**Name of Journal: *World Journal of Gastrointestinal Surgery***

**ESPS Manuscript NO: 21067**

**Manuscript Type: Minireviews**

**Chronic haemorrhagic radiation proctitis: A review**

Nelamangala Ramakrishnaiah VP *et al*. Radiation proctitis review

**Vishnu Prasad Nelamangala Ramakrishnaiah, Srinivasan Krishnamachari**

**Vishnu Prasad Nelamangala Ramakrishnaiah,** **Srinivasan Krishnamachari,** Department of Surgery, Jawaharlal Institute of Postgraduate Medical Education and Research, Puducherry 605006, India

**Author contributions:** Nelamangala Ramakrishnaiah VP and Krishnamachari S equally contributed to the minireview in designing and writing the aims, writing the methods, searching the literature and in writing the discussion and conclusions.

**Conflict-of-interest** **statement:** None.

**Open-Access:** This article is an open-access article which was selected by an in-house editor and fully peer-reviewed by external reviewers. It is distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/

**Manuscript source:** Invited manuscript

**Correspondence to: Vishnu Prasad Nelamangala Ramakrishnaiah, MS, DNB, MNAMS, FMAS, MCh (GI Surgery), Professor** of Surgery, Department of Surgery, Jawaharlal Institute of Postgraduate Medical Education and Research, Dhanvantri Nagar, Gorimedu, Puducherry 605006, India. vprasad285@gmail.com

**Telephone:** +91-413-2277586

**Received:** June 28, 2015

**Peer-review started:** July 5, 2015

**First decision:** August 9, 2015

**Revised:** April 29, 2016

**Accepted:** May 10, 2016

**Article in press:**

**Published online:**

**Abstract**

Chronic haemorrhagic radiation proctitis (CHRP) is a difficult problem faced by the patients following radiation for pelvic malignancy. There is no standard treatment for this condition, but many methods of treatment are available. The aim of this study was to review the literature to see whether there is an improvement in the available evidence in comparison with previously published systematic reviews in treating patients with CHRP. The PubMed/Medline database and Google Scholar search was selectively searched. Studies, which treated patients with rectal bleeding due to chronic radiation proctitis or chronic haemorrhagic radiation proctitis, were included. Seventy studies were finally selected out of which 14 were randomized controlled clinical trials. Though these studies could not be compared, it could be seen that there was an improvement in the methodology of the studies. There was an objective assessment of symptoms, signs and an objective assessment of outcomes. But, still, there were only a few studies that looked into the quality of life following treatment of CHRP. To increase recruitment to trials, a national registry of cases with established late radiation toxicity would facilitate the further improvement of such studies. Some of the conclusions that could be reached based on the available evidence are 4% formalin should be the first line treatment for patients with CHRP. Formalin and argon plasma coagulation (APC) are equally effective, but formalin is better for severe disease. Refractory patients, not responding to formalin or APC, need to be referred for hyperbaric oxygen therapy or surgery. Radio-frequency ablation is a promising modality that needs to be studied further in randomized trials.

**Key words**: Radiotherapy; Proctitis; Complications; Systematic review; Formalin

© **The Author(s) 2016**. Published by Baishideng Publishing Group Inc. All rights reserved.

**Core tip:** The aim of this study was to review the literature to see whether there is an improvement in the available evidence in comparison with the previously published systematic reviews in treating patients with chronic haemorrhagic radiation proctitis (CHRP). The PubMed/Medline database and Google Scholar search was selectively searched. Seventy studies were finally selected out of which 14 were randomized controlled clinical trials. It could be seen that there was an improvement of the methodology of the studies though they were not comparable. Based on the available evidence, 4% formalin should be the first line treatment for patients with CHRP.

Nelamangala Ramakrishnaiah VP, Krishnamachari S. Chronic haemorrhagic radiation proctitis: A review. *World J Gastrointest Surg* 2016; In press

**INTRODUCTION**

One to five percent of patients who receive radiotherapy as adjuvant or neoadjuvant therapy for pelvic malignancy will develop chronic haemorrhagic radiation proctitis (CHRP). In one of the recently published series, it was noted that 1319 patients received radiation for carcinoma of cervix over a period of 22 mo and 124 similar patients during the same period needed treatment for CHRP in the same centre[1]. The meaning of the above sentence shows the magnitude of the problem of CHRP. Newer methods of radiotherapy like three-dimensional conformal radiation therapy and intensity-modulated radiation therapy can use higher doses of radiation to the target tissues with less exposure to adjacent normal tissues. Protons and neutrons, so-called particle radiation, are also being tested but the long-term outcomes of these modalities are not known, and these are expensive. The use of brachytherapy is also found to be associated with fewer complications. Thus, the incidence of CHRP is related to the dose of radiation, the area of exposure, methods of delivery, the use of cytoprotective agents and other factors[2].

　　Because the rectum has a fixed position in the pelvis, it becomes more susceptible to radiation injury. Acute radiation injury of rectum occurs within three months of starting radiotherapy. It is an inflammatory process of rectal mucosa with a loss of microvilli, oedema, and ulceration. It is self-limiting and manifests as abdominal pain, tenesmus, diarrhoea, incontinence and urgency and resolves within three months[3]. Unlike acute radiation proctitis, chronic radiation proctitis takes a period of 3 mo after pelvic radiation, but usual median time is 8-12 mo. It can also continue from acute phase[3]. It is due to obliterative endarteritis, submucosal fibrosis, and neo-vascularization (Figure 1). Chronic radiation proctitis can present with rectal bleeding, tenesmus, mucus discharge, diarrhoea, incontinence, and urgency. It may be asymptomatic also. The diagnosis of radiation proctitis should be suspected if a patient presents with the above -mentioned symptoms and gives a history of pelvic radiation. The diagnosis is confirmed by sigmoidoscopy or colonoscopy that shows pale, friable mucosa with telangiectasia. Rectovaginal, recto-urethral, recto-vesicular fistulizing disease is a late-presenting sign. There is no role of biopsy to confirm the diagnosis since it may produce complications.

There is no standard treatment for CHRP. However many treatments are available like amino salicylates, butyric acid enema, steroid enemas, formalin, argon plasma coagulation (APC), hyperbaric oxygen, radiofrequency ablation and even surgical therapy. The outcome of any of these medical and surgical treatment can be disappointing[1]. There are not many good-quality placebo-controlled trials.

In this study, our aim was to review the literature to see whether there is an improvement in the available evidence in comparison with previously published systematic reviews in treating patients with CHRP. The PubMed/Medline literature database was selectively searched for articles with the keywords “Proctitis/drug therapy”[Mesh] OR “Proctitis/radiotherapy”[Mesh] OR “Proctitis/surgery”[Mesh] AND “radiotherapy”, “Management of CHRP” “Related Review articles”. In addition Google search and Google Scholar search was also made using key words “Radiation proctitis” “Formalin” “Endoscopic therapy” “Argon plasma coagulation” “Radiofrequency ablation” “cryotherapy” “Hyperbaric oxygen therapy” and “surgery”. The literature search was mostly limited to articles in English and human patients. No limitations for the year of publication were applied. All the studies that treated patients with rectal bleeding due to chronic radiation proctitis or CHRP were included in the review. Studies of patients treating acute radiation proctitis were excluded.

We could find 142 articles in total. After removing the duplicates and studies on acute radiation proctitis, there were about 86 articles. Out of these 86, 16 were further excluded because of various reasons such as anecdotal studies. Various studies that were found to be relevant are summarized below. Importance was given to randomized controlled clinical trials.

**STUDIES USING ANTI-INFLAMMATORY DRUGS, STEROIDS, SUCRALFATE AND PENTOSAN POLYPSULPHATE**

Sulfasalazine or 5-aminosalicylates, steroids are the drugs used initially for treating CHRP. Their mechanism of action is by inhibition of prostaglandin synthesis. It may also be due to inhibition of folate-dependent enzymes[4].Sucralfate stimulates epithelial healing and forms a protective barrier[5]. Sucralfate is shown to be better than anti-inflammatory agents[6].Pentosan polysulphate is similar to sucralfate. There are more than seven to eight publications using these drugs.

In a prospective double-blind, randomized controlled trial involving 37 consecutive patients with radiation-induced proctosigmoiditis[6], there were 36 females treated for cervical cancer and one male treated for prostate cancer. The mean duration after completion of the radiotherapy was 8.3 mo. These patients were randomized to receive either 3 g oral sulfasalazine plus 20 mg twice daily of rectal prednisolone enemas(group I, *n* = 18) or 2 g of rectal sucralfate enema plus oral placebo (group II, *n* = 19) for four weeks. These two groups were comparable with respect to demography, clinical symptoms and endoscopic staging of the disease. Patients in Sucralfate enema showed a better clinical response although endoscopically the response was not statistically significant. Follow-up was limited to 4 wk.

Rougier *et al*[7,8], in their randomized trial, compared betamethasone enema (5 mg bd) with hydrocortisone mousse (90 mg bd) and concluded that hydrocortisone group had a better outcome. There were 32 patients with CHRP in this study. The outcomes used were bowel activity, tenesmus, rectal bleeding and endoscopic grading. Follow-up was limited to 4 wk.

In another randomized study by Cavcić *et al*[9], compared combination of oral metronidazole (400 mg tds), mesalamine (1 g tds) and rectal betamethasone to oral mesalamine and rectal betamethasone and found that the rectal bleeding and ulcers were significantly lower in the metronidazole group. In this study, there were sixty patients randomized into either group. The efficacy of metronidazole was assessed on the basis of rectal bleeding, diarrhoea and proctosigmoidoscopy in all patients. The follow-up was up to 12 mo. Grigsby *et al*[10] prospectively showed the benefit of oral pentosan polysulphate given for a period of 1 year in 13 patients.

**STUDIES USING** **SHORT-CHAIN FATTY ACID ENEMAS**

Short-chain fatty acids (SCFA) stimulate the growth of colonic mucosa. The vasodilatation effect may improve the blood flow of colonic mucosa[11]. Butyric acid is the main SCFA. There are more than six studies using SCFA. Many of them are case series.

Two randomized studies showed non-significant improvement of symptoms and signs but both the studies were underpowered[12,13]. Talley *et al*[12] in their randomized double-blind placebo-controlled cross over trial of 15 patients treated one group with the butyric acid enema and another group with normal saline placebo. Symptoms score, endoscopic scores and even histology were compared.

Similarly, Pinto *et al*[13] in their randomized prospective double blind controlled trial of 19 patients treated one group with SCFA enema and another group with placebo. In this study apart from symptoms and endoscopic features, biopsies for mucosal DNA and protein content were also measured. Patients were followed up to 6 mo.

Though we were treating patients with CHRP in our institute since 1985, study on chronic haemorrhagic proctitis were started in 1999. The first study on CHRP in our institute, done by Senthil Kumar *et al*[14] in 2001, compared sucralfate-steroid enema (25 mg of prednisolone and 1gm of sucralfate twice daily for 14 d) with butyric acid retention enema (60 mL containing 40 mmol of butyric acid twice daily for 14 d) in a double-blind randomized controlled trial. There were thirty patients randomly allocated. They were followed up to 4 wk. Outcomes were measured by the improvement in the colonoscopic grading of severity and clinical symptoms. Histopathological improvements were also compared by taking the biopsy before and after treatment. The conclusion was that both the methods of treatment were equally effective since there was relief of symptoms of radiation proctitis in both the methods of treatment without improvements in endoscopic scores or histology. However, the sucralfate-steroid enema was easier to prepare[14]. No toxicities were reported in any of these studies.

**STUDIES USING FORMALIN THERAPY**

Formalin scleroses and seals fragile neovasculature in tissues damaged due to radiation and prevents further bleeding. In 1986, Rubinstein was the first to use formalin for a CHRP patient to get a good response[15]. Following this, there are several reports in the literature[16-30]. But the majority are retrospective in nature, a few are prospective studies. The technique and the concentration of formalin used in these studies also differ. The two main methods of using it are 4% solution as irrigation or as soaks. There are reports of using 10% solution of formalin also[31].There are four Randomized trials using formalin for CHRP.

Ours is one of the first published randomized trial comparing the efficacy of the 4% formalin dab with Sucralfate-steroid retention enema (100 mg of prednisolone and 1 gm sucralfate in 100 mL of normal saline twice daily for 14 d)[1]. In this study, 102 patients were randomly allocated to either of the treatment arms. This study objectively assessed the symptoms scores using the radiation proctopathy system assessment scale (RPSAS) and also the sigmoidoscopic grade (Modified Chi grading) before and after treatment and found that Formalin dab is superior to sucralfate-steroid enema in treating CHRP involving only the rectum. It was also observed that a single session of formalin dab can effectively treat CHRP in 90% of the patients, and multiple sessions could effectively treat 99% of the patients whereas sucralfate-steroid enema was effective only in 75% of patients. These patients were followed up to 9 mo. There was no complications or toxicity.

Following this Yeoh *et al*[32] showed in their randomized study that APC and topical formalin had comparable efficacy in the durable control of rectal bleeding associated with chronic radiation proctitis but had no beneficial effect on anorectal dysfunction. In this study thirty patients were randomized into each group. Anorectal symptoms, (modified LENT-SOMA questionnaire) anorectal manometry and anorectal morphology by endorectal ultrasound were assessed before and after treatment.

Guo *et al*[33] in their randomized trial showed that 10% formalin is associated with complications and 4% formalin should be the choice for treating CHRP. In this study 122 patients were randomized into 4% or 10% formalin application. Outcomes were compared with symptoms score and rectoscope scores. Follow-up was up to 1 year. Wong *et al*[34] from their prospective database, after a decade of experience of treating patients with radiation proctitis, have shown that formalin is more effective than APC in treating patients with CHRP. APC has the potential to complement topical formalin application and can be used to treat the proximal and distal rectum concurrently.

The contrary report has been published by Alfadhil *et al*[35] in their retrospective comparative study of 22 patients who received formalin application or APC. Improvement in Hb% was used to assess the outcome. The severity of the proctitis was not assessed before the treatment. The lag time between radiation and endoscopic treatment was not known. The study was underpowered, and the groups were not comparable. The details of the adverse events not mentioned. They concluded that APC is more effective than formalin and has less adverse effects.

Sahakitrungruang *et al*[36] in their randomized controlled trial comparing colonic irrigation with oral antibiotics administration *vs* 4% formalin application for treatment of CHRP have shown that the former method is better than 4% formalin application. Fifty patients were randomly allocated to each arm. Daily self-administered colonic irrigation of 1 liter tap water and a 1-wk period of oral antibiotics-ciprofloxacin and metronidazole were given in one arm. Four percent formalin application for 3 min was done in another arm. Patient’s satisfaction was surveyed. The limitation was that the study was a 2-armed design without a crossover trial. Hence, it could not illustrate whether the antibiotics and irrigation were equally important. Some of the adverse events noted in the literature regarding the use of formalin for CHRP are the rectal stricture, worsening of incontinence, anococcygeal pain, and formalin colitis[24,30].

**STUDIES USING THERMAL COAGULATION THERAPY**

Endoscopic coagulation with a variety of devices has been reported to be effective for CHRP[7]. The technique involves coagulation of a bleeding point rather than the entire friable mucosa. Several treatment sessions are often required[7]. The modalities include heater probe, bipolar Electrocoagulation, neodymium:yttrium-aluminium-garnet (Nd: YAG) laser, potassium titanyl phosphate (KTP) laser, argon laser and APC. Simple heater probe and APC are preferred for their better safety profile[37].

Both the heater probe and bipolar cautery are contact probes. The heater probe has a Teflon-coated heating element at its tip that delivers standardized energy over set times. Bipolar electrocautery probe has a pair of electrodes at its tip through which current is passed using the tissue for conduction[38]. Jensen *et al*[39] in his randomized study showed that 21 patients treated either with a heater probe, or bipolar cautery showed benefits without much difference between the two modalities[39]. A mean of four sessions was needed in each arm during treatment in this study. Patients were followed up to one year. The increase in haematocrit, endoscopic resolution, and patient satisfaction were compared. No complications were noted.

Nd:YAG laser is the first endoscopic laser used for treating CHRP. Some of the complications reported with this are transmural necrosis, fibrosis, necrosis, stricture formation and recto-vaginal fistula. Nd:YAG laser use for CHRP has declined due to several reasons, firstly its cost, second, the need to aim directly at telangiectasias and the possibility of severe endoscopic damage if the laser strikes the endoscope in retroflection[40]. Taylor *et al*[41] used KTP laser for treating 26 patients with bleeding secondary to CHRP using 4-10 W and a median of two sessions. They reported a symptomatic improvement in 65% patients while there was no change in 7 (30%), and symptom like hematochezia increased in 1 (5%). Similarly, there are only case series using argon laser for treating CHRP.

There are more than 15 published reports of APC for CHRP. Many are retrospective studies, and some of them are prospective case series. Many of the series report unsuccessful medical treatment before going for APC. In APC bipolar diathermy current is applied using inert argon gas as a conducting medium. It can be applied tangentially and radially.

Karamanolis *et al*[42] showed in their prospective study treating more than 56 patients, that APC was successful in all patients with mild and in almost all patients with moderate CHRP. In contrast, APC failed in 50% of patients, wherever the presence of severe mucosal damage was present. The grading of severity was based on endoscopic criteria taking into consideration telangiectasia distribution and surface area involved. For APC application, a 2.3 mm diameter front firing APC probe inserted through the working channel of the flexible sigmoidoscope was used. The argon flow rate and the electrical power were set at 2 L/min and 40 W, respectively. Patients were followed up for a mean of 17 mo. Patients required 1-2 sessions of APC for mild proctitis while patients with moderately to the severe form required a statistically significantly higher number of APC sessions. In cases of severe and diffuse involvement of the rectum, multiple treatments sessions are required, and success is less certain as shown by other reports also[43-51]. In many of these series, the response is objectively scored using bleeding severity score, haematological parameters, and endoscopic scores.

Chruscielewska-Kiliszek *et al*[52] in their randomized, double-blind trial comparing oral sucralfate or placebo following APC for CHRP have shown that additional sucralfate treatment after APC did not influence the clinical or endoscopic outcomes. One hundred and seventeen patients completed the treatment protocol, 57 in the sucralfate group and 60 in the placebo group. Patients were graded clinically and endoscopically according to the Chutkan and Gilinski scales before and at 8 and 16 wk after initial APC treatment (1.5-2 L/min, 25-40 W) and after 52 wk (clinical only)[52]. Complications (1%-15%) following APC, such as pain, ulceration, perforation, explosion, extensive necrosis and rectal stricture have been cited in the literature[42].

**STUDIES USING RADIO-FREQUENCY ABLATION**

There are more than five reports of case series and retrospective studies using radio-frequency ablation (RFA) for CHRP. Many case series have shown, using BARRx Halo90 electrode catheter that was fit on the distal end of the flexible sigmoidoscope, an energy density of 12 J/cm2 at a power density of 40 W/cm2, hemostasis could be obtained after 1 to 2 sessions[53-56].

Several benefits RFA have been claimed, these include squamous re-epithelialization, lack of stricturing and ulceration. Using RFA much broader area of tissue can be treated simultaneously compared to the point by point approach by other methods[37]. The radio-frequency unit is mobile and can be used in different rooms of an endoscopy unit. Zhou *et al*[54] have used real-time endoscopic optical coherence tomography (EOCT) to visualize epithelialization and subsurface tissue microvasculature pre- and post-treatment RFA in their case series and have shown the potential of EOCT for follow-up assessment of endoscopic therapies.

**STUDIES USING CRYOABLATION**

Cryoablation is similar to APC and involves the non-contact application of liquid nitrogen or carbon-dioxide to tissues for superficial ablation. It is possible to treat a larger surface area like in RFA. Its effect is due to ischemic necrosis which can be immediate or delayed.

There are only case series of 20 patients. During cryoablation, a decompressive rectal tube has to be inserted because of the risk of over insufflation and perforation. Cryotherapy units are less mobile. Unlike in Radiofrequency ablation, the depth of tissue penetration may be more here. This may lead to strictures. However, colonic lavage is not necessary since there is no risk of explosion. The number of required sessions range from one to four[57-59].

**STUDIES USING HYPERBARIC OXYGEN THERAPY**

There are more than 12 published studies using hyperbaric oxygen therapy (HBOT) for CHRP**.** New reports have started appearing in the literature regarding the efficacy of HBOT**.**

Clark *et al*[60] in their randomized controlled double-blind crossover trial (150 patients) with a long-term follow-up, up to 5 years, showed that in patients with refractory CHRP, HBOT had a significant healing response. Primary outcome measures involved were the late effect in normal tissue-subjective, objective, management, analytic (SOMA-LENT) score and standardized clinical assessment. The secondary outcome was the change in the quality of life[60].

In one of the largest Australasian study using HBOT for chronic radiation injuries, Tahir *et al*[61] showed a clinical response rate for CHRP of 95%, where around half of the cases had a durable major response, with some patients experiencing symptom relief lasting as long as seven years.

At pressure greater than atmospheric pressure and using 100% oxygen, HBOThas an angiogenic effect and has been shown to cause an eight to nine-fold increase in the vascular density of soft tissues over air-breathing controls[7]. HBO acts to stimulate collagen formation and re-epithelialization**.** There is no uniformity in the methods of treatment using HBOT[42,61-65]. Although it can be perceived from the studies that HBOT is useful in refractory radiation proctitis, there is marked variation between the studies. There are no major adverse effects. Minor adverse event recorded is transient aural barotrauma. The reported number of HBOT sessions for a successful treatment range from 12 to 90. The cost of HBOT is high, and hence, it is not widely applicable.

***Other interventions***

Oxidative stress is thought to be one of the mechanisms in the development of chronic radiation proctitis and antioxidants have been used to treat CHRP. Use of vitamin C and E have been reported. Kennedy *et al*[66] treated twenty consecutive patients with CHRP. They used a combination of vitamin E at a dose of 400 IU tid and vitamin C at a dose of 500 mg tid. They assessed the response by symptom index and lifestyle questionnaire. A good number of study patients in the study seem to benefit. This pilot study was not studied further.

Retinol palmitate (vitamin A) has been shown to increase wound healing because of increased collagen cross-linking. This has been used in a randomized study to show improvement of symptoms of chronic radiation proctopathy by Ehrenpreis *et al*[67]. They randomized 19 patients, 10 patients to retinol palmitate group and nine to the placebo group. Five placebo nonresponders were crossed over to the retinol palmitate. The RPSAS scores before and every 30 d for 90 d were measured. The definition of response was a reduction in two or more symptoms or by at least two RPSAS[67]. Therewas a significant improvement in symptoms in the treatment group compared with the control and also when the controls were crossed over to treatment. But the study was underpowered.

***Surgical interventions***

Surgery is the last resort in patients with CHRP. Around 10%-25% of patients with CHRP finally need surgery[68]. Intractable bleeding, perforation, stricture, and fistula are some of the indication for surgery in patients with chronic radiation proctitis. There are case reports of non-surgical dilatation for strictures for this condition[69]. Significant improvement of bleeding by diversion has been shown by one of the retrospective study[70]. Fistula with the adjacent structures may need resection or resection with reconstruction with a diverting stoma. Whenever surgical treatment became a necessity, studies report poor outcomes with high complications (15%-80%) and mortality (3%-9%)[70-73]. Since nonoperative interventions are commonly used nowadays in managing patients with CHRP, There are no recently published series on surgical interventions on this issue**.**

***Discussion***

Evidence-based medicine requires the systematic and critical evaluation of published and unpublished trials[74]. In 2002, when Denton *et al*[7] first published their systemic review of the non-surgical intervention of late radiation proctitis, they could identify only six randomized controlled trials. The majority of the evidence available was either one individual’s or one center’s experience with a specific intervention without comparison to a control or another agent. This is probably due to the low incidence of chronic radiation proctitis in the majority of centers and the difficulties that co-exist in compiling a series large enough to be randomized between therapies[34].

Thirteen years later we could identify a total of 14 randomized controlled trials treating 804 patients with CHRP. In many of these studies, we could get the details of the reason for radiation therapy and the dosage. The diagnosis was based on the history and the endoscopic findings. At present, there is no validated score for CHRP, which can be used universally for grading the severity. There can be the inter-observer difference of the same findings. Tissue biopsy may not be conclusive. Patients may not tell their exact symptoms unless directed questions are asked. The severity of the radiation proctitis was graded in many of the studies objectively using symptoms score like RPSAS[1,67] or LENT-SOMA scale[32,60] and intraluminal findings by the sigmoidoscopic or colonoscopic grade (modified Chi grading[1] or Chutkan and Gilinski scales[52]). But different studies used different severity scores and hence the inter-institutional comparison of data is still difficult. The same is true with the outcome measures. Yeoh *et al*[32] have tried to see the rectal functions as well as morphology by using anorectal manometry and endorectal ultrasound. Zhou *et al*[54] have shown the most efficient objective assessment of the response to treatment by using EOCT. There were only a few studies that surveyed the quality of life following treatment[36,60]. However, unlike the previous studies, follow-up of recent studies is fairly long and is usually more than 9 to 12 mo[1,33,36,52,60]. With these randomized trials it is possible to say that there is evidence to make the following judgments:(1) Sucralfate enema appears to have a better effect than anti-inflammatory agents; (2) Anti-inflammatory drugs appear to have a better effect if used with oral metronidazole; (3) Rectal hydrocortisone appears to have a better effect than rectal betamethasone; (4) Sucralfate-steroid retention enema and short chain fatty acid enema are both equally but moderately effective in treating CHRP, but sucralfate-steroid enema is easy to prepare; (5) Four percent formalin is more effective than sucralfate-steroid retention enema and can be effective in 99% of the patients of CHRP; (6)Four percent formalin should be preferred over 10% formalin in treating patients with CHRP since 10% formalin is likely to cause adverse events; (7) Heater probe and bipolar cautery are equally effective in treating patients with CHRP; (8) Both APC and formalin don’t improve the rectal dysfunction but only stop the bleeding; (9) Both APC and formalin are equally effective, but formalin may be better in severe disease; (10) Additional oral treatment after APC will not improve the outcomes; (11) Radiofrequency ablation is a promising upcoming modality of treating CHRP but more robust data in the form of randomized trials needed; (12) HBOT is the only treatment modality, currently, which addresses the underlying problem and effective in treating CHRP patients but is costly and available in a few centers; (13) Vitamin A and other modalities have to be kept in mind while treating these patients since some report shows its efficacy. Further trials and robust data needed to show its efficacy; and (14) Surgical intervention is to be kept as a last resort in patients not responding to any of the methods described above.

Looking at the available evidence, it is clear that there is some improvement in the methodology of these studies. There is an objective assessment of symptoms and signs and also the objective assessment of the outcomes in some of these studies. The major drawback is that the objective assessment is not uniform, different studies using different scores. Also, not much importance is given to the quality of life assessment following treatment. It has been felt by the previous reviewers that one study, even if well conducted, will not be able to modify the changes in practice[7]. It has been felt by Denton *et al*[7] that in order to increase recruitment to trials a national registry of CHRP cases would facilitate multicenter trials with uniform entry criteria, uniform baseline and uniform therapeutic assessments providing standardized outcome data[75].

Limitations of this review: The search was limited to PubMed/Medline, Google and Google Scholar and was not complete and was limited to English language journals only. Individual authors were not contacted.

**CONCLUSION**

Based on this evidence, it can be concluded that the first line treatment of a patient with CHRP, the most effective way of treating CHRP, should be 4% formalin application. Since it is cheap, easily available, can be applied easily and effective in 99% of the patients. If the radiation proctitis extends beyond rectum, then APC will be a better alternative. Alternatively, both formalin and APC can be used as complementary methods. Those patients who are refractory to formalin or APC may be referred for treatment with HBOT. In centers where radiofrequency ablation is available, further randomized studies should be done to see the efficacy of it in treating patients with CHRP. Those patients with CHRP who do not respond to any of the modality may need surgery in the form of diversion colostomy. Those patients with CHRP presenting with complications like stricture, fistula or other complications like obstructions may need surgery at presentation.

**REFERENCES**

1 **Nelamangala Ramakrishnaiah VP**, Javali TD, Dharanipragada K, Reddy KS, Krishnamachari S. Formalin dab, the effective way of treating haemorrhagic radiation proctitis: a randomized trial from a tertiary care hospital in South India. *Colorectal Dis* 2012; **14**: 876-882 [PMID: 22356304 DOI: 10.1111/j.1463-1318.2012.03008.x]

2 **Do NL**, Nagle D, Poylin VY. Radiation proctitis: current strategies in management. *Gastroenterol Res Pract* 2011; **2011**: 917941 [PMID: 22144997 DOI: 10.1155/2011/917941]

3 **Agrawal PP**, Bansal N, Bahadur AK, Singh K, Rathi AK. Management of chronic hemorrhagic radiation proctitis. *Asia-Pac J Clin Onco* 2007; **3**: 19-29 [DOI: 10.1111/j.1743-7563.2006.00084.x]

4 **Peppercorn MA**. Sulfasalazine. Pharmacology, clinical use, toxicity, and related new drug development. *Ann Intern Med* 1984; **101**: 377-386 [PMID: 6147110 DOI: 10.7326/0003-4819-101-3-377]

5 **Nagashima R**. Mechanisms of action of sucralfate. *J Clin Gastroenterol* 1981; **3**: 117-127 [PMID: 6798100]

6 **Kochhar R**, Patel F, Dhar A, Sharma SC, Ayyagari S, Aggarwal R, Goenka MK, Gupta BD, Mehta SK. Radiation-induced proctosigmoiditis. Prospective, randomized, double-blind controlled trial of oral sulfasalazine plus rectal steroids versus rectal sucralfate. *Dig Dis Sci* 1991; **36**: 103-107 [PMID: 1670631 DOI: 10.1007/BF01300096]

7 **Denton AS**, Andreyev HJ, Forbes A, Maher EJ. Systematic review for non-surgical interventions for the management of late radiation proctitis. *Br J Cancer* 2002; **87**: 134-143 [PMID: 12107832 DOI: 10.1038/sj.bjc.6600360]

8 **Rougier P**, Zimmerman P, Pignon JP, Kac J, Crespon B, Zrihen E, Charbit MF. Rectitesradiques: Efficatecomparee de deux types de corticoids adminstrelocalement. *Med Chir Dig* 1992; **21**: 91-93

9 **Cavcić J**, Turcić J, Martinac P, Jelincić Z, Zupancić B, Panijan-Pezerović R, Unusić J. Metronidazole in the treatment of chronic radiation proctitis: clinical trial. *Croat Med J* 2000; **41**: 314-318 [PMID: 10962052]

10 **Grigsby PW**, Pilepich MV, Parsons CL. Preliminary results of a phase I/II study of sodium pentosanpolysulfate in the treatment of chronic radiation-induced proctitis. *Am J Clin Oncol* 1990; **13**: 28-31 [PMID: 1689537 DOI: 10.1097/00000421-199002000-00008]

11 **Aalkjaer C**. Short chained fatty acids and the colon: how do they cause vasodilatation? *J Physiol* 2002; **538**: 674 [PMID: 11826156 DOI: 10.1113/jphysiol.2001.015115]

12 **Talley NA**, Chen F, King D, Jones M, Talley NJ. Short-chain fatty acids in the treatment of radiation proctitis: a randomized, double-blind, placebo-controlled, cross-over pilot trial. *Dis Colon Rectum* 1997; **40**: 1046-1050 [PMID: 9293933 DOI: 10.1007/BF02050927]

13 **Pinto A**, Fidalgo P, Cravo M, Midões J, Chaves P, Rosa J, dos Anjos Brito M, Leitão CN. Short chain fatty acids are effective in short-term treatment of chronic radiation proctitis: randomized, double-blind, controlled trial. *Dis Colon Rectum* 1999; **42**: 788-795; discussion 795-796 [PMID: 10378604 DOI: 10.1007/BF02236937]

14 **Senthil Kumar R**, Vivekanandan S, Pai D, Sahai A, Narasimhan R. Prospective randomised double blind controlled trial to compare rectal sucralfate-steroid combination enema and short chain fatty acid enema in patients with radiation induced proctosigmoiditis. A dissertation submitted to Pondicherry University in partial fulfillment of the requirement for the award of the degree of M.S. Surgery. 2001

15 **Rubinstein E**, Ibsen T, Rasmussen RB, Reimer E, Sørensen BL. Formalin treatment of radiation-induced hemorrhagic proctitis. *Am J Gastroenterol* 1986; **81**: 44-45 [PMID: 3484606]

16 **Seow-Choen F**, Goh HS, Eu KW, Ho YH, Tay SK. A simple and effective treatment for hemorrhagic radiation proctitis using formalin. *Dis Colon Rectum* 1993; **36**: 135-138 [PMID: 8425416 DOI: 10.1007/BF02051168]

17 **Mathai V**, Seow-Choen F. Endoluminal formalin therapy for haemorrhagic radiation proctitis. *Br J Surg* 1995; **82**: 190 [PMID: 7749685 DOI: 10.1002/bjs.1800820216]

18 **Biswal BM**, Lal P, Rath GK, Shukla NK, Mohanti BK, Deo S. Intrarectal formalin application, an effective treatment for grade III haemorrhagic radiation proctitis. *Radiother Oncol* 1995; **35**: 212-215 [PMID: 7480824 DOI: 10.1016/0167-8140(95)01565-X]

19 **Saclarides TJ**, King DG, Franklin JL, Doolas A. Formalin instillation for refractory radiation-induced hemorrhagic proctitis. Report of 16 patients. *Dis Colon Rectum* 1996; **39**: 196-199 [PMID: 8620787 DOI: 10.1007/BF02068075]

20 **Roche B**, Chautems R, Marti MC. Application of formaldehyde for treatment of hemorrhagic radiation-induced proctitis. *World J Surg* 1996; **20**: 1092-1094; discussion 1094-1095 [PMID: 8798370 DOI: 10.1007/s002689900166]

21 **Chapuis P**, Dent O, Bokey E, Galt E, Zelas P, Nicholls M, Yuile P, Mameghan H. The development of a treatment protocol for patients with chronic radiation-induced rectal bleeding. *Aust N Z J Surg* 1996; **66**: 680-685 [PMID: 8855923 DOI: 10.1111/j.1445-2197.1996.tb00717.x]

22 **Yegappan M**, Ho YH, Nyam D, Leong A, Eu KW, Seow C. The surgical management of colorectal complications from irradiation for carcinoma of the cervix. *Ann Acad Med Singapore* 1998; **27**: 627-630 [PMID: 9919329]

23 **Counter SF**, Froese DP, Hart MJ. Prospective evaluation of formalin therapy for radiation proctitis. *Am J Surg* 1999; **177**: 396-398 [PMID: 10365878 DOI: 10.1016/S0002-9610(99)00072-0]

24 **Pikarsky AJ**, Belin B, Efron J, Weiss EG, Nogueras JJ, Wexner SD. Complications following formalin installation in the treatment of radiation induced proctitis. *Int J Colorectal Dis* 2000; **15**: 96-99 [PMID: 10855551 DOI: 10.1007/s003840050240]

25 **Ismail MA**, Qureshi MA. Formalin dab for haemorrhagic radiation proctitis. *Ann R Coll Surg Engl* 2002; **84**: 263-264 [PMID: 12215030]

26 **Luna-Pérez P**, Rodríguez-Ramírez SE. Formalin instillation for refractory radiation-induced hemorrhagic proctitis. *J Surg Oncol* 2002; **80**: 41-44 [PMID: 11967906 DOI: 10.1002/jso.10095]

27 **Ouwendijk R**, Tetteroo GW, Bode W, de Graaf EJ. Local formalin instillation: an effective treatment for uncontrolled radiation-induced hemorrhagic proctitis. *Dig Surg* 2002; **19**: 52-55 [PMID: 11961356 DOI: 10.1159/000052006]

28 **Chautems RC**, Delgadillo X, Rubbia-Brandt L, Deleaval JP, Marti MC, Roche B. Formaldehyde application for haemorrhagic radiation-induced proctitis: a clinical and histological study. *Colorectal Dis* 2003; **5**: 24-28 [PMID: 12780922 DOI: 10.1046/j.1463-1318.2003.00396.x]

29 **Parikh S**, Hughes C, Salvati EP, Eisenstat T, Oliver G, Chinn B, Notaro J. Treatment of hemorrhagic radiation proctitis with 4 percent formalin. *Dis Colon Rectum* 2003; **46**: 596-600 [PMID: 12792434 DOI: 10.1007/s10350-004-6614-1]

30 **de Parades V**, Etienney I, Bauer P, Bourguignon J, Meary N, Mory B, Sultan S, Taouk M, Thomas C, Atienza P. Formalin application in the treatment of chronic radiation-induced hemorrhagic proctitis--an effective but not risk-free procedure: a prospective study of 33 patients. *Dis Colon Rectum* 2005; **48**: 1535-1541 [PMID: 15933799 DOI: 10.1007/s10350-005-0030-z]

31 **Haas EM**, Bailey HR, Farragher I. Application of 10 percent formalin for the treatment of radiation-induced hemorrhagic proctitis. *Dis Colon Rectum* 2007; **50**: 213-217 [PMID: 17080283 DOI: 10.1007/s10350-006-0707-y]

32 **Yeoh E**, Tam W, Schoeman M, Moore J, Thomas M, Botten R, Di Matteo A. Argon plasma coagulation therapy versus topical formalin for intractable rectal bleeding and anorectal dysfunction after radiation therapy for prostate carcinoma. *Int J Radiat Oncol Biol Phys* 2013; **87**: 954-959 [PMID: 24113059 DOI: 10.1016/j.ijrobp.2013.08.034]

33 **Guo GH**, Yu FY, Wang XJ, Lu F. A randomized controlled clinical trial of formalin for treatment of chronic hemorrhagic radiation proctopathy in cervical carcinoma patients. *Support Care Cancer* 2015; **23**: 441-446 [PMID: 25128068 DOI: 10.1007/s00520-014-2401-2]

34 **Wong MT**, Lim JF, Ho KS, Ooi BS, Tang CL, Eu KW. Radiation proctitis: a decade's experience. *Singapore Med J* 2010; **51**: 315-319 [PMID: 20505910]

35 **Alfadhli AA**, Alazmi WM, Ponich T, Howard JM, Prokopiw I, Alaqeel A, Gregor JC. Efficacy of argon plasma coagulation compared to topical formalin application for chronic radiation proctopathy. *Can J Gastroenterol* 2008; **22**: 129-132 [PMID: 18299729 DOI: 10.1155/2008/964912]

36 **Sahakitrungruang C**, Patiwongpaisarn A, Kanjanasilp P, Malakorn S, Atittharnsakul P. A randomized controlled trial comparing colonic irrigation and oral antibiotics administration versus 4% formalin application for treatment of hemorrhagic radiation proctitis. *Dis Colon Rectum* 2012; **55**: 1053-1058 [PMID: 22965404 DOI: 10.1097/DCR.0b013e318265720a]

37 **Rustagi T**, Mashimo H. Endoscopic management of chronic radiation proctitis. *World J Gastroenterol* 2011; **17**: 4554-4562 [PMID: 22147960 DOI: 10.3748/wjg.v17.i41.4554]

38 **Johnston MJ**, Robertson GM, Frizelle FA. Management of late complications of pelvic radiation in the rectum and anus: a review. *Dis Colon Rectum* 2003; **46**: 247-259 [PMID: 12576899 DOI: 10.1007/s10350-004-6530-4]

39 **Jensen DM**, Machicado GA, Cheng S, Jensen ME, Jutabha R. A randomized prospective study of endoscopic bipolar electrocoagulation and heater probe treatment of chronic rectal bleeding from radiation telangiectasia. *Gastrointest Endosc* 1997; **45**: 20-25 [PMID: 9013165 DOI: 10.1016/S0016-5107(97)70298-0]

40 **Taylor JG**, Disario JA, Bjorkman DJ. KTP laser therapy for bleeding from chronic radiation proctopathy. *Gastrointest Endosc* 2000; **52**: 353-357 [PMID: 10968849 DOI: 10.1067/mge.2000.107726]

41 **Taylor JG**, DiSario JA, Buchi KN. Argon laser therapy for hemorrhagic radiation proctitis: long-term results. *Gastrointest Endosc* 1993; **39**: 641-644 [PMID: 8224685 DOI: 10.1016/S0016-5107(93)70215-1]

42 **Karamanolis G**, Psatha P, Triantafyllou K. Endoscopic treatments for chronic radiation proctitis. *World J Gastrointest Endosc* 2013; **5**: 308-312 [PMID: 23858374 DOI: 10.4253/wjge.v5.i7.308]

43 **Sebastian S**, O'Connor H, O'Morain C, Buckley M. Argon plasma coagulation as first-line treatment for chronic radiation proctopathy. *J Gastroenterol Hepatol* 2004; **19**: 1169-1173 [PMID: 15377295 DOI: 10.1111/j.1440-1746.2004.03448.x]

44 **Silva RA**, Correia AJ, Dias LM, Viana HL, Viana RL. Argon plasma coagulation therapy for hemorrhagic radiation proctosigmoiditis. *Gastrointest Endosc* 1999; **50**: 221-224 [PMID: 10425416 DOI: 10.1016/S0016-5107(99)70228-2]

45 **Dees J**, Meijssen MA, Kuipers EJ. Argon plasma coagulation for radiation proctitis. *Scand J Gastroenterol Suppl* 2006; **(243)**: 175-178 [PMID: 16782638 DOI: 10.1080/00365520600664300]

46 **Villavicencio RT**, Rex DK, Rahmani E. Efficacy and complications of argon plasma coagulation for hematochezia related to radiation proctopathy. *Gastrointest Endosc* 2002; **55**: 70-74 [PMID: 11756918 DOI: 10.1067/mge.2002.119877]

47 **Lam MC**, Parliament M, Wong CK. Argon plasma coagulation for the treatment of hemorrhagic radiation colitis. *Case Rep Gastroenterol* 2012; **6**: 446-451 [PMID: 22933988 DOI: 10.1159/000339462]

48 **Kwan V**, Bourke MJ, Williams SJ, Gillespie PE, Murray MA, Kaffes AJ, Henriquez MS, Chan RO. Argon plasma coagulation in the management of symptomatic gastrointestinal vascular lesions: experience in 100 consecutive patients with long-term follow-up. *Am J Gastroenterol* 2006; **101**: 58-63 [PMID: 16405534 DOI: 10.1111/j.1572-0241.2006.00370.x]

49 **Swan MP**, Moore GT, Sievert W, Devonshire DA. Efficacy and safety of single-session argon plasma coagulation in the management of chronic radiation proctitis. *Gastrointest Endosc* 2010; **72**: 150-154 [PMID: 20493484 DOI: 10.1016/j.gie.2010.01.065]

50 **Rotondano G**, Bianco MA, Marmo R, Piscopo R, Cipolletta L. Long-term outcome of argon plasma coagulation therapy for bleeding caused by chronic radiation proctopathy. *Dig Liver Dis* 2003; **35**: 806-810 [PMID: 14674672 DOI: 10.1016/S1590-8658(03)00454-7]

51 **Taïeb S**, Rolachon A, Cenni JC, Nancey S, Bonvoisin S, Descos L, Fournet J, Gérard JP, Flourié B. Effective use of argon plasma coagulation in the treatment of severe radiation proctitis. *Dis Colon Rectum* 2001; **44**: 1766-1771 [PMID: 11742159 DOI: 10.1007/BF02234452]

52 **Chruscielewska-Kiliszek MR**, Regula J, Polkowski M, Rupinski M, Kraszewska E, Pachlewski J, Czaczkowska-Kurek E, Butruk E. Sucralfate or placebo following argon plasma coagulation for chronic radiation proctitis: a randomized double blind trial. *Colorectal Dis* 2013; **15**: e48-e55 [PMID: 23006660 DOI: 10.1111/codi.12035]

53 **Nikfarjam M**, Faulx A, Laughinghouse M, Marks JM. Feasibility of radiofrequency ablation for the treatment of chronic radiation proctitis. *Surg Innov* 2010; **17**: 92-94 [PMID: 20504783 DOI: 10.1177/1553350610365701]

54 [**Zhou C**](http://www.ncbi.nlm.nih.gov/pubmed/?term=Zhou%20C%5BAuthor%5D&cauthor=true&cauthor_uid=20593010), [Adler DC](http://www.ncbi.nlm.nih.gov/pubmed/?term=Adler%20DC%5BAuthor%5D&cauthor=true&cauthor_uid=20593010), [Becker L](http://www.ncbi.nlm.nih.gov/pubmed/?term=Becker%20L%5BAuthor%5D&cauthor=true&cauthor_uid=20593010), [Chen Y](http://www.ncbi.nlm.nih.gov/pubmed/?term=Chen%20Y%5BAuthor%5D&cauthor=true&cauthor_uid=20593010), [Tsai TH](http://www.ncbi.nlm.nih.gov/pubmed/?term=Tsai%20TH%5BAuthor%5D&cauthor=true&cauthor_uid=20593010), [Figueiredo M](http://www.ncbi.nlm.nih.gov/pubmed/?term=Figueiredo%20M%5BAuthor%5D&cauthor=true&cauthor_uid=20593010), [Schmitt JM](http://www.ncbi.nlm.nih.gov/pubmed/?term=Schmitt%20JM%5BAuthor%5D&cauthor=true&cauthor_uid=20593010), [Fujimoto JG](http://www.ncbi.nlm.nih.gov/pubmed/?term=Fujimoto%20JG%5BAuthor%5D&cauthor=true&cauthor_uid=20593010), [Mashimo H](http://www.ncbi.nlm.nih.gov/pubmed/?term=Mashimo%20H%5BAuthor%5D&cauthor=true&cauthor_uid=20593010). Effective treatment of chronic radiation proctitis using radiofrequency ablation. *Therap Adv Gastroenterol* 2009; **2**: 149-156 [PMID: 20593010 DOI: 10.1177/1756283X08103341]

55 **Eddi R**, Depasquale JR. Radiofrequency ablation for the treatment of radiation proctitis: a case report and review of literature. *Therap Adv Gastroenterol* 2013; **6**: 69-76 [PMID: 23320051 DOI: 10.1177/1756283x12456895]

56 **Dray X**, Battaglia G, Wengrower D, Gonzalez P, Carlino A, Camus M, Adar T, Pérez-Roldán F, Marteau P, Repici A. Radiofrequency ablation for the treatment of radiation proctitis. *Endoscopy* 2014; **46**: 970-976 [PMID: 25290097 DOI: 10.1055/s.0034.1377756]

57 **Ritch CR**, Katz AE. Update on cryotherapy for localized prostate cancer. *Curr Urol Rep* 2009; **10**: 206-211 [PMID: 19371478 DOI: 10.1007/s11934-009-0035-6]

58 **Hou JK**, Abudayyeh S, Shaib Y. Treatment of chronic radiation proctitis with cryoablation. *Gastrointest Endosc* 2011; **73**: 383-389 [PMID: 21295650 DOI: 10.1016/j.gie.2010.10.044]

59 **Moawad FJ**, Maydonovitch CL, Horwhat JD. Efficacy of cryospray ablation for the treatment of chronic radiation proctitis in a pilot study. *Dig Endosc* 2013; **25**: 174-179 [PMID: 23362977 DOI: 10.1111/j.1443-1661.2012.01355.x]

60 **Clarke RE**, Tenorio LM, Hussey JR, Toklu AS, Cone DL, Hinojosa JG, Desai SP, Dominguez Parra L, Rodrigues SD, Long RJ, Walker MB. Hyperbaric oxygen treatment of chronic refractory radiation proctitis: a randomized and controlled double-blind crossover trial with long-term follow-up. *Int J Radiat Oncol Biol Phys* 2008; **72**: 134-143 [PMID: 18342453 DOI: 10.1016/j.ijrobp.2007.12.048]

61 **Tahir AR**, Westhuyzen J, Dass J, Collins MK, Webb R, Hewitt S, Fon P, McKay M. Hyperbaric oxygen therapy for chronic radiation-induced tissue injuries: Australasia's largest study. *Asia Pac J Clin Oncol* 2015; **11**: 68-77 [PMID: 25382755 DOI: 10.1111/ajco.12289]

62 **Warren DC**, Feehan P, Slade JB, Cianci PE. Chronic radiation proctitis treated with hyperbaric oxygen. *Undersea Hyperb Med* 1997; **24**: 181-184 [PMID: 9308141]

63 **Girnius S**, Cersonsky N, Gesell L, Cico S, Barrett W. Treatment of refractory radiation-induced hemorrhagic proctitis with hyperbaric oxygen therapy. *Am J Clin Oncol* 2006; **29**: 588-592 [PMID: 17148996 DOI: 10.1097/01.coc.0000236004.95384.5b]

64 **Jones K**, Evans AW, Bristow RG, Levin W. Treatment of radiation proctitis with hyperbaric oxygen. *Radiother Oncol* 2006; **78**: 91-94 [PMID: 16337705 DOI: 10.1016/j.radonc.2005.11.004]

65 **Dall'Era MA**, Hampson NB, Hsi RA, Madsen B, Corman JM. Hyperbaric oxygen therapy for radiation induced proctopathy in men treated for prostate cancer. *J Urol* 2006; **176**: 87-90 [PMID: 16753375 DOI: 10.1016/S0022-5347(06)00491-5]

66 **Kennedy M**, Bruninga K, Mutlu EA, Losurdo J, Choudhary S, Keshavarzian A. Successful and sustained treatment of chronic radiation proctitis with antioxidant vitamins E and C. *Am J Gastroenterol* 2001; **96**: 1080-1084 [PMID: 11316150 DOI: 10.1111/j.1572-0241.2001.03742.x]

67 **Ehrenpreis ED**, Jani A, Levitsky J, Ahn J, Hong J. A prospective, randomized, double-blind, placebo-controlled trial of retinol palmitate (vitamin A) for symptomatic chronic radiation proctopathy. *Dis Colon Rectum* 2005; **48**: 1-8 [PMID: 15690650 DOI: 10.1007/s10350-004-0821-7]

68 **Jao SW**, Beart RW, Gunderson LL. Surgical treatment of radiation injuries of the colon and rectum. *Am J Surg* 1986; **151**: 272-277 [PMID: 3946764 DOI: 10.1016/0002-9610(86)90086-3]

69 **Johansson C**. Endoscopic dilation of rectal strictures: a prospective study of 18 cases. *Dis Colon Rectum* 1996; **39**: 423-428 [PMID: 8878503 DOI: 10.1007/BF02054058]

70 **Ayerdi J**, Moinuddeen K, Loving A, Wiseman J, Deshmukh N. Diverting loop colostomy for the treatment of refractory gastrointestinal bleeding secondary to radiation proctitis. *Mil Med* 2001; **166**: 1091-1093 [PMID: 11778411]

71 **Pricolo VE**, Shellito PC. Surgery for radiation injury to the large intestine. Variables influencing outcome. *Dis Colon Rectum* 1994; **37**: 675-684 [PMID: 8026234 DOI: 10.1007/BF02054411]

72 **Anseline PF**, Lavery IC, Fazio VW, Jagelman DG, Weakley FL. Radiation injury of the rectum: evaluation of surgical treatment. *Ann Surg* 1981; **194**: 716-724 [PMID: 7305485 DOI: 10.1097/00000658-198112000-00010]

73 **Lucarotti ME**, Mountford RA, Bartolo DC. Surgical management of intestinal radiation injury. *Dis Colon Rectum* 1991; **34**: 865-869 [PMID: 1914719 DOI: 10.1007/BF02049698]

74 **Denton AS**, Bentzen SM, Maher EJ. How useful are observational reports in the evaluation of interventions for radiation morbidity?: an analysis of formalin therapy for late radiation proctitis. *Radiother Oncol* 2002; **64**: 291-295 [PMID: 12242117 DOI: 10.1016/S0167-8140(02)00138-X]

75 **Henson C**. Chronic radiation proctitis: issues surrounding delayed bowel dysfunction post-pelvic radiotherapy and an update on medical treatment. *Therap Adv Gastroenterol* 2010; **3**: 359-365 [PMID: 21180615 DOI: 10.1177/1756283x10371558]

**P- Reviewer:** Jongen J, Linard C, Pigo F **S- Editor:** Gong XM

 **L- Editor:** **E- Editor:**



Figure 1b

Figure 1a

**Figure 1 Chronic radiation proctitis.** A: Chronic radiation proctitis (low power view).This picture shows the mucosa with severe oedema, non-specific inflammation, lymphocytosis, hyalinization in the stroma and fibrin thrombi in the postcapillary venules (Hematoxylin and Eosin stain, 10×); B: Chronic radiation proctitis (High power view). This picture shows two veins in the lamina propria, one with patchy occlusive fibrin thrombus. The wall shows thickening and hyalinization. A dense non-specific inflammation including few eosinophils also seen (Hematoxylin and Eosin stain, 40×).