Cronkhite-Canada Syndrome Showing Good Early Response to Steroid Treatment

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**Case:** A 43-year-old female visited our hospital with complaints of abdominal pain and diarrhea. The patient had diffuse postprandial abdominal pain, dyspepsia, loss of taste, and repeated watery diarrhea for a duration of one month. She was in poor general condition and unable to consume regular diet. During that period, she gradually experienced hair loss, nail dystrophy of both hands and feet, and skin hyperpigmentation of both upper and lower limbs (Fig. 1). She had no remarkable previous medical, surgical, and family history; she underwent screening esophagogastroduodenoscopy seven months ago, which revealed mild gastritis, and had no history of colonoscopy. She was a non-smoker and

![Image of the Month](image.png)

**Fig. 1.** Clinical manifestation of the patient at the initial presentation. (A) Hair loss. (B) Skin pigmentation and nail dystrophy. (C) Tongue swelling and mucosal atrophy.
non-drinker. She had been running her own restaurant business for the past year and had been under a lot of stress.

The initial vital signs were stable; she appeared chronically ill. Her height was 147 cm and weight was 42 kg (body mass index, 19.4 kg/m²). The initial laboratory data were as follows: white blood cell count 8,620/μL, hemoglobin 14.7 g/dL, platelet count 308,000/μL, protein 5.8 g/dL, albumin 3.8 g/dL, blood urea nitrogen 7.1 mg/dL, creatinine 0.57 mg/dL, and C-reactive protein 0.16 mg/dL. Serum antinuclear antibody and immunoglobulin G4 (IgG4) were absent. Carcinoembryonic antigen and cancer antigen 19-9 were within normal range. Upper and lower endoscopy were performed, which revealed numerous sessile polyps with mucosal edema and hyperemia involving the stomach, duodenum, and colon, except the esophagus (Fig. 2). The size of polyps was variable; however, there was no polyp with large enough size that required endoscopic resection. Forceps biopsies revealed hyperplastic polyps in the stomach and juvenile polyps in colon, accompanied by inflammatory cell infiltration with dilated gland, cystic dilatation of crypt, and edema of lamina propria (Fig. 3). Tissue IgG4 staining result was negative. Rapid urease test result was positive. Abdomino-pelvic computed tomography and small bowel series showed no significant polyp-like lesion in the jejunum and ileum.

Based on these findings, the patient was finally diagnosed with Cronkhite-Canada syndrome (CCS), and admission for steroid treatment with nutritional support was recommended. However, the patient wanted ambulatory treatment for personal reasons. Thus, an oral corticosteroid (prednisolone, 40 mg per day) was administered in an outpatient setting. After 3 weeks, she regained appetite, sense of well-being, and formed stool, with improvement of skin hyperpigmentation.

Fig. 2. Initial upper and lower endoscopic findings. Numerous variable-sized sessile polyps with mucosal edema and hyperemia were observed in the (A, B) stomach, and (C, D) colon.
and tongue swelling; however, her hair loss continued. Considering the favorable treatment response, steroid dose was gradually tapered by 5 mg per week. Diarrhea improved after 6 weeks. Follow-up upper endoscopy and colonoscopy after 3 months from the initial diagnosis showed marked regression of gastrointestinal polyposis with remaining mild
mucosal edema and hyperemia (Fig. 4). Forceps biopsies revealed nonspecific inflammation. Hair loss, skin pigmentation, and nail dystrophy improved at 5 months after the start of steroid treatment (Fig. 5). Steroid dose was successfully tapered, and eradication of the *Helicobacter pylori* was performed. The patient remained stable without recurrence during the 9-month follow-up period from the initial diagnosis.

**Diagnosis: Cronkhite-Canada syndrome (CCS)**

CCS is a rare non-hereditary disorder characterized by 1) gastrointestinal polyposis and 2) characteristic ectodermal changes (hair, skin, nail, and tongue). Since the first report of two patients by Cronkhite and Canada, more than 400 cases have been reported worldwide, and the majority of them were in Japan. Until now, 15 cases of CCS have been reported in Korea. According to previous reports, the mean age of patients at the time of diagnosis was between late 50s and early 60s, with greater occurrence in males (male to female ratio was 1.3-1.84:1). Pathophysiology of CCS is still unclear, but some authors have suggested mental stress and physical fatigue as possible precipitating factors. Common complications are malnutrition, gastrointestinal bleeding, and infection. The overall prognosis is poor, and the mortality rate is reported to be as high as 55%, if not properly treated. Recently, the autoimmune process is considered as the main pathophysiologic factor of CCS due to the good response in steroid treatment and IgG4-positive cells in tissue biopsy. Common presenting symptoms of CCS are diarrhea, weight loss, abdominal pain, and anorexia. Characteristic ectodermal changes, such as nail changes, hair loss, and skin hyperpigmentation are present in more than 70% of cases. Polyps can involve the entire gastrointestinal tract, from the esophagus to the rectum; however, the esophagus is usually rarely involved (2-12%). Histologic findings of polyps are usually nonspecific, showing hyperplastic polyp, hamartomatous or juvenile polyp, inflammatory polyp, and tubular adenoma. At the initial presentation or during the follow-up period, gastrointestinal malignancies, such as colon or gastric cancer, are reported (15-25%). However, whether polyps in CCS possess malignant potential has been controversial due to the difficulties distinguishing between premalignant adenomas and multiple inflammatory pseudopolyps.

Given its rarity, standard treatment of CCS has not yet been established. However, steroid administration accompanied by nutritional support is known as the mainstay of treatment. Regression of polyposis and achievement of sustained endoscopic remission are the purpose for treating CCS. Watanabe et al. suggested that good early response (regression of polyposis within 6-12 months after the start of treatment or initial diagnosis) and good long-term response (maintenance of endoscopic remission over a 3-year period after initial treatment) are related with favorable prognosis. Administration of histamine H2 receptor antagonist, proton pump inhibitor, 5-aminosalicylic acid, antibiotics, and *Helicobacter pylori* eradication also showed good results in previous reports. During the follow-up period, CCS can recur even in patients with successful discontinuation of steroid and clinical remission; thus, minimal annual endoscopic surveillance is
usually recommended and maintenance treatment using a low-dose steroid, immunomodulator, and anti-tumor necrosis factor should also be considered.

In conclusion, CCS may be associated with significant complications, such as gastrointestinal bleeding, infection, and malignancy. However, early detection and proper steroid administration can achieve rapid clinical improvement and regression of polyposis, which can bring better overall outcome of CCS, as we have seen in this case. During long-term follow-up, recurrence of polyposis and CCS-related cancer can develop; thus, regular endoscopic surveillance is recommended.

REFERENCES


