randomly from the afferent limb, ATZ and the pouch body and submitted in separate containers. Further, any abnormal mucosal area, such as polyps, deep ulcers, and stricture, needs to be sampled and submitted separately [40].

Histologic evaluation of surveillance biopsies plays an essential role in the management of patients with pouch neoplasia. Every pouch surveillance biopsy should be evaluated using
the same criteria used for the surveillance colonic biopsy from patients with IBD. In the landmark paper by Riddell et al (1983) [41], a standardized classification of dysplasia was proposed for the evaluation of biopsy specimens from patients with IBD. The pathology report for pouch surveillance biopsy should clearly state the presence or absence of dysplasia. While most cases can be readily classified either as negative for dysplasia or positive for dysplasia, the authors acknowledged a category of “mucosa indefinite for dysplasia” defined as “epithelial changes impossible to be classified as either unequivocally negative or positive for dysplasia”. Like colonic mucosal biopsies from IBD, the diagnosis of pouch dysplasia is based on the presence

| Entity | Duration of symptoms | Clinical response to antibiotic treatment | Pre-pouch ileum | Pouch | Rectal cuff | Other features |
|--------|----------------------|------------------------------------------|----------------|-------|-------------|---------------|
| Pouch inflammation of no clinical significance | N/A | Should not be treated with antibiotics | +/- | Variable degree of acute and chronic inflammation | +/- | Asymptomatic |
| Acute pouchitis | Acute onset, < 4 weeks symptom duration | Antibiotic- response | Variably involved | Acute inflammation, crypt abscess, chronic inflammation (+/-) | +/- |
| Chronic pouchitis | > 4 weeks symptom duration and/or > 3 attacks in a 12-month period | Antibiotic- response, antibiotic-dependent, or antibiotic-refractory | +/- | Acute inflammation, crypt abscess, chronic inflammation, PGM (+/-) | +/- |
| Secondary infectious pouchitis | Variable | Antibiotic-refractory* | +/- | Acute inflammation, crypt abscess, chronic inflammation (+/-), PGM (+/-), granulomatous inflammation (in some fungal infection) | +/- | Viral inclusion, fungal organisms on special stain, or positive stool C. difficile toxin results |
| Ischemic pouchitis | Variable | Antibiotic-refractory | Relatively normal | Acute inflammation, crypt abscess, chronic inflammation (+/-), PGM (+/-), variable fibrosis | Relatively normal | Asymmetric and well demarcated inflammation of the pouch by endoscopy, hematoidin or hemosiderin deposits |
| Autoimmune pouchitis | Variable | Antibiotic-refractory | +/- | Acute inflammation, crypt abscess, chronic inflammation, villous blunting, PGM (commonly present) | +/- | Prominent deep crypt apoptosis |
| Crohn’s disease of the pouch | Variable | Antibiotic-refractory | Variably involved | Acute and chronic inflammation, crypt abscess, villous blunting, PGM (common), non-caseating granuloma (10-12% cases) | Variably involved | Stricture or ulceration away from staple lines, and/or fistula occurring ≥ 3 months after ileostomy reversal, involvement of upper gastrointestinal tract |
| Idiopathic pre-pouch ileitis | Variable | Antibiotic-refractory | Acute and chronic inflammation, villous blunting, PGM (common) | Normal | Normal |
| Cuffitis | Variable | Antibiotic-refractory | Relatively normal | Relatively normal | Chronic active colitis pattern |

N/A: not applicable. PGM: pyloric gland metaplasia. *C. difficile pouchitis may respond to anti-C. difficile antibiotic treatment.
of a combination of microscopic features, including 1) architectural alterations exceeding that resulting from repair in chronic pouchitis, and 2) cytologic abnormalities, after excluding the possibility of inflammatory and regenerative changes that may affect the mucosa in chronic pouchitis.

Epithelium negative for dysplasia

In normal pouch mucosa, the villous architecture is maintained and the crypts are regularly spaced and touch the upper edge of the muscularis mucosae (Fig. 1a). Mononuclear inflammatory cells are present in the lamina propria, but should not noticeably expand it. Neither neutrophilic inflammation nor cytologic atypia are present. In normal ATZ biopsy, the epithelium can be small bowel mucosa or normal rectal mucosa with or without anal squamous epithelium and should lack inflammation or atypia. In some cases, the intestinal mucosa may show architectural abnormalities and metaplasia (PGM in small bowel mucosa or Paneth cell metaplasia in ATZ mucosa) without obvious inflammation or atypia (quiescent colitis equivalent). Sometimes, the biopsy may show a pattern of chronic active enteritis or chronic active colitis and in these cases, the damaged and regenerative epithelium may show villous blunting (for small bowel mucosa) and loss of mucin, features also seen in dysplastic cells. However, the presence of chronic active inflammation, surface and crypt epithelial damage, and the lack of significant nuclear enlargement, hyperchromasia, and pleomorphism should allow an interpretation of negative for dysplasia in this scenario (Fig. 3). The resolving or regenerative phase of pouchitis may pose the greatest diagnostic challenge with respect to the interpretation of dysplasia as there may no longer be neutrophilic crypt injury and the crypts may show varying degrees of nuclear enlargement, hyperchromasia, and nuclear stratification. Mitotic activity may be brisk. However, the presence of at least partial surface maturation (i.e., the nuclear enlargement and hyperchromasia in the regenerative crypts do not extend onto the surface mucosa), and the increased mononuclear inflammatory cells in the lamina propria are clues to regenerative changes. In difficult cases, knowing that the patient has been recently treated for an acute episode of pouchitis may be helpful.

Epithelium positive for dysplasia

This category is subdivided into low-grade dysplasia (LGD) and high-grade dysplasia (HGD) based on the degree of architectural and cytologic abnormalities.

Positive for dysplasia, LGD

Most biopsy specimens of LGD have crypts lined by epithelium with enlarged and hyperchromatic nuclei. Nuclear stratification is present but typically confined to the basal half of the epithelium. The nuclei in LGD maintain normal polarity; that is, their long axes are perpendicular to the basement membrane. Most cases of LGD do not exhibit surface nuclear maturation; in other words, the aforementioned atypical nuclear features should involve both crypt and surface epithelium (Fig. 4). In one report of 22 patients with initial diagnosis of pouch LGD, six (27.3%) had persistence or progression after a median follow-up of 9.5 years [42].

Positive for dysplasia, HGD

HGD exhibits enlarged nuclei with marked nuclear hyperchromasia, pleomorphism, high nuclear to cytoplasmic ratio, and loss of nuclear polarity (Fig. 5). Abnormal mitotic figures
may occasionally be seen (Fig. 5b). In addition to these cyto-
nlogic/nuclear features, HGD may exhibit increased archite-
ctural complexity which may manifest as crowded or cribriform
glands or villiform/papillary surface configuration. Like LGD,
atypical nuclear features in HGD should involve both crypt
and surface epithelium. In one report of 12 patients with pouch
HGD, 25% persisted or progressed after the initial manage-
ment during a median follow-up of 5.4 years [42].

**Epithelial changes indefinite for dysplasia**

This category refers to ambiguous epithelial alterations that
cannot be classified either as negative or positive for dysplasia
with certainty. One study reported a rate of 2.3% indefinite for
dysplasia in the ATZ and pouch in a cohort of 932 patients with
IPAA for IBD [43]. Progression of IND to definite dysplasia is
not common. For example, during a mean follow-up of 19.3
months, two (of 21, 9.5%) patients progressed: one patient de-
volved LGD and one developed HGD [43]. In practice, there
are a myriad of histologic patterns that may be categorized as
indefinite for dysplasia. Most commonly, there are atypical cy-
tologic features in the setting of florid inflammation or ulcer-
ation; in this setting, it may be difficult to distinguish regenera-
tive changes from LGD or sometimes HGD. Another setting is
the presence of severe nuclear abnormalities in the crypt bases
when the surface mucosa cannot be evaluated due to ulceration
or mechanical denudation. In addition, suboptimal embedding
with tangential section or various cytologic features that arise
as effects of suboptimal staining or fixation artifacts may lead
to an interpretation of indefinite for dysplasia.

**Pouch adenocarcinoma**

Anatomic alterations resulting from the surgical procedure
make the determination of the exact origin (pouch body vs.
rectal cuff/ATZ) or cell lineage (intestine vs. anal gland) of
pouch adenocarcinoma extremely difficult. In the largest se-
ries of 12 pouch and pre-pouch adenocarcinoma cases, eight
patients had UC and four had CD including two with de novo
CD of the pouch. The mean age at IBD diagnosis was 40 years
and the mean duration from IBD diagnosis to pouch cancer
was 25.6 years. The mean duration from pouch construction
to pouch cancer was 10 years (range: 2 - 20 years) [44]. The
tumor was located in the ATZ in eight cases (out of 12, 67%),
in the pouch in one case, in the ATZ and pouch in two, and in
non-specified peri-pouch region in one case. Pouch and peri-
pouch adenocarcinoma has morphologic similarities to UC-as-
associated colonic adenocarcinoma with a comparable frequency
of tumor infiltrating lymphocytes (66.7%), lack of dirty nec-
crosis (91.6%), mucin differentiation (58.3%) (Fig. 6), signet
ring cell differentiation (25%), heterogeneity (20%), well dif-
ferentiation (25%) (Fig. 6), and poor differentiation (41.6%).
Fifty percent of pouch and peri-pouch adenocarcinomas have
Crohn-like reaction. Pouch and peri-pouch adenocarcinoma
shows expression of CK20. Approximately 73% of pouch and
peri-pouch adenocarcinoma cases had CDX2 expression and
55% CK7 expression [44]. Of note, preoperative detection rate
of pouch dysplasia in pouch surveillance biopsies in patients
with pouch adenocarcinoma is low: out of nine patients who
underwent surveillance, only three had dysplasia on surveil-
lance biopsies [44].

**Other miscellaneous benign mass/lesion and malignancies
in pouch**

Other benign mass/lesion can affect the pouch. For example,
large pedunculated inflammatory cap polyp has been reported
in the ileal pouch causing intermittent dyschezia [45]. Histol-
ogically, the polyp is composed of chronically and actively
inflamed granulation tissue with exuberant regenerative and
hyperplastic epithelium. Lymphoma arising in an ileal pouch
after immunomodulatory therapy for IBD and squamous cell
carcinoma of the ileal pouch have been reported [40, 46].

**Cuffitis and Pre-Pouch Ileitis**

**Cuffitis**

During the stapled restorative proctocolectomy, a 1.5 - 2.0
cm cuff of diseased columnar epithelium remains at the location proximal to the ATZ [47] and is at risk for developing symptomatic inflammation. Cuffitis has been conventionally considered a form of residual UC; it is defined as inflammation of the rectal cuff in the area between the anastomosis and dentate line on endoscopy and histology, with or without minimal inflammation of the pouch body [48]. It has histologic features similar to UC such as crypt distortion, crypt abscess, and neutrophilic and mononuclear inflammatory cell infiltration. According to the response to 5-aminosalicylate (5-ASA)/steroid response, it can be classified as 5-ASA-responsive, -dependent, and -refractory cuffitis [48]. In patients with 5-ASA/steroid-refractory cuffitis, subsequent investigation may reveal evidence of CD (32.8%) or surgery-associated ATZ complications (24.1%).

Pre-pouch ileitis

Inflammation in the neo-terminal ileum proximal to the pouch is observed in patients after restorative proctocolectomy, with or without association with pouchitis. Pouchitis-associated pre-pouch ileitis typically responds to antibiotic therapy. However, pre-pouch ileitis in the presence of normal pouch typically does not respond to antibiotics and is considered as idiopathic pre-pouch ileitis. Preoperative backwash ileitis has been reported to be a risk factor for the development of pre-pouch ileitis in one study [49]. Biopsy from the neo-terminal ileum in patients with pre-pouch ileitis shows a pattern similar to pouchitis including variable degrees of villous blunting, chronic inflammation in the lamina propria, and neutrophil-mediated epithelial injury including variable degree of erosion/ulceration [49]. Crypt rupture-associated granulomas may be seen. Patients with idiopathic pre-pouch ileitis can develop strictures despite medical therapy and may require surgical resection and endoscopic dilatation of the stricture [49]. Other causes of pre-pouch ileitis include NSAIDs-related injury and ischemia induced by surgery [30, 49]. In cases with ischemia as a potential etiology, there may be stricture, submucosal fibrosis, and capillary ectasia [49]. In some institutions, ulceration of the small bowel, including the afferent limb, without diffuse pouchitis which persists after at least 4 weeks of antibiotic therapy may prompt a diagnosis of pouch CD [30].

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Figure 6. Pouch adenocarcinoma. This adenocarcinoma is well differentiated, contains abundant extracellular mucin, and invades the muscularis propria (a, H&E stain, × 20; b, H&E stain, × 40).
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