Microsatellite instability in colitis associated colorectal cancer

As molecular pathways of colon carcinogenesis continue to be defined for high risk hereditary colon cancer syndromes and for average risk sporadic colon cancers, it seems that colon carcinogenesis in the setting of chronic idiopathic inflammatory bowel disease (IBD) may be unique. Although to some extent this notion seems intuitive because of the substrate of chronic inflammation from which these cancers arise, it is nevertheless curious that despite the setting of chronic inflammation, colon cancers occurring in patients with colitis share several features in common with those that arise in patients with hereditary non-polyposis colorectal cancer (HNPCC). For example, in both conditions there is a tendency for the colorectal cancers to affect young individuals, be multifocal, show a proximal colonic distribution, and display mucinous, signet-ring cell or undifferentiated histology. In addition, HNPCC colorectal cancers often have a rim of surrounding inflammation, referred to as a Crohn’s-like reaction. These clinicopathological similarities have prompted some investigators to explore whether colitis associated cancers might in fact share a common molecular pathogenesis with HNPCC colorectal cancers.

Colorectal cancers in patients with HNPCC occur by virtue of a defective ability to repair DNA base pair mismatches. This results in DNA replication errors that have the net effect of altering genes that are critical for maintaining normal growth and behaviour of colonic epithelial cells. Such replication errors can be identified by using markers that detect short repetitive sequences, or microsatellites, located throughout the genome. Errors in replicating these sequences result in a phenotype termed microsatellite instability (MI). Several genes act in concert to orchestrate normal repair of DNA base mismatches, including hMLH1, hMSH2, hMSH3, hMSH6, hPMS1, and hPMS2. Except for hMSH3, germline mutations of any one of these genes are known to give rise to tumours of the colorectum and other organs in patients with HNPCC, with mutations of hMLH1 and hMSH2 accounting for the vast majority of known HNPCC families.

In this issue (see page 367), Cawkwell and colleagues investigated the frequency of MI in colitis associated colorectal cancers to determine whether these tumours might share a similar molecular pathogenesis with HNPCC. They analysed 46 colitis associated colon adenocarcinomas for MI using a panel of four markers in a fluorescence PCR-based MI assay, and also stained tumours by using immunohistochemistry for loss of hMLH1 and hMSH2. Six (15%) of 41 cases demonstrated MI at one or more marker, whereas only one (2.4%) case had MI at two or more of the four markers. The latter case was also the only one that demonstrated loss of immunohistochemical staining of hMSH2, a useful surrogate marker of higher rates of MI in tumours. The authors suggest that MI is uncommon in colitis associated colon cancers, thus distinguishing these cancers from HNPCC where the frequency of MI positivity is about 85%, but perhaps somewhat similar to sporadic colon cancers which have a 15% MI positivity rate.

Before accepting that MI is uncommon in colitis associated neoplasms, it is worth considering that other investigators have reported a somewhat higher frequency of MI in colitis associated dysplasias and cancers. Using a panel of five microsatellite markers, Suzuki et al noted MI of at least one marker in 17/63 (27%) patients, and 15/120 (8.3%) dysplastic and cancerous lesions manifested MI of two or more markers. These investigators also reported that some colitis associated neoplasms manifested mutations of transforming growth factor β1 type II receptor, a target gene that is often affected by MI in HNPCC. Brentnall and coworkers used a panel of 13 markers and observed MI at least one marker in 40% of cancers and 85% of high grade dysplasias, and MI of at least two markers in 40% of cancers and 46% of high grade dysplasias. In that study, MI was quite frequent even in colitic mucosa that was non-dysplastic, regardless of whether the patient had colitis of short (less than three years) or long (more than eight years) duration, and MI was not found in acute, self limited forms of colitis. Furthermore, Heinen and colleagues used a panel of six microsatellite markers and noted MI at one locus in 13% non-dysplastic, 5% dysplastic, and 17% cancerous lesions in colitis, and curiously, only non-dysplastic mucosa manifested MI at two loci.

How can we reconcile the differences among these studies? To be sure, the studies differ with regard to the number and types of neoplastic tissues analysed and to some extent the method of analysing MI. However, the crucial difference relates to the choice and number of markers used in the various studies. Because MI testing has been so variable between laboratories, a National Cancer Institute workshop established consensus recommendations in 1998 suggesting that a reference panel of five markers be used to classify a tumour as having either a high (MI-H), low (MI-L), or stable (MS) microsatellite phenotype. If the recommended five loci are analysed, a tumour is classified as MI-L if one locus exhibits instability, or MI-H if two or more loci are unstable. If more than the five markers are used, MI-L status would apply to tumours manifesting instability at less than 30–40% of loci, whereas tumours with instability of more than 30–40% of loci would be considered MI-H. Importantly, included among the five reference markers are BAT25 and BAT26, which recognise mononucleotide repeats and are therefore considered more sensitive markers. It is possible that if Cawkwell and coworkers had used these more sensitive markers, the frequency of MI in their colitis associated cancers might in fact be higher than reported.

As we await additional studies using standard markers, a common theme seems to already be emerging from the existing literature: microsatellite instability, especially the MI-L phenotype, may in fact be fairly common in ulcerative colitis, both in neoplastic and non-neoplastic tissue. Some hypothesise that if the DNA repair machinery is defective in ulcerative colitis mucosa, the oxidative stress that accompanies chronic inflammation can damage DNA and overwhelm the ability of the colonocytes to repair DNA, resulting in the MI phenotype of even non-dysplastic mucosa. Although the biological and clinical significance of low level
How should endoscopic accessories be selected: trial or error?

Before being put on the market, medicines undergo stringent evaluation of their efficacy and safety, satisfying the regulatory bodies of individual countries. The costs of these activities are borne by manufacturers or developers who may later be held responsible for subsequent shortcomings. By contrast, medical devices, as for example those used in endoscopic procedures, require no such prior assessment in many countries. In the United Kingdom, since June 1998, all new products must satisfy the requirements of the Medical Devices Agency,¹ the UK Competent Authority, to be awarded the CE mark allowing use throughout the European Community. Endoscopists and assistants commonly learn of new products from commercial stands at exhibitions held alongside scientific meetings or from manufacturers’ representatives or sales staff. The apparent virtues and novelty of products are exalted whereas objective clinical evidence of efficacy or safety is usually not available. Phase III development is not a clearly defined prerequisite before device marketing. Why should devices be less well investigated and evaluated than drugs when offered for sale? The initiative and ingenuity of manufacturers in developing and producing new equipment is to be warmly commended, but enthusiasm for sales ahead of rigorous clinical evaluation must be seriously questioned.

Endoscopists are confronted with an overwhelming range of equipment from competing companies and at present choices are based upon previous experience, recommendations from colleagues and an inspection of the product. This situation is unsatisfactory and devices must be subjected to the same rigorous evaluation as medicines. Who should organise such trials? Ideally, the manufacturers in necessary collaboration with clinicians, obeying the rules of controlled trials with ethical approval. The expense would be considerable but less than in the case of medicines. Moreover the time between development and commercial release would be similarly prolonged—but would this necessarily be a bad thing? Many products currently offered for sale would not reach the market place.

One alternative approach is for trials to be organised by clinicians, perhaps with commercial sponsorship, for independent publication in peer-reviewed journals. The paper by England et al in this issue (see page 395) is an excellent example of such collaboration, investigating two designs of stents for palliation of malignant obstruction of the bile duct. The study was supported by the equipment manufacturers and the findings were analysed by a professional monitoring group according to European Good Clinical Practice guidelines.

Bacterial colonisation and encrustation of plastic biliary stents has been shown by in vitro studies to be affected by the surface characteristics of the stent material and in vivo studies by the structure of the stent used. Much interest has focused on the potential for reducing encrustation by removal of side holes from the stent design—for example, the “tir tree” or Tannenbaum stent. A previous non-randomised study showed significantly improved patency compared with the pig tail design.¹² Patency rates for 12F and 10F straight stents are similar whereas 7F and 8F stents are more prone to occlusion.¹³¹⁴ Use of plastic stents in the palliation of malignant obstructive jaundice has been challenged by the superior performance of large bore expanding metal stents with improved patency rates revealed in four prospective randomised trials.¹⁵¹⁶ However, at detection, the majority of such strictures are caused by pancreatic malignancy often with a median survival less than the median patency of plastic stents. Many of those who develop locoregional jaundice are not restented with patients’ wishes and compliance rather than procedural logistics proposed as the major factors.¹⁷ Diagnostic uncertainty at the outset reserves a role for an easily removable and exchangeable stent—factors inherent in plastic constructions. Whether the choice of stent affects overall survival is less clear.

England et al have looked at a remaining issue of whether the 10F Teflon Tannenbaum stent or the more commonly used 10F polyethylene Cotton-Leung stent with side holes, achieved longer patency. In a well planned prospective multicentre study they randomised at deep cannulation 134 patients presenting with a malignant common duct stricture to palliation using either the Tannenbaum or the Cotton-Leung design. All patients received prophylactic antibiotics at stenting. Respective groups of 65 and 69 patients were similar in age, type of malignancy (over 60% pancreatic), length of stricture, and American Society of Anesthesiologists’ classification. There was an overall generous use of needle knife papillotomy at 35%, but this was evenly distributed between the groups. No statistical difference was found in the time to restenting or death with jaundice.

MI is not yet clear, this issue is receiving increasing attention. For example, germline hMSH6 mutations have recently been reported to give rise to an attenuated HNPCC phenotype.⁹ Moreover, based upon observations made in yeast models, inactivation or inhibition of hMSH6 might result in errors at mononucleotide tracts, but not dinucleotide or longer repeat elements.⁹ This raises the possibility that mutations of hMSH6 or perhaps silencing of the gene by methylation might contribute to the neoplastic process in ulcerative colitis, a prospect worthy of further investigation.

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Combination of interferon α therapy with non-steroidal anti-inflammatory drugs in chronic hepatitis C

Chronic infection with hepatitis C virus (HCV) affects 2–15% of the world’s population. In the USA there are more than 4 million chronically HCV infected people and the estimates for Europe are similar. In the European Community chronic hepatitis C is viewed as a major health problem and efforts are being made to develop both preventive and new therapeutic strategies (Key Action 2 of the 5th Framework Programme 1998–2002 of Quality of Life and Management of Living Resources).

Treatment of chronic hepatitis C is currently based on interferon α (IFN-α) with limited efficacy, only achieving a sustained virological response in less than 25% of patients with a favourable profile treated for one year.1 To improve the response rate of standard IFN-α monotherapy, several regimens have been tried combining IFN-α three times weekly for six to 12 months with various other agents. These agents include those with antiviral activity like ribavirin, amantadine,2 and azithromycin; others with immunomodulatory properties such as thymosin; those with liver enzyme lowering activity like ursodeoxycholic acid and ribavirin;2 and others with miscellaneous properties such as ofloxacin, N-acetylcysteine, glycyrrhizin, and non-steroidal anti-inflammatory drugs (NSAIDs). The aim of all these efforts was to increase IFN efficacy by decreasing the number of non-responders and relapsed responders as well as to reduce the dose of IFN-α required for a sustained response and consequently its side effects. The most rewarding of all these combinations with IFN was that of ribavirin which in large randomised controlled trials1 proved to be far superior to IFN-α monotherapy of six and 12 months duration. In fact, the overall sustained response rate achieved with combination therapy in these two large trials reached 33% and 41% over six and 12 months of treatment, respectively, compared with 6% and 16% achieved with IFN monotherapy over the same time periods. The increase in sustained response was even more dramatic in patients with HCV genotypes 1 and 4–6 than in patients with the more easily eradicable genotypes 2 and 3. The sustained virological response doubled in the latter from 30% to 65% but tripled in the former from 9% to 30%. However, despite these exciting results most treated patients with chronic hepatitis C fail to attain a sustained virological response. This is particularly true for infection with genotypes 1 and 4, with 70% of infected patients failing to respond even to a 12 month combination therapy regimen. It is therefore conceivable that further efforts are needed in order to improve the efficacy of combination therapy. To this end different types of interferon-like compounds, new formulations of IFN-α with attachment of polyethylene glycol, use of daily IFN dosing, higher doses and induction regimens aiming at rapid decrease of HCV replication are being tried. Combination of all these IFN...
Interferon α and NSAIDs in hepatitis C

directed efforts with adjuvant drugs is also under evaluation.

NSAIDs combined with IFN has been tried in naive and non-responsive patients with chronic hepatitis C with little or no success.5,6 However, existing evidence suggests that NSAIDs with their cyclooxygenase inhibiting activity possess an IFN potentiating effect by enhancement of 2,5'-oligosynthetase activity.7 Negative results from the few early clinical studies of combined treatment with NSAIDs and IFN and case reports of NSAID induced hepatotoxicity8 in chronic hepatitis have been discouraging.

In this issue (see page 427), Muñoz et al report on a prospective randomised controlled trial evaluating the efficacy of combined treatment with IFN and ketoprofen versus IFN alone. The authors have managed to study a fairly large, for a single centre, number of patients with well defined inclusion criteria of chronic hepatitis C. The patients were allocated into three treatment groups receiving either: (1) IFN alone at 3 million units three times weekly for six months; (2) IFN combined with a small dose of the NSAID ketoprofen, 200 mg given three hours before the thrice weekly injections of IFN; or (3) IFN combined with daily ketoprofen at a dose of 200 mg given twice a day. Ketoprofen was administered as a slow-release form. Both groups also received ranitidine (150 mg/day) prophylactically. Ketoprofen was selected over other NSAIDs because it has been used in the past in patients with hepatitis without serious side effects, it has a high bioavailability, its pharmacokinetics are not influenced by cirrhosis, and it inhibits both the cyclooxygenase and lipoxygenase inflammatory pathways. The patients were monitored carefully clinically, biochemically, and virologically. HCV RNA was measured by RT-PCR and negative cases were retested by a nested PCR with a detection limit of 1000 copies/ml. Both doses of ketoprofen were well tolerated with similar side effects to the IFN monotherapy arm except perhaps for a greater drop in haemoglobin concentrations. The virological response rate at the end of follow up (sustained virological response) was significantly higher in the daily ketoprofen group (26%) compared with the small thrice weekly dose and to the IFN monotherapy groups (0% and 10%, respectively) and 9% in the two groups combined, p=0.009). The response rate of the few cirrhotic patients in group 3 was similar. Moreover the 'flu-like side effects of IFN-α were less frequent in the ketoprofen treated patients.

This is the first time that ketoprofen has been given to patients with chronic hepatitis C twice daily in a slow-release form, similar to the doses recommended for the treatment of osteoarthritits and rheumatoid arthritis. This probably explains the major difference in the efficacy of this regimen compared with the lower dose scheme and with the negative results of a similar study using a different ketoprofen dosage.4 Independently of these encouraging results, there are some points of concern in this study that have also been recognised by the authors. Firstly, as expected, the number of patients studied, although large for a single centre study (n=70), is still very small for a three arm study and both type 1 and type 2 errors are possible. In fact the difference between groups 1 and 3 is not statistically significant. Furthermore, because of the small number of patients an intent-to-treat analysis weakens the differences significantly. Secondly, it is difficult to explain the very large discrepancy between the rate of biochemical and virological response at the end of treatment observed in all three groups. Even in the IFN monotherapy group the biochemical response rate was 52% and the virological rate 10%. Thirdly, it is not conceivable how the group on the small ketoprofen dose could have had a response rate below that of the IFN monotherapy group. Fourthly, again as expected, owing to the small number of patients studied, the effect on the efficacy of therapy of several baseline variables such as viral load, HCV genotypes, and extent of liver fibrosis was not evaluated. These and other emerging questions should be answered in future trials of large numbers of patients.

Regardless of the study’s limitations, this report of a significant increase in the sustained virological response rate achieved at a small additional financial cost should herald new clinical trials of combined treatment with ketoprofen and IFN. The postulate that NSAIDs may potentiate the action of IFN has received timely independent support from a recent study in hepatoma cell lines, indicating that blockade of arachidonic acid metabolism by NSAIDs, activates a signalling pathway potentiating IFN-α dependent gene activation.9 Clearly the last word on the combination of NSAIDs with IFN has still to be written.

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