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<tr>
<td>Author(s)</td>
<td>Tan, VPY; Chan, P; Hung, IFN; Pang, R; Wong, BCY</td>
</tr>
<tr>
<td>Citation</td>
<td>The 5th Annual Conference of the Organisation for Oncology and Translational Research, Macau, China, 20-21 February 2009. In Expert Opinion on Investigational Drugs, 2010, v. 19 n. s1, p. S57-S66</td>
</tr>
<tr>
<td>Issued Date</td>
<td>2010</td>
</tr>
<tr>
<td>URL</td>
<td><a href="http://hdl.handle.net/10722/132478">http://hdl.handle.net/10722/132478</a></td>
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Chemoprophylaxis in Colorectal Cancer: Current Concepts and a Practical Algorithm for Use

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Abstract

Background: Colorectal cancer (CRC) is the second most common lethal cancer in the Western world. There is a 10 to 20 years lead time from normal mucosa to carcinoma which offers a window of opportunity to modify and prevent the outcome of CRC, with its incipient morbidity and mortality.

Objective: To review the evidence for chemoprophylaxis in CRC, identify currently utilized agents and determine their role in the current management algorithm of CRC.

Methods: Review of large cohort-control and randomized controlled trials in the most studied chemoprophylaxis agents.

Results/conclusion: Currently, the role for chemoprophylaxis in CRC remains a niche area, celecoxib is the only recommended agent for utilization in patients with familial polyposis syndromes. However, the role of chemoprophylaxis is likely to grow significantly in the next decade as understanding of the stepwise tumorigenesis cascade becomes better understood and currently conducted clinical trials are completed.

Keywords
Chemoprophylaxis, Chemoprevention, Colo-Rectal Carcinoma, Adenoma, Adenomatous Polyps, COX-2 Inhibitors, NSAIDS, Vitamin D, Calcium, 5-ASA

1 Introduction

1.1 Epidemiology
Colorectal cancer (CRC) is the second most common lethal cancer after lung cancer, accounting for 55,000 deaths per year in the United States and its incidence is rapidly rising in Asia. Overall, the incidence of CRC in Asian countries including Hong Kong, Singapore, Japan and Korea is approaching that of the West. The detection rate of advanced neoplasms is between 4.5-7.8% in both Asian and Western populations. However, despite a declining mortality rate for CRC in the West, Asia is observing a rising mortality. Thus, CRC is a widely prevalent disease associated with considerable mortality and morbidity in both the East and West.

The reason for the rising incidence of CRC in Asia and the high rate seen in the West is clearly related to a complex interplay between genetics and the environment. The adoption of the ‘westernized lifestyle’ which includes a diet high in fat and protein but low in fibre is cited as a cause of CRC development. However the influence of family history and racial differences seen between different ethnic communities in Asia underscore the influence of genetic factors in the etiology of CRC development. In countries like Singapore and Malaysia where the ethnic population consists of Chinese, Malay and Indians, the incidence of CRC is consistently lower in Indians and Malays than among the Chinese. Overall, studies in Asia have found that Japanese, Chinese
and Koreans have a higher incidence of advanced neoplasms compared to other Asians.

1.2 Adenoma Carcinoma Sequence

Unlike lung cancer, CRC is one of the most preventable cancers, due to the long lead times involved in the step-wise progression from normal mucosa to adenomatous polyp to carcinoma, and the therapeutic interventions including colonoscopy available to definitively treat adenomas prior to malignant change. Polypectomies to remove adenomas during screening colonoscopies may lower the mortality rate from CRC by as much as 30-40%.

The natural evolution of normal mucosa to adenomatous polyp to overt carcinoma, spans on average 10-20 years. This long lead time presents an opportunity in terms of providing a window of opportunity for prevention and intervention; however it also complicates the study of the impact of pharmacological agents, necessitating studies of considerable duration and patient size to detect a significant effect. Thus as the epidemiology of adenomas closely resembles that of CRC itself, prevention of adenomas has been used as a surrogate end-point as it is reasoned that preventing adenomas will most likely also prevent CRC.

1.3 Risk Factors in CRC Development

Many of the risk factors associated with CRC development including age, sex, ethnicity and family history are not modifiable. Older age particularly over the age of 50 is associated with an accelerated rate of CRC development, hence the current recommendation is that CRC screening commences at the age of 50 in an average risk individual. Consistently, the incidence of CRC development in males is higher than females, moreover a study found that pregnancy was associated with a reduced risk of CRC, leading to a hypothesis that female sex hormones may be protective factor for CRC development. As previously mentioned, Japanese, Korean and Chinese ethnic groups have a higher incidence of CRC when compared to other Asians, even when the ethnic population resides in the same country. Finally, a positive family history confers increased risk of CRC development. A prospective study showed that the age adjusted risk for CRC in a person with an affected first degree relative is 1.72. A meta-analyses of 27 case-control studies found that first degree relatives of patients with CRC have an increased risk of colon cancer of 2.42 and the relative risk is increased to more than 5 fold if the index case is under 45 years.

Modifiable risk factors include obesity, smoking and ethanol intake. A meta-analyses found a relative risk of CRC of 1.37 for overweight and obese men, probable causative factors cited include hyper-insulinaemia and the role of oxidative stress initiated by hyperglycaemia. Smoking increases the risk of CRC, more so for rectal cancers than for colon cancer where the risk is between 1.43-2.64 depending on the number of cigarettes
per day. Ethanol intake of more than 7 standard drinks per week was associated with an increased risk of CRC of 1.72, moreover the increased risk of smoking and ethanol intake was additive. Despite the plethora of detrimental health effects caused by obesity, smoking and excess ethanol and despite public health campaigns to raise awareness, the prevalence of these modifiable CRC risk factor continues to grow, with some even calling the global rise in obesity an epidemic.

1.4 CRC and health seeking behaviour

CRC screening via facial occult blood testing, flexible sigmoidoscopy and colonoscopy is the cornerstone of prevention. However its efficacy is limited due to low compliance with screening guidelines. In the United States, Canada, Europe and Australia, there is widespread scientific recognition of the value of CRC screening and there are national guidelines which support the screening program. Despite this, access to colonoscopic polypectomy is not yet widely available even in developed countries and even when available, the uptake of screening for CRC in those eligible remains poor. In Japan, only 17% of the eligible population underwent screening, whereas in the West, eligible uptake rates were consistently less than 50% (see Table 1). Even in patients with a family history of CRC, a study found that of patients with positive genetic testing 71% of patients had a screening colonoscopy within 12 months of diagnosis. In Asia, only several countries including Singapore, Japan, Taiwan, Korea and Australia have a national guideline regarding CRC screening, and overall governmental financial and general support for CRC is very limited. So it is not surprising the rates of screening are low. Despite the significant development of diagnostic and therapeutic tools to detect and treat pre-cancerous lesions in the colon and rectum, there are significant barriers to screening both in an average risk population and in higher risk patients, hence the need for the role of chemoprophylaxis in CRC to emerge.

2 Aim of Chemoprophylaxis

Chemoprophylaxis is a concept in the prevention of CRC which aims to interfere with the process of carcinogenesis by the targeting of key molecular pathways via pharmacologic agents to prevent, arrest or cause regression of adenomas, which are presumed to be precursors to CRC. Chemoprophylaxis can be utilized in a number of population cohorts. Chemoprophylaxis may be given as primary prevention, either to patients with average risk of CRC or only to those selected population sub-groups at higher risk of CRC, including relatives of patients with familial polyposis coli syndromes, or CRC. Chemoprophylaxis may also be utilized as secondary prevention, whereby it should be utilized in conjunction with CRC screening in patient sub-groups such as patients with detected adenomatous polyps, a prior history of CRC and family members of patients with CRC. The ideal chemoprophylaxis agent should be potent, inexpensive, widely available, and suitable to long term administration and most importantly, free from serious side effects.
2.1 Chemoprophylaxis Candidates & Mechanism of Action

The 10-20 years lead time in the development of CRC provides a window during which chemoprophylaxis may be utilized. However, it also complicates the studies of emergent pharmacologic agents, as treatment may have to be given for prolonged periods of time, particularly for primary prevention, before a therapeutic effect may or may not be seen. Moreover, large numbers of patients would need to be involved with prolonged follow-up which incurs problems such as escalating drop-out rates over time. Finally, studied patients would be subjected to additional risk involved with surveillance colonoscopies and polypectomies. The bulk of current evidence in chemoprophylaxis lies in studies of secondary prevention of either CRC or adenomas which require a shorter period of follow-up to assess the effect of chemoprophylaxis.

A large number of agents have been identified as potential chemoprophylaxis agents, predominantly from findings of observational studies. However, the results of most adenoma primary prevention trials have been disappointing. Studies of antioxidant vitamins, fiber, diet (median follow-up 3 years) and hormone replacement therapy, conducted in average-risk populations have all been negative. The apparent discordance between the findings of observational studies and the concomitant randomized controlled studies is explainable by the fact that the former are more prone to confounding and bias than the latter, especially with regard to the assessment of preventive behaviors that may be difficult to detect and control for. Studies of chemoprophylaxis in the secondary prevention of adenomas or CRC have yielded better results. The best studied agents include Vitamin D and calcium, non-steroidal anti-inflammatory drugs (NSAIDS) and cyclo-oxygenase 2 inhibitors (COX-2).

2.2 Calcium & Vitamin D

The focus on calcium and Vitamin D as possible chemoprophylaxis agents in CRC came from observational studies which revealed an inverse relationship between calcium and vitamin D intake with risk of CRC and recurrent polyps. The mechanisms underlying calcium’s anti-carcinogenic effects in the large bowel are not clear. One hypothesis is based on its capability to bind to and precipitate bile acids and soluble fatty acids, rendering them relatively inert in the bowel lumen, whereas more recent studies suggest that extra cellular calcium can affect cell proliferation and differentiation via the calcium sensing receptor, a cell surface receptor that is expressed both in normal colon and colon cancer cell lines. With regard to vitamin D, in vitro and in vivo studies have shown that vitamin D and vitamin D analogs can inhibit cell proliferation, induce differentiation, and promote apoptosis.

Primary prevention of CRC through treatment with calcium and/or vitamin D has yielded results which generally show a reduction of CRC. (see Table 2). A large randomized, double-blind, placebo-controlled trial involving over 30,000 postmenopausal women treated with calcium and vitamin D (dose 200IU twice daily) versus placebo for an
average of 7.0 years found the incidence of invasive colorectal cancer did not differ significantly between women assigned to calcium plus vitamin D supplementation and those assigned to placebo, hazard ratio 1.08 (95% confidence interval (CI), 0.86 to 1.34, P = 0.51). The ongoing 5 year extension follow-up will assess the longer-term effect of this intervention, as 7 years may not have been sufficient time to demonstrate the effect of calcium and vitamin D, however the longer follow-up time may or may not result in a different outcome as the dose of Vitamin D given to the study patients is lower than the dose usually utilised for CRC prevention. However a pooled analysis of 10 cohort studies, a comprehensive analysis involving 534,536 patients, assessed dietary consumption and total calcium intake (diet plus supplements) supported a reduction in the incidence of colorectal cancer of 10 to 15 percent, the relative risk for the highest versus the lowest quintile of intake was 0.86 (95% CI = 0.78 to 0.95; P(trend) =0.02).

More recently, two studies from Japan have found positive associations between calcium and/or vitamin D intake and lowering of the risk of CRC. Mizoue et al’s study found that the odds ratio for the highest versus the lowest quintile for calcium intake was 0.64 (95% confidence interval, 0.45-0.93). Higher levels of dietary vitamin D were significantly associated with decreased risk of colorectal cancer among those who had fewer chances of sunlight exposure at work or in leisure (P for trend=0.02). Ishihara et al’s group undertook a prospective cohort study involving 74, 639 patients and found the multivariate hazard ratio in the highest quintile of dietary calcium intake compared with the lowest was 0.71 (95% CI: 0.52, 0.98) among men. No association was seen for Vitamin D intake.

For secondary prevention, randomized clinical trials that found that calcium supplementation lowered the incidence of recurrent colorectal polyps to some degree, with one report demonstrating that this protection was confined to subjects with higher endogenous vitamin D levels.

Previous trials demonstrating beneficial effects of calcium and/or vitamin D supplementation in CRC prevention, have led to the use of these agents in risk-reduction strategies. However, the benefits of primary prevention are mildly positive and for secondary prevention, the reductions have only been in the development of recurrent adenomas, the surrogate marker for development of CRC.

5 Amino-Salicylates
The protective association between 5-aminosalicylates (5-ASA) and CRC emerged from observations of patients with inflammatory bowel disease (IBD), particularly ulcerative colitis (UC) who were treated with 5-ASA. The mechanism by which 5-ASA exerts its chemoprotective effect is unclear at present although protean mechanisms have been suggested. Proposed mechanisms include 5-ASA’s anti-inflammatory action on the bowel, action as a free-radical scavenger and it’s improvement of cellular replication fidelity amongst others.

5-ASA has been extensively studied in the reduction of CRC only in patients with IBD, in particular UC. A recent meta-analysis of 9 observational studies (3 cohort, 6 case-
control) involving a total of 1,932 subjects with UC found a protective association between the regular use of 5-aminosalicylates and CRC (OR=0.51; 95% confidence interval (CI): 0.37-0.69) and a combined endpoint of CRC/dysplasia (OR 0.51; 95% CI: 0.38-0.69) 53. A large nested case-control study involving 18,969 patients with IBD also supports the meta-analysis finding that regular 5-ASA usage dramatically reduced the risk of CRC compared to irregular usage (Adjusted OR 0.60; 95% CI 0.38-0.96) 54 as do a number of smaller studies 55 56.

Unfortunately, at present further studies are needed before 5-ASA could be recommended to be utilised as chemo-prophylactic agent, both in the IBD and general populace. Questions which will need to be answered include the optimal dose of 5-ASA, whether all 5-ASA are equal in their effect and finally, the role of 5-ASA when patients are already stabilized on an alternate agent including azathioprine 53.

2.3 Non-Steroidal Anti-Inflammatory Drugs

The suggestion of NSAIDS as possible chemoprophylaxis arose from the consistent concordance of data from more than 40 observational studies suggesting that NSAIDS reduce the incidence of colorectal adenomas, colorectal cancer, and deaths from colorectal cancer 57. A leading hypothesis explaining the mechanism of NSAIDS in preventing CRC and adenomas is based on the presence of tumorigenic COX-2 within adenomas but not in normal intestinal tissue. COX-2 mediates the production of prostaglandin E2 (PGE2) in epithelial tissues, resulting in activation of signaling pathways that promote cell proliferation and inhibit cell death 58, 59. COX-2 is over expressed in 40-50% of adenomas and 85% of CRC 60. NSAIDS, which includes sulindac and aspirin, the two most studied NSAIDS as CRC chemoprophylactic agents, are non selective cyclo-oxygenase inhibitors (see Table 2). Emergent evidence suggest sulindac may exert it chemoprophylactic action via mechanisms other than via COX-2 61

Sulindac, a nonselective cyclooxygenase inhibitor, was previously reported to cause complete or nearly complete regression of rectal adenomas in uncontrolled studies 62-65, and in a small, placebo-controlled, drug-crossover trial of patients with familial adenomatous polyposis 65. Regression of rectal adenomas, though of lesser magnitude, was reported in two subsequent placebo-controlled studies 3, 66 However, rapid recurrence of adenomas was observed from three months after discontinuation of drug therapy 3, 65 and evidence of long-term efficacy of sulindac is still lacking, with case reports of tumor progression in patients receiving sulindac 67.

Randomized trials of aspirin demonstrated substantial chemopreventive effect in secondary prevention, with reductions of approximately 20-50 percent among patients in whom recurrent adenomas developed. 23, 68, 69 Aspirin was also associated with more substantial reductions in the risk of advanced lesions than in that of non-advanced lesions suggests that the effects of aspirin may be greater in later stages of the adenoma–carcinoma sequence 68. These findings are tempered somewhat by the observation in one study that low-dose, but not high dose, aspirin had an antitumor effect 68. There was an increase in the risk of stroke among patients who were randomly assigned to receive
aspirin (P=0.06), a finding consistent with previous studies of populations at low cardiovascular risk 68.

The sulindac and aspirin studies have had generally consistent results, with benefits after factors including age, race, sex, and location of the tumor in the colon or the rectum have been controlled for, however, the gastrointestinal toxicity associated with NSAIDS at large may limit their long-term use for cancer prevention 70. Almost all studies indicate that the reduction in the risk of colorectal cancer or adenomas declines after aspirin therapy is stopped 71. Long-term studies, as well as direct comparisons of selective and nonselective cyclooxygenase inhibition, could further define the relative clinical benefits and adverse effects of these agents so that NSAID place in the management of CRC may be further clarified 72 73.

2.4 Cyclo-Oxygenase-2 Inhibitors

Selective COX-2 inhibitors were developed as a safer alternative to nonselective NSAIDs, with respect to gastrointestinal bleeding, for the treatment of pain and inflammation. These agents preferentially inhibit COX-2, an inducible enzyme mediating inflammation and tumorigenesis, and not COX-1, the constitutively expressed enzyme responsible for protective mechanisms in the gastric mucosa and renal vasculature. Selective COX-2 inhibitors have fewer effects on gastric mucosa or platelet function than do the nonselective NSAIDs and, as a result, may be associated with fewer ulcers and hemorrhagic complications 74. COX-2 inhibitors may also have effects on other enzymes, such as induction of 15-lipoxygenase-1 expression which could mediate its biologic effects 75. The best studied COX-2 inhibitors to date include celecoxib and rofecoxib (see Table 2).

It was noted that in patients with familial adenomatous polyposis (FAP), celecoxib showed antitumor activity, with regression, in addition to suppression of established adenomas 72. In a six-month study in patients with FAP, treatment with a celecoxib 400mg twice daily was associated with statistically significant 28% regression of colorectal adenomas as compared to a reduction of 4.5% in the placebo group (p=0.003) 72. Significant regression was not associated with the dose of 100 mg twice daily 72. These findings suggest that celecoxib could serve as an adjunct to current best management in the FAP syndromes by suppressing polyp formation in patients with residual rectum after total colectomy or in patients with an intact colon who are awaiting colectomy.

Three landmark studies on COX-2 inhibitors on sporadic adenomas have demonstrated the significant effect of COX-2 inhibitors on secondary CRC and/or adenoma prevention. The Adenomatous Polyp Prevention on Vioxx (AP-PROVe) trial utilizing rofecoxib, the Prevention of Sporadic Adenomatous Polyps (PreSAP) trial and the Adenoma Prevention with Celecoxib (APC) trial both utilizing celecoxib compared to placebo, found statistically significant reductions in recurrent adenomas in the treatment arm over placebo, and moreover found that a greater effect was observed in advanced adenomas 75-78.
Tempering the significant impact of COX-2 inhibitors on the recurrence of adenomas are safety issues concerning COX-2 inhibitors. In both the APPROVe and APC studies there was a significant increase in cardiovascular events including myocardial infarction stroke, stroke and heart failure. The potential addition of COX-2 inhibitors to an optimal endoscopic surveillance program must be weighed against the known risk of serious cardiovascular events. In the APPROVe trial a total of 46 patients in the rofecoxib arm had a confirmed thrombotic event compared to 26 patients in the placebo group (RR, 1.96) whilst in the APC trial, a significant dose-response excess of major cardiovascular events for celecoxib 200mg and 400mg of 2.5 (95% CI, 1.0-6.4) and 3.5 (95% CI, 1.4-8.5) respectively, lead to premature discontinuation of both studies. Secondary analysis of the APC study indicates that the risk was for those patients with a baseline history of atherosclerotic heart disease over a median on 3.1 years. Interestingly the PreSAP trial which was also discontinued due to safety concerns did not find any statistically significant excess cardiovascular events in the active treatment arm with celecoxib.

The studies of COX-2 are important as they demonstrate the relationship between COX-2 and colonic adenomas. COX-2 inhibition does have a place in the secondary prevention of CRC. At present, given the long term safety data of COX-2 inhibitors, celecoxib is the only approved agent for chemoprophylaxis in high risk patients with FAP.

### 3 Proof of Concept

The place of chemoprophylaxis in CRC has been an area of interest for some time and provides a number of instructive points. Firstly, it demonstrates the importance of observational studies as flags to possible beneficial agents for prevention of CRC, but more importantly, this story highlights the absolute necessity to subject the findings of observational studies to rigorous randomized, controlled trials to determine the actual contribution of a proposed chemoprophylactic agent, consider anti-oxidant vitamins and hormone replacement therapy, whilst controlling for possible confounding factors. Secondly, it demonstrates the absolute necessity to demonstrate the safety of the ideal CRC chemoprophylaxis agent over the long term, as COX-2 inhibitors are safe over the short term when used as treatment for pain and inflammation and their long term safety issues have only come to light when utilized long term. Thirdly, the search for the ideal chemoprophylaxis agent has yielded a number of agents including calcium and vitamin D, aspirin, sulindac and celecoxib which could be used as an adjunct to currently employed screening methods, particularly given possibility of overlooked adenomas, since up to 15-25 percent of small polyps may be missed in a single colonoscopy. 5-ASA’s role as a chemoprophylaxis agent is at present unresolved. Finally, the search for the ideal chemoprophylaxis agent for CRC prevention has yielded further insight into the steps in tumourigenesis, mainly the proof of concept of the role of COX-2 in colonic adenoma development.
4 Conclusion

With each iteration of studies in the quest for the ideal CRC chemoprophylaxis agent, more knowledge is generated about CRC and the process of tumourigenesis. Although current chemoprophylaxis agents are imperfect, it is evident that selected patients may benefit from usage of calcium and vitamin D, aspirin and sulindac for secondary prevention of adenomas and/or CRC. In patients at high risk patients due to FAP, celecoxib is recommended. However, attention needs to be paid in balancing potential cardiovascular risk associated with COX-2 inhibitors. Newer chemoprophylaxis agents such as selenium, difluoromethylornithine and ursodexycholic acid are on the horizon and are currently under investigation. Combination chemoprophylactic therapies hold promise to herald in a new era of reduction in the morbidity and mortality associated with CRC.

5 Expert opinion

The role of chemoprophylaxis in the prevention of adenomas and ultimately CRC continues to evolve. Current utility is in the moderate to high risk populations which include patients with Familial Adenomatous Polyposis Coli and Hereditary Non-Polyposis Colorectal Cancer or Lynch’s Syndrome (see figure 1). In these patients, appropriate screening/surveillance remains the cornerstone of management with prophylactic surgery an important therapeutic adjunct. Chemoprophylaxis utilising celecoxib is recommended as it has been shown to arrest and possibly cause the regression of adenomas that develop in the interim between lower gastro-intestinal endoscopies. However in patients with cardio-vascular risk factors and/or history sulindac should be substituted for celecoxib. For patients with moderate CRC risk, the benefits of chemoprophylaxis must be balanced with the likely adverse effects of the chemoprophylactic agent. In this group with moderate risk for CRC utilization of a chemoprophylactic agent which is able to treat other co-morbid conditions, strengthens the indications for use, in particular, the utilization of calcium with or without Vitamin D in a patient with osteoporosis, or aspirin in a patient with concomitant cardiovascular disease, would deliver increased mileage for the chemoprophylactic agent whilst mitigating the possible adverse effects. The exception being in patients with a detected advanced adenoma, here it would be reasonable to administer celecoxib until the next surveillance colonoscopy (usually within 3 years), to treat any synchronous adenomas that had been missed or only partially resected. In patients with average risk of CRC, there are a multitude of modalities over and above chemoprophylaxis which have been shown to decrease the risk of CRC (see Figure 2), the most important of which are to manage lifestyle factors and participate in CRC screening. Here, there needs to be a relatively high threshold for utilizing chemoprophylaxis because of the high risk of doing harm.

These are exciting times in the primary and secondary prevention of adenomas and CRC, the results from the Women’s Health Initiative and the results of the newer chemoprophylactic agents may change the treatment paradigm of CRC within the next 10 years.
### Uptake of CRC Screening

**Table 1**

<table>
<thead>
<tr>
<th>Study</th>
<th>Country</th>
<th>CRC Screening</th>
<th>% Uptake *</th>
<th>Year</th>
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<tr>
<td>Myers et al 82</td>
<td>USA</td>
<td>FOBT</td>
<td>41%</td>
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<tr>
<td>Hardcastle et al 83</td>
<td>United Kingdom</td>
<td>FOBT</td>
<td>38.2%</td>
<td>1996</td>
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<tr>
<td>Robinson et al 84</td>
<td>United Kingdom</td>
<td>FOBT</td>
<td>37.7%</td>
<td>1996</td>
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<tr>
<td>Cole et al 85</td>
<td>Australia</td>
<td>FOBT</td>
<td>32-40.7%</td>
<td>2002</td>
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<tr>
<td>Seeff et al 86</td>
<td>USA</td>
<td>FOBT and/or lower GI endoscopy</td>
<td>42.5%</td>
<td>2004</td>
</tr>
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<td>Saito et al 21</td>
<td>Japan</td>
<td>FOBT</td>
<td>17%</td>
<td>2006</td>
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<tr>
<td>Zarychanski et al 87</td>
<td>Canada</td>
<td>FOBT and/or lower GI endoscopy</td>
<td>17.6-23.5%</td>
<td>2007</td>
</tr>
<tr>
<td>Deutekom et al 88</td>
<td>Netherlands</td>
<td>FOBT</td>
<td>49%</td>
<td>2009</td>
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</table>

* proportion of patients who take up the CRC screening when eligible to do so  
** self reported answer to questionnaire about screening for CRC  
CRC = colorectal cancer  
FOBT = Faecal occult blood testing

### Summary of Chemoprophylaxis Trial Outcomes

**Table 2**

<table>
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<th>Study</th>
<th>Agent</th>
<th>Type of Prevention*</th>
<th>Relative Risk Ratio</th>
<th>Odds Ratio</th>
<th>Hazard Ratio</th>
<th>95% CI</th>
<th>P-Value</th>
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<tr>
<td>Wactawski-Wende et al 43</td>
<td>Calcium + Vitamin D</td>
<td>Primary CRC</td>
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<td>0.86-1.34</td>
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<td>Baron et al 48</td>
<td>Calcium</td>
<td>Secondary Sporadic</td>
<td>0.76</td>
<td>0.6-0.96</td>
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<td>Bonithon-Kopp et al 89</td>
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<td>Secondary Sporadic</td>
<td>0.66</td>
<td>0.3-1.17</td>
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<td>Cho et al 44</td>
<td>Calcium</td>
<td>Primary CRC</td>
<td>0.86</td>
<td>0.78-0.95</td>
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<td>Grau et al 50</td>
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<td>Giardiello et al 3</td>
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<td>FAP</td>
<td>0.56</td>
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<td>Study</td>
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<tr>
<td>Sandler et al 23</td>
<td>Aspirin</td>
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<td>0.65</td>
<td>0.46-0.91</td>
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<tr>
<td>Baron et al 68</td>
<td>Aspirin 81mg</td>
<td>Secondary</td>
<td>0.81</td>
<td>0.6-0.96</td>
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<td>Aspirin 325mg</td>
<td>Sporadic</td>
<td>0.96</td>
<td>0.8-1.13</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benamouzig et al 69</td>
<td>Aspirin</td>
<td>Secondary</td>
<td>0.73</td>
<td>0.52-1.04</td>
<td>0.08</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arber et al 75</td>
<td>Celecoxib</td>
<td>Secondary</td>
<td>0.64</td>
<td>0.56-0.75</td>
<td>&lt;0.001</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baron et al 77</td>
<td>Rofecoxib</td>
<td>Secondary</td>
<td>0.76</td>
<td>0.69-0.83</td>
<td>&lt;0.0001</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bertagnolli et al 78</td>
<td>Celecoxib 200mg</td>
<td>Secondary</td>
<td>0.67</td>
<td>0.59-0.77</td>
<td>&lt;0.001</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Celecoxib 400mg</td>
<td>Sporadic</td>
<td>0.55</td>
<td>0.48-0.64</td>
<td>&lt;0.001</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Refers to Primary or Secondary prevention of CRC and/or adenomas, FAP, CRC or sporadic denotes whether the study population was confined to patients with Familial Adenomatous Polyposis, prior history of Colorectal Carcinoma, or sporadic adenomas respectively

** Only for patients with Vitamin D levels above the median 29.1ng/ml

# Hazard ratio and CI refers to calcium in male study patients, no association seen between Vitamin D and a reduction in CRC development

## Odds ratio and CI refers to calcium, association also seen study patients with higher vitamin D intake or sunlight exposure p= for trend 0.02

**References:**