IBD—Impact of Colonic pH, Onset of Action and other Factors in Modern Therapeutic Approach

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Abstract
Pharmacotherapy rediscovered the management of ulcerative colitis and Crohn’s disease. Novel drug delivery systems are popular in treating IBD. The treatment of IBD with novel drug delivery system and other medicines involves complicated mechanism. The onset of action, pH conditions of the colon in normal and diseased condition of the patient and adverse effects of the drugs makes the greatest impact in designing successful IBD therapy.

Keywords: Inflammatory bowel disease; Crohn’s disease; Diarrhea; Hypokalemia

Introduction

Inflammatory bowel disease (IBD) is an idiopathic disease. Novel drug delivery systems are very successful in the treatment of colonic diseases. The colonic environment can be explored for local and systemic delivery of drugs and other bioactive compounds such as hormones, insulin, vasopressin and other plant ingredients. The colon is the major region of the GIT. The common colonic disease such as diverticular inflammatory bowel disease (IBD) which includes Crohn’s disease and ulcerative colitis, colitis ulcerosa diversional colitis, ischemic colitis, colon cancer and lymphoma of the colon can be treated successfully by modern therapeutic approaches. In past decades the general pathophysiology of inflammatory bowel disease was described on the basis of clinical manifestation. The investigators and clinicians are struggled to provide the effective therapy for IBD due to its dismaying clinical manifestation. The causes of inflammatory bowel disease is multi factorial and may be resulted from inappropriate activation of mucosal immune system, inflammatory responses, generic factors, candidate genes, chromosome location etc.

The infectious organism such as Escherichia coli, measles virus, cytomegalo virus and factor like saturated fats, milk products, allergic foods may also be the cause of the IBD. General pathophysiology of ulcerative colitis and Crohn’s disease is limited to large intestine often inflammation and ulcers occurs in the inner lining of the large intestine or in mucosal layer. The ulcerative colitis resulting in diarrhea, blood and pus. Crohn’s disease, otherwise called regional enteritis. Crohn’s disease involves any part of the gastrointestinal tract from mouth to anus with the inflammation extending through the bowel wall to the serosal surface. Both the diseases used to have waxing and waning intensity and severity [1-3].

Crohn’s disease and ulcerative colitis significantly differ from each. The treatment methods are common to both the diseases. Many extra intestinal manifestations are shared by both the diseases occurs in adults and children. The signs and symptoms of ulcerative colitis involves diarrhea with the presence of blood and mucus. Weight loss, abdominal pain, painful bowel movements, abdominal cramps and extra intestinal symptoms like arthritic knees may be observed in youngsters. The pathophysiology of ulcerative colitis showed an increased amount of colonic sulfate reducing bacteria. This may be due to the result of higher concentration of hydrogen sulfide toxic gas. Some reports suggested that sulfur containing red meat, alcohol consumption also increased the disease relapsing in patients in remission. The ulcerative colitis occur in 38-100 for every 100,000 in the US. The disease occurs predominantly in northern countries 0.1% population. Ulcerative colitis has no known cause and it is treated as autoimmune disease.

General pathophysiology of IBD

The pathophysiology of IBD is yet to be revealed. The IBD is still under active research for its cause. The general pathway is the inflammation of the mucosal lining of the intestinal tract, which causes ulceration, edema, bleeding associated with fluid and electrolyte loss. The researches have been struggling to identify the gene involved in beginning of IBD. The early linkage discovered was on chromosome 16 (IBD 1 gene). The identification of 'IBD 1 gene' leads to the identification of NOD2 gene also called as CARD 15 which was declared as the first gene clearly associated with IBD. This gene also has the susceptibility to Crohn’s disease. Studies also identified the susceptibility genes on chromosome 5 and 6 (5Q31, 6P21, 19P). CARD 15 is a polymorphic gene involved in the innate immune system, which show 60 different variations. Three of these variants played a vital role in IBD development. This means that these genes allowing the IBD to occur, but not a causative source. The main point to be considered here is a presence of genes does not develop the disease, but may allow the disease to occur (Figure 1).

Diagnosis and endoscopy

H and E biopsy of colonic tissue usually showing the crypts. Abscess is an indicator for ulcerative colitis. The initial blood diagnostic tests involves blood count test for the assessment of anemia, thrombocytosis, high platelet count, the renal function test, test for chronic diarrhea associated with hypokalemia, hypomagnesemia, liver function test, primary sclerosing cholangitis, X ray, urine analysis also used as a diagnostic tool to investigate ulcerative colitis. Endoscopy is the best available tool to diagnose for ulcerative colitis. The endoscopy reveals the loss of vascular appearance of the colon and also identifies
erythema and damaged mucosa, superficial ulceration and pseudopolyps in the colon [4].

Figure 1: Inflammatory bowel disease.

Management of IBD

The treatment approaches depending on the severity of the disease. Pharmacotherapy is significantly successful in the treatment of ulcerative colitis. The optimized medical approach differs with a physician. The main objective of the treatment is to induce the remission followed by the maintenance therapy to prevent relapse. The drug treatment involves aminosalicylates such as sulfasalazine cortico steroids (Prednisolone), immune suppressive agents (azathioprine) and biological agents such as infliximab. Ulcerative colitis generally be cured by surgical removal of the large intestine, which is not recommended in the early stages. Drug delivery systems with modified and targeted drug delivery to the site and prodrugs approach can also successfully used in IBD management. The list of the drugs approved commonly for ulcerative colitis and Crohn’s are given below [5] (Table 1).

Table 1: List of the drugs approved commonly for ulcerative colitis and Crohn’s.

<table>
<thead>
<tr>
<th>Corticosteroids</th>
<th>hydrocortisone, methylprednisolone, Prednisolone as I.V route, oral or retention enemas.</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-Aminosalicylic acid compounds</td>
<td>Sulphasalazine combination of sulphapyridine and 5-aminosalicylic acid joined by an ace - bond. Poorly absorbed, split by bacteria in the colon. 5-ASA part is the active moiety; sulphapyridine part can cause sulfonamide toxicity. Reduces relapses in UC and used for treatment of exacerbations.</td>
</tr>
<tr>
<td>Mesalamine</td>
<td>5-ASA. Delayed release, pH-dependent preparations to allow release in the colon. Better tolerated than sulphasalazine, but still carries risk of hematological side effects.</td>
</tr>
<tr>
<td>Olsalazine</td>
<td>2 molecules of 5-ASA linked by azo-bond split by colon bacteria.</td>
</tr>
<tr>
<td>Balsalazide</td>
<td>prodrg of 5-ASA</td>
</tr>
<tr>
<td>Immunosuppressants</td>
<td>Azathioprine</td>
</tr>
<tr>
<td>Metronidazole</td>
<td>Perianal Crohn’s disease.</td>
</tr>
<tr>
<td>Infliximab</td>
<td>For severe active Crohn’s disease refractory to treatment with steroids .</td>
</tr>
<tr>
<td>Monoclonal Antibody</td>
<td>Inhibits the pro-inflammatory cytokine TNF-α.</td>
</tr>
<tr>
<td>NSAID</td>
<td>Ibuprofen, Diclofenac, Indomethacin Analgesic – rapid (full effect within 1 week) Anti-inflammatory (full effect within 3 weeks)</td>
</tr>
</tbody>
</table>

Various studies on colonic pH conditions

The various studies conducted worldwide by following different techniques registered the pH changes in normal and IBD colon. The Table 2 displays the colonic pH conditions in the normal healthy volunteer patient with ulcerative colitis and Crohn’s disease studied by various researchers. Changes in the intestine and colonic pH also considered as an important factor in the treatment of ulcerative colitis and IBD. The design and development of novel treatment and novel drug delivery systems are also significantly influenced by the changes in the luminal pH conditions and that should be considered in the treatment of IBD. The formulation also developed to deliver an active agent directly to the inflammation site. This approach reduces the absorption of drugs in the upper GI tract as well as the systemic side effects. This method involves pH dependent drug delivery systems (Asacol, Mesacol and Salofalk). Another common technique involves bacterial enzymatic metabolism (sulphasalazine, olsalazine and balsalazide) which also affected by changes in colonic pH.

Treatment methods for IBD and its limitations

Treatment usually based on reports of clinical history, physical parameters endoscopy, radiology, histology and regular laboratory tests. The study on these reports gives the clear idea about IBD and also distinguishes the ulcerative colitis and Crohn’s disease, but it is very difficult to distinguish ulcerative colitis and Crohn’s disease in at least 10% of the population for them IBD is limited to the colon [17,18]. Long time management of inflammatory bowel disease involves the drug therapy and lifestyle management. The therapy may be started with antidiarrheal in the beginning to give the symptomatic relief and the treatment should be focused on reducing the inflammation once the symptoms are subsiding.
<table>
<thead>
<tr>
<th>S.no</th>
<th>Study</th>
<th>Patient</th>
<th>Small bowel pH</th>
<th>Colonic pH</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Proximal</td>
<td>Distal</td>
</tr>
<tr>
<td>1</td>
<td>[6]</td>
<td>39 normals</td>
<td>6.4</td>
<td>7.3</td>
</tr>
<tr>
<td>2</td>
<td>[7]</td>
<td>13 normals</td>
<td>6.4</td>
<td>7.4</td>
</tr>
<tr>
<td>3</td>
<td>[8]</td>
<td>7 normals</td>
<td>6.6</td>
<td>7.4</td>
</tr>
<tr>
<td>4</td>
<td>[9]</td>
<td>2 normal+7 GI disorder cases</td>
<td>5.5-7.5</td>
<td>6.5 -7.5</td>
</tr>
<tr>
<td>5</td>
<td>[10]</td>
<td>66 normals</td>
<td>6.6</td>
<td>7.5</td>
</tr>
</tbody>
</table>

**Patient with Ulcerative Colitis**

| 6    | [11]  | 7 active | 6 inactive | 6.1 | 5.9-6.6 | 7.2 | 4.7 | 4.9-5.5 | - |
| 7    | [12]  | 3 active | 3 very active | Normal | Normal | Normal | Normal | Normal | - |
| 8    | [13]  | 6 active | 7.3 | 8.3 | 6.7 | 6.7 |
| 9    | [14]  | 7 active | 4 inactive | 6.8 | 6.6 | 8.2 | 7.9 | 6.5 | 6.5 |
| 10   | Eve   | 4 active | 6.5 | 6.8 | 5.5 | 7.5 |

**Patient with Crohn’s disease**

| 11   | [7]   | 9 with ileo cecal resection | 6.3 | 7.3 | 6.7 | NA |
| 12   | [16]  | 3 active | 1 inactive | 7.2 | 7.8 | 5.3 | 5.3 |
| 13   | [15]  | 12 active | 6.5 | 7.5 | 6.2 | 6.5 |
| 14   | [14]  | 5 active | 7 inactive | 6.5 | 6.8 | 7.9 | 6.5 | 6.5 | 6.5 |

| NA- Data not available |

### Table 2: pH conditions in normal and inflamed colon.

A correct diet and nutritions are advisable as a supporting measures for the successful IBD management. A drugs such as 5 amino salicylic acid (Mesalamine) were used in the treatment of IBD [19]. 5–ASA widely replaces the sulfasalazine for its safety and less adverse effects. Mesalamine (5-ASA) not considered as the very potent anti-inflammatory agent, but shown to be effective in IBD patients, but also fails to show significant improvement in the set of patients Affected with IBD.

**Important considerations of amino salicylates**

- Amino salicylates are unstabslets in gastric acid
- Rapidly absorbed in the small intestine.
- The safety and tolerability of amino salicylates are the reason behind the withdrawal of these agents.
- The greater number of withdrawals was reported with sulfasalazine 3 g/day than balsalazide 6.75 g/day [20,21].
- Balsalazide has a more adverse drug reactions than delayed release Mesalamine [22].
- The unstabslets nature of Mesalamine in the upper GI region paves the way for developing them into novel drug delivery systems such as delayed release formulation based on enteric coating pH dependent release system which breaks at ileal or colonic PH or environment. Prodrug based systems, microflora activated system based on poly surcharges such as Pectin, Guar gum, Chitosan, timed release system, etc [23,24].

**Corticosteroids**

- Corticosteroids are recommended when 5–ASA compounds are inadequate in producing the expected results.
- Topical corticosteroids (enemas) used in the patient with ulcerative colitis, Prednisolone 60 mg/day is used orally in the treatment of ulcerative colitis and Crohn’s disease. The mechanism of action of corticosteroids was well known and is
acted by inhibiting the several inflammatory pathways and stimulation of lymphocyte apoptosis [25].

- Corticosteroids are known for its systemic side effects in which adrenal suppression and osteoporosis, corticosteroids induced hypertension and diabetes are well noted.

- Intravenous administration of Prednisolone also recommended when the condition of the patient is severe [26,27].

- The main disadvantage is the usage of these corticosteroids are difficult in colonic disease than ulcerative colitis due to the variation in colonic pH, transit time and bacterial metabolism [30,31].

**Immune modulatory therapy**

- Azathiopurine and Mercaptopurine (6-mercaptopurine) are the most commonly used immune modulator. These are the derived products of thioguanines. This recommended when corticosteroids cannot be withdrawn from the patient.

- Although azathiopurine earlier reported for side effects and producing an increased risk of lymphoma, which is becoming a highly recommended immune suppressive agent [32-35].

- The efficacy of the drug was mostly depend on the dose, and the optimal dosage was 2.0-2.5 mg/kg/day and 1.0-1.5 mg/kg/day was found optimal for Azathiopurine and Mercaptopurine respectively [36].

- The side effects of these agents are limited the usage of Azathiopurine and Mercaptopurine. So that Azathiopurine should be prescribed with caution and step wise approach. The serious adverse effects include bone marrow suppression, variation in the white cell counts [37,38].

- Cyclosporine is recommended in the patient with steroid refractory ulcerative colitis [39]. Cyclosporin inhibits the cellular immune response by blocking cytokine production by T lymphocytes through calcineurin dependent pathway [40,41].

- Cyclosporins provide the rapid onset of action by intravenous route with significant clinical improvement about a week [42] cyclosporins should not be recommended in the treatment of Crohn’s disease which may cause severe personal or cutaneous fistula [42]. Although cyclosporins are considered as a alternative for corticosteroid therapy their adverse effects limits the usage.

- Tacrolimus and mycopholate mofetil also effective in the treatment of IBD. Tacrolimus is a macrolide immune suppression which inhibits the immune response through a calcineurin dependent pathway [43] Tacrolimus can induce remission in adults and children [44,45] Recent days studies supported the usage in the treatment of corticosteroids dependent Crohn’s disease in remission and relapsing [46-49].

- It can be given as weekly injection 15 mg and 25 mg weekly by IM and SC respectively. The mechanism of action of this drug is unclear and also known for its side effects such as immune suppression, interstitial pneumonitis associated with non-productive cough, dyspnea and hepatic fibrosis.

**Biological agents**

- Biological agents such as infliximab (protypical anti – TNF agent) made the significant advancements in the treatment of crohns disease. The mechanism of action of infliximab yet to be explained completely. It is a chimeric monoclonal antibody. It is an agent which shows the significant results in the treatment of crohn’s disease but not in ulcerative colitis [50,51].

- Basiliximab was showing significant clinical remission in ulcerative colitis in pilot scale study [52].

- Daclizumab infusion showed the decreased clinical activity score in refractive ulcerative colitis patients. Natalizumab demonstrated a significant clinical response in active ulcerative colitis patients [53]. The rapid immuno modulatory agents are known for its severe adverse effects which sometimes fatal.

**Probiotics and antibiotics in IBD**

The studies worldwide supported the recognition of antibiotics in the treatment of IBD [54]. Other reports on probiotic also revealed that probiotic can be used as a supporting agent in the treatment of IBD. The usage of antibiotics in the treatment of ulcerative colitis is limited. In crohn’s disease metronidazole (750 mg/day/tid) found to be effective. The side effects of metronidazole such as neurotoxicity to be taken into consideration before and during the treatment. Metronidazole is effective in the treatment of Crohn’s disease, but showed no significant response in the treatment of ulcerative colitis.

The administration of probiotics is an excellent supporting approach in the management of crohn’s disease which is free of any side effects [55,56]. Lactobacillus acidophilus LA1 have the reported effect in immune enhancement adherent to human intestinal cells and balancing the microflora. Lactobacillus acidophilus NCFB is effective is effective in lowering faecal enzyme activity decreased fecal mutagenecity in the treatment of rotavirus diarrhea Crohn’s disease and antagonistic against carcinogenic bacteria. Lactobacillus casei shirota is effective in prevention of intestinal disturbances balancing intestinal bacteria and immune enhancement. Bifido bacterium bifidum is effective in treatment of rotavirus diarrhea and balancing intestinal microflora. Lactobacillus gasseric (ADH) is effective in the treatment of IBD [57-60].

**Importance of onset of Action in IBD Treatment**

- Pharmacotherapy brought the excellent changes in the management of ulcerative colitis and crohn’s disease. The onset of action plays a major role in various treatment. In case of the IBD onset of action is most important in management, maintenance of remission of disease.

- In multicentre trial revealed that onset of action of balsalazide was earlier than Mesalamine [61].

- The patient with Sulfasalazine intolerance can be successfully treated with Balsalazide, Olsalazine or Mesalazine.

- Although corticosteroids have role in maintenance and remission of ulcerative colitis. Their efficacy relying on rapid onset of action and anti-inflammatory activity which gives the consistency in treatment with corticosteroids in ulcerative colitis.

- The study on immunomodulatory agents revealed that the clinical benefits of thioguanine derivatives desired only after 4 months of therapy in Crohn’s disease [62].

- The clinical reports demonstrated that infliximab is effective with rapid onset of action that gives the improvement within days in
Crohn's disease but it is not effective in ulcerative colitis treatment [63,64].

The onset of action of therapy is most important in the treatment of IBD especially in the treatment of ulcerative colitis. Although novel biological agents, immune modulator and other novel agents are available, therapy is dominated by amino salicylates and corticosteroids. The importance of onset of action in IBD or UC therapy determines the efficacy of therapeutic agents and therapeutic strategies (Table 3).

**Advanced drug delivery systems in IBD**

- Targeting the drugs to the colon gaining the importance in treating GI disorders. The local disorder such as IBD, irritable bowel syndrome (IBS), carcinoma can be successfully treated by colonic deliveries.
- The advanced drug delivery systems prepared for the treatment of IBD mainly based on pH, transit time and micro flora activation.
- The coated systems with pH dependent polymers such as polymethacrylic acid derivatives (Eudrgists) are widely used for this purpose [65]. Polysachhrides such as Pectin, Chitosan, amylose and Guar gum can be successfully explored as colon drug delivery systems. The polysaccharide systems are found to be more successful because of their practicality and the abundant microflora of the colon.
- By combining the knowledge of threshold pH of the polymer and their solubility in different pH environments, duly, the system has been designed to release the drugs at target site exclusively on the colon [66,67].
- According to the various studies worldwide revealed and suggested fluctuation in the pH of the colon is due to various reasons [65].
- Some reports suggested that change in the G.I profile may occur in patient with IBD, which should be considered in developing delayed release formulations [65].
- Apparently the colon has lower pH value (6.5) than the small intestine (pH 7.0-7.8). The behavior of various pH sensitive polymers coated marketed products (Pentasa®, Asacol®, Salofalk®,) with human subjects indicated that there was a marked individual variation in urinary recovery of these drugs [66].

<table>
<thead>
<tr>
<th>Drug</th>
<th>Marketed product</th>
<th>Polymers technology used (or)</th>
<th>Site of release</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mesalamine</td>
<td>Asacol</td>
<td>Eudrigt S coating dissolves at pH&gt;7</td>
<td>Distal part of intestine and colon</td>
</tr>
<tr>
<td>Mesren</td>
<td></td>
<td>Eudrigt S coating dissolves at pH&gt;7</td>
<td>Distal part of intestine and colon</td>
</tr>
<tr>
<td>Salofalk</td>
<td></td>
<td>Eudrigt L coated tablet dissolves at pH&gt;6</td>
<td>Middle and Distal part of intestine and colon</td>
</tr>
<tr>
<td>Pentasa</td>
<td></td>
<td>Ethyl cellulose coated granules membrane controlled release</td>
<td>Stomach to colon</td>
</tr>
<tr>
<td>Budesonide</td>
<td>Entocort</td>
<td>Eudragit L 100-55 coated ethyl cellulose granules dissolves at pH&gt;5.5</td>
<td>Proximal intestine and colon</td>
</tr>
<tr>
<td>Sulfasalazine</td>
<td>Salazopyrine</td>
<td>Azo bond cleaved by colonic bacteria</td>
<td>Colon</td>
</tr>
</tbody>
</table>

Table 3: Various marketed products of IBD drugs.

- When observed in patients after administration of pentasa®, tablets there was an individual variation in urinary recovery [67].
- Mesalamine tablets manufactured by different companies have a different release profile when tested in various pH media.
- Research reports have clearly stated that the pH sensitive polymer-based colonic deliveries may not release the drug in the colon as expected.
- There are possibilities of the drug being released in advance, prior to entry of the terminal part of the G.I tract and or poor cumulative percentage of drug release occurring due to variation in the colonic pH during IBD [68]. Various studies proved that there was a fall in colonic pH in the IBD diseased colon [69].

**Conclusion**

Treatment and management of IBD is always a challenging aspect. The pharmacotherapy approaches moderately successful in the treatment of ulcerative colitis. The medical approach to treat this disease extensively differs with physician. The condition of patient and severity of the disease also influence the treatment method. The main objective of the IBD therapy is to induce the remission and to prevent the relapse. Since, there is no tailor made approach for this disease various factor to be consider before designing the treatment method to the patient. The marketed products are dominated by pH sensitive polymer based mechanism but it has its own drawbacks which were discussed in this work. The treatment methods such as amino salicylates, corticosteroid, immunosuppresssive agent, biological agents, surgical methods have reported disadvantages in the treatment of IBD. The longtime management of IBD must be well designed with the proper life style management. The onset of action plays a vital role in the management and treatment of ulcerative colitis and it has to be considered as an important factor in pharmacotherapy. All the advance drug delivery systems are not completely perfect devices. Study reports says that drug delivery systems in the treatment of IBD has to be monitored carfully. Before selecting the drug delivery system, a physician has to understand the mechanism of drug release in drug delivery system. The various natural polymers based drug delivery system also under research. Which can be considered as a alternative to pH sensitive devices. This review discussed some crucial parameters to be considered in the treatment of better IBD therapy.

**References**

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