ABSTRACT: Because the glucocorticosteroid receptor seems to be uniform in the human body, there is currently no support for a possibility of separating the therapeutic and adverse glucocorticosteroid actions at the receptor level. However, based on a new generation of glucocorticosteroids characterized by a high first pass metabolism in the liver, it seems possible today to reach a more selective topical therapy of inflammatory bowel disease. The properties of three new glucocorticosteroids are presented: the highly potent budesonide, fluticasone propionate and tixocortol pivalate — the latter with only low topical potency. Their properties can be exemplified by budesonide, which is currently the best documented compound. The topical potency of budesonide is 200 and 15 times higher than those of hydrocortisone and prednisolone, respectively. This means that there is a high potential for anti-inflammatory and immunosuppressive actions on rectal and bowel mucosa. The compound is metabolically stable in the bowel compartment, which allows full retention of glucocorticosteroid activity in the target organ. However, when absorbed and distributed to the liver, there is a 90% first pass hepatic metabolism to metabolites of very low potency. This suggests that after topical application to rectal or bowel mucosa, glucocorticosteroid activity in the systemic circulation is low. This is in contrast to prednisolone, which has a hepatic first pass metabolism of just 20%. Can J Gastroenterol 1990;4(7):407-414 (pour résumé, voir page 408)

Key Words: Budesonide, Corticosteroids, Fluticasone, Tixocortol pivalate

SINCE THEIR INTRODUCTION AS general anti-inflammatory agents in the early 1950s, the glucocorticosteroids are one of the mainstays in the therapy of active ulcerative colitis and Crohn’s disease. It was found early that symptomatic improvement could be attained by daily treatment with 30 mg or more of prednisolone (or its equivalent) administered via the oral and rectal routes. The therapeutic efficacy was confirmed by placebo controlled trials for ulcerative colitis in the mid 1950s, while it took a much longer time to attain such verification for Crohn’s disease (1-3).

DEVELOPMENT OF CURRENT GLUCOCORTICOSTEROID THERAPY IN IBD
The glucocorticosteroid era started with the introduction of cortisone, which is now known to be a prodrug with a very low affinity for the
Les derniers glucocorticostéroïdes et les maladies inflammatoires de l'intestin

RESUME: Parce que le récepteur des glucocorticostéroïdes semble uniforme dans tout l'organisme humain, il semble impossible de séparer les actions thérapeutiques et indésirables des glucocorticostéroïdes au niveau des récepteurs. Pourtant, grâce à une nouvelle génération de glucocorticostéroïdes caractérisés par un haut métabolisme de premier passage hépatique, on semble entrevoir l'éventualité d'une thérapie locale plus sélective dans les maladies inflammatoires de l'intestin. Les propriétés de trois de ces nouveaux médicaments sont présentées: le puissant budesonide, le propionate de fluticasone et le pivalate de tixocortol − ce dernier ne manifestant qu'une faible activité locale. Le mieux établi des trois, le budesonide sera examiné à titre d'exemple. L'activité locale du budesonide est de 200 à 15 fois supérieure à celle de l'hydrocortisone et de la prednisolone, respectivement. Cela signifie qu'il offre un potentiel anti-inflammatoire et immunsuppresseur élevé au niveau de la muqueuse rectale et intestinale. Le composé est métaboliquement stable dans le compartiment intestinal, ce qui autorise la pleine rétention de l'action glucocorticostéroïde au niveau de l'organe cible. Cependant, lorsqu'il est absorbé et parvient au foie, le produit est dégradé en métabolites de très faible puissance. Cela suggère qu'après application locale à hauteur du rectum et de l'intestin, l'activité systémique est faible. Ce résultat est à comparer à l'action de la prednisolone dont le métabolisme de premier passage est de l'ordre de 20%.

Figure 1) Schematic overview of glucocorticosteroid development

glucocorticosteroid receptor. Cortisone (or its close derivative, prednisone) has to be reduced in the liver to the corresponding 11-OH compound − hydrocortisone and prednisolone, respectively − to reach hormonal activity at the glucocorticoid and mineralocorticoid receptors. The chemical development of improved glucocorticosteroids for oral or other routes of systemic therapy culminated early with the introduction of prednisolone and 6-methylprednisolone, both having enhanced glucocorticoid but reduced mineralocorticoid activity. No major differentiation of the various glucocorticosteroid actions has been possible in systemic therapy, because all of these actions seem to be mediated via the same type of glucocorticosteroid receptor. To reduce the extent of serious adverse effects (eg, Cushing's syndrome, emotional disturbances, blocked hypothalamic-pituitary-adrenal axis, osteoporosis and connective tissue atrophy), high dose treatment with systemic glucocorticosteroid has to be limited to a few weeks and then tapered off to doses much lower than the 30 mg/kg mentioned in the introduction.

It was shown early that proctitis and distal colitis could be treated topically using retention enemas or foams containing hydrocortisone or its 21-ester (4-7), or prednisolone or its 21-ester (8,9). The doses chosen for local therapy were similar to those used in systemic therapy (for hydrocortisone 100 mg and for prednisolone 20 to 40 mg). Therefore, it is not astonishing that it soon became apparent that this therapy was not just local, but − verified by depressed plasma cortisol levels − provoked marked systemic actions (10-14). Somewhat better selectivity for the rectal and sigmoid mucosa may be obtained by using prednisolone metasulphobenzoate, which is a larger molecule with a probably slower release of prednisolone (15). However, the site of that release is not yet known with certainty (15) and the conjugate may induce sulpho-like side effects (3).

Today, it can be judged that if the medical aim is local therapy of bowel or rectal mucosa, hydrocortisone or prednisolone (or their esters) is not the right choice, as both hormones have too high systemic availability.

The aim of this paper is to describe the novel potential for a more selective topical therapy due to the forthcoming introduction of new glucocorticosteroids with high first pass metabolism in the liver. This drug development has not been primarily directed to inflammatory bowel disease (IBD) but to an improved topical (inhalation) therapy of asthma and rhinitis. The main steps are schematically outlined in Figure 1. The first step was an increment of oral
glucocorticosteroid potency of the prednisolone type of structure leading to potent but very biostable compounds like betamethasone, dexamethasone and triamcinolone. It was then possible to enhance the poor topical anti-inflammatory potency of these compounds by the introduction of lipophilic constituents in the 17-alpha and/or 16-alpha,17-alpha positions, resulting in the first topically potent 'skin' steroids betamethasone 17-valerate and triamcinolone acetonide. During that development the vasoconstriction (blanching) test on human skin was evolved as a rapid and relevant test in the judgement of topical glucocorticosteroid potency in man (16). Little attention was paid to the biotransformation rates and routes of these lipophilic glucocorticosteroids due to their restricted percutaneous absorption through the stratum corneum barrier. However, when betamethasone 17-alpha-valerate and the derivative beclomethasone 17-alpha, 21-dipropionate were inhaled for the topical treatment of asthma and rhinitis, the systemic glucocorticosteroid side effects were found to be surprisingly minute (17). This results from the fact that the liver, via its drug metabolizing enzymes (including the cytochrome P450 system), can more easily inactivate this lipophilic type of glucocorticosteroid structure than the less lipophilic structures of the parent molecules betamethasone, dexamethasone and triamcinolone (17,18).

For glucocorticosteroid esters a clear distinction must be made between 17-alpha and 21-esters. Esterification in the 17-alpha position enhances glucocorticosteroid potency markedly (16), while 21-esters have unaltered or even reduced activity. Furthermore, the 17-alpha but not the 21-ester bond is rather stable in extrahepatic tissue, eg, airways and intestinal walls (18). This suggests that at the site of local application, 17-alpha-esters can exert strong glucocorticosteroid activity, which on the other hand can be reduced in the systemic compartment via hepatic biotransformation. This improved profile of 17-alpha over 21-esters has been demonstrated also for the rectal mucosa. Enemas containing betamethasone 21-phosphate provoked systemic actions even at the minimum dose for therapeutic efficacy (19). On the other hand, enemas containing betamethasone 17-alpha-valerate (20) or beclomethasone 17-alpha,21-dipropionate (21-24) were reported to induce less plasma cortisol depression than that obtained with therapeutically equipotent prednisolone or betamethasone 21-phosphate enemas.

As shown in Figure 1, the developmental line that started with betamethasone valerate and beclomethasone dipropionate has now been reinforced with new topical glucocorticosteroids - eg, budesonide (25,26) and fluticasone propionate (27,28). The latter two glucocorticosteroids have from the beginning been designed and selected for a high first pass metabolism in the liver, and their properties are described in more detail below.

**SELECTION OF OPTIMAL GLUCOCORTICOSTEROID PROFILE FOR TOPICAL THERAPY OF BOWEL AND RECTUM**

The general opinion is that most specific glucocorticosteroid actions are mediated via triggering of glucocorticoid receptors. When a glucocorticoid agonist is bound by the receptor, there is a conformational change in the steroid-receptor complex and a translocation of the complex to glucocorticoid-responsive elements in the genome. By modulation at these sites there is enhanced or reduced transcription of mRNA for glucocorticoid-steroid sensitive proteins (29). The transcription and translation of some peptides or proteins are strongly suppressed by glucocorticosteroid (eg, most interleukins) while other proteins are reinforced (eg, beta-receptors, surfactant, angiotensin converting enzyme, growth hormone and metallothionin). The varying combination of enhanced and reduced transcription leads to the wide spectrum of glucocorticosteroid actions, eg, anti-inflammatory actions, immunosuppression, local or systemic catabolic actions, etc. The glucocorticosteroid receptor seems to be uniform in the human body (30), which means that the same receptor type - depending on its cellular location - can mediate therapeutic as well as adverse glucocorticosteroid reactions. Thus, there is currently no possibility of separating the various glucocorticosteroid actions via subtypes of glucocorticosteroid receptors. However, for organs accessible to local therapy, a better way to achieve separation is to deposit a restrictive quantity of selected glucocorticosteroids directly in the organ, and thus reduce the adverse effects originating from bulk glucocorticosteroids distributed from a high systemic bolus. One recent example is glucocorticosteroid inhalation for the topical treatment of asthma and rhinitis (17,26), which has more or less revolutionized the therapy of moderate and severe asthma. No corresponding clinical development has occurred for the topical treatment of IBD, which - at least to some extent - may depend on the wrong glucocorticosteroid having been exploited. In theory, with the right glucocorticosteroid the prospects are better for the development of a selective topical therapy for the bowel than for the airways. This is due to the fact that these glucocorticosteroids are inactivated by biotransformation mainly in the liver (18,31,32). When absorbed from the bowel compartment, nearly all of the drug has to pass through the liver and its specialized drug metabolizing systems before being distributed in the systemic circulation. However, when the inhaled drug is absorbed from the airway mucosa it reaches the heart first and only approximately one-third will have first pass access to the liver.

Thus in theory, the optimal glucocorticosteroid profile for topical anti-inflammatory immunosuppressive therapy of IBD is a compound with high topical glucocorticosteroid potency and with rather high metabolic stability in the bowel and rectum compartments - the latter property to secure full activity in the target organ. On the other hand, the compound should have 100% first pass metabolism in the liver.

The new glucocorticosteroids will
be judged according to these criteria. Topical potency is estimated by two parameters. One is the relative affinity for glucocorticosteroid receptors, determined in subcellular systems with little possibility of metabolic inactivation. As determined in rat models, in a series of eight glucocorticosteroids not inactivated in the target organ, there is a close correlation between receptor affinity and topical anti-inflammatory potency (Spearman’s rank order correlation coefficient 0.98) (Figure 2). The other estimation of topical glucocorticosteroid potency is activity in the vasoconstriction test (16). In this test, ethanolic solutions of the glucocorticosteroid are applied to human skin under occlusion to facilitate percutaneous absorption.

What level of glucocorticosteroid potency is optimally required for the topical treatment of IBD? The efficacy of enemas with hydrocortisone (4-7) and prednisolone (8,9,13) suggests that low glucocorticosteroid potency is sufficient to alleviate the symptoms of proctitis and left-sided colitis. However, little is as yet known regarding to what extent intense glucocorticosteroid treatment by more potent enema preparations will induce a more complete and long lasting healing of lesions. The study by Danielsson et al (13) illustrates that it is possible to reach better symptomatic and objec-

tive healing with a potent preparation (budesonide) than with prednisolone.

Furthermore, the most exciting medical aim for improved topical IBD therapy is to reach topical (local) selectivity for the ileal and colonic bowel walls via retarded oral formulations. Ideally, such a formulation should release the active glucocorticosteroid just within the bowel segments most often affected by Crohn’s disease and ulcerative colitis. It is still poorly known how deep the submucosal penetration must be for optimal ‘topical’ therapy. In Crohn’s disease it has been proposed that the tissue infarction starts with vascular injury and arterial occlusion at the level of the muscularis propria (33), which would suggest that high anti-inflammatory activity is desired also in the deep compartments of the bowel wall. A retarded oral formulation for the selective treatment of the bowel should therefore release a potent glucocorticosteroid with sufficient water and lipid solubility to allow its dissolution as well as its effective absorption deep into the bowel wall. Ideally, the glucocorticosteroids should have a high affinity for the bowel tissue and thus stay long in that compartment. Another reason for selecting highly potent glucocorticosteroids for retarded oral formulations is that, due to the risk of present or forthcoming luminal strictures in Crohn’s disease, the tablets should be small.

However, to achieve improved topical therapy of IBD, the main medical aim is to select a glucocorticosteroid with a much higher first pass metabolism than that of prednisolone (Figure 1). A high first pass metabolism in the intestinal-liver compartment is reflected as a low systemic availability via the rectal or oral routes. Therefore, in the following discussion, oral or rectal bioavailability is used as a parameter of unwanted systemic activity.

PROFILE OF NEW GLUCOCORTICOSTEROIDS

Table 1 describes the structures and topical potencies of the new glucocorticosteroids. The topical potency (receptor affinity and vasoconstriction activity) is given in relation to

![Figure 2](image-url)  
Close correlation between affinity for glucocorticosteroid receptor (in vitro) and topical antiedema potency (in vivo). Based on studies in rats (52,53)

![Figure 3](image-url)  
Oral bioavailability (%) in humans. Based on Table 7 in reference 55, which summarizes the human studies published (hydrocortisone one study, prednisolone seven, methylprednisolone two, dexamethasone four, budesonide two)
TABLE 1
Newer glucocorticosteroid substances for inflammatory bowel disease

<table>
<thead>
<tr>
<th>Substitution position</th>
<th>D-ring</th>
<th>Name</th>
<th>Relative potencies</th>
<th>Receptor affinity</th>
<th>Topical vasoconstriction</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sat H H</td>
<td></td>
<td>Hydrocortisone</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Sat H H</td>
<td></td>
<td>Tixocortol pivalate</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unsat H H</td>
<td></td>
<td>Prednisolone</td>
<td>13</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Unsat Cl H</td>
<td></td>
<td>Beclomethasone dipropionate</td>
<td>80</td>
<td>600</td>
<td></td>
</tr>
<tr>
<td>Unsat H H</td>
<td></td>
<td>Budesonide</td>
<td>-195</td>
<td>-1000</td>
<td></td>
</tr>
<tr>
<td>Unsat F F</td>
<td></td>
<td>Fluticasone propionate</td>
<td>-</td>
<td>-1200</td>
<td></td>
</tr>
</tbody>
</table>

Structure and topical glucocorticosteroid potencies of glucocorticosteroids of current interest in the topical treatment of inflammatory bowel disease. The topical glucocorticosteroid potency is given as affinity for the glucocorticosteroid receptor and as topical vasoconstriction activity - both in relation to hydrocortisone. Sat Saturated bond in the 1-2 position; Unsat Unsaturated (double) bond in the 1-2 position. Based on the following references: tixocortol pivalate (34), prednisolone (in the files of Draco and 54), beclomethasone dipropionate (17,42,48,54), budesonide (42,54), fluticasone propionate (48).

hydrocortisone. Hydrocortisone has a low topical potency (Table 1) and its oral bioavailability is approximately 50% (Figure 3). The latter figure means that half of orally (and probably rectally) administered hydrocortisone passes intact into the systemic circulation. Prednisolone has a somewhat higher receptor affinity (approximately 10 times that of hydrocortisone) (Table 1) but also a clearly higher oral bioavailability (approximately 80%) than that of hydrocortisone (Figure 3). This means that with prednisolone the prospects of a selective therapy of the bowel are even more lacking.

Tixocortol pivalate is a late development from the hydrocortisone structural line. By changing the 17-beta side chain of the D-ring from –COCH2OH to –COCH2SH and esterifying the thiol group with pivalinic acid, the thioester tixocortol pivalate was obtained (Table 1). This change has not enhanced receptor affinity over that of hydrocortisone (34), and as an enema preparation even higher doses (250 mg) than those in the latter reference are used (35-37). The interesting property of the substance is its high first pass metabolism, leading to a bioavailability of 10 to 20%. In man the acute rectal administration of 500 to 2000 mg tixocortol pivalate did not markedly affect plasma or urinary cortisol levels (39). It should be added that these doses were given in the morning, when the body is less sensitive to external glucocorticosteroids than compared to the late evening. However, under the experimental conditions used, a 5 mg betamethasone 21-phosphate enema strongly reduced cortisol levels (39). By saturation and reduction of the A-ring and rearrangement of the thiol bond, tixocortol pivalate is biotransformed into metabolites with low glucocorticosteroid activities (38,40). Animal studies demonstrate that the plasma clearance of tixocortol pivalate is very high, demonstrating an extensive extrahepatic metabolism (38). This is supported by in vitro studies showing that the erythrocyte enzyme S-methyltransferase can rearrange the thiol bond of tixocortol pivalate (38). This rapid extrahepatic metabolism may result in partial inactivation in the target organ, which might lead to even lower potency in the bowel wall than the low figure given in Table 1. Restricted intestinal absorption may also contribute to low therapeutic activity, since 2 to 42% of the dose given was found unchanged in human feces (38). Thus, in theory tixocortol pivalate has an improved profile for the selective glucocorticosteroid treatment of proctitis and left-sided colitis, but more studies are required to judge its therapeutic potency on that indication. Its very low topical potency excludes its use via rectal formulations.

The 17-alpha substituted glucocorticosteroids, beclomethasone dipropionate, budesonide and fluticasone propionate (Table 1) all have intrinsic
activities at least 100 times greater than those of hydrocortisone and tixocortol pivalate, and all three are inactivated by liver biotransformation alone. Beclomethasone dipropionate enemas are reported to be active in proctosigmoiditis at doses of 2 to 5 mg with little systemic activity (14,21-24). Beclomethasone dipropionate and its first hydrolysis product - the corresponding 17-alpha monopropionate with an unusual fluoromethyl carbothioate group into metabolites with low glucocorticosteroid potencies (48,49). After intravenous injection into humans its plasma clearance is approximately 0.9 L/min (48). The oral bioavailability is proposed to be very low, depending on hepatic first pass metabolism, but also on low (less than 50%) absorption from the intestines leading to marked excretion of intact substance. The partial intestinal absorption may be related to low water solubility, impairing dissolution and transport into the mucosal surface.

CONCLUSIONS

Figure 4 summarizes the prospects for more selective topical IBD therapy by new glucocorticosteroids. The figure is exemplified by budesonide, which is currently the best documented compound. Via the rectal or oral route it is possible to deliver to the bowel mucosa a glucocorticosteroid with a topical anti-inflammatory potency approximately 200 times that of hydrocortisone. Based on the T_max in plasma the dwell time in the bowel compartment is estimated to be one to a few hours, which is a sufficiently long period for the triggering of important anti-inflammatory actions (50,51). When passing through the liver, there is a 90% first pass metabolism delivering only about 10% of intact glucocorticosteroid to the systemic circulation.

Enhanced drug selectivity for the bowel compartment can be exploited therapeutically in two ways. The reduced risk of serious adverse reactions outside the target area makes protracted therapeutic or even prophylactic treatment more possible. En-
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