

ABOUT THE AUTHOR



Serre-Yu Wong, MD, PhD

Dr. Wong is a physician-scientist and Assistant Professor of Gastroenterology at the Icahn School of Medicine at Mount Sinai. She received her MD and PhD degrees from the NYU School of Medicine and completed her residency and gastroenterology fellowship at the Icahn School of Medicine at Mount Sinai. Dr. Wong's research integrates host-microbe science, cross-disciplinary care, and global partnership to improve outcomes and quality of life for patients with perianal fistulizing Crohn's disease. She co-founded the Mount Sinai multidisciplinary perianal Crohn's disease clinic and is an active member of the international TOpClass perianal consortium.

Affiliations: Assistant Professor, Division of Gastroenterology, Icahn School of Medicine at Mount Sinai, New York, NY

Isolated Perianal Fistulas: When and How Should I Investigate for Inflammatory Bowel Disease?

Serre-Yu Wong, MD, PhD

Key Takeaways:

- Approximately, 5–10% of all perianal fistulizing Crohn's disease (PFCD) patients will have isolated PFCD. High or complex tracts, multiple internal openings, chronicity, and refractoriness to treatment—along with patient factors—should raise suspicion for PFCD (isolated or not).
- A negative initial luminal evaluation does not exclude CD — surveillance is key. Up to 25% of patients presenting initially with isolated complex fistulas develop luminal CD over time (median 2.5 years). Periodic reassessment with imaging, endoscopy, and symptom monitoring is critical to avoid missed or delayed diagnosis.
- Diagnosis and management of isolated PFCD requires a multidisciplinary, patient-centered approach. TOpClass criteria offer practical diagnostic guidance using clinical, radiologic, and histologic features. For patients with significant symptoms and complex isolated PFCD, anti-TNF therapy may be considered, though evidence is limited and optimal duration remains unclear.

Isolated Perianal Fistulas in Context

Perianal fistulas are a challenging manifestation of Crohn's disease (CD), affecting approximately one in five patients.^{1,2} Perianal fistulizing Crohn's disease (PFCD) is associated with a complex disease course, distinct symptom burden, frequent need for surgical intervention, reduced quality of life, and increased healthcare utilization and costs.³⁻⁵ Timely recognition and diagnosis are critical. Management strategies, both medical (e.g., anti-tumour necrosis factor (TNF) agents as first-line therapy) and surgical, differ significantly from those used for luminal CD alone and may prevent disease progression.⁵⁻⁸

Most PFCD cases present concurrently with or after a diagnosis of luminal CD.^{9,10} However, in approximately 10% of patients, perianal fistulas appear in the absence of luminal inflammation.¹¹ Of these patients, we estimate that one-quarter will eventually manifest luminal CD, while 5–10% will remain as isolated PFCD.^{10,11} Given that over 90% of perianal fistulas without luminal disease are cryptoglandular in origin, distinguishing PFCD in this context is diagnostically challenging.¹²

Cryptoglandular fistulas typically exhibit a simple anatomy—superficial, low-lying tracts with minimal sphincter involvement—and are more likely to heal.^{2,12} In contrast, CD-related fistulas are often more complex, originating higher in the anal canal or rectum, with branching or multiple tracts, and are commonly refractory to standard treatment.^{2,12} Nonetheless, overlap exists: cryptoglandular perianal fistulas can be complex, and CD perianal fistulas can be simple. Importantly, no objective test currently exists to definitively distinguish CD-related from cryptoglandular fistulas.¹³ This raises an important question: when—and how—should we evaluate for underlying inflammatory bowel disease (IBD)?

Evaluate the Nature of the Fistula

The first step in assessing a patient with an isolated perianal fistula is to carefully evaluate the nature of the perianal disease itself (**Table 1**). Features that should raise concern for PFCD include fistulas that originate high in the anal canal or rectum, have multiple internal openings, exhibit branching morphology, or present as multiple discrete fistulas. In addition to anatomy, fistula behaviour can also signal risk: fistulas that are chronic, recurrent, or refractory to treatment

may be more likely associated with CD. The presence of other forms of perianal disease—such as strictures, ulcers, or fissures—further supports this suspicion, provided there are no alternative explanations such as infection, prior obstetric injury, or iatrogenic causes (e.g., from cancer-related procedures). Taken together, the anatomic complexity, clinical course, and associated perianal findings should all be considered in evaluating for potential PFCD.^{13,14}

Assess Patient-level Risk Factors for CD

Beyond local findings, patient-level factors are essential in determining the likelihood of underlying IBD (**Table 2**). Younger age at fistula diagnosis, particularly under age 40, has been associated with an increased risk of CD in some studies.^{11,13} A thorough clinical history should explore both current and past gastrointestinal symptoms, prior perianal disease, and any autoimmune or immune-mediated conditions, including extraintestinal manifestations of IBD and comorbidities such as hidradenitis suppurativa.^{11,13,14} A detailed surgical history, including intestinal and perianal operations, as well as a family history of IBD, can provide further diagnostic clues.

During the physical examination, clinicians should assess for signs commonly associated with IBD, including ophthalmic and oral findings, and perform a comprehensive perianal exam to identify non-fistulizing manifestations such as skin tags, ulcers, or fissures. In selected patients, fecal calprotectin may serve as a useful adjunct.¹³ While a normal result does not exclude CD in patients with high clinical suspicion, an elevated calprotectin level may prompt further evaluation in those with a lower pre-test probability of CD.

Comprehensive Luminal Evaluation

Once the decision is made to evaluate for CD (**Table 1**), the diagnostic workup should aim to definitively confirm or exclude the presence of luminal disease. This distinction matters: if CD is diagnosed, anti-TNF therapy is recommended as the first-line biologic treatment.

Ileocolonoscopy with segmental biopsies is the cornerstone for evaluating luminal disease—even in areas that appear endoscopically normal, as histologic inflammation may precede visible disease. We have observed cases of isolated perianal fistulas wherein we found histologic

Fistula characteristics	Other patient characteristics
Origin high in anal canal or rectum	Age <40 at fistula onset
Multiple internal openings	Family history of IBD
Branching or multiple tracts	IBD-related extraintestinal manifestations
Chronic, recurrent, or refractory course	Coexisting autoimmune or immune-mediated inflammatory diseases
Presence of non-fistulizing perianal disease (e.g. strictures, fissures, ulcers)	Prior intestinal or perianal surgeries
	Recurrent oral or genital lesions

Table 1. Fistula and patient characteristics to evaluate for when considering whether to evaluate for CD in patients who present with isolated perianal fistula; *courtesy of Serre-Yu Wong, MD, PhD.*

TOpClass consensus criteria for isolated perianal Crohn's disease
The following findings are sufficient for considering diagnosis of isolated perianal CD: • Histologically-confirmed disease: epithelioid granuloma in fistula or surrounding perianal tissue (excluding cryptolytic or foreign-body granulomas) • Crohn's perineum: anorectal stricture or inflammatory fissures or ulcers in the absence of another cause Alternatively, consider isolated perianal Crohn's disease if score ≥ 5 based upon: Major criteria (3 points each): • Advanced fistula complexity • Family history of IBD in 1st or 2nd degree relative • Confirmed diagnosis of IBD-related extraintestinal manifestation or orofacial granulomatosis Minor criteria (1 point each): • Unconfirmed diagnosis of IBD-related extraintestinal manifestation (potential, past, or prior) • Suspected oral or genital CD • Presence of hidradenitis suppurativa • Minor perianal disease (single >1 cm edematous skin tag, ≥ 3 small skin tags, non-fistulizing perianal skin inflammation, or natal cleft ulceration) • Recurrence following fistula repair or lay-open procedure with curative intent

Table 2. TOpClass consensus criteria for isolated perianal Crohn's disease in patients presenting with perianal fistula and no luminal inflammation; *courtesy of Serre-Yu Wong, MD, PhD.*

evidence of inflammation that later manifested clinically and endoscopically as luminal CD. Other modalities that can be used include video capsule endoscopy, intestinal ultrasound, and magnetic resonance enterography.¹⁵ Using a combination of these complementary tests may increase diagnostic yield, depending on the resources available at a given institution.¹⁴

Importantly, luminal disease may not be evident at initial presentation. In a case series from our institution, 25% of patients with isolated complex perianal fistulas developed luminal CD over time, with a median time to diagnosis of 2.5 years, and a range extending up to 10 years.¹¹ Therefore, a single negative evaluation should not be considered definitive. The TOpClass

consortium emphasized the need for ongoing surveillance—though no consensus was reached on the optimal surveillance interval, with recommendations ranging from symptom-guided re-evaluation to routine annual screening.¹⁴

What if No Luminal CD is Found?

Between 5–10% of patients with PFCD will remain without evidence of luminal disease.^{10,11} Historically, establishing a definitive diagnosis of isolated PFCD in such patients has not been clear. To address this gap, the international perianal disease TOpClass Consortium—a multidisciplinary panel of IBD gastroenterologists, surgeons, and radiologists—recently conducted

a systematic review and published consensus recommendations.¹⁴ While not yet fully validated, these proposed diagnostic criteria offer practical guidance for clinical use (**Table 2**). According to these guidelines, the presence of diagnostic histopathologic features in fistula tissue or the surrounding area, as well as severe associated perianal disease, can independently establish a diagnosis of isolated PFCD. A total score of ≥ 5 —achievable through either two major criteria, one major plus two minor criteria, or five minor criteria—is considered sufficient to support the diagnosis.

Effective diagnosis and management of isolated PFCD requires multidisciplinary collaboration. While gastroenterologists typically lead the evaluation for luminal disease, colorectal surgeons often have a clinical gestalt about whether a fistula's characteristics are more suggestive of CD rather than a cryptoglandular origin.

Managing Isolated PFCD

Shared-decision making is essential for managing isolated PFCD, and patients should be informed about both the knowns and unknowns of the disease. For patients experiencing significant perianal symptoms or whose fistulas are unlikely to heal with surgery alone, a trial of biologic therapy—typically anti-TNF agents—can be considered, provided the patient is amenable. Anti-TNFs may help reduce inflammation, support fistula healing, and facilitate surgical interventions.¹⁶⁻¹⁸ However, it should be noted that the data supporting their efficacy is limited. One study, for example, reported lower remission rates in patients with complex idiopathic perianal fistula compared to those with confirmed PFCD.¹⁹

If a patient shows a positive response to optimized anti-TNF therapy, this treatment

may be continued with regular monitoring. Yet, there is no consensus on the optimal treatment duration after clinical and radiologic remission—recommendations range from 3 months to lifelong therapy, reflecting the lack of data in this area. If there is no therapeutic response, anti-TNF therapy should be discontinued.¹⁴ At that point, the diagnosis of isolated PFCD should be re-evaluated, and consideration given to initiating a second-line biologic.

Conclusion

Perianal fistulas without overt evidence of luminal CD present a clinical dilemma. While most are cryptoglandular in origin, a minority herald PFCD. Identifying these cases is important, as their medical and surgical management differs substantially from that for idiopathic fistulas. Comprehensive screening and luminal evaluation—including histology, imaging, and ongoing surveillance—are essential components of care. Yet, questions remain: Is isolated PFCD a distinct clinical entity? What constitutes the best treatment strategy? Further research is needed to clarify its natural history, guide treatment, and improve outcomes for this enigmatic subset of IBD.

Correspondence

Serre-Yu Wong, MD, PhD

Email: serre-yu.wong@mountsinai.org

Financial Disclosures

S.W.: Advisory board: Bristol Myers Squibb;

Research grant: Takeda and Eli Lilly

References

1. Tsai L, McCurdy JD, Ma C, Jairath V, Singh S. Epidemiology and natural history of perianal Crohn's disease: a systematic review and meta-analysis of population-based cohorts. *Inflamm Bowel Dis*. 2022;28(10):1477–1484. doi:10.1093/ibd/izab287
2. Zhou Z, Ouboter LF, Peeters K, Hawinkels L, Holman F, Pascutti MF, et al. Crohn's disease-associated and cryptoglandular fistulas: differences and similarities. *J Clin Med*. 2023;12(2):466. doi:10.3390/jcm12020466
3. Adegbola SO, Dibley L, Sahnan K, Wade T, Verjee A, Sawyer R, et al. Burden of disease and adaptation to life in patients with Crohn's perianal fistula: a qualitative exploration. *Health Qual Life Outcomes*. 2020;18(1):370. doi:10.1186/s12955-020-01622-7
4. Panes J, Reinisch W, Rupniewska E, Khan S, Fornis J, Khalid JM, et al. Burden and outcomes for complex perianal fistulas in Crohn's disease: systematic review. *World J Gastroenterol*. 2018;24(42):4821–4834. doi:10.3748/wjg.v24.i42.4821
5. Chen G, Pedarla V, Null KD, Cazzetta SE, Khan QR, Schwartz DA. Health care costs and resource utilization among patients with Crohn's disease with and without perianal fistula. *Inflamm Bowel Dis*. 2022;28(6):870–877. doi:10.1093/ibd/izab198
6. Adler J, Gadepalli S, Rahman M, Kim S. Early tumour necrosis factor antagonist treatment prevents perianal fistula development in children with Crohn's disease: post hoc analysis of the RISK study. *Gut*. 2025;74(4):539–546. doi:10.1136/gutjnl-2024-333280
7. Cho WJ, Wong SY. A guide through the tunnel: updates in the approach to classification and management of perianal fistulizing Crohn's disease. *Curr Gastroenterol Rep*. 2025;27(1):46. doi:10.1007/s11894-025-00998-0
8. Hanna LN, Anandabaskaran S, Iqbal N, Geldof J, LeBlanc JF, Dige A, et al. Perianal fistulizing Crohn's disease: utilizing the topclass classification in clinical practice to provide targeted individualized care. *Clin Gastroenterol Hepatol*. 2025;23(6):914–926. doi:10.1016/j.cgh.2024.06.047
9. Molendijk I, Nuij VJ, van der Meulen-de Jong AE, van der Woude CJ. Disappointing durable remission rates in complex Crohn's disease fistula. *Inflamm Bowel Dis*. 2014;20(11):2022–2028. doi:10.1097/mib.0000000000000148
10. Schwartz DA, Loftus EV, Jr., Tremaine WJ, Panaccione R, Harmsen WS, Zinsmeister AR, et al. The natural history of fistulizing Crohn's disease in Olmsted County, Minnesota. *Gastroenterology*. 2002;122(4):875–880. doi:10.1053/gast.2002.32362
11. Sangmo L, Quraishi B, Rajauria P, Giselbrecht E, Colombel JF, Ungaro R, et al. Natural history of clinically suspected isolated perianal fistulizing Crohn's disease. *Clin Gastroenterol Hepatol*. 2025;23(4):671–673.e672. doi:10.1016/j.cgh.2024.09.016
12. Chin Koon Siw K, Engel J, Visva S, Mallick R, Hart A, de Buck van Overstraeten A, et al. Strategies to distinguish perianal fistulas related to Crohn's disease from cryptoglandular disease: systematic review with meta-analysis. *Inflamm Bowel Dis*. 2022;28(9):1363–1374. doi:10.1093/ibd/izab286
13. Munster LJ, Hanna LN, Hart AL, Tozer PJ, Buskens CJ, van der Bilt JDW. Diagnosing Crohn's disease in presumed cryptoglandular perianal fistulas: an expert Delphi consensus on early identification of patients at risk of Crohn's disease in perianal fistulas (PREFAB). *J Crohns Colitis*. 2025;19(1):jjaf002. doi:10.1093/ecco-jcc/jjaf002
14. Hanna LN, Munster LJ, Joshi S, Wendelien van der Bilt JD, Buskens CJ, Hart A, et al. Isolated perianal Crohn's disease: a systematic review and expert consensus proposing novel diagnostic criteria and management advice. *Lancet Gastroenterol Hepatol*. 2025. doi:10.1016/s2468-1253(25)00007-x
15. McCurdy JD, Weng R, Parlow S, Dawkins YM, Brar G, Oliveira L, et al. Video capsule endoscopy can identify occult luminal Crohn's disease in patients with isolated perianal fistulas. *J Crohns Colitis*. 2023;17(10):1624–1630. doi:10.1093/ecco-jcc/jjad078
16. Dige A, Nordholm-Carstensen A, Hagen K, Hougaard HT, Krogh K, Agnholt J, et al. Effectiveness of infliximab treatment of complex idiopathic anal fistulas. *Scand J Gastroenterol*. 2021;56(4):391–396. doi:10.1080/00365521.2021.1879246
17. Fiske HW, Tse CS, Al-Bawardy B, Magavi P, Konijeti GG, Mao E, et al. Clinical course of isolated recurrent, persistent complex perianal fistulas without luminal Crohn's disease: a multicenter case series of 24 patients. *Crohns Colitis*. 2024;6(4):otae065. doi:10.1093/crocol/otae065
18. Rinawi F, Greer MC, Walters T, Church PC, Ricciuto A, Langer JC, et al. Anti TNF treatment of complex perianal fistulas in children without luminal Crohn's disease: Is it an option? *J Pediatr Surg*. 2022;57(11):569–574. doi:10.1016/j.jpedsurg.2022.03.031
19. McCurdy JD, Parlow S, Dawkins Y, Samji K, Rhee GG, Oliveira L, et al. Tumor necrosis factor inhibitors may have limited efficacy for complex perianal fistulas without luminal Crohn's disease. *Dig Dis Sci*. 2020;65(6):1784–1789. doi:10.1007/s10620-019-05905-y