Alimentary and pancreas

Intestinal function and intestinal blood supply: a 20 year surgical study

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SUMMARY The experience of The Middlesex Hospital in the investigation and management of chronic intestinal arterial occlusion (IAO) over a 20 year period is presented. Of some 100 patients referred only 28 were found to have IAO and these were critically evaluated in terms of insorptive and exsorptive function. No relation was found between intestinal performance and the degree of potential ischaemia suggested by angiography. Twenty two patients underwent reconstruction of the coeliac axis and mesenteric arteries, with three perioperative deaths. The remainder were followed for periods ranging between six months and 10 years. Of these, 15 are asymptomatic, one is unchanged, and one is subjectively worse. The role of elective surgery in relation to the visceral arteries is critically discussed in the light of these results, and in the perspective of the world experience.

It is known that the origins of the three main visceral arteries are frequently narrowed or blocked by atheroma, but the relationship between this lesion and the development of abnormalities in the alimentary tract remains unclear. Whereas intestinal infarction, if not relieved, is always fatal, it is not easy to identify the pre-infarctive state, which might be amenable to treatment. This study addresses the following questions: (1) Does it happen that the blood supply to the gut becomes so reduced that symptoms occur, before the point of infarction is reached, and if so, how frequently? (2) Can this situation be identified clinically, or by laboratory tests of intestinal function? (3) Given the existence of symptomatic occlusion of the visceral arteries, what is the risk of a fatal infarction occurring in the individual patient? (4) What is the role of elective surgical reconstruction of the intestinal blood supply?

The ideal way of resolving these questions would be to measure gut blood flow in a control population of normal people, and in patients with intestinal arterial occlusion (IAO), and to investigate alimentary function in both groups, so as to identify any significant difference. The subjects would then be watched over a period of years, without treatment, in order to compare symptoms and eventual mortality. Obviously such a study would be both unethical and impractical. The investigator is constrained by the needs of his patients, and by considerations of money and time, quite apart from the fact that at the present moment we have no method of measuring intestinal blood flow in the intact human being which is acceptably non-invasive. Total splanchnic blood flow can, and has been measured, but this is a different question, as is later discussed. The problem has to be approached indirectly, and this has been done in three ways.

Methods

ANIMAL STUDIES The mesenteric circulation of the experimental animal has been extensively studied over the last century and this work is reviewed elsewhere. Much of it is probably irrelevant to human illness, but from the mass of data collected two observations stand out. (1) The intestine takes about one fifth of the resting cardiac output, and after a meal this
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Proportion rises by about 15%. (2) Chronic reduction of the blood supply to the gut does not have any discernible effect on structure or function, until infarction occurs, which brings about a catastrophic series of disturbances, leading to death.

These findings derive from methods which are at best invasive, and often lethal, and have no clinical use. It is reasonable to suggest, however, that the human gut may respond in the same way as do other mammals to ischaemic insult.

AUTOPSY STUDIES
The incidence of arterial stenoses and occlusions in a given population can be determined by postmortem examination. Our group recently reviewed this problem and produced revised data which in the light of 203 consecutive autopsies, indicated that although minor degrees of ostial stenosis are common, 'critical stenosis' as defined by careful analysis of external and internal diameters of the visceral arteries, in relation to accepted estimates of flow, is unusual. This series (which is the largest so far reported) corrects previous data which suggested that major ostial lesions were frequent. There is a clinical parallel in that in a series of 2000 mesenteric arteriograms the number of identifiable stenoses was less than 5%. The more interesting conclusion from our study, however, was that no relationship could be found between IAO (as seen postmortem) and symptoms of digestive disease experienced during life.

CLINICAL STUDIES
These mainly depend on arteriography. A qualitative guide to intestinal bloodflow can be made by measuring the cross sectional area of the vessels radiologically. Such studies have, however, never been related to intestinal function. Most surgeons reporting series of operations to restore circulation to the intestine have been content to plan treatment on the basis of angiography, and to estimate success on subjective grounds. Bearing in mind the prevalence of asymptomatic arterial lesions, and the placebo effect of major surgery, the indications for this type of operation, which is expensive and not without risk, remain unclear. Certainly, disturbances of function after successful revascularisation of the acutely ischaemic bowel have been studied in depth but this is a quite different clinical situation. Whether or not the insorptive and exsorptive functions of the GI tract are affected by IAO is disputed, because some authors describe consistent abnormalities in relation to reduced blood supply, while others deny the usefulness of all such measurements.

The most direct approach to this problem has been made by Hansen's group in Denmark who estimated total splanchnic bloodflow (SBF) using hepatic clearance of indocyanine green, in 15 persons, distributed between four controls (that is patients with abdominal pain but no arterial lesion), five suspects (patients with abdominal pain and a minor stenosis shown radiographically) and six patients who had both symptoms and major angiographic lesions and underwent surgery. Hansen showed that resting splanchnic blood flow was comparable in all groups, but that the six identified patients were unable to increase total splanchnic flow when challenged by a meal. After reconstruction of the visceral arteries, the patient group exhibited an increased fasting flow, equivalent to the controls, and regained the capacity to expand flow after the test. Measurement of liver blood flow, however, bears an uncertain relationship to what happens in the gut. The liver receives blood via the hepatic artery (which was not studied here) and the portal vein, via the stomach, spleen, the diaphragm, and innumerable collateral pathways which expand when the normal routes of supply are blocked. What we cannot at present measure is intestinal blood flow, which is the main determinant of eventual infarction. Although Hansen's group could correlate splanchnic blood flow with later clinical performance, no operative nor long term data are included, and the authors modestly describe their important work as a preliminary study. More recently Fleming has developed a method of fractionating splanchnic blood flow using labelled sulphur colloid, and this may prove useful in the investigation of suspected chronic ischaemia.

PRESENT STUDY
Because of the lack of knowledge of the relationship between radiologically demonstrated lesions of the visceral arteries (IAO) and the symptoms and natural history of the individual patient, a prospective study of the problem was started in 1965 in the Department of Surgical Studies of The Middlesex Hospital, and the data reported here extend to January 1984.

We have, deliberately, avoided the use of terms such as 'intestinal angina', 'chronic intestinal ischaemia' or 'abdominal claudication', because such phrases assume the reality of a syndrome which it is our purpose to evaluate. Nonetheless, some sort of shorthand is useful and intestinal arterial occlusion (IAO) here refers to lesions seen on an angiogram, with no conclusions being drawn as to their clinical importance.

From 1962 to 1984 approximately 100 patients with abdominal pain were referred to The Middlesex Hospital as possible cases of intestinal arterial...
occlusion. This total is necessarily inexact, because as already reported a number of the patients were not investigated further and are now untraceable, either because their symptoms had disappeared when they arrived at the clinic, or because their problems did not seem in any way to resemble those related to intestinal arterial occlusion and hence were not followed up. A further group were found on investigation to have other causes of abdominal pain, as is shown in Table 1, and were not submitted to angiography. Once these conditions had been excluded, the aorta and visceral arteries were opacified. In 21 the findings were normal. If the angiogram showed stenosis or occlusion of the main trunks (CA, SMA, IMA), it seemed ethical to submit the patient to a more exacting diagnostic procedure, in order to decide whether major surgery was justified. All patients in this series were received because of the complaint of abdominal pain, and our study does not relate to other reported series where vascular operations have been carried out on patients without alimentary symptoms, but in whom intestinal arterial occlusion was incidentally discovered on an angiogram undertaken for other reasons.

PATIENTS
Table 1 shows the total group. Following the exclusions already defined, 28 patients underwent detailed investigation of their alimentary function, with a view to possible vascular surgery. This group comprised 14 men and 14 women (age range 23 to 77 years, mean age 49). They were evaluated as follows.

1 CLINICAL APPRAISAL
(a) Symptoms
By definition, all our patients complained of abdominal pain related to meals, and reported loss of weight and an alteration in bowel habit (either constipation or diarrhoea). Nausea and vomiting was noted in nine, and 20 gave a history relating to vascular occlusion elsewhere in the body.

(b) Physical signs
The 20 patients with arterial symptoms had objective evidence of arterial disease, as evidenced by absent lower limb pulses, or asymptomatic arterial bruits. An abdominal systolic bruit, though not always specifically sought, but was found in 14. (The significance of this sign is in any case doubtful8). Apart from weight loss, there were no other relevant abnormalities on examination.

(c) Baseline investigations
Physical evaluation was followed by baseline tests comprising chest radiograph, ECG, blood film with indices and routine serum biochemistry, in order to eliminate concomitant disease and to determine fitness for possible surgery.

2 ARTERIOGRAPHY
In the first years of this study this was by the translumbar route. Subsequently the preferred technique was retrograde catheterisation via the femoral artery, with free injection into the aorta and selective opacification of the visceral trunks. Typical findings are illustrated in Figures 1–4.

3 TESTS OF INTESTINAL FUNCTION
Once an arterial obstruction had been identified, the patient was further studied with regard to intestinal function. The tests applied necessarily reflect a series which has extended over 20 years (see Table 2), and nowadays would be somewhat different. The following functions were investigated:

(a) Iron metabolism and $B_{12}$ absorption
This comprised serum iron concentration (Fe), total iron binding capacity (TIBC), $B_{12}$ and folate levels, Schilling test for $B_{12}$ absorption.

(b) Fat metabolism
This comprised serum $\beta$-lipoprotein and fasting triglyceride concentrations, mean daily faecal fat excretion over five days, and also serum carotene concentrations which have been reported in the

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Table 1  Cases of IAO referred to The Middlesex Hospital, 1962–84

| ±100 patients referred with suspected intestinal angina | ±37 Not investigated further (Clearly unrelated problems or symptoms disappeared) |
| ±63 Investigated | 14 found to have other pathology (Gall stones, peptic ulcer, carcinoma of stomach, colon or pancreas, abdominal aortic aneurysm) |
| 49 Aortograms | 21 no arterial lesion – not investigated further |
| 28 Arterial lesions | 6 no operation |
| | 1 refused surgery |
| | 3 arteries |
| | 1 fibromuscular hyperplasia |
| | 1 unacceptable operative risk |
| 22 Reconstructions | Followed from 6 to 120 months |
| | 3 operative deaths |
| | 2 MI (8, 16 days) |
| | 1 bowel infarct (4 days) |
| | 2 later deaths |
| | 1 CVA (3 ms) |
| | 1 bowel infarct (9 ms) |
| | 15 asymptomatic |
| | 1 unchanged |
| | 1 worse |
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past,\textsuperscript{10 13 21 22} as reflecting malabsorption of vitamin A in chronic intestinal ischaemia.

(c) Liver function
By means of serum albumen and globulin concentrations, bilirubin, alkaline phosphatase, and AST.

(d) Carbohydrate absorption and metabolism
Comprising random blood glucose concentration, glucose tolerance test (GTT) following a 50 g dose and d-xylose excretion after a 5 g dose.

(e) Intestinal protein loss
By measurement of faecal activity, following an I/V dose of Cr\textsuperscript{51} labelled albumen.

(f) Barium studies
These were initially by conventional upper GI series, later by the intubated duodenal technique.

(g) Imaging
The series was begun before fibrendoscopy of the alimentary tract, or ultrasound examination, were easily available, but later patients were studied with these techniques.

Results
The results of these studies are set out in Table 2. It will be seen that the great majority of patients had normal absorptive and exsorptive function as measured by the tests used. Abnormalities did occur (deficient fat absorption in two, carbohydrate absorption in four and an exudate protein-losing enteropathy in two) but these were sporadic and bore no consistent relation either to the patients' symptoms, nor to the arterial lesion. Because of this, it seemed pointless to repeat the tests after operation.

As in all previous case series, management was decided on clinical criteria as no laboratory test gave sufficient discrimination. In practice this meant that significant intestinal arterial occlusion – that is, that requiring operation – was identified by exclusion, so
that only those patients who continued to complain of pain in the presence of a known arterial block, when a full clinical assessment had not produced any other explanations, were submitted to reconstructive surgery.

The types of operation used are set out in Table 2 and illustrated in Fig. 5. Case 1 was operated upon by Mr Ian Ranger and reported\textsuperscript{23} in 1962. The remaining operations were carried out by the principal authors (AM:20, JMFC:2). Side-to-side anastomosis between the aorta and the SMA was performed in six cases (Fig. 6). This technique has the advantage of being possible through a purely abdominal approach. Patch angioplasty of the CA, SMA or both was chosen in 10 cases, and reimplantation of the SMA in one. For these procedures a full thoraco-abdominal incision is preferred. In two patients all that was necessary was to free the origin of the CA from the surrounding tough fibrous tissue including the median arcuate ligament of the diaphragm and sympathetic and lymphatic fibres. In each case a gradient was measured across the arterial lesion, which was abolished by the procedure. In one case a tight stricture of the IMA was corrected by a Dacron patch. It was not considered justified to carry out routine postoperative angiograms, until late in the series when a digital facility became available and it was possible to opacify the visceral arteries using an intravenous injection. In the four cases so studied, three reconstructions were patent and in one an occluded graft was accompanied by return of symptoms. It will be seen that of the patients operated on three died in the immediate postoperative period, one six months later from a stroke, and one nine months later, because of total infarction of the midgut requiring enterectomy, which was followed by a fistula. Of the remainder, 13 were free of symptoms for variable periods, two were unchanged and one was substantially worse (this poor result was considered to be because of an error in psychological assessment before the operation). The follow up period is given in each case and

Fig. 2 Coeliac and superior mesenteric occlusion with stenosis of the inferior mesenteric artery.
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The operations are exacting but recovery is on the whole smooth and free of complications. In particular we have not encountered haemorrhage from suture lines, graft or wound infection, pulmonary embolus or wound dehiscence in any of our surgically treated cases.

Discussion

The diagnosis of intestinal ischaemia remains a challenge, Dunphy in 1936 showed that seven out of 12 patients dying from acute occlusion of the superior mesenteric artery have a prodromal history of abdominal pain. Others have reported that asymptomatic patients with occluded main visceral arteries may be precipitated into acute intestinal infarction by a fall in central arterial pressure, however caused. Some surgeons therefore favour prophylactic correction of any such occlusion seen on an angiogram, whilst others would only operate on those who are symptomatic.

Clearly, diagnostic tests which could accurately identify the patient who is at risk of infarction would be valuable. Arteriography reveals the vascular lesion, but the clinical relevance of such appearances are not certain enough to allow a surgical approach.

ranges from six months to 10 years.

As already explained, because of the unreliable correlation of preoperative tests of intestinal function with symptoms and radiological findings, there seemed no point in repeating these costly and uncomfortable examinations, following surgery. Our results are therefore assessed symptomatically, and the defects of this approach are admitted. The only objective criterion of improvement is recovery of lost weight, which occurred in 13 patients. Two fatal intestinal infarcts occurred in the operative group, but this must be viewed against the fact that the incidence of infarction in untreatable intestinal arterial occlusion is yet to be discovered. Whereas, of course, it is well established that gut necrosis is often preceded by postcibal abdominal pain. The improvement in quality of life in those patients with patent reconstructions was, however, most impressive, and although they are difficult to identify, such people can be enormously benefited by appropriate surgery.

Fig. 3 Triple vessel occlusion.

Fig. 4 Carcinoma of the pancreas, showing CA obstruction and SMA stenosis.
## Table 2. Patients with confirmed IAO. Clinical and laboratory findings and results of treatment

<table>
<thead>
<tr>
<th>Case no.</th>
<th>Sex &amp; Age</th>
<th>Date of presentation</th>
<th>Duration of symptoms (months)</th>
<th>Arterial lesion CA SM IM</th>
<th>Associated vascular disease</th>
<th>Abnormalities detected</th>
<th>Operation</th>
<th>Course</th>
</tr>
</thead>
<tbody>
<tr>
<td>1*</td>
<td>M54</td>
<td>1962</td>
<td>24</td>
<td>o o o</td>
<td>Bilateral claudication angina</td>
<td>Diabetic GTT</td>
<td>Ileocaecal/aortic anastomosis</td>
<td>Symptom-free for 4 years. Death from myocardial infarction. Anastomosis patent at autopsy</td>
</tr>
<tr>
<td>2</td>
<td>M60</td>
<td>1967</td>
<td>9</td>
<td>o o o</td>
<td>Nil</td>
<td>Nil</td>
<td>Patch CA</td>
<td>Pain persisted following surgery. Lost to follow up after one year</td>
</tr>
<tr>
<td>3</td>
<td>F61</td>
<td>1969</td>
<td>6</td>
<td>o o o</td>
<td>Myocardial infarct</td>
<td>Protein-loss (2-3%)</td>
<td>Aorto SMA Anastomosis</td>
<td>Alive and well 8 years after operation. Faecal protein loss reverted to normal post operation</td>
</tr>
<tr>
<td>4</td>
<td>M48</td>
<td>1970</td>
<td>24</td>
<td>o o o</td>
<td>Nil</td>
<td>Nil</td>
<td>Patch CA</td>
<td>Symptoms persist at 9 years</td>
</tr>
<tr>
<td>5</td>
<td>F65</td>
<td>1971</td>
<td>18</td>
<td>o o o</td>
<td>Mitral stenosis</td>
<td>Glucose (8 mmol/l) GTT (diabetic) Protein loss (2-19%)</td>
<td>None</td>
<td>Refused operation. Alive with same symptoms at 10 years</td>
</tr>
<tr>
<td>6</td>
<td>M27</td>
<td>1971</td>
<td>12</td>
<td>o o o</td>
<td>Nil</td>
<td>Ischaemic stricture seen on barium study</td>
<td>Patch CA</td>
<td>Symptom-free 10 years after operation</td>
</tr>
<tr>
<td>7</td>
<td>M61</td>
<td>1972</td>
<td>12</td>
<td>o o o</td>
<td>Angina</td>
<td>Nil</td>
<td>Aorto SMA anastomosis</td>
<td>Symptom-free for 6 months, then death from cerebrovascular accident</td>
</tr>
<tr>
<td>8</td>
<td>F38</td>
<td>1974</td>
<td>6</td>
<td>o o o</td>
<td>Hypertension carotid and vertebral stenosis</td>
<td>Nil</td>
<td>Aorto SMA anastomosis</td>
<td>Death from bowel infarct one week after operation</td>
</tr>
<tr>
<td>9</td>
<td>M64</td>
<td>1974</td>
<td>9</td>
<td>o o o</td>
<td>Hypertension claudication</td>
<td>Nil</td>
<td>Aorto SMA anastomosis</td>
<td>Symptom-free at 6 years</td>
</tr>
<tr>
<td>10</td>
<td>M51</td>
<td>1976</td>
<td>24</td>
<td>o o o</td>
<td>Nil</td>
<td>Nil</td>
<td>None</td>
<td>Alive and well at 3 years, some persistent symptoms</td>
</tr>
<tr>
<td>11</td>
<td>M40</td>
<td>1977</td>
<td>36</td>
<td>o o o</td>
<td>Ileal infarct resected previously</td>
<td>Faecal Fat (8 g/day). Carotene (0.69 µmol/l). d-Xylose (25%) previous resection on barium study</td>
<td>Aorto SMA anastomosis</td>
<td>Further laparotomy 1979 – blind loop resected. Symptom-free 6 years post reconstruction</td>
</tr>
<tr>
<td>12</td>
<td>F44</td>
<td>1977</td>
<td>24</td>
<td>o o o</td>
<td>Nil</td>
<td>Nil</td>
<td>None</td>
<td>Subsequent psychiatric referral following suicide attempt</td>
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<tr>
<td>13</td>
<td>F46</td>
<td>1977</td>
<td>42</td>
<td>o o o</td>
<td>Nil</td>
<td>Nil</td>
<td>Freeing CA</td>
<td>Symptom-free at 6 months – subsequently lost to follow-up</td>
</tr>
<tr>
<td>No.</td>
<td>Code</td>
<td>Year</td>
<td>Age</td>
<td>Vessel</td>
<td>Cause</td>
<td>Treatment</td>
<td>Complication</td>
<td></td>
</tr>
<tr>
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</tr>
<tr>
<td>14</td>
<td>M44</td>
<td>1977</td>
<td>12</td>
<td>o o o</td>
<td>Myocardial infarction</td>
<td>Nil</td>
<td>Aorto SMA anastomosis</td>
<td>Death 3 days post operation from myocardial infarct</td>
</tr>
<tr>
<td>15</td>
<td>F23</td>
<td>1980</td>
<td>24</td>
<td>o o o</td>
<td>Small vessel disease</td>
<td>Nil</td>
<td>None</td>
<td>Symptoms controlled on steroids at 2 years.</td>
</tr>
<tr>
<td>16</td>
<td>F35</td>
<td>1981</td>
<td>30</td>
<td>o o o</td>
<td>Nil</td>
<td>Nil</td>
<td>Patch angioplasty CA</td>
<td>Renal function deteriorating</td>
</tr>
<tr>
<td>17</td>
<td>F45</td>
<td>1981</td>
<td>18</td>
<td>o o o</td>
<td>Angina claudication BK amputation</td>
<td>Fe (12 μmol/l) TIBC (103 μmol/l)</td>
<td>Patch angioplasty SMA</td>
<td>Symptom-free one year post operation</td>
</tr>
<tr>
<td>18</td>
<td>M47</td>
<td>1981</td>
<td>30</td>
<td>o o o</td>
<td>Splenic flexure resected for ischaemia</td>
<td>Nil</td>
<td>Laparotomy only</td>
<td>Symptom-free for 6 months, then re-occlusion.</td>
</tr>
<tr>
<td>19</td>
<td>F56</td>
<td>1981</td>
<td>24</td>
<td>o o o</td>
<td>Hypertension</td>
<td>Nil</td>
<td>Patch angioplasty CA</td>
<td>Infarction. Resection. Fistula. Death 9 months post operation</td>
</tr>
<tr>
<td>20</td>
<td>F61</td>
<td>1981</td>
<td>18</td>
<td>o o o</td>
<td>Claudication BK amputation</td>
<td>Nil</td>
<td>Patch angioplasty CA and SMA</td>
<td>Symptom-free for 6 months, then recurrence of mild discomfort</td>
</tr>
<tr>
<td>21</td>
<td>F38</td>
<td>1981</td>
<td>48</td>
<td>o o o</td>
<td>Aortitis previous SMA surgery</td>
<td>Nil</td>
<td>None</td>
<td>Severe distal arteritis. Symptoms persist on steroids</td>
</tr>
<tr>
<td>22</td>
<td>F65</td>
<td>1981</td>
<td>12</td>
<td>o o o</td>
<td>Type II aortic dissection</td>
<td>d-Xylose (16%) gallstones on ultrasound</td>
<td>Reimplantation of SMA</td>
<td>Symptoms relieved. Lost to follow-up a few weeks post operation</td>
</tr>
<tr>
<td>23</td>
<td>M54</td>
<td>1981</td>
<td>6</td>
<td>o o o</td>
<td>Hypertension</td>
<td>DU on endoscopy. Gallstones on ultrasound</td>
<td>Patch angioplasty CA and IMA</td>
<td>Re-operation for small bowel volvulus at one week. Cholecystectomy at 6 months.</td>
</tr>
<tr>
<td>24</td>
<td>M73</td>
<td>1982</td>
<td>12</td>
<td>o o o</td>
<td>Hypertension angina myocardial infarct</td>
<td>Nil</td>
<td>None</td>
<td>Symptoms continue at one year. Considered unacceptable operative risk</td>
</tr>
<tr>
<td>25</td>
<td>F77</td>
<td>1983</td>
<td>6</td>
<td>o o o</td>
<td>Angina</td>
<td>Nil</td>
<td>None</td>
<td>Symptoms continue at 6 months</td>
</tr>
<tr>
<td>26</td>
<td>M61</td>
<td>1983</td>
<td>18</td>
<td>o o o</td>
<td>Previous aortic surgery, angina</td>
<td>Fe (6 μmol/l) TIBC (105 μmol/l)</td>
<td>Patch angioplasty SMA</td>
<td>Symptoms-free at 6 months</td>
</tr>
<tr>
<td>27</td>
<td>F66</td>
<td>1983</td>
<td>12</td>
<td>o o o</td>
<td>Hypertension</td>
<td>Pyloric ulcer on endoscopy</td>
<td>Aorto SMA bypass</td>
<td>Symptom-free for 9 months. Graft then occluded and pain recurred. Patency restored and confirmed angiographically. Well to date.</td>
</tr>
<tr>
<td>28</td>
<td>M55</td>
<td>1984</td>
<td>36</td>
<td>o o o</td>
<td>Previous aortic surgery</td>
<td>Faecal fat (113G/day) d-Xylose (10%)</td>
<td>Patch SMA</td>
<td>Died at 10 days – myocardial infarction</td>
</tr>
</tbody>
</table>

* See ref 30. o = Normal vessel. o = Stenosis. o = Occlusion.
A decision needs to be made. This is because asymptomatic intestinal arterial occlusion is common\textsuperscript{3,25} and how much it constitutes a threat to life has yet to be decided. Almost certainly, resting blood flow in such patients is within the normal range, and the pain which some of them experience after eating represents failure of the gut to produce the hyperaemia which the processes of propulsion and digestion require. This would equate with what is observed to happen in other vascular territories such as the heart and the legs. There is ample evidence that resting blood flow in the claudicating limb is normal, and deficiencies appear only when the circulation is challenged by exercise. If the alimentary tract behaves in the same way it is not surprising that tests of intestinal function carried out in the

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Fig. 5  Surgical techniques for reconstruction of the visceral arteries.
resting hospital patient are unremarkable, even given gross mesenteric arterial occlusion.

By means of various compensatory mechanisms to redistribute blood flow in the wall, the absorptive area appears capable of working well up to the near the point of irreversible damage. Because of available collateral circulation (particularly in the coeliac and inferior mesenteric territories) major occlusion in the visceral arteries can be compensated, to the extent that the gut is not ischaemic and no symptoms occur. This compensation may additionally be effected by internal switching mechanisms at submucosal or mucosal level, and by the operation of the counter current exchange mechanism within the villus. In any event, the tips of the villi which are the main absorptive area only receive some 8% of total flow. As arterial input is still further reduced, then pain arises from the ischaemic muscle which does not receive the increased flow required by peristalsis, but mucosal flow is preserved and absorptive/exsorptive capacity remains impaired. Hensen et al showed that total splanchnic flow rises in (arteriographically) normal subjects after a meal, and that this rise fails to occur in patients with intestinal arterial occlusion, but can be restored by suitable surgery.

As vascular disease progresses, resting flow eventually falls below the level required to maintain the mucosal defences against osmolar and bacterial challenge, and infarction (focal or massive) results. One might postulate four stages: (0) normal; (1) an arterial lesion is present, but compensated, so that both resting and post-cibal flow are unimpaired, and there are no symptoms; (2) the arterial disease has progressed to the point where resting flow is normal but reactive hyperaemia cannot occur. This is signalled by the occurrence of postcibal pain; (3) The blood supply is so deficient that resting flow is reduced and perhaps minor impairments of cellular function and small, focal infarcts are occurring. This is analogous to 'rest pain' in the ischaemic limb; (4) the bowel in infarcted.

It is the identification of stage 3 which has been the object of so much clinical and laboratory endeavour over the last 80 years, since Schnitzler first postulated the concept of 'abdominal claudication', and it must be admitted that little progress has been made. Many authors have commented that tests of intestinal function have little predictive value, though no formal prospective study has up till now been carried out. Nonetheless, it has often been stated that chronic intestinal arterial occlusion is associated with malabsorption. Our data suggest that this is not the case, and that the weight loss which occurs is because of diminished intake. Quite different, of course, is the patient who sustains an acute ischaemia and is rescued by emergency surgery from a threatened infarction. Here the bowel is seriously damaged, perhaps by liberated superoxide radicles and a prolonged period of malfunction follows while the mucosa regenerates.

To return to the questions posed in the introduction, it is possible to claim, in the light of our experience and that previously reported, that reduction in blood supply to the gut can give rise to symptoms before infarction occurs, but these are of a variable and inconstant nature and difficult to identify. They tend to occur in an age group where other causes of abdominal pain abound, many of them more easily treated, and it is important to exclude these before deciding on a vascular origin for the pain. Moreover, arterial occlusions can occur without symptoms and probably without influencing blood flow, and can coexist with or be the result of, malignant tumours (Fig. 4).

The syndrome of intestinal angina is rare, and the academic interest which it has aroused is not entirely
justified. This is shown by the fact that a unit particularly interested in the problem and having wide sources of referral, but using strict criteria, has authenticated only 28 cases in 20 years. Hertzer and Rogers report a similar experience.

No non-invasive test is of use in identifying such a patient, nor in predicting who is at risk from infarction. It is occasionally possible to relieve abdominal pain and restore weight loss by reconstructing the visceral arteries, and in the most favourable cases this can be a spectacularly successful surgical exercise. These ambitious operations are not devoid of risk, however, even in centres accustomed to carrying them out. It is likely that failures are under reported.

For the future, what is needed is a simple non-invasive method of measuring intestinal blood flow before and after challenge, which can be correlated with the results of the non-invasive radiological methods now available.

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