Excessive alcohol consumption favours high risk polyp or colorectal cancer occurrence among patients with adenomas: a case control study

M Bardou, S Montembaut, V Giraud, A Balian, E Borotto, C Houdayer, F Capron, J-C Chaput, S Naveau

Background and aims: Excessive alcohol consumption is a risk factor for developing colorectal adenomas. This study aimed to investigate the influence of excessive alcohol consumption on the occurrence of high risk polyps (adenoma ≥10 mm, villous component, high grade dysplasia) or colorectal cancer among patients with at least one colonic adenoma.

Patients and methods: Three groups of patients with at least one colorectal adenoma were included in a case control study: 401 heavy drinkers (group HD, mean daily alcohol intake 117 (SD 4) g/day for a mean duration of 22 (SD 0.6) years), aged 57 (0.5) years (78% men); 152 patients suffering from irritable bowel syndrome (IBS), aged 61 (0.9) years (57% male); and 108 patients with a family history (FH) of colorectal adenoma or cancer, aged 55 (1) years (64% male). Exclusion criteria were: anaemia, haematochezia, personal history of colorectal adenoma or cancer, and for groups HD and IBS a family history of colorectal adenoma and/or cancer. Relative risks were estimated by the odds ratio (OR) using a logistic regression model and were expressed with 95% confidence interval (CI).

Results: After age and sex adjustment, the likelihood of having an adenoma ≥10 mm was higher in group HD than in the IBS group (OR 1.8, 95% CI (1.2–2.7)) and the likelihood of having high risk adenomas or cancer was higher in group HD compared with the IBS group (OR 1.6, 95% CI (1.2–2.1)) and the FH group although this was not significant (OR 1.6, 95% CI (0.97–2.6) (p=0.081); 90% CI (1.03–2.4)). After age and sex adjustment, the likelihood of having an adenoma with high grade dysplasia or cancer was higher in group HD than in the IBS group (OR 1.7, 95% CI (1.02–2.8)) or group FH, although this was not significant (OR 3.7, 95% CI (0.98–15) (p=0.076); 90% CI (1.10–12.47)).

Conclusion: In patients with at least one colorectal adenoma, excessive alcohol consumption increases the likelihood of developing high risk adenomas or colorectal cancer.

MATERIAL AND METHODS

Patients
All patients older than 40 years who were referred to our unit between January 1987 and December 1996 for excessive alcohol consumption (heavy drinkers (group HD) >50 g of alcohol/day for the year before admission) underwent screening colonoscopy if they had no contraindications. Two other groups of patients who drank less than 30 g/day were included during the same period: patients who were undergoing colonoscopy because of a family history (group FH) of colorectal cancer and patients whose relatives suffered from colorectal cancer were included in the FH group. The control group consisted of patients with irritable bowel syndrome (IBS) who had undergone colonoscopy because of a family history of colorectal cancer. Patients of the IBS group did not present any personal history of colorectal cancer either.

Conclusion: The risk of developing colorectal cancer increased with alcohol consumption in case-control studies.

Abbreviations: IBS, irritable bowel syndrome; OR, odds ratio.
cancer or adenomatous polyps (no patient had a family history of adenomatous polyposis coli) and patients who were suffering from irritable bowel syndrome (IBS). IBS was diagnosed using the Rome II criteria.20

Information on alcohol consumption was recorded using a specific standardised 25 item questionnaire, which has recently been revalidated.21 Patients were asked about their drinking habits a few days before colonoscopy during consultation for outpatients or during hospitalisation for inpatients. The questionnaires were completed in the same standardised way by a senior gastroenterologist in our unit. We did not use a self administrated questionnaire as French people are not familiar with them. Nalpas and colleagues26 described recently in a French population that anonymous self administrated questionnaires were associated with a high level of incomplete questionnaires or with responses that were unusable. This 25 item questionnaire was used in the same way for outpatients and inpatients in the three groups. The same senior gastroenterologists interviewed outpatients and inpatients. Patients were asked about their drinking habits over the previous five years before admission to hospital for alcoholic disorders and total duration of alcohol abuse. The average daily intake was calculated for each beverage (beer, wine, aperitifs) and expressed in grams of pure ethanol. Total daily consumption of ethanol was obtained by adding the amounts consumed for each type of beverage. It took approximately 15 minutes to complete a questionnaire. The patient’s family was also interviewed if possible.

All patients in the three groups were asked about their personal and family history of colorectal adenomas or cancer, and about their bowel habits, based on the Rome criteria, to avoid overlapping of the three groups. Criteria for exclusion were anaemia, haematochezia, familial polyposis, inflammatory bowel disease, personal history of colorectal adenomas or cancer and, except for the FH group, family history of colorectal adenomas or cancer. Alcoholic patients were not included if they had uncompensated cirrhosis. Patients were included in the study only if colonoscopy reached the caecum. During the 10 year period of the study, among the 4517 patients who were referred to our unit for a screening colonoscopy without any clinical or biological symptoms of intestinal disease, 661 patients were entered into the study because of the presence of at least one colorectal adenoma. A total of 401 patients were included in the HD group (mean daily alcohol intake 117 (SD 4) g/day for a mean duration of 22 (SD 0.6) years), 152 in the IBS group, and 108 in the FH group. Age and sex distributions were different between the three groups (table 1).

### Cases and controls

As the exposing factor being studied was excessive alcohol consumption, three case control studies were performed.

- In the first study, cases (n=84) were patients with at least one adenomatous polyp larger than 10 mm, and controls (n=489) were those with adenomatous polyps <10 mm and without villous contingent high grade dysplasia or cancer.
- In the second study, cases (n=172) were patients with at least one high risk colorectal polyp (≥10 mm and/or with a villous component and/or with high grade dysplasia) or a histologically proved colorectal cancer, and controls (n=489) were those with adenomas <10 mm and without high grade dysplasia, a villous component, or colorectal cancer.
- In the third study, cases (n=37) were patients with high grade dysplasia or colorectal cancer, and controls (n=624) were those with adenomatous polyps, irrespective of size, and no high grade dysplasia or colorectal cancer.

All colonoscopies had to reach at least the caecum and were performed by a senior endoscopist. In all patients with at least one adenoma of the colon, the location and number of adenomas was recorded as was the location of high risk adenomas or cancer.

Histopathological diagnosis was performed in the anatomo-pathology unit of our hospital and the degree of dysplasia was classified according to the World Health Organisation classification system.23

### Statistical analysis

Univariate analysis studies were performed using the $\chi^2$ method for dichotomous variables and the Student’s $t$ test for continuous variables.

Relative risks in the three groups of patients (IBS, FH and HD) of cases and controls were compared using the global $\chi^2$ method.

Relative risks of adenomas ≥10 mm, high risk adenoma or cancer, and high grade dysplasia or cancer, according to study group, were estimated by odds ratios (ORs) adjusted for age and sex using a stepwise logistic regression. In this model, if the value of the probability level is less than some predefined alpha level, say 0.1, then the variable is said to be statistically significant. The statistical significance of each studied variable was tested using the maximum likelihood method.

### RESULTS

#### Characteristics of the adenomas

Adenomas ≥10 mm were found in 122 patients: 84 of these 122 patients had adenomas ≥10 mm without villous contingent or high grade dysplasia, nine patients in the IBS group, 16 in the FH group, and 59 in the HD group. Adenomas with a villous component were found in 57 patients (13, 5, and 39 patients in the IBS, FH, and HD groups, respectively). Adenomas with high grade dysplasia were found in 28 patients (three, two, and 23 patients in the IBS, FH, and HD groups, respectively). High risk adenoma (≥10 mm or with a villous contingent or with severe dysplasia) were found in 172 patients (25, 23, and 124 patients in the IBS, FH, and HD groups, respectively).

Colorectal cancer was found in 12 patients (three, none, and nine patients in the IBS, FH, and HD groups, respectively). Most (68.64%) of the high risk colorectal adenomas were located in the distal colon (left colon, sigmoid, and rectum). A total of 90% of patients had a single high risk colorectal adenoma while 3% had three or more high risk adenomas.
Results of analysis

In the first case control study, we examined the relative frequency in the three risk groups (IBS, FH, HD) of patients with adenomas ≥10 mm but without high grade dysplasia or a villous contingent (cases) and patients without adenomas ≥10 mm and without high grade dysplasia or a villous contingent (controls). The relative frequency was significantly different (p<0.01) between cases (n=84) and patients with adenomas <10 mm (controls, n=489). Heavy drinking was more frequent (70%) in cases than in controls (57%) (table 2).

After age and sex adjustment, the likelihood of having an adenoma ≥10 mm was higher in group HD than in group IBS (OR 1.6 (95% CI 0.97–2.6) (p=0.081), 90% CI (1.02–2.3)).

Table 2 Group distribution between cases and controls for adenomas >10 mm

<table>
<thead>
<tr>
<th>Cases</th>
<th>Controls</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenomas &gt;10 mm (n=84)</td>
<td>Adenomas &lt;10 mm (n=489)</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Age [y] (mean [SD])</td>
<td>59 (1)</td>
<td>57 (0.5)</td>
</tr>
<tr>
<td>Male/female ratio</td>
<td>61/23 (73%)</td>
<td>343/146 (70%)</td>
</tr>
<tr>
<td>Patient groups</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IBS</td>
<td>9 (11%)</td>
<td>127 (26%)</td>
</tr>
<tr>
<td>FH</td>
<td>16 (19%)</td>
<td>85 (17%)</td>
</tr>
<tr>
<td>HD</td>
<td>59 (70%)</td>
<td>277 (57%)</td>
</tr>
</tbody>
</table>

The p value is given for the comparison by global χ² method of the relative frequency in the three groups of patients (irritable bowel syndrome [IBS], family history [FH], and heavy drinkers [HD] groups) of cases and controls. The number of patients (cases plus controls=571) is less than the total number of patients (601) because cases with adenomas >10 mm and high grade dysplasia or a villous contingent were excluded from this part of the study.

Table 3 Group distribution between cases and controls for high risk adenomas or colorectal cancer

<table>
<thead>
<tr>
<th>Cases</th>
<th>Controls</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenomas ≥10 mm</td>
<td>Adenomas &lt;10 mm</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Age [y] (mean [SD])</td>
<td>59 (0.9)</td>
<td>57 (0.5)</td>
</tr>
<tr>
<td>Male/female ratio</td>
<td>126/46 (73%)</td>
<td>343/146 (70%)</td>
</tr>
<tr>
<td>Patient groups</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IBS</td>
<td>25 (15%)</td>
<td>127 (26%)</td>
</tr>
<tr>
<td>FH</td>
<td>23 (13%)</td>
<td>85 (17%)</td>
</tr>
<tr>
<td>HD</td>
<td>124 (72%)</td>
<td>277 (57%)</td>
</tr>
</tbody>
</table>

The p value is given for the comparison by global χ² method of the relative frequency in the three groups of patients (irritable bowel syndrome [IBS], family history [FH], and heavy drinkers [HD] groups) of cases (patients with at least one high risk adenoma or a colorectal cancer) and controls (patients with at least one colorectal adenoma but without high risk adenomas).

Table 4 Group distribution between cases and controls for adenomas with high grade dysplasia or colorectal cancer

<table>
<thead>
<tr>
<th>Cases</th>
<th>Controls</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenomas with high grade dysplasia or cancer</td>
<td>Adenomas without high grade dysplasia or cancer</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>Age [y] (mean [SD])</td>
<td>61 (2)</td>
<td>57 (0.5)</td>
</tr>
<tr>
<td>Male/female ratio</td>
<td>30/7 (81%)</td>
<td>439/185 (70%)</td>
</tr>
<tr>
<td>Patient groups</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IBS</td>
<td>5 (14%)</td>
<td>147 (24%)</td>
</tr>
<tr>
<td>FH</td>
<td>2 (5%)</td>
<td>106 (17%)</td>
</tr>
<tr>
<td>HD</td>
<td>30 (81%)</td>
<td>371 (59%)</td>
</tr>
</tbody>
</table>

The p value is given for the comparison by global χ² method of the relative frequency in the three groups of patients (irritable bowel syndrome [IBS], family history [FH], and heavy drinkers [HD] groups) of cases (patients with at least one adenoma with high grade dysplasia or a cancer) and controls (patients with at least one colorectal adenoma but without high grade dysplasia or cancer).

Influence of cirrhosis

In the HD group, we evaluated the influence of cirrhosis in the three case control studies and found that cirrhotic patients were more numerous between cases and controls in any of the three studies performed (table 3).

DISCUSSION

We have shown a relationship between alcoholism and the risk of high risk adenomas or colorectal cancer. To our knowledge, this is the first study that has investigated high risk adenomas or cancer among alcoholic patients with endoscopically proved adenomas compared with two other populations, one with a risk higher than in the general population (patients with a family history of colorectal tumours) and one with a risk thought to be at least equal to the general population (those suffering from IBS). The association between colorectal cancer and alcohol consumption is controversial. A meta analysis of the relation between alcohol and colorectal cancer found only a small but significant effect.

A recent case control study in Argentina found an OR for colorectal cancer of 2.8 (1.6–5.1) for drinkers in comparison with alcohol abstainers, with a significant trend in risk with dose. The association was observed for wine, as well as for beer and spirits.

The association between alcohol consumption and large adenomas has already been proved by Boutron and colleagues. They concluded that alcohol may be involved in the second step of the adenoma-carcinoma sequence by...
promotion of growth of the adenoma, and that alcohol acts as an early promoter of colorectal carcinogenesis but plays no role in malignant changes.

The mechanism by which alcohol might promote colorectal cancer is still controversial. Alcohol in itself would seem to have no direct carcinogenic effect. Its metabolite acetaldehyde has been proved to be cytotoxic. A direct action of acetaldehyde on the faecal microflora has been suggested. Seitz et al have demonstrated in an animal model that the intestinal flora may be involved in acetaldehyde production as germ free rats had significantly lower acetaldehyde concentrations in the rectum (84 (11) v 234 (33) nmol/g/colon; p<0.01) and in the caecum (59 (13) v 121 (33) nmol/g/colon; p<0.05) compared with conventional animals, and that alcohol feeding induced rectal but not caecal hyperregeneration. Acetaldehyde can react with various intracellular and extracellular proteins to form both stable and unstable condensation products. Several proteins involved in these reactions have been shown to form cross links with acetaldehyde, including albumin, tubulin, haemoglobin, collagen, and cytochrome P-450IIE1. Acetaldehyde-protein adduct formation may lead to cellular damage or dysfunction through alteration of biological protein function or adduct triggered immunological effects. Holstege et al have hypothesised that only on their extracellular location, acetaldehyde modified epitopes could serve as neoantigens and thereby elicit humoral or cellular immune responses. Ki-ras mutations are thought to be early events in the carcinogenic process leading to colon tumours. It has been reported that alcohol may be involved in the disease pathway of colon tumours represented by specific Ki-ras mutations.

Ethanol may increase the risk of adenomatous polyps by interfering with dietary habits, and the relation between ethanol and adenomatous polyps could be the result of confounding by unspecified dietary factors such as folate or vitamin C as alcoholic patients often have a low fruit and vegetable diet. Folate, which is plentiful in vegetables and fruits, vitamin C as alcoholic patients often have a low fruit and vegetable diet. Folate, which is plentiful in vegetables and fruits, may be protective against colorectal cancer. A recent nested case control study by Kato and colleagues found that the risk of colorectal cancer in patients within the highest quartile for serum folate was half that of those in the lowest quartile. They also found that the risk of colorectal cancer was almost twice as high in subjects with below median serum folate and above median total alcohol intake compared with those with above median serum folate and below median alcohol consumption. This may suggest that alcohol intake and folate deficiency have a synergistic action in the promotion of colorectal carcinoma, particularly in alcoholic patients who usually have a low folate and vitamin C intake and a high fat intake. This has recently been confirmed in a large epidemiological study. Multivitamin use and folate intake have been reported to have a protective effect in colon cancer in women only after long use. We might hypothesise that the risk of colorectal high risk adenomas would be increased only after many years of a low folate diet. Our alcoholic patients were drinking more than 50 g/day for a mean duration of 22 (0.6) years.

We did not specifically study the role of tobacco but this is another risk factor in these results, as we previously found that alcoholic patients are more likely to be current smokers than patients suffering from IBS. This is in agreement with Faivre and colleagues who found that the proportion of large adenomas in men which could be attributed to combined alcohol and tobacco consumption was 78%.

We did not study the role of cholesterol but it needs to be discussed as we have previously observed that serum cholesterol concentration was lower in patients with adenomas than in patients without adenomas. In the same study, we showed that alcoholism was independently associated with the risk of having colorectal adenomas.

The ORs after adjustment for age and sex were in agreement with the findings that alcoholism may be involved in the severity of colonic adenomas as the relative risk of heavy drinkers having a large adenoma, a high risk adenoma or cancer, or a high grade dysplasia or cancer was significantly increased compared with IBS patients (ORs 1.8, 1.6, and 1.7, respectively). Comparison between the HD and FH groups showed that there was a trend towards an increased risk for high risk adenomas or cancer and for high grade dysplasia or cancer in group HD compared with group FH, but this was not statistically significant at the 5% level.

In conclusion, in our study alcoholism was a risk factor for the development of high risk adenomas or colorectal cancer. This suggest that alcoholic patients may benefit from a screening policy of colonic adenomatous polyps when their liver function allows.

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**Table 5** Relation between cirrhosis and: adenomas >10 mm; high risk adenomas or cancer; and high grade dysplasia or cancer among alcoholic patients

| Circumstance | Adenomas >10 mm (n=59) | Adenomas <10 mm (n=276) | p Value | NS
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Cirrhosis</td>
<td>20 (34%)</td>
<td>105 (38%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cirrhosis</td>
<td>High risk adenoma or cancer (n=124)</td>
<td>No high risk adenoma or cancer (n=277)</td>
<td>106 (38%)</td>
<td>NS</td>
</tr>
<tr>
<td>Cirrhosis</td>
<td>High grade dysplasia or cancer (n=30)</td>
<td>No high grade dysplasia or cancer (n=371)</td>
<td>143 (39%)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Per cent values represent the number of patients with proven cirrhosis (cirrhosis was assessed either on a liver biopsy or using clinical and biological signs) among the total number of patients in each subgroup of the heavy drinkers.

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