Probiotics in inflammatory bowel disease: is it all gut flora modulation?

S Ghosh, D van Heel, R J Playford

Understanding probiotic action may permit modulation of the immune system, both locally and systemically

There is considerable public, media, and scientific interest in “natural” products, including probiotics, in modulating intestinal inflammation and health. Intestinal microflora are intimately involved in the generation of immunocompetent cells and tuning the balance between T helper 1 (Th1) and Th2 immunity during the development of the gut associated immune system. It is now generally accepted that the intestinal bacterial flora contributes significantly to the pathogenesis of inflammatory bowel disease (IBD) along with mucosal immune dysregulation and genetic susceptibility. Considerable research is focused on modifying the intestinal flora with probiotic bacteria to attenuate inflammatory activity and prevent relapses in ulcerative colitis, Crohn’s disease, and pouchitis.

A rationale for the use of probiotics in IBD stems from reports of dysbiosis in the intestinal flora in ulcerative colitis, Crohn’s disease, and pouchitis, either by conventional anaerobic culture or by analysis using molecular probes. It is however unclear whether such alterations in intestinal flora drives the inflammation or is a consequence of it. The practical application of probiotic strategy has been especially encouraged by the positive results of a trial in its use for the prevention and treatment of pouchitis. The multispecies probiotics used pose special challenges in identifying precise mechanism of action, although alterations in faecal flora have been demonstrated. Despite some positive trials, generalization from pouchitis to their use for all forms of IBD appears somewhat premature, however, as for example, a trial of administration of Lactobacillus GG after surgical resection for Crohn’s disease proved ineffective in preventing relapse. Further studies are therefore required in ulcerative colitis and Crohn’s disease before firm recommendations may be made.

Lactobacilli are a major constituent of the intestinal microflora and are frequently used as probiotics, often in the health food industry. Among the Lactobacillus genus, different species of bacteria induce distinct mucosal cytokine profiles in the gut immune system of BALB/c mice. For example, an increase in the Th2 cytokines interleukin (IL)-10 and IL-4 was observed in mice fed Lactobacillus delbrueckii subspecies Bulgaricus and Lactobacillus casei whereas, in contrast, a significant induction of the Th1 cytokines IL-2 and IL-12 was observed with Lactobacillus acidophilus. It is therefore important that notice is taken of which specific bacteria are being used.

Various knockout, transgenic, and adoptive transfer murine and rodent models of IBD have been generated and the requirement for bacterial colonisation to induce a IBD phenotype is virtually universal, despite the complexities of the immune network. In contrast, in the IL-10 knockout mouse model of colitis, probiotic therapy with Lactobacillus species and Bifidobacterium species are frequently used, the optimum use of probiotics in IBD requires greater understanding of their effects on the immune system.

In this issue of the Gut, researchers from Cork, Ireland, challenge the conventional hypothesis of mechanisms of probiotic efficacy by administering Lactobacillus salivarius subcutaneously to IL-10 knockout mice [see page 694]. The anti-inflammatory effect of subcutaneous administration was not specific as it was also seen in a murine model of arthritis. Non-viable bacteria could not be tested as the group receiving heat treated L. salivarius had 100% mortality by week 10. No change in faecal microflora occurred as a result of this subcutaneous administration of L. salivarius, suggesting a mechanism of action distinct from colonic floral modulation. Various indicators of altered immune function were seen with decreased tumour necrosis factor (TNF) and IL-12 levels being obtained from splenocytes that had been stimulated by Salmonella typhimurium. In contrast, transforming growth factor (TGF)-β levels were maintained. Such systemic anti-inflammatory activity is counter-intuitive to a simplistic model of gut flora modulation, and leads to speculation about the molecules involved in driving immunomodulation.

The findings of the current study are not the first to suggest that probiotics have more than a local anti-inflammatory effect by modulating the flora. For example, Lactobacillus casei or Lactobacillus bulgaricus reduced the inflammatory response induced by coculture of bacteria with mucosal explants from Crohn’s disease affected intestinal tissue. In this study, a significant reduction of proinflammatory cytokines such as TNF was noted. Such anti-inflammatory effect might even be systemic, as shown by the bacteria CpG DNA experiments discussed later.

Probiotics may also influence mucosal cell-cell interactions and cellular “stability” by actions such as enhancement of intestinal barrier function by modulating cytoskeletal and tight junctional protein phosphorylation. For example, live probiotics such as Lactobacillus acidophilus or Streptococcus thermophiles protect in vitro intestinal epithelial cell lines (HT29, Caco-2) from pathogen invasion and adhesion by enteroinvasive Escherichia coli. Similarly, the non-pathogenic E coli strain Nissle 1917 inhibited adhesion and invasion of intestinal epithelial cell line (intestine 407) by adherent invasive E coli strains isolated from patients with Crohn’s disease. In “the age of the genome”, it is not surprising that much time and attention has been spent on studying the importance of the detailed bacterial DNA sequences in these effects. Bacterial DNA contains non-methylated CpG motifs which bind to toll-like receptor 9 (TLR-9). TLR-9 signalling is dependent on the adaptor protein MyD88. Such immunostimulatory DNA sequences (ISS-DNA or CpG DNA) of bacterial origin have been shown to reduce inflammation in rodent IBD models such as DSS induced colitis, hapten induced colitis in BALB/c mice, and the IL-10 knockout mice model of colitis. This reduction in inflammation was accompanied by inhibition of
proinflammatory cytokine and chemokine production and suppression of induction of matrix metalloproteinases in the colon.16

Further evidence of the central role of bacterial DNA has come from novel experiments where both intragastric and subcutaneous administration of probiotic and E. coli DNA attenuated the severity of DSS induced colitis.17 The form that this DNA takes appears crucial, as methylated probiotic DNA, calf thymus DNA, and DNase treated probiotics were ineffective.

Given this complexity, do we need live bacteria, dead bacteria, or just the DNA? Unfortunately, the data are confusing and sometimes contradictory. TLR-9 and the adaptor protein MyD88 appear essential in signalling, and in their presence even non-viable bacteria can signal. In TLR-9 deficient mice, unlike TLR-2 or TLR-4 deficient mice, intragastric γ irradiated (that is, non-viable) probiotics had no effect on DSS induced colitis. Mice deficient in MyD88 did not respond to γ irradiated probiotics.17

The immune modulatory function of DNA has also been demonstrated in a study of peripheral blood mononuclear cells from healthy donors where Bifidobacterium genomic DNA caused induction of secretion of the anti-inflammatory IL-10.18

Given the high GC content of Bifidobacterium chromosomal DNA, it will be of interest to assess the effect of its subcutaneous administration in the IL-10 knockout model of colitis.

The immune modulatory properties of the various probiotic bacteria may differ, and this becomes problematic for the use of multispecies preparations. Furthermore, not all immunostimulatory oligonucleotides have the palindromic CpG motif. In one study, chromosomal DNA was purified from nine strains of Lactobacillus delbrueckii subsp. Bulgaricus and six strains of Streptococcus thermophilus derived from yoghurt starter cultures. Only DNA from L. bulgaricus NIAI B6 induced significant GC content of Bifidobacterium chromosomal DNA, it will be of interest to assess the effect of its subcutaneous administration in the IL-10 knockout model of colitis.

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What about the host? Identification of NOD2 mutations associated with Crohn’s disease provides further support for the central role of bacteria in the pathogenesis.20 Three NOD2 mutations located near or in the leucine rich repeats involve a frameshift mutation (Leu1007fsinsC) or two missense mutations (Gly908Arg and Arg702Trp). These alterations are associated with increased risk of development of Crohn’s disease and result in defective induction of nuclear factor κB (NFκB) activation by bacterial peptidoglycan and muramyl dipeptide (MDP). MDP induced activation of NFκB in mononuclear cells is absent in patients with Crohn’s disease homozygous for the Leu1007fsinsC mutation.21 22 It is therefore interesting to speculate that NOD2 mutations associated with Crohn’s disease result in defective sensing of some bacteria which may precipitate inappropriate diffuse activation of NFκB and inflammation through non-NOD2 mechanisms. Repeating the experiments with Crohn’s mucosal explants23 from patients with homozygous, heterozygous, and double heterozygous NOD2 mutations, and appropriate controls cocultured with Lactobacilli, may provide interesting data using TNF readouts.

Our attempts to understand how bacteria modulate the immune system will undoubtedly yield novel therapeutic targets and therapeutic agents. Irrespective of whether or not dysbiosis is reproducibly confirmed in IBD, we need to consider probiotic therapy in terms of specific molecules modulating defined targets in the gut mucosal and systemic immune system, and move

Figure 1  Pathogen associated molecular patterns (PAMPs) derived from bacteria (including probiotics) are recognised by pattern recognition receptors (PRRs, such as Toll-like receptors). Initiation of dendritic cell (DC) maturation starts after ligation of PRRs, which also requires adaptor proteins such as MyD88 for signalling. Type of PAMPs determines the selective priming of DCs for production of Th1, Th2, and Treg lymphocyte polarising factors. Different PAMPs ligate to specific corresponding PRRs. IL, interleukin; TNF, tumour necrosis factor; TGF-β, transforming growth factor β.
away from a simplistic concept of re-
populating the intestinal flora with "friendly" bacteria. Bacteria are only "friendly" in the context of the desired immune modulation required for a specific disease. The environment to which the immune system is exposed to determines DC phenotype and state of activation and eventually drives the balance between effector and regulatory T cells (Fig 1). Understanding probiotic action may permit modulation of the immune system, both locally and systemically. The article by Shell and colleagues is the right step towards stimulating further research where immunologists, microbiologists, and gastroenterologists can collaborate.


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Crohn’s disease

Fistulising Crohn’s disease: MMPs gone awry
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Broadening our understanding of the role of matrix metalloproteinases in Crohn’s disease

Fistulae are a common complication of Crohn’s disease (CD), and their most common perianal manifestation is present in 14–38% of CD patients in referral populations. Despite advances in conservative treatment, fistulae rarely heal, while surgical resection is effective but frequently does not prevent local recurrence or fistulising disease at other sites. Remodelling of the extracellular matrix (ECM) is a key event in chronic bowel inflammation, especially in CD which is characterised by both active fibrogenesis (that is, ECM production and deposition), leading, for example, to stricture formation, and by enhanced fibrolysis (that is, breakdown and removal of ECM), such as occurs in fistula formation. While fibroblasts and myofibroblasts, and to a minor degree endothelial and epithelial cells, produce the intestinal ECM, many more cell types are involved in ECM breakdown by releasing a broad spectrum of enzymes that can degrade essentially every ECM component, such as the various collagenas, non-collagenous (glyco-) proteins, glycosaminoglycans, and proteoglycans. The most important of these enzymes are the matrix metalloproteinases (MMPs), a structurally related class of at least 20 zinc dependent proteases. MMPs are classified...
according to their primary specificities and structural features into interstitial collagenases (MMP-1, MMP-8, MMP-13, MMP-18), gelatinases (degrading denatured and basement membrane collagens: MMP-2, MMP-9), stromelysins (degrading a broad spectrum of ECM substrates: MMP-3, MMP-7, MMP-10, MMP-11), elastases (MMP-12), and membrane-type (transmembrane) MMPs (with broad substrate specificities: MMP-14, MMP-15, MMP-16, MMP-17, MMP-24, MMP-25).

As uncontrolled MMP activity would virtually lead to dissolution of organs, MMP activity is strictly controlled, not only at the transcriptional and translational levels, but also by the requirement of proteolytic activation of thezymogens and by several protease inhibitors. Proteolytic activation occurs by the plasminogen activator-plasmin cascade and by certain active MMPs themselves, such as MMP-3, MMP-10, MMP-14, and MMP-24. The most important physiological inhibitors are the tissue inhibitors of metalloproteinases (TIMP), mainly TIMP-1, which blocks almost all MMPs, and TIMP-2, which is specific for MMP-2. A fifth level of control is localisation of secreted MMPs either at cell membrane compartments that are involved in local ECM proteolysis (for example, by binding to ECM receptors that mediate cellular locomotion through the matrix) or on certain ECM components, such as collagens.

Interestingly, it is preferentially the inactive proforms of MMPs that bind to collagen which become released from the ECM on proteolytic activation. This illustrates that both homeostasis and proteolytic remodelling of the ECM are highly sophisticated and tightly regulated processes, and that simply measuring upregulation or downregulation of MMPs and their inhibitors will give a very incomplete picture of ECM remodelling at best. In this line, no characteristic difference is found between procollagen mRNA levels as a measure of fibrogenesis and MMP and TIMP mRNA levels as a measure of fibrolysis in inflamed colonic specimens from patients with CD and patients with UC colitis. This was initially unexpected as CD often leads to stenosis due to excess ECM deposition while fibrosis is uncommon in ulcerative colitis. But it comes as no surprise when one considers the complex regulation of fibrolysis in the transmural inflammation of CD.

The paper by Kirkegaard and colleagues1 in this issue of Gut brings our understanding of the role of MMPs in CD a step further [see page 701]. Using complementary techniques, they identified and investigated MMP-3 and MMP-9 as key players in fistulae of CD and other causes. At first glance the occurrence of fistulæ, a manifestation of enhanced fibrolytic activity, in a fibrostenotic intestinal disease may appear paradoxical. However, severe inflammation which always leads to release and activation of MMPs by cells of the inflammatory infiltrate can drive both processes: fistulæ will form when there is no rapid compensatory fibrogenic response to fill up the defect, perhaps favoured by a too quick re-epithelialisation or re-endothelialisation; fibrosis will develop when the fibrogenic response is strong and quick, leading to scar formation. Both processes can obviously coexist in close proximity. What are the factors determining these divergent pathways? Recent studies demonstrated that at least early activation of MMPs is necessary to allow for a fibrogenic response,1,2 either by destroying basement membranes which have a role in maintaining cellular quiescence and differentiation in the gut8 with subsequent mesenchymal activation, or by allowing mesenchymal cell migration and proliferation by removal of constraining ECM. Thus the cellular source and temporospatial pattern of MMP release and activation probably explain these divergent pathologies. As mesenchymal cells (fibroblasts, myofibroblasts, and to a lesser extent endothelial cells and inflammatory cells (mainly macrophages, monocytes, and neutrophils) are the major sources of MMPs and TIMPs, there must be a differently tuned interplay of these cells driving either fistula or fibrosis. Kirkegaard et al observed that acute fistulising inflammation in CD and other aetiologies is characterised by high expression of MMP-3 and MMP-9 coupled with high activity of MMP-2 and MMP-9 in inflammatory cells, while in chronic fistulæ, MMP-9 almost disappeared but MMP-3 expression was maintained, with a shift to mesenchymal, mainly myofibroblastic, cells.1 The finding that TIMPs remained low in all fistula specimens is of particular interest. The TIMP-1/3 ratio is a major determinant of fibrogenesis, as exemplified in liver fibrosis.17 Low expression of TIMP-1 is also characteristic of osteoarthritis, the hallmark of which is unopposed MMP activation in the joint leading to destruction of cartilage and bone, somewhat reminiscent of fistula formation in the gut.18 Also, it is osteoarthritis that has led to a novel concept of chronic fibrolytic inflammation that is driven by altered synovial fibroblasts, the equivalent of which could be myofibroblasts in chronic fistulising diseases. These fibroblasts/myofibroblasts can secrete atypical chemokines attracting fibrolytic inflammatory cells on the one hand, and have a fibrolytic and migratory phenotype themselves on the other.19 MMP-3, which has a broad substrate specificity and which acts as a proactivator (in addition to plasmin), at least of MMP-1, MMP-2, and MMP-9, has been identified as a particularly aggressive enzyme in intestinal inflammation which favours invasiveness.20 This does not exclude an important contribution by MMP-12, MMP-13, MMP-14, and MMP-19 which are expressed in inflamed intestine21 but were not considered in this study.

Can we derive therapeutic concepts from this and related studies on intestinal inflammation? A drug that effectively prevents emerging or eliminates chronic fistulæ would be most welcome. Anti-inflammatory, antibiotic, and immunosuppressive approaches can lead to long term fistula control or even closure in 25–50% of patients, roughly doubling the placebo response which is unsatisfactory.1 Anti-tumour necrosis factor (TNF-α) strategies can lead to good short term and occasionally long term results in cases that are refractory to these therapies but treatment is costly and may have grave side effects.1 None the less, this approach is rational as TNF-α is a potent inducer of MMP-3 in intestinal myofibroblasts.22 Based on this and other studies, four novel strategies aimed at inhibiting intestinal inflammation and remodelling in inflammatory bowel disease in general, or in fistulising CD in particular, merit further exploration. These are the use of: (1) inhibitors of growth factors or cytokines, such as epidermal growth factor, hepatocyte growth factor, insulin-like growth factor (IGF-1 and IGF-II), fibroblast growth factor (aFGF, bFGF), interleukin 1, and stem cell factor that induce MMPs in epithelial and mesenchymal cells;23 (2) synthetic inhibitors of kinases that trigger MMP production, especially p38 kinase;24 (3) synthetic MMP inhibitors that were shown to ameliorate experimental colitis;25 and (4) integrins (that is, cellular receptors for ECM proteins that can bind to and/or induce certain MMPs).26–28 Inhibition of growth factors/cytokines and kinases always carries the risk of side effects because of their importance in normal tissue regeneration and of their ubiquitous expression in other cells and organs. Inhibition of MMPs appears more promising, especially as the first generation of broad spectrum inhibitors which have multiple side effects are becoming replaced by orally available subtype specific agents.29 However, the
necessity for cell-type specific targeting (for example, to MMP-3 expressing myofibroblasts of fistulae) remains an unsolved problem. Therefore, it is attractive to target integrins that promote MMP-expression as many of these receptors are cell-type specific. Thus MMP-1, MMP-2, and MMP-3, MMP-14, MMP-1 and MMP-3, and MMP-9 are induced and activated by integrins α2β1, α4β1, α6β1, and αvβ3, respectively, which are predominantly found on activated (myo-) fibroblasts. Such inhibitors have (in part) been developed, and orally available agents are in phase I and II clinical studies (for example, for metastatic tumour disease). It remains to combine the most promising of these agents, alone or in combination, with established therapies in preclinical and clinical studies, for our patients with complicated inflammatory bowel disease.

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transcription-polymerase chain reaction showed predominant expression in liver. The peptide was called LEAP-1 (liver-expressed antimicrobial peptide).\(^1\) The cDNA sequence predicted an 84 amino acid pre-propeptide with two cleavage sites. The first predicts cleavage of the N terminal endoplasmic reticulum targeting signal sequence (amino acids 1–24). The second consensus cleavage site for prohormone converses would yield the central prodomain (amino acids 25–59) plus the 25 amino acid C terminal peptide (also called hepcidin-23; amino acids 60–84). A slightly truncated 20 amino acid peptide was also found (hepcidin 20; amino acids 63–84). Independently, Park and colleagues\(^1\) isolated the same cysteine rich peptide from human urine and named it hepclin, reflecting its hepatic expression and bactericidal properties. In addition to hepcidin-20 and hepclin-25, a minor 22 amino acid form was found (hepcidin-22; amino acids 63–84). The antimicrobial activity of urinary hepcidin was inhibited by 100–150 mM NaCl,\(^1\) similar to the normal range for plasma sodium (\(\sim 135–145\) mM). Whether circulating plasma hepcidin plays a significant antimicrobial role in vivo therefore remains unclear.

Experiments in mice have shown that hepcidin plays a role in iron regulation. Pigeon et al identified mouse liver hepcidin mRNA because it was induced by carboxyl iron overload.\(^5\) The mouse has two hepcidin genes in tandem (probably arising from a duplication event), located adjacent to the upstream stimulatory factor 2 (Usf2) transcription factor gene.\(^5\) Mice that lack hepatic hepcidin expression, due to targeted disruption of the adjacent Usf2 gene, developed predominant iron loading of parenchymal cells of the liver and pancreas with relative sparing of the macrophage rich spleen.\(^6\) Similar patterns of iron overload in Hfe knockout and Usf2 (hepcidin deficient) knockout mice expressed that hepclin and Hfe may function in the same regulatory pathway. Hepclin was proposed to be a negative regulator (that is, “inhibitor”) of iron absorption in the duodenum and of iron release from macrophages.\(^7\) This was supported by demonstration that transgenic mice constitutively expressing liver hepcidin died within a few hours of birth with decreased body iron levels and severe microcytic hypochromic anaemia.\(^7\)

Further observations implicate hepcidin in the anaemia of chronic disease. This is the most common anaemia of hospitalized patients, often associated with infection, cancer, and autoimmune diseases. There is a diminished response of erythroid precursors to erythropoietin, decreased survival of erythrocytes, impaired iron absorption, and retention of iron in macrophages. The net effect would be to decrease the iron available to erythroid precursor cells.\(^8,9\) Two patients with severe iron refractory anaemia associated with hepatic adenoma showed spontaneous resolution of anaemia after removal of the tumour.\(^9\) The adenomatous tissue expressed inappropriately high levels of hepcidin mRNA. On the basis of these observations, it was proposed that hepcidin may be important in the pathogenesis of anaemia of chronic disease.\(^8\) Analysis of urinary hepcidin excretion in patients with anaemia due to chronic infection or severe inflammatory disease provided some confirmatory evidence of increased hepcidin levels in response to inflammation in humans.\(^9\) In these patients, hepcidin excretion correlated with serum ferritin concentration which, like hepclin, increases in response to both elevated body iron stores and inflammation. Interestingly, injection of erythropoietin into mice was shown to decrease dramatically liver hepcidin expression. Therefore, the efficacy of erythropoietin is mediated in part by suppression of hepclin expression, leading to increased plasma iron levels.\(^10\)

Regulation of hepclin expression has been investigated in cultured cells. Surprisingly, iron loading of primary human hepatocytes with either ferric ammonium citrate or dextriferrin decreased hepclin mRNA levels.\(^10\) This is in contrast with experiments in mice in which dietary or parenteral iron loading increased hepclin mRNA levels.\(^11\) This suggested that during iron overload in the whole animal, a signal from non-parenchymal cells may induce hepclin to express hepclin.\(^11\) Mouse liver hepcidin mRNA is induced not only by iron overload but also by treatment with bacterial lipopolysaccharide, hinting that iron and inflammatory cytokines may have a common signalling pathway.\(^11\) Hepclin mRNA in primary human hepatocytes was induced either by medium conditioned by monocytes exposed to lipopolysaccharide or by interleukin 6 (IL-6).\(^11\) At the level of regulation by transcription factors, the promoters of human and mouse hepclin genes contain at least one functional binding site for CCAAT/enhancer binding protein C (CEBPx), a transcription factor highly expressed in adult liver which plays a role in stimulating hepclin expression. Hepatocyte nuclear factor 4x (HNF4x) appears to reduce hepclin expression.\(^12\)

The medical importance of hepclin is emphasised by recent studies in human iron overload. Homozygous mutations of hepclin have been demonstrated in two families with juvenile haemochromatosis.\(^13\) This disorder is characterised by a rapid rate of iron loading, usually causing presentation before the age of 30 years. There is a high frequency of endocrine and cardiac iron overload, and failure of early diagnosis and treatment may be fatal.\(^14\) Juvenile haemochromatosis is genetically heterogeneous with a second form mapping to chromosome 1q21. Whether this second gene will encode another protein in the hepclin-Hfe pathway remains to be resolved.\(^15\) The potential interaction between hepclin and Hfe has also been suggested by the description of haemochromatosis associated with one mutated allele of both hepclin and Hfe, termed “digenic inheritance”.\(^16\)

The classical Hfe related form of the disease accounts for more than 90% of well characterised haemochromatosis patients in the UK.\(^17\) In Hfe related haemochromatosis and in Hfe knockout mice, there is failure to upregulate hepclin levels despite hepatic iron overload. This indicates that the Hfe protein is involved in regulation of hepclin levels in response to iron. Disruption of the normal hepclin response to body iron stores may contribute to the pathogenesis of iron overload seen in Hfe related haemochromatosis.\(^18–20\)

Recognition of liver produced hepclin as an iron regulatory factor has challenged the hypothesis that the control of iron absorption in relation to body stores is mediated in duodenal crypts. Hepclin has revived the concept that some regulation may be mediated by the liver. Rats switched from a control to an iron deficient diet showed changes in hepclin expression in close temporal relationship with changes in duodenal iron transporter expression. This suggested that hepclin may act directly on villous enterocytes, pointing to the liver as a regulatory site.\(^21\) Certainly, siderosis in Kupper cells, often on a background of mild iron deposition in hepatocytes, is a familiar feature of common liver diseases such as hepatitis C infection and alcoholic liver disease.\(^22\) Whether increased concentration of hepclin, perhaps in response to inflammation, has a role in the hepatic siderosis observed in viral and alcoholic liver disease remains to be demonstrated. To date, investigation of hepclin in various clinical conditions has been impeded by the lack of anti-hepcidin antibodies and suitable assays. Recently, a western blot densitometric assay was reported using an antibody raised
against hepcidin-25, allowing quantitation of human hepcidin in serum samples. In this issue of Gut, Kulaksiz and colleagues describe antisera raised against two unique synthetic peptides corresponding to the predicted human hepcidin prodomain (two antisera to the same peptide) and the mature hepcidin C-terminal peptide (see page 735). All three antisera detected immunoreactive bands estimated at 10 and 20 kDa on western blots of guinea pig and human liver and HepG2 cells. The 10 kDa band only was detected faintly in human serum. Immunohistochemistry of guinea pig and human liver with all three antisera showed staining in hepatocytes with a basolateral expression pattern. This confirms and extends earlier mRNA results, and is consistent with basolateral release of hepcidin into the liver sinusoids.

A new enzyme-linked immunosorbent assay (ELISA) was developed using the prodomain antisera. The ELISA was used to investigate serum levels of prohepcidin immunoreactivity in controls and patients with haemochromatosis, renal anaemia, and chronic renal insufficiency. The mean level of pro-hepcidin immunoreactivity detected in the serum of healthy volunteers was an order of magnitude greater than levels of hepcidin detected in human urine. Pro-hepcidin immunoreactivity was increased in chronic renal insufficiency patients who had normal haemoglobin levels compared with the healthy control group, despite erythropoietin treatment, which has been observed to decrease hepatic hepcidin expression. These observations are therefore consistent with renal metabolism and/or elimination of peptide. Pro-hepcidin was decreased in HFE-related haemochromatosis patients (C282Y homozygous) compared with healthy controls, consistent with earlier mRNA studies, and indicating defective hepcidin regulation in HFE-related haemochromatosis.

Questions and inconsistencies remain regarding the size and identity of hepcidin immunoreactive bands on western blots. The sequence of hepcidin-25 predicts a mass of 2.7 kDa, corresponding to the result determined by mass spectrometry. Furthermore, the oxidised synthetic hepcidin-25 co-migrates with native hepcidin-25 in analytical high pressure liquid chromatography and capillary electrophoresis. On western blots however, serum hepcidin-25 immunoreactivity has been detected with an apparent mass of 10 kDa, co-migrating with the synthetic peptide, possibly reflecting aggregation of hepcidin monomers. Kulaksiz et al detected bands of 10 kDa and 20 kDa with their three antisera on western blots. They attributed the 10 kDa band to prohepcidin on the grounds that it was detected by antisera to both the mature hepcidin peptide and to the prodomain. However, the predicted mass of pro-hepcidin is 6.9 kDa (dExPASY programme: http://ca.expasy.org/tools/peptide-mass.html). It might be informative to compare the mobility of synthetic peptides versus the bands observed in human serum in this system. Perhaps attention to denaturation conditions and accurate verification of experimentally determined molecular masses may help to clarify the discrepancies.

Is pro-hepcidin an active player in iron regulation or merely a precursor and an indirect measure of hepcidin potential? Interestingly, a homozygous point mutation of the human hepcidin prodomain has been detected, predicting a threonine to methionine substitution at amino acid 31 (T31M). This individual had normal serum iron indices, normal haemoglobin, and normal mean corpuscular volume. However, this residue is not conserved in evolution: both wild-type mouse hepcidin genes have a methionine at the position equivalent to human threonine 31. The normal phenotype associated with homozygous T31M mutation therefore does not resolve the role of pro-hepcidin.

In summary, hepcidin is a peptide hormone with important biological effects on iron metabolism. It has sequence homology to antimicrobial peptides, although whether this function is conserved in vivo remains to be demonstrated. Hepcidin is derived by cleavage of a pre-propeptide; questions remain about the processing and activity of the peptide products. It has been implicated as a regulator of body iron stores and mutations have been found in some families with juvenile haemochromatosis. Iron balance is probably sensed by Kupffer cells and signalled to hepatocytes by cytokines such as IL-6. Hepcidin expression is also induced by inflammatory agents, giving it a potential role in the anaemia of chronic disease. The reagents and assays described will permit investigations of the role of hepcidin in clinical samples in relation to the siderosis observed in viral and alcoholic liver diseases. This peptide is a fascinating addition to the family of molecules involved in iron metabolism. Future studies of its action and regulation in vivo and in vitro should fill in some of the tantalising gaps in our understanding of this increasingly complex field.

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COMMENTARY

Gastrointestinal symptoms

Bringing science to the art of diagnosis

P Moayyedi

Which gastrointestinal symptoms are useful in distinguishing organic from functional disease?

Patients and clinicians are becoming increasingly intolerant of diagnostic uncertainty. This is reflected in the rising demand for endoscopic procedures implicitly suggesting that gastrointestinal symptoms are an unreliable indicator of serious pathology. Unfortunately, the growing array of tests that are being demanded for patients is placing further pressures on already stretched health care budgets. The natural reaction to this is to evaluate whether we can improve on the value of the history to diagnose gastrointestinal disease. This process was started over 30 years ago when researchers such as Card and colleagues and de Dombal and colleagues evaluated the use of computers to aid the clinician in making diagnoses in patients with upper gastrointestinal symptoms. The enthusiasm for this approach faded when data suggested that computer improved diagnostic accuracy of computers was not sufficient to prevent investigations. The paucity of subsequent research in this field is disappointing as it would be useful to know what information computers were using that enhanced diagnostic acumen. The article by Hammer and colleagues in this issue of Gut [see page 666] is therefore refreshing as it prospectively evaluates a wide range of gastrointestinal symptoms to establish which are useful in distinguishing organic from functional disease. The strength of this study is that it evaluates a large relatively unselected group of patients, reducing spectrum and selection bias. It also divided patients into those with and without organic disease rather than subdividing the data into different diagnoses. This improves the power of the study and gives the clinician overall information on whether the patient is likely to have an organic disease.

Their data suggest that aspects of the history are valuable diagnostic tools for lower gastrointestinal disease. They identify three ‘alarm features’ that were important independent predictors of organic lower gastrointestinal pathology. These were age over 50 years, male sex, and blood on the toilet paper. They also identified five ‘non-alarm’ features (frequent abdominal pain, radiating pain, pain with loose bowel movements, reflux, and absence of diarrhoea) that independently predicted functional bowel disease. The approach of positively diagnosing functional bowel disease with non-alarm symptoms or positively diagnosing organic disease with alarm symptoms performed equally well. The interesting observation is that putting these two pieces of information together improved the accuracy of the model. This is intuitive, as a clinician is likely to weigh up the absence of alarm symptoms and the presence of positive features indicating a functional bowel disorder before deciding that the patient is at low risk of significant organic disease.

These data will provide useful information for the Rome group when they next consider guidelines for the assessment of irritable bowel syndrome (IBS). They have highlighted symptoms similar to the classical Nanning criteria that are important in making a positive diagnosis of functional gastrointestinal disease but also other characteristics such as radiating and frequent pain that make IBS more likely. Unfortunately, the model is probably not sufficiently accurate to be used to exclude patients from investigation in clinical practice. Using the prevalence of lower gastrointestinal organic disease in the article, the model had a positive predictive value of 79% and a negative predictive value of 86%. The possibility of missing organic pathology in 14% of patients would lead most clinicians to opt for investigating even if the model predicted a low probability of disease. The accuracy of the model may have been improved if non-gastrointestinal sympotms such as backache and frequency of micturition had been included but the concern about using models to predict organic disease is emphasised by the conflicting messages from other studies. For example, three papers also found that rectal bleeding predicted organic disease whereas two studies found no statistically significant association. Hammer et al found that weight loss and anorexia were not independent predictors of disease whereas others reported these symptoms were associated with organic pathology. Finally, Hammer and colleagues found that severe abdominal pain was associated with a reduced risk of finding organic pathology in the univariate analysis. This observation is supported by one study whereas another suggested
abdominal pain is associated with colorectal disease.15

These inconsistent findings probably relate to differences in the design and populations of the different studies. The quality of the data collection on gastrointestinal symptoms, sample size, and prevalence of underlying organic disease varied between studies. In particular, Hammer et al had very few colorectal cancers in their sample and the main organic disease driving the model was inflammatory bowel disease. If ulcerative colitis was the main disease found, then rectal bleeding and absence of pain will predict pathology and weight loss may not be an important feature in the model. It is interesting that the study12 that also had few colorectal cancers found similar results, whereas the study16 that evaluated a larger number of colorectal cancers found that features such as weight loss and abdominal pain were important. The case mix of disease in the underlying population is likely to have a large influence on the symptoms that computers models will predict are important.

Upper gastrointestinal symptoms are a good example of this. Hammer et al found that weight loss, dysphagia, age, and sex were not predictors of organic pathology whereas other researchers have found these to be significantly associated with the risk of upper gastrointestinal cancer.18 Indeed, it is almost inconceivable that male sex and increasing age would not be risk factors for upper gastrointestinal malignancy given what is known about the epidemiology of these cancers.1 Hammer et al did not find that these were important because the majority of their patients had gastro-oesophageal reflux and peptic ulcer and these diseases were driving the model. It is not surprising therefore that heartburn symptoms and aspirin use were found to predict pathology whereas alarm symptoms did not.

Although alarm symptoms may be of more value than is suggested by this paper, the overall message is consistent with other researchers. Factors that are found to be predictive of functional or organic disease are not very useful clinically. The model used by Hammer et al found it very difficult to distinguish between groups and as most patients had organic disease it generally defaulted to assigning patients to the disease group. Others have found the same problem: individual factors may be statistically significantly associated with disease but when they are combined in a model they generally perform very poorly.16 This is particularly true when the models are tested prospectively or transferred from a hospital to a primary care setting.22

Science has a great deal to offer the clinician in helping determine a diagnosis from the patient history. Statistical techniques can identify symptoms and signs that are important to consider and those that do not contribute to making a diagnosis. The problem is that a single study is not sufficient. After all, we do not rely on one study to definitively tell us that smoking causes lung cancer or Helicobacter pylori causes distal gastric adenocarcinoma. There is a need for a range of diagnostic studies conducted in different settings using different designs. The common feature of all should be the careful prospective collection of a detailed symptom assessment.23 When sufficient data have been accumulated, a picture will emerge on what are the important features that identify a patients as having organic disease. The problem that is addressed also has to be amenable to this approach. It will be difficult to use the history to guide us in patients with upper gastrointestinal disease but the situation with lower gastrointestinal pathology looks more promising. Insights provided by diagnostic studies may allow us to avoid invasive tests in some patients and identify others as having a high risk of serious colorectal disease that warrants urgent investigation. It is not sufficient to show the sensitivity and specificity of this approach as the ultimate goal is to provide more cost effective health care to patients.24 Randomised controlled trials will therefore be required to assess whether a more thorough clinical history will reduce costs or improve patient outcomes.


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