On the pathogenesis of the irritable bowel syndrome: The irritable bowel or the irritable patient?

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ABSTRACT: The traditional perspective of irritable bowel syndrome (IBS) as a behavioural problem has tended to downplay the role of gastrointestinal dysfunction. Contrary to predictions based on the traditional philosophy, a recent study has shown that IBS patients have increased pain tolerance compared to healthy subjects. This profile of pain tolerance is similar to that seen in chronic organic disease of the gut (eg, Crohn's disease), raising the possibility that IBS patients may experience pain resulting from gastrointestinal dysfunction. The recent finding of increased airway responsiveness to inhaled methacholine in certain IBS patients provides an objective and quantifiable measurement of tissue dysfunction in that syndrome, and focuses attention on possible mechanisms underlying the altered responsiveness of hollow organs in patients with IBS; these mechanisms are discussed. Can J Gastroenterol 1990;4(1):33-38

Key Words: Asthma, Behaviour, Inflammation, Motility, Smooth muscle, Stress
La pathogenèse du syndrome du côlon irritable : Côlon ou patient irritable?

RESUME: A cause de la perspective traditionnelle qui consiste à voir le syndrome du côlon irritable (IBS-SGI) comme un problème de comportement avant tout, on a eu tendance à accorder une importance moindre à la compréhension des causes du dysfonctionnement gastroentestinal dans cette affection. Or, contrairement aux prévisions conformes à cette philosophie, une étude récente démontre que les patients souffrant de SGI manifestent une tolérance accrue à la douleur, comparés aux sujets en bonne santé. Ce profil de tolérance à la douleur est semblable à celui que l'on observe dans le cas d'autres maladies organiques chroniques de l'intestin (maladie de Crohn, par exemple); il serait donc possible que les patients atteints de SGI souffrent bien de douleurs résultant d'une dysfonction gastroentérale. Des résultats récents montrent, chez certains de ces patients, une faculté de réponse accrue des voies aériennes aux inhalations de méthacholine — fournissant une mesure objective et quantifiable de la dysfonction des tissus dans ce syndrome et attirant l'attention sur les mécanismes qui sont peut-être à l'œuvre dans la faculté de réponse altérée des organes creux chez ces patients; ces mécanismes sont examinés.

Inappropriate study design
Incomplete understanding of basis for gastrointestinal symptoms
Limited understanding of drug action and the pharmacology of gut motility
Actual absence of efficacy

Another reason for the failure to demonstrate drug efficacy against gastrointestinal symptoms in IBS is the limited understanding of the mechanisms underlying these symptoms. Although there are considerable data illustrating motility disturbances throughout the gut in IBS patients, the relationship between symptoms and abnormal motor patterns is incompletely understood. This is perhaps most pronounced in relation to abdominal pain; it is likely better understood in the case of constipation and diarrhea. A further problem rests with the complexities of the pharmacology of gastrointestinal motility (14). These issues are serious ones in that they may not only mislead one into believing that potentially useful drugs have no place in the treatment of IBS, but may also prompt one to dismiss the notion that gastrointestinal dysfunction exists as a basis for symptoms in IBS.

A different argument that mitigates in favour of a primary behavioural abnormality in IBS is the demonstration that symptoms similar to those reported primarily behavioural etiology. However, there exist several other reasons why a drug might exhibit poorly demonstrable efficacy against these symptoms (Table 2). The first and most obvious reason is that traditional methods for evaluating drug efficacy are inappropriate for a condition that is heterogeneous not only in its clinical presentation, but most likely in its pathogenesis. Methods for evaluating drug efficacy require patient homogeneity, a relatively low placebo effect and a predictable and stable course of the disease over time. They do not take into account the considerable heterogeneity of the patient population, the very high placebo rate, or the spontaneous relapses and remissions that characterize IBS. This subject has recently been reviewed in depth (13).
in IBS are experienced, but not reported, by as much as 14% of the population (15,16). This finding, taken in conjunction with a report of a greater prevalence of learned illness behaviour in IBS patients compared to those with peptic ulcer disease (17), suggests that IBS patients are intolerant of ‘normal’ gastrointestinal sensations and use this experience to seek attention.

Since pain is the symptom that has been shown to be the most common reason for IBS patients to seek attention, one might surmise that such patients select themselves from the general population by virtue of poor pain tolerance. If one assumes, for a moment, that gut function is normal in IBS, then previous reports of intolerance to balloon distension of the rectum in IBS patients support this notion (18). However, the results of a recent study seriously weaken this argument.

Pain perception and reporting were examined in IBS patients using electrocutaneous stimulation over the dorsum of the hand (Figure 1) (19). Results were compared to those obtained in healthy controls and in a group of patients with chronic abdominal pain due to Crohn’s disease. Both IBS and Crohn’s disease patients had significantly (P = 0.016) higher pain thresholds than normals, and these thresholds were similar in the IBS and Crohn’s disease groups. In addition, touch thresholds were higher in these groups compared to the controls, and were significantly higher in the IBS patients than in those with Crohn’s disease. These results indicate that IBS patients are less sensitive to low intensity nonpainful stimuli (touch) and have a higher threshold for painful stimuli than normal subjects. The results suggest that IBS patients do not select themselves from the general population by virtue of a generalized reduction in pain tolerance. It follows that reports of pain by IBS patients should not be disregarded as a manifestation of learned illness behaviour.

Since the ability to tolerate higher levels of pain is usually associated with painful chronic conditions (20-22) and can be induced in animals subjected to pain (23), the results suggest that IBS patients experience chronic pain on the basis of gastrointestinal dysfunction.

IRRITABLE BOWEL?

There is an emerging literature which demonstrates that sensory perception within the gut is altered in IBS patients, suggesting that the bowel may be truly ‘irritable’. Although the emphasis has been placed on an altered motor function in IBS, it is likely that the exaggerated motor responses observed in IBS may reflect, at least in part, altered sensory input.

Earlier studies reporting intolerance to balloon distension of the rectum in IBS patients (17) support the notion that there is altered sensory input from the gut, particularly if one accepts that these patients are able to tolerate larger amounts of pain than normal subjects (19). The demonstration of abnormal vagal activity in IBS patients (24) may also reflect this, particularly as the majority of the vagal fibres are afferent. Recently reported studies provide further evidence of ‘irritability’ in the gastrointestinal tract of IBS patients.

One study extended previous work by showing that intolerance to balloon distension is evident in the stomach of patients with idiopathic functional dyspepsia, which may represent a subgroup of the IBS population (25). Another study examined the ability of various luminal stimuli to induce pain and/or colonic motor responses in IBS patients and controls. The luminal stimuli consisted of balloon distension, infusion of 15 mM deoxycholic acid (DCA) and of a mixture of short chain fatty acids (SCFA) (70 mM acetic acid and 50 mM lactic acid) delivered in random order and interspersed with saline infusion (26). DCA infusion reproduced the familiar pain in seven of eight IBS patients but in only one of five controls. SCFA infusion produced similar responses in five of seven IBS patients but in none of the five controls. The threshold for pain induced by balloon distension of the distal colon was lower in IBS patients compared to the control threshold (75.7 versus 171 mL,
P<0.001). Although the motor responses to the acid infusions were larger in IBS patients compared to the controls, the differences were not significant. These results suggest that there is an increased sensitivity of afferent nerves responding to mechanical or chemical stimulation in the gut of IBS patients, which contrasts sharply with the demonstration of increased pain thresholds outside the gut in IBS patients. The implication is that the gastrointestinal tract is indeed abnormal in IBS patients.

**GASTROINTESTINAL DYSFUNCTION IN IBS**

**Altered motor function:** Previous discussions regarding the nature of gastrointestinal dysfunction in IBS have tended to focus on motor abnormalities, and there has even been speculation on the existence of a primary disorder of smooth muscle in the gut in IBS (27). This was prompted in part by demonstrations of altered myoelectric activity recorded in vivo from the unstimulated colon of IBS patients, and in particular the suggestion that there may be an altered slow wave frequency with a higher incidence of 3 cycles/min activity in IBS (28-30). Since slow waves are generated by oscillations in membrane potential, changes in slow wave frequency may be regarded as manifestations of a fundamental alteration in smooth muscle cell (31). However, this finding has not proven to be robust and several other workers have failed to demonstrate it (32-33).

This has as much to do with the difficulties in obtaining and processing electrical signals from the human colon (34) as it does with the inherent complexity and variability of the electrophysiologic control of colonic motility (35). The question of whether there is a primary abnormality in smooth muscle cell function in IBS remains open, but it is this author's opinion that if such an abnormality exists, it does so only in a subpopulation of patients; as already mentioned, IBS is likely to reflect more than one pathogenetic process.

In spite of the uncertainty regarding a primary role for smooth muscle dysfunction in IBS, there is little doubt that stimulated motor activity is abnormal in IBS. Several studies have shown that motor responses to nutrients (36,37), to drugs (38), to hormones (39) and to bile acids (40) are exaggerated in IBS patients versus controls. Since some of these stimuli were delivered via the gut lumen, the exaggerated motor response may reflect, in part, an altered sensory input. However, in certain instances, stimuli such as cholecystokinin and parasympathomimetic drugs were delivered parenterally and would be presumed to act on the motor apparatus directly. The extent to which these responses reflect changes in enteric (afferent) nerves or smooth muscle cells remains to be determined.

**Altered epithelial function:** Altered gastrointestinal function in IBS is not restricted to the neuromuscular tissues. A study from Denmark has shown that the intestinal epithelium from patients with diarrhea-predominant IBS exhibits net secretory characteristics compared to controls when challenged with bile acids (41). In addition, there is a study in which patients with IBS and a specific food intolerance produced more prostaglandin E₂ in the gut lumen when challenged in a double blind manner (42). Although the source of the prostaglandins could not be identified, it was most likely produced by the epithelium and is unlikely to have originated in the deep muscular layers of the gut wall.

**IRRITABLE BODY?**

The prevalence of extragastrointestinal symptoms in IBS patients (43,44) has prompted the investigation of dysfunction in hollow organs outside the gut in IBS patients. Abnormal urodynamics have been recorded in patients with constipation due to colonic inertia (45) and in other IBS patients (46). Others have reported significantly lower blood pressures in a group of IBS patients compared to controls (47-49). Some of these data have been interpreted as providing evidence of altered smooth muscle function (46,47,49). However, vascular and urinary bladder responses may reflect hormonal and neural as well as muscular factors, and data obtained from such studies are difficult to quantitate accurately.

To pursue this issue further, and to overcome some of the obstacles found in previous studies, airway responsiveness was recently examined in IBS patients. In humans, the measurement of the volume of air expired in 1 s under maximal effort following methacholine or histamine inhalation (FEV₁) is a reliable method of assessing bronchial airway calibre and reactivity (50-52). The response to inhaled methacholine is generally considered to reflect an interaction with muscarinic receptors on airway smooth muscle (53). IBS patients were found to be significantly more sensitive to the bronchoconstricting effects of methacholine than a group of healthy subjects or those with organic diseases of the gastrointestinal tract (54). Significantly larger changes in FEV₁ were recorded in IBS patients and occurred following the administration of significantly smaller amounts of methacholine than was required in other groups. The changes in FEV₁ were not, however, of the magnitude observed in asthmatic patients, and none of the IBS patients studied were atopic.

What are the implications of these results? First, they provide bona fide evidence of altered smooth muscle function outside the gut in IBS patients. The alteration in smooth muscle function may be primary secondary to altered innervation of the muscle or, as in asthma, the consequence of an inflammatory process in the airways. Second, these results raise the spectre of a convenient screening test for IBS patients, although it must be emphasized that the results were obtained in a highly selected IBS group and it is not known whether the findings can be extrapolated to the IBS population at large. Finally, the results prompt comparisons between IBS and asthma.

**IBS AS 'THE ASTHMA OF THE GUT'?**

Could IBS, or subgroups of IBS, patients reflect the same spectrum of pathophysiological processes that are believed to produce asthma? Why not? There are broad similarities between the digestive and respiratory tracts, and several striking similarities between the two conditions (Table 3).

Let us recall that asthma was considered initially to be a largely psychogenic
TABLE 3

Similarities between asthma and IBS

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<thead>
<tr>
<th>Asthma</th>
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<tr>
<td>Yes</td>
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<td>(53)</td>
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<td>Yes*</td>
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*Primarily in childhood asthma, but originally in all asthma. **In the context of post-infectious IBS (49) and the demonstration of increased prostaglandin production in IBS patients with specific food intolerance.

Asthma

- Hyperresponsiveness on pharmacologic stimulation
- Psychological factors implicated in the pathogenesis
- Mast cells implicated
- Inflammation implicated

IBS

- Altered smooth muscle function
- Inflammation implicated
- Interference with immune function
- Psychological factors implicated

Disorder in which pulmonary function, but not structure, was altered (55-57), much along the same lines as IBS is now traditionally considered (3,4,5,17). However, the perception of asthma changed considerably as research uncovered the mechanisms underlying the increased responsiveness of bronchial smooth muscle (53). An understanding of these mechanisms has in turn led to a more rational and efficacious pharmacotherapy for asthma. The lesson to be learned is that one must persist in investigating this common and often frustrating clinical problem. To dismiss it as a psychogenic disorder is to overlook an increasing body of evidence which points to altered function in tissues inside and outside the gut. We must continue to investigate the mechanisms whereby the gut becomes ‘irritable’ or ‘hyperresponsive’, bearing in mind that these mechanisms may have their origins in the brain, the gut and the environment.

REFERENCES

33. Bueno I, Fioramonti J, Ruckebusch Y, Frexinos J, Coulom P. Evaluation of...


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