Guidelines for the management of hereditary colorectal cancer from the British Society of Gastroenterology (BSG)/Association of Coloproctology of Great Britain and Ireland (ACPGBI)/United Kingdom Cancer Genetics Group (UKCGG)

Kevin J Monahan, Nicola Bradshaw, Sunil Dolwani, Bianca Desouza, Malcolm G Dunlop, James E East, Mohammad Ilyas, Asha Kaur, Fiona Laloo, Andrew Latchford, Matthew D Rutter, Ian Tomlinson, Huw J W Thomas, James Hill

ABSTRACT
Heritable factors account for approximately 35% of colorectal cancer (CRC) risk, and almost 30% of the population in the UK have a family history of CRC. The quantification of an individual’s lifetime risk of gastrointestinal cancer may incorporate clinical and molecular data, and depends on accurate phenotypic assessment and genetic diagnosis. In turn this may facilitate targeted risk-reducing interventions, including endoscopic surveillance, preventative surgery and chemoprophylaxis, which provide opportunities for cancer prevention. This guideline is an update from the 2010 British Society of Gastroenterology (BSG)/Association of Coloproctology of Great Britain and Ireland (ACPGBI) guidelines for colorectal screening and surveillance in moderate and high-risk groups; however, this guideline is concerned specifically with people who have increased lifetime risk of CRC due to hereditary factors, including those with Lynch syndrome, polyposis or a family history of CRC. On this occasion we invited the UK Cancer Genetics Group (UKCGG), a subgroup within the British Society of Genetic Medicine (BSGM), as a partner to BSG and ACPGBI in the multidisciplinary guideline development process. We also invited external review through the Delphi process by members of the public as well as the steering committees of the European Hereditary Tumour Group (EHTG) and the European Society of Gastrointestinal Endoscopy (ESGE). A systematic review of 10 189 publications was undertaken to develop 67 evidence and expert opinion-based recommendations for the management of hereditary CRC risk. Ten research recommendations are also prioritised to inform clinical management of people at hereditary CRC risk.

OBJECTIVE
To provide a clear strategy for the management of people at hereditary risk of colorectal cancer (CRC), which includes diagnosis, endoscopic management, prevention and surgical care.

AIMS AND METHODS
An estimated 35% of CRC is due to heritable factors, with approximately 29% of the UK population having a family history of a first-degree relative (FDR) or second degree relative (SDR) with CRC. While highly penetrant syndromes such as Lynch syndrome (LS), familial adenomatous polyposis (FAP) and other polyposis syndromes account for only 5–10% of all CRC diagnoses, advances in genetic diagnosis, improvements in endoscopic surgical control, and medical and lifestyle interventions provide opportunities for CRC prevention and effective treatment in susceptible individuals.

The purpose of this guideline is to provide an evidence-based framework for the optimal management of hereditary CRC for clinicians involved in their management, including gastroenterologists, nurse practitioners, physicians, colorectal surgeons, clinical geneticists, genetic counsellors and pathologists. This guideline was commissioned by the Clinical Services and Standards Committee (CSSC) of the British Society of Gastroenterology (BSG), via the colorectal section, and a guideline chair selected. It is an update of the previous iteration of the BSG/Association of Coloproctology of Great Britain and Ireland (ACPGBI) guideline published in 2010 and developed in accordance with the BSG National Institute for Health and Care Excellence (NICE)-compliant guideline process.

The Guideline Development Group (GDG), which included gastroenterologists from the BSG, clinical geneticists from United Kingdom Cancer Genetics Group (UKCGG), colorectal surgeons from the ACPGBI, a pathologist, a genetic counsellor and a patient representative, was selected to ensure wide-ranging expertise across all relevant disciplines. Members of the GDG, and participants in the eDelphi process, completed a Declaration of Conflict of Interests (COI) form which was reviewed and vetted by the BSG.

A scoping meeting was held on 13 October 2017, and in advance of this meeting the GDG was asked to develop key priorities and questions.

The GDG determined that the primary measure of effectiveness of any intervention was a reduction of the lifetime risk of CRC, and the following secondary outcome measures:
i. Reduction in the incidence of advanced adenomas at colonoscopy
ii. Prevention of CRC
iii. Reduced morbidity related to CRC, or morbidity secondary to complications of surveillance and treatment
iv. Improved identification of hereditary CRC syndromes
v. Improved pathways from diagnosis to treatment in susceptible populations.

We sought a consistent approach in our assessment of the relative effectiveness of interventions. In principle we agreed that surveillance should only be offered to individuals who remain at higher risk of developing CRC than the general population. As CRC risk is not always clearly defined, as a surrogate we accepted that advanced adenoma yield on surveillance should be approximately double that in susceptible populations compared with the average risk population.

A relative threshold for genetic testing was agreed for people with a 10% or greater probability of having a germline pathogenic variant in a cancer susceptibility gene in accordance with previous UK guidelines. However the GDG agreed that the arbitrary nature of this threshold meant that it could be modified in cases where objective risk assessment was difficult to attain, and clinicians had sufficient clinical suspicion of risk.

Key questions we sought to cover included the following:
1. Which aspects of the previous guidelines require updating?
2. What is the lifetime CRC risk and optimal surveillance for those with a family history of CRC (where LS and polyposis syndromes have been excluded)?
3. What is the diagnostic yield of genetic testing and/or surveillance for high-risk populations?
4. What is the optimal gastrointestinal (GI) surveillance for patients at hereditary risk GI cancer?
5. What is the impact of high-quality endoscopy in patients with known or suspected hereditary cancer syndromes?
6. Should we develop gene- or gender-specific guidelines for surveillance?
7. What is the optimal diagnostic assessment and surveillance interval for ‘Lynch-like’ syndrome patients?
8. How can we improve recognition, diagnosis and treatment of patients at hereditary risk of CRC?
9. Which diagnostic genetic tests should we offer serrated polyposis syndrome (SPS), multiple colorectal adenoma (MCRA) and early onset CRC (EOCRC) patients (if any)?
10. When should colonoscopic surveillance for familial risk patients stop, because it is no longer necessary, or because the patient should be referred for surgery?
11. Which are the optimal surgical approaches in patients with hereditary CRC syndromes?
12. What is the evidence for chemoprophylaxis in patients who are at hereditary risk of CRC?
13. What is the evidence for the effect of lifestyle modification on hereditary risk of CRC?
14. What information do we need to share with our patients at inherited risk of GI cancer?

Twenty-three PICOs (Patients, Interventions, Controls and Outcomes) were developed which considered these questions. The Appraisal of Guidelines for Research and Evaluation (AGREE II) instrument provided a methodological framework.

A literature search was commissioned externally, with search strategies agreed, and was performed by the Yorkshire Healthcare Consortium, which returned 10189 publications. Returned abstracts were reviewed for relevance. Additional references were obtained by cross-referencing and by recommendation from the GDG. Relevant published national and international guidelines were also scrutinised. After each round of Delphi, and before the guideline was finalised, the search was repeated and any important studies published since the initial evidence search were incorporated.

A modified electronic Delphi process was used to develop and refine statements. Initial draft statements formulated by the writing committee were reviewed by the GDG to allow for modification and to identify additional references. After a preliminary discussion, formal anonymous voting rounds were undertaken using SurveyMonkey. Each statement was scored by each member of the GDG using a five-point scale. We also invited key national and international opinion leaders from the UKCGG steering group, ACPGBI, BSG, the European Hereditary Tumour Group (EHTG) and the European Society of Gastrointestinal Endoscopy (ESGE) to participate in the modified Delphi process. We included additional patient and public involvement in the Delphi process by inviting participants through the national charities Bowel Cancer UK and Lynch Syndrome UK. Consensus required at least 80% agreement, and consensus of over 70% was accepted if the GDG felt a statement was required for clinical practice. Where consensus was not reached, feedback from the GDG members was disseminated after each round to allow members to reconsider their original position. Where appropriate, revisions to statements were made and a further voting round was undertaken in second and third rounds. A final (fourth) round of voting for statements where consensus had not been reached for 11 statements was performed within the GDG only.

Surveillance and molecular testing recommendations are summarised in table 1 and table 2 respectively. The GDG also developed 10 research recommendations (online supplementary file 1) which were prioritised by electronic voting.

The GRADE (Grading of Recommendations, Assessment, Development and Evaluations) tool was used to evaluate the strength of evidence and the strength of recommendations made (see executive summary). The GRADE system specifically separates the strength of evidence from the strength of a recommendation. While the strength of a recommendation may often reflect the evidence base, the GRADE system allows for occasions when this is not the case—for example, where it seems good sense to make a recommendation despite the absence of high-quality scientific evidence such as a large randomised controlled trial (RCT).

**EXECUTIVE SUMMARY OF KEY RECOMMENDATIONS**

**Service provision, communication and management principles**
- We recommend that the moderate risk category of family history of CRC (FHCC) is the minimum threshold for referral from primary care (GRADE of evidence: very low; Strength of recommendation: strong)
- We recommend that individuals with a FHCC, which meets this referral criteria, be referred to a specialist familial CRC clinic in secondary or tertiary care (GRADE of evidence: low; Strength of recommendation: weak)
- We recommend that patients should be referred to a specialist service which includes access to constitutional genetic testing in the presence of either deficient mismatch repair (MMR) (with no evidence of MLH1 promoter methylation or BRAF V600E), or polyposis. (GRADE of evidence: low; Strength of recommendation: strong)
- There are insufficient clinical data to develop specific guidance for patients with very rare conditions such as polymerase proofreading associated polyposis (PPAP), or NTHL1-associated polyposis (NAP); therefore, we suggest patients with these syndromes should be referred to

---


---

Put the text into a Latex file.
Table 1  Summary of surveillance recommendations

<table>
<thead>
<tr>
<th>Indication for surveillance</th>
<th>Category</th>
<th>Modality</th>
<th>Age to commence (years)</th>
<th>Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>Family history of CRC</td>
<td>Average risk</td>
<td>National screening</td>
<td>As defined by national screening</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Moderate risk</td>
<td>Colonoscopy</td>
<td>55</td>
<td>Post-polypectomy guidelines</td>
</tr>
<tr>
<td></td>
<td>High risk*</td>
<td>Colonoscopy</td>
<td>40</td>
<td>5 yearly until age 75 years</td>
</tr>
<tr>
<td>Lynch syndrome</td>
<td>MMR gene pathogenic variant carriers</td>
<td>Colonoscopy</td>
<td>25</td>
<td>2 yearly until age 75 years</td>
</tr>
<tr>
<td></td>
<td>*MLH1 and MSH2 gene carriers</td>
<td>Colonoscopy</td>
<td>35</td>
<td>2 yearly until age 75 years</td>
</tr>
<tr>
<td></td>
<td>Stomach, small bowel and pancreas</td>
<td>Not indicated outside a clinical trial</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lynch-like syndrome</td>
<td>Individuals with deficient MMR tumours without hypermethylation/BRAF pathogenic variant and no pathogenic constitutional pathogenic variant in MMR genes, and no evidence of biallelic somatic MMR gene inactivation (and their unaffected FDRs)</td>
<td>Colonoscopy</td>
<td>25</td>
<td>2 yearly until age 75 years</td>
</tr>
<tr>
<td>Serrated polyposis syndrome</td>
<td>Affected individuals (WHO 2019)</td>
<td>Colonoscopy</td>
<td>From age of diagnosis</td>
<td>1–2 yearly until age 75 years</td>
</tr>
<tr>
<td>Multiple colorectal adenomas (MCRAs)</td>
<td>10 or more adenomas without constitutive pathogenic variants in APC or MUTYH</td>
<td>Colonoscopy</td>
<td>From age of diagnosis</td>
<td>1–2 yearly until age 75 years</td>
</tr>
<tr>
<td>Familial adenomatous polyposis (FAP)</td>
<td>APC pathogenic variant carriers</td>
<td>Colonoscopy</td>
<td>12 to 14</td>
<td>1–3 yearly depending on phenotype</td>
</tr>
<tr>
<td></td>
<td>Gastroscopy and duodenoscopy</td>
<td>25</td>
<td>As per Spigelman classification</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sigmoidoscopy/pouchoscopy</td>
<td>From time of colectomy</td>
<td>1–3 yearly depending on phenotype</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Individuals with an FDR with a clinical diagnosis of FAP (ie, “at-risk”) and in whom a constitutional pathogenic variant has not been identified</td>
<td>Colonoscopy</td>
<td>12 to 14</td>
<td>5 yearly until national screening age</td>
</tr>
<tr>
<td></td>
<td>Gastroscopy and duodenoscopy</td>
<td>Commence only if clinical diagnosis made of colorectal polyposis phenotype</td>
<td>As per Spigelman classification</td>
<td></td>
</tr>
<tr>
<td>MUYTH-associated polyposis (MAP)</td>
<td>MUYTH gene pathogenic variant carriers</td>
<td>Colonoscopy</td>
<td>18 to 20 years</td>
<td>Annual</td>
</tr>
<tr>
<td></td>
<td>Gastroscopy and duodenoscopy</td>
<td>35</td>
<td>As per Spigelman classification</td>
<td></td>
</tr>
<tr>
<td>Peutz-Jeghers syndrome (PJS)</td>
<td>STK11 gene pathogenic variant carriers</td>
<td>Upper gastrointestinal endoscopy, colonoscopy and video capsule endoscopy</td>
<td>8</td>
<td>see main text</td>
</tr>
<tr>
<td>Juvenile polyposis syndrome (JPS)</td>
<td>SMAD4 and BMP1R1A pathogenic variant carriers</td>
<td>Colonoscopy</td>
<td>15</td>
<td>1–3 yearly depending on phenotype</td>
</tr>
<tr>
<td></td>
<td>SMAD4 pathogenic variant carriers</td>
<td>Gastroscopy and duodenoscopy</td>
<td>18</td>
<td>1–3 yearly depending on phenotype</td>
</tr>
<tr>
<td></td>
<td>BMP1R1A pathogenic variant carriers</td>
<td>Gastroscopy and duodenoscopy</td>
<td>25</td>
<td>1–3 yearly depending on phenotype</td>
</tr>
</tbody>
</table>

*Amsterdam criteria families where MMR testing is not possible may be offered surveillance as per Lynch syndrome families and/or additional constitutional testing. CRC, colorectal cancer; FDR, first degree relative; MMR, mismatch repair.

Guidelines

► We recommend that hospitals which diagnose or manage patients at hereditary CRC risk should ensure clinical pathways to facilitate their care, and processes to monitor the quality of the service. (GRADE of evidence: low; Strength of recommendation: weak)
► We recommend that individuals at increased familial CRC risk receive specialist knowledge and are aware of patient/support organisations and discussion with regard to lifestyle and participation in research projects. (GRADE of evidence: very low; Strength of recommendation: strong)

Family history of CRC (FHCC)
► We recommend that for all patients referred from primary care for assessment for a FHCC, MMR status should be assessed in tumour tissue from a close affected family member. (GRADE of evidence: moderate; Strength of recommendation: strong)
► We recommend that a reported family history of polyposis should be verified by review of histopathology and/or endoscopy reports which confirm the presence of a minimum of 10 adenomas or serrated lesions in a FDR. (GRADE of evidence: low; Strength of recommendation: strong)
► We recommend that patients with a moderate familial CRC risk should have a one-off colonoscopy at age 55 years.
We recommend that all surveillance colonoscopies are performed by endoscopists who consistently achieve BSG colonoscopy KPI (key performance indicators) minimum standards, specifically caecal intubation rate, adenoma/polyp detection rate and comfort score. (GRADE of evidence: low; Strength of recommendation: strong)

► We suggest high-quality, high-definition white light endoscopy as the preferred modality for colonoscopy surveillance. Chromoendoscopy (virtual or dye-based) does not offer a clear advantage over high definition white light examination for colonoscopic surveillance, apart from in the context of determining the multiple polyp phenotype. (GRADE of evidence: moderate; Strength of recommendation: weak)

► We suggest a repeat colonoscopy performed by an expert endoscopist is indicated in the event of a previously failed colonoscopy, with efforts made to both improve patient experience and to ensure procedure completion, given the advantages of colonoscopic surveillance. If colonoscopy is inadequate or if the examination is incomplete then a repeat colonoscopy should be performed at a 2 yearly interval for all LS patients. (GRADE of evidence: moderate; Strength of recommendation: strong)

Prevention and lifestyle modification in familial CRC
► We recommend that individuals with LS should be advised that regular use of daily aspirin reduces CRC risk. (GRADE of evidence: moderate; Strength of recommendation: strong)

► We suggest that people with LS should be offered research opportunities to take aspirin daily at different dosages. If they decline research participation they may be advised on their choices regarding dose of aspirin, risks and benefits of long-term aspirin use and ensure their medical practitioner is aware of their intake. (GRADE of evidence: low; Strength of recommendation: weak)

► There is insufficient evidence of the benefit of chemoprophylaxis in polyposis syndromes. (GRADE of evidence: moderate; Strength of recommendation: strong)

► We suggest that individuals at increased familial risk of CRC should be strongly encouraged not to smoke, to maintain a normal body mass index (BMI), to moderate their consumption of red and processed meat, and to exercise regularly. (GRADE of evidence: low; Strength of recommendation: weak)

Quality and advanced endoscopic imaging in colonoscopic surveillance
► We recommend that colonoscopy is the gold standard diagnostic and preventative method of surveillance for people with hereditary risk of CRC. (GRADE of evidence: moderate; Strength of recommendation: strong)

► We recommend that all surveillance colonoscopies are performed by endoscopists who consistently achieve BSG colonoscopy KPI (key performance indicators) minimum standards, specifically caecal intubation rate, adenoma/polyp detection rate and comfort score. (GRADE of evidence: low; Strength of recommendation: strong)

► We recommend that for all people when first diagnosed with CRC, testing using immunohistochemistry (IHC) for MMR proteins or microsatellite instability is used to identify tumours with deficient DNA MMR, and to guide further sequential testing for LS. (GRADE of evidence: moderate; Strength of recommendation: strong)

► We recommend that colonoscopic surveillance should be performed at a 2 yearly interval for all LS patients. (GRADE of evidence: moderate; Strength of recommendation: strong)

► We recommend that age of onset of surveillance colonoscopy should be stratified according to the LS-associated gene. We recommend colonoscopy from age 25 years for MLH1 and MSH2 mutation carriers and 35 years for MSH6 and PMS2 mutation carriers. There are insufficient data to support stratifying age of onset of surveillance by gender.

Table 2  Molecular testing strategies in hereditary colorectal cancer (CRC)

<table>
<thead>
<tr>
<th>Risk category</th>
<th>Somatic or constitutive testing</th>
<th>Eligibility</th>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Family history of CRC</td>
<td>Somatic</td>
<td>Moderate-risk or high-risk family history</td>
<td>dMMR/pMMR</td>
</tr>
<tr>
<td></td>
<td>Constitutive</td>
<td>Amsterdam criteria families where MMR testing is not possible</td>
<td>Panel testing of affected individuals or unaffected testing</td>
</tr>
<tr>
<td>CRC</td>
<td>Somatic</td>
<td>Universal testing</td>
<td>dMMR/pMMR and subsequent testing as defined by NICE DG27 guideline</td>
</tr>
<tr>
<td>Early onset CRC (EOCRC)</td>
<td>Constitutive</td>
<td>Diagnosis of CRC at 30 years and under</td>
<td>Panel testing determined by MMR status</td>
</tr>
<tr>
<td>Lynch-like syndrome</td>
<td>Somatic</td>
<td>dMMR tumours without hypermethylation/BRAF pathogenic variant and no</td>
<td>Somatic testing panel</td>
</tr>
<tr>
<td>serrated polyposis syndrome</td>
<td>Constitutive/ somatic</td>
<td>constitutional pathogenic variant in MMR genes</td>
<td>Excluded known predisposition syndromes</td>
</tr>
<tr>
<td>Multiple colorectal adenoma</td>
<td>Constitutive</td>
<td>MCRAs under 60 years of age with ≥10 adenomas, or patients over 60 years of</td>
<td>Gene panel testing</td>
</tr>
<tr>
<td>(MCRAs)</td>
<td></td>
<td>age with ≥20 adenomas, or ≥10 with a family history of multiple adenomas or</td>
<td></td>
</tr>
</tbody>
</table>

⁎ dMMR, MMR proficient; MMR, mismatch repair; NICE, National Institute for Health and Care Excellence; pMMR, MMR deficient.
We suggest that people diagnosed with CRC under age 50 years, where hereditary CRC syndromes have been excluded, undergo standard post-CRC surveillance for 3 years, then continue yearly colonoscopic surveillance until the age they are eligible for national screening. (GRADE of evidence: low; Strength of recommendation: weak)

Guidelines

Serrated polyposis syndrome (SPS)

We recommend a diagnosis of SPS should be made in accordance with the new WHO 2019 criteria for SPS. Since causative gene pathogenic variants for SPS have not been identified, a definitive diagnosis of SPS should be phenotype-driven. (GRADE of evidence: moderate; Strength of recommendation: strong)

Other intestinal polyposis syndromes may present with serrated lesions. If (i) the patient is under 50 or (ii) there are multiple affected individuals within kindred or (iii) there is dysplasia within any of the polyps, then we suggest that other polyposis syndromes should be excluded by gene panel testing before making a definitive diagnosis of SPS. (GRADE of evidence: very low; Strength of recommendation: weak)

We recommend the cumulative number of serrated polyps from all endoscopic examinations should be used when applying the WHO 2019 diagnostic criteria for SPS. (GRADE of evidence: moderate; Strength of recommendation: strong)

We recommend that patients with SPS should have colonoscopic surveillance yearly once the colon has been cleared of all lesions >5 mm in size. If no polyps ≥10 mm in size are identified at subsequent surveillance examinations the interval can be extended to 2 yearly. (GRADE of evidence: moderate; Strength of recommendation: strong)

We recommend all FDRs of patients with SPS on the basis of the new WHO 2019 SPS criteria, one or two should be offered an index colonoscopic screening examination at age 40 years or 10 years before the diagnosis of the index case. (GRADE of evidence: moderate; Strength of recommendation: strong)

We suggest all FDRs of SPS patients have a surveillance examination every 5 years unless polyp burden indicates an examination is required earlier according to post-polypectomy surveillance guidelines. (GRADE of evidence: low; Strength of recommendation: strong)

Multiple colorectal adenoma (MCRA) patients

We suggest an individualised approach to germline testing of patients with MCRA (defined as having 10 or more metachronous adenomas). Consider this testing for:
- Patients under 60 years of age with lifetime total of ≥10 adenomas or
- Patients from 60 years of age with lifetime total of: ≥10 adenomas, or
- ≥10 adenomas and a FHCC or polyposis

(GRADE of evidence: low; Strength of recommendation: weak)

We suggest that patients with a finding of 10 or more polyps (adenomas or serrated lesions) should, at their next colonoscopy, have a high-quality colonoscopic assessment with pancolonic dye spray in order to define accurately the multiple polyp phenotype. (GRADE of evidence: very low; Strength of recommendation: weak)

We suggest that the endoscopic management of patients with 10 or more metachronous adenomas, without MUTYH or APC gene mutations, should be individualised according to phenotype. (GRADE of evidence: very low; Strength of recommendation: weak)

We suggest annual colonoscopic surveillance for patients with 10 or more metachronous adenomas after the colon has been cleared of all lesions >5 mm in size. If no polyps 10 mm or greater in size are identified at subsequent surveillance
examinations the interval can be extended to 2 yearly. (GRADE of evidence: very low; Strength of recommendation: weak)

**Familial adenomatous polyposis (FAP)**
- We recommend that colonic surveillance should normally commence age 12–14 years in those confirmed to have FAP on predictive genetic testing. (GRADE of evidence: low; Strength of recommendation: strong)
- We suggest that for those with FAP, intervals between surveillance colonoscopy may be individualised depending on colonic phenotype every 1–3 years. (GRADE of evidence: low; Strength of recommendation: weak)
- We suggest that colonoscopy screening is performed for patients with congenital hypertrophy retinal pigment epithelium (CHRPE) for a specialist ophthalmic review. Patients with bilateral and multiple CHRPE lesions should be referred for screening for FAP and considered for genetic testing and colonoscopy. (GRADE of evidence: low; Strength of recommendation: weak)

**FAP: Surgery, and desmoid disease**
- We recommend that for patients with FAP who are undergoing colonoscopic surveillance, relative indications for surgery are: polyps >10 mm in diameter, high grade dysplasia within polyps and a significant increase in polyp burden between screening examinations. (GRADE of evidence: low; Strength of recommendation: strong)
- We recommend that absolute indications for immediate colorectal surgery in FAP include: documented or suspected cancer or significant symptoms attributable to the polyposis. (GRADE of evidence: low; Strength of recommendation: strong)
- We suggest that FAP patients should be counselled about the risk of postoperative desmoid disease formation. (GRADE of evidence: low; Strength of recommendation: weak)
- We suggest, for FAP patients before colectomy, determining genotypes or family history of desmoid disease which may be predictive of desmoid formation. (GRADE of evidence: very low; Strength of recommendation: weak)
- We suggest that sulindac in combination with high-dose selective oestrogen receptor modulators may be effective in FAP patients with intra-abdominal desmoids and desmoids located at the abdominal wall. (GRADE of evidence: low; Strength of recommendation: weak)
- We recommend the role of elective surgery for intra-abdominal desmoids should be restricted to treating secondary effects of the desmoid disease, and this surgery should be performed in expert centres. (GRADE of evidence: low; Strength of recommendation: strong)

**MUTYH-associated polyposis (MAP)**
- We recommend that colorectal surveillance is commenced in MAP commencing age 18–20 years. If surgery is not undertaken then annual surveillance is suggested. (GRADE of evidence: moderate; Strength of recommendation: strong)
- We recommend that for monoallelic MUTYH pathogenic variant carriers, the risk of CRC is not sufficiently different to population risk to meet thresholds for screening and routine colonoscopy is not recommended. (GRADE of evidence: moderate; Strength of recommendation: strong)

**Peutz-Jeghers syndrome (PJS)**
- We suggest that in an asymptomatic patient with PJS, GI surveillance by upper GI endoscopy, colonoscopy and video capsule endoscopy commence at age 8 years. We recommend that small bowel surveillance should continue 3 yearly. If baseline colonoscopy and oesophago-gastro-duodenoscopy (OGD) are normal, then they can be safely deferred until age 18 years; however, if polyps are found at baseline examination, then they should be repeated 3 yearly. Earlier investigation of the GI tract should be performed in symptomatic patients. (GRADE of evidence: low; Strength of recommendation: weak)

**Juvenile polyposis syndrome (JPS)**
- We suggest colonoscopic surveillance should commence from the age of 15 years or earlier if symptomatic. The surveillance interval should be 1–3 yearly, personalised according to colorectal phenotype. (GRADE of evidence: low; Strength of recommendation: weak)
- We suggest that for those with a confirmed clinical or genetic diagnosis, upper GI endoscopic surveillance should start at the age of 18 years for SMAD4 mutation carriers and 25 years for BMPRIA mutation carriers and those without an identified constitutional. The surveillance interval should be 1–3 yearly, personalised according to upper GI tract phenotype. (GRADE of evidence: low; Strength of recommendation: weak)
- We suggest that for those with an FDR with a clinical diagnosis of JPS and in whom a mutation has not then been identified, screening of the upper GI tract is not required routinely but should be initiated if/when a clinical diagnosis is made on the basis of colonic phenotype. It may, however, be considered if there is a family history suggestive of hereditary haemorrhagic telangiectasia (HHT), even in the absence of colonic polyps. (GRADE of evidence: low; Strength of recommendation: weak)
 ► We suggest that patients with SMAD4 pathogenic variant should be evaluated for HHT, and that those at risk of, or with a confirmed diagnosis of, HHT are best managed in conjunction with a specialist HHT centre. (GRADE of evidence: low; Strength of recommendation: weak)

 ► Patients with JPS and a microdeletion involving BMPRIA and PTEN are at risk of the clinical manifestations of both JPS and PTEN-hamartoma tumour syndrome (PHTS). We suggest that they should be referred to their local genetics centre for further advice and to coordinate their surveillance needs. (GRADE of evidence: low; Strength of recommendation: weak)

**SERVICE PROVISION, COMMUNICATION AND MANAGEMENT PRINCIPLES**

**We recommend that moderate risk of FHCC is the minimum threshold for referral from primary care**

(GRADE of evidence: very low; Strength of recommendation: strong)

Consensus reached: 100% agreement.

We recommend that individuals with an FHCC, which meets this referral criteria, be referred to a specialist familial CRC clinic in secondary or tertiary care.

(GRADE of evidence: low; Strength of recommendation: weak)

Consensus reached: 82% agreement.

We recommend that patients should be referred to a specialist service which includes access to constitutional genetic testing in the presence of defective MMR (with no evidence of MLH1 promoter methylation or BRAF V600E), or polyposis.

(GRADE of evidence: low; Strength of recommendation: strong)

There are insufficient clinical data to develop specific guidance for patients with very rare conditions such as polymerase proofreading associated polyposis (PPAP), or NTHL1 associated polyposis (NAP), therefore patients with these syndromes should be referred to multidisciplinary expert centres for clinical management.

(GRADE of evidence: low; Strength of recommendation: weak)

Consensus reached: 91% agreement.

We recommend that hospitals that diagnose or manage patients at hereditary CRC risk should ensure clinical pathways to facilitate their care, and processes to monitor the quality of the service.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 100% agreement.

We recommend that individuals at increased familial CRC risk receive specialist knowledge and are aware of patient support organisations and discussion with regard to lifestyle and participation in research projects.

(GRADE of evidence: very low; Strength of recommendation: strong)

Consensus reached: 95% agreement

People at hereditary CRC risk require coordinated, timely and high-quality care to reduce their cancer risk and should have access to a full range of management options that minimise the risk of morbidity and mortality. A structured referral pathway may ensure better inter-specialty communication and timely, efficient management of hereditary risk from primary through to tertiary care provision. It also facilitates an audit trail and subsequent monitoring of performance. Patients should have access to a full range of management options that minimise the risk of morbidity and mortality.

Moreover, studies consistently report that high quality screening and surveillance services result in a reduction of CRC incidence and mortality in individuals with FAP and LS. Registries of high-risk patients should be linked to robust quality assurance mechanisms for interventions, such as colonoscopy, with effective recall mechanisms in place to ensure high-risk individuals receive surveillance procedures on schedule.

Awareness of hereditary conditions may be inadequate, resulting in an inconsistent approach to the management of these individuals. Many patients do not have personalised management strategies and there is a failure to provide adequate follow-up. Patient advocacy organisations recommend improvements in the detection of pre-cancerous polyps, early diagnosis of CRC, and personalised treatment options for LS individuals, who should be seen by a team of specialists. The relevance of genomics is growing in clinical practice, and is increasingly relevant across the cancer multidisciplinary teams, with improving access to constitutional genetic testing. Genetic testing and/or counselling may resolve uncertainty about personal and familial cancer risk and provide information to guide and personalise decisions about future health care in anyone with an FHCC. It has been recommended that a dedicated clinical champion for hereditary CRC should be established in each colorectal multidisciplinary team (MDT) to oversee service coordination and to ensure patient pathways. The establishment of these champions will be another critical component in establishing equity of care. We recommend the establishment of family cancer specialist services by CRC teams in secondary care to ensure local pathways for patients at hereditary risk of CRC, and which can arrange testing of relatives for MMR status. This service should be supported by commissioners and incorporate a multidisciplinary approach involving geneticists, gastroenterologists and colorectal surgeons with links between secondary and tertiary care. Adherence to surveillance recommendations should be monitored at least annually. We suggest a minimum standard of ≥90% compliance. Non-compliant cases should be reviewed to determine whether reason for deviation from surveillance recommendations was clearly documented and clinically appropriate. Thus patients with a family history of CRC may be managed by their local hospital, and patients who require constitutional gene testing be managed by a tertiary care clinic, for example, in clinical genetics, either locally or regionally.

**FAMILY HISTORY OF CRC (FHCC)**

**Definitions and terminology**

A substantial proportion of the UK population have an FHCC without evidence of an inherited CRC syndrome. These individuals have a moderately increased relative risk (RR) of CRC (2–6 fold) compared with the general population. Lifetime CRC risk may be inferred from the age of onset of CRC in affected relatives, and familial aggregation, that is, the number of family members affected with CRC.

This section refers to asymptomatic patients referred for optimal management of a family history of either CRC or multiple polyps. The GDG agreed three categories of familial risk (in the absence of known hereditary CRC syndromes) which were determined according to lifetime CRC risk and the diagnostic yield of colorectal surveillance (box 1). Familial clusters (or aggregations) are of affected family members with CRC who are FDRs of each other. The individual referred for assessment should be an FDR of at least one affected member of such families.

Patients with average risk include those without an FHCC, or with an FHCC which does not significantly increase their lifetime CRC risk, that is, below the level of the moderate risk.
Guidelines

Box 1 Categories of risk in patients with a family history of colorectal cancer (FHCC)

Categories of risk – FHCC

► Average risk: No FHCC, or a FHCC which does not fulfil moderate or high-risk categories.
► Moderate risk FHCC:
  – One FDR diagnosed with CRC under 50 years, or
  – Two FDRs (in first degree kinship) diagnosed with CRC at any age, of whom the patient under assessment is an FDR of at least one affected individual.
► High risk FHCC: Families with a cluster of at least three affected FDRs with CRC diagnosed at any age, across at least two generations, of whom the patient under assessment is an FDR of at least one affected individual.

CRC, colorectal cancer; FDR, first degree relative.

population. For the average risk populations surveillance may be effectively managed via national bowel cancer screening programmes. Those people in moderate- or high-risk categories require additional surveillance above and beyond national screening however (figure 1).

Assessment of tumours in the affected relatives of those with an FHCC

We recommend that for all patients referred from primary care for assessment for an FHCC, MMR status should be assessed in tumour tissue from a close affected family member.

(GRADE of evidence: Moderate; Strength of recommendation: strong)

Consensus reached: 82% agreement.

We recommend that a reported family history of polyposis should be verified by review of histopathology and/or endoscopy reports which confirm the presence of a minimum of 10 adenomas or serrated lesions in an FDR.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 90% agreement.

Histopathological confirmation of CRC alters management of familial CRC surveillance in 20% of UK patients through verification of a diagnosis of CRC, multiple adenomas or other relevant features. Similarly review of endoscopy reports may assist in identification of patients with suspected familial risk such as those with polyposis syndromes.

When LS and Lynch-like tumours are excluded in families, their lifetime risk of CRC decreases. To quantify familial CRC risks associated with MMR deficient (dMMR) or MMR proficient (pMMR) tumours, a UK group analysed 2941 population-based cases of CRC. CRC risks in FDRs were strongly associated with dMMR tumours, early-onset disease and more than one affected FDR.

In a study by Bapat et al of 3143 CRC patients, dMMR tumours were associated with increasing numbers of FDRs with CRC (p=0.002); this association disappeared, however, when dMMR cases meeting Amsterdam criteria were removed from the analysis.

A multicentre international registry based study assessed MMR status in 33 496 FDRs of 4853 cases of CRC. In comparison with the FDRs of pMMR CRC cases the FDRs of CRC

Figure 1 Management of people with a family history of colorectal cancer. BSG, British Society of Gastroenterology; CRC, colorectal cancer; FHCC, family history of colorectal cancer; FDR, first degree relative; MMR, mismatch repair.
cases with suspected ‘Lynch-like’ syndrome and with LS had a higher risk of CRC, but not those with dMMR non-LS. There was a greater risk of CRC in FDRs if CRC cases were diagnosed under 50 years of age, or if the tumours had clinicopathological features suggestive of LS.

**Surveillance for colorectal neoplasia in those with a moderate risk FHCC**

*We recommend that patients with a moderate familial CRC risk should have a one-off colonoscopy at the age of 55 years.*

*(GRADE of evidence: moderate; Strength of recommendation: strong)*

Consensus reached: 85% agreement.

*We recommend that subsequent colonoscopic surveillance should be performed as determined by post-polypectomy surveillance guidelines.*

*(GRADE of evidence: moderate; Strength of recommendation: strong)*

Consensus reached: 95% agreement.

An important question is whether the adenoma detection rate in those with an FHCC is higher than the detection rate in the general population. Most CRCs develop from adenomas and “advanced” adenomas (AAs, defined as either an adenoma size of at least 10 mm, villous architecture of at least 25%, or high grade dysplasia) are considered to be the precursors of CRC. The term “advanced neoplasia” (AN) refers to the identification of either AAs or CRC.

The effectiveness and requirement for familial risk surveillance may be best determined by comparing the long-term CRC risk of a defined cohort of at-risk patients not undergoing surveillance with that of the general population. Theoretical relative risks of CRC <2 may be dominated by other genetic or environmental effects (and may require complex and validated risk modelling tools to determine suitability for surveillance). Where long-term CRC data are not available, the findings at surveillance may be used as a surrogate means to determine the need for post-polypectomy surveillance, although this method is inferior. In this context, that surveillance procedure may not have been warranted where the AA yield on that surveillance was less than doubled compared with a comparable yield in a control population.

There is a low prevalence of CRC in studies of surveillance in familial risk populations. There are limited data suggesting that metachronous CRC risk may be higher in patients at moderate familial risk versus population risk. As AAs are strongly associated with CRC development, AAs may be considered a proxy for CRC risk. In studies of patients with a moderate familial risk there is considerable heterogeneity in the prevalence of AA with a typical prevalence of between 8% and 10%; approximately double that of those without a family history (online supplementary tables 1-GRaDE table 1). There is evidence from observational studies that the diagnostic yield of colonoscopy has increased in familial CRC risk cohorts over the past two decades, consistent with improvements in endoscopic technique, equipment and quality assurance.

In the German bowel cancer screening population (an average risk population) the prevalence of AAs measured between 2003 and 2012 increased from 7.4% to 9.0% among men, and from 4.4% to 5.2% among women. In meta-analysis, the prevalence of adenomas is significantly higher in individuals with an FHCC than in controls (OR 1.7, 95% CI 1.4 to 3.5). Many observational control studies of surveillance colonoscopy for familial risk report a lower prevalence of AAs in the average risk population compared with the data from this German screening cohort. This may be related to lower ages of familial risk populations studied, and that many of the studies pre-date improvements in colonoscopic quality standards.

There is some observational evidence that colonoscopic surveillance mitigates this increased risk. In a German case-control study of CRC patients with an FHCC, those who had had a prior colonoscopy had a lower CRC risk that individuals without a family history who had not undergone colonoscopy. In the E3N French prospective study of 92 076 women, 692 CRCs were diagnosed after a median follow-up of 15.4 years; women with FHCC who had not had a previous colonoscopy had a 80% higher CRC risk than those without FHCC. In women who had had a previous colonoscopy, CRC risk was similar in women with and without FHCC.

**Age and risk of AAs in those with an FHCC**

Projected annual transition rates from advanced adenomas to CRC strongly increase with age, with annual transition rates increasing from 2.6% in patients in their 50s to >5% in their 80s. In an influential prospective study from 1994 FDRs of CRC patients had a risk of CRC at the age of 40 years equivalent to that of the average risk population aged 50 years. Notably, this historical study did not exclude patients with LS, and the increased risk conferred was predominantly in individuals with an affected FDR diagnosed under of 45 years.

Age is a strong predictor for adenomas and AAs in both familial risk individuals and controls in a series of observational studies. The incidence of AAs in patients aged 40–49 years in a surveillance cohort is equivalent to the general population, although the age of diagnosis of CRC in the affected FDR is not a predictor of risk of AAs. While adenomas but not AAs are more common in these studies, this may reflect the natural history of the adenoma to carcinoma sequence, whereby patients derive more benefit from surveillance colonoscopy from the age of 50 onwards, due to resection of AAs.

The subdivision of FHCC risk into those with 2× FDRs who were affected over or under 60 years of age is associated with any difference in the diagnostic yield on colonoscopic surveillance. Although this age criterion was used in previous iterations of this guideline to subdivide into “low-moderate” and “high-moderate” risk, no evidence was identified to support this differentiation.

Several studies of cohorts of patients with moderate FHCC have demonstrated a negligible incidence in AAs in colonoscopic surveillance before the age of 50 years, but an increased risk after this age. There is little evidence in case-control series of significant differences in AA incidence between patients with one FDR diagnosed under the age of 50 years, and those families with a cluster of two FDRs diagnosed at any age.

The prevalence of AAs under the age of 50 years is not significantly increased in patients under surveillance for FHCC compared with the average risk population. This supports commencing colonoscopic surveillance at the age of 50–55 years for those at moderate familial risk. As the incidence of CRC is increasing in younger patients, this age recommendation may need to be reviewed in future guideline iterations—pending further relevant data specifically in those with a FHCC.

**Surveillance in patients with a moderate familial risk of CRC**

To exclude LS in those with an FHCC, a close affected relative’s tumour should undergo MMR tumour testing. A pathology review of the relative’s tumour should also be undertaken to ensure that there is no evidence of multiple polyps. After this risk
assessment a decision about colonoscopic intervention should be considered.

The risk of AAs in surveillance colonoscopy (ie, after index/ screening procedure) is largely determined by the presence of advanced neoplasia at the index procedure.

In a surveillance programme from St Mark’s Hospital, London, with well-organised recall of high and moderate risk families, AAs and cancer were more common in families who fulfilled Amsterdam criteria than those at moderate risk (on initial colonoscopy 5.7% and 0.9%, respectively). In families with moderate risk, advanced pathology was particularly uncommon under the age of 45 (1.1% and 0%) and on follow-up colonoscopy if AAs were absent initially (1.7% and 0.1%). With colonoscopic surveillance the incidence of CRC was substantially lower (80% in families with moderate risk (p=0.00004) and 43% in families with LS (p=0.06)) than the expected incidence in the absence of surveillance.

Registry data from other populations also suggest a benefit in selected moderate familial risk populations undergoing surveillance colonoscopy. These studies also confirm an association of AAs at the index procedure with advanced neoplasia in subsequent surveillance procedures (GRADE online supplementary tables 1-3; GRADE table 1). In a Swedish moderate familial risk cohort, the risk of future AAs was associated with the prevalence of advanced lesions at the screening colonoscopy (multivariate analysis OR 5.22, 9% CI 2.3 to 9.94). It is of interest that adenomas and advanced lesions were not associated with the same risk factors: family history was predictive of advanced adenomas but not adenomas at the index screening colonoscopy.

The FACTS (Familial CRC Surveillance) randomised controlled trial compared intervals of surveillance in familial CRC. Individuals aged between 45 and 65 years with moderate familial CRC risk, where LS had been largely excluded, were randomly assigned to either a colonoscopy at 6 years or a colonoscopy at 3 and 6 years. Intention-to-treat analysis showed no significant difference in the proportion of patients with AAs (the primary outcome measure) at the first follow-up examination at 6 years (6.9%) versus 3 years (3.5%). The presence of AAs at the index colonoscopy was the only significant predictor for the presence of AAs at first follow-up (OR 5.2, 95% CI 1.6 to 16.87). Thus a 6 yearly interval was non-inferior to a 3 yearly surveillance interval, with the exception being that an AA at index colonoscopy predicts further advanced neoplasia at 3 years.

Surveillance for colorectal neoplasia in those with a high familial risk of CRC

We suggest that in high-risk families (a cluster of 3× FDRs with CRC across >1 generation) a 5 yearly colonoscopy should be performed from the age of 40 years until the age of 75.

(Grade of evidence: low; Strength of recommendation: weak)

Consensus reached: 86% agreement.

Families who fulfil Amsterdam criteria but who do not have evidence of dMMR do not share the same cancer incidence as families with LS (ie, hereditary MMR deficiency). Relatives in such families were found to have a lower incidence of CRC than those in families with LS, and incidence was not increased for other cancers. These families should not be described or counselled as having LS. To facilitate distinguishing these entities, the designation of “familial CRC type X” was suggested by Lindor et al to describe this type of familial aggregation of CRC.

In a prospective surveillance study of a high familial risk population there was no significant difference in the prevalence of AAs in LS individuals versus FCC-X individuals. However on follow-up there were no incident cancers in the FCC-X group versus 4.4% CRC in Lynch patients, indicating lower risk of interval CRC in FCC-X despite equivalent AA risk.

In a prospective pooled cohort study of 1585 patients from eight international centres families were classified as FCC type X if they fulfilled the original Amsterdam criteria and late onset (LOFCC) if they fulfilled the Amsterdam criteria apart from not having a cancer diagnosed aged under 50. The results for FCC type X and LOFCC were very similar. At baseline, 22 prevalent asymptomatic CRCs were diagnosed, 120 (7.6%) individuals had high-risk adenomas and 223 (14.2%) simple adenomas. On follow-up high-risk adenomas were detected in 92 (8.7%) and multiple adenomas were detected in 20 (1.9%) individuals, from approximately 35 years of age onwards. Again the presence of AA at index colonoscopy was predictive of advanced neoplasia at subsequent procedures—33% of patients with an AA at index colonoscopy had an AA or cancer on follow-up.

This study by Mesher et al indicated patients at high familial CRC risk should be managed similarly with 5 yearly colonoscopies undertaken from between 30 and 40 years of age with more intensive surveillance in individuals developing multiple or high-risk adenomas.

Amsterdam criteria families, where MMR testing of a CRC from an affected individual is not possible, may be offered surveillance as per LS. However such patients should be reviewed by a specialist service who may consider alternative testing strategies such as panel testing of affected individuals, or unaffected testing.

PREVENTION AND LIFESTYLE MODIFICATION IN FAMILIAL CRC

We recommend that individuals with LS should be advised that regular use of daily aspirin reduces CRC risk.

(Grade of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 90% agreement.

Long-term data from the CAPP2 RCT suggests that aspirin reduces this risk by approximately half as compared with placebo. The benefits of regular aspirin intake take at least 3 to 5 years to become evident. Taking aspirin for less than 2 years’ duration does not seem to confer any benefit in reducing the incidence of cancer, or increasing survival in patients with LS.

We recommend that people with LS should be offered research opportunities to take aspirin daily at different dosages. If they decline research participation they may be advised on their choices regarding dose of aspirin, risks and benefits of long-term aspirin use and ensure their medical practitioner is aware of their intake.

(Grade of evidence: low; Strength of recommendation: strong)

Consensus reached: 90% agreement.

There is uncertainty about the optimum dosage of aspirin to recommended to individuals with LS. There is some evidence that long-term intake of daily 600 mg aspirin can reduce the risk of all cancers including CRC in LS from the CAPP2 randomised control trial. There is no other high quality evidence for any other dose of aspirin/ length of treatment but there are studies ongoing (CAPP3 trial) which aim to identify optimum dosage. Evidence for the optimum dose will inform awareness and education among health professionals to mitigate the reluctance to prescribe higher doses of aspirin within primary care. In the

Guidelines


interim clinicians may consider 150 mg aspirin in the context of LS outside of a clinical trial, with 300 mg doses in those with a BMI above 25 kg/m².

There is insufficient evidence of the benefit of chemoprophylaxis in polyposis syndromes.

(GRADE of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 100% agreement.

Non-steroidal anti-inflammatory drugs (NSAIDs) have been the most commonly studied chemoprophylaxis agents in patients with FAP, predominantly for lower GI tract disease, with some RCTs. Aspirin and tiracoxib have been found to be ineffective, although sulindac, celecoxib and rofecoxib have been demonstrated to reduce adenoma burden in the short term.54-57 A number of small series provide further support for these drugs and have also shown benefit from topical indomethacin.58 However long-term cancer prevention as an end point has not been adequately addressed.59 60 In the largest cohort of 54 patients, published in abstract form, 10% developed cancer while on chemoprophylaxis.61

Other classes of agents have also been assessed, such as omega 3 fish oils, but again the results are only short term with reduction in polyp size and number as the main endpoint.62

Celecoxib and the combination of sulindac and etoriflavin have been reported as being beneficial63 64 for those with FAP and advanced duodenal disease; this was a short-term study using polyp number and size as the primary endpoint. A cohort study reported outcomes of the use of Evien dep, which was observed to reduce polyp number and size.65 However, no studies have demonstrated an effect on duodenal cancer prevention in FAP.

Cyclooxygenase-2 (COX-2) expression may be increased in JPS.66-67 There exists a theoretical potential benefit in the use of selective COX-2 inhibitors in JPS or PJS, but to date there are no trials demonstrating efficacy.

We recommend that individuals at increased familial risk of CRC should be strongly encouraged not to smoke, to maintain a normal BMI, to moderate their consumption of red and processed meat, and to exercise regularly.

(GRADE of evidence: low; Strength of recommendation: moderate)

Consensus reached: 81% agreement.

Diet and lifestyle factors are well established as significant contributors to up to half of all CRCs.68 70 A systematic review of epidemiological studies investigating the associations between nutritional factors, FHCC and CRC risk71 suggests that combinations of FHCC and higher consumption of alcoholic beverages, red or processed meat, or overweight/obesity increases the risk of CRC. There is evidence that LS individuals who smoke (particularly males with MLH1 mutations) have an increased risk of CRC. Data suggest current smokers are at significant increased risk of CRC irrespective of the age of initiation of smoking. Risk in former smokers decreased with each non-smoking year.72-75

CAPP2 study data from 29 month follow-up also indicate that overweight individuals with LS were more likely to develop CRC than those normal/underweight.76-79 Though modifiable environmental risk factors such as weight and exercise80 are common to both sporadic and familial CRC, individuals with familial risk may benefit from discussion about modifiable factors in order to potentially reduce their level of risk.81 82 There is emerging evidence of the benefit of targeted lifestyle modification in those with an FHCC.83

QUALITY AND ADVANCED ENDOSCOPIC IMAGING IN COLONSCOPY SURVEILLANCE

We recommend that colonoscopy is the gold standard diagnostic and preventative method of surveillance for people with a hereditary risk of CRC.

(GRADE of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 100% agreement.

We recommend that all surveillance colonoscopies are performed by endoscopists who consistently achieve BSG colonoscopy KPI minimum standards, specifically caecal intubation rate, adenoma/polyp detection rate and comfort score.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 94% agreement.

We suggest a repeat colonoscopy performed by an expert endoscopist is indicated in the event of a previously failed colonoscopy, with efforts made to both improve patient experience and to ensure procedure completion, given the advantages of colonoscopic surveillance. If colonoscopy is not possible then consider CT colonography.

(GRADE of evidence: low; Strength of recommendation: weak)

Consensus reached: 90% agreement.

We suggest that if the bowel preparation for colonoscopy is inadequate or if the examination is incomplete then a repeat colorectal surveillance procedure should be arranged within 3 months.

(GRADE of evidence: moderate; Strength of recommendation: weak)

Consensus reached: 95% agreement.

We suggest high-quality, high-definition white light endoscopy as the preferred modality for colonoscopy surveillance. Chromoendoscopy (virtual or dye-based) does not offer a clear advantage over high definition white light examination for colonoscopic surveillance, apart from in the context of determining the multiple polyp phenotype.

(GRADE of evidence: moderate; Strength of recommendation: weak)

Consensus reached: 89% agreement.

High quality colonoscopy has been recognised as a core element of successful cancer prevention in sporadic patients.84 There are limited data that this may also be relevant to cancer prevention in LS.85 Therefore colonoscopic quality indicators in endoscopists performing surveillance in LS patients should at least reach if not exceed the KPIs required for sporadic colonoscopy, using validated measures, in particular caecal intubation rate, adenoma/polyp detection rate and, given that patients may require serial colonoscopic procedures, comfort score.86 87

Colonoscopy is less effective for cancer prevention if the procedure is not complete to the caecum or the bowel preparation is inadequate. Where caecal intubation is not achieved, a repeat examination with an expert colonoscopist is appropriate. Inadequate bowel preparation reduces adenoma and advanced adenoma detection rates.88-89 Inadequate preparation at initial colonoscopy led to a threefold increase in miss rate in adenomas 5 mm or smaller.90 Therefore, repeat colonoscopy within 3 months seems appropriate for individuals at high familial risk.

Advanced imaging techniques have been proposed to help reduce missed lesions, especially small and flat lesions. More non-polypoid lesions are found in LS compared with sporadic patients.91 Chromoendoscopy, both dye-based and virtual, was recommended in recent ESGE guidelines on colonoscopic surveillance in LS.92
Tandem studies with chromoendoscopy show a consistent benefit (online supplementary 1 GRADE table 2)93-98; however, a study comparing a second pass with chromoendoscopy to a second white light pass did not show improved adenoma detection.94 Meta-analysis of data in sporadic patients shows an OR for at least one neoplastic lesion of 1.53 (95% CI 1.31 to 1.79).99 Real world cohort data comparing white light to chromoendoscopy provides some support for improved detection in high familial risk patients (39% LS) (15/24 (63%) adenomas with chromoendoscopy versus 15/77 (19%) white light endoscopy).100 A recent large, multicentre, Spanish randomised parallel group study, using high definition endoscopes and with high adenoma-detecting endoscopists, did not demonstrate a significant increase in adenoma detection with chromoendoscopy in 256 Lynch patients (OR 1.34, 95% CI 0.79 to 2.28).101 A similar sized multi-centre parallel group RCT study in the Netherlands had similar results using chromoendoscopy in the proximal colon (overall adenoma detection rate (ADR) 33% vs 27% with white light endoscopy).102 At 2 year follow-up there was no difference in ADR between the two groups, but there were four cancers in the chromoendoscopy group versus one in the white light endoscopy group. Simply combining the results for ADR of these two studies gives a risk ratio of 1.23 (95% CI 0.94 to 1.60, p=0.14; Fisher exact, n=497; our calculation) suggesting a limited clinical benefit in terms of ADR for considerable extra effort, which may not translate into improved cancer prevention.

Virtual chromoendoscopy, narrow band imaging (NBI, second generation, Olympus) and I-SCAN (Pentax) have shown some benefit in tandem studies (online supplementary tables 1 GRADE table 3); however, this improved detection is not consistent with meta-analysis data from sporadic patients.103 and NBI performed less well than chromoendoscopy in a cohort, tandem study.102 In a study comparing different forms of chromoendoscopy, chromoendoscopy was superior to white light colonoscopy, autofluorescence imaging, and narrow-band imaging for detection of diminutive colorectal lesions in adenomatous polyposis.103

Advanced colonoscopic imaging can assist in making a diagnosis of polyposis by revealing additional lesions required to meet diagnostic criteria: diagnoses of adenomatous polyposis may be missed if dye spray is not used,104 and there is similar evidence from the CONSCOP study that the identification of serrated polyps is enhanced through the use of pancolonic dye spray.105

In summary, a high quality, high definition white light colonoscopic examination, by an endoscopist who meets all colonoscopy KPIs, seems adequate for LS and high familial risk patients, with the exception of those with multiple polyps where chromoendoscopy may help define the phenotype.

Non-invasive surveillance methods for people with FHCC

There is insufficient evidence to recommend other methods of surveillance for those with familial CRC risk such as FIT, MR or CT colonography.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 95% agreement.

In meta-analysis of 12 published studies of patients at increased risk of CRC (predominantly familial risk), the average sensitivity of FIT for advanced neoplasia was 48% (95% CI 39% to 57%) and the average specificity was 93%.106 A subgroup analysis of patients with familial risk only was performed, and the sensitivity for CRC was 86% and for advanced neoplasia 46%. Thus, although FIT may be close to equivalence to colonoscopy for the detection of CRC, AAs would be missed by surveillance with FIT alone.

Patients with an FHCC no longer on colonoscopic surveillance should participate in national bowel cancer screening programmes designed for the average risk population. Some indirect evidence suggests that FIT or other forms of stool testing methods alongside colonoscopy may potentially be a useful adjunct in surveillance in patients after discharge from colonoscopic surveillance.107

Other methods such as colon capsule108 or MR colonography109 lack efficacy in this population. Although there is some evidence of the efficacy of CT colonography,110 it is not clear how effective CT may be in the identification of serrated or non-polypoid lesions. Repeated CT scanning in particular may be inappropriate due to the risk of radiation damage to patients with inherited DNA repair defects. However, if total colonoscopy is not possible despite expert referral, low radiation-dose CT colonography with quality assurance111 is the preferred modality, as it has similar test performance to colonoscopy for CRC.112

**LYNCH SYNDROME**

We recommend that for all people when first diagnosed with CRC, testing using immunohistochemistry (IHC) for MMR proteins or microsatellite instability is used to identify tumours with deficient DNA MMR, and to guide further sequential testing for LS.

(GRADE of evidence: strong; Strength of recommendation: strong)

Consensus reached: 100% agreement.

LS is a condition defined by the presence of pathogenic variants in the coding sequence or regulatory domains of the four MMR genes: MLH1, MSH2, MSH6 and PMS2. Patients with EPCAM mutations which embrace the regulatory domain of MSH2 should be managed as those with MSH2 pathogenic variants. Selection methods based on family history of cancer, or other clinical parameters, such as the Amsterdam or Bethesda criteria,112 113 were used historically to identify high-risk patients who may benefit from interventions and/or genetic testing; however, advances in constitutional testing in the past two decades have facilitated the accurate genetic diagnosis of LS. A comprehensive review performed by NICE of the clinical- and cost-effectiveness of universal diagnostic testing for LS was published in 2017.18 We recommend the use of colonoscopy biopsies as the preferred source material for tumour MMR testing.

**Colonoscopic surveillance and LS**

We recommend that colonoscopic surveillance should be performed at a 2 yearly interval for all LS patients.

(GRADE of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 85% agreement.

Surveillance colonoscopy in LS does not completely eradicate the risk of CRC, with the well-recognised phenomenon of interval cancers related to multiple factors including adherence and timeliness of colonoscopy. However, the optimal interval for surveillance colonoscopy is yet to be established.

The literature around colonoscopic surveillance is mixed with few studies reporting on recognised key performance indicators including adenoma detection rate, or caecal intubation rate or compliance with the screening interval. A study in the UK identified that hospital recall systems, clinician or patient related issues affect compliance with LS surveillance intervals.115 In this study variable colonoscopy quality indicators were highlighted with a
caecal intubation rate was 92%, and approximately 10% had inadequate bowel preparation. In a retrospective, two centre Dutch study 31 interval CRCs were diagnosed in 29 patients with LS, within 2 years of previous colonoscopy, all of whom were MLH1 and MSH2 pathogenic variant carriers, and 84% were located in the proximal colon. In three of a total of five patients where colon examination was not achieved during the previous colonoscopy, the interval CRC was found in the unexamined proximal segment. In six of nine patients with a previous adenoma, the interval CRC was detected in the same colon segment, raising the possibility of incomplete endoscopic resection.

There is some evidence that earlier tumour stage may be observed in those with more frequent colonoscopy. More recent prospective data have described cancer incidence and survival in LS in patients undergoing surveillance. The incidence of CRC was influenced by the LS-associated gene; cumulative CRC incidence at 70 years by gene was greater in those with MLH1 (46%) or MSH2 (35%) constitutional pathogenic variants compared with those with MSH6 (20%) or PMS2 (0%) pathogenic variants (although with wide confidence intervals). The prognosis from interval CRC was good, or (20%) 10–20 years. The data regarding differences between the LS-associated genes are not sufficiently robust such that the surveillance interval can be stratified by LS gene. A lower penetrance of CRC in those with an associated gene pathogenic variant. Lynch syndrome: gene and gender specific guidelines

We recommend that age of onset of surveillance colonoscopy should be stratified according to the LS-associated gene. We recommend colonoscopy from the age of 25 years for MLH1 and MSH2 mutation carriers and 35 years for MSH6 and PMS2 mutation carriers. There are insufficient data to support strati- fying age of onset of surveillance by gender.

(GRADE of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 95% agreement.

Outcomes from the Dutch Hereditary Non-Polyposis Colorectal Cancer (HNPCC) registry showed that only 2/246 (0.8%) patients with LS developed CRC before the age of 20 years and another two between the age of 20 and 25 years. A number of other studies have confirmed that the risk of developing CRC before the age of 25 years is very low. It is largely based on these studies that most groups have recommended starting surveillance colonoscopy at the age of 20–25 years. However, a meta-analysis of the data for MLH1 and MSH2 pathogenic variant carriers has questioned whether surveillance colonoscopy was justified before the age of 30 years.

The US multi-society task force guidelines recommend starting surveillance colonoscopy at 20–25 years (or 2–5 years younger than youngest affected relative if diagnosed <25 years) but to consider starting at 30 and 35 years for MSH6 and PMS2 patho- genic variant carriers, respectively.

Ten Broeke et al described a series of 377 patients with constitutional PMS2 pathogenic variants. They observed that the median age at first CRC was 52 years (26–86 years), and noted gender differences in CRC risk. The cumulative risk (%) in men and women up to the age of 69 years was 18.8% and 10.5%, respectively. They recommended that commencement of colonoscopy surveillance could be deferred. A nationwide French study of patients with LS supports a genotype-phenotype correlation. There were no PMS2 pathogenic variant carriers in this cohort. Overall there was a cumulative CRC risk by 70 years of 38% in males and 31% in females. When analysed by genotype the cumulative CRC risks were 46% MLH1, 48% MSH2 and 12% MSH6. Furthermore, age (years) at CRC also varied by genotype: MLH1 45 (15–90), MSH2 44 (16–95), MSH6 54 (24–85). These data are supported by analyses of a prospective LS database. These reports observed that there was genotype specific penetrance. Furthermore, within each genotype the incidence of CRC was age dependent and CRC occurred as a stochastic event. In the report addressing first cancer incidence and survival in patients with LS receiving surveillance, Moller et al observed cumulative CRC cancer incidence to the age of 70 years was 46% MLH1, 35% MSH2, 20% MSH6 and 0% PMS2. Overall, there was no difference between gender. The most recent report from this prospective database describes cancer risk and survival up to age 75 years, analysing the data both by gene and gender. The cohort includes 3119 patients with 24,475 observation years. The previously described genotype dependent

Lynch syndrome: gene and gender specific guidelines

We recommend that age of onset of surveillance colonoscopy should be stratified according to the LS-associated gene. We recommend colonoscopy from the age of 25 years for MLH1 and MSH2 mutation carriers and 35 years for MSH6 and PMS2 mutation carriers. There are insufficient data to support strati- fying age of onset of surveillance by gender.

(GRADE of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 95% agreement.

Outcomes from the Dutch Hereditary Non-Polyposis Colorectal Cancer (HNPCC) registry showed that only 2/246 (0.8%) patients with LS developed CRC before the age of 20 years and another two between the age of 20 and 25 years. A number of other studies have confirmed that the risk of developing CRC before the age of 25 years is very low. It is largely based on these studies that most groups have recommended starting surveillance colonoscopy at the age of 20–25 years. However, a meta-analysis of the data for MLH1 and MSH2 pathogenic variant carriers has questioned whether surveillance colonoscopy was justified before the age of 30 years.

The US multi-society task force guidelines recommend starting surveillance colonoscopy at 20–25 years (or 2–5 years younger than youngest affected relative if diagnosed <25 years) but to consider starting at 30 and 35 years for MSH6 and PMS2 patho- genic variant carriers, respectively.

Ten Broeke et al described a series of 377 patients with constitutional PMS2 pathogenic variants. They observed that the median age at first CRC was 52 years (26–86 years), and noted gender differences in CRC risk. The cumulative risk (%) in men and women up to the age of 69 years was 18.8% and 10.5%, respectively. They recommended that commencement of colonoscopy surveillance could be deferred. A nationwide French study of patients with LS supports a genotype-phenotype correlation. There were no PMS2 pathogenic variant carriers in this cohort. Overall there was a cumulative CRC risk by 70 years of 38% in males and 31% in females. When analysed by genotype the cumulative CRC risks were 46% MLH1, 48% MSH2 and 12% MSH6. Furthermore, age (years) at CRC also varied by genotype: MLH1 45 (15–90), MSH2 44 (16–95), MSH6 54 (24–85). These data are supported by analyses of a prospective LS database. These reports observed that there was genotype specific penetrance. Furthermore, within each genotype the incidence of CRC was age dependent and CRC occurred as a stochastic event. In the report addressing first cancer incidence and survival in patients with LS receiving surveillance, Moller et al observed cumulative CRC cancer incidence to the age of 70 years was 46% MLH1, 35% MSH2, 20% MSH6 and 0% PMS2. Overall, there was no difference between gender. The most recent report from this prospective database describes cancer risk and survival up to age 75 years, analysing the data both by gene and gender. The cohort includes 3119 patients with 24,475 observation years. The previously described genotype dependent
penetration was confirmed; of note the number of PMS2 carriers in this cohort is small.

Further studies are required to further stratify risk for MSH6 and PMS2 pathogenic variant carriers. Until such data are available, based on the current sparse literature it appears reasonable to defer initiation of surveillance for these patients. The current literature is sufficiently robust to recommend that MSH6 and PMS2 pathogenic variant carriers should have different ages of onset for colorectal surveillance (figure 2).

Surgery in LS patients with CRC
We suggest that for LS patients with MLH1 or MSH2 mutations who develop colon cancer or colonic neoplasia not amenable to endoscopic control, the decision to perform segmental versus total/near total colectomy should balance the risks of metachronous cancer, the functional consequences of surgery, the patient’s age and patient’s wishes.

(GRADE of evidence: Moderate; Strength of recommendation: Strong)
Consensus reached: 100% agreement. We recommend that for LS patients with MSH6 or PMS2 mutations there is insufficient evidence for oncological benefit of extended colectomy over segmental resection.

(GRADE of evidence: low; Strength of recommendation: strong)
Consensus reached: 89% agreement.

When abdominal-perineal excision can be avoided, a standard low anterior resection is a reasonable option to treat rectal cancers in LS patients, even though the residual colon is at high risk of metachronous neoplasia.

(GRADE of evidence: low; Strength of recommendation: weak)
Consensus reached: 88% agreement.

The decision to perform a total/subtotal colectomy or segmental colectomy involves consideration of:

i. The risk of metachronous CRC

ii. Survival from metachronous CRC

iii. Functional consequences and quality of life (QoL) following surgery

This decision regarding which operation is preferable should be made on the basis of individual patient factors and preferences, with special emphasis on the risk of metachronous CRC, age and the preparedness of the patient to continue colonoscopic surveillance. High-quality patient information and shared decision-making between patient and surgeon will facilitate this decision.

Risk of metachronous cancer
Parry et al found the cumulative risk of metachronous CRC to be 16% at 10 years, 41% at 20 years and 62% at 30 years after segmental colectomy. In contrast none of 50 subjects who had extensive colectomy developed metachronous CRC. They calculated that the risk of metachronous CRC was reduced by 31% for every 10 cm of large bowel removed. Kalady et al reported 296 patients (253 with segmental colectomy and 43 with total colectomy/ileorectal anastomosis). Of the 253 segmental colectomy patients, 55 patients (25%) developed a second CRC at a median of 69 months after index surgery. Stages of the metachronous cancers were I-16, II-18, III-12, and IV-2. By comparison, four of 38 patients (11%) who underwent total colectomy developed subsequent high-risk adenomas and only three (8%) developed metachronous cancer.

A Finnish study reported the cumulative risk of subsequent CRC to be 20% within 10 years and 47% within 25 years after standard resection and 4% and 9% after extended surgery. A further study showed metachronous CRC in 6.3% of pathogenic variant carriers treated with total/subtotal colectomy compared with 27% treated by segmental colectomy. In meta-analysis much of the excess risk of metachronous CRC appears to be in carriers of pathogenic variants in the MLH1 and MSH2 pathogenic carriers, with insufficient data to suggest excess risk in MSH6 or PMS2 variant carriers.
Survival from metachronous cancer

Moller et al in an international collaborative study found the cumulative incidence of metachronous CRC was 36% from 40 to 70 years. Five and 10 year crude survival after colectomy for metachronous CRC was 94% and 91%, respectively, with no significant difference between the pathogenic variants of the different genes.19

Natarajan et al found no significant difference in survival time between patients undergoing extended colectomy and limited resection in patients with LS.138 The potential health effects in terms of life expectancy for patients undergoing subtotal colectomy or hemicolectomy for CRC were analysed. The 10 year risk of CRC after subtotal colectomy was 4% and after hemicolectomy was 16% and stages of CRCs detected within a 2 year surveillance interval were 32% Dukes’ A, 54% Dukes’ B, and 14% Dukes’ C (derived from two cohort studies). The overall LE gain of subtotal colectomy compared with hemicolectomy was 14% Dukes’ C (derived from two cohort studies). The overall LE gain of subtotal colectomy compared with hemicolectomy was 14% Dukes’ C (derived from two cohort studies). The overall LE gain of subtotal colectomy compared with hemicolectomy was 14% Dukes’ C (derived from two cohort studies).

The authors concluded that unless surveillance results improve, subtotal colectomy was the preferred treatment for CRC in LS

Functional consequences and QoL

In a Dutch study, no difference in global QoL was noted between 51 LS patients who underwent partial colectomy and 53 patients who underwent subtotal colectomy, although functional outcome (stool frequency and social impact) was worse after subtotal colectomy than after segmental colectomy.141

Maeda et al constructed a state-transition (Markov) model to compare segmental colectomy and total abdominal colectomy with ileorectal anastomosis.142 Quality-adjusted life years (QALYs) were calculated based on utility states for patients based on the colectomy they received. Multiple sensitivity analyses were planned to examine the impact of each assumption on model results. For young (30-year-old) patients with LS, mean survival was slightly better with total colectomy than with segmental resection (34.8 vs 35.5 years). When QALYs were considered, the two strategies were approximately equivalent, with QALYs per patient of 21.5 for segmental colectomy and 21.2 for total colectomy. They suggested that with advancing age, segmental colectomy becomes a more favourable strategy.

The decision regarding which operation is preferable should be made on the basis of individual patient factors and preferences, with special emphasis on the risk of metachronous CRC, age and the preparedness of the patient to continue colonoscopic surveillance.

Patients presenting with rectal cancer

A standard low anterior resection or abdominal perineal resection is a reasonable option to treat rectal cancers in LS patients, even though the residual colon is at high risk of metachronous neoplasia. A retrospective study of 79 LS patients with rectal cancer who had undergone proctectomy found a cumulative risk of metachronous colon cancer to be 19% at 10 years, 47% at 20 years, and 69% at 30 years after surgical resection.143 Kalady et al followed 50 HNPPC patients with a primary diagnosis of rectal cancer treated by proctectomy. Forty-eight high-risk adenomas developed in 13 patients (39.4%) and five patients (15.2%) developed metachronous adenocarcinoma at a median of 6 years (range 3.5–16) after proctectomy, including three at an advanced stage. Overall 17 of 33 patients (51.5%) developed high-risk adenoma or cancer after proctectomy.144

Upper GI, pancreatic and small bowel risk management in LS

We recommend that gastric, small bowel, or pancreatic surveillance in LS patients is only performed in the context of a clinical trial.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 100% agreement.

We recommend screening for H pylori in patients with LS and subsequent eradication therapy if indicated.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 100% agreement.

Gastric cancer

In a retrospective study, Capelle et al estimated lifetime risks of gastric cancer in 2014 patients with LS from the Dutch Hereditary Cancer Registry.145 Gastric cancer was diagnosed in 32 patients (1.6%); 22 of these patients (69%) had a negative family history of gastric cancer. The lifetime risk of gastric cancer was reported to be 4.8% and 9% for MLH1 and MSH2 pathogenic variant carriers, respectively. None of the 378 MSH6 pathogenic variant carriers developed gastric cancer. The median age of diagnosis was 55 years (range 27–82 years).

In a recent prospective study, Moller et al reported gene-specific prospective cumulative cancer risks (up to the age of 75 years) for gastric cancer in 3119 patients with LS.146 The risk of gastric carcinoma was reported to be the highest for MLH1 and MSH2 pathogenic variant carriers: 7.1% (95% CI 3.5% to 10.8%) for MLH1, 7.7% (95% CI 1.9% to 13.6%) for MSH2, and 5.3% (95% CI 0.0% to 13.1%) for MSH6. No gastric cancers were observed in PMS2 pathogenic variant carriers. The study also reported a 5 year overall survival rate of 61% (95% CI 33% to 81%) for LS-associated gastric cancer. This compares favourably with the prognosis in unscreened patients with resectable gastric cancer, who have been reported to have an overall 5 year survival of 10–30%.147

A Finnish study reported the characteristics of 62 gastric cancers that occurred in 570 family members from the Finnish HNPCC registry. There was an overrepresentation of intestinal gastric cancers, which was found in 79% of cases, with only 13% of cases being diffuse gastric cancers. As the development of intestinal gastric cancers is thought to be closely associated with H. pylori-associated chronic gastritis, the authors investigated whether atrophic gastritis and H. pylori infection could be markers of gastric cancer risk in patients with LS. Twenty percent of the LS-associated gastric cancers were H. pylori positive. The median age at diagnosis was 56 years.147

Renkonen-Sinisalo et al assessed the diagnostic yield of upper endoscopy in a series of 73 MMR pathogenic variants carriers (median age 47 years).148 The authors found no early gastric cancers or premalignant lesions (diagnostic yield 0). A single screened-detected small bowel cancer was identified: an advanced stage duodenal cancer. No additional studies evaluating the benefit of gastric surveillance in LS were identified from the literature.

The cumulative lifetime risk of gastric cancer is relatively low. Aspirin chemoprophylaxis may reduce the risk of all LS-associated cancers in patients with LS. In conclusion, there is no convincing evidence to support the utility of gastric surveillance in patients with LS.
Small bowel cancer

The cumulative lifetime risk of developing small bowel cancer has been estimated to be 4.2% in patients with constitutional MLH1 and MSH2 pathogenic variants. Small bowel cancers have rarely been reported in patients with constitutional MSH6 and PMS2 pathogenic variants. A recent study has reported gene-specific prospective cumulative cancer risks (up to the age of 75 years) for duodenal carcinoma in 3119 patients with LS. The risk of duodenal carcinoma was reported to be the highest for MLH1 pathogenic variant carriers (6.5%, 95% CI 1.7% to 10.2% for MLH1; 2.0%, 95% CI 0.1% to 4.0% for MSH2), and no small bowel cancers were observed in patients with constitutional MSH6 or PMS2 pathogenic variants. LS-related small bowel carcinomas have an earlier onset compared with sporadic tumours with a median age of onset of 52 years (range 23–69 years). LS-related small bowel carcinomas most commonly occur in the duodenum (49%), and decrease in frequency from the jejunum (29%) to the ileum (12%).

Patients with LS who develop small bowel cancers have been demonstrated to have a better prognosis compared with patients who develop sporadic tumours. Moller et al, in a recent study, reported a 5 year survival rate of 67% (95% CI 28% to 88%) in LS patients with small bowel cancers diagnosed below the age of 65. This compares favourably with the 5 year survival of sporadic small bowel adenocarcinoma, which has been estimated to be 25–30%. Several studies have investigated the diagnostic yield of surveillance for small bowel cancers in patients with LS. In 2010 Saurin et al compared the use of CT enteroclysis and video-capable endoscopy (VCE) in 35 asymptomatic patients with LS. Histologically confirmed small bowel tumours were identified in three patients (diagnostic yield of 8.6%): one jejunal adenocarcinoma (T3N0M0) and two adenomas with low-grade dysplasia. VCE identified all three tumours, but CT enteroclysis would have missed the two adenomas. In a Dutch study, Haanstra et al investigated the prevalence of small bowel tumours in 200 asymptomatic pathogenic variant carriers (aged 35–70 years) using VCE. Caecal visualisation was achieved in 95% of procedures. Histologically confirmed small bowel tumours were identified in two patients (diagnostic yield of 1%): one adenocarcinoma (TisN0Mx) and one adenoma, both located in the duodenum. In addition, another patient was diagnosed with a duodenal carcinoma (T2N0Mx) 7 months after a negative VCE (incidence of 1.5%). This suggests that VCE may miss some small bowel tumours. In a follow-up study, asymptomatic LS patients who underwent a VCE were invited to undergo a second VCE procedure 2 years later. A total of 155 (78%) of the initial 200 pathogenic variant carriers underwent a second VCE. Potentially significant lesions were identified in 17 patients (11%), which required further investigations: eight gastroduodenoscopies and nine balloon-assisted endoscopies were carried out, but no small bowel tumours were identified.

In the CAPP2 randomised trial, Burn et al examined the effect of aspirin in 861 patients with LS who were randomly assigned in a two-by-two factorial design to 600 mg of aspirin or aspirin placebo or 30 g of resistant starch or starch placebo, for up to 4 years. The primary outcome was the incidence of new primary CRCs, but the incidence of all LS cancers was also examined. Following a re-analysis of the data at a mean follow-up of 55.7 months, for participants completing 2 years of intervention (258 aspirin; 250 aspirin placebo), per-protocol analysis yielded a hazard ratio (HR) of 0.45 (95% CI 0.26 to 0.79; p=0.005) for all LS-related cancers.

The absolute lifetime risk of small bowel cancer in patients with LS is 4.2%; this risk is likely to be most significant for MLH1 pathogenic variant carriers. LS-related small bowel cancers have been demonstrated to have a better prognosis than sporadic small bowel cancers. Aspirin chemoprophylaxis has been demonstrated to significantly reduce the risk of all LS-associated cancers in patients with LS. Video capsule endoscopy has been used to successfully identify small bowel tumours in patients with LS. However, there have been a limited number of studies and the diagnostic yield has been variable (1–8.6%). VCE may have missed some cancers. No small bowel tumours were detected on follow-up surveillance after an interval of 2 years. In addition, a high false positive rate (11%) has been reported, resulting in asymptomatic patients requiring time consuming and invasive tests.

Pancreatic cancer

In a retrospective study, Kastrinos et al estimated pancreatic cancer risks in individuals (n=6342) from 147 families with constitutional MLH1, MSH2 and MSH6 pathogenic variants. Forty-seven pancreatic cancer cases were reported in 31 families: 18 patients had a negative family history (38%). The cumulative risk for pancreatic cancer up to the age of 70 was reported to be 3.7% (95% CI 1.45% to 5.88%). This represented an 8.6-fold increased risk compared with the general population. Most of the pancreatic cases were observed in individuals from MSH2 families (31/47) and MLH1 families (13/47). Only three of the pancreatic cancers were diagnosed in the MSH6 families. The median age of pancreatic cancer diagnosis was 51.5 years (range 19–85).

Prospective lifetime risks stratified by the MMR gene have now also been reported for pancreatic cancer. Moller et al reported a cumulative risk of pancreatic cancer (up to the age of 75) for MLH1 of 6.2% (95% CI 2.6% to 9.8%), for MSH2 of 0.5% (95% CI 0.0% to 1.5%) and for MSH6 of 1.4% (95% CI 0.0% to 4.2%). No pancreatic cancers were observed in PMS2 pathogenic variant carriers. The authors also calculated 5 year overall survival for the LS patients with pancreatic cancer. The poor prognosis of pancreatic cancer patients is well established, and none of the affected carriers was alive at 5 years.

Surveillance for pancreatic cancer has been recommended in high-risk groups (defined as >5%), including patients from familial pancreatic cancer pedigrees with an affected FDR, patients with PJS, and CDKN2A (P16), BRCA2 and MMR pathogenic variant carriers with at least one affected FDR. The goal of screening is to identify and treat early stage pancreatic cancers (T1N0M0) and high-risk precursor lesions, high-grade pancreatic intraepithelial neoplasia (PanIN-3) and intraductal papillary mucinous neoplasia (IPMN) with high-grade dysplasia. Signoretti et al conducted a systematic review and meta-analysis of 16 pancreatic surveillance studies in high-risk groups. A relatively low diagnostic yield of pancreatic cancers and relevant precursor lesions was reported (3.3%) using endoscopic ultrasound and MRI as first line screening tests. A significant proportion (25%) of the screen-detected cancers were also unresectable or metastatic. Some patients underwent surgery for precursor lesions and were not found to have high-risk precursor lesions. A significant morbidity (up to 40%) and mortality (0.5–6%) have been reported for the surgical treatment of suspicious pancreatic findings. Pancreatic surveillance has not been demonstrated to reduce pancreatic-cancer specific mortality in patients with LS.

The cumulative lifetime risk of pancreatic cancer is relatively low for MLH1 pathogenic variant carriers, and low (<5%) for
MSH2, MSH6 and PMS2 pathogenic variant carriers. Aspirin chemoprophylaxis has also been demonstrated to significantly reduce the risk of all LS-associated cancers in patients with LS. The utility of pancreatic screening remains unproven, and there is a danger of overtreatment with patients undergoing surgery for benign or low-risk lesions. There is a significant morbidity and mortality associated with pancreatic surgery.

**LYNCH-LIKE SYNDROME**

We recommend that deficient MMR tumours without hypermethylation/BRAF pathogenic variant and no pathogenic constitutional pathogenic variant in MMR genes should undergo somatic tumour testing with a CRC gene panel.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 100% agreement.

We recommend that if double somatic MMR pathogenic variants are identified, manage proband and their FDRs based on the FHCC.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 95% agreement.

We suggest that if no or one somatic pathogenic variant is identified, the proband and their FDRs should be managed as per LS.

(GRADE of evidence: low; Strength of recommendation: weak)

Consensus reached: 100% agreement.

LSL describes a subgroup of patients with CRC or other LS-related tumours that manifest MMR deficiency (microsatellite instability (MSI) and/or loss of MMR protein expression) that is neither explained by somatic MLH1 promoter hypermethylation, BRAF pathogenic variant or a detectable pathogenic constitutional variant in an MMR gene or EPCAM LCS cases, and therefore cannot be readily assigned to either the sporadic or inherited MMR deficiency categories, respectively.

Combining data from published studies, using current diagnostic approaches, an estimated 59% (95% CI 55% to 64%) of dMMR CRC cases are unexplained and categorised as LLS.

**Cancer risks**

Three studies have investigated CRC risks for patients with LLS and their relatives. Overbeek et al compared the characteristics of 76 families with a constitutional variant in a DNA MMR gene with those of 18 families with unexplained dMMR tumours. Although the mean age of CRC onset of the index case was comparable at 44 years, a significantly higher proportion of the families with LS fulfilled the Amsterdam II criteria compared with the families with LLS (66% vs 11%; p<0.0001).

Rodríguez-Soler et al were the first to quantify the risk of CRC for FDRs of CRC cases with LLS. This population-based study examined the risk of CRC in the FDRs of CRC cases with LLS, LS and sporadic pMMR tumours. The authors found the highest risk of CRC for the FDRs of the LS cases (standardised incidence ratio (SIR) for LS 6.04, 95% CI 3.58 to 9.3), an intermediate risk for the FDRs of LS cases (SIR for LLS 2.12, 95% CI 1.16 to 3.56), and the lowest risks for the FDRs of sporadic pMMR CRC cases (SIR for sporadic 0.48, 95% CI 0.27 to 0.79; p<0.001).

In a large family cohort study, Win et al subsequently confirmed that FDRs of LLS cases have a higher CRC risk compared FDRs of sporadic pMMR CRC cases, but a lower risk compared with FDRs of LS cases.

Compared with FDRs of sporadic pMMR CRC cases, a higher risk of CRC was estimated for FDRs of LLS cases (HR 2.06, 95% CI 1.59 to 2.67), and an even higher risk for the FDRs of LS cases (HR 5.37, 95% CI 4.16 to 6.94).

**Potential aetiologies**

Studies have shown that up to 70% of patients with LLS have double (biallelic) somatic pathogenic variants in the MMR genes. In a study of 25 LLS cases with unexplained immunohistochemical absence of the MLH1 or MSH2 protein, Mesenkamp et al identified biallelic somatic pathogenic variants (one pathogenic sequence variant and loss of heterozygosity) in 13 cases (52%: 8/18 in MLH1 and 5/7 in MSH2). Other studies have subsequently confirmed the presence of a high proportion of biallelic somatic pathogenic variant LLS cases. In a study of 36 LLS cases, Geurts-Giele et al found double somatic pathogenic variants in 21 cases (58%; 16/21 in MLH1 and 5/12 in MSH2).

In a study of 32 LLS cases, Haraldsdottir et al identified double somatic pathogenic variants in 22 cases (69%; seven for MLH1, 11 for MSH2, three for MSH6 and one for PMS2).

Biallelic somatic pathogenic variants may be in the form of point pathogenic variants coupled with loss of heterozygosity, or another point pathogenic variant. Double variant somatic pathogenic variants are likely to be in trans (one on each allele), and therefore if identified provide a potential explanation for the tumour MMR deficiency in patients with LLS. A recent study has shown that a significant proportion of LLS cases may also be explained by false positive screening results, caused by an incorrect interpretation of MMR immunohistochemistry (IHC) results. The authors found discordant findings on IHC and MSI in six out 32 LLS cases (19%). However, there remains a possibility that some patients with LLS may actually have LS, due to constitutional pathogenic variants in the MMR genes that are not detected using current diagnostic methods, for example, pathogenic variants within regulatory/promoter regions and complex structural variants. Finally some patients with LLS may carry constitutional or somatic pathogenic variants in other genes, which may explain the deficient MMR tumour phenotype and/or occur in conjunction with somatic pathogenic variants in the MMR genes. Jasen et al investigated 62 LLS cases using gene panel sequencing including the POLE, POLD1 and MMR genes. The authors found somatic (n=7) or constitutional variants (n=2) in the POLE/POLD1 exonuclease domain in nine tumours (14.5%), which showed an ultramutated phenotype. Six of these cases also were found to carry somatic MMR variants. In a study by Morak et al, MUTYH diagnostic testing was carried out in 85 patients with LLS and biallelic constitutional pathogenic variants were found in one patient (1.18%). Consequently, LLS cases have been found to have heterogeneous aetiologies, ranging from biallelic somatic pathogenic variants to unidentified constitutional pathogenic variants in MMR genes. This is likely to explain the intermediate cancer risks, which have been reported for the FDRs of LLS cases, and has implications for the clinical management of LLS families.

Thus double somatic pathogenic variants in the MMR genes may explain up to 70% of LLS cases. Therefore, tumour sequencing of the DNA MMR genes should be undertaken in LLS cases, as this would help guide genetic counselling and the management recommendations for LLS cases and their FDRs (figure 3). If double somatic pathogenic variants are identified, we would recommend that these patients and their FDRs be managed based on the FHCC, and not as LS. However, if no somatic pathogenic variants or only one somatic pathogenic variant or loss of heterozygosity of one allele is identified we...
Guidelines

Figure 3  Management of Lynch-like syndrome. CRC, colorectal cancer; FDRs, first degree relatives; FHCC, family history of colorectal cancer; MMR, mismatch repair.

would recommend that these cases and the FDRs be managed as potential LS cases and follow colorectal surveillance guidelines for LS. This is based on the possibility that these cases could have LS due to an unidentifiable constitutional pathogenic variant in an MMR gene. These recommendations are consistent with the NCCN (National Comprehensive Cancer Network) guidelines170 and may help reduce variability in practice in the UK.171

EARLY ONSET CRC

We recommend that in patients under 30 years of age with dMMR CRC, an LS constitutional panel test should be performed, followed by tumour testing for somatic testing if constitutional testing is negative.

(GRADE of evidence: low; Strength of recommendation: strong)
Consensus reached: 91% agreement.

We recommend that in patients under 30 years of age with pMMR CRC, a constitutional CRC multiple gene panel test should be performed.

(GRADE of evidence: low; Strength of recommendation: strong)
Consensus reached: 100% agreement.

Several studies have sought to define the prevalence and spectrum of constitutional cancer predisposition gene pathogenic variants among patients diagnosed with early-onset CRC (EOCRC), defined most commonly as CRC onset below age 50 years.172-174

In a retrospective cohort of very young patients (age 35 or younger) with CRC referred for genetic evaluation, syndrome-specific genetic testing guided by patient phenotype and family history, identified highly penetrant CRC syndromes in 33% of patients (67 of 193; 23 LS, 22 pathogenic variant-negative LS, 16 FAP, two constitutional MMR deficiency (CMMRD), two MAP, and one Li-Fraumeni syndrome). LS (23.3%; including patients with unexplained MMR deficiency, variants of uncertain significance, and incomplete LS investigations) and FAP (8.3%) were the most commonly identified CRC syndromes.173

In a prospective cohort, panel testing of 25 cancer predisposition genes (APC, ATM, BARD1, BMP11A, BRCA1, BRCA2, BRIPI, CDH1, CDK4, CDKN2A, CHEK2, EPCAM, MLH1, MSH2, MSH6, MUTYH, NBN, PALB2, PMS2, PTEN, RAD51C, RAD51D, SMAD4, STK11, TP53) was performed in 450 patients diagnosed with early-onset CRC (aged <50). Forty-eight patients (10.7%) had dMMR tumours and 402 (89.3%) had pMMR tumours. Sixteen percent of patients (72 of 450) were found to have a pathogenic variant in at least one cancer predisposition gene.172

Of the 48 patients with dMMR tumours, 40 patients (83.3%) had at least one constitutional pathogenic variant: 37 patient had LS (13 MLH1, 16 MSH2, one MSH2/monallelic MUTYH, two MSH6, five PMS2), one patient had the low penetrant APC c.3920T>A, p.I1307K variant and a PMS2 variant of uncertain significance, and two patients had biallelic MUTYH pathogenic variants. Of the 402 patients with pMMR tumours, 32 (8%) had at least one constitutional pathogenic variant in a cancer predisposition gene: nine (2.2%) patients had pathogenic variants in high penetrant CRC predisposition genes (five APC, one APC/PMS2, two biallelic MUTYH pathogenic variants), 13 patients had pathogenic variants in genes not traditionally linked with CRC susceptibility (three ATM, one TM/CHEK2, two BRCA1, four BRCA2, one CDKN2A and two PALB2), three patients had the low penetrant APC c.3920T>A, p.I1307K variant, and seven patients had monoallelic MUTYH pathogenic variants.172

A constitutional pathogenic variant in a high penetrance CRC gene was identified in 22% of patients with CRC diagnosed between 20–29 years (4/18): two MSH2 pathogenic variants in patients with dMMR tumours and two APC pathogenic variants in patients with pMMR tumours.172 Of note, in those patients with a pathogenic variant identified in a high penetrance gene over the age of 30 years, all either had a significant family history, multiple primaries or exhibited dMMR on tumour testing.

A recent retrospective evaluation of patients with early-onset CRC (aged <50) reported comparable findings. In this study,
In conclusion

- The prevalence of cancer predisposition gene pathogenic variants in patients with early onset CRC (age <50) has been reported to range from approximately 15% to 20%. LS is the most common genetic diagnosis, accounts for up to 68% of cancer predisposition pathogenic variants in patients with early onset CRC (age <50 years), and the vast majority will have dMMR tumours.

- In patients with pMMR tumours, the most common genetic diagnosis is an adenomatous polyposis syndrome (FAP and MAP), and the prevalence of associated constitutional pathogenic variants has been reported to reach the 10% threshold for patients diagnosed with CRC below the age of 30.

- In patients with pMMR tumours between the age of 30 and 50 years at diagnosis of CRC, a genetic diagnosis may be suspected on the basis of FHCC or other clinical parameters. Altering the threshold for genetic testing in this patient cohort also needs to be balanced between clinical need and the ethical and logistical considerations of population testing.

Colonoscopic surveillance in EOCR

We suggest that people diagnosed with CRC under the age of 50 years, where hereditary CRC syndromes have been excluded, undergo standard post-CRC surveillance for 3 years, then continue 5 yearly colonoscopic surveillance until the age they are eligible for national screening.

(GRADE of evidence: very low; Strength of recommendation: weak)
Consensus reached: 80% agreement.

The risk of metachronous CRC is highest during the 36 months after surgery; subsequently this risk decreases. The most plausible explanation is that many early, apparently metachronous cancers are actually due to prevalent cancers or advanced adenomas missed at the time of the primary CRC diagnosis. As a consequence, new guidelines (in development) for sporadic CRC may recommend that standard post-CRC colonoscopic surveillance cease after 3 years.

Recently published, large, population-based cancer registry studies, including ones that specifically excluded patients with LS, recommend closer surveillance in high-risk populations, however.

There have been no studies specifically assessing metachronous CRC risk or the role of colonoscopic surveillance in EOCR patients. However, when patients with hereditary CRC syndromes are excluded from CRC cohorts there is no evidence that metachronous CRC risk is significantly different from the sporadic population. Nevertheless, it may not be appropriate to discharge EOCR cases from surveillance after 3 years in the absence of published surveillance outcome data. Thus, it may be considered prudent to offer 5 yearly colonoscopic surveillance until they are eligible for national screening in patients where LS and polyposis have been excluded.

SERRATED POLYPOSIS SYNDROME

Definition

- 2019 Updated WHO clinical criteria for the diagnosis of serrated polypsis

Criterion 1

- At least 5 serrated lesions/polyps proximal to the rectum all being ≥ 5 mm in size, with 2 or more ≥ 10 mm in size

Criterion 2

- More than 20 serrated lesions/polyps of any size distributed throughout the large bowel, with at least 5 proximal to the rectum

Any histological subtype of serrated lesion/polyp (hyperplastic polyp, sessile serrated lesion without or with dysplasia, traditional serrated adenoma, and unclassified serrated adenoma) is included in the final polyp count. The polyp count is cumulative over multiple colonoscopies

Diagnosis

We recommend a diagnosis of SPS should be made in accordance with the new WHO 2019 criteria for SPS. Since causative gene pathogenic variants for SPS have not been identified, a definitive diagnosis of SPS should be phenotype-driven.

(GRADE of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 95% agreement.

Other intestinal polyposis syndromes may present with serrated lesions. If (i) the patient is under 50 or (ii) there are multiple affected individuals within a kindred or (iii) there is dysplasia within any of the polyps, then other polyposis syndromes should be excluded by gene panel testing before making a definitive diagnosis of SPS.

(GRADE of evidence: very low; Strength of recommendation: weak)
Consensus reached: 90% agreement.

We recommend the cumulative number of serrated polyps from all endoscopic examinations should be used when applying the WHO 2019 diagnostic criteria for SPS.

We recommend the cumulative number of serrated polyps from all endoscopic examinations should be used when applying the WHO 2019 diagnostic criteria for SPS.

(GRADE of evidence: moderate; Strength of recommendation: strong)
Consensus reached: 94% agreement.

SPS most probably comprises a phenotypically and genetically heterogeneous group of diseases. The phenotypic criteria for a diagnosis of SPS have recently been revised by the WHO in 2019 to include the following: (1) at least five serrated polyps proximal to the rectum, all >5 mm in size with at least two >10 mm in size; (2) at least 20 serrated polyps (of any size) with at least five located proximal to the rectum.174 Fulfilment of either criterion is sufficient for a diagnosis of SPS. Importantly, the previous WHO 2010 criterion 2, which required the presence of just one serrated lesion in a patient who has an FDR with SPS, has been removed. Given that the prevalence of serrated lesions in the Western population may be up to 39%, most of which will be sporadic, the chances of a false positive diagnosis using this criterion were high.

The prevalence of SPS in the West is generally considered to be around 1:3000 in screening populations. This may change as awareness of the condition among clinicians increases. Egoavil et al.179 found that patients with multiple serrated polyps but failing to fulfil the WHO criteria had a similar risk of CRC as those who did fulfil the criteria. Dovetailing with this to some degree (and possibly explaining this), Crowder et al.180 concluded that SPS was underdiagnosed due to failure to consider cumulative polyp numbers (rather than at a single episode). By evaluating “cumulative pathology” in a cohort of 927 consecutive patients undergoing colonoscopy, they found that up to 1.8% of patients with a serrated lesion at first colonoscopy eventually fulfilled the criteria for SPS.

Patients with SPS have an overall lifetime risk of CRC of approximately 7–70%.181 and there is an increased risk in FDRs of patients with SPS. The risk of CRC thus may be reduced with appropriate management.181 182 There is an increased risk of CRC in FDRs of patients with SPS, but it does not follow a classical Mendelian pattern of inheritance. A single causative gene has not been identified and there is phenotypic overlap with some well characterised intestinal polyposis syndromes (MUTYH-associated polyposis, hereditary mixed polyposis syndrome, attenuated FAP). Most, but not all, of the tumours arising in SPS follow the serrated neoplasia pathway characterised by microsatellite instability (due to MLH1 promoter methylation) and BRAF pathogenic variant.183 This raises the possibility of accelerated tumorigenesis as well as phenotypic overlap with LS. Recently, constitutional truncating pathogenic variant in RNF43 pathogenic variant has been identified and shown to segregate with phenotype in one kindred with SPS.184 185 The same pathogenic variant was also identified in two patients in a separate study.

In the absence of a single causative gene (and the possibility that some forms of SPS may be polygenic in nature), the diagnosis of SPS should be phenotype-driven. Given the overlap with other well-defined syndromes, these should be excluded in all patients in whom a diagnosis of SPS is made. A specific gene panel for SPS may include testing for RNF43 and GREM1; however the frequency of these pathogenic variants is too low to recommend routinely.186

Consensus reached: 94% agreement.

Colonoscopic surveillance is currently recommended following detection and resection for adenomas in the colorectum due to an increased risk of advanced adenomas and CRC in such patients; however, there is another major pathway to CRC “serrated pathway” which accounts for 15–30% of CRC and has serrated lesions as cancer precursors. SPS is common in bowel cancer screening programmes which use guaiac faecal occult blood test (gFOBT) or FIT as a screening test, with estimates of SPS prevalence ranging from 1:150 to 1:300.187 188 A recent Spanish FIT based cohort followed up all their patients with proximal serrated polyps, tripling the number of additional cases of SPS, for a final prevalence of 1:100.189 Therefore, especially when using FIT in bowel cancer screening, colonoscopists should be alert to a diagnosis of SPS.

The BSG position statement on serrated polyps in the colon and rectum recommended 1–2 yearly surveillance for patients meeting the WHO criteria for serrated polyposis syndrome.190 This recommendation was on the basis that in early cohorts future risk of CRC was elevated as much as 7% at 5 years191 192; however in larger cohorts with rigorous surveillance performed every 1–2 years, with all lesions larger than 5 mm in size resected, at academic centres, the risk appeared much lower with CRC only diagnosed at 1.9 cases per 1000 years of patient follow-up.181 182 In a US study following up SPS patients which extended surveillance intervals for SPS patients to 2 years after colon clearance with no lesion ≥10 mm found at surveillance, no cancer developed or surgery was needed.193 A similar multicentre European study that investigated surveillance after colonic clearance to 1 or 2 year follow-up dependent on lesion size (≥10 mm), number and pathology also showed no difference in advanced neoplasia detection with a 2 year surveillance interval once the colon was cleared.194

No new data directly relevant to this area have been published since the BSG position statement. However, a study of patients with multiple serrated polyps and adenomas, not fulfilling the criteria for SPS, also noted that their risk for CRC was equivalent to patients who met the WHO definition of SPS, and that their FDRs had a comparable risk of CRC.179

Colonoscopic surveillance in unaffected FDRs of patients with SPS

We recommend all FDRs of patients with SPS on the basis of the new WHO 2019 SPS criteria 1 or 2 should be offered an index colonoscopic screening examination at the age of 40 years or 10 years before the diagnosis of the index case.

(Grade of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 89% agreement.

We suggest all FDRs of serrated polyposis patients have a surveillance examination every 5 years unless polyph burden indicates an examination is required earlier according to post-polypectomy surveillance guidelines.

(Grade of evidence: low; Strength of recommendation: strong)

Consensus reached: 84% agreement.

SPS may be a familial condition with between 1.3% and 7.7% of index cases having an FDR who meets the original WHO criteria 1 or 3 for SPS.195 196 A further 14.3%–24.4% of FDRs meet original WHO criterion two for SPS. Current guidelines recommend a one-off screening examination of all FDRs to detect these familial SPS cases; however, the majority of FDR will not meet the original WHO criteria for SPS after this initial examination. The risk of CRC for FDRs of SPS patients is estimated to be substantially elevated above that of the general
population by three- to fivefold.\textsuperscript{179, 197, 198} This risk may also apply to those with multiple serrated polyps not meeting WHO criteria for SPS. It is unclear how this risk is distributed in the SPS FDR population.

In three series the mean or median age at diagnosis for CRC in FDRs of SPS patients ranged from 5.5 to 62. CRC was very rare in those aged less than 40 years with the youngest case being aged 25. Authors have recommended a screening examination starting at 35 or between 40 and 50 for FDRs of SPS cases, or starting 5–10 years before the index case\textsuperscript{179, 195–197}.

In a follow-up study of 78 patients, those who did not meet any WHO criteria for SPS seem to have low risk for subsequent polyp or advanced neoplasia development; however, those who meet WHO criteria 2 do seem to develop polyps with five of 14 patients developing three or more adenomas and two others of the 14 developing a serrated lesion \(\geq 10\text{mm}\) in size.\textsuperscript{196} High rates of polyps in patients who met WHO 2 criteria were also seen by Hazewinkel et al.\textsuperscript{195} Therefore a significant proportion of WHO 2 SPS patients would meet criteria for 3 yearly surveillance according to current guidelines. No cancers developed. No data are available to look at the colonoscopic yield in FDRs of index patients who meet WHO criteria 2, that is, a second degree relative of a patient with SPS WHO 1 or 3, although the risk of CRC in second degree relatives of index cases of SPS is only slightly elevated (SIR 1.38, 95%CI 1.01 to 1.91).\textsuperscript{198}

The WHO definition of SPS has recently changed with criteria 2 abandoned completely so that FDRs with serrated polyps are not considered to meet the definition for SPS. Therefore a blanket recommendation for colonoscopic surveillance for FDRs of SPS every 5 years based on the three- to fivefold increase in SIR of CRC is suggested from age 40 or 10 years before the diagnosis for the index case, with more frequent surveillance in line with sporadic recommendations if additional polyps are detected.

FDRs of patients with multiple serrated polyps (10 or more polyps in total of which 50% are serrated) but who do not meet the 2019 WHO criteria for SPS might be considered for similar surveillance approaches for FDRs.

The risk of CRC in FDRs of SPS patients is sufficiently elevated that they should be offered colonoscopic screening and regular surveillance.

**MULTIPLE COLORECTAL ADENOMAS**

We suggest an individualised approach to germline testing of patients with MCRA (defined as having 10 or more metachronous adenomas). Consider this testing for:

- Patients under 60 years of age with lifetime total of \(\geq 10\) adenomas;
or
- Patients from 60 years of age with lifetime total of:
  - \(\geq 20\) adenomas, or
  - \(\geq 10\) adenomas and an FHCC or polyposis

*(GRADE of evidence: low; Strength of recommendation: weak)*

Consensus reached: 91% agreement.

We suggest that patients with a finding of 10 or more polyps (adenomas or serrated lesions) should, at their next colonoscopy, have a high-quality colonoscopic assessment with panaromic dye spray in order to accurately define the multiple polyp phenotype.

*(GRADE of evidence: very low; Strength of recommendation: weak)*

Consensus reached: 89% agreement.

We suggest that the endoscopic management of patients with 10 or more metachronous adenomas, without MUTYH or APC gene mutations, should be individualised according to phenotype.

*(GRADE of evidence: very low; Strength of recommendation: weak)*

Consensus reached: 91% agreement.

We suggest annual colonoscopic surveillance for patients with 10 or more metachronous adenomas after the colon has been cleared of all lesions >5mm in size. If no polyps 10mm or greater in size are identified at subsequent surveillance examinations the interval can be extended to 2 yearly.

*(GRADE of evidence: very low; Strength of recommendation: weak)*

Consensus reached: 80% agreement.

Patients with multiple adenomas but without classical familial adenomatous polyposis are frequently encountered in clinical practice. Approximately 1.1% of patients undergoing colonoscopy in the English bowel cancer screening programme have 10 or more adenomas.\textsuperscript{199} There is an association of adenoma multiplicity with metachronous advanced adenoma and/or CRC risk, with the degree of risk correlating with increasing adenoma number and size. With less than three adenomas as reference, Cubiella et al.\textsuperscript{(n=5401)} reported a statistically increased risk for AN when three or four adenomas (14.8%), and also when five to nine adenomas (18.4%), were present at index colonoscopy.\textsuperscript{200}

In the largest study to date of individuals with 10–19 adenomas,\textsuperscript{201} 3789 patients with 10 or more colorectal polyps underwent constitutional testing of the prevalence of pathogenic variants with a 17-gene panel. The diagnostic yield of pathogenic variants remained above 5% in all ages and cohorts, despite a decrease with age. In the multiple adenoma cohort, the yield was higher in those patients with a personal or family history of cancer. In 1342 patients with 10–19 adenomas, 7.8% had a pathogenic variant in one of the panel genes, with 2.2% in “traditional” polyposis predisposition genes, and 2.8% in MMR genes. Thus an unbiased multi-gene panel test approach may be associated with a higher diagnostic yield.

In a study of 7225 individuals with MCRA and oligopolyposis, pathogenic variants in APC or common European founder pathogenic biallelic variants in MUTYH were identified in 87/970 (9%) individuals with 10–19 adenomas and 559/3253 (17%) individuals with 20–99 adenomas.\textsuperscript{202} There was an incremental increase in the odds of a pathogenic variant with an increasing number of adenomas and earlier age at adenoma diagnosis, particularly under the age of 50 years.

In addition to APC and MUTYH there is an evolving range of other multiple adenoma susceptibility genes including NTHLI, GREM1, POLE, POLD1 and MSH3. Spier et al in 2015 demonstrated POLE pathogenic variants in 1.5% of individuals with greater than 20 synchronous or 40 metachronous adenomas although this percentage increased with a family history.\textsuperscript{202} There is also some evidence of polygenic risk contributing to this attenuated polyposis phenotype.\textsuperscript{203}

In summary, it is advisable to consider a diagnosis of a highly penetrant syndrome in patients with multiple adenomas. A nuanced approach to patient selection for constitutive testing should incorporate patient age and personal and family history.

With regard to colonoscopic surveillance in the multiple adenoma patient population, the GDG suggests constitutional gene testing to rule out known hereditary syndromes. Subsequently an individualised approach should be taken depending on adenoma size and number, with the goal of clearance of colorectal adenomas >5 mm in size. However, surgical resection should be considered where surveillance is not feasible.
In patients with 10–99 adenomas without APC or MUTYH pathogenic variants, the clinical phenotype is similar to that of attenuated polyposis, with both an upper GI (21% with duodenal adenomas) and colorectal phenotype. In a study of 83 patients also of 10–99 adenomas without APC or MUTYH pathogenic variants from the UK and Holland, the upper GI phenotype was reported in a minority of patients (9.6%), all of whom had Spigelman I or II disease (ie, early stage).

For FDRs of MCRA patients, clinicians may consider a colonoscopic assessment of risk at the time of referral, and/or a repeat assessment at age 50 years. This is in order that patients who may be at risk of high penetrance cancer predisposition syndromes may be effectively identified (for example, the children of individuals with unidentified mosaic pathogenic variants of the APC gene). However the GDG did not reach consensus on this recommendation, and recommend that multicentre outcome data in these patients be collected and published to help guide future recommendations.

FAMILIAL ADENOMATOUS POLYPOSIS
FAP is defined by the presence of pathogenic variants in the APC gene with a prevalence of about 1/8500. It is a dominantly inherited multisystem cancer predisposition syndrome with a characteristic phenotype characterised by colorectal and upper GI polyposis.

Colorectal surveillance in FAP
We recommend that colonic surveillance should normally commence at the age of 12–14 years in those confirmed to have FAP on predictive genetic testing.

(GRADE of evidence: low; Strength of recommendation: strong)
Consensus reached: 100% agreement.

We suggest that for those with FAP, intervals between surveillance colonoscopy may be individualised depending on colonic phenotype every 1–3 years.

(GRADE of evidence: low; Strength of recommendation: weak)
Consensus reached: 94% agreement.

We suggest that colonoscopy screening is performed for individuals who have an FDR with a clinical diagnosis of FAP (ie, “at-risk”) and in whom an APC mutation has not been identified, starting at the age of 12–14 years, and should continue on 5 yearly surveillance until either a clinical diagnosis is made and they are then managed as FAP, or they reach the age at which they can enrol in national screening.

(GRADE of evidence: very low; Strength of recommendation: weak)
Consensus reached: 95% agreement.

Current international guidelines recommend starting colonoscopy surveillance or screening at the age of 12–14 years, after diagnostic genetic testing has been performed in at-risk children. There are no new data to recommend any change to this recommendation of age at which to start. The risk of CRC under the age of 20 years is very small and under 15 years extremely rare. In patients with FAP-related symptoms such as rectal bleeding, diarrhoea or mucus discharge should lead to a colonoscopy at any age, particularly in those with a constitutional pathogenic variant at codon 1309, which is associated with a greater risk of a more severe colorectal phenotype.

Colonoscopic surveillance has been shown to lead to a reduction in CRC and CRC-associated mortality. Data detailing the results of polyposis registries have shown that in symptomatic patients the incidence of CRC was 50–70% compared with 3–10% in those that were identified by registry initiated surveillance.

Surveillance and prophylactic intervention has reduced CRC associated mortality.

Colonoscopic surveillance enables assessment of adenoma burden and distribution, which can guide the timing of and type of prophylactic surgery required. Previous guidelines recommended the use of flexible sigmoidoscopy. However, most endoscopic procedures are performed under general anaesthesia in children and young teenagers; therefore a full colonoscopy is recommended in an affected patient, to better determine their phenotype, as polypl distribution is not uniform and the rectum and sigmoid may be normal despite the presence of more proximal adenomas. It is not recommended therefore for flexible sigmoidoscopy to be used routinely in screening/surveillance in FAP.

Current guidelines advocate annual endoscopic assessment. There is no evidence for accelerated carcinogenesis in FAP and the rather scant data that are available do not indicate a rapid increase in polyp number in teenagers/young adults. Therefore it does not seem logical to mandate an annual assessment for all affected patients, particularly given that there is significant variability in colorectal phenotype. Those with an attenuated phenotype (<100 adenomas) may not require such frequent colonoscopic surveillance as those with a classical phenotype (>100 adenomas). In addition, if an individual only has adenomas of 1–2 mm their surveillance could perhaps be longer than those with larger polyps, for example, 8–9 mm. Personalising surveillance interval according to phenotype, 1–3 yearly would appear safe as long as families are not lost to follow-up and this would concord with current paediatric guidelines. Extending the interval, however, should be for those with an attenuated phenotype, in the setting of good quality colonoscopy with robust systems in place to ensure appropriate recall.

There are a few patients with a particularly attenuated phenotype. In this group, primary endoscopic management by surveillance and polypectomy may be considered either to defer surgery or possibly to avoid the need for surgery altogether. However, there are no robust data to support this approach.

At risk patients, where predictive genetic testing is not possible, should be screened by colonoscopy every 5 years from the age of 12–14 years. If adenomas are identified, the patient should undergo repeat colonoscopy at a frequency depending on the colorectal phenotype as described above. If no phenotype has been observed by the age of 50 years, FAP is unlikely and so the patient could be discharged from routine colonoscopic screening to have continued screening under the auspices of the national bowel cancer screening programme.

Upper GI surveillance in patients affected with FAP
We recommend upper GI surveillance for FAP patients starting at the age of 25 years.

(GRADE of evidence: low; Strength of recommendation: strong)
Consensus reached: 90% agreement.

We suggest that for those considered at risk, where predictive genetic testing is not possible, screening with upper GI endoscopy is not routinely recommended but should be started if/when a clinical diagnosis of FAP is made based on colorectal phenotype.

(GRADE of evidence: very low; Strength of recommendation: weak)
Consensus reached: 89% agreement.

Lifetime risk of duodenal polyposis approaches 100% in FAP. The absolute lifetime risk of developing duodenal cancer...
in FAP is estimated to be around 5%. Because of this, surveillance has been recommended and survival benefit for those diagnosed with duodenal cancer by surveillance compared with those who presented symptomatically has been demonstrated.

The Spigelman classification (Table 3) is the system that is most widely used for staging non-ampullary duodenal disease; it has been shown to correlate with cancer risk and is recommended to determine surveillance intervals. There is debate about how to incorporate ampullary disease in this classification system. A staging system for ampullary disease has been proposed which in one series correlated to the development of ampullary cancer. A surveillance interval determined by the combination of these staging systems may be the most helpful and reliably replicated clinical means of managing duodenal surveillance (see Figure 1). A duodenoscope is required to reliably assess and/or biopsy the periampullary region and ampulla itself. Although there are reports of chromoendoscopy increasing duodenal adenoma detection, its utility in clinical practice is not established. It may increase the number of adenomas detected but there remains no evidence that dye spray increases the pick-up of larger, more clinically meaningful lesions. There are no data to suggest that it alters the need for endoscopic or surgical intervention. The concern is that it may increase the pick-up of small lesions and may artificially “upstage” the duodenal disease but without reflecting a higher cancer risk, compared with white light endoscopy. Recent data suggest that it is polyp size and the presence of high-grade dysplasia that are the most important predictors of cancer risk, polyp multiplicity. It should be noted that the Spigelman classification system was developed and validated using white light endoscopy, without high definition.

The role of endoscopic therapy in the duodenum and ampullary is not well established. It has an acceptable safety profile; however, there are no long-term data to demonstrate it reduces cancer risk or indeed prevents or delays the need for prophylactic surgery. Nevertheless, it is widely performed in centres dealing with a large cohorts of patients with FAP. It would be most prudent for patients being considered for endoscopic therapy for duodenal disease to be referred to their local specialist centre, so that assessment and work up to decide as to the appropriateness of endoscopic and surgical intervention can be performed.

The data regarding endoscopic management of ampullary disease are rather scant. However, it appears to be less safe with respect to haemorrhage, pancreatitis and perforation, morbidity rates up to 43% and with high rates of recurrence, up to 58%. It cannot be routinely recommended. Referral to a specialist hepatopancreaticobiliary (HPB) centre should be made for those in whom endoscopic ampullectomy is being considered.

Gastric lesions are also common in adult patients with FAP. Fundic gland polyps (FGPs) are seen in up to 80% of patients with FAP and although there is some debate they are likely to be an entirely benign entity without malignant potential. Gastric adenomas are being seen more commonly, as indeed is gastric cancer. Although historically gastric cancer risk was not thought to be elevated in patients with FAP, there are reports that gastric adenomas and cancer are becoming an important clinical problem. There are no data published regarding outcomes of endoscopic therapy for gastric adenomas in FAP. Referral to a specialist centre for assessment and management seems prudent given the lack of evidence and absence of consensus guidelines.

### Table 3: Staging the duodenum and ampulla and recommended OGD surveillance intervals

<table>
<thead>
<tr>
<th>Points allocated</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of polyps</td>
<td>1–4</td>
<td>5–20</td>
<td>&gt;20</td>
</tr>
<tr>
<td>Polyp size (mm)</td>
<td>1–4</td>
<td>5–10</td>
<td>&gt;10</td>
</tr>
<tr>
<td>Histological type</td>
<td>Tubular</td>
<td>Tubulovillous</td>
<td>Villous</td>
</tr>
<tr>
<td>Degree of dysplasia</td>
<td>Mild</td>
<td>Moderate</td>
<td>Severe</td>
</tr>
</tbody>
</table>

OGD, oesophago-gastro-duodenoscopy.

<table>
<thead>
<tr>
<th>Total points</th>
<th>Spigelman stage</th>
<th>Recommended follow-up interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>5 years</td>
</tr>
<tr>
<td>1–4</td>
<td>I</td>
<td>3 years</td>
</tr>
<tr>
<td>5–6</td>
<td>II</td>
<td>Annual and consider endoscopic therapy</td>
</tr>
<tr>
<td>7–8</td>
<td>III</td>
<td>6–12 months and consider endoscopic or surgical therapy</td>
</tr>
<tr>
<td>9–12</td>
<td>IV</td>
<td>6–12 months and consider endoscopic or surgical therapy</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Ampulla size</th>
<th>Normal ampulla</th>
<th>Minor polyposis</th>
<th>Major polyposis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Villous histology</td>
<td>n/a</td>
<td>Less than 1 cm</td>
<td>More than 1 cm</td>
</tr>
<tr>
<td>Degree of dysplasia</td>
<td>n/a</td>
<td>Mild</td>
<td>Moderate or severe</td>
</tr>
</tbody>
</table>

### Conventional hypertrophy retinal pigmentation epithelium

We suggest that patients with congenital hypertrophy retinal pigmentation epithelium (CHRPE) be referred for a specialist ophthalmic review. Patients with bilateral and multiple CHRPE lesions should be referred for screening for FAP and considered for genetic testing and colonoscopy.

(Grade of evidence: low; Strength of recommendation: weak)
Consensus reached: 100% agreement.
Up to two thirds of patients with FAP have CHRPE identified at ophthalmoscopy, compared with a prevalence in the general population of 1–4%. CHRPE lesions associated with FAP are most often multiple, bilateral (in 86% of cases) and oval or pisciform in shape. Multiple retinal lesions have a very high specificity as a phenotypic marker for FAP. Referral for a specialist ophthalmic review will assist in characterising those with lesions which need to be considered screening for FAP. Such patients should be referred to a specialist centre for consideration of screening for FAP by genetic testing and thereafter colonoscopy, age 12–14 years, or at time of diagnosis if diagnosed at an older age.

### FAP and surgery

We recommend that for patients with FAP who are undergoing colonoscopy surveillance, relative indications for surgery are polyps >10mm in diameter, high grade dysplasia within polyps and a significant increase in polyp burden between screening examinations.

(Grade of evidence: low; Strength of recommendation: strong)
Consensus reached: 82% agreement.
For most patients the choice of surgery will be between total colectomy with ileorectal anastomosis (IRA) and proctocolectomy and ileal pouch anal anastomosis (IPAA). The choice of surgery will depend on rectal polyg number, size and presence of high-grade dysplasia, genotype and the functional consequences of the surgical procedure. IPAA should also be considered.
for those patients who are likely to be poorly compliant with follow-up surveillance.

Total proctocolectomy with end ileostomy can be considered for patients with poor sphincter function, incontinence, distal rectal cancer, cancers requiring radiation, or for those who desire to avoid the functional consequences of an ileal pouch.

In patients with FAP, colon cancer will inevitably develop if the colon is not removed. Total colectomy will prevent colon cancer in FAP patients. Prophylactic surgery can usually be planned at a time which is suitable to the patient, based on the risk of cancer as assessed colonoscopically. The timing and choice of surgical procedure should take into account the educational, social, family planning and emotional development of the patient and their reliability for attending follow-up evaluations.

Indications for surgery include polyps >10 mm diameter, polyps with high-grade dysplasia, and marked increases in polyp number between examinations. Symptoms from polyps should be rare in those undergoing regular screening and surgery should be performed before symptoms from polyps develop.

Total colectomy with ileorectal anastomosis (TAC-IRA) can be offered to patients with relative rectal sparing (<20 polyps) if all rectal adenomas are <5 mm in diameter and any polyps >5 mm can be endoscopically removed. However, flexibility with regard to a threshold polyp number should be employed with the advent of high-definition or chromo-endoscopic techniques.

The decision to retain the rectum is made based on future rectal cancer risk, polyposis phenotype in the rectum, functional considerations and the genotype.

Bülow et al evaluated 776 patients who had IRA, including 576 before the ileorectal pouch era, and 200 after the ileorectal pouch became available in these centres. The cumulative risk of rectal cancer by Kaplan-Meier analysis was 10% in the pre-pouch era versus 2% in the pouch era.

A cohort study from Church et al of 213 patients with FAP included 165 patients who had rectal-sparing surgery, with 128 of these having <20 polyps and 37 having >20 polyps. The rectal cancer incidence was 1.6% in the patients with <20 polyps, compared with 10.8% in the patients with >20 polyps.

Predicting future rectal excision may also aid decision making about choice of surgery. A study from St Marks hospital of 427 patients who underwent IRA found that by the age of 60 years, half of the patients retained their rectum. Rectal polyp count exceeding 20, APC pathogenic variant codon 1250–1450, colonic polyp count ≥500 and age <25 years at the time of surgery were independent predictors of progressive rectal disease. Church et al also found rectal polyp count >20 to predict future rectal excision.

Genotype can also be utilised specifically when discussing surgical options. A study of four national polyposis registries included 475 polyp patients with a previous colectomy. Cumulative risks of secondary proctectomy 20 years after primary colectomy were 10%, 39% and 61% in the attenuated, intermediate and severe genotype groups, respectively (p<0.03, groups compared separately). Patients with a severe genotype have a high-risk of rectal excision after primary colectomy and may be better directed to proctocolectomy with IPAA. Influence of genotype on survival has also been reported by Newton et al.

For patients undergoing IRA, 1–3 flexible sigmoidoscopies are required as per FAP patients with intact colons.

There have been no randomised trials comparing the functional outcome of IRA versus IPAA. The study by Aziz et al provides a fair summary of the literature. They performed a meta-analysis on 12 non-randomized studies published between 1991 and 2003 and containing over 1000 patients (47% IRA vs 53% IPAA). Bowel frequency, nocturnal defaecation and use of incontinence pads were significantly less in the ileorectal group, although faecal urgency was reduced with the ileal pouch. There was no significant difference between the techniques in terms of sexual dysfunction, dietary restriction or postoperative complications.

The fecundity of women with FAP before operation and after colectomy with ileorectal anastomosis has been reported to be similar to that of the general population. However, fecundity dropped to 54 per cent following proctocolectomy with ileal pouch-anal anastomosis. It is recommended that the significant reduction in female fecundity after IPAA should be communicated to young women with FAP before surgery.

The risk of developing postoperative fertility problems is not associated significantly with the type of surgery, indication for surgery, complications or other comorbid conditions. Postoperative fertility problems appear to be more common among women who had their first surgical procedure at a younger age.

Indications for rectal excision following IRA include: the development of rectal cancer, polyps >10 mm diameter, polyps with high-grade dysplasia, and marked increases in polyp number between examinations. When conversion of an IRA to IPAA is required, functional outcomes appear similar to primary IPAA procedures. Complication rates and pouch failure rates are reported to be similar, but conversion to IPAA will not be possible in a small percentage of patients.

Proctocolectomy with IPAA is the treatment of choice in the presence of: rectal cancer, a large rectal polyp burden (>20 synchronous adenomas, adenoma with high-grade dysplasia, large (>10 mm) adenomas) or a severe phenotype (>1000 synchronous adenomas).

Risk of neoplasia and cancer in the pouch

There is a very small risk of adenocarcinoma after an IPAA. Most cancers develop in residual rectal or in the anal transitional zone (ATZ) mucosa. Cancer can also develop within the ileal component of the pouch. Currently nearly all pouches are constructed with the use of stapling devices which results in the ATZ mucosa being preserved. Von Roon reported on 206 patients with a median follow-up of 10.3 years. The risk of adenoma of the IPAA at 10 years was 51% after stapled IPAA; no patient developed cancer. van Duijvendijk et al reported the risk of developing a polyp of the IPAA after 7 years was 31% after stapled IPAA.

Of 212 patients followed up in the Dutch polyposis registry, the cumulative risk of developing an adenoma in the pouch at 10 year follow-up was 45%. Twenty-five patients (11.8%) developed an adenoma with advanced pathology and four (1.9%) developed a carcinoma. The cumulative risk of developing a pouch carcinoma at 10 year follow-up was 1%. A 2013 review of 24 studies reporting 92 pouch-related cancers found that 23 of 92 cancers (25%) developed in the pouch mucosa and 69 (75%) in the ATZ. A Mayo clinic study on 117 patients showed a median time to development of dysplasia was 149 months. Adenocarcinoma developed in one patient after 284 months. Risk of dysplasia at 10, 20 and 25 years was 17%, 45% and 69%, respectively.

Thus, although the risk of severe mucosal dysplasia and cancer is low, annual endoscopic surveillance of any remaining rectal mucosa, ATZ mucosa and ileal pouch are recommended for life.
FAP and desmoid disease

We suggest that FAP patients should be counselled about the risk of post-operative desmoid disease formation.

(GRADE of evidence: low; Strength of recommendation: weak)

Consensus reached: 95% agreement.

For FAP patients before colectomy, consider determining genotypes or family history of desmoid disease which may be predictive of desmoid formation.

(GRADE of evidence: low; Strength of recommendation: weak)

Consensus reached: 78% agreement.

We suggest that sulindac in combination with high-dose selective oestrogen receptor modulators may be effective in FAP patients with intra-abdominal desmoids and desmoids located at the abdominal wall.

(GRADE of evidence: very low; Strength of recommendation: weak)

Consensus reached: 100% agreement.

We recommend the role of elective surgery for intra-abdominal desmoids should be restricted to treating secondary effects of the desmoid disease, and this surgery should be performed in expert centres.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 95% agreement.

Desmoid tumours will develop in around 15% of patients with FAP. Risk factors include abdominal surgery, positive family history for desmoids and site of the pathogenic variant.248 249 In individuals with APC pathogenic variants in the desmoid region 3' to codon 1399, abdominal surgery was associated with a 65% risk of developing mesenteric desmoids.249 As desmoids can cause significant complications and pose a low risk of death, genotypes predictive of desmoid formation and family history of desmoids should be determined for all patients before colectomy. Identification of at-risk patients will enable appropriate counselling and consent before surgery and allow informed decision about the timing of surgery. This decision should sensibly balance the risk of malignancy with the risk of mesenteric desmoid disease.

Some desmoid tumours spontaneously stop growing, some regress and others remorselessly increase in size. In a small proportion, this increase may be rapid and uncontrollable. There are no proven predictors of growth pattern.

The distinction between intra- and extra-abdominal desmoids is important. Abdominal wall desmoids cause pain (variable) and mass effect. Intra-abdominal lesions, in addition, are at risk of causing secondary effects: ureteric and small bowel obstruction, fistula formation or small bowel ischaemia. The quality of data on which to make decisions about treatment of abdominal wall and intra-abdominal desmoid disease is limited and largely consists of small, non-controlled studies. As such, the treatment of patients with desmoid tumours remains controversial. For abdominal wall lesions, there is an expanding role of percutaneous ablative treatments and in some institutions ablation has become the primary treatment.250

Church et al 251 combined symptoms (pain, restriction, hospitalisation and sensation of a mass) with size and growth rate in a proposed staging system for intra-abdominal desmoid disease. Decisions about treatment are partly based on size and symptoms, but are more often determined by secondary effects of the desmoid on the urinary and GI tract. Following diagnosis, if no secondary effects are present, serial imaging with CT or MRI scanning every 6–12 months depending on growth rate is reasonable. Radiological screening at 12 months following surgery for those who are at high-risk of desmoid development might be appropriate.

If there is concern about the size or growth rate of desmoid disease, first line treatment should be with high-dose selective oestrogen receptor modulators and sulindac, according to the regime outlined by Quast et al.252 In this observational series all desmoid patients treated and followed at their institution had completed at least 1 year of treatment.252 Response was defined as stable size or regression of desmoid size between two CT or MRI scans. Of the 134 patients included, half had a confirmed diagnosis of FAP. Eighty-five per cent of patients showed regression or had stable desmoid size. The mean time to reach at least stable size was 14.9±9.1 months. After regression or stabilisation, medication was tapered in 60% of the treated patients with only one long-term recurrence after >10 years.

In patients with progressive intra-abdominal desmoid disease that does not respond to this treatment, chemotherapy (eg, doxorubicine and dacarbazine or methotrexate and vinblastine)253 254 or radiation therapy is indicated.

Surgery should be reserved for secondary effects of the desmoid disease.

MUTYH-ASSOCIATED POLYPOSIS

Colorectal surveillance in MAP

We recommend that colorectal surveillance is commenced in MAP starting at the age of 18–20 years. If surgery is not undertaken then annual surveillance is suggested.

(GRADE of evidence: low; Strength of recommendation: strong)

Consensus reached: 83% agreement.

We recommend that for monallelic MUTYH pathogenic variant carriers, the risk of CRC is not sufficiently different to population risk to meet thresholds for screening, and routine colonoscopy is not recommended.

(GRADE of evidence: moderate; Strength of recommendation: strong)

Consensus reached: 94% agreement.

MAP is a recessively inherited cancer and polyposis predisposition syndrome caused by pathogenic variants in the MUTYH base-excision repair gene, with significant phenotypic overlap with FAP. It is classically said to be associated with a more attenuated phenotype than FAP but there is significant phenotypic variation. Past guidelines have suggested that colonoscopy should commence at the age of 18–20 years, on the basis that CRC in MAP is rare below the age of 30 years207 and be performed up to 2 yearly, as the colonic phenotype is usually more attenuated. This attenuated phenotype also lead the authors to comment that endoscopic management may suffice if there is an attenuated phenotype.

CRC in MAP is more likely to be right sided and synchronous.255 256 A cumulative CRC risk of 63% at 60 years has been reported.257 The median age of CRC seems dependent on the underlying genotype.256 MAP accounts for up to 29% of those patients with 10–100 adenomas but also up to 29% of those with a classical phenotype (100–1000 adenomas). Biallelic pathogenic variants have also been reported in patients with <10 adenomas and also in patients with CRC but no adenomas.258 259 There are further data to suggest that CRC risk in MAP may not correlate to adenoma number.257

Colonoscopy surveillance may not be effective in MAP and it has been postulated that there may be accelerated carcinogenesis. Nieuwenhuis and colleagues reported that of those who presented with a polyposis phenotype but without CRC, 98%...
Guidelines

developed CRC during 5 years of surveillance. If the presentation was with CRC, then they observed a 5-year metachronous cancer risk of 11%. In light of these data, annual colonoscopy would appear appropriate if colonoscopy surveillance is pursued, but it appears that surgery may be the more appropriate management strategy, balanced against age, co-morbidity and expected functional outcome.

Upper GI surveillance in patients affected with MAP

We suggest that upper GI surveillance should be considered in MAP, starting at the age of 35 years. We recommend that the surveillance interval is determined as outlined for FAP.

(Grade of evidence: low; Strength of recommendation: weak)

Consensus reached: 94% agreement.

The data regarding the upper GI tract phenotype in MAP is far less extensive than in FAP. It is generally suggested that the two conditions should be managed in a similar manner. Duodenal adenomas have been reported in 17–34% of those with MAP. The median age at which duodenal adenomas are diagnosed has been reported as 50 years. Two patients were diagnosed with duodenal or ampullary cancer in one series, age 83 and 63 years, respectively; both were diagnosed at index upper GI endoscopy but no cancer arose in patients who were on a surveillance programme. Polyps appear to progress through the development of villous features and increasing size in this series, rather than progression in number or dysplasia. Certainly the phenotype does appear to be different and polyp multiplicity in MAP does not seem to be observed in the same manner as it is in FAP. There appears to be scientific evidence to suggest that MAP duodenal polyps may behave differently to those in FAP. A higher burden of somatic pathogenic variants in MAP duodenal adenomas is reported, compared with FAP, despite a lower Spigelman stage disease. It is postulated that this may reflect an increased cancer risk in the context of apparently less severe benign disease; however, long-term data to support this hypothesis are lacking.

Because of the reduced frequency of duodenal adenomas, compared with FAP and the later onset of duodenal adenomas, it has been suggested that starting upper GI endoscopic surveillance could be deferred until the age of 35 years. In the absence of more robust data, determining surveillance intervals according to duodenal and ampullary staging, as has been recommended in FAP, appears to be a pragmatic approach.

There are few data regarding gastric lesions in MAP. FGGPs have been reported in 6% of a cohort with MAP and gastric adenomas in 3%. This would suggest that gastric polyposis is seen less commonly than in FAP. In the same series gastric cancer was observed but the incidence was not statistically different to the risk for the general population. Further data regarding the gastric phenotype are required to determine risk and how best to manage such lesions.

PEUTZ-JEGHERS SYNDROME

PJS is an autosomal dominant genetic disorder characterised by the development of benign hamartomatous polyps in the GI tract and hyperpigmented macules on the lips and oral mucosa (which fade with age). The prevalence of PJS is between 1/8300 and 1/29 000 and most patients have pathogenic variants in the STK11 (also called LKB1) gene.

Diagnostic criteria: A diagnosis of PJS in an individual may be made when any one of the following is present:

- Two or more histologically confirmed PJ polyps
- Any number of PJ polyps detected in one individual who has a family history of PJS in a close relative
- Characteristic mucocutaneous pigmentation in an individual who has a family history of PJS in a close relative
- Any number of PJ polyps in an individual who also has characteristic mucocutaneous pigmentation
- A pathogenic variant in STK11.

The risk of GI tract cancer in PJS

GI tract carcinogenesis in PJS is controversial. The malignant potential of the PJ polyp is unclear. A hamartoma–adenoma–carcinoma sequence has been proposed but robust supporting data are lacking. There are data suggesting that the PJ polyp is non-neoplastic, including descriptions of polyclonality and the rarity of dysplasia in PJ polyps.

It is widely accepted that there is an increased risk of malignancy in PJS. However, the risk is difficult to quantify, with the majority of the literature being small cohort studies with inherent bias, potentially leading to overestimation of risk. A meta-analysis has been performed by Hearle et al creating a cohort of 419 patients with PJS. This offers the most comprehensive data for cancer risk with a luminal GI cancer risk of 1%, 9%, 15%, 33% and 57% at 30, 40, 50, 60 and 70 years, respectively. More recent data, although from a smaller cohort, support the proposition that GI cancers are less of a clinical problem and that pancreatic and breast cancers are the most commonly diagnosed malignancies in PJS.

GI surveillance and PJS

We suggest that in an asymptomatic patient with PJS, GI surveillance by upper GI endoscopy, colonoscopy and VCE commence at the age of 8 years. We recommend that small bowel surveillance should continue 3 yearly. If baseline colonoscopy and OGD are normal, then they can be safely deferred until the age of 18 years; however, if polyps are found at baseline examination, then they should be repeated 3 yearly. Earlier investigation of the GI tract should be performed in symptomatic patients.

(Grade of evidence: low; Strength of recommendation: weak)

Consensus reached: 82% agreement.

Small bowel polyps resulting in intussusception is the major clinical problem affecting children with PJS. The cumulative intussusception risk is estimated at 50–68% during childhood, with 15–30% requiring surgery before the age of 10 years with a median age of the first intussusception of 10–16 years. These data include patients who preceded routine small bowel surveillance. They also combine patients with an established diagnosis of PJS undergoing surveillance, with those not under surveillance in whom the diagnosis of PJS was made at presentation with an intussusception. The recommendation to start small bowel surveillance at the age of 8 years, and earlier if symptomatic, is based on these data.

Gastroscopy and colonoscopy are the preferred investigation to assess the upper GI tract and colon, respectively. Barium follow through has been replaced by VCE and MRI enterography, both of which are better tolerated by patients, have a similar accuracy in detecting clinically significant polyps (> 1 cm) and avoid the need for repeated ionising radiation. VCE and MR enterography may be complimentary. While VCE may be better at detecting smaller polyps, MRI enterography is better at localisation and accurate sizing.

Double balloon enteroscopy (DBE) should not be recommended as a surveillance tool as it is technically challenging, limited by size of the abdomen and requires both oral and rectal approaches;
therefore a general anaesthetic is required to reliably visualise the whole length of the small bowel. The main role of DBE is therapeutic, for targeted polypectomy.

Role of endoscopic polypectomy in PJS
We suggest elective polypectomy to prevent polyph related complications. Small bowel polyps >1.5–2 cm in size (or smaller if symptomatic) should be considered for elective resection to prevent intussusception.

(GRADE of evidence: low; Strength of recommendation: weak)
Consensus reached: 88% agreement.

Polypectomy at surveillance is recommended in PJS. It is not clear whether this approach modifies cancer risk and indeed, given the lack of dysplasia in PJS polyps, it is unlikely to do so. The main role for polypectomy therefore is to prevent polyph-related complications. Prevention of anaemia and bleeding, however, is difficult to quantify and there are no data regarding this. Intussusception is the most important clinical problem in children with PJS, with an accumulative risk of 50–68% in childhood.268 269

Existing guidelines suggest intervening in small bowel polyps which are 1.5–2 cm, earlier if symptomatic.270 This is supported by data which report a median polyp size of 35 mm (15–60 mm) in cases of confirmed intussusception.269 In addition, in a cohort in whom routine surveillance was performed, with intervention planned as outlined above, no patient required emergency surgery for intussusception during 683 patient-years follow-up.266

Options to remove a PJS polyp include endoscopy, surgery or combined approaches. There are no data to state that one modality is superior to another and choice will be dependent on patient (eg, age, past abdominal surgery), polyp factors (size, location and multiplicity) and local availability of techniques. For those who undergo surgery for small bowel polyps, intraoperative enteroscopy and a “clean sweep” is recommended and has been demonstrated to reduce the need for subsequent laparotomy.277 Polyp clearance by intraoperative enteroscopy in the PJS may be effective in the long term.272 278

JUVENILE POLYPOSIS SYNDROME
JPS is an autosomal dominant condition characterised by predisposition to hamartomatous polyps in the GI tract. The prevalence is between 1/100 000 and 1/160 000, with the following diagnostic criteria:279:

- >5 juvenile polyps in the colon or rectum
- Juvenile polyps in other parts of the GI tract
- Any number of juvenile polyps and a positive family history
- A pathogenic variant in SMAD4 or BMPRIA.

About 40% of families have mutations in SMAD4 and a further 40% in BMPRIA. Some patients have large chromosomal deletions encompassing both PTEN and BMPRIA genes. There is a requirement to develop an evidence base for the personalised management of JPS-related cancers, which is beyond the scope of this guideline.

JPS and colorectal surveillance
We suggest colonoscopic surveillance should commence from the age of 15 years or earlier if symptomatic. The surveillance interval should be 1–3 yearly, personalised according to colorectal phenotype.

(GRADE of evidence: low; Strength of recommendation: weak)
Consensus reached: 83% agreement.

The polyps in JPS are thought to have the potential for malignant transformation with dysplasia present in 8% of resected polyps in one series.280 The risk of CRC is ill-defined, being based on small series, with inherent bias. Such cohorts have reported CRC in 14–22% of patients,280 281 with a reported lifetime CRC risk of 39–68%.280 282 CRC in childhood is not a clinically significant problem. The mean age of CRC in two of the larger studies was 44 years.280 282

The risk of CRC is undoubtedly elevated, and the current consensus is that colonoscopic surveillance is required, with the aim of preventing or detecting early cancers. Colonoscopy and polypectomy also has the potential to prevent polyph-related morbidity (bleeding, anaemia or abdominal pain). There are few data regarding the outcomes of surveillance in JPS. In one of the largest cohorts, colonoscopy was safe. Two patients developed CRC on surveillance—one in the setting of carpeting of polyps, and the other arose following what was almost certainly an incomplete colonoscopy in the mid 1970s.280

For those with a confirmed genetic diagnosis of JPS, or individuals with a clinical diagnosis of JPS in whom molecular genetic test results were uninformative, colonoscopic surveillance can be delayed until the age of 15 years if asymptomatic.

There are no data to guide the most appropriate interval for colonoscopy surveillance in JPS. There are no data to support accelerated carcinogenesis in JPS. A personalised approach with the surveillance interval (1–3 yearly) based on individual colonic phenotype appears to be a pragmatic approach and is in line with recent paediatric polyposis guidelines.279 Colonic resection may be considered for those with a polyph burden not manageable endoscopically. For those unaffected individuals from families with a clinical but not molecular diagnosis, yearly colonoscopy would seem adequate. If normal, the interval to be reviewed if a JPS phenotype is found or other sporadic polyps are identified.

Upper GI surveillance and JPS
We suggest that for those with a confirmed clinical or genetic diagnosis, upper GI endoscopic surveillance should start at the age of 18 years for SMAD4 pathogenic variant carriers and the age of 25 years for BMPRIA pathogenic variant carriers and those without an identified mutation. The surveillance interval should be 1–3 yearly, personalised according to upper GI tract phenotype.

(GRADE of evidence: low; Strength of recommendation: weak)
Consensus reached: 83% agreement.

We suggest that for those with an FDR with a clinical diagnosis of JPS and in whom a mutation has not been identified, screening of the upper GI tract is not required routinely but should be initiated if/when a clinical diagnosis is made on the basis of colonic phenotype. It may however be considered if there is a family history suggestive of hereditary haemorrhagic telangiectasia (HHT), even in the absence of colonic polyps.

(GRADE of evidence: low; Strength of recommendation: weak)
Consensus reached: 83% agreement.

Upper GI endoscopy is not required in those unaffected individuals in whom the diagnosis in the family is clinical without a confirmed molecular diagnosis. It is clear that the colon is the site most likely to be affected in JPS and therefore any upper GI tract assessment can be deferred until a colonic phenotype of JPS has been established in that individual.

There is a paucity of data evaluating the upper tract in JPS. Historical series report the incidence of gastric polyps between 63% and 83% and duodenal polyps between 14% and 33%.283 284 More recently, 28/41 patients had neither macroscopic nor microscopic features of upper GI tract polyps and 13/41 had confirmed histological upper GI tract involvement at a median age of 33 years.286 In this cohort five patients underwent
OGD before the age of 15 years and three had changes consistent with JPS, but none had dysplasia. In another series, gastroduodenal polyps were seen in 37%; in this cohort none of the six patients who had upper GI endoscopy before the age of 18 years had gastroduodenal polyps. 280

Due to limited data, it is difficult to determine the exact lifetime risk of gastric cancer in JPS. There were two gastric cancers in a cohort of 44 patients with JPS (median age 36 years) 280 and a further two (age not specified) in a cohort of 56 patients who appear to have been mostly JPS but some of whom had phenotypic features overlapping with Cowden syndrome. 286 In another series, 3/42 patients had either high grade dysplasia or cancer in the stomach 285 and a further two patients had prophylactic gastrectomy for benign gastric polyp burden.

The risk of severe gastric polyposis and gastric cancer phenotype appears to be increased in patients with SMAD4 pathogenic variants. 287 288 All patients with advanced gastric polyposis were SMAD4 pathogenic variant carriers in one cohort. Aretz et al reported a significantly higher risk of gastric polyposis in SMAD4 pathogenic variant carriers (73%) vs 8%; p<0.001) and again all cases of gastric cancer in this cohort occurred in patients with SMAD4 pathogenic variants. 288

Current published guidelines for the age at which to start upper GI surveillance ranges from 12 years 261 to 25 years of age. 25 Given the emerging genotype–phenotype association and the lack of significant pathology being found in childhood, it is recommended that those with a SMAD4 mutation undergo upper GI tract surveillance 1–3 yearly from the age of 18 years, and those with a BMPR1A mutation from 25 years of age.

Additional investigations required in patients with a SMAD4 mutation

We suggest that patients with a SMAD4 pathogenic variant should be evaluated for haemorrhagic telangiectasia (HHT), and that those at risk of, or with a confirmed diagnosis of HHT are best managed in conjunction with a specialist HHT centre.

(GRADE of evidence: low; Strength of recommendation: weak) Consensus reached: 90% agreement.

The coexistence of JPS and HHT was reported early in the 1980s. 299 It is now recognised that this is due to SMAD4 mutations. Up to 76% of patients with JPS due to SMAD4 mutations have features of HHT. 290 Thoracic aortic disease and mitral valve dysfunction have also been reported. 290 291 Aortopathy has been described in 38% of patients with a SMAD4 mutation, irrespective of the JPS phenotype. 291

Patients may lack overt clinical symptoms of HHT but are at risk of asymptomatic arteriovenous malformation (AVMs) which can result in life threatening complications. Patients with a SMAD4 pathogenic variant are probably best managed in conjunction with a specialist centre with experience in evaluating and managing HHT patients. The recent British Thoracic Society guidelines on pulmonary AVMs may act as a useful resource. 292

PTEN-hamartoma syndrome overlap and additional investigations indicated in a patient with microdeletions involving BMPR1A

Patients with JPS and a microdeletion involving BMPR1A and PTEN are at risk of the clinical manifestations of both JPS and PTEN-hamartoma tumour syndrome (PHTS). We suggest that they should be referred to their local genetics centre for further advice and to coordinate their surveillance needs.

(GRADE of evidence: low; Strength of recommendation: weak) Consensus reached: 89% agreement.

PHTS encompasses four major clinically distinct syndromes associated with constitutional pathogenic variants in the tumour suppressor PTEN. These are associated with macrocephaly, dysmorphism, developmental delay and an increased risk of extra-intestinal cancers and possibly intestinal cancers. BMPR1A is located in the same chromosomal region as PTEN and deletions involving both genes have been reported. Microdeletions involving both genes have important clinical implications. There are numerous case reports suggesting that if the pathogenic variant (microdeletion) is found in both genes, these patients may present with a severe form of JPS with onset in early childhood and may also have an increased risk of the extra-intestinal manifestations of PHTS. 294–295

Author affiliations

1Family Cancer Clinic, St Mark’s Hospital, London, UK
2Faculty of Medicine, Imperial College, London, UK
3Clinical Genetics, West of Scotland Genetics Services, Glasgow, UK
4Gastroenterology, Cardiff and Vale NHS Trust, Cardiff, UK
5Clinical Genetics, Guy’s and St Thomas’ NHS Foundation Trust, London, UK
6C GCC, University of Edinburgh, Edinburgh, UK
7Translational Gastroenterology Unit, John Radcliffe Hospital, Oxford, UK
8Oxford NIHR Biomedical Research Centre, University of Oxford, Oxford, UK
9Faculty of Medicine & Health Sciences, Nottingham University, Nottingham, UK
10Head of Policy and Campaigns, Bowel Cancer UK, London, UK
11Genetic Medicine, Central Manchester University Hospitals Foundation Trust, Manchester, UK
12Polyposis Registry, St Mark’s Hospital, London, UK
13Gastroenterology, University Hospital of North Tees, Stockton-on-Tees, UK
14Northern Institute for Cancer Research, Newcastle University, Newcastle upon Tyne, UK
15Nuffield Department of Clinical Medicine, Wellcome Trust Centre for Human Genetics, Birmingham, UK
16Cancer Research Centre, University of Edinburgh, Edinburgh, UK

Twitter Kevin J Monahan @kevinjmonahan and Sunil Dolwani @sdolwani

Collaborators Hereditary CRC guidelines eDelphi consensus group (professional societies or other roles in brackets) Toni Sepplà (EHTG, INSIGHT), Sue Clark (EHTG, BSG, INSIGHT, ACPGBI, ESCP), Omar Faiz (ACPGBI, ESCP), Francesca Baluguer (AEG, EHTG), Monique van Leerdam (EHTG, ESGE), D Gareth Evans (UKCCG, EHTG), Rodrigo Jover (EHTG, ESGE), Marc Trischkowitz (UKCCG), Marc Hansson (UKCCG), Helen Hanson (UKCCG), Sarah Gibson (UKCCG), Amy Taylor (UKCCG), Gabriella Moeslein (EHTG, ESCP), Anja Wagner (EHTG), JC Saurin (ESGE), Tracy Smith (PPI, Lynch Syndrome UK), Jane Ashford (PPI, Lynch Syndrome UK), Jennifer Martin (PPI, Lynch Syndrome UK), Jennifer Cunningham (PPI, Polyposis representative), Mark Cooper (PPI, Lynch Syndrome UK).

Contributors The document was written by all members of the guideline development group (GDG) and edited by the lead author. The guideline development process was contributed by all members of the GDG as outlined in the methods section.

Funding This study was funded by British Society of Gastroenterology. Prof. James East was funded by the National Institute for Health Research (NfH) Oxford Biomedical Research Centre (BRC). The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health.

Competing interests KJM: Medical advisory board of Bowel Cancer UK, Lynch Syndrome UK. JH and FL: FAP trial (now closed) with funding awarded to NHS trust research facility. JE: Advisory board Lumendi, Boston Scientific; Speaker fees Olympus, Falk. MDR: Speaker fees: SwissSCWeb, Pentax; Research Grant: Olympus; Consultancy: Norgine.

Patient consent for publication Not required.

Provenance and peer review Commissioned; externally peer reviewed.

Data availability statement All data relevant to the study are included in the article.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iDs

Kevin J Monahan http://orcid.org/0000-0002-7918-4003
Matthew D Rutter http://orcid.org/0000-0001-9507-0295
Ian Tomlinson http://orcid.org/0000-0003-3037-1470

Guidelines
Guidelines

1. Introduction

2. Pathogenesis of Familial Adenomatous Polyposis

3. Clinical Manifestations

4. Diagnosis

5. Management

6. Conclusion

References


et al.: 1–34. doi:10.1136/gutjnl-2019-319915


