Gender and the Pathogenesis of Gastrointestinal Diseases: The Role of Steroid Sex Hormones in the Development

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Abstract
The pathogenesis of gastrointestinal diseases are unclear. The strategy of any medical regimen for management of any disease is usually dependent on our understanding of the mechanism by which this disease has developed.

Different incidences of different gastrointestinal diseases in women from those in men suggest possible role of steroid sex hormones in the promotion or prevention of the development of these diseases. The present review is aimed at finding any clue that could help in improving our understanding and the outcome of management accordingly.

Conclusion: Steroid sex hormones play crucial role in the pathogenesis of several gastrointestinal diseases. Based on our findings in this survey, further studies are needed in order to determine how we could operate these findings in our clinical practice.

Keywords: Gastrointestinal diseases; Inflammatory bowel diseases; Functional bowel diseases; Fistula-in –ano; Idiopathic constipation; Peptic ulcer; Pathogenesis; Gender; Sex hormones; Epidemiology; Incidences

Introduction
Gastrointestinal diseases are recognized, as being dominant in women (Irritable bowel diseases, chronic constipation), but others, as being more common in men (Anal fistulae). These observations raise concerns about the involvement of sex steroid hormones in the pathogeneses of these diseases.

In order to examine this probability, we conducted a survey on a number of main common gastrointestinal diseases.

The outcome of this investigation was as followings:

Peptic ulcers and sex hormones
There is, sometimes, no clear evidence on potential role of sex hormones in inducing or modulating pathogenesis of gastrointestinal disorder. A hypothesis that they may act throughout potentiating the effect of other co-factors is possible. On the basis of experimental and clinical observations, it is known that ulcers of the gastroduodenal mucosa develop in a sexually dependent manner. For example, in the fertile age, peptic ulcer disease occurs more frequently among men than in women [1-3]. This sex- dependent differences may also be found in experimental models of mucosal damage, as oral administration of ethanol generated more severe gastric erosions in male rats than in females [4,5]. In this model of mucosal damage, gonadectomy protected the stomach against ethanol- induced injury only in male rats, but not in females [4].

Moreover, the administration of the testosterone receptor blockers, cyproterone acetate attenuated gastric haemorrhagic erosions in intact male rats following ethanol challenge.

In contrast, in a different model of gastroduodenal ulcers (as induced by restraint and forced exercise) the stomach of male rats was less sensitive to mucosal damage compared to that of females [6].

In conjunction to these findings, it can be suspected that sex hormones might possibly play a role in the development of gastroduodenal mucosal ulcer.

The actual sexual hormonal status of females may also be related to the generation of gastroduodenal ulcers. Pregnancy and lactation in rats markedly reduced steroid- and cysteamine-induced gastroduodenal lesions [7,8]. A lower incidence of peptic ulcers is also present in pregnant women [3]. Moreover, during pregnancy, the frequency of haemorrhage and/or perforation from gastroduodenal ulcers is lower compared to its high incidence in peripuerium. Thus the elevation of endogenous progesterone plasma levels may be responsible for pregnancy -induced protection against mucosal ulcers, since early pregnant rats (with increased plasma oestrogen level) were less sensitive to gastroduodenal ulcers [8].

In addition, exogenous administration of 17 B estradiol into intact female rats increased mucosal damage induced by cysteamine [9].

Bowel motility and sex steroid hormones
The frequency of bowel movements in young women per week during the first half of the menstrual cycle where estrogen predominates was found almost half of those in males [10]. Yet, the colonic transit time was more prolonged during the second half where estrogen and progesterone predominated. Also, less often incidents of defecation per week in pre-menopausal than in post-menopausal women and in the last group than in men suggest some role of feminizing hormones in the pathogenesis of constipation.

Evidence exists from a few numbers of small studies that changes
in gastrointestinal function occur in a cyclic pattern in menstruating women and may be related to cyclic variations in the ovarian hormones [11-15]. Even some constipated patients developed diarrhea just before menstruating. [12] Simmons et al reported prolonged gastrointestinal transit time during the luteal phase of the menstrual cycle where estrogen and progesterone predominate, when they compared their measurements with those recorded during the follicular phase where estrogen only predominates [16]. They also reported that the colonic motility in five groups of rats they found higher duration of maximum electric activity in pregnant rats when compared with those in controls, ovariecetomized, ovariecetomized and treated with estrogen, and ovariecetomized and treated with progesterone.

In another study, Wald and others [20] found that intestinal transit time as was measured by breath hydrogen level after lactulose ingestion, was longer on Days 18-20 (mid luteal) than on Days 8-10 (mid follicular). These findings were further confirmed by Heathkemper's report in 1992 [21]. It was a small study aimed at assessing the intestinal motility in 34 young females aged between 19-37 years, of who nine were on contraceptive pills, and they were followed for two successive menstrual cycles by Heathkemper's team. Heathkemper et al found differences in bowel patterns between midfollicular and luteal phases except in the subgroup that were on oral contraceptive pills. They also found an increase in stool consistency scores at menses when estrogen and progesterone levels were low or decreasing [21].

Stool pattern in women on contraceptive pills, whose ovarian hormones levels supposedly stable, remain relatively stable across the two phases of the menstrual cycle. They showed the greatest cycle-related changes in stool consistency scores at menses. These findings suggest that important factor that influences stool pattern may be the amplitude of change (increase/decrease) in ovarian hormones levels, which occurs in normal menstrual cycle and in those on contraceptive pills.

A possible correlation between circulating levels of motilin and progesterone in normal pregnancy and early postpartum was investigated in a Chinese study [22]. Qiu XH reported that progesterone levels increased along with gestational age whereas motilin levels decreased. A significant negative correlation was found between the two hormones. He suggested that progesterone appeared to have a profound inhibitory effect on motilin. The decreased motilin level might partially be responsible for gastrointestinal hypomotility during pregnancy. These Chinese results were confirmed further by others' findings [23].

**Anal fistula development and gender**

The discrepancy in the incidence of non-specific fistula-in-ano between the adult males and females is well acknowledged. Sainio [24] in his population study over ten year period reported an incidence of 5.6/100 000 for females and 12.3/100 000 for males, with a male: female ratio of 1.8:1. Other quotes an even higher ratio, reaching 9:1 in some series [25]. This discrepancy suggests some protective role of female sex hormones against the development of idiopathic peri-anal fistula in women.

Acute infection of anal glands often results in the formation of a chronic intrasphincteric collection that may drain down to the perineum, up into the supralveolar compartment or across the sphincter into the ischiorectal fossa [26-28]. Abscess complicating an infection in an anal gland are caused by intestinal organisms and are variably associated with an internal opening at the outset. Drainage consequently often results in the creation of a track or fistula between the skin and anal mucosa at the dentate line. The conducted prospective study by Eykyn & Grace in 1986 [29] evidently provides the most informative data on the causative agents in these cases. The participant patients had an initial examination under anaesthesia and a further examination, one week later, by an experienced surgeon to inspect the presence of a fistula. This clinical assessment was harmonized by an enormously inclusive microbiological survey. The significant microbiological findings are summarized in Table 1. This table provides an objective evidence to support the idea that gram negative bacteria are mainly responsible for the development of fistulae in those with anal abscess.

But whether sex hormonal receptors are exist on the surface of the anal tissue or not. This was answered by Oettling G and Franz HB [30]. In 1998, Oettling G and Franz HB [30] investigated the expression of androgen, oestrogen and progesterone receptors (ARs, ERs, PRs) in the tissues of the anal continence organ in 23 patients (seven men, seven premenopausal women and nine postmenopausal women) using immunohistochemical techniques. In their report, they concluded that specific immunostaining for ARs, ERs and PRs were found over cell nuclei. ARs were found in the smooth muscle cells of the internal anal sphincter in all but one of the females (10/11) and all males (7/7), ERs were found in 12/12 females and 4/7 males, and PRs were found in 4/10 females and 1/7 males. The squamous epithelium exhibited a similar pattern of immunostaining.

Two oestrogen receptor (ER) subtypes, named ERα and ERβ have been isolated and copied [31-33]. Vascularsmooth muscle and endothelial cells have been reported to express ERα and Erβ protein & mRNA [34-36]. Oestrogen has been proved to have anti-inflammatory effect by acting against inflammatory promoters, such as bacterial cell wall component, lipopolysaccharide (LPS) via activating ERα [37-42].

For investigating a probability that LPS (inflammatory promoter) could induce attenuation of ER expression by endothelial cells and this act could possibly be antagonized by oestrogen, a laboratory study was conducted by Holm and others [43]. They found that treatment with LPS (10µg/ml) for 4 days attenuated iNOS protein expression by about 60%, and reduced ERα mRNA expression by about 50% and ERβ mRNA expression by about 60%. But stimulation with LPS (10µg/ml) for shorter time (16 hours) had no effect.

**Table 1: Type of organisms isolated from anorectal abscesses with and without fistulae.**

| Type of organism | Fistula present (n=53) | No fistula (n=27) | P  |
|------------------|------------------------|------------------|----|
| Number of abscesses yielding anaerobes | 49 (92.5%) | 8 (29.6%) | <0.0001 |
| Escherichia coli | 45 (84.9%) | 5 (18.5%) | <0.0001 |
| Staph. aureus | 1 (1.9%) | 8 (29.6%) | P=0.0012 |
| Anaerobes 'Gut-specific Bacteroids' | 47 (88.7%) | 5 (18.5%) | <0.0001 |
| Anaerobes not 'gut-specific' (only) | 2 (3.8%) | 17 (63%) | P=0.0001 |
| Gut aerobes* 'gut-specific anaerobes' | 45 (85%) | 4 (14.8%) | <0.0001 |

Includes E. coli, Klebsiella spp, Proteus spp, Citrobacter spp, Salmonella sp, Str. faecalis

* Plus 2 colonies isolated.
Holm and colleagues also found that dexamethasone (1uM) had no effect on the induced down regulation of ERα mRNA expression by LPS. This means that NF-B signalling pathway is not involved in LPS induced attenuation of ER gene expression (dexamethasone is known as an inflammatory inhibitor by acting against NF-B dependent transcription of pro-inflammatory genes) [44]. They, also, found that oestrogen increased mRNA expression in the presence of LPS. This may explain the lower incidences of anal fistulae in females than in males and raises a probability that oestrogen may be a promising effective therapy for these cases.

**Inflammatory bowel diseases & gender**

In general there are no striking differences in the sex incidence of colitis or Crohn's disease. Variation of 1.5:1 [45-53] for excess of men or women, with either disease, with no general consistency from one country, or study group, to another. Given environmental association with smoking or probably with oral contraceptive use, it is difficult to believe that examination of the sex rates of the two diseases will yield significant clues to causes of disease frequency [54].

**Fissure in ano and gender**

There is often no apparent disorder which predisposes adult patients to anal fissure. Childbirth seems to be an associated factor in some women [55-57]. Martin in 1953 [57] reported that acute anal fissure occurred in association with recent childbirth in 11 % of patients. Gough women [55-57]. Martin in 1953 [57] reported that acute anal fissure to anal fissure. Childbirth seems to be an associated factor in some

**Implications in research and clinical practice**

How to operate the findings in this survey in our current medical regimen necessitates further studies.

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