Familiar drugs may prevent cancer

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Abstract
Despite positive results in large scale chemoprevention trials, many physicians are unaware of the potential cancer preventive properties of drugs in common usage. The antiestrogen tamoxifen and the selective cyclo-oxygenase-2 inhibitor celecoxib have been licensed in the USA for the chemoprevention of breast and colorectal cancers respectively in selected high risk individuals. Similarly, folate and retinol have been shown to decrease the incidence of colorectal cancer and squamous cell carcinoma of the skin respectively in large scale intervention trials. Other retinoids have proved efficacious in the tertiary chemoprevention of cancers of the breast and head/neck. Epidemiological evidence also exists in favour of aspirin, non-steroidal anti-inflammatory drugs, and angiotensin converting enzyme inhibitors preventing certain cancers. Phytochemicals may represent less toxic alternatives to these agents. Although some of these drugs are available without prescription and most are not yet licensed for use in cancer chemoprevention, physicians and patients of individuals should be aware of this accumulating evidence base. Practitioners should be amenable to patient referral to discuss complex issues such as risk estimation or potential benefit from intervention.

Keywords: cancer chemoprevention; tamoxifen; folate; retinoids; cyclo-oxygenase

Cancer incidence and mortality continues to increase, and it has now overtaken heart disease as the commonest cause of death in Britain and Ireland. Although tamoxifen may have attenuated the rising mortality rates from breast cancer, chemotherapy has displayed a disappointing lack of impact on the prognosis from solid malignancies in general. Alternative strategies (see fig 1) have developed since the “war on cancer” was first announced by US President Nixon in 1971. One involves pharmacological intervention to arrest, inhibit or reverse carcinogenesis, and is termed cancer chemoprevention. Its definitions are shown in fig 2.

Two drugs have already been licensed in the USA for use in cancer chemoprevention (see below) and results of large European clinical trials are eagerly awaited. Despite the absence of licensed drugs in countries such as Britain, physicians are often asked questions about cancer prevention by patients’ relatives and “high risk” individuals, not least because of the frequently held fears evoked by this common disease and its media coverage. Such questions pertain to some of the drugs in common clinical usage for other diseases. Unlike the prevention of cancer, primary and secondary prevention are well established in other diseases such as dental caries, heart attacks, and stroke, and play a prominent part in medical education. It should be noted that any physician directly or indirectly involved with cancer screening is increasingly likely to encounter such questions, since many of the tests used will detect high risk patients with premalignant disease.

The aim of this review is to inform practising physicians of positive large scale chemoprevention trials, and why these drugs may prevent cancer. From our experience, patients tend to ask about specific drugs, and we will therefore

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**Figure 1** Simplified diagram of intervention strategies employed in the prevention and treatment of cancer. Carcinogenesis is shown using the multistep model (modified from Fearon and Vogelstein”). Dietary intervention is exemplified by UK Government advice on red meat intake based on epidemiological data. Reviews on immunoprevention (for example, vaccination) and advances in the treatment of cancer have been published recently. For overlap between chemoprevention and chemotherapy, see fig 2.
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Figure 2 The levels of cancer chemoprevention, as defined by subject group (modified from Gescher et al." and De Flora et al.). Drug toxicity is least acceptable in primary chemoprevention, where large numbers of subjects are required for results to reach statistical significance. 

Tamoxifen
Successful chemopreventive agents may be developed from their proven role in palliative or adjuvant chemotherapy. An example is the oestrogen analogue tamoxifen, which was initially found to improve survival in patients with metastatic breast cancer, and subsequently found to decrease the incidence of tumours in the contralateral breast after surgery. Forty per cent of newly diagnosed female cancers in the USA arise in tissues responsive to hormones, and it is known that the incidence of endometrial, ovarian, and breast cancers rises in an age dependent fashion similar to other cancers only until the menopause, after which time there is a distinct slowing of the rate of rise. Epidemiological evidence for the role of oestrogen as a primary stimulant of breast cell proliferation is supported by preclinical and clinical studies, and unlike the response in the uterus, the proliferation of breast cells is probably augmented by progestogens. Receptor antagonism is therefore an important mechanism of intervention during breast carcinogenesis, and the demonstration of the occurrence of the oestrogen receptor-β in prostate, colon, and ovary offers similar potential for these tissues.

Significant controversy exists between North America and Europe regarding the proved clinical efficacy of the two antioestrogenic agents, tamoxifen and raloxifene, both synthesised more than 20 years ago, well before the cloning and mechanistic elucidation of oestrogen receptors. The Breast Cancer Prevention Trial (BCPT) study in the US involved 13 388 women of “relatively high risk” (that is, with a substantially greater than 1.66% chance of developing breast cancer within five years), but free of detectable invasive breast cancer at study recruitment. Risk was determined using a mathematical model previously published, and included factors such as age (any woman 60 or more years old was eligible), benign breast disease, nulliparity, and family history. Tamoxifen reduced relative risk for subsequent development of breast cancer in the treated group as a whole by 45%, and may also have prevented the progression of established preneoplastic lesions. These results led the Food and Drug Administration to approve the use of tamoxifen (20 mg/day) by women deemed at increased risk of developing breast cancer, albeit with consideration of its toxicities. Tamoxifen causes hot flushes, vaginal bleeding or discharge, fluid retention and amenorrhoea, increases risk for venous thromboembolic events and can cause endometrial hyperplasia, dysplasia, and carcinoma. Extrapolating from data on its effect in the adjuvant chemotherapy setting, one might speculate that tamoxifen continues to reduce the incidence of primary breast cancer for at least five years after treatment stops. However, caution is advised since no reduction in mortality has yet been demonstrated in the BCPT study, perhaps because this drug may prevent only oestrogen receptor positive tumours, which may respond to hormonal manipulation even once carcinoma is established.

Results favouring the prevention of breast cancer have also been published in a trial of raloxifene in postmenopausal women with known osteoporosis, but not considered at increased risk of breast cancer. A larger breast cancer prevention trial of raloxifene is ongoing, enrolling 20 000 women and due for publication in 2006. Raloxifene, licensed in most countries only for the chemoprevention of osteoporosis, is generally better tolerated than tamoxifen. It may also cause hot flushes and peripheral oedema, and does increase risk for thromboembolism, but may not possess the endometrial stimulatory properties of tamoxifen.

In contrast to the BCPT study, two smaller European trials of tamoxifen in healthy women, with lower average risk factors, have not demonstrated any protective effects. In the Italian study, post-hysterectomy patients were included, and the British study allowed patients to continue supplemental oestrogen therapy; the mean age of participants for both trials was lower than the BCPT study. Consequently, in Europe the results of the International Breast Cancer Intervention study (IBIS) are awaited, in which more than 4000 women with at least a twofold increased risk of developing breast cancer have been recruited. The controversy regarding the benefit of agents such as tamoxifen in preventing breast cancer in the US compared with Europe illustrates the importance of subject selection in large scale chemoprevention trials. It is feasible that high risk subgroups, perhaps with particular molecular defects (see below) or a certain receptor status in preneoplastic lesions, must be studied for statistical significance to be demonstrated. Indeed such individuals may have the most favourable benefit-toxicity ratios.
Aspirin and NSAIDs

Retrospective epidemiological studies suggest a decreased incidence of cancers of the oesophagus, stomach, colon, and rectum in regular users of non-steroidal anti-inflammatory drugs (NSAIDs). The most convincing evidence exists for regular usage of aspirin, which may reduce the incidence of colorectal cancer by up to 50%, although there may be a delay of approximately one decade before the benefits of daily usage are seen. Colorectal adenomas have been regarded as the quintessential precursor lesions of cancer since the 1970s, and are present in a third of the general population by the age of 50 years and approximately half the population by the age of 70 years. Mutations in the APC gene, first described in the inherited syndrome of familial adenomatous polyposis (FAP), are found in 80% of all colorectal adenomas and carcinomas. Although FAP accounts for only 0.5% of all colorectal cancers, this disease may therefore represent a useful model of the more common sporadic form of this cancer.

In individuals with FAP, the presence of the APC gene defect confers a nearly 100% lifetime risk of developing colorectal cancer. At present such individuals are advised to undergo regular colonoscopy with or without colectomy. It has been known for many years that aspirin and NSAIDs, such as sulindac, can cause regression of FAP polyps, although adenomas do recur and regrow when treatment is curtailed. The postulated mechanism is their ability to inhibit the cyclo-oxygenase (COX) enzyme involved in prostaglandin synthesis. COX consists of two isoforms, and aspirin’s irreversible acetylation of COX-1 is thought to account predominantly for its gastrointestinal toxicity. Highly selective inhibitors of COX-2, such as celecoxib, have therefore been developed. COX-2 has been implicated in the pathogenesis of human cancers of the colorectum, breast, head/neck, lung, pancreas, stomach, and prostate. Celecoxib, originally licensed in Europe and

North America for the treatment of arthritis, was recently approved by the US Food and Drug Administration as an adjunct to usual care for patients with FAP. This decision was based on the results of a double blind, placebo controlled trial in 83 patients with FAP, which demonstrated that 400 mg of celecoxib twice daily for six months resulted in polyp incidence 28% lower than placebo. Other “surrogate” biomarkers, such as prostaglandin synthesis or COX-2 expression, were not measured. Such biomarkers allow prediction of effects on cancer mortality based on scientific hypotheses, for example colorectal cancer develops in adenomas, COX-2 levels in adenomas represent carcinogenic progression, and suppression of COX-2 levels may represent efficacy of intervention.

The long term safety of selective COX-2 inhibitors such as celecoxib remains unknown, but early indications are that their side effect profile is similar to that of traditional NSAIDs, although peptic ulceration appears to be less likely than for the older agents. These agents are likely to maintain selectivity for COX-1 at the dose range used in the published study. Irreversible inhibition of COX-1 by aspirin accounts for its antiplatelet effect since these cells do not possess nuclei and are

Box 1: Key points
- There is increasing epidemiological evidence that certain drugs may decrease the incidence of cancer.
- Two drugs have been licensed in the USA for the chemoprevention of breast and colorectal cancers in selected high risk individuals.
- Folate and retinol have been shown to decrease the incidence of colorectal and skin squamous cell cancers respectively in large scale intervention trials.
- Aspirin, NSAIDs, and ACEIs may also reduce cancer incidence, as may certain phytochemicals.
- Practitioners should be amenable to referral of individuals in whom there is objective evidence of high risk that they may develop certain cancers.

Box 2: Five key references

Box 3: Useful websites for monitoring chemoprevention trials ongoing
- http://clinicaltrials.gov/
- http://cancerinet.nid.nih.gov/cgi-bin/srcchgi.exe
- http://iarc.fr/pageroot/UNITS/CHP.HTM
- http://cancerindex.org/clinks4t.htm
Questions (see answers on p 496)

1. Which two drugs have been licensed by the US Food and Drug Administration for the chemoprevention of breast and colorectal cancers respectively, in which highly selected individuals?

2. According to the Nurses’ Health Study, how much folate must be taken as a supplement for how long in order to decrease the incidence of colorectal cancer?

3. (A) In the pathogenesis of which human cancers has COX-2 been implicated? (B) Which drugs inhibit the COX enzyme?

4. Why are biomarkers useful in cancer chemoprevention trials?

5. (A) In what disease does differentiation therapy with all-trans retinoic acid lead to complete remission in the vast majority of patients? (B) Which skin cancer might retinol prevent?

6. In which dietary components are the following putative cancer chemopreventives found: EGCG, curcumin, genistein, resveratrol?

Folate is central to methyl group metabolism, and as such may influence both methylation of DNA and the available nucleotide pool for DNA replication and repair. DNA hypomethylation is an early step in colon carcinogenesis. Vitamin B_9_, cofactor in this pathway, and the val/val polymorphism of the methylene-tetrahydrofolate reductase (MTHFR) gene may also influence the association between folate intake and the development of carcinoma from adenoma.

The degree of benefit from taking folate supplements may be greater than that from its consumption in the diet. The Nurses’ Health Study began in 1976 and followed 121 700 married, registered female nurses of ages 30–55 years prospectively by questionnaire. In one subgroup, supplementation with folate was protective against colorectal cancer, with the greatest risk reduction among women taking daily doses of more than 400 μg folate; however, this reduction became statistically significant only after 15 years of use. It is also becoming apparent that the protective role of folate supplementation may be greatest for those genetically predisposed to colorectal cancer, and that the benefit conferred by MTHFR genotype may be offset by a methyl deficient diet.

Retinoids

Like folate, retinol (vitamin A) is available in the diet, particularly from green leafy vegetables, liver, eggs, and milk. Retinol is the precursor of all physiologically occurring retinoids, and is required for normal vision and reproduction. The oxidation products of retinol are essential for the maintenance of normal epithelial differentiation (reviewed in Hansen et al [39]).

It has been hypothesised that retinoids, at optimal or supraphysiological levels, inhibit the development of epithelial carcinogenesis. This activity is utilised in acute promyelocytic leukaemia, in which treatment with all-trans retinoic acid leads to complete remission in up to 95% of patients. In the Skin Cancer Prevention-Acnic Keratosis trial, 25 000 IU of retinol was taken daily and primary prevention of squamous and basal cell carcinomas of the skin were the two endpoints measured. The trial involved 2297 subjects deemed to be at moderate risk, and the treatment was found to prevent squamous cell carcinoma significantly. A drawback of this study is the fact that increasing dietary ingestion of retinol is unlikely to deliver more retinol to skin; it merely leads to an accumulation of retinyl esters in liver tissue. It has therefore been proposed that direct administration of retinoids to target tissues may be more effective chemoprevention than oral supplementation.

It is vital that any agent under consideration for chemoprevention in healthy high risk individuals over prolonged periods of time should not cause more harm than benefit. Although the retinoids, isotretinoin and retinol palmitate, have been demonstrated to prevent second primary cancers in patients with malignancies of the lung and head/neck,


**Answers**

1. Tamoxifen has been licensed for the primary chemoprevention of breast cancer in high risk individuals after consideration of its toxicities. Risk can be determined mathematically, using factors such as age, benign breast disease, nulliparity, and family history. Celecoxib, the selective COX-2 inhibitor, has been licensed as an adjunct to standard treatment of patients with familial adenomatous polyposis. This represents secondary chemoprevention (see fig 2).

2. In one subgroup of this large trial, supplementation with folate was protective against colorectal cancer, reaching significance in women taking daily doses of more than 400 µg folate for 15 years.

3. (A) COX-2 has been implicated in the pathogenesis of human cancers of the colorectum, breast, head/neck, lung, pancreas, stomach, and prostate. (B) Aspirin, NSAIDs, selective COX-2 inhibitors, and certain phytochemicals (for example curcumin, resveratrol) inhibit the COX enzyme.

4. If cancer mortality acts as the only endpoint, chemoprevention trials must study time periods of 5–15 years in order to reach significance, with no indication of beneficial or detrimental effects. “Surrogate” biomarkers allow prediction of the efficacy of intervention based on scientific hypotheses of carcinogenesis.

5. (A) Acute promyelocytic leukaemia. (B) Squamous cell carcinoma of the skin.

6. They are found in tea, the spice turmeric, soya, and wine respectively.

Compliance is often problematic on account of toxicity. Similarly the synthetic retinoid, fenretinide, significantly decreased the risk of breast cancer in premenopausal women in a tertiary chemoprevention study (see fig 2), but night blindness and erythema proved prohibitive at higher doses. Newer retinoids that selectively bind to retinoid X receptors appear highly chemopreventive in preclinical models of epidermal and mammary carcinogenesis, and do not possess the toxicity/teratogenicity profile of the classical retinoids.

**Others**

Treatment of hypertensive patients with captopril, an angiotensin-I converting enzyme inhibitor (ACEI), is associated with a reduced risk of developing malignancy, particularly lung and breast cancers. This may relate to the ability of ACEIs to inhibit angiogenesis, the formation of new blood vessels vital to growth of cancers beyond 1–2 mm³. As well as inhibiting chemotaxis of capillary cells, captopril is a free sulphydryl donor which can lead to the generation of antiangiogenic compounds in vitro.

Captopril also inhibits matrix metalloproteinase activity, integral to the neovascularisation process.

Finally, certain phytochemicals offer a minimally toxic form of intervention during carcinogenesis with similar mechanisms to those described above. An example is provided by curcumin, a potent antioxidant derived from the spice turmeric. This compound appears innocuous in doses up to 8 g/day, yet possesses biological activity in micromolar concentrations very similar to that of aspirin. In addition, it appears to inhibit other processes linked to carcinogenesis, such as lipid peroxidation and angiogenesis. Other such putative chemopreventive agents include epigallocatechin gallate (EGCG) in green and black tea, the isoflavone genistein found in soya, resveratrol from wine, and micronutrients such as selenium and vitamin D. Both curcumin and EGCG have been shown to affect processes pivotal to cell signalling, such as kinases, cell cycle regulatory proteins, and downstream elements of cellular signalling cascades crucial for cell proliferation (reviewed in Gescher et al’). A convergence is thus developing between the targets identified for cancer chemopreventive agents and those for chemotherapeutic drugs. It is conceivable that modification of these targets may prove more efficacious in intervention when fewer cellular components are malfunctioning (chemoprevention) than treatment when dysregulation of many pathways is already established (chemotherapy).

**Conclusions**

Physicians should be aware of the potential cancer chemopreventive properties of commonly used drugs. Tamoxifen was found to reduce breast cancer development in high risk women in one large controlled intervention study, and the results of a similar study in Europe are awaited. Consideration of its toxicities must be weighed up against the cancer risk for each individual. Raloxifene, already used in the chemoprevention of osteoporosis, may offer a less toxic alternative to tamoxifen. A large prospective intervention study has found that folate supplementation may decrease the incidence of colorectal cancer after 15 years of daily use. NSAIDs and celecoxib have been shown to cause regression of adenomas, considered the premalignant lesions in this disease. Retrospective epidemiological surveys have suggested that aspirin may significantly reduce the incidence of colorectal cancer after at least nine years of daily usage. As demonstrated by studies of celecoxib and folate, consideration of risk ratios and genetic predisposition is increasingly important in recruitment and subset analysis of chemoprevention trials. Retinoids have been shown to be efficacious in the primary prevention of skin squamous cell carcinoma, and the tertiary chemoprevention of cancers of breast and head/neck, but toxicity has proved limiting. ACEIs and phytochemicals may also prevent certain cancers, and the latter may represent less toxic...
alternatives to conventional drugs. There are complex issues in assessing one’s risk of developing cancer and the potential benefit from intervention; practitioners should be amenable to patient referral should individuals wish to discuss these issues with specialist oncologists.

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41. www.postgradmed.com