Introduction

Cryptogenic multifocal ulcerous stenosing enteritis (CMUSE) is a rare condition characterised by chronic or relapsing moderate ileous episodes resulting from multiple small intestinal strictures, multiple shallow ulcers of the small bowel and beneficial therapeutical effect of glucocorticosteroids (1, 2). Japanese gastroenterologists call this syndrome chronic non-specific multiple ulcers of the small intestine (CNSU) (3). The aetiology of CMUSE has not been clarified yet and pathogenesis is still poorly understood. Some authors even doubt the real existence of this entity.

Patients with CMUSE are often referred for surgery because of symptomatic small intestinal strictures and several resections of the small bowel had to be performed. We report three cases of CMUSE diagnosed within the past 10 years at a single tertiary centre, where more than eight thousand GI endoscopies are performed per year. Review of available relevant literature is provided.

Case reports

Case 1

A 35-year-old woman was admitted to our department because of chronic diarrhoea, colicky abdominal pain, weight loss (16 kg during previous five years) and repeated moderate ileous episodes. She was cachectic (body-mass index 12.8 kg/m²). Laboratory tests showed iron-deficiency anaemia (haemoglobin 96 g/L), thrombocytosis (459.10⁹ g/L) and proteino-energy malnutrition (serum prealbumin 0.17 g/L). Abdominal ultrasound revealed strenuous small intestinal peristalsis and a large volume of fluid in the intestinal loops. Enteroctasis provided no further information. Gastroscopy and colonoscopy were normal. Push-enteroscopy found multiple shallow ulcers of the small intestine (CNSU) (3). The aetiology of CMUSE has not been clarified yet and pathogenesis is still poorly understood. Some authors even doubt the real existence of this entity.

Coeliac disease, lymphoma, Crohn's disease and vasculitis were excluded. No infective agent was found. Total parenteral nutrition and glucocorticosteroids improved her general condition including nutritional status (serum albumin 35.8 g/L, prealbumin 0.29 g/L, body-mass index 13.4 kg/m²). The patient was discharged on enteral nutrition, glucocorticosteroids (prednisone was gradually replaced by budesonide) and 5-aminosalicylates. She was followed-up at regular controls. Nine months later the patient had to be admitted because of a worsened general condition (frequent watery stools, fatigue, oedemas of lower extremities, fluidothorax and ascites). Laboratory tests revealed hypo-
kalaemia (2.2 mmol/L), hypomagnesaemia (0.51 mmol/L), low serum albumin (15.7 g/L) and prealbumin (0.10 g/L), elevated liver enzymes and high C-reactive protein (136 mg/L). Capsule endoscopy was performed and multiple ulcers of the duodenum, jejunum and ileum were found. Despite no evident stenosis at repeated preceding enteroclysis, asymptomatic capsule retention in the distal ileum occurred. Her general condition improved by means of systemic glucocorticosteroids and total parenteral & enteral nutrition within two months again. Nevertheless she was not fit for surgery. The patient died after further 7 months as a result of fulminating bilateral bronchopneumonia complicated by sepsis. Six short stenosing fibrous strictures of the small intestine and disintegrated video capsule were found at autopsy.

**Case 2**

A 60-year-old woman was referred to our department because of severe malnutrition, weight loss (10 kg per month), diarrhoea, low-grade fever, fatigue and breathlessness. Bilateral fluidothorax, pericardial effusion, ascites and severe oedemas of low extremities were found on admission. Iron-deficiency anaemia (94 g/L), leukocytosis (24.10⁹/L) and thrombocytosis (426.10⁹/L) were found. Biochemical tests showed low serum albumin (15.2 g/L),

![Fig. 1: Push-enteroscopy. Multiple shallow ulcers were found on the top of transverse folds in the proximal jejunum.](image1)

![Fig. 2: Double balloon enteroscopy. A large but shallow ulcer is seen in front of tight fibrous stricture of the distal jejunum.](image2)

![Fig. 3: Prevailing infiltration with plasmatic cells was identified in all layers of the small intestine. Immunohistochemistry, anti-CD138 staining, magnification 100x.](image3)

![Fig. 4: Increased content of collagen was found in interstitial tissue of the small bowel. Small intestinal epithelium, impaired by extensive infiltration with plasmatic cells and lymphocytes. Optical histology, haematoxilin-eosin staining, magnification 40x.](image4)
prealbumin (0.06 g/L) and high C-reactive protein (165 mg/L). Gastroscopy revealed multiple ulcers of the duodenum, shallow ulcers were also found in the caecum and ascending colon at colonoscopy. Capsule endoscopy identified segmental ulcerative involvement of the small intestine (different from that seen in Crohn’s disease). Double balloon enteroscopy showed multiple shallow ulcers of the duodenum and jejunum and multiple jejunal stenoses (Fig. 2). Non-specific inflammation of biopsy specimens was found at histology, consisting of neutrophils, eosinophils and both B and T lymphocytes. There was a marked infiltration of all layers by plasmatic cells (Fig. 3) and overproduction of collagen was recognised (Fig. 4). Flow cytometry of a small intestinal biopsy specimen revealed CD3⁺CD8⁺ T cells, no subset of CD3⁺CD8– T lymphocytes were identified. Infective aetiology, vasculitis and malignancy were excluded. Complex therapy including total parenteral & enteral nutrition and systemic glucocorticosteroids improved her general condition and laboratory parameters (albumin 34.4 g/L). During the subsequent 3 months the patient remained symptom free despite persisting multiple small intestinal stenoses at control enteroclysis (Fig. 5). Control double balloon enteroscopy confirmed healing ulcers and three tight jejunal stenoses were dilated endoscopically up to 18 mm in diameter. Replacement of prednisone by budesonide was unsuccessful, acute moderate ileus episode occurred and forced a return to prednisone. Switching of systemic glucocorticosteroids to methotrexate was tried (parallel treatment was administered for 8 weeks). However, the moderate ileus status occurred again three weeks after withdrawal of the prednisone. At this moment, corticoidependence of the disease has been definitely confirmed and a prednisone dose of 20 mg daily is used as the basic treatment.

**Case 3**

A 50-year-old woman suffered for five years from colicky abdominal pain, recurrent moderate ileus episodes and collapses during difficult defecation. Repeated colonoscopies, enteroclyses and laparoscopy were performed elsewhere without any diagnostic progress. She was referred to our department at this point. Body mass index was 19.7 kg/m², laboratory findings (including nutritional parameters) were normal. Capsule enteroscopy detected erosions and small ulcerations in the ileum. Double balloon enteroscopy followed, multiple shallow ulcers were seen in the terminal ileum. Non-specific inflammatory histology was found with neutrophils, eosinophils, plasmatic cells and increased content of collagen. Gastrointestinal infection, Crohn’s disease, vasculitis and malignancy were excluded. Budesonide (9 mg per day) was sufficient to achieve full remission. The patient remained symptom free during 9 months of subsequent follow-up.

None of these three patients took any non-steroidal anti-inflammatory drugs.

**Discussion**

We present three cases of CMUSE collected within the past 10 years at our department. The first descriptions of this rare condition probably came from the late 50’s and early 60’s (4, 10, 12, 22, 32, 41), followed by further case reports or small series (2, 3, 7, 18, 46, 48). Matsumoto et al. (33) informed that Okabe and Sakimura reported first cases of CMUSE in Japan in 1968. However, all these early reports are difficult to evaluate nowadays as diagnostic potential was limited at that time. Nowadays, owing to double balloon enteroscopy we are able to investigate the entire small intestine (9,29), take numerous biopsy specimens and utilise other advanced diagnostic methods like flow cytometry and/or immunohistochemistry. That is why several distinct entities can be newly recognised.

Only about fifty cases of CMUSE have been published so far (9, 33, 34, 37, 46, 42) but this entity has probably been considerably underdiagnosed or misdiagnosed, mostly with Crohn’s disease and non-steroidal anti-inflammatory drugs (NSAIDs) induced enteropathy. Perlemuter et al. (37) reported 12 cases of CMUSE hospitalised between 1965 and 1993. This was a retrospective analysis of medical records based on queries sent to 220 French gastroenterology departments. Despite surgery, symptoms re-occurred in seven of ten patients and recurrence of strictures was reported in four. Inflammatory infiltrate was made of neutrophils and eosinophils. Steroid therapy was effective but caused dependence (37). Chang et al. (9) collected 2 cases of CMUSE diagnosed by means of double balloon enteroscopy in 48 patients (a series from 6 university hospitals in Korea). Both individuals suffered from chronic recurrent abdominal pain. One patient was presented with recurrent melena (for 41 months), the other one was referred for surgery because of retention of a capsule endoscope in the ste-

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**Fig. 5:** Enteroclysis. Multiple persisting stenoses of the small intestine caused by fibrous strictures.
notic site. There were multiple small intestinal stenoses, shallow ulcers and mixed inflammatory infiltrate at histology (plasma cells, monocytes, neutrophils and eosinophils). Mesenteric arteriography did not demonstrate any evidence of arteritis (9).

Immunopathological pathogenesis of CMUSE is supported by the favourable therapeutical effect of glucocorticosteroids. Most of these patients become corticodependent (37, 44). All our three patients positively responded to this treatment and indeed 2 of 3 became dependent on glucocorticosteroids. The key to the solution of aetiology and understanding pathogenesis might be overstimulated production of fibrous tissue. This is a principal sign of CMUSE, responsible for chronic or relapsing moderate ileous episodes. Fibroblast proliferation can be augmented by pro-inflammatory cytokines (IL-6, IL-8, TNF-alpha), fibroblast growth factors (FGF II), granulocyte/macrophage colony-stimulating factor (GM-CSF), transforming growth factor beta (TGF-beta), platelet-derived growth factor (PDGF) (47) but also by endotoxin (lipopolysaccharide) (49). Other growth factors, such as the connective tissue growth factor (CTGF), which is secreted by fibroblasts and endothelial cells, also promote formation of fibrous tissue (47). Collagen is degraded by a family of matrix metalloproteinases that includes the collagenases. Matrix cells, neutrophils, and macrophages all secrete these proteinases. In healthy people, the degradation of collagen can be very rapid, and begins immediately after collagen is produced (1). We can hypothesise that multifocal small intestinal disturbance of collagen degradation in CMUSE might play a crucial role in pathogenesis. In CMUSE, fibrous tissue formation can be associated with low or even absent systemic inflammatory response (37).

We do not agree with Perlemuter et al. (37, 38) that CMUSE could be termed as a type of “atypical vasculitis”. Any type of vasculitis of any cause, if found, should be assigned simply as “vasculitis” not CMUSE. Small intestinal involvement can be found in Churg-Strauss syndrome (20, 27, 36), systemic lupus erythematosus (43), Buerger’s disease (30), Wegener’s granulomatosis (11), Weber-Christian disease (28) and systemic sclerosis (13). Vasculitis could be associated with multiple small intestinal ulcers and their complications (including perforation) but quite rarely with multifocal stenoses of the small bowel. Furthermore, vasculitis mostly represents a systemic involvement (kidneys, skin, joints, lungs etc.). None of our three patients revealed any extra-intestinal involvement and small intestinal histology found no signs of vasculitis. Matsumoto (33) did not reveal any arteritis in his series as well. Perlemuter et al. (38) suggested the association of CMUSE with C2 complement deficiency; however, this was not confirmed by other authors (44). Recently, Fraile et al. (16) reported an association of CMUSE with X-linked recessive reticulate pigmentary disorder. Hussey et al. (23) published a case of CMUSE as a manifestation of enterocolic venopathy.

Chronic or relapsing moderate ileous episodes resulting from multiple small intestinal strictures are a leading clinical symptom (33, 37). All our three patients suffered from such episodes. Wireless capsule endoscopy was complicated by retention of the capsule in one of our cases of CMUSE. Spontaneous disintegration of a retained video capsule was reported by our group recently (45). Multiple small intestinal fibrous strictures were previously resected surgically (44), nowadays they can be treated endoscopically by means of double balloon enteroscopy (39). Three tight tandem stenoses of one of our patients with CMUSE were solved by balloon dilatation within double balloon enteroscopy.

In differential diagnosis of CMUSE, first of all Crohn’s disease (9,15), NSAIDs-induced enteropathy (9, 35), tuberculosis and other infections of the small bowel (8, 9, 20), Behcet disease (19, 21, 24) and malignancies must be excluded. It is a well-known fact, that the majority of extra-nodal malignant lymphoma involves the gastrointestinal tract. Not only polypoid and diffuse types of this disease, but also the ulcerative one can be detected in the small bowel (50). Usually either diffuse large B cell lymphoma or mucosa-associated lymphoid tissue (MALT) lymphoma are confirmed by histological methods (31).

It is also obligatory to distinguish CMUSE from other non-frequent pathological conditions. We are convinced that CMUSE is distinct from chronic ulcerative jejunitis (14,25), collagen sprue (17) and from autoimmune or eosinophilic enteritis (9,40). Similarly, non-specific small intestinal ulcers (5, 6, 9, 26) should not be considered to be CMUSE if multiple stenoses of the small bowel are absent.

Prognosis of CMUSE remains uncertain. Patients were previously referred for surgery because of symptomatic small intestinal strictures. However, the postoperative recurrence rate is high. Matsumoto (33) reports a single case of CMUSE followed up for 40 years since 1963. This patient was operated on seven times because of recurrence of tight stenoses and several resections of the small bowel had to be performed. Nowadays, endoscopic treatment of symptomatic stenoses should be tried first. Therapy with glucocorticosteroids is a treatment of choice. However, most patients develop corticosteroid dependence.

In conclusion, CMUSE, although a rare condition affecting the small bowel, should always be considered when chronic or relapsing moderate ileous episodes result from multiple small intestinal strictures and multiple shallow ulcers of the small bowel are found (in the absence of Crohn’s disease, NSAID use or small intestinal infection). Double balloon enteroscopy enables precise diagnostic work, possible endoscopic treatment of stenoses, may obviate the need for surgery and prevent excessive small bowel resections.

The study was supported by research project MZO 00179906 from the Ministry of Health.

References


