Effect of long-term placebo controlled calcium supplementation on sigmoidal cell proliferation in patients with sporadic adenomatous polyps

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Abstract
A long-term, double blind intervention trial was undertaken in patients with sporadic adenoma treated by polypectomy to investigate the putative role of calcium as a protective factor in colon carcinogenesis. The aim of the study was to assess the effect of a daily dietary supplementation of 2 g calcium over nine months on cell proliferation measured as proliferation index in colonic mucosa. A total of 48 patients were entered into the study of which 30 were fully compliant. After intervention proliferation index (mean (SEM)) in colonic epithelium was decreased in both the calcium (13.5 (1.5) to 11.4 (1.2)) and the placebo group (13.7 (0.9) to 10.8 (1.1)). The difference in the change between the two groups was not significant (p<0.01). Changes in proliferation index % of crypt compartments were also not significantly different between the two groups. A significantly positive correlation between soluble calcium in faeces and the total proliferation index % in colonic epithelium at baseline and after intervention (r=0.54, p<0.01, r=0.50, p<0.01 respectively) suggests that an increase of free luminal calcium alone is insufficient for inhibition of cellular proliferation.

Methods
Study design
Participants were recruited over a period of nine months from patients attending endoscopic units in eight hospitals in Heidelberg and surrounding areas with one physician being responsible for recruiting patients and biopsying in each hospital. Patients with newly diagnosed colorectal adenomas who were 30–75 years of age were eligible for the study. Additional inclusion criteria were a histologically confirmed adenomatous polyp larger than 4 mm, informed consent about participation in the study, and biopsy for determination of cell proliferation. Biopsy specimens were obtained at the initial colonoscopy from macroscopically normal mucosa at the rectosigmoidal junction (about 20 cm from the anal verge) but at least 5 cm from any neoplastic lesion, using standard forceps. Number, size, and localisation of polyps and whether or not they were removed, were recorded.

Exclusion criteria included conditions predisposing to colorectal cancer other than sporadic adenomas, contraindications to calcium supplementation or to biopsy sample excision, conditions indicating severe restriction of general health, or drug use that may interfere with the study. In particular, exclusion criteria were as follows: FAP; inflammatory bowel disease; chronic alcohol abuse; intestinal malabsorption syndromes; cancer other than non-melanoma skin cancer; renal insufficiency; gastrectomy, ileostomy, colectomy; vegetarian diet; consumption of more than 1 litre milk per day or more than 600 g dairy products per day (milk-alkali-syndrome); renal stones, haematuria; sarcoidosis; hyperparathyroidism; hypercalcaemia; abnormal serum calcium or creatinine concentrations at the time of initial

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colonooscopy; cholestyramine treatment; calcium or vitamin D supplementation; anticoagulant treatment, problems in understanding the nature of the study; and place of residence more than 100 km from the study centre.

Eligible patients were followed up immediately after the initial polypectomy and subsequently every three months at their private residences. Before entering the actual treatment phase, participants were asked to collect 48 hour stool and 24 hour urine samples. The usual dietary intake was assessed with a self administered food frequency questionnaire. Study participants were asked to continue their normal Western style diet. Before randomisation, a run in phase of three months was applied in which all participants received placebo. Less than 80% compliance to the treatment protocol during the run in phase resulted in exclusion from randomisation. Compliant participants were then randomly allocated to either calcium or placebo treatment for another nine months. At every follow up visit, two 24 hour recalls were conducted to monitor short-term changes in dietary habits. Treatment compliance was assessed by pill count. Drug intake besides calcium supplementation was assessed at entry into the study and every three months during follow up. Faecal (48 hour) and 24 hour urine samples were again collected after nine months. After a total of 12 months - that is, three months run in phase and nine months calcium supplementation - a control colonoscopy was performed during which biopsy samples were again obtained at 20 cm proximal to the anal verge for final examination. The study was approved by the ethical committee of the University of Heidelberg.

Medication and randomisation
Calcium was applied in the form of effervescent tablets as calcium-glucono-lactate, with each tablet containing 1000 mg calcium. Participants were requested to take one with their breakfast and one with their evening meal, resulting in a daily supplementation dose of 2000 mg calcium. Placebo tablets consisted mainly of citrate, fructose, and sucrose. Both calcium and placebo medication were kindly provided by Sandoz AG, Nürnberg, Germany.

Randomisation was conducted in advance. Each box of medication covered three months intervention and was labelled by randomly selected numbers. Thus, the study coordinator and interviewer were blinded throughout the study. Participants allocated to calcium treatment did not detect any difference in medication appearance when calcium supplements were introduced after the run in phase.

Collection and preparation of faecal and urine samples
Faecal samples were defecated into plastic bowls and placed immediately on dry ice (−79°C). The complete 48 hour sample was stored at −80°C until further analysis. For aliquoting, samples were thawed overnight at 4°C, pooled, and homogenised at room temperature in a rotating homogeniser (Heidolph, Kelheim, Germany) at 600 rpm for five minutes. Aliquots of 40 g were freeze dried, reweighed, and ground to a fine, homogeneous powder. Urine was collected over 24 hours, aliquoted, and stored at −20°C until further analysis.

Calcium analysis
To 100 mg of freeze dried faeces 2 ml of concentrated HNO₃ was added in a tightly sealed universal bottle. The bottles were placed in a heating block at 120°C for one hour. On cooling 10 ml 1 M HCl was added prior to analysis of total calcium by atomic absorption spectrophotometry (AAS). Soluble calcium was measured by adding 10 ml of distilled water to 100 mg of freeze dried faeces and incubating overnight at room temperature. Supernatants were again analysed by AAS.

Dietary analysis
During the basic examination a food frequency questionnaire for self administration was distributed to each participant. This questionnaire asked for the frequency and portion size of 121 food items. The resulting quantity in food consumption was converted into nutrient intake by using a German food composition table (Federal Food Code of Germany). This Food Code has been developed by the Federal Health Office and encompasses a coding system of more than 10 000 different food codes with the nutrient data for each food code. This version of the food frequency questionnaire was not formally validated. However, subsequent versions showed similar characteristics in terms of validity and reliability compared with other published food frequency questionnaires (H Boeing, unpublished data).

During supplementation the 24 hour recall method was used. At each follow up visit participants were asked about their food intake on the previous day. For each single food item consumed, the amount was estimated by use of household measures or portion size photos. A cross check at the end of each 24 hour recall was applied to search for previously omitted items. The detailed responses were coded according to the Federal Food Code and subsequently converted into nutrient intake.

Determination of proliferation index
For the first five patients who entered the study, cell proliferation was measured by the BrdU method. To facilitate recruitment of patients from external hospitals, however, a proliferation marker based on an endogenous cell cycle related antigen was required to circumvent immediate incubation of viable tissue. Therefore, PCNA (clone PC10) immunohistochemistry was applied for the subsequent study participants. In any one patient, the same method was used before and after intervention. After clinical examination and, as for BrdU incubation of biopsy samples,
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the specimens were fixed in absolute methanol, transported to the laboratory, and processed within 24 hours. Both BrdU and PCNA protocols have been described and compared elsewhere.20

All sections were examined by one observer who had no knowledge of the clinical features, the histological diagnosis or the random allocation to treatment groups. In a previously conducted analysis of variance,20 the number of crypts and number of biopsy samples that must be evaluated to classify a subject within a certain interval was estimated. According to these findings, five crypts of each biopsy sample were evaluated whenever three samples were used for the evaluation, eight crypts of each of two biopsy samples, and 30 crypts when only one biopsy sample was available to characterise a subject at each point in time.

A crypt was analysed if its entire length was visible in the longitudinal section and the base contacted the muscularis mucosae. In each crypt column (which is the single column of epithelial cells lining one side of a sectioned crypt meeting the criteria) the total number of cells and the number of labelled cells and their position in the crypt column relative to the base of the crypt were noted. Total number of cells, number of labelled cells, and PIs, which is the ratio of labelled cells to the total number of cells, were determined for each crypt column. PI% were also calculated separately for each of five compartments of equal size into which each crypt column was divided. The same procedure was repeated for percentiles of the crypt column of standardised length, and a running mean over five percentiles was calculated to smooth the distribution.

Sample size considerations
For sample size calculations the two sided t test for unpaired samples was used to test for significance of the difference in the mean change of PI% from the baseline value after intervention between the two treatment groups. The standard deviation (SD) of the major end point ‘change in PI% after intervention as compared to baseline’ was estimated to be around 2, based on data by Gregoire et al.,15 who gave a standard deviation of the mean change after intervention in the placebo group of 1.7 and in the calcium group of 2.3. With α=0.05 and β=0.1, the number of patients required to detect a difference of 2 in the change of PI% between two groups was calculated to be 21 patients in each group.21 To account for drop outs during the study, a total of 50 patients was planned to be recruited for the study.

Statistical analysis
Baseline correlation analysis was conducted treating age, number of adenomas, size of the largest adenoma, and faecal calcium concentration as continuous variables and grade of dysplasia (tubular/tubulo-villous/villous) as a categorical variable.

For comparison between the two groups, dietary data were adjusted for sex. For each nutrient (j), a regression model was calculated with sex (x) as dependent variable (coded as 1 and 2): yij=αj+βjx. Intercept (αj) and slope (βj) parameters were used to adjust each patient’s intake values (i) for each nutrient, thereby simulating a group of balanced sex:

\[ y_{ij} = \beta_j + (y_{ij} - \frac{\beta_j}{\alpha_j} \alpha_j) \]

Subjects were regarded as compliant and included in the final evaluation of the effect of calcium intervention if the pill count yielded an 80% adherence to the treatment and the increase in faecal calcium excretion was greater than 1500 mg.

Mean PI% of colonic epithelial cell proliferation was determined for each group at each time point. The treatment effect was estimated and tested with Wilcoxon’s rank sum test on the difference between the mean changes of PI% in each group after intervention compared with baseline. Average scores were used for ties. Results were considered statistically significant at α=0.05. Multiple regression analysis was used to adjust for confounding variables. The PI% as a function of cell position relative to crypt column base was examined separately for each group at each time point.

Results
Of 48 patients who were entered into the calcium intervention study, four were not compliant during run in phase, one died, six were not compliant during the main intervention phase and four either refused or were not able to undergo their final examination – that is, biopsy samples could not be obtained. Each of the remaining 33 participants showed a compliance greater than 80% at each pill count evaluation. In three of these participants, however, calcium excretion in faeces indicated low compliance. They were therefore excluded from the final analysis. The final statistical analysis is based on 30 participants (15 in each group) who completed the study in a full compliant manner (>96% compliance). Table 1 gives characteristics of the study group with respect to age, sex, and usual dietary intake. The average age of the final study group was 62 (range 46–75), 14 women and 16 men. Mean age of both groups was similar, but distribution of sex

| TABLE 1 Age, sex, and dietary data prior to intervention in calcium and placebo group (mean (SD)) |
|-------------------------------------------------|-------------------------------------------------|
| Placebo group (n=15)                               | Calcium group (n=15)                             |
| Age                                             | Age                                             |
| 63-3 (6-1)                                       | 60-9 (9-2)                                      |
| No of female patients                           | No of female patients                           |
| 10                                              | 4                                               |
| Dietary intake*                                 | Dietary intake*                                 |
| Energy (kJ/d)                                    | Energy (kJ/d)                                    |
| 10410 (2169)                                    | 11678 (5788)                                    |
| Protein (g/d)                                    | Protein (g/d)                                    |
| 93 (37)                                         | 99 (50)                                         |
| Fat (g/d)                                       | Fat (g/d)                                       |
| 97 (38)                                         | 105 (51)                                        |
| Carbohydrates (g/d)                             | Carbohydrates (g/d)                             |
| 253 (52)                                        | 293 (160)                                       |
| Fibre (g/d)                                     | Fibre (g/d)                                     |
| 38 (7)                                          | 35 (20)                                         |
| Calcium (mg/d)                                  | Calcium (mg/d)                                  |
| 933 (416)                                       | 922 (370)                                       |
| Vitamin D (µg/d)                                | Vitamin D (µg/d)                                |
| 4 (4)                                           | 4 (3)                                           |

*As determined by food frequency questionnaire before intervention, adjusted for sex.
was unbalanced (Table I). For comparison, dietary data were therefore adjusted for sex. In the calcium group usual energy, protein, fat, carbohydrate, and fibre intake were slightly higher than in the placebo group, whereas the calcium intake was lower. However, a wide variation of mean values rendered these differences insignificant. Dietary habits during the study ascertained by repeated 24 hour recalls were stable in both groups (detailed data not shown). Drugs or dietary supplements that could potentially affect cell proliferation were only taken in the calcium group (two participants with vitamin E and C supplements and one with β-sitosterol supplement for at least three months; anti-inflammatory drugs were never taken by any participant).

Multivariate analysis showed age of patients and size of largest adenoma at initial examination to be significantly positively associated with initial PI% (r=0.49, p<0.01; r=0.51, p<0.01 respectively). No association could be found between cell proliferation and number of adenoma, grade of dysplasia in removed adenoma or distance of rectosigmoidal junction to the nearest adenoma. The method of measuring cell proliferation was included in each multivariate model, because BrdU generally yielded lower PI% than PCNA. At baseline examination as well as after intervention, a significant positive correlation was seen between concentration of soluble calcium in faeces and PI% in sigmoidal mucosa (r=0.54, p<0.01, r=0.50, p<0.01). Subdivided into crypt compartments, this correlation between soluble calcium and PI% was only significant for compartment 1 and 2.

The power of the study was diminished for two reasons. Firstly, only 15 patients completed the study according to protocol in each group rather than the intended 21. Secondly, standard deviation of the major end point 'change in PI% after intervention as compared to baseline' was larger than anticipated. The minimum detectable difference of PI% between both groups in this study with n=15 in each group and SD=4.75 was 4.8. Given an expected no effect in the placebo group, a decrease of PI% in the calcium group to approximately 7% of that found in healthy controls could have been detected as statistically significant compared with the placebo group.

After nine months of intervention a decrease of total PI% in sigmoidal mucosa was evident in both calcium (from 13.5 before to 11.4 after intervention) and placebo group (from 13.7 to 10.7) (Table II). The difference between the two groups regarding the change of PI% after intervention compared with baseline, was not statistically significant (p=0.7). The observed decrease in the calcium group could therefore not be ascribed to calcium intervention. Age, sex, and adenoma size did not influence this result when included as covariates in a regression analysis. The method of measuring cell proliferation had no influence on the change in PI% after intervention – that is, did not interfere with the main result. Five of 15 patients in the calcium group but only three of 15 in the placebo group experienced an increase in PI% (Fig 1). There was a tendency in the calcium group for initially high values to decrease and initially low values to increase.

Proliferative activity in crypt compartments was highest in compartment 2 in both groups.
of patients at both time points (Table II). PI% decreased in both groups during intervention in compartments 2, 3, and 4, whereas for the basal compartment 1, a decrease was only evident in the placebo group (Table II). This indicates a slight downward shift of the proliferative zone in the crypt due to calcium supplementation. However, the differences between the calcium and placebo groups regarding the change in each compartment were not statistically significant. In compartment 5, PI% remained unchanged after intervention. Also, total number of cells per crypt column was not affected by intervention.

The two groups were also compared at the two different time points regarding the continuous distribution of labelled cells along the crypt (Fig 2). Before intervention, a steep rise in proportion of labelled cells occurred to a maximum at the 20th percentile in the calcium group and the 18th percentile in the placebo group. After intervention the peak PI reverted to the 18th percentile of crypt height in the calcium group, whereas in the placebo group the maximum PI% could be found at the 20th percentile. In both groups there was a decrease of proliferative activity after intervention compared with baseline almost along the whole length of the crypt, except for the uppermost 20% and the basal 10% of the crypt. However, these differences were not significant at any cell position.

An intention to treat analysis of all randomised subjects was not possible, because five non-compliant and four compliant subjects refused to or were not able to undergo final examination – that is, biopsy samples could not be obtained. Because the study was conducted in symptom free subjects, it is very unlikely that compliance rate is related to the outcome measure. Four subjects with less than ideal compliance (<80%) did have a final examination and biopsy. Including their data in the analysis did not materially effect the results.

Daily calcium excretion in faeces increased significantly both as total calcium and soluble calcium in the calcium group compared with the corresponding change in the placebo group (Table III). The increase in faecal calcium output of 1807 mg in the calcium group corresponds closely to 2000 mg calcium by which the diet was supplemented.

Calcium excretion in urine before intervention was in both groups on average 85 mg/day (Table III), which is well below the obligatory amount of renal calcium loss under conditions of calcium balance (114–116 mg/day), showing a negative calcium balance in these patients. After intervention with 2 g calcium per day for six months renal calcium excretion stayed below this level in the calcium group and the increase was not significantly different from that in the placebo group.

**Discussion**

This study shows that independent of intervention, cell proliferation (measured predominantly by PCNA immunohistochemistry) in colonic mucosa of adenoma patients correlates positively with patient age and size of the largest diagnosed adenoma. This association with known risk factors for colorectal

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**Figure 2: Proliferation index as a function of the percentile distance from the base of the crypt column before and after nine months of calcium intervention separate for calcium and placebo group (curve smoothed by a running mean of five).**

**TABLE III. Effect of six month calcium (2 g/dl) intervention on calcium excretion in faeces and urine in adenoma patients (mg/d) (mean (SEM)).**

<table>
<thead>
<tr>
<th></th>
<th>Placebo group (n=15)</th>
<th>Calcium group (n=14)†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Before intervention</td>
<td>6 Months after intervention</td>
</tr>
<tr>
<td><strong>Faeces</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total calcium</td>
<td>914 (141)</td>
<td>723 (78)</td>
</tr>
<tr>
<td>Water soluble calcium</td>
<td>114 (23)</td>
<td>100 (16)</td>
</tr>
<tr>
<td><strong>Urine</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total calcium</td>
<td>68 (9)</td>
<td>74 (11)</td>
</tr>
</tbody>
</table>

*Wilcoxon rank sum test on the difference between the changes in both groups. †No collection of faeces and urine in one participant.
neoplasia supports the idea of the measurement of cell proliferation to be an adequate biomarker for increased colon cancer risk.

The major intention of the study was to investigate whether or not a longterm calcium supplementation would decrease cell proliferation in the sigmoid mucosa of patients treated by polypectomy. A decrease in cell proliferation was evident in both calcium and placebo groups after nine months of intervention. The power of the study permits the conclusion that a longterm calcium intervention in patients treated by polypectomy is unable to normalise the raised proliferative status in the sigmoid colon of these patients. Both patient recruitment and compliance problems were encountered in this study. Forty eight instead of the intended 50 patients were entered into the study, of which only 30 were fully compliant.

The most plausible explanation for low compliance seems to be that the traumatic experience of a polypectomy has only a short-term effect on most patients’ perception of illness. During the course of a one year intervention, a number of other medical, psychological, and social problems emerge, which allows the memory of a previous diagnosis to pale by significance. The longer the study period the more participants began to consider it less important to continue intervention and drop out. Compliance in short-term studies therefore tends to be higher.

An obvious disadvantage of the study was that two different proliferation markers were used. The method used for an individual patient was shown to influence only the level of PI% at a particular point in time, but not the change in PI% after intervention compared with baseline. The use of two different markers is therefore very unlikely to have distorted the results.

There are several possible reasons as to why a decrease in cell proliferation in both groups was found independent of medication. Participation in a preventive trial may induce participants to change their health related behaviour, such as diet. Based on the dietary information ascertained at onset and during the course of this study, it is evident, that participants in both groups continued their normal diets. Alternatively, our findings may be a result of statistical regression to the mean. However, a prerequisite for regression to the mean to occur, which is a positive correlation between values measured at the two points in time is not met in our data. Another possible explanation for the decrease in cell proliferation in both patient groups is that removal of the neoplastic lesion from the colon may cause general reduction of cellular proliferation along the entire colon. There is some support for this because Risio et al. found a significant decrease in colon proliferation as a function of time of a polyp free colon after two years. This may explain why calcium intervention in patients treated with polypectomy without a control group seems effective.

In most calcium intervention studies cell proliferation was measured in the rectum, and results were extrapolated to the risk of developing a neoplastic lesion in the colon. The authors justify this by reference to studies of Terpstra et al. and Ponz de Leon et al. Both showed increased cell proliferation in high risk patients not only in mucosa adjacent to a neoplastic lesion but in the entire colon and rectum. A more recent study where an in vivo injection of BrdU was carried out in 75 patients with adenocarcinoma, showed no difference in proliferative activity between caecum, transverse colon, descending colon, sigmoid, and rectosigmoidal junction. Comparison of these colonic sites with the rectum showed a significantly lower PI% in the rectal mucosa. The distribution of the label along the crypt was also significantly different. Therefore, it cannot be excluded that the inhibiting effect of calcium on cell proliferation may be different in the rectum from that in the colon.

A postulated mechanism by which calcium may exhibit its protective effect on colonic epithelium is by binding free fatty acids and secondary bile acids in the colonic lumen. Analysis of faecal lipids before and after six months of intervention in the same study group essentially showed evidence of increased precipitation of calcium salts of fatty acids but this did not lead to a decrease in the concentration of free fatty acids in this study (Weisgerber et al., submitted data). This finding is in agreement with no effect of calcium on cell proliferation.

Another postulated mechanism by which calcium may act on colonic epithelium is direct stimulation of cell differentiation. An insight into this can be obtained from the results on calcium excretion in faeces and urine. Renal calcium excretion in these patients was lower than obligatory renal calcium loss under conditions of calcium balance. Renal saving dominating obligatory renal loss is probably caused by a negative calcium balance at entry into the study despite reasonably high calcium intake; the reason for this being a comparatively low calcium absorption. During intervention, almost the entire amount of supplemented calcium was excreted via faeces, again implying a low calcium absorption capacity in this study group.

As shown by the positive correlation between the soluble calcium concentration in faeces and mucosal PI%, it may be necessary for calcium to enter the colonocyte to exhibit its inhibiting effect on crypt cell proliferation. In patients with limited calcium absorption and low calcium uptake into the intestinal cell, high calcium supplementation would not decrease cell proliferation because passive transport through the colonic epithelium is paracellular. Decreasing calcium absorption with increasing age was a well recognised phenomenon and may be one reason why elderly persons exhibit a higher proliferative activity in the intestinal mucosa. Supplementation of 2 g/day calcium in young volunteers (mean age 27 years) does cause urinary calcium to increase significantly from 134 mg to 172 mg on average, indicating increased net calcium absorption.
As calcium absorption in the colon is dependent on 1,25-dihydroxyvitamin D, calcium supplements alone may not be effective in patients with low calcium absorption. Calcium intervention in 79 adenoma patients (mean age 61 years) failed to reduce crypt cell production rate after 12 months compared with placebo.19 The only longer-term trial that showed reduction of epithelial proliferation after six months of intervention with 1500 mg/day was conducted in FAP patients (mean age 38 years), whose blood concentrations of vitamin D metabolites were above the normal range, indicating sufficient vitamin D supply.18 Nevertheless, several short-term studies in elderly high risk patients have shown a protective effect of calcium on cell proliferation in sigmoidal and rectal mucosa.12-14 It is possible that calcium exerts only a transient effect on intestinal proliferation and that in the long term limited calcium absorption hinders a protective effect. It may be useful for on-going intervention studies33 to measure serum concentrations of vitamin D metabolites as a possible discriminating determinant between 'responders' and 'non-responders' to calcium intervention.

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4 Deschner EE, Maskens AP. Significance of the labeling index and labeling distribution as kinetic parameters in colorectal mucosa of cancer patients and DMM treated patients. Cancer Res 1982; 50: 1136-41.