Update on the Management of Refractory Coeliac Disease

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Abstract

True Refractory Coeliac Disease (RCD) is being currently defined as persisting or recurring villous atrophy with crypt hyperplasia and increased intraepithelial lymphocytes in spite of a strict gluten free diet for more than 12 months or when severe persisting symptoms necessitate intervention independent of the duration of the dietary therapy. Currently two categories of RCD are being recognized: type I without aberrant T-cells and type II with aberrant T-cells detected by immunophenotyping by flowcytometric analysis or immunohistology of the intestinal mucosa. Establishing the diagnosis of RCD requires exclusion of other causes of villous atrophy and malignancies that may complicate coeliac disease. In contrast to patients with a high percentage of aberrant T cells, patients with RCD type I seem to profit from an immunosuppressive treatment. In cases of RCD II with persistent clinical symptoms and/or high percentage of aberrant T cells in intestinal biopsies in spite of a corticosteroid treatment, more aggressive therapeutic schemes should be considered. However, no therapy seems to be curative in RCD II. Cladribine (2-CDA) seems to have some role in the management of these patients. More recently, high dose chemotherapy followed by autologous stem cell transplantation has been used in patients resulting in a dramatic improvement in the clinical, laboratory, histopathological and immunological parameters. This review provides an overview of the currently available diagnostic and therapeutic methods in a complicated form of coeliac disease.

Key words

Coeliac disease – refractory coeliac disease – pathogenesis – diagnosis - therapy

Introduction

Coeliac disease (CD) is a life-long inflammatory condition of the GI tract that affects the small intestine in genetically susceptible individuals. On small bowel biopsy there is a characteristic, although not specific, mucosal lesion that impairs nutrient absorption by the involved bowel. Prompt improvement of nutrient absorption and healing of the characteristic intestinal mucosal lesion is seen upon withdrawal of dietary gluten.

Non-responsive celiac disease (NCD) can be described in terms of the clinical scenario of a lack of initial response to a prescribed gluten free diet (GFD), or the recurrence of symptoms despite maintenance of GFD in a patient who responded initially to GFD (2). Although clinical improvement is usually followed by histological improvement most of the time, on occasions there is evidence for histological improvement with persistence of clinical symptoms that could be related to other causes (3,4). Clinical improvement is usually evident within the first few weeks of starting GFD; however, it might take up to two years before a complete restoration of intestinal mucosa is evident (5).

Specific definition of refractory coeliac disease (RCD) is missing in the literature. We define true RCD as persisting or recurring villous atrophy with crypt hyperplasia and increased intraepithelial lymphocytes (IEL’s) in spite of a strict GFD for more than 12 months or when severe persisting symptoms necessitate intervention independent of the duration of the GFD (6). RCD may not respond primarily or secondarily to GFD (7). All other causes of malabsorption must be excluded and additional features supporting the diagnosis of CD must be searched for, including the presence of antibodies (tTG) in the untreated state and the presence of coeliac-related HLA-DQ markers. Currently two categories of RCD are being recognized: type I without aberrant T-cells and type II with aberrant T-cells detected by immunophenotyping by flowcytometric analysis or immunohistology of the intestinal mucosa (5). Arbitrarily, a percentage of aberrant cells CD7+CD3- of CD103+ IEL or cytoplasmic CD3+ surface CD3- % of CD103+ IEL of = 10%

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has been regarded as normal, and more than 20% as definitively abnormal.

Pathogenesis of RCD

Genetic and environmental factors

The environmental factor is mainly ingestion of gluten, while several genes contribute to the genetic predisposition (8). The main genetic factors, as mentioned before, are given HLA-DQ genes, i.e. the genes encoding DQ2 or DQ8 in the HLA complex on 6p21. Approximately 95% of coeliacs have a DQ2 comprised of DQB1*02 and DQA1*03. A small number of individuals lacking either of those heterodimers, have DQB1*02 or DQA1*05 alone (9,10). Gene dosage also affects CD susceptibility, heterodimer comprised of DQB1*02 and DQA1*05 and most of the remaining 5% have a DQ8 heterodimer. Homozygous individuals who carry DQB1*02 and DQA1*05 in cis on both chromosomes have a greater risk of developing complicated forms of coeliac disease (11). Non-HLA complex genes seem to contribute, but the nature and effects of these genes are less well known. The identification and knowledge of the function of additional genetic factors should improve the understanding of the actual pathogenesis of CD and lead to new diagnostic strategies in case-finding and screening high risk groups.

Diagnostic approach to RCD

Revision of the initial diagnosis coeliac disease

In a patient with villous atrophy refractory to a GFD, the first step requires the reassessment of the initial diagnosis of CD, in order to exclude other diseases, such as giardiasis, tropical sprue, postinfectious diarrhoea, collagenous sprue, protein intolerance, autoimmune enteropathy - associated T lymphoma (EATL).

Exclude other causes of diarrhoea with/without villous atrophy

In case of persisting diarrhoea despite demonstrable improvement in the histologic lesion and exclusion of dietary errors, other associated disorders should be considered. Well known causes responsible for symptoms principally include microscopic colitis and more rarely intermittent pancreatic insufficiency in CD, secondary lactase deficiency, bacterial overgrowth, coexisting inflammatory bowel disease, irritable bowel syndrome but also anal incontinence (4,13). There are many other causes of villous atrophy besides CD. Clinical history should investigate longer stays near the equator for detection of tropical sprue. Small bowel enteropathy seems to occur often in southern parts of Africa. Giardiasis should be excluded by immunofluorescence of stool samples and may be diagnosed by duodenal histology. Crohn’s disease with involvement of the duodenum may mimic or even coexist with CD. The term collagenous sprue should be used with caution, as this disease is not an established independent entity. A subepithelial matrix broader than 10–20 μm should point to the diagnosis of collagenous sprue. Deposition of excess of extracellular matrix underneath the basement membrane is an unspecific reaction, which can be seen in gluten-responsive CD, as well as in several other entities of RCD and also in enteropathy - associated T lymphoma (EATL).

Collagenous band-like structures regress to a large part in responsive CD. Autoimmune enteropathy is seen mainly in children and young adults, but may occur also in elderly patients. The histological picture often shows a diminished number of paneth cells. The number of IEL’s often is normal and patients present frequently with concurrent autoimmune diseases.

We have to realize that villous atrophy has also been reported in association with the presence of a thymoma, with protein intolerance, in conjunction with common variable immunodeficiency syndromes and eosinophilic enteritis. In common variable immunodeficiency, antibody testing for CD-associated antibodies is not useful. Only histological and clinical improvement on a strict GFD may reveal underlying CD in single cases of common variable immunodeficiency.

Exclude malignant complications of coeliac disease

Unexplained weight loss, abdominal pain, fever and night sweating should alarm physicians of an overt EATL. Other markers for overt EATL may be positive stool blood tests, increased LDH or beta2-microglobulin (14,15). In patients on GFD, EATL need not necessarily be accompanied by
duodenal villous atrophy (16). A high index of suspicion for an overt lymphoma should lead to an extensive work-up including upper and lower endoscopy, ear-nose-throat (ENT)-workup, CT-scan of thorax and abdomen with enteroclysis, video-capsule enteroscopy (VCE) and double balloon enteroscopy (DBE) in order to obtain histological specimens. In some cases laparotomy, intra-operative enteroscopy and full thickness biopsies are necessary, as the operative procedure may come to an earlier diagnosis what may be essential. New advances in small bowel imaging including CT scan and MR enteroclysis can improve the diagnostic accuracy in these patients (17).

PET-scan has been investigated in patients with EATL and RCD. Hadithi et al (18), in a prospective cohort of 8 EATL patients and 30 patients with RCD, demonstrated that PET can visualize in all patients the sites affected by EATL as confirmed on biopsy.

The diagnosis of overt T-cell lymphoma is based on histological and immuno-histochemical features with mainly evidence of large or medium size T-cell proliferation expressing a CD3+ CD8+/- and CD103+ phenotype. The majority presents as CD3+, CD8-, CD30+ large cell lymphoma; however small cell lymphomas often are CD3+, CD8+, CD30- (14,15).

Diagnosis of small bowel adenocarcinoma may even be more difficult than lymphoma. Especially tumours located in the jejunum and ileum, which are not reached by standard endoscopic techniques, require extensive investigations. Diagnosis may often be made only after operative procedures. Obscure gastrointestinal bleeding, obstructive symptoms, stenotic lesions on radiological examinations and VCE retention should raise the suspicion of these malignancies (author’s own experience).

The evolving role of double balloon enteroscopy

First described by Yamamoto and colleagues (19) in 2001, DBE is a new endoscopic technique with the potential to allow complete visualization of the entire small bowel.

In a European retrospective study, enteroscopy was used for the diagnosis of all patients suspected of having RCD (20).

Establishing the diagnosis of RCD

Finally, RCD is a diagnosis of exclusion, defined as a persisting villous atrophy that does not respond to a strict GFD. The demonstration of an aberrant clonal intraepithelial T-cell population and/or loss of antigen on IELs seem to characterize patients as having a high risk for the development of overt lymphoma and differentiates RCD II from RCD I, which shows low or almost absent aberrant T cells. RCD II is also referred to as a cryptic intestinal T-cell lymphoma; sprue-like intestinal T-cell lymphoma. Detection of a clonal T cell population by testing for TCR rearrangement is thought to be highly predictive of EATL development. However, oligo or monoclonal IEL’s population can be detected in the majority of both RCD I and RCD II patients, also in patients that do not develop an EATL. Clonality is therefore of limited use in establishing the diagnosis of RCD and predicting the development of EATL.

RCD I versus RCD II

Clinical and biological behaviour

Patients with RCD I may represent an earlier stage of the disease than RCD II and the prognosis may be better, while the risk of developing an overt lymphoma could be almost non-existent. In RCD I adherence to the GFD should be carefully investigated since a strict GFD may induce remission in some patients. In RCD I patients often develop concomitant autoimmune diseases, infectious and thrombembolic complications. Retrospective data from our patients suggest that RCD I patients have a mortality rate, which is not different from that of the general population (author’s own experience) The presence of mucosal ulcerations (ulcerative jejunitis) should alert the doctor for the possible presence of an early EATL (21).

RCD II is observed mostly in adults and the mean age at diagnosis of RCD II is between 50 and 60 years, but younger cases may be observed. Most of the patients develop severe malabsorption with weight loss, abdominal pain and diarrhoea. Some patients may also have skin lesions mimicking pyoderma gangrenosum or ulcerations mostly in legs, arms and face, chronic chest or sinusoidal infections or unexplained fever. The link between CD and RCD II is usually suggested by the detection of circulating antigliadin, anti-EMA or anti-TG antibodies before the initiation of the GFD in almost two-thirds of patients, an HLADQ2 or DQ8 status in almost all patients (11) and an initial response on GFD in about one-third of patients with RCD II.

Endoscopic and radiological features

Usually, in RCD I and II, the same pattern of villous atrophy is observed as in classical active CD. The finding of mucosal ulcerations, mostly in the jejenum, defines the clinical picture of ulcerative jejunitis (21). In some cases of RCD II also stomach and/or colonic ulcerations may be found (22). Enteroscopy using push- or DBE or VCE should be performed in such patients with RCD II in order to search for overt lymphoma and ulcerative jejunitis CT-scan with MRI-enteroclysis may be useful to exclude overt lymphoma and may demonstrate a mesenteric cavitation syndrome and hypoosplenisim (volume<100 cm3) in 30% of cases (author’s own experience). Enlarged mesenteric lymph nodes often accompany RCD, without necessarily being specific for a T-cell lymphoma. Staging investigations as recommended for non-Hodgkin lymphomas should be performed including bone marrow aspiration, CT-scan of the thorax and abdomen, sonography of the neck as well as ENT-workup especially in RCD II patients.
HLA-DQ typing
Typing of HLA-DQA1* and DQB1* alleles can be performed on whole blood samples. In our immunology laboratory, this typing is being performed with a combined single stranded conformation polymorphism/heteroduplex method by a semi-automated electrophoresis and gel staining method on the Phastsystem (Amersham-Pharmacia-Biotech, Uppsala, Sweden) (11).

We found a highly significant correlation between HLA-DQ2 homozygosity and the development of serious complications of CD, in particular RCD II and EATL.11. The link between HLA-DQ2 homozygosity and development of RCD II and CD-associated lymphoma of intraepithelial origin thus suggests that the strength of the gluten-specific T-cell response in the lamina propria directly or indirectly influences the likelihood of RCD II and lymphoma development. It has been reported earlier by Vander et al (23) that HLA-DQ2 homozygous antigen-presenting cells induce higher T-cell proliferation and cytokine secretion than HLA-DQ2/non-DQ2 heterozygous antigen-presenting cells. This may explain the strongly increased risk for disease development in HLA-DQ2 homozygous individuals. This would indicate that the adherence to a GFD is particularly important for CD patients who are HLA-DQ2 homozygous.

Small intestinal biopsies
Upper gastrointestinal endoscopy should be performed in all patients. At least 10 duodenal biopsies are to be taken for histological, immunohistochemical and flow cytometric examination. Four to six biopsies are fixed and preserved in 10% formalin for histopathological and immunohistochemical evaluation. Three - four biopsies for TCR gene rearrangement studies are taken separately, preserved on histocon and frozen at –20°C. For immunophenotypical evaluation 3-4 biopsies are taken and preserved in RPMI medium.

Immunophenotyping of IEL
Lymphocytes and enterocytes are isolated from 3-4 small intestinal biopsies. Intraepithelial localisation of lymphocytes is confirmed by surface expression of CD103 (αEβ7 integrin, a gut homing receptor for E-cadherin).

Arbitrarily, a percentage of aberrant cells CD7+CD3- of CD103+ IEL or cytoplasmic CD3 of CD103- % of CD103+ IEL of = 10% has been regarded as normal, and more than 20% as definitely abnormal.

TCR gene rearrangement study
Clonality assessment for TCR-γ gene rearrangements is achieved using the BIOMED-2 multiplex TCR-PCR protocol. Detection of a clonal T cell population by testing for TCR rearrangement is thought to be highly predictive of EATL development. However, oligo- or monoclonal IEL's populations can be detected in the majority of both RCD I and RCD II patients and also in patients that do not develop an EATL. Clonality is therefore of limited use in establishing the diagnosis of RCD and to predict the development of EATL (24,25).

RCD I vs. RCD II
In RCD I, histology of the small bowel mucosa in most cases is indistinguishable from active untreated CD. The number of IELs may be lower than in RCD II and active CD although this has not been proven in prospective studies. The IEL phenotype is normal with the expression of surface CD3 associated with surface CD8 and TCR-β as in classical active CD. Number of CD8 and TCR-β positive IELs should exceed 50% of IELs and lower expression seems to be quite sensitive for differentiation from RCD II but not very specific.

In RCD II an abnormal IEL clonal population may be observed in 80% of patients with RCD on small intestinal biopsies. Although these IELs have a normal cytological feature, they exhibit an abnormal IEL phenotype with the expression of intracytoplasmic CD3e, surface CD103 and the lack of classical surface T-cell markers such as CD8, CD4 and TCR-αβ. Furthermore the abnormal IEL phenotype is associated with clonal TCR gene rearrangement. This abnormal IEL population usually represents more than 50% of the IELs and may also be observed in gastric and/or colonic epithelium in around 2/3 of patients and may be found in the PBL in 1/3. It may be also detected in skin lesions or in the chest in patients, suggesting that RCD II is a diffuse gastrointestinal disease. The use of CD3 and CD8 on fixed biopsies is a very reliable method in order to assess the presence of this abnormal IEL phenotype, even retrospectively.

More recently, it has been shown on cell lines derived from clonal abnormal IEL that recurrent chromosomal abnormalities including a recurrent 1q trisomy may be found in these patients. The diagnostic yield of these cytogenetic features has not been evaluated so far. These chromosomal abnormalities, the clonality of the T-cell receptor gene and the loss of antigens on IEL’s, together with the frequent diffusion of the abnormal lymphocytes indicate, in spite of their normal cytology and low in situ proliferative capabilities that this clonal IEL population can be considered as a cryptic intraepithelial lymphoma. This hypothesis is sustained by follow-up studies.

Medical treatment options
Treatment of RCD I
In contrast to patients with a high percentage of aberrant T cells, patients with RCD I seem to profit from an immunosuppressive treatment. According to the data of Goerres et al (26) , azathioprine should be first line therapy after induction of clinical remission with corticosteroids. Dose and duration of treatment with azathioprine are not established.

In contrast to RCD II, long-term treatment with corticosteroids or locally acting budesonide may be considered only in patients who have contraindications to other immunosuppressants. Cyclosporine A, infliximab and tacrolimus have been reported to be effective in case reports, but further data are required particularly in the light of severe limited use.
side effects (27). These agents should only be considered in cases of clinical deterioration despite corticosteroid therapy or intolerance to azathioprine. Intestinal absorption of cyclosporine A is worse than that of tacrolimus, what has to be considered for treatment of acute cases. Also parenteral administration of tacrolimus has to be considered. Close monitoring of renal function is inevitable.

Treatment with infliximab may induce prompt clinical and histological response but this effect has to be weighed against its possible acute allergic and chronic immunosuppressive side effects (28).

**Treatment of RCD II**

RCD II is usually resistant to medical therapies. Response to corticosteroid treatment does not exclude underlying EATL, which has been shown in single cases. In cases of RCD II with persistent clinical symptoms and/or high percentage of aberrant T cells in intestinal biopsies in spite of a corticosteroid treatment, more aggressive therapeutic schemes should be considered. However, no therapy seems to be curative in RCD II.

Some patients may benefit from azathioprine (26). Caution is required when initiating immunosuppressive therapy, as this may induce a high risk of progression to an overt lymphoma. In most cases CHOP-like regimens have been applied, but also other agents used for nodal NHL may be applied. Maurino et al (29) reported the results of treating 7 RCD II patients with azathioprine. Clinical and histologic improvement was noted in 5 of the 7 treated patients, although 3 patients died (1 from leukopenic fever and 2 died early). However, in their follow-up report on treating 13 patients with azathioprine, they reported a 46% mortality rate. A recent report on the anti–tumor necrosis factor agent infliximab for treatment of RCD has been published, but no data were provided on aberrant T cells (T flow cytometry or immunohistology) (30).

Recognizing that some patients with RCD II, and especially with ulcerative jejunitis, are suffering from a low-grade EATL, we treated a group of these patients with cytotoxic chemotherapy. Cladribine (2-chlorodeoxyadenosine [2-CDA]) is a synthetic purine nucleoside with cytotoxic activity. Cladribine is of proven value in the treatment of hairy cell leukemia. Pathologic cells in hairy cell leukemia are CD103 positive as in T cells in RCD II. In the last decade of the twentieth century, stem cell transplantation (SCT) has become an increasingly accepted treatment option for patients with severe autoimmune diseases refractory to conventional treatment. The applicability of autologous SCT (ASCT) in a selected group of refractory coeliacs with aberrant T cells (32). Between March 2004 and March 2006, 7 patients were transplanted. EATL was excluded by endoscopic examination, computed tomography, body positron emission tomography and bone marrow biopsy.

Stem cells were harvested from the peripheral blood after mobilization using G-CSF. The conditioning regimen consisted of T-cell depletion with fludarabine and myeloablation with melphalan. On follow up, our patients showed improvement in the small intestinal histology, together with impressive clinical improvement as demonstrated by disappearance of diarrhoea and abdominal pain, normalization of serum albumin, electrolytes and haemoglobin, increase in BMI and improvement of the performance status. Two years after transplantation, our first patient is showing further improvement in his immunopathology status as demonstrated in further decline in the percentage of aberrant T cells to 3% and histologically improved from Marsh III-A to Marsh-I. We propose that enhanced apoptosis of activated but aberrant T cells has led to this late but remarkable decline. Our most recent patient with clinically short bowel syndrome is showing remarkable clinical, endoscopic and immunological improvement. Furthermore, the first 3 patients showed a significant increase in the percentage of CD8+ lymphocytes, which is seen as a marker of lymphocyte regeneration after ASCT. Absence of a demonstrable improvement in the surface expression of CD8 on the IEL might be regarded as a poor prognostic indicator of response; this can only be proved or disapproved through a longer term follow up.
Although the short term results in these patients are promising, follow up at present is too short to allow firm conclusions as to efficacy. The selection of patients for this treatment should be restricted to those patients with a substantial population of aberrant T cells, even after therapy with 2- CDA who have a greater tendency to progress to highly lethal EATL.

**Follow-up and overt lymphoma in RCD**

RCD II is a serious disorder with a 5-year survival less than 50% and the most frequent cause of death is the occurrence of an overt T-cell lymphoma and recurrent infections. The presence of an abnormal clonal IEL population is significantly associated with a poor survival and a high risk of progression to overt lymphoma. The same clonal TCR-gene rearrangement initially identified in patients with clonal RCD may be subsequently observed in lymphomatous specimens suggesting a continuum between RCD and high-grade lymphoma. The risk of developing an overt T-cell lymphoma in patients with RCD II seems to be favoured by immunosuppressive drugs.

ASCT in RCD II patients seems to promising. Whether a close monitoring with video capsule and/or PET-scan are capable of detecting earlier lesions in RCD before the development of an overt lymphoma which results in a better outcome, remains to be answered.

**References**

25. Blumberg RS, Yockey CE, Gross GG, Ebert EC, Balk SP. Human intestinal intraepithelial lymphocytes are derived from a limited number of T cell clones that utilize multiple V beta T cell receptor genes. J Immunol 1993;150:5144-5153.