Suppressive effect of aspirin on aberrant crypt foci in patients with colorectal cancer

B Shpitz, E Klein, G Buklan, D Neufeld, A Nissan, H R Freund, M Grankin, J Bernheim

Background and aims: Aspirin and other non-steroidal anti-inflammatory drugs have been shown to reduce the risk of colorectal cancer (CRC). Animal models have shown that aspirin is also effective in reducing the density of aberrant crypt foci (ACF). The aim of the study was to evaluate the effect of chronic administration of aspirin on the distribution pattern and histological characteristics of ACF in patients with CRC.

Methods: Our study compared the distribution patterns and histomorphological characteristics of ACF between a group of CRC patients treated with low dose aspirin (n = 59) and a control group without aspirin (n = 135). ACF were visualised on methylene blue stained macroscopically normal mucosa, micro-dissected, and serially cut.

Results: ACF were found in 75.8% of mucosal samples from the control group and in 36% of mucosal samples from the aspirin treated group, indicating a 47% decline in prevalence of ACF in colonic samples of patients treated with aspirin. A significant reduction from 92.5% to 40% (p < 0.0001) was found in distal large bowel samples containing one or more ACF. Similarly, the aspirin treated group showed a reduction in ACF density of 64% and 82%, respectively, in both proximal and distal parts of the colon, indicating a significant reduction in ACF/cm² in distal colonic samples (p < 0.01). The aspirin treated group displayed a 52% reduction in dysplastic ACF although this difference was not statistically significant.

Conclusions: Our study has provided evidence of the effective chemopreventive action of low dose aspirin on ACF in humans.

Materials and methods

Study group

Normal appearing colorectal mucosal samples were collected from surgical resection specimens obtained from 194 patients who underwent large bowel resection for sporadic CRC. Patients with familial syndromes of CRC (familial adenomatoid polyposis and hereditary non-polyposis colon cancer), patients on NSAID treatment other than aspirin, as well as those who received preoperative radiation and/or chemotherapy were excluded from the study. Information regarding administration of drugs other than aspirin or NSAIDs and history of accompanying diseases were available in 50% of patients. Therefore, these details were not included in the data analysis. Dietary habits of the patients were not available. Mucosal samples were collected from surgical resection specimens obtained from 194 patients who underwent large bowel resection for sporadic CRC.

Tissue collection, sampling, and processing

For identification and collection of ACF, mucosal samples were spread flat and fixed in 4% buffered formalin.

Abbreviations: ACF, aberrant crypt foci; CRC, colorectal cancer; NSAIDs, non-steroidal anti-inflammatory drugs; PPAR, peroxisome proliferator activated receptor.
formalin. ACF were screened under low power magnification following staining in 1% methylene blue. The identified ACF were counted, microdissected, embedded in paraffin, sectioned, and stained with haematoxylin-eosin. According to the histopathological pattern, ACF were classified as non-hyperplastic, hyperplastic, and dysplastic (fig 1). A total of 291 ACF were microdissected.

Data analysis included comparison of the prevalence, distribution density, multiplicity (number of foci per ACF), and histopathological characteristics of ACF in patients with and without aspirin treatment. The distribution pattern of ACF in both groups was compared separately for the proximal and distal colon. The anatomical boundary between the distal and proximal parts of the large bowel was the splenic flexure.

Table 1 Duration of aspirin administration in the treatment group

<table>
<thead>
<tr>
<th>Time on aspirin (months)</th>
<th>No of patients (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>12–60</td>
<td>35 (59.3)</td>
</tr>
<tr>
<td>61–120</td>
<td>21 (35.6)</td>
</tr>
<tr>
<td>&gt;120</td>
<td>3 (5.1)</td>
</tr>
</tbody>
</table>

Table 2 Density of aberrant crypt foci (ACF) in different age groups in the control group (expressed as ACF/cm²)

<table>
<thead>
<tr>
<th>Site of colon</th>
<th>Age group (y)</th>
<th>&lt; 50</th>
<th>51–60</th>
<th>61–70</th>
<th>71–80</th>
<th>&gt;80</th>
</tr>
</thead>
<tbody>
<tr>
<td>Right</td>
<td></td>
<td>0.04</td>
<td>0.04</td>
<td>0.01</td>
<td>0.02</td>
<td>0</td>
</tr>
<tr>
<td>Left</td>
<td></td>
<td>0.12</td>
<td>0.09</td>
<td>0.12</td>
<td>0.1</td>
<td>0.16</td>
</tr>
</tbody>
</table>

Statistical analysis

Comparison between groups was performed with the Mann-Whitney U test and the χ² test, as appropriate, using the SPSS statistical package (SPSS Inc, Chicago, Illinois, USA). All values are expressed as mean (SD), unless indicated otherwise.

RESULTS

A total of 206 mucosal samples were obtained from 194 patients. More than one specimen was obtained from some patients who underwent subtotal large bowel resection. Fifty nine patients had been receiving aspirin treatment for at least one year. All patients in the study group were treated with 100 mg/day aspirin except for three who were treated with buffered aspirin 325 mg/day. Average time on aspirin was 48 months (range 12–108 months). Data on duration of aspirin treatment are shown in table 1.

There were 65 (48%) women and 70 (52%) men in the control group and nine (16%) women and 50 (84%) men in the study (aspirin) group. Mean ages of the control and study groups were 72.6 (range 35–86) years and 69.6 (range 62–82) years, respectively.

Baseline ACF density in the control group was calculated separately for the right and left colons. No significant difference in ACF density was found between the different age groups (table 2). ACF were found in 75.8% of mucosal samples from the control group and in 36% of those from aspirin treated patients, indicating a 47% decline in prevalence of ACF in “treated” colonic samples. There was a reduction in samples that contained at least one ACF from 46% to 32% in the proximal (p = 0.9) and from 92.5% to 40% (p<0.0001) in the distal large bowel (table 3).

The distribution density of ACF in the control and aspirin treated groups is shown in table 4. The aspirin treated group showed a reduction of 64% and 82% in ACF density in both proximal and distal parts of the colon, respectively, indicating a significant reduction in ACF/cm² in the distal but not in the proximal colon samples (p = 0.9 for the proximal colon; p<0.01 for the distal colon) (table 4).

The effect of aspirin on the histological characteristics of microdissected ACF is shown in table 5. Aspirin treatment resulted in a redistribution pattern of dysplastic ACF: the aspirin treated group displayed a 52% reduction in dysplastic ACF compared with the control group, although this difference was not statistically significant (table 5).
Our findings showed that low dose aspirin treatment, administered for at least one year before surgery, resulted in reduction of overall density of ACF. We further demonstrated that significantly fewer mucosal samples in the aspirin treated group contained ACF. This is the first study in humans to demonstrate a chemopreventive effect of low dose aspirin at the earliest stages of colon tumorigenesis.

Aspirin plays a major role in primary and secondary prevention of cardiovascular events such as myocardial infarction and cerebrovascular accidents. It is therefore not surprising that a substantial percentage of elderly patients with CRC take aspirin as a chemopreventive agent for cardiovascular events. This fact has enabled us to collect prospective data on a group of patients who had been receiving chronic aspirin treatment and to evaluate the effect of aspirin on the distribution and histological characteristics of ACF by comparing them with a group of patients who were not taking aspirin. The design of our study warrants special comment. Patients in our study group had already had CRCs, giving rise to concerns that many of the patients might be resistant to the chemopreventive action of aspirin. This could be relevant, at least for those patients who had been taking aspirin for a long period of time (longer than a decade). In fact, the median time on aspirin in our study group was relatively short (48 months) and only a small proportion of our patients (5%) had been taking aspirin for more than 10 years. Although Greenberg and colleagues demonstrated a relatively short (48 months) and only a small proportion of our patients (5%) had been taking aspirin for more than 10 years. Although Greenberg and colleagues demonstrated a lower risk for adenomas even after one year on aspirin treatment, another study found that a substantial reduction in CRC risk could be demonstrated only after at least 10 years of aspirin use. Interestingly, our results indicate that even among patients with CRC, aspirin treatment was capable of exerting suppressive action at the earliest stages of colorectal tumorigenesis.

Two factors that could have influenced the distribution patterns, prevalence, and histological subtypes of ACF in our study were diet and medications. Different dietary components have been shown to affect the prevalence of ACF in carcinogen induced colon tumorigenesis models in rodents. High fat diets containing mixed lipids increased the total number of ACF while high energy and fat restricting diets exerted a suppressive effect on advanced ACF. Similarly, dietary supplementation of wheat germ extract and fermented brown rice effectively inhibited ACF development. However, as no studies have been conducted in humans to evaluate the possible effects of various dietary components on ACF, the potential effect of diet is largely unknown. Data on dietary restrictions were not available in our patients with cardiovascular diseases; therefore, although theoretically a low fat diet recommended for patients with cardiovascular diseases could influence the distribution patterns of ACF, the real influence of these components on the prevalence of ACF in our study remains unknown. Similarly, no studies in humans regarding the effect on ACF of cardiovascular or other medications, except NSAIDs, have been reported in the literature.

Aspirin and other NSAIDs were shown to be effective chemopreventive agents during the initiation and post-initiation stages of colon carcinogenesis in rodents. Low dose aspirin was effective in suppressing ACF formation but was unable to prevent cancer. No specific data from these studies have demonstrated how aspirin specifically affected dysplastic ACF. To the best of our knowledge no studies have been published on the effect of aspirin on early morphological biomarkers of colonic carcinogenesis in humans. A recent study using magnification colonoscopy evaluated the effect of sulindac on ACF and showed that this agent was indeed effective in reducing the density and prevalence of ACF in the rectum, one year after initiation of treatment. Although this study demonstrated complete disappearance of ACF in seven of 11 treated patients and reduction of ACF density in the rest, no specific data on dysplastic foci were provided. As only one patient in this group actually had colon cancer, it was probably unreliable to search specifically for the effect of sulindac on dysplastic lesions.

Even though we were able to demonstrate a general ACF suppressive effect, the more critical issue in evaluation of chemopreventive action of low dose aspirin would be the potential effect specifically on dysplastic ACF which are truly...
neoplastic microscopic lesions and have been recognised as the most important lesions in the ACF-adenoma-carcinoma sequence. While we were able to demonstrate a general ACF suppressing effect, our data showed that aspirin treated patients displayed a reduced proportion of dysplastic ACF (from 13.3% in the control group to 6.3% in treated patients), the results were not statistically significant. Surprisingly, there are no data addressing this important issue.

The vast majority of our patients in the aspirin group had been on low dose aspirin. The minimal effective aspirin dose in early colonic tumorigenesis is unknown. Ruffin et al suggested that a single 81 mg dose of aspirin taken daily should be sufficient to significantly reduce colorectal mucosal prostaglandins E2 and F2α. Based on these data, the above dose was recommended for future chemopreventive studies in CRC. Although an aspirin dose close to that reported in the latter study could explain the effect of aspirin on ACF observed in our study, it is not yet known whether this dose would be sufficient to suppress colorectal carcinogenesis.

Our present study showed that a greater reduction in ACF density was noted in the distal compared with the proximal colon. In fact, the only statistically significant reduction in the percentage of samples containing ACF as well as a reduction in ACF density was found in the distal large bowel. This is in accordance with findings in a previous study in a rodent model11 that demonstrated a distal predilection of nabumetone suppressive action on ACF. The cause of this differential suppression of ACF in different parts of the large bowel is unclear although higher levels of cyclooxygenase 2 expression as well as stronger induction of peroxisome proliferator activated receptor (PPAR) in distal parts of premalignant rat colons have been found.12 It is noteworthy that both PPAR and cyclooxygenase 2 are molecular targets of NSAIDs.13-14

In conclusion, this study has provided the first evidence of a chemopreventive effect of low dose aspirin on ACF in humans. Further studies should be carried out to evaluate the potential mechanisms of the differential effect of aspirin and other NSAIDs on ACF in humans as well as their effect on dysplastic ACF.

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