Sporadic duodenal adenoma is an uncommon finding. In a large series from Germany, only 6.9% of 378 duodenal polyps found at 25 000 upper gastrointestinal endoscopies were adenomatous. Another series similarly found that only 0.4% of 584 endoscopy patients had duodenal polyps, of which 7% were adenomatous. Most adenomas were found incidentally at endoscopy but occasionally caused bleeding or obstruction of the duodenum or ampulla of Vater. The distribution of small bowel adenomas shows a predominance at the ampulla and periamputillary region, with decreased numbers proximally in the duodenum and distally in the small bowel, with a small peak in the distal ileum.

Adenomas of the small and large intestine are neoplasias—that is, they lie along a multistep pathway to carcinogenesis—effected by numerous genetic and epigenetic events. Curiously however, despite numerous phenotypic similarities between small and large intestinal epithelia, including a very high cellular turnover, small intestinal neoplasia is very rare compared with its colorectal counterpart. None the less, small intestinal carcinoma is associated with colorectal carcinoma and vice versa, but not gastric or oesophageal carcinoma.

The extent to which a duodenal adenoma is associated with colorectal neoplasia is not well described. A previous clinicopathological study described 21 cases of duodenal adenomas, of which 11 underwent colonoscopy. Four of these 11 cases were classified as having familial adenomatous polyposis. Of the remaining seven cases of sporadic duodenal adenomas, four (37%) were found to have colorectal neoplasia.

We aimed to see if finding a sporadic duodenal adenoma was a sign of associated colorectal neoplasia. Identifying such an association would allow earlier detection of colorectal neoplasia, as well as providing some insight into duodenal and colorectal carcinogenic pathways. We determined the frequency of colorectal neoplasia in a relatively large group of patients with sporadic duodenal adenomas and compared this with a matched control group of symptomatic patients presenting for endoscopic procedures. Furthermore, we also compared the frequency of colorectal cancer in patients with sporadic duodenal adenomas with the incidence of colorectal cancer in the population.

**PATIENTS AND METHODS**

Patients with a diagnosis of duodenal adenoma were identified using the pathology coding databases of the two major metropolitan university teaching hospitals (Sir Charles Gairdner Hospital and Royal Perth Hospital) in Perth, Western Australia, during the period 1992–2002. Endoscopic reports of the identified patients were retrieved to confirm the location of the adenomas and to complete the descriptive record of the cases. Patients with, or belonging to, familial adenomatous polyposis (FAP) and hereditary non-polyposis colorectal cancer (HNPCC) kindreds were excluded. Clinical records for identified patients were examined for associated synchronous or metachronous diagnoses of colorectal neoplasia. Only the most histologically advanced colorectal lesion was recorded for each patient.

Duodenal adenoma location, size, and histology were recorded. Similar details were recorded for colorectal neoplasms, when identified. Patients with multiple colorectal neoplasms were categorised according to the most advanced lesion found. Advanced adenomas (duodenal and colorectal) were defined as having a size ≥10 mm, the presence of a villous component, or high grade dysplasia. Indications

**Objective:** The objective of this study was to assess the association between colorectal neoplasia and sporadic duodenal adenoma.

**Methods:** A retrospective case control study was conducted using the databases of two major teaching hospitals in Western Australia. The frequency of colorectal neoplasia in patients with sporadic duodenal adenomas was compared with that in a control group of patients presenting for endoscopies. The frequency of colorectal cancer in duodenal adenoma patients was also compared with the population incidence.

**Results:** Of 56 sporadic duodenal adenoma patients, 34 (61%) had been colonoscoped. When comparing the findings between patients with sporadic duodenal adenoma and an endoscoped control group, all colorectal neoplasias were significantly more common in the duodenal adenoma group (56% vs 33%; odds ratio (OR) 2.4 (95% confidence intervals (CI) 1.1–5.4)). Although finding either advanced colorectal adenoma or cancer was also more common in duodenal adenoma patients (38% vs 19%; OR 2.3 (95% CI 1.0–5.2)), as was finding colorectal cancer alone (21% vs 8%; OR 3.0 (95% CI 1.0–9.1)), the results were not statistically significant. However, the incidence of colorectal cancer was much greater in duodenal adenoma patients than in the population (p<0.001).

**Conclusions:** Sporadic duodenal adenoma has a clinically important association with colorectal neoplasia. Thus patients with duodenal adenomas should undergo colonoscopy to detect colorectal neoplasia.
for upper gastrointestinal endoscopy and colonoscopy were recorded.

Findings in patients with duodenal adenomas were compared with: (1) endoscoped control patients, who were randomly selected age and sex matched patients presenting to Sir Charles Gairdner Hospital for both upper endoscopy and colonoscopy; and (2) population statistics, in which the incidence of colorectal cancer in the age and sex matched population was obtained from published Australian statistics. These statistics are compiled by the Australian Institute of Health and Welfare and are available on the internet (http://www.aihw.gov.au/publications/can/ca98).11

Conditional logistic regression derived odds ratios (OR) and 95% confidence intervals (CI) were used to compare the rates of colorectal neoplasia in the sporadic duodenal adenoma group with those of the endoscoped controls, and the χ² test with continuity correction was used to compare duodenal adenoma patients with the population.

RESULTS
From January 1992 until June 2002 (inclusive), 57 618 upper gastrointestinal endoscopies were performed on 39 784 patients. A total of 100 patients were identified on the basis of the pathology code for duodenal adenoma. All identified specimens were obtained endoscopically. However, 44 patients were excluded because of FAP (24), HNPCC (1), invasive carcinoma (10), and other diagnoses (9).

Sporadic duodenal adenomas
Of the remaining 56 patients with sporadic duodenal adenomas, 31 were males and 25 were females. Mean age of the patients was 67 years (males 71 years; females 62 years), with a range of 38–91 years. Indications for upper gastrointestinal endoscopy are shown in table 1, and were predominantly for anaemia, abdominal pain, and indications requiring endoscopic retrograde cholangiopancreatography.

A single duodenal adenoma was identified in each of the 56 patients. Adenomas were located in the duodenal cap (6 (11%)), second part of the duodenum (32 (57%)), or ampullary region (18 (32%)). Adenoma size was accurately reported in 44 patients, with a mean and median of 15 mm (range 3–70 mm). Large adenomas (>10 mm) were found in at least 28 patients (50%). In 10 of 12 patients where adenoma size was not precisely reported, the ampulla was involved. Villosity was reported in part or in toto in 21 adenomas (38%). High grade dysplasia was reported in two adenomas (4%). Thus in at least 34 patients (61%) the adenomas were advanced. There were no statistically significant relationships between adenoma location, size, or histology within this study group.

Colorectal neoplasia associated with duodenal adenomas
Of the 56 patients with duodenal adenomas, 34 (61%) had at least one colorectal adenoma. Colorectal neoplasia was found in 19 of these 34 patients (56%). Colorectal cancer was found in seven patients (21%), advanced colorectal adenoma was found in six patients (18%), and non-advanced colorectal adenoma was found in the remaining six patients (18%). Thus 13 of the 34 patients with colorectal neoplasia (38%) had either colorectal cancer or an advanced adenoma. Of the seven patients with colorectal cancer, eight cancers were actually found, with four located in the caecum, one in the ascending colon, two in the sigmoid, and one in the rectum. There was no significant relationship between different locations of the duodenal adenomas and the presence or nature of colorectal neoplasia. There was also no significant relationship between whether a duodenal adenoma was advanced or non-advanced and the presence or nature of colorectal neoplasia.

To confer clinical perspective to our identified rate of colorectal neoplasia in patients with sporadic duodenal adenomas, we compared each case with three randomly selected age and sex matched controls, consisting of symptomatic patients undergoing endoscopy and colonoscopy. Control patients were also obtained from our endoscopic database. Of 102 endoscoped control patients, 34 were found to have colorectal neoplasia (33%). This consisted of colorectal cancer in eight patients (8%), advanced adenomas in 11 patients (11%), and non-advanced adenomas in 15 patients (15%). In all, 19 patients (19%) had either colorectal cancer or an advanced colorectal adenoma. Importantly, patients with sporadic duodenal adenomas, when compared with endoscoped controls, had a significantly greater risk of associated colorectal neoplasia (55% v 33%; p = 0.03; OR 2.4 (95% CI 1.1–5.4)). Although duodenal adenoma patients when compared with endoscoped controls were more likely to have advanced colorectal neoplasia—that is, advanced adenoma or cancer (38% v 19%; p = 0.05; OR 2.3 (95% CI 1.0–5.2)) and colorectal cancer alone (21% v 8%; p = 0.05; OR 3.0 (95% CI 1.0–9.1))—the results were not statistically significant because the confidence intervals included 1.0 (tables 2 and 3).

Colorectal cancer in patients with duodenal adenoma compared with the population
We further compared the risk of colorectal cancer between patients with sporadic duodenal adenoma and that of age and sex matched population controls, based on Australian cancer statistics.11 The cumulative incidence in 34 matched population controls yielded an expected finding of 0.1 cancers (0.3%) in this group. This contrasts with the observed findings of seven cancers in 34 patients with duodenal

Table 1  Indications for upper gastrointestinal endoscopy in sporadic duodenal adenoma patients

<table>
<thead>
<tr>
<th>Indication</th>
<th>n</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain</td>
<td>14</td>
<td>Reflux (5), dyspepsia (5), epigastric pain (4)</td>
</tr>
<tr>
<td>Anaemia</td>
<td>13</td>
<td>Anaemia or iron deficiency (13)</td>
</tr>
<tr>
<td>Melena</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>ERCP</td>
<td>14</td>
<td>Stones (6), abnormal LFT (2), pancreatitis (3), jaundice (3)</td>
</tr>
<tr>
<td>Abnormal x ray</td>
<td>2</td>
<td>Duodenal lesion (1), gastric ulcer (1)</td>
</tr>
<tr>
<td>Dysphagia</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Diarrhoea</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Nausea and vomiting</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Not recorded</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>56</td>
<td></td>
</tr>
</tbody>
</table>

ERCP, endoscopic retrograde cholangiopancreatography; LFT, liver function tests.
Sporadic duodenal adenoma is associated with colorectal neoplasia 263

than the general population. This was confirmed by our already having a much higher incidence of colorectal neoplasia. It is noteworthy that being symptomatic, the control patients resolve the issue more precisely. It is none the less significant. A larger series, albeit of a rare condition, may suggest that our series lacked power to establish statistical significance. When compared with an age and sex matched group of controls, and the population (n = 34)

endoscoped controls, and the population

<table>
<thead>
<tr>
<th>Colonic neoplasia found</th>
<th>Controls (n = 102)</th>
<th>p Value</th>
<th>OR</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>All colorectal neoplasia</td>
<td>34 (33%)</td>
<td>0.03</td>
<td>2.5</td>
<td>1.1–5.4</td>
</tr>
<tr>
<td>Cancer or advanced adenoma</td>
<td>19 (19%)</td>
<td>0.05</td>
<td>2.7</td>
<td>1.0–5.2</td>
</tr>
<tr>
<td>Cancer</td>
<td>8 (8%)</td>
<td>0.05</td>
<td>3.0</td>
<td>1.0–9.1</td>
</tr>
<tr>
<td>Non-advanced adenoma</td>
<td>11 (11%)</td>
<td>0.44 NS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DA v endoscoped controls</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DA v population (n = 34)</td>
<td></td>
<td>&lt;0.001</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Patients were classified according to the most advanced lesion found at colonoscopy.


denomas (21%) and was highly statistically significant (p<0.001).

**Indications for colonoscopy (table 4)**

Of the 34 patients with duodenal adenomas who were colonoscopy, 15 (44%) of the procedures were for blood loss indications—namely, anaemia (13) or frank bleeding (2). This was not significantly different from 52 of 102 patients (51%) in the endoscoped control group (p = 0.49). Importantly, 15 patients underwent colonoscopy purely on the basis of identification of a duodenal adenoma and seven (47%) of these were found to harbour colorectal neoplasia. One patient was found to have colorectal cancer five years after identification of an ampullary villous adenoma, and another patient was found to have a sigmoid carcinoma having presented with rectal bleeding three months after incidental identification of a sporadic duodenal advanced adenoma.

**DISCUSSION**

In this study we have demonstrated a strong association between sporadic duodenal adenomas and colorectal neoplasia. When compared with an age and sex matched group of symptomatic patients undergoing upper endoscopy and colonoscopy, the yield of colorectal neoplasia was significantly greater for patients with duodenal adenoma (55% v 33%; p = 0.03). Advanced colorectal adenoma or cancer was also more common among patients with duodenal adenomas (38% v 19%; p = 0.05; OR 2.3 (95% CI 1.0–5.2)), as was the finding of cancer alone (21% v 8%; p = 0.05; OR 3.0 (95% CI 1.0–9.1)), although by definition a confidence interval including 1.0 is not statistically significant. The odds ratios of 2.3 and 3.0 are clinically important in magnitude, but the wide confidence intervals (1.0–5.2 and 1.0–9.1, respectively) suggest that our series lacked power to establish statistical significance. A larger series, albeit of a rare condition, may resolve the issue more precisely. It is none the less noteworthy that being symptomatic, the control patients already have a much higher incidence of colorectal neoplasia than the general population. This was confirmed by our finding that the rate of colorectal cancer in patients with duodenal adenomas was substantially greater than the age and sex matched cumulative incidence rates, as determined by population statistics (21% v 0.3%; p<0.001).

The present study is the largest of its type, involving the highest number of sporadic duodenal adenomas. Our findings add to those of a previous small study identifying colorectal neoplasia in four of seven patients (57%) with sporadic duodenal adenomas. Previous studies have shown that the diagnosis of small intestinal carcinoma increases the risk of finding colorectal cancer but not other gastrointestinal tract malignancy. For instance, Neugut and Santos revealed that after a diagnosis of primary adenocarcinoma of the small intestine, the relative risk of finding a primary adenocarcinoma of the colon or rectum was 5.0 (95% CI 2.3–9.4) in men and 3.7 (95% CI 1.3–8.0) in women compared with expected numbers derived from population based tumour registries. We excluded duodenal or ampullary carcinomas from our analysis because these were often found at advanced stages with poor prognosis, and patients were therefore not subjected to further investigations such as colonoscopy. Furthermore, separating analyses of duodenal adenomas from carcinomas may yield important pathogenetic insights on neoplastic initiation and promotion as only a small subset of adenomas progresses to carcinoma.

A possible explanation for the strong association between duodenal adenomas and colorectal neoplasia in this study may be that some patients had undiagnosed FAP. Although attenuated FAP is a possibility, the commoner FAP phenotypes are unlikely given that multiple polyps, if not gross carpeting, of the colonic mucosa is frequently seen. Duodenal adenomas in FAP are also frequently multiple, a feature absent in this patient series. Recently, biallelic mutations in the MYH gene encoding a base excision-repair protein have been described, in which patients develop multiple colorectal neoplasias and occasional associated duodenal adenomas. It is possible that some of our cases were due to this recessive syndrome. Some of the patients in our series may also have undiagnosed HNPCC, which increases the risk of small intestinal cancer by 25-fold over the population average.

| Table 2 Proportion of duodenal adenoma patients and endoscoped controls with identified colorectal neoplasia |
|---------------------------------|---------|--------|
| Colorectal neoplasia found*     | Duodenal adenoma (n = 34) | Controls (n = 102) | p Value |
| All colorectal neoplasia        | 19 (56%) | 34 (33%) | 0.03   |
| Cancer or advanced adenoma      | 13 (38%) | 19 (19%) | 0.05   |
| Cancer                          | 7 (21%)  | 8 (8%)  | 0.05   |
| Advanced adenoma                | 6 (18%)  | 11 (11%) | 0.44 NS|
| Non-advanced adenoma            | 6 (18%)  | 15 (26%) | 0.51 NS|

*Patients were classified according to the most advanced lesion found at colonoscopy.
these cancers, approximately one third involve the duode-
um.14 Indeed, spontaneous mutations can occur in the mis-
mismatch repair genes in the absence of a family history of HNPCC associated cancer. Future examination for microsatellite
instability and MLH1/MSH2 protein expression in duodenal
adenomas in this series would be useful to identify
potential undiagnosed HNPCC cases.

Another explanation for the association between duodenal
and colonic neoplasia may be that they share common
pathogenetic pathways. This may include genetic and/or
environmental factors. The data on this issue are currently
unclear. For instance, analyses of genetic mutational steps in
duodenal and colorectal carcinogenesis show differences in
the frequency and site of APC gene mutations20–21 while
showing some positive correlations for late events, such as
p53 mutations.20 21 Furthermore, positive correlations in
protein and fat consumption have been found between the
small intestinal and colorectal carcinoma.9

Given that many of the colorectal neoplasias identified in
patients with duodenal adenomas were found on investiga-
tion for colonic symptoms, is it appropriate to reserve
colonoscopy only for those duodenal adenoma patients who
have, or develop, symptoms referable to colonic disease? This
study found that seven of 15 patients (47%) who underwent
colonoscopy purely on the basis of the finding of a duodenal
adenoma were subsequently found to harbour colorectal
neoplasia. Although the number of patients was small, we
believe that colonoscopy is indicated for all duodenal
adenoma patients. An analogous situation arises when
asymptomatic patients found to have left colonic adenomas
on sigmoidoscopy are recommended to have complete
colonoscopy due to a 30% incidence of synchronous proximal
colonic neoplasia.22–24 Importantly, if colonoscopy was
automatically recommended in patients with a sporadic duodenal
adenoma in this study, one sigmoid cancer would have been
discovered three months prior to presenting with rectal
bleeding, while another cancer may have been prevented,
by detection of an advanced neoplastic lesion five years earlier.
Furthermore, as neither specific location nor histology of the
duodenal adenoma was useful in determining who would derive greater benefit from colonoscopy, it seems reasonable
to recommend colonoscopy for all patients identified with a
duodenal adenoma.

In summary, this study confirms a significant association
between duodenal adenoma and colorectal adenoma and
carcinoma outside of recognised hereditary colorectal cancer
syndromes. We therefore recommend colonoscopy in all
patients found to have a duodenal adenoma and consider it
prudent to maintain ongoing colonoscopic surveillance for
these patients, in a similar fashion to those with identified
colorectal neoplasia.

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Bruce Latham for extracting histopathology data; and Dr Karen Byth
for biostatistics advice.

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REFERENCES
1 Hochter W, Weingart J, Sebh J, et al. Duodenal polyps. Incidence,
histologic substrate and significance. Dig Med Wochenschr
2 Jepsen JM, Persson M, Jakobsen NO, et al. Prospective study of prevalence
and endoscopic and histopathologic characteristics of duodenal polyps in
patients submitted to upper endoscopy. Scand J Gastroenterol
3 Sillier F. Investigations on the significance of the adenoma-carcinoma
4 Muto T, Bussey HJR, Marson BC. The evolution of cancer of the colon and
5 Fearon ER, Vogelstein B. A genetic model for colorectal tumorigenesis.
6 Perzin KH, Bridge MF. Adenomas of the small intestine: A clinicopathologic
review of 51 cases and a study of their relationship to carcinoma. Cancer
7 Arber N, Neugut AI, Weinstein IB, et al. Molecular genetics of small bowel
8 Neugut AI, Santos J. The association between cancers of the small and large
9 Lowenfels AB, Sonnt A. Distribution of small bowel tumors. Cancer Lett
10 Seifert E, Schulte F, Stolle M. Adenoma and carcinoma of the duodenum and
papilla of Vater: a clinicopathologic study. Am J Gastroenterol
1992;87:37–42.
11 Atkins WS, Marson BC, Cuzick J. Long-term risk of colorectal cancer
13 Australian Institute of Health and Welfare (AIHW) and Australasian
Canberra: AIHW (Cancer Series No 17), 2001:AIIHW cat No CAN 12
sequence in the duodenum of patients with familial adenomatous polyposis.
15 Yao T, Iida M, Ohsato K, et al. Duodenal lesions in familial polyposis of the
16 Sieber OM, Lipton L, Croah M, et al. Multiple colorectal adenomas, classic
adenomatous polyposis, and germ-line mutations in MTH. N Engl J Med
17 Watson P, Lynch HT. Extracolonic cancer in hereditary nonpolyposis
bowel carcinoma in hereditary nonpolyposis colorectal cancer. Cancer

Table 4 Indications for colonoscopy in duodenal adenoma patients and endoscoped
control patients

<table>
<thead>
<tr>
<th>Indication</th>
<th>DA (n = 34)</th>
<th>Colorectal neoplasia found for indication</th>
<th>Endoscoped controls (n = 102)</th>
<th>Colorectal neoplasia found for indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anemia</td>
<td>13 (38%)</td>
<td>7 (54%)</td>
<td>27 (26%)</td>
<td>12 (44%)</td>
</tr>
<tr>
<td>Frank bleeding</td>
<td>2 (6%)</td>
<td>2 (100%)</td>
<td>25 (25%)</td>
<td>9 (36%)</td>
</tr>
<tr>
<td>Altered bowel habit</td>
<td>2 (6%)</td>
<td>2 (100%)</td>
<td>14 (14%)</td>
<td>4 (29%)</td>
</tr>
<tr>
<td>Diarrhoea</td>
<td>2 (6%)</td>
<td>0</td>
<td>5 (5%)</td>
<td>0</td>
</tr>
<tr>
<td>Constipation</td>
<td>2 (6%)</td>
<td>0</td>
<td>2 (2%)</td>
<td>0</td>
</tr>
<tr>
<td>Abnormal x ray</td>
<td>2 (6%)</td>
<td>0</td>
<td>2 (2%)</td>
<td>0</td>
</tr>
<tr>
<td>Weight loss</td>
<td>3 (3%)</td>
<td>1 (33%)</td>
<td>3 (3%)</td>
<td>1 (33%)</td>
</tr>
<tr>
<td>Pain</td>
<td>3 (3%)</td>
<td>2 (67%)</td>
<td>3 (3%)</td>
<td>2 (67%)</td>
</tr>
<tr>
<td>Family history CRC</td>
<td>2 (6%)</td>
<td>1 (50%)</td>
<td>6 (6%)</td>
<td>3 (50%)</td>
</tr>
<tr>
<td>Follow up polyp</td>
<td>2 (6%)</td>
<td>1 (50%)</td>
<td>6 (6%)</td>
<td>3 (50%)</td>
</tr>
<tr>
<td>Duodenal adenoma</td>
<td>15 (44%)</td>
<td>7 (47%)</td>
<td>102</td>
<td>34 (33%)</td>
</tr>
<tr>
<td>Total</td>
<td>34</td>
<td>19 (56%)</td>
<td></td>
<td>102</td>
</tr>
</tbody>
</table>

DA, duodenal adenoma patients; CRC, colorectal cancer.


Sporadic duodenal adenoma is associated with colorectal neoplasia

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