The three autosomal dominant inherited polyposis syndromes, familial adenomatous polyposis, juvenile polyposis, and Peutz-Jeghers polyposis predispose to colorectal cancer as does hereditary non-polyposis colorectal cancer syndrome. Uncovering the genetic background of these four cancer traits provides the possibility for genetic testing of the family members of an affected patient. Before testing identification of the underlying family specific pathogenic mutation is mandatory. This is possible in about 60% to 95% of families. Endoscopic surveillance can be safely discontinued in mutation negative family members and surveillance or prophylactic surgery can be targeted to mutation positive members alone. Testing requires genetic counselling and written informed consent to prevent misunderstanding and to minimise untoward effects such as anxiety. Permanent surveillance and adequate prophylactic treatment for all mutation positive subjects and families is best ensured in national or regional polyposis registries with the capacity to take care of long term follow up from generation to generation.

There are three main polyposis syndromes that share the features of autosomal dominant mode of inheritance and an increased risk of gastrointestinal cancer, especially colorectal cancer: (1) familial adenomatous polyposis (FAP), (2) juvenile polyposis (JP), and (3) Peutz-Jeghers polyposis (P-JP). A fourth closely related autosomal dominant trait is hereditary non-polyposis colorectal cancer syndrome (HNPPC). It has several clinical and genetic aspects in common with the polyposis syndromes, but also some important differences, considering the clinical management and cancer prevention measures available on the basis of the hereditary nature of these cancer predisposition conditions.

The molecular genetic background of the three polyposis syndromes and HNPPC were largely uncovered during the past 15 years. The first step was the localisation of the adenomatous polyposis coli (APC) gene on chromosome 5q followed by the characterisation of the gene in 1991. Two years later two genes associated with the HNPPC susceptibility were localised and later identified and followed by discovery of further DNA mismatch repair genes with proved or suspected association with the HNPPC syndrome. Lastly, the genes responsible for a major part of families with JP and P-JP were discovered (table 1).

The discovery of the genetic causation of the polyposis syndromes and HNPCC offers the possibility for predictive genetic testing of the descendants of the affected family members who have the theoretical risk of 50% of carrying the pathogenic mutation. The test can relieve the unaffected half of the family members from the cancer threat, endoscopic surveillance, and anxiety involved. On the other hand, the mutation positive family members can be identified with certainty, and appropriate clinical surveillance and prophylactic treatment can be offered for them to ensure optimal cancer prevention. The strategies of prevention and their efficacy vary in the four conditions, and many clinical and genetic problems still await for adequate solution. This review gives some guidelines for the management of polyposis and HNPCC families with the advent of predictive genetic testing.

**Clinical features and the cancer risk in polyposis syndromes and HNPCC**

**Familial adenomatous polyposis MIM (Mendelian inheritance in man) no 175100, FAP and Gardner syndrome; 276300, Turcot syndrome**

FAP is characterised by multiple adenomatous polyps that tend to progress to adenocarcinoma. In classic FAP the number of polyps is 100 at least but an attenuated form has been distinguished with a fewer number of adenomas. Turcot syndrome is a variant associated with a brain tumour, medulloblastoma. The estimated incidence of FAP varies around 1 per 10 000 newborns or from 1 to 2 per 1 000 000 per annum. Between 30% to 50% of new patients are solitary cases probably representing new mutations of the APC gene. Mutations of this gene located in the long arm of chromosome 5 (5q 21–22) cause the disease.

The clinical diagnosis of FAP requires demonstration of 100 colorectal adenomas. Histological examination of several adenomas is necessary. In the context of a definite family history the detection of fewer adenomas is sufficient as well as in the case of an attenuated disease form. Final diagnosis is achieved by discovery of a mutated APC gene but the detection rate has been about 60% to 95% of all FAP families. Well known extracolonic manifestations of the disease such as epidermoid cysts, osteomas of the jaws, desmoid...
starting at the age of 12 to 15 years at intervals of a few years. This requires examination of all children of an affected parent for about 20 years. P-JP is characterised by mucocutaneous melanin pigmentation and hamartomatous intestinal polyposis. The polyps preferentially affect the small intestine but they occur in the stomach and large intestine as well. Peutz-Jeghers syndrome is caused by mutations of the LKB1 gene on the short arm of chromosome 19, 19p 13.3. The incidence of P-JP is of the same range as observed in JP probably about 1 per 100 000 newborns. The polyps are originally hamartomas but epithelial dysplasia has been demonstrated to develop rarely and adenocarcinoma may occur in contiguity with the Peutz-Jeghers polyp. Consequently, there is an 80-fold to 500-fold excess of gastrointestinal cancer in the P-JP syndrome.

Since the recognition of the cancer risk involved in P-JP, endoscopic surveillance with polypectomies has been recommended for the affected patients by two year intervals using both upper gastrointestinal and colonoscopic approaches ("top and tail endoscopy"). Unfortunately, the small bowel remains mainly beyond the reach of endoscopies. Therefore, at operations needed because of bleeding episodes or small bowel obstruction due to intussusception the entire small intestine should be cleared up from the polyps using intraoperative endoscopy and enterotomies. The problem of an excess of many non-gastrointestinal cancers such as breast, endometrial, ovarian, or lung cancer with risk ratios from 15 to 30 is difficult to cover with any simple preventive programme.

### Hereditary non-polyposis colorectal cancer syndrome (HNPCC)

**MIM no 120435–6**

HNPCC or Lynch syndrome is an autosomal dominant cancer predisposition syndrome without clear clinical signs preceding cancer except for solitary colorectal adenomas. The predominant tumours are colorectal and endometrial cancer. Several other cancer types are, however, also occurring in excess such as carcinoma of the stomach, ovary, ureter and renal pelvis, bile ducts, small ducts, duodenum, and brain tumours. Mutations of three DNA mismatch repair genes have been associated with the HNPCC trait, MSH2, MLH1, and MSH6. The role of other mismatch repair genes (for example, PMS1, PMS2, and MSH3) is still under evaluation. HNPCC is more common than the polyposis syndromes and it explains up to 3% of all cases of colorectal cancer.

Colorectal cancer develops at young age (mean 45 years), located predominantly in the proximal colon, and synchronous or metachronous second tumours occur in more than a third of patients. The DNA content of the tumours is diploid and microsatellite instability is a characteristic feature. In histological examination tumours exhibit high mucin content, Crohn’s disease-like inflammatory reaction around, and the differentiation grade is commonly poor. The clinical or pathological features, however, are not specific and do not alone allow a diagnosis of HNPCC to be done. The family history has a central role in the identification. According to the Amsterdam criteria the following conditions should be fulfilled: (1) at least three patients with colorectal cancer of whom one is a first degree relative of the other two, (2) affected family members in two generations, at least, (3) within the same range as observed in JP probably about 1 per 100 000 newborns. The polyps are originally hamartomas but epithelial dysplasia has been demonstrated to develop rarely and adenocarcinoma may occur in contiguity with the Peutz-Jeghers polyp. Consequently, there is an 80-fold to 500-fold excess of gastrointestinal cancer in the P-JP syndrome.

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### Table 1 Genetic background of polyposis syndromes and HNPCC

<table>
<thead>
<tr>
<th>Disease</th>
<th>Chromosome</th>
<th>Gene(s)</th>
<th>Discovered (ref)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAP</td>
<td>5q21</td>
<td>APC</td>
<td>Bodmer et al 1987 (1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Groden et al 1991 (2)</td>
</tr>
<tr>
<td>JP</td>
<td>18q 21</td>
<td>SMAD4/DPDC4</td>
<td>Howe et al 1998 (8)</td>
</tr>
<tr>
<td></td>
<td>10q 21.22</td>
<td>BMPR1A/ALK3</td>
<td>Howe et al 2001 (9)</td>
</tr>
<tr>
<td>P-JP</td>
<td>19p 13.3</td>
<td>LKB1</td>
<td>Hemminki et al 1998 (7)</td>
</tr>
<tr>
<td>HNPCC</td>
<td>2p 21.23</td>
<td>MSH2</td>
<td>Peltonäkki et al 1993 (3)</td>
</tr>
<tr>
<td></td>
<td>3p 21</td>
<td>MLH1</td>
<td>Lindblom et al 1993 (4)</td>
</tr>
<tr>
<td></td>
<td>2p 16</td>
<td>MSH6</td>
<td>Miyaki et al 1997 (6)</td>
</tr>
</tbody>
</table>

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respectively. The corresponding lifetime risks for other cancer types remain between 13% and 2%.2,27 Cancer prevention in HNPCC consists of repeated colonoscopy with polypectomies at two to three year intervals and of endometrial suction biopsies possibly combined with endovaginal ultrasonography.35 Prophylactic surgery (colectomy, hysterectomy) remains an alternative in selected cases. Prevention and early detection of other tumour types is problematic because of their relative rarity and when there are no suitable screening methods.

**PREDICTIVE GENETIC TESTING**

The polyposis syndromes can be identified on the basis of their typical clinical features—that is, by demonstration of the intestinal polyps in endoscopy and their histological typing after polypectomies. This is the standard method in symptomatic patients who usually are the probands of a new family or solitary cases. Endoscopic screening of all first degree family members is consequently indicated to achieve early diagnosis before the development of cancer. In this evaluation the extracolonic manifestations of the disease, such as melanin pigmentation in P-JP or retinal pigmentation, epidermoid cysts of the jaw in FAP, may offer additional clues.

Clinical screening, however, is unreliable depending on the age of the subject, severity of the polyposis associated with the particular mutation of the family, and the syndrome in question. There is variation in the phenotypic expression even between members of the same family affected with FAP.28 Some FAP families also present with late onset of adenomas and mild phenotype, attenuated FAP, often associated with APC gene mutations localised either in the 5' or 3' part of the gene or in exon 9.37-38 Therefore, negative endoscopic findings at a certain age cannot exclude the possibility of later disease expression. The issue is even more difficult in HNPCC where there are no clear clinical indicators of the disease at all. Thus, knowledge of the pathogenic mutation can greatly help the organisation of family surveillance in polyposis syndromes. In HNPCC this knowledge is almost irreplaceable. Unfortunately, despite typical criteria of a specific polyposis syndrome or HNPCC the underlying mutation remains undetected in some 20% to 40% of families, as is shown in the data of the Finnish HNPCC Registry (table 2).

Table 2. Detection rate of pathogenic mutations in the Finnish Polyposis and HNPCC registry (December, 2001)

<table>
<thead>
<tr>
<th>Test</th>
<th>FAP (%)</th>
<th>JP (%)</th>
<th>P-JP (%)</th>
<th>HNPCC (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mutation detected</td>
<td>57 (77)</td>
<td>6 (75)</td>
<td>7 (54)</td>
<td>92 (77)</td>
</tr>
<tr>
<td>No mutation detected</td>
<td>17</td>
<td>2</td>
<td>6</td>
<td>28</td>
</tr>
<tr>
<td>Net examined</td>
<td>31</td>
<td>4</td>
<td>2</td>
<td>20</td>
</tr>
<tr>
<td>Families in total</td>
<td>105</td>
<td>12</td>
<td>15</td>
<td>140</td>
</tr>
</tbody>
</table>

After identification of the pathogenic mutation the predictive testing of the family members has high accuracy, practically 100%. Mutation positive subjects can accordingly be advised to appropriate surveillance or prophylactic treatment, while the follow up of the mutation negative subjects can be discontinued. Omission of endoscopic surveillance in half of the descendants accordingly cuts the costs of family screening. Even though the cost of one mutation test of a known mutation (£50) is higher than a single sigmoidoscopy (£120) or colonoscopy (£320), the total costs are reduced when repeated endoscopies of mutation negative subjects become unnecessary. On the other hand, the cost of the search for a new germline mutation is considerably higher, about £2500. After all, the economic aspects of the genetic compared with endoscopic screening play a secondary part considering the far greater accuracy of genetic screening. It should be noted that the prices presented above reflect the situation in year 2001 at the Helsinki University Hospital. The prices vary with time and may be quite different elsewhere.

Mutation data have also been suggested to serve as a guide in the planning of prophylactic surgery or clinical surveillance. The idea comes from the observations about genotype phenotype correlation. In a Dutch series of FAP patients treated with colectomy and ileorectal anastomosis the patients with mutations 3' to codon 1250 had a higher rate of rectal cancer than those with mutations 5' to this codon.41 It was thought that the choice of the method of surgery, colectomy with ileorectal anastomosis versus proctocolectomy with ileal pouch-anal anastomosis, should be determined on the basis of the APC gene mutation type. The genotype phenotype correlations, however, seem more complicated. For example, mild phenotype occurs in patients with mutation at each end of the APC gene or in exon 9,37,38 and other considerations such as the desmoid tumour tendency, partly determined by the mutation site, might also influence on decisions.42 At present, mutation analysis has limited value for surgical decisions in FAP and the clinical features of individual patients and families have a more central role.

**ETHICAL AND LEGAL ASPECTS OF GENETIC TESTING**

Recognition of a hereditary cancer syndrome in a family provokes anxiety in family members. The possibility of genetic testing for the diagnosis of mutation status of the relatives may cause ambivalent feelings. A mutation negative test result naturally is reassuring but the finding of a pathogenic mutation may increase anxiety, even though it enables appropriate surveillance and treatment. In a recent study on predictive testing for at risk members of HNPCC families the anxiety scores expectedly increased in mutation carriers and decreased in mutation negative individuals, but the difference disappeared within one month.43 This result was obtained in the context of an well organised genetic counselling programme with a follow up for one year. Despite careful individual counselling the mutation positive subjects tended to misunderstand the significance of the test result and the consequent cancer risk quite often.44 The necessity of individual counselling and obtaining an informed consent for testing is clear because of possible untoward effects of testing even besides anxiety. There may also be effects on eligibility for insurance or employment even though there is common agreement that genetic testing should not cause discrimination and the test result should be kept entirely confidential. The practices in different countries considering insurance companies and employers may be different than in the European Union.

The appropriate age for testing depends on the average expression of the disease, which occurs already during teenagers in FAP and possibly earlier in P-JP and JP. In HNPCC the cancer incidence begins to rise significantly first after the age of 20 to 25 years, when the surveillance programme should start. At that age there is usually no problem in obtaining consent after genetic counselling. However, in FAP and other polyposis syndromes the information should be given to both
children and their parents. In this context there may occur instances where an affected person does not permit information about the inherited condition to be distributed to other relatives or even to the children thus preventing the situation of the medically indicated cancer screening programme. Such a situation causes a serious ethical problem where repeated discussions may eventually give a solution. The finding of a mutation positive result in polyposis or HNPCC causes the worry about organisation of permanent surveillance and proper prophylactic treatment throughout the rest of the life including screening and testing of all family members involved. An individual clinician faces many practical problems if the family is large and scattered around the country and even abroad. In such instance polyposis registries give the best guarantee of continued care of surveillance, genetic counseling, identification, genetic counselling and testing, and even research cooperation. In maintenance of updated patient follow up data the registries need at present informed consent of the family members as has been provided by a European Union directive, the Personal Data Act. In many registries including the Finnish Polyposis and HNPCC Registry achieving a permanent position and independent financial status still awaits final solution, a problem that will hopefully be managed in near future considering the established excellent results in cancer prevention.

**FUTURE DEVELOPMENT**

Knowledge about the genetic causation of the polyposis syndromes or HNPCC has thus far given not much hope for cure of the underlying genetic abnormality by gene therapy, for example, given the very short half time of the cells in the main target organ, intestinal epithelium. The only really curative treatment is induction of abortion after intrauterine molecular genetic diagnosis, a procedure that has not been taken in general use, thus far, despite its applicability. Increasing knowledge about the exact function of the genes involved may help in the development of targeted medical treatments, which normalise the disturbed function. Such development could at best obviate the need for prophylactic surgery or, at least, help in prevention of the poorly treatable extracolonic complications such as desmoid tumours or duodenal adenomas and cancer in FAP.

**REFERENCES**