Association of hidradenitis suppurativa with Crohn’s disease

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Abstract

Hidradenitis suppurativa (HS) is a chronic inflammatory skin disorder characterized by recurrent nodules, abscesses, and sinus tracts. Crohn’s disease (CD) is characterized by inflammation of the entire digestive tract and belongs to the group of inflammatory bowel diseases, and there are many extraintestinal manifestations, among which hidradenitis suppurativa is one of the rare extraintestinal manifestations. There appears to be a strong association between CD and HS based on clinical and histological similarities (sinus tract development, granulomatous inflammation, and scarring), intersections in pathogenesis (genetic loci, immune dysregulation mechanisms, and microbiome changes), and commonality in treatment. In this review, we summarize recent studies on the association between HS and CD.

Key Words: Hidradenitis suppurativa; Crohn's disease; Epidemiology; Clinical features; Pathogenesis; Treatment

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**INTRODUCTION**

Hidradenitis suppurativa (HS), known as acne inversa, is a chronic inflammatory skin disorder characterized by recurrent nodules, abscesses, and sinus tracts. HS most commonly affects the axillary, inguinal, and anogenital regions. The course of the disease can persist for decades.

The chronic and recurrent inflammation of HS, which is similar to that in Crohn’s disease (CD), finally leads to fistula and sinus tracts. Ostlere et al.[1] first noticed a high risk of HS in patients with CD in the 1990s. Subsequently, several reported cases indicated similar pathologic features[2,3] and genetic susceptibility[4] between HS and CD. Later, patients with HS with simultaneous CD symptoms were significantly relieved by treatment with anti-tumor necrosis factor-alpha inhibitors (anti-TNF-α). Anti-TNF-α therapy not only promoted great progress in the treatment of HS but also manifested the possible common pathogenesis of HS and CD[5]. In addition, some studies found that both HS and CD shared the same immune dysregulation mechanism, such as significantly increased interleukin-1 (IL-1), IL-6, IL-17, IL-23, and TNF[6,7]. According to similar clinical and histological characteristics, as well as the intersections of pathogenesis and treatment, it is widely accepted that there may be a strong association between CD and HS. In this review, we summarize the recent studies focused on the association between HS and CD.

**Epidemiology**

The incidence of HS ranges from 0.03% to 4%[8-11]. However, due to the sparse clinical epidemiological data and misdiagnosis of HS, the actual incidence may be higher. With the improvement in diagnosis, recent research data have shown that the incidence of HS has been increasing to 10 per 100000[12].

Numerous studies have indicated that patients with CD have a significantly higher risk of HS than ordinary people. In a population-based cohort study from Minnesota, the incidence of HS among 679 patients with inflammatory bowel disease (IBD) was 1.2%. The relative risk of HS in patients with IBD is 9 times that of the general population, while the risk of HS in patients with CD is higher than that of ulcerative colitis (UC) patients[13]. Three cohort studies from the Netherlands revealed that the incidence of HS in patients with CD was 17% (17/102), 15% (96/634), and 26% (181/688)[14-16]. Interestingly, the literature results showed that the incidence of HS in patients with CD was higher than that of CD in patients with HS. According to Garg et al[17], the incidence of CD in patients with HS was 2.0% (1025/51340).

In addition, HS with CD is more common in women and blacks. A large cohort study indicated that female sex was the most independent risk parameter for HS formation in IBD (odds ratio [OR] = 3.494, P < 0.001)[16]. Kamal et al[18] retrospectively analyzed 18 patients with HS combined with IBD (CD vs UC = 15:3). Among the 15 patients with CD, 11 (73%) were female and 9 (60%) were black. However, related studies remain inadequate. Other future studies are needed to further explore the risk parameters.

**ETIOLOGY AND PATHOLOGY**

At present, the pathogenesis of HS and CD remains unclear. It is believed that HS is a multifactor-related systemic inflammatory disease that may be caused by autoimmune inflammation[19]. Studies have shown that HS and CD carry similar risk factors, immune dysregulation mechanisms, genetic loci, and microbiome changes.

It is accepted that smoking increases the risk of CD and reduces treatment effects. Recent studies have also found a strong association between smoking and HS. Garg et al[20] retrospectively measured the prevalence of newly diagnosed HS in nearly 4 million smokers over three years. Among tobacco smokers, the overall odds of new HS diagnosis was increased by 91% (OR = 1.91, 95%CI: 1.85–1.97) compared with...
nonsmokers ($P < 0.001$). The potential pathogenic effects of smoking in HS include epidermal hyperplasia and follicular plugging$^{[21]}$; neutrophil chemotaxis$^{[22]}$; TNF-α secretion$^{[23]}$; and induction of Th17 cell differentiation in keratinocytes$^{[24]}$. Prospective cohort studies have shown that smoking cessation is the safest and least expensive option for HS. However, it is difficult to change a smoking habit according to clinical results.

The TNF-α and IL-23/Th17 signaling pathways were associated with HS and CD, suggesting the same immune-mediated origins (Figure 1). TNF-α, associated with many other inflammatory diseases such as psoriasis and IBD, is secreted by innate and adaptive immune cells. Several studies have shown that TNF-α increases in the serum and lesion tissues of patients with HS. As an effective therapeutic target, adalimumab was the first approved anti-TNF-α agent for the treatment of HS. Similarly, increased proinflammatory IL-23/Th17 cells are associated with many chronic inflammatory diseases. IL-12 and IL-23 were largely expressed by macrophages in the lesion skin of HS, accompanied by Th17 and CD4+ T cell infiltration. As a cell activator and inflammatory regulator of keratinization, IL-17 cells were produced in the lesions and surrounding skin of HS. It has been clinically confirmed that the IL-12 and IL-17 inhibitor ustekinumab has certain efficacy in the treatment of HS$^{[25]}$.

Few studies have focused on the correlation between HS and CD at the microbiota and gene levels. The microbiota influences the immunologic and physiologic homeostasis of the skin epithelia and gut mucosa by activating Toll-like receptors to recognize pathogens and repair damage. However, a variety of environmental factors can alter microbial balance, leading to a decrease in microbial diversity. Such alterations in the microbiota may cause immune dysregulation and susceptibility to HS and IBD$^{[26]}$. Gower-Rousseau et al$^{[3]}$ reported three cases of HS occurring in two first-degree relatives of patients with CD, which might suggest common genetic susceptibility for the two diseases. Janse et al$^{[16]}$ identified the protective gene ELOVL7 and the risk-related genes SULT1B1 and SULT1E1 in HS combined with IBD. However, these genetic associations need further exploration.

### CLINICAL FEATURES

HS, with the characteristics of chronicity and recurrence, usually involves the axilla, inguina, perianal area, perineum, buttocks, and the area under the breast fold with typical skin lesions, including deep painful nodules, abscesses, sinuses, and bridge scars. Extensive sinus tracts and scars are the result of continuous disease progression and recurrent attacks. Usually, the Hurley stage is used to assess the severity of HS and guide treatment during clinical practice (Table 1)$^{[27,28]}$: Stage I (mild)-abscess (single or multiple) without sinus tracts and cicatrization/scarring; stage II (moderate)-recurrent abscesses with tract formation and cicatrization, single or multiple, and widely separated lesions; stage III (severe)-diffuse or near-diffuse involvement or multiple interconnected tracts and abscesses across the entire area.

Sometimes, it is difficult to differentiate perianal fistulizing CD (PFCD) from HS only by clinical features and pathological manifestations. Both HS and PFCD have abscesses or sinus tracts with perianal pain, redness, itching, bleeding, and increased purulent secretions. HS lesions are usually far from the anal canal and rectum. Most fistula lesions do not extend to the dentate line of the anal canal (no apocrine glands at or above the dental line). These may be helpful for identification of HS. On the other hand, patients with PFCD may have gastrointestinal symptoms or unexplained biological abnormalities such as abdominal pain, diarrhea, anemia, hypoferritinemia, and elevated C-reactive protein (CRP). This may put doctors on the alert and distinguish them from HS. However, some patients may have fistulas connecting to the anal canal due to the apocrine glands at the distal end of the anal canal. Approximately 17%-40% of patients with CD have perianal HS, where symptoms often overlap with PFCD (Figures 2 and 3). In addition, the histological features of CD and HS may all be granulomas and lymphatic follicles. Hence, imaging assessment, especially magnetic resonance imaging (MRI), is extremely important. Monnier et al$^{[29]}$ compared pelvic MRI characteristics of 23 patients with HS and 46 patients with CD with anoperineal disease. The results showed that the specific characteristics of HS mainly included a predominance of the ‘absence of features’ in the perianal area, the absence of rectal wall thickening, and bilaterality of features. Other features included subcutaneous edema, less involvement of the anal sphincter, and sinus passages without connection to the anorectal canal.
Table 1 Hurley stages

| Grade | Clinical manifestations |
|-------|-------------------------|
| I     | Abscess (single or multiple) without sinus tracts and cicatrization/scarring |
| II    | Recurrent abscess with sinus tracts and scarring, single or multiple widely separated lesions |
| III   | Diffuse or almost diffuse involvement, or multiple interconnected sinus tracts and abscesses across the entire area |

Figure 1 Common immune mediated origins of hidradenitis suppurativa and Crohn’s disease. DC: Dendritic cell; CD: Crohn’s disease; Th: T helper cell; IL: Interleukin; TNF: Tumor necrosis factor; TGF: Transforming growth factor; IFN: Interferon; HS: Hidradenitis suppurativa. Figure courtesy of Dr. Bo-Lin Yang.

Figure 2 A 40-year-old man with perianal Crohn’s disease and lumbar hidradenitis suppurativa, Hurley stage III. Photo courtesy of Dr. Bo-Lin Yang.

Patients with simultaneous CD and HS usually present with severe pathogenetic conditions and increased colonic involvement of CD. Kamal et al retrospectively analyzed 15 patients with CD with HS[18]: 47% of patients presented with colonic CD, 53% with ileocolonic CD, there were no cases of isolated ileal disease or upper gastrointestinal CD, and 67% with perianal lesions. The average age at HS diagnosis was 34 years old. The lesion sites of HS were axillary (53%), inguinal (47%), and perianal/perineal (73%). Thirteen (93%) patients had Hurley II or III disease. One patient developed squamous cell carcinoma.

TREATMENT

Like CD, HS is incurable and few treatments are available to maintain long-term remission, which imposes a huge medical and social burden. Treatment is even more difficult when patients also have CD. Traditional HS therapies include antibiotics,
hormonal therapy, and especially surgical therapy. However, the surgical wounds of patients with CD combined with HS are usually difficult to heal. Fortunately, a variety of drugs and surgical methods have been shown to alleviate the disease and prevent new lesions with the rapid clinical application of biological agents. Thus, systemic drug therapy should be used first. Steroids are not recommended in the acute phase of CD-related anal fistula and HS because of limited efficacy and increased risk of infection. Once local sepsis is effectively controlled, the application of immunosuppressive agents and biological agents needs to be evaluated. On the basis of effective control of active intestinal lesions, surgical treatment can be considered.

**General treatment**

Smoking and obesity are strongly correlated with the severity of HS. Smoking is a risk factor for CD onset and recurrence. It is strongly recommended that patients with HS quit smoking. Loose clothing is preferred to avoid mechanical stress and friction[30,31].

**Drug therapy**

**Antibiotics:** Antimycobacterial therapy has shown no effects on induction and maintenance of remission or mucosal healing in patients with CD and should not be used as primary therapy[32]. There are currently no high-quality data on the treatment of HS with antibiotics, especially patients with CD and HS. Antibiotics are routinely used for the treatment of infection. The combination of clindamycin (300 mg, 2/d) and rifampicin (300 mg, 2/d) is currently used by clinicians. In a prospective study, clindamycin and rifampicin were combined for 12 wk and the patients were followed for 1 year. Clinical response was observed in 73% (19/26) of patients at the outset but decreased to 41% (7/17) after the 1-year follow-up. The mean recurrence time was 4.2 mo. The most common side effects were diarrhea and nausea[33]. Although many other antibiotic application studies have been carried out clinically, antibiotics are generally not a practical solution to HS.

**Biological agents:** The application of biological agents has provided new therapeutic approaches and significantly reduced the incidence[34]. TNF-α, as the initiating factor of inflammatory progression, is mainly concentrated in the serum and skin lesions of patients with HS and is significantly higher than that of normal people. Kelly et al[35] confirmed the increased levels of TNF-α, IL-17, and IL-1 in HS lesions. Despite the wide variety of inflammatory cytokines associated with the pathogenesis of HS, anti-TNF-α therapy can effectively control most proinflammatory cytokines.

Adalimumab: As an effective therapeutic target, adalimumab is the first and only anti-TNF-α agent approved by the Food and Drug Administration for the treatment of moderate to severe HS[36,37]. The standard regimen was 160 mg at week 0, 80 mg at week 2, and then 40 mg weekly by subcutaneous injection. The efficacy of adalimumab in the treatment of HS reported in the literature varies greatly. In an open clinical trial, ten patients with HS received adalimumab treatment at week 0 at a dose of 160 mg, 80 mg at week 1, and then 40 mg every 2 wk (every other week, EOW) for 12 wk. The primary endpoint was HiSCR-50 (the counts of abscesses and inflammatory nodules...
were reduced by 50% compared with baseline). At the end of the study, no patients were considered to be responsive to the treatment, and four participants quit due to lack of efficacy[38]. However, a double-blind, placebo-controlled randomized clinical trial (RCT) showed that 21 patients receiving adalimumab [80 mg at baseline and 40 mg every other week (EOW) for 12 wk] had better efficacy than the placebo group. The efficacy endpoint was a change in Sartorius score and Hurley stage at 12 wk. The most significant reduction in the Sartorius score was observed at the 12th week (P = 0.07)[39]. In general, adalimumab is more effective for the treatment of severe HS than ustekinumab[45].

Infliximab (IFX): The efficacy for HS was first observed in patients with CD treated with IFX[5]. An RCT involving 38 patients determined the efficacy of IFX in the treatment of HS, with standard treatment regimens of 0, 2, and 6 wk of 5 mg/kg followed by maintenance therapy every 8 wk for 22 wk. The primary endpoint was that the Hidradenitis Suppurativa Severity Index (HSSI) score decreased by 50% from baseline after treatment. The results showed that the HSSI, erythrocyte sedimentation rate, CRP, and pain were significantly relieved. Among 15 patients receiving IFX treatment, the HSSI decreased by more than 50% in 26.7% of patients, 60% of patients had a 25%-50% decrease, and 13.3% of patients had less than 25% decrease. The HSSI improvement was less than 25% in most patients who received placebo (88.9%)[44]. A comparative study of IFX and adalimumab showed that IFX had better assessment indicators, including the Dermatology Life Quality Index, clinical evaluation, and efficacy duration. This study indicated that IFX might be more suitable for the treatment of severe HS than adalimumab[45].

Ustekinumab: Studies have shown that specific genetic variants of IL-12 and IL-23 receptor subunits are associated with severe HS. In a prospective study, 17 patients with HS received ustekinumab (≤ 100 kg: 45 mg; > 100 kg: 90 mg) and were followed for 40 wk. Twelve (70%) patients completed the treatment. At the 40th week, 8 (47%) patients obtained a clinical response (total abscesses and inflammatory nodule counts were reduced by 50%; abscesses or drainage fistula counts did not increase); 14 (82%) had a modified sartorius score that was moderately improved[25]. In 2019, Cline et al[46] reported that a 39-year-old woman with CD was unable to effectively control HS with IFX, adalimumab, and azathioprine treatment. After a combination of adalimumab (80 mg loading dose followed by 40 mg/wk maintenance) and ustekinumab (90 mg/8 wk), HS was effectively controlled for the first time at the follow-up 1 mo later.

**Surgical treatment**

Chronic sinuses and scars formed by HS usually require surgical intervention. A systematic review[47] showed the recurrence rate following various surgical procedures for HS: The total recurrence rate of extensive resection was 13%, direct suture was 15%, transfer flap was 8%, and skin graft was 6%. The local resection recurrence rate was 22%, and the window drainage recurrence rate was 27%. Although surgery has removed a large amount of tissue, it cannot prevent recurrence.

**Incision and drainage**: Incision and drainage are not effective in treating HS. When fluctuating abscesses occur locally in patients with HS, incision and drainage can quickly relieve pain. Some patients may be cured, but the recurrence rate is quite high. Incision and drainage are usually ineffective for inflammatory nodules and sinus tracts. For patients with active CD with HS, simple incision and drainage is a relatively rational treatment that can control local inflammation, avoid sepsis, and provide opportunities for the application of steroids, immunological agents, or biological agents.

**Deroofing (local or extensive incision)**: Deroofing (local or extensive incision) can be used in the treatment of moderately severe HS (Hurley stage II or III). Under local or regional anesthesia, inflammatory nodules or sinus passages are fully incised. The skin covering the surface of each sinus tract should be completely open to explore additional sinus openings at its base and edges. The granulation and necrotic tissue in the open inflammatory nodules and sinus passages are cleared with coarse gauze or a curette. The wound should be completely open or make a marsupialization for secondary healing (Figure 4).

**Extensive resection + reconstruction**: After the failure of drug treatment or conservative surgical treatment, serious HS (Hurley III) may require extensive resection of lesions. Extensive excision generally removes hair follicles and sebaceous gland tissues while preserving normal subcutaneous fat. When deeper sinus passages are
encountered, more skin with accessory organs, sinuses, and scar tissue is resected simultaneously. Wide excision with skin closure is controversial and depends mainly on the extent of excision of the lesion. According to the resected wound surface, the skin can be closed by free and direct flap sutures (usually with extension-reduction sutures), local flap transfer, and skin grafts. Although HS is not technically curable, most patients can control their clinical symptoms for a long time after extensive resection.

**Multimodal therapy**

HS usually requires multimodal therapy—a combination of surgery and drugs. Biologics could potentially augment the surgical intervention results. A retrospective study showed that a combination of surgery with biologics was significantly better than single surgical treatment for recalcitrant HS. The recurrence rates (19% vs 38.5%; \( P < 0.01 \)) were significantly lower. Disease progression (18% vs 50%; \( P < 0.001 \)) among patients with adjuvant biologic therapy was continued for at least 6 mo\cite{48}. There is no current literature regarding adverse events integrating biologic therapy and surgery in patients with HS.

Kamal et al\cite{18} analyzed the treatment of 15 patients with CD with perianal HS at Icahn Medical College from 2003 to 2013. The results were as follows: 14 patients (93%) received medication for CD, including antibiotics (100%), steroids (64%), mesalamine (71%), azathioprine/6-mercaptopurine (78%), and anti-TNFs (80%). Among the 12 patients treated with anti-TNFs [IFX (\( n = 11 \)) and adalimumab (\( n = 4 \))], 75% required an increased dose (based on weight gain or shorter intervals) and 11 (92%) exhibited a response to anti-TNF treatment. In addition, 67% of patients required surgery during treatment. These results suggest that compared with patients with simple HS, CD-associated HS has a poor response to anti-TNF treatment. A combination of surgery and medication may be necessary.

Based on the existing clinical evidence, we propose a clinical strategy for HS combined with CD (Figure 5). Meanwhile, we present a case of multimodal treatment of HS combined with CD: A 37-year-old man with CD, Hurley stage III. Due to recurrent perianal abscesses and fistulas, the patient received drainage and seton of the perianal lesion and IFX therapy after surgery. At the 1-mo follow-up, the wound healed well, and the local pain drainage, CRP, and white blood cell count were significantly reduced. Abdominal symptoms are almost relieved (Figure 6).

**CONCLUSION**

Both HS and CD can influence the quality of life and daily activities of patients. According to the consistency in pathogenesis, clinical manifestations, and treatment between HS and CD, HS may be another extraintestinal manifestation of CD.

When physicians encounter patients with HS accompanied by digestive symptoms or unexplained biological abnormalities, such as anemia, hypoferritinemia, and elevated CRP, gastrointestinal examination should be performed to determine the intestinal lesions. Similarly, in patients with CD with accompanying HS-like skin lesions, especially in the perianal and perineum areas, further attention is required to differentiate HS from anal fistula.
Although there is a large amount of literature on the treatment of the two diseases, there is still no unified conclusion on the treatment of patients with HS combined with CD. Antibiotics have limited therapeutic effects on CD and HS unless the bacterial culture of the lesions is positive or some immunomodulatory antibiotics (such as tetracycline and ciprofloxacin) are applied. The application of biological agents in the treatment of HS combined with CD has opened up new treatment methods and greatly relieved symptoms. As wounds are difficult to heal, surgical treatment is usually used as an adjunct to medication in patients with HS and CD. Multimodal treatment, surgical approach, and medical treatment can help postoperative healing. This may be the main direction for the future treatment of comorbidities.
In summary, early diagnosis and timely comprehensive treatment are important for patients with CD-related HS to improve quality of life. Clinicians should be aware of this link to avoid delayed diagnosis and ensure adequate management and follow-up.

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